



Dedicated to my father  
Prof.dr. J.Th.G.Overbeek (1911-2007)  
who encouraged me to be curious.

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# Detection of congenital anomalies before or after birth; does it make a difference?

*Diagnose van congenitale afwijkingen voor of na  
de geboorte; maakt het uit?*

## **Proefschrift**

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## **PROMOTIECOMMISSIE**

**Promotoren**      Prof. jhr. dr. J.W. Wladimiroff  
                         Prof. dr. D. Tibboel

**Overige leden**    Prof. dr. N.M.A. Bax  
                         Prof. dr. G.J. Bonsel  
                         Prof. dr. J. Deprest

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# 1

## Introduction and research objectives

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# Introduction

Stuart Campbell produced the first ultrasound image of a congenital anomaly in the human fetus in 1972<sup>1</sup>. This revolutionised Obstetrics and Gynecology. Reliable determinations of gestational age<sup>2</sup> and cervical competence<sup>3</sup> can be made and information on fetal and utero-placental perfusion<sup>4</sup>, fetal growth<sup>5</sup> and a vast number of fetal anomalies can now be obtained<sup>6</sup>. Major fetal anomalies are associated with preterm delivery<sup>7</sup>, perinatal mortality and morbidity<sup>8</sup>, unwarranted obstetric surgery and prolonged hospitalisation<sup>9</sup>. About 2 to 3% of newborns have detectable congenital anomalies<sup>10,11</sup>, of which 20% will result in perinatal death<sup>9,12</sup>. When ultrasound techniques became sufficiently powerful, many countries introduced a second trimester ultrasound screening test to detect these anomalies<sup>13,14</sup>.

## Screening-universal or risk based?

When addressing screening one should distinguish between universal screening and targeted screening<sup>15</sup>. Targeted screening refers to the examination of women at known increased risk of having off-spring with congenital anomalies. Two risk situations can be distinguished for such an indication-based only ultrasound scan.

Firstly, there is the situation that risk factors are known prior to the pregnancy, i.e. a previously affected infant, a parent with a congenital anomaly, maternal type 1 Diabetes Mellitus, maternal use of antiepileptic drugs. In this subset of women, 18-22 weeks is the ideal time for a thorough fetal anomaly scan. We found an overall prevalence of major fetal anomalies in this increased risk subset of 4-5%<sup>16</sup>.

In another group the ultrasound scan for suspected fetal anomalies is based on abnormal obstetric findings that appear during the current pregnancy. Whereas in early pregnancy it is fetal nuchal translucency which is closely associated with chromosomal abnormalities, notably trisomy 21 and cardiac anomalies<sup>17</sup>, obstetric markers for fetal congenital anomalies later in pregnancy include poly- and oligohydramnios, severe fetal growth restriction and fetal cardiac arrhythmias<sup>18</sup>. In this subset of women the prevalence of fetal congenital anomalies approximates 40 to 60%<sup>16</sup>. Until recently (1<sup>st</sup> January 2006) in the Netherlands anomalies were often detected beyond the second half of the second trimester due to the lack of a population-based 18-22 week fetal screening program. Population-based universal screening involves low-risk populations not known to have clinical risk factors. This examination includes determination of gestational age, number of fetuses, fetal viability, placental location and a search for fetal congenital anomalies. The rationale for such a universal screening test is that approximately 75% of infants with congenital anomalies are born out of low risk pregnancies<sup>19</sup>.



## The 18-22 week screening ultrasound scan

The 18-22 week period is determined by the high quality sonographic images available at this time of pregnancy and the possibility to detect the majority of structural anomalies. The legal upper limit of 24 weeks of gestation for termination of pregnancy in case of a major fetal congenital anomaly may also play a role but fortuitously the opportunities for detection by ultrasound are also diminished beyond this period. The acoustic window, provided by amniotic fluid, becomes relatively smaller due to fetal growth and increased mineralization of the fetal skeleton creates ever more shadows with advancing gestational age. The clear visualization of fetal structures is therefore limited in later gestation. It has been reported that in two-third of prenatally diagnosed fetal anomalies clinical symptoms may develop as late as 23-24 weeks of gestation<sup>18</sup>. Prenatal screening at 18-22 weeks has now become a routine part of antenatal care in most European countries<sup>20</sup>. In case of detection of a fetal anomaly compatible with life, there will be the option of adjusting obstetric management in terms of timing, mode and location of the delivery<sup>21</sup>. In some instances intrauterine treatment may be contemplated<sup>22</sup>.

## Requirements for screening tests

In general a screening test should meet a number of conditions. The prevalence of the anomaly should be high enough to ensure that the application of the test – providing it has an acceptable sensitivity and specificity – will result in a measurable and cost-effective improvement in health outcome. There has to be an appropriate hospital infrastructure allowing women with a positive screening result to be referred for further examination. Additionally there have to be adequate therapeutic options and the test itself has to be safe without any side effects.

## The efficacy of screening

The pick up rate at routine ultrasonographic screening in the Eurofetus Study, in which 2262 malformed fetuses are registered, was 56.2%. Sensitivity was higher for major anomalies (73.7%) compared to minor anomalies (45.7%). Within the subset of major anomalies, detection was high for central nervous system anomalies (88.3%) and urinary tract anomalies (88.5%), and lower for heart and great vessel anomalies (38.8%)<sup>23</sup>. Various studies reported a specificity for the test between 99.5% and 99.9%<sup>24,25</sup>. No side effect has been demonstrated<sup>26</sup> and studies in Finland and England have shown the test to be cost effective<sup>27,28</sup>. In the Netherlands with 8 academic centers and 13 satellite centers the infrastructure for further examinations within a week following a positive screening test is present as well, although legal and political objections led to a relatively late introduction compared with other countries in Europe.

Therapeutic options following a positive screening result include termination of pregnancy in the presence of a major or lethal anomaly. In case of an ongoing pregnancy adjustment of obstetric management may contribute to a favorable outcome. In addition, parents are counseled so that they can prepare themselves for the birth of a child facing multiple problems. A 50% reduction in perinatal mortality has been demonstrated with routine 2<sup>nd</sup> trimester ultrasound. But this was the result of parents requesting a termination of pregnancy and should strictly speaking not be referred to as a reduction in mortality but rather a temporal shift<sup>29</sup>. Large screening studies failed to reveal an effect on morbidity, as measured by the proportion of babies with an Apgar score <7 at 1 minute<sup>29</sup>. Each anomaly carries its own specific type of morbidity and end-points such as the Apgar score may not be able to demonstrate a difference in morbidity between prenatally and postnatally diagnosed anomalies. Only very few studies have been able to determine a reduction in morbidity for infants with a specific anomaly detected prenatally versus detection at birth, such as transposition of the great vessels, complete atrioventricular septal defect, tetralogy of Fallot and pulmonary atresia<sup>30,31</sup>. More premature births have been demonstrated for infants with a prenatally as opposed to a postnatally detected gastrointestinal malformation<sup>32</sup>, without further information on morbidity and outcome, however.

## **Late introduction of screening as a research opportunity**

There was no nationally implemented screening at 18-22 weeks in the Netherlands until the beginning of 2006<sup>33</sup>. Fetal anomalies were therefore often diagnosed late in pregnancy (beyond 24 weeks of gestation) or were only detected at birth. The previous non-screening situation in the Netherlands set the scene for the research presented in this thesis. In the absence of the 18-22 week fetal screening test one can expose the impact of the indication-based only scan approach on the prenatal detection rate and outcome of major fetal congenital anomalies compatible with life but associated with severe handicap. A non screening situation also allows an answer to the question as to whether prenatal detection of less severe fetal anomalies amenable to postnatal treatment would improve postnatal outcome when compared with detection only at birth. Prenatal detection of milder fetal congenital anomalies may assist in determining which newborn need to be referred for further postnatal investigation and treatment. It may further optimise postnatal management.

## The following research objectives were defined.

1. To determine the impact of prenatal detection of fetal spina bifida on outcome in a non-screening setting (Chapter 2.1)
2. To study the outcome of prenatally versus postnatally detected anomalies, that are amenable to postnatal treatment: (i) fetal duodenal atresia (Chapter 3.1), (ii) fetal gastroschisis (Chapter 3.2), (iii) fetal omphalocele (Chapter 3.3), (iv) fetal talipes equinovarus (Chapter 4.1)
3. To establish prenatal cut-off levels for postnatal referral of mild renal pyelectasis (Chapter 5.1) and to determine the impact of prenatally diagnosed fetal unilateral multicystic dysplastic kidney on postnatal optimisation of diagnosis and treatment (Chapter 5.2)

## REFERENCES

- 1 Campbell S, Johnstone FD, Holt EM, May P. Anencephaly: early ultrasonic diagnosis and active management. *Lancet* 1972; 2: 1226-1227.
- 2 Robinson HP. Sonar measurement of fetal crown-rump length as means of assessing maturity in first trimester of pregnancy. *Br. Med. J.* 1973; 4: 28-31.
- 3 Heath VC, Southall TR, Souka AP, Elisseo A, Nicolaides KH. Cervical length at 23 weeks of gestation: prediction of spontaneous preterm delivery. *Ultrasound Obstet. Gynecol.* 1998; 12: 312-317.
- 4 Griffin D, Cohen-Overbeek T, Campbell S. Fetal and utero-placental blood flow. *Clin. Obstet. Gynaecol.* 1983; 10: 565-602.
- 5 Snijders RJ, Nicolaides KH. Fetal biometry at 14-40 weeks' gestation. *Ultrasound Obstet. Gynecol.* 1994; 4: 34-48.
- 6 Pilu G., Nicolaides KH. *Diagnosis of fetal abnormalities.* New York: The Parthenon Publishing Group, 1999.
- 7 Grandi C, Luchtenberg G, Rittler M. The contribution of birth defects to spontaneous preterm birth. *Am. J. Perinatol.* 2007; 24: 487-492.
- 8 Buitendijk S, Zeitlin J, Cuttini M, Langhoff-Roos J, Bottu J. Indicators of fetal and infant health outcomes. *Eur. J. Obstet. Gynecol. Reprod. Biol.* 2003; 111 Suppl 1: S66-S77.
- 9 Chung CS, Myrianthopoulos NC. Congenital anomalies: mortality and morbidity, burden and classification. *Am. J. Med. Genet.* 1987; 27: 505-523.
- 10 Reerink JD, Hengreen WP, Verkerk PH, Ruys JH, Verloove-Vanhorick SP. Congenital disorders in the first year of life. *Ned. Tijdschr. Geneesk.* 1993; 137: 504-509.
- 11 De Galan-Roosen AE, Kuijpers JC, Meershoek AP, van VD. Contribution of congenital malformations to perinatal mortality. A 10 years prospective regional study in the Netherlands. *Eur. J. Obstet. Gynecol. Reprod. Biol.* 1998; 80: 55-61.
- 12 Prevalence of congenital malformations in the Northern Netherlands 1981-2006. 2007; <http://www.rug.nl/umcg/faculteit/disciplinegroepen/MedischeGenetica/Eurocat/professionals/tabellen>
- 13 Stoll C, Tenconi R, Clementi M. Detection of Congenital Anomalies by Fetal Ultrasonographic Examination across Europe. *Community Genet.* 2001; 4: 225-232.
- 14 Smith-Bindman R, Hosmer W, Feldstein VA, Deeks JJ, Goldberg JD. Second-trimester ultrasound to detect fetuses with Down syndrome: a meta-analysis. *Jama* 2001; 285: 1044-1055.
- 15 Wladimiroff Juriy W. Routine Ultrasonography for the detection of fetal structural anomalies. In *When to screen in Obstetrics and Gynecology*, Wildschut Hajo I. J., Weiner Carl P., Peters Tim J. (eds). Saunders Elsevier: Philadelphia, 2006; 244-252.
- 16 Wladimiroff JW, Cohen-Overbeek TE, Ursem NT, Bijma H, Los FJ. Twenty years of experience in advanced ultrasound scanning for fetal anomalies in Rotterdam. *Ned. Tijdschr. Geneesk.* 2003; 147: 2106-2110.
- 17 Nicolaides KH, Heath V, Cicero S. Increased fetal nuchal translucency at 11-14 weeks. *Prenat. Diagn.* 2002; 22: 308-315.
- 18 Hegge FN, Franklin RW, Watson PT, Calhoun BC. An evaluation of the time of discovery of fetal malformations by an indication-based system for ordering obstetric ultrasound. *Obstet. Gynecol.* 1989; 74: 21-24.
- 19 Levi S. Ultrasound in prenatal diagnosis: polemics around routine ultrasound screening for second trimester fetal malformations. *Prenat. Diagn.* 2002; 22: 285-295.
- 20 Garne E, Loane M, Dolk H, De VC, Scarano G, Tucker D, Stoll C, Gener B, Pierini A, Nelen V, Rosch C, Gillerot Y, Feijoo M, Tincheva R, Queisser-Luft A, Addor MC, Mosquera C, Gatt M, Barisic I. Prenatal diagnosis of severe structural congenital malformations in Europe. *Ultrasound Obstet. Gynecol.* 2005; 25: 6-11.
- 21 Boyd PA, Bhattacharjee A, Gould S, Manning N, Chamberlain P. Outcome of prenatally diagnosed anterior abdominal wall defects. *Arch. Dis. Child Fetal Neonatal Ed* 1998; 78: F209-F213.
- 22 Jani JC, Nicolaides KH, Gratacos E, Vandercruys H, Deprest JA. Fetal lung-to-head ratio in the prediction of survival in severe left-sided diaphragmatic hernia treated by fetal endoscopic tracheal occlusion (FETO). *Am. J. Obstet. Gynecol.* 2006; 195: 1646-1650.
- 23 Grandjean H, Larroque D, Levi S. The performance of routine ultrasonographic screening of pregnancies in the Eurofetus Study. *Am. J. Obstet. Gynecol.* 1999; 181: 446-454.
- 24 Brand IR, Kaminopetros P, Cave M, Irving HC, Lilford RJ. Specificity of antenatal ultrasound in the Yorkshire Region: a prospective study of 2261 ultrasound detected anomalies. *Br. J. Obstet. Gynaecol.* 1994; 101: 392-397.
- 25 Antsaklis AJ. Debate about ultrasound screening policies. *Fetal Diagn. Ther.* 1998; 13: 209-215.
- 26 Salvesen KA. Epidemiological prenatal ultrasound studies. *Prog. Biophys. Mol. Biol.* 2007; 93: 295-300.
- 27 Leivo T, Tuominen R, Saari-Kemppainen A, Ylostalo P, Karjalainen O, Heinonen OP. Cost-effectiveness of one-stage ultrasound screening in pregnancy: a report from the Helsinki ultrasound trial. *Ultrasound Obstet. Gynecol.* 1996; 7: 309-314.
- 28 Long G, Sprigg A. A comparative study of routine versus selective fetal anomaly ultrasound scanning. *J. Med. Screen.* 1998; 5: 6-10.
- 29 Bucher HC, Schmidt JG. Does routine ultrasound scanning improve outcome in pregnancy? Meta-analysis of various outcome measures. *BMJ* 1993; 307: 13-17.

- 30 Bonnet D, Coltri A, Butera G, Fermon L, Le BJ, Kachaner J, Sidi D. Detection of transposition of the great arteries in fetuses reduces neonatal morbidity and mortality. *Circulation* 1999; 99: 916-918.
- 31 Fuchs IB, Muller H, Abdul-Khalik H, Harder T, Dudenhausen JW, Henrich W. Immediate and long-term outcomes in children with prenatal diagnosis of selected isolated congenital heart defects. *Ultrasound Obstet. Gynecol.* 2007; 29: 38-43.
- 32 Garne E, Loane M, Dolk H. Gastrointestinal malformations: impact of prenatal diagnosis on gestational age at birth. *Paediatr. Perinat. Epidemiol.* 2007; 21: 370-375.
- 33 Hoogervorst H. Beleidskader naar aanleiding van het RIVM rapport perinatale sterfte. 2005; <http://www.rivm.nl/pns/down-seo/beleidskader/index.jsp>



# 2

## Diagnosis of a neural tube defect

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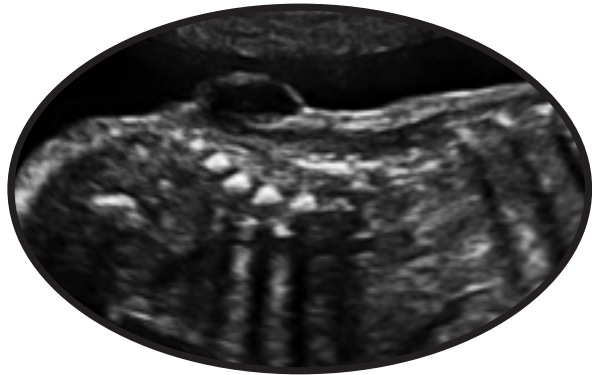
This chapter deals with the poor detection rate of spina bifida in a non-routine ultrasound screening setting. Spina bifida is compatible with life, but for the vast majority of infants it has a devastating impact on quality of life. When diagnosed before 24 weeks of gestation most couples will opt for termination of pregnancy



1



3



2

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- 1 Longitudinal image through the normal fetal spine at 20 weeks' gestation. The spine is bordered by the intact skin.
  - 2 Longitudinal section through the fetal spine with the fluid-filled myelomeningocele.
  - 3 Image of a newborn infant with a myelomeningocele at the lower end of the back.



# 2.1

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## Audit of prenatal and postnatal diagnosis of isolated open spina bifida in three university hospitals in the Netherlands

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M.A.G. Olde Scholtenhuis (1), T.E. Cohen-Overbeek (2), M. Offringa (3), P.G. Barth (4), Ph. Stoutenbeek (5), R.H. Gooskens (6), J.W. Wladimiroff (2), C.M. Bilardo (7)

- 1 Department of Obstetrics and Gynecology, Isala Clinics, Zwolle, the Netherlands
- 2 Department of Obstetrics and Gynecology, Dijkzigt Academic Hospital, Rotterdam, the Netherlands
- 3 Department of Neonatology Emma Children's Hospital, Academic Medical Center, Amsterdam, the Netherlands
- 4 Department of Pediatric Neurology, Academic Medical Center, Amsterdam, the Netherlands
- 5 Department of Obstetrics and Gynecology, University Medical Center, Utrecht, the Netherlands
- 6 Department of Child neurology, University Medical Center, Utrecht, the Netherlands
- 7 Department of Obstetrics and Gynecology, Academic Medical Center, Amsterdam, the Netherlands

# Abstract

**Objective** To audit the current Dutch policy of prenatal detection of isolated open spina bifida based on offering detailed ultrasound examination only on indication.

**Methods** A retrospective analysis of prenatally diagnosed isolated spina bifida cases and of newborns diagnosed with this condition was carried out in three university hospitals. The data were collected from databases and clinical records of the departments of prenatal diagnosis, obstetrics, neonatology, child neurology and neurosurgery of the three centers.

**Results** Between January 1996 and December 1999, 88 cases of isolated open spina bifida were diagnosed prenatally by ultrasound investigation. Thirty-eight cases (43%) were diagnosed before the 24<sup>th</sup> week of gestation. Of these, 35 (92%) ended in termination of the pregnancy at the parents' request. Of the remaining 50 cases (57%) diagnosed after the 24<sup>th</sup> week of gestation, eight (16%) pregnancies were terminated beyond the legal limit for termination due to the severity of the condition. Of the 88 cases of isolated spina bifida, 25 infants (28%) were still alive at the age of 4 years. In the same audit period 112 newborn infants with isolated open spina bifida were admitted to the neonatology, child neurology, or neurosurgery ward of the three centers. Of these cases, 47 (42%) had been diagnosed prenatally and 65 (58%) were an unexpected finding at birth. In 24 infants (21%) surgical treatment was withheld because of the severity of the condition and predicted poor outcome, whereas the remaining 88 (79%) infants underwent surgical repair.

**Conclusion:** The current practice in the Netherlands of offering ultrasound screening to high-risk patients only leads to early detection of a minority of cases of spina bifida. Most cases are diagnosed either after the 24<sup>th</sup> week of gestation or they remain undiagnosed until after birth. When spina bifida is diagnosed before the 24<sup>th</sup> week of gestation the vast majority of parents opts for termination. In order to reduce the birth prevalence of spina bifida in the Netherlands the introduction of a policy of routine ultrasound screening should be considered.

# Introduction

Neural tube defects (anencephaly and spina bifida) are the most common severe congenital defects of the central nervous system. Their prevalence varies greatly, depending on ethnic, racial and socio-economic backgrounds<sup>1</sup>. The mortality and morbidity rates associated with spina bifida (rachischisis, myelocoele and myelomeningocele) are high: 20% of affected infants die within the first year, 37% within the first decade, 44% within the second and only 50% survive beyond the third decade<sup>2</sup>. A total of 88% of the surviving infants suffer of variable degrees of life-long disabilities and are at risk for psychosocial maladjustment<sup>3,4</sup> although the intellectual performance may be normal in the vast majority of cases<sup>5</sup>.

In the 1970s, measurement of maternal serum  $\alpha$ -fetoprotein (AFP) at around 16 weeks' gestation was introduced as a screening method for open neural tube defects in countries with a high prevalence of this anomaly<sup>6</sup>. Definitive diagnosis was based on the AFP level in amniotic fluid. In the 1980s ultrasonography became the diagnostic method of choice, replacing amniocentesis almost entirely<sup>7,8</sup>. In the last decade, most European countries have adopted ultrasound examination as a routine part of antenatal care<sup>9</sup>. The combined effects of this screening strategy and use of periconceptional folic acid has led to a dramatic fall in the incidence of spina bifida at birth in many countries in and beyond Europe<sup>1,10</sup>. The practice is different in the Netherlands where both routine second-trimester ultrasonography and maternal serum AFP screening are not part of routine prenatal care. An ultrasound scan is either performed for specific obstetric indications or because of a suspected or known increased risk of fetal anomalies. The latter two categories are referred to tertiary centers for specialized ultrasonography. A baby born with spina bifida in the Netherlands is usually referred to a tertiary center for further investigations and is managed by a team of specialists. At the request of the Ministry of Health, a working party of the Dutch Health Council has evaluated the necessity of screening for neural tube defects in the Netherlands. This working party initiated the present study, which aimed to audit the current Dutch policy and its consequences in three referral areas<sup>11</sup>. The frequency of open spina bifida based on the number of defects diagnosed prenatally and at birth, parents' choices, pregnancy outcome and post-natal outcome up to a maximum of 4 years of age were documented.

## Patients and Methods

### Prenatal diagnosis

Data were collected from the files of all patients scanned between January 1996 and December 1999 at the departments of prenatal diagnosis at the Academic Medical Center in Amsterdam, the Dijkzigt Academic Hospital in Rotterdam and the University Medical Center in Utrecht. There were 88

patients in whom the ultrasound diagnosis of isolated spina bifida (spina bifida with or without meningocele) was made prenatally. The following indications prompted a scan in these pregnancies: raised AFP in maternal serum or amniotic fluid (10 cases); increased anamnestic risk for congenital defects (four cases); Rhesus isoimmunization (two cases) and use of antiepileptic drugs (one case). The indication for referral of the remaining 71 patients was suspicion of an anomaly at a scan performed elsewhere or an obstetric condition such as oligohydramnios, polyhydramnios or fetal growth restriction entailing an increased risk of congenital structural defects. Information on subsequent management and outcome of pregnancy was obtained from obstetric records. Infant survival was documented from the medical records of the departments of neonatology, child neurology and neurosurgery from a minimum of 6 months up to a maximum of 4 years of age for the children born in 1996.

## Postnatal diagnosis

The group in which the diagnosis of isolated open spina bifida (myelocele, myelomeningocele, rachischisis) was made or confirmed at birth consisted of 112 infants admitted to the neonatology, child neurology, or neurosurgery wards of the three university hospitals between January 1996 and December 1999. In 47 of these cases (42%) the condition was already known prenatally. This group included also the 25 fetuses of the above mentioned prenatal group who survived. The remaining 22 fetuses in which the diagnosis was made prenatally were referred to the above mentioned departments after birth for further treatment.

Information on the timing of diagnosis (whether prenatally or at birth), parents' choices with regard to subsequent management, pregnancy outcome, clinical management and outcome of the children up to a maximum of 4 years of age was recorded from the medical records. In the Netherlands, spina bifida teams consisting of a neonatologist, a child neurologist, a neurosurgeon, a rehabilitation physician, an orthopedic surgeon, a child urologist and a social worker assess the affected newborns and counsel the parents on the prognosis and appropriateness of surgical treatment. The criteria for judging the prognosis may have differed slightly between the three centers, but consisted mainly of the criteria described by Lorber<sup>12</sup>. Poor prognostic factors are considered to be defects higher than L2-L3, presence of hydrocephalus at birth, or presence of a gross kyphosis or major associated defects. The latter have been excluded from the study group.

# Results

## Prenatal diagnosis

During the study period 88 cases of isolated open spina bifida were detected by prenatal ultrasound examination. In only 17 (19%) cases was the indication for the scan an increased *a priori* risk of neural tube defect or other structural anomalies. The findings in the other 71 cases (81%) were unexpected.

Of the 88 prenatally diagnosed cases of isolated open spina bifida, 38 (43%) were observed before the 24<sup>th</sup> week of gestation. Thirty-five of these cases (92%) were terminated at the parents' request. In the other three cases (8%) the parents chose to continue the pregnancy but requested no surgical intervention after birth. One of these children died soon after birth and two were still alive at the end of the study's follow-up period.

From the 88 prenatally diagnosed cases of isolated open spina bifida, 50 (57%) were diagnosed after the 24<sup>th</sup> week. In eight of these 50 cases (16%) the pregnancy was terminated. This was possible even beyond the usual Dutch legal term for termination (23 completed weeks), because the severity of the condition was judged by a panel of specialists as 'life threatening'. In 20 (40%) of these late diagnoses, the parents, based on the negative advice of the spina bifida team, requested no surgical intervention after birth. A total of 23 (46%) of these 50 neonates were still alive at the end of the observation period (Table 1). The overall survival rate was 28%.

## Postnatal diagnosis

In the study period a total of 112 newborns with isolated open spina bifida were admitted to the neonatology, child neurology, or neurosurgery department of the three university hospitals. These included cases diagnosed elsewhere and referred after birth for postnatal assessment. Forty-seven (42%) of these 112 infants had been diagnosed prenatally. In 65 (58%) of these 112 infants, spina bifida was an unexpected finding at birth. Eight (12%) of these 65 cases had actually been overlooked at a third-trimester ultrasound scan performed for an obstetric indication. In 15 (32%) of the prenatally diagnosed cases and in nine (14%) of the postnatally diagnosed cases surgical repair after birth had been withheld due to the expected extremely poor prognosis (Table 2). Surgical repair was undertaken in 32 (68%) of the prenatally diagnosed cases and in 56 (86%) of the postnatally diagnosed cases. Of the 88 infants that underwent surgery, 75 (85%) had a myelomeningocele and 13 (15%) a meningocele. Hydrocephaly was present in 72 (82%) of these infants and in 69 (78%) a ventricle-peritoneal drain was inserted (Table 3). The overall survival rate in this group was 79%. Because of the still relatively young age of these 112 infants at the end of the study period, it is difficult to draw a definitive conclusion on the number and severity of handicaps and various disabilities. According to their condition

and development at the last visit a tentative prognosis could be made. The expectations was that approximately 33% would have a variable degree of mental retardation, 42% would be wheel-chair dependent, and 88% would suffer bladder dysfunction and 57% bowel dysfunction (Table 4).

## Discussion

The present audit has revealed that in these three Dutch university centers the majority of spina bifida cases occurred in couples with no apparent *a priori* risk factors. In most cases the diagnosis was made at birth or at a prenatal stage beyond the legal limit for termination. Survival was strikingly different depending on the stage of diagnosis. In the case of an ultrasound diagnosis before 24 weeks of gestation, 92% of the parents opted for termination of pregnancy (TOP). The overall survival rate of the prenatally diagnosed group was 28%, whereas in the postnatally detected cases the survival rate was 79%. This latter category constitutes the major source of morbidity.

### The potential role of ultrasound diagnosis

Of all congenital anomalies, neural tube defects are the most amenable to prenatal identification<sup>9,13</sup>. At present, ultrasound is the most accurate method of detecting neural tube defects in the second trimester of pregnancy, with a sensitivity of about 71% (95<sup>th</sup> CI, 60-80) and a specificity close to 100%<sup>14</sup>. Due to the continuous improvement of ultrasound techniques and the introduction of the ultrasonographic cranial signs associated with spina bifida ('banana-sign' and 'lemon-sign'), the diagnostic sensitivity of ultrasound has steadily increased even in units without the expertise of a tertiary center<sup>14,15</sup>. The importance of standardizing ultrasound scans at around 20 weeks' gestation-the most favorable moment for diagnostic purposes-is demonstrated by the fact that in the audit eight cases of spina bifida were overlooked on ultrasound examination carried out in the third trimester. This may have partly been attributable to lack of experience of the sonographer, or to the presence of unfavorable conditions such as maternal obesity, oligohydramnios or breech presentation masking a myelomeningocele. Hydrocephaly was not present in any of the misdiagnosed cases.

Apart from the role in diagnosing major structural anomalies<sup>9</sup> the impact of this technique on perinatal outcomes is still subject of debate<sup>16,17</sup>. The only randomised study which thus far has proven an impact of routine ultrasound on perinatal mortality has only been able to show a shift from perinatal death to TOP in cases in which structural anomalies were diagnosed during pregnancy without any clear impact on neonatal morbidity<sup>18</sup>. This argument has thus far discouraged health planners in the Netherlands

from investing resources in a screening program. Although the general feeling of obstetricians and midwives is that ultrasound is an invaluable diagnostic tool, given the widespread use of this technique and the fact that most women highly value it, it has become impossible to carry out a large randomised controlled trial to provide evidence that ultrasound screening may be justified.

## **Living with spina bifida**

Despite the improvement in surgical management of spina bifida in recent decades resulting in prolonged life expectancy, no improvement in the degree of disabilities or quality of life of affected individuals has been demonstrated<sup>2,3,5</sup>. Spina bifida patients rarely reach independence, have a satisfactory job or manage to have a partner or start a family<sup>4,19</sup>. In the USA the lifelong cost of an individual born with spina bifida has been estimated to be about \$300.000<sup>20,21</sup>. However, besides the medical and social costs it is impossible to quantify in monetary terms the psychological burden of the condition for the affected individual and his/her family. This is clearly reflected by the fact that parents, when faced with a diagnosis of spina bifida and with the option to decide on the future of the pregnancy, mostly opt for TOP. In a multicenter study conducted in several countries, Mansfield reported a termination rate associated with prenatally diagnosed spina bifida of between 20% and 100% with an average of 64%<sup>22</sup>. In the present study the termination rate for spina bifida was 92%, similar to that for Down syndrome reported in the study of Mansfield<sup>22</sup>. Also striking is the difference in survival rate among newborns in which the condition was known prenatally (28%) or only after birth (79%). The explanation for this is that those detected prenatally, even at late scans, included the most severe cases presenting with hydrocephaly, breech presentation or polyhydramnios, as clearly suggested by the higher incidence of hydrocephaly in this group.

## **Effects of prevention on the prevalence of spina bifida in the Netherlands**

Given the regional character of the audit we cannot extrapolate the obtained data to a national prevalence figure for isolated spina bifida. According to an estimate based on data from the obstetrical and pediatric registry and from the EUROCAT registry of congenital malformation in two geographical areas, the yearly prevalence of isolated spina bifida in the Netherlands is supposed to be about 0.5‰<sup>11</sup>. However, underestimation of the true prevalence, due to incomplete registration of severe cases resulting in early neonatal deaths from home deliveries, cannot be ruled out. The introduction of periconceptional folic acid supplements has been successful in the Netherlands from 1995 onwards and the estimate is that about 36%

of pregnant women are currently using folic acid<sup>23</sup>. Yet national data have thus far failed to show a significant decrease in the incidence of neural tube defects<sup>23</sup>. This may be in keeping with the suggestion that folic acid may not be all that effective in preventing the occurrence of new cases of spina bifida in low-risk women, and only shows an effect in areas with poor socioeconomic status and low dietary intake<sup>24,25</sup>

In Britain the incidence of live births, stillbirths and pregnancies terminated because of a fetal neural tube defect has fallen steadily from 1972 to 1992 from 215:100.000 to 38:100.000 and has stabilised ever since<sup>10,26</sup>. It is possible that the effect of an improved diet during pregnancy, together with a successful folic acid campaign, has reduced the prevalence of the condition to a minimum beyond which further reduction is not achievable. As a result of routine ultrasound screening 'spina bifida teams' in Britain have been made redundant as a liveborn infant with a severe form of spina bifida is now seldom seen.

The situation in the Netherlands is clearly different. A lower ethnic predisposition to spina bifida and a higher and more homogeneous socioeconomic level have been associated with a lack of a significant decline in the incidence of the condition after the introduction of folic acid supplements.

## New therapeutic initiatives

A promising new development in the therapy of spina bifida has been suggested by recently published experiences on *in utero* repair of isolated spina bifida<sup>27</sup>. The procedure is still in an early stage of development but when performed before 24 weeks of gestation it appears to improve neurologic outcome<sup>28,29</sup>. Early detection of the disease will provide parents with more choices than simply TOP

## Conclusion

An ultrasound screening policy for congenital anomalies based on 'indications' only inevitably carries a poor pick-up rate for structural anomalies. This audit shows that the present Dutch screening policy enables early prenatal detection of only a minority of cases of isolated spina bifida. The success of prenatal screening programs in most European countries and recent promising attempts to intrauterine repair of the condition should encourage a reassessment of the Dutch policy.



**TABLE 1 Indication for ultrasound scan and outcome of prenatally diagnosed isolated spina bifida**

	<24 weeks (N=38) (N(%))	≥24 weeks (N=50) (N(%))	Total (N=88) (N(%))
<b>Indication</b>			
Specific indication	17	0	17 (19)
Incidental finding	21	50	71 (81)
<b>Management</b>			
Termination	35 (92)	8 (16)	43 (49)
Intervention*	0	22 (44)	22 (25)
Non-intervention	3 (8)	20 (40)	23 (26)
<b>Outcome</b>			
Stillbirth#	35 (92)	8 (16)	43 (49)
Neonatal death	1 (3)	17 (34)	18 (21)
Alive at the end of the study period	2 (5)	23 (46)	25 (28)
Lost to follow-up	0	2 (4)	2 (2)

\* Surgical repair after birth; # Including pregnancy terminations.

**TABLE 2 Number of infants with isolated open spina bifida admitted to three university centers, and management**

	Prenatal diagnosis (N(%))	Postnatal diagnosis (N(%))	Total (N(%))
Infants	47 (42)	65 (58)	112 (100)
Non-intervention	15 (32)	9 (14)	24 (21)
Surgery	32 (68)	56 (86)	88 (79)

**TABLE 3 Clinical features and need for ventricular-peritoneal drain, in relation to the time of diagnosis, in infants undergoing surgery for isolated spina bifida**

Surgical repair	Prenatal (N=32) (68%) (N(%))	Postnatal (N=56) (86%) (N(%))	Total (N=88) (79%) (N(%))
Meningocele	5	8	13 (15)
Myelomeningocele	27	48	75 (85)
Hydrocephaly	29 (91)	43 (77)	72 (82)
Ventricular-peritoneal drain	26 (81)	43 (77)	69 (78)

**TABLE 4 Expectations concerning mental development, ability to walk and bladder/bowel dysfunction in the 112 surviving infants with isolated spina bifida**

<b>Expectation</b>	<b>N</b>	<b>%</b>
<b>Mental development</b>		
Normal development	50	57
Mental retardation	29	33
Unpredictable	7	8
Unknown	2	2
<b>Ambulation</b>		
Walk without support	16	18
Walk with instruments	25	28
Wheelchair	37	42
Unpredictable	9	10
Unknown	1	1
<b>Bladder/bowel dysfunction</b>		
Bladder dysfunction	77	88
Bowel dysfunction	50	57
Unpredictable	27	31
Continence	11	12
Unknown	2	2

## REFERENCES

- International Centre for Birth Defects, EUROCAT. World Atlas of Birth Defects. Geneva: World Health Organization, 1998; 20-31.
- Hunt GM, Poulton A. Open spina bifida: a complete cohort reviewed 25 years after closure. *Dev. Med. Child Neurol.* 1995; 37: 19-29.
- Date I, Yagyu Y, Asari S, Ohmoto T. Long-term outcome in surgically treated spina bifida cystica. *Surg. Neurol.* 1993; 40: 471-475.
- Zurmohle UM, Homann T, Schroeter C, Rothgerber H, Hommel G, Ermett JA. **Psychosocial adjustment of children with spina bifida.** *J. Child Neurol.* 1998; 13: 64-70.
- Hagelsteen JH, Lagergren J, Lie HR, Rasmussen F, Borjeson MC, Lagerkvist B, Mutilainen M, Taudorf K, Kohler L. Disability in children with myelomeningocele. A Nordic study. *Acta Paediatr. Scand.* 1989; 78: 721-727.
- Knight, G. J. and Palomaki, G. E. Maternal serum alpha-fetoprotein and the detection of open neural tube defects. In *Maternal Serum Screening for Fetal Genetic Disorders*, Elias S (ed.) New York, NY: Churchill Livingstone, 1992; 41-58.
- Wald, N. J. and Cuckle, H. S. Open neural-tube defects. In *Antenatal and Neonatal Screening*, Wald NJ (ed). Oxford: Oxford University Press, 1984; 25-73.
- Ennever FK, Lave LB. Parent preferences and prenatal testing for neural tube defects. *Epidemiology* 1995; 6: 8-16.
- Grandjean H, Larroque D, Levi S. The performance of routine ultrasonographic screening of pregnancies in the Eurofetus Study. *Am. J. Obstet. Gynecol.* 1999; 181: 446-454.
- Morris JK, Wald NJ. Quantifying the decline in the birth prevalence of neural tube defects in England and Wales. *J. Med. Screen.* 1999; 6: 182-185.
- Health Council of the Netherlands. Prenatal Screening: Down Syndrome, Neural Tube defects, Routine Ultrasonography. Publication No. 200/11. The Hague: Health Council of the Netherlands, 2001.
- Lorber J. Results of treatment of myelomeningocele. An analysis of 524 unselected cases, with special reference to possible selection for treatment. *Dev. Med. Child Neurol.* 1971; 13: 279-303.
- Boyd PA, Chamberlain P, Hicks NR. 6-year experience of prenatal diagnosis in an unselected population in Oxford, UK. *Lancet* 1998; 352: 1577-1581.
- Vos JM, Offringa M, Bilardo CM, Lijmer JG, Barth PG. Sensitive and specific screening for detection of spina bifida by echography in the second trimester; systematic review and meta-analysis. *Ned. Tijdschr. Geneesk.* 2000; 144: 1736-1741.
- Nicolaides KH, Campbell S, Gabbe SG, Guidetti R. Ultrasound screening for spina bifida: cranial and cerebellar signs. *Lancet* 1986; 2: 72-74.
- Ewigman BG, Crane JP, Frigoletto FD, LeFevre ML, Bain RP, McNellis D. Effect of prenatal ultrasound screening on perinatal outcome. RADIUS Study Group. *N. Engl. J. Med.* 1993; 329: 821-827.
- Eik-Nes SH, Salvesen KA, Okland O, Vatten LJ. Routine ultrasound fetal examination in pregnancy: the 'Alesund' randomized controlled trial. *Ultrasound Obstet. Gynecol.* 2000; 15: 473-478.
- Saari-Kemppainen A, Karjalainen O, Ylostalo P, Heinonen OP. Ultrasound screening and perinatal mortality: controlled trial of systematic one-stage screening in pregnancy. The Helsinki Ultrasound Trial. *Lancet* 1990; 336: 387-391.
- Staal-Schreinemachers AL, Vos-Niel JM, Begeer JH. Future prospects for children with spina bifida aperta. *Ned. Tijdschr. Geneesk.* 1996; 140: 1268-1272.
- Botto LD, Moore CA, Khoury MJ, Erickson JD. Neural-tube defects. *N. Engl. J. Med.* 1999; 341: 1509-1519.
- Waitzman, N. J, Scheffler, R. M., and Romano, P. S. An assessment of total costs and policy implications. In *The Cost of Birth Defects: Estimates of the Value of Prevention*, Waitzman NJ (ed.). Lanham, MD: University Press of America: 1996; 145-147.
- Mansfield C, Hopfer S, Marteau TM. Termination rates after prenatal diagnosis of Down syndrome, spina bifida, anencephaly, and Turner and Klinefelter syndromes: a systematic literature review. *European Concerted Action: DADA (Decision-making After the Diagnosis of a fetal Abnormality)*. *Prenat. Diagn.* 1999; 19: 808-812.
- van der Pal-de Bruin KM, Buitendijk SE, Hirasings RA, den Ouden AL. Prevalence of neural tube defects in births before and after promotion of periconceptional folic acid supplementation. *Ned. Tijdschr. Geneesk.* 2000; 144: 1732-1736.
- Berry RJ, Li Z, Erickson JD, Li S, Moore CA, Wang H, Mulinare J, Zhao P, Wong LY, Gindler J, Hong SX, Correa A. Prevention of neural-tube defects with folic acid in China. China-U. S. Collaborative Project for Neural Tube Defect Prevention. *N. Engl. J. Med.* 1999; 341: 1485-1490.
- Kalter H. Folic acid and human malformations: a summary and evaluation. *Reprod. Toxicol.* 2000; 14: 463-476.
- Kadir RA, Sabin C, Whitlow B, Brockbank E, Economides D. Neural tube defects and periconceptional folic acid in England and Wales: retrospective study. *BMJ* 1999; 319: 92-93.
- Aaronson OS, Tulipan NB, Cywes R, Sundell HW, Davis GH, Bruner JP, Richards WO. Robot-assisted endoscopic intrauterine myelomeningocele repair: a feasibility study. *Pediatr. Neurosurg.* 2002; 36: 85-89.
- Kitano Y, Flake AW, Crombleholme TM, Johnson MP, Adzick NS. Open fetal surgery for life-threatening fetal malformations. *Semin Perinatol.* 1999; 23: 448-461.
- Walsh DS, Adzick NS, Sutton LN, Johnson MP. The Rationale for in utero repair of myelomeningocele. *Fetal Diagn. Ther.* 2001; 16: 312-322.



# 3

## Diagnosis of gastro-intestinal anomalies and abdominal wall defects

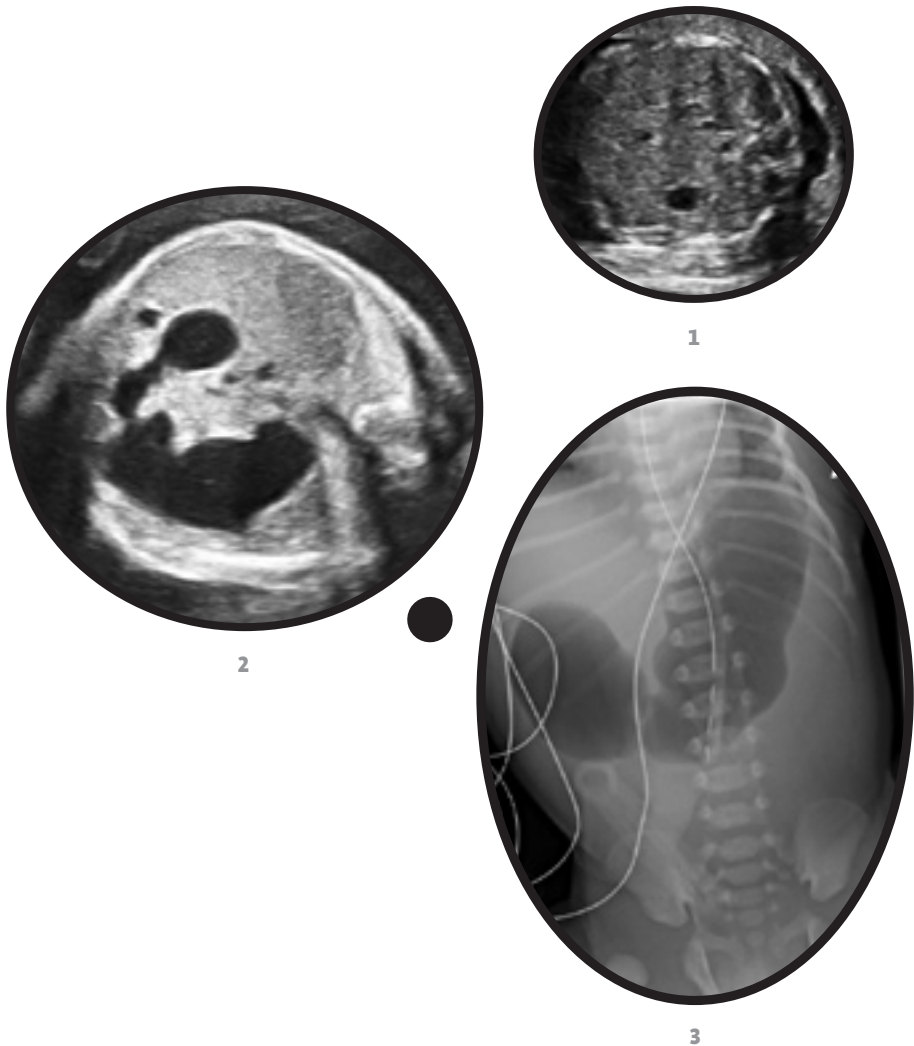
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In this chapter, perinatal outcome following prenatal diagnosis versus postnatal diagnosis is presented for three major fetal anomalies at abdominal level, notable duodenal obstruction (chapter 3. 1), gastroschisis (chapter 3. 2), and omphalocele (chapter 3. 3).

Whereas the latter two anomalies should be picked up at an 18-22 week ultrasound scan, this is not the case for duodenal obstruction, which is usually detected as late as the third trimester of pregnancy.

Gastroschisis and omphalocele are both abdominal wall defects. Gastroschisis is usually diagnosed as a simple anomaly for which perinatal outcome depends on the subsequent occurrence of intestinal complications.

Omphalocele is clearly associated with other anomalies and numerical chromosomal abnormalities, such as trisomy 18 and 13.



- 1 Transverse plane of section through the normal fetal abdomen with the spine at 3 o'clock and stomach at 6 o'clock.
- 2 Transverse plane of section through the fetal abdomen showing the dilated stomach and duodenum (double-bubble phenomenon). A connection between the two dilated structures is visible, indicating an obstruction of the duodenum.
- 3 X-ray of newborn infant with duodenal obstruction showing the air-filled stomach and duodenum

# 3.1

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## Isolated or non-isolated duodenal obstruction: perinatal outcome following a prenatal diagnosis or a diagnosis only after birth

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T.E. Cohen-Overbeek (1), E.W.M. Grijseels (1), N.D. Niemeijer (1),  
W.C.J. Hop (2), J.W. Wladimiroff (1), D. Tibboel (3)

1 Department of Obstetrics and Gynecology, Division of Obstetrics and Prenatal Medicine, Erasmus MC, Rotterdam, the Netherlands

2 Department of Epidemiology & Biostatistics, Erasmus MC, Rotterdam, the Netherlands

3 Department of Pediatric Surgery, Erasmus MC, Rotterdam, the Netherlands

# Abstract

**Objectives** To determine whether the pre- or postnatal diagnosis of either isolated or non-isolated duodenal obstruction (DO) is associated with different outcomes.

**Methods** A single center retrospective analysis was carried out of 91 cases diagnosed with a DO between January 1991 and June 2003. Data on the diagnosis, treatment and outcomes of the cases were gathered, and differences between the groups were analyzed.

**Results** 28 cases were diagnosed before and 63 after birth. Of 15 presumed isolated cases in the prenatal group four revealed associated or chromosomal anomalies after birth. The prenatally (N=11) and postnatally (N=27) detected subsets of isolated DO were significantly different for the type of obstruction. The prenatal subset displayed a lower median gestational age at delivery, lower median birth weight and a higher prematurity rate (8/11 versus 8/27). After delivery the detection of DO occurred significantly later in the postnatal subset. In the non-isolated cases with DO no difference existed for the type of chromosomal or associated anomaly or the type of obstruction between the prenatal (N=17) and postnatal subset (N=36). Trisomy 21 was present in 7/17 (41%) versus 22/36 (61%) cases. Two terminations and three intrauterine deaths occurred in the prenatal non-isolated subset. The liveborn infants from the prenatally (N=12) detected non-isolated subset showed a significantly higher prematurity rate (9/12 versus 14/36), lower median birth weight and earlier diagnosis after delivery. After surgery outcome was similar between both subsets of either isolated or non-isolated DO. All infants with an isolated DO survived. Neonatal death occurred in three prenatally and five postnatally diagnosed cases with non-isolated DO.

**Conclusions** Outcome of prenatally and postnatally diagnosed DO is not essentially different despite more prematurity and a lower birth weight. Of the prenatal cases of DO assumed isolated, 25% revealed additional chromosomal or associated anomalies after delivery, which influenced outcome.



# Introduction

Duodenal obstruction (DO) is a congenital abnormality which occurs in 1:3 per 10,000 live births<sup>1,2</sup> and is the small-bowel obstruction most commonly detected during fetal life<sup>3</sup>. Approximately half of the cases is associated with other abnormalities<sup>4</sup> (non-isolated DO), particularly cardiac, vertebral, renal and gastrointestinal such as anal atresia and tracheoesophageal atresia<sup>5</sup>. DO is complicated in 30% with trisomy 21<sup>5</sup>.

Intrinsic DO is presumed to result from failed canalisation of the lumen in the 9<sup>th</sup>-10<sup>th</sup> week of embryonic life leading to stenosis or atresia<sup>6</sup>. It is still not clear in what way disturbed intercellular contact or abnormal resolution forms an etiological basis for intrinsic duodenal obstruction, as appropriate animal models are lacking for this specific anomaly. Compression from a surrounding annular pancreas or peritoneal bands<sup>5</sup> may be the cause of extrinsic DO. The sonographic image of DO is characterized by a 'double-bubble' appearance as a result of dilatation of the stomach and duodenum. A connection between the two dilated structures can always be demonstrated. Often polyhydramnios develops during the late second or third trimester of pregnancy<sup>7</sup>. A timely prenatal diagnosis of DO would allow proper counselling of the couple<sup>2</sup> regarding additional karyotyping and adjustment of obstetric management. In case of severe associated anomalies and/or aneuploidy, there may be the option of pregnancy termination if detected early in pregnancy. However, only just over half of the cases of DO are diagnosed prenatally<sup>3</sup> and often beyond 24 weeks of gestation. This may be due to the fact that only small amounts of amniotic fluid are swallowed by the fetus during the first half of pregnancy<sup>8</sup>. The reduced reabsorption capacity in duodenal atresia and the increase in swallowing of amniotic fluid as pregnancy progresses leads to the 'double-bubble' appearance and polyhydramnios. At delivery bile stained amniotic fluid may be present due to intrauterine vomiting, sometimes mistaken as meconium containing amniotic fluid. Few studies have compared the outcome of a DO detected before birth and a diagnosis first made after birth. Neonatal morbidity was reduced in the prenatally diagnosed subset of DO in one study<sup>9</sup>. In two other studies, the age of the newborn at diagnosis and at surgery was significantly earlier resulting in less metabolic complications<sup>10</sup> and significantly earlier start of intestinal feeding<sup>11</sup> in cases diagnosed prenatally. However, no impact on overall outcome could be demonstrated by others which was mainly attributed to the high incidence of associated major anomalies<sup>4,12</sup>. In these and other studies<sup>13-15</sup> the outcome of fetuses and infants with or without associated anomalies has not been assessed separately. For parental counselling and obstetric management this information is of paramount importance as there is a suggestion that outcome in the absence of associated anomalies (isolated DO) may be excellent<sup>4</sup>. In our study the following questions were addressed: (i) what is

the impact of prenatal diagnosis on outcome of isolated and non-isolated DO; (ii) is there a difference in outcome between prenatally diagnosed and postnatally diagnosed isolated or non-isolated DO in liveborn infants; (iii) is there a difference in the type of obstruction between the prenatally and postnatally established subsets.

## Methods

The Erasmus University Medical Center serves as the single referral center for both fetal anomaly scanning and pediatric surgery in the South West of the Netherlands, a referral area with 35.000 newborns/year. A retrospective data analysis was carried out of all fetuses and infants from singleton pregnancies diagnosed with a congenital DO either pre- or postnatally during the period January 1991-June 2003. Cases were collected from the ultrasound database of the Division of Prenatal Medicine and the patient database of the Department of Pediatric Surgery. Inclusion criteria were prenatal diagnosis and postnatal treatment in our center or postnatal treatment in our center without a prenatal diagnosis elsewhere.

Additional investigations included fetal karyotyping and serial sonographic follow-up of fetal growth and amniotic fluid volume. DO was considered non-isolated if other anomalies were also present. This included, but was not limited to, aneuploidy. In all other cases DO was classified as isolated. Fetal growth restriction (FGR) was defined as a fetal upper abdominal circumference below the 10<sup>th</sup> centile for gestational age. Polyhydramnios was defined as an amniotic fluid volume above the 95<sup>th</sup> centile<sup>16</sup> for gestational age. Amniotic fluid drainage was performed in case of massive polyhydramnios and extreme maternal discomfort. Prematurity was defined as a delivery before 37 weeks of gestation. Infants with a birth weight below the 10<sup>th</sup> centile adjusted for gender<sup>17</sup> were considered small-for-gestational age (SGA). Counselling by a pediatric surgeon was offered to all couples during the prenatal period. All women with a prenatal diagnosis of duodenal obstruction were delivered vaginally unless obstetric reasons dictated otherwise.

In the majority of cases whether presented before or after birth the diagnosis was confirmed by plain abdominal X-ray. Occasionally, a contrast swallow examination, an ultrasound examination of the abdomen or a combination of the above methods was needed. When a termination of pregnancy (<24 weeks) or intrauterine death had occurred the obstruction was determined at post mortem examination if permission was granted.

Surgery was planned as an elective procedure as soon as the metabolic status had stabilized and presence or absence of associated anomalies was assessed. All newborns were evaluated by a consultant clinical geneticist and karyotyping was performed in case of suspicion of syndromal (Down syndrome in most cases) or other dysmorphic features. At surgery the nature

of the duodenal obstruction, i.e. atresia, stenosis or membrane as well as the presence or absence of an annular pancreas and malrotation was established. Atresia was defined as complete obstruction of the duodenum. A limited passage of the duodenum was considered a stenosis. Depending on the type and level of obstruction a duodenoduodenostomy, a duodenojejunostomy, a duodenogastrostomy or a membrane resection was performed. The necessity of more than one day artificial ventilation was recorded. Due to motility disorders of the bowel, infants were primarily given parenteral nutrition (TPN) and enteral feeding was instituted as soon as stomach retentions diminished. Adverse neonatal outcome was defined as neonatal death or complications resulting from the abnormality itself or the subsequent surgical procedure.

Notes were reviewed for fetal sonographic data, neonatal and surgical outcome with a minimal follow up of 12 months. Statistical analysis of comparing groups was performed using the Mann-Whitney test or Fisher's exact test of continuous or categorical data, respectively.  $P=0.05$  (two-sided) was considered the limit of statistical significance.

## Results

The total study group consisted of 109 cases, of which 30 were prenatally and 79 postnatally diagnosed. Of the prenatal group, two were delivered and treated elsewhere and were therefore excluded. Sixteen out of 79 cases in the postnatal group were excluded because of a prenatal diagnosis elsewhere. Of the remaining 28 cases in the prenatal group, 15 were initially considered isolated. However, after delivery additional anomalies were recorded in three cases involving a membranous ventricular septal defect and coarctation of the aorta, a case of hemivertebra at C2-C3, combined with only 10 right sided ribs and a case of esophageal atresia without tracheoesophageal fistula. One patient presented at 36 weeks with polyhydramnios and a 'double-bubble' phenomenon. Amniocentesis was performed but trisomy 21 without associated anomalies was revealed only after delivery at 38 weeks of gestation.

### **Prenatally diagnosed isolated versus non-isolated DO (N=28)**

Details of the prenatal group comparing the isolated (N=11) and the non-isolated subset (N=17) of DO are given in Table 1. The median gestational age at detection of the DO for the isolated subset (33.3 wks) was significantly ( $p<0.05$ ) higher than for the non-isolated subset (31.0 wks). Polyhydramnios was present in 9/11 (82%) isolated cases and in 13/17 (76%) non-isolated cases, and mostly established after 24 weeks of gestation. Amniotic fluid drainage was required in four cases of the isolated subset and in one case of the non-isolated subset.

The nature of the associated anomalies in the non-isolated subset was mainly cardiac, gastro-intestinal and skeletal (vertebral). In five cases of the non-isolated subset additional anomalies in the gastrointestinal tract were only diagnosed after delivery. These included one case of anal atresia and sacral vertebral anomalies, one case of apple peel atresia of the proximal jejunum and intestinal malrotation and three cases of esophageal atresia, twice associated with a tracheoesophageal fistula. Karyotyping was carried out in 10/11 isolated cases with normal results. However, the karyotype was abnormal in 7/17 (41%) non-isolated cases, each representing trisomy 21. Only two patients presented with a 'double-bubble' before 24 weeks of gestation and in both cases a termination of pregnancy was requested and performed. Trisomy 21 was established in one case without associated structural anomalies. The second case revealed an additional esophageal atresia without a tracheoesophageal fistula at the post mortem examination. Three cases of intrauterine death occurred at 33.4, 33.6 and 38 weeks of gestation, respectively. In the first case an additional esophageal atresia with fistula was diagnosed at autopsy. The latter two were associated with trisomy 21. One showed a grossly dilated duodenum with stenosis and growth below the 10<sup>th</sup> centile at autopsy, while in the other case autopsy was refused.

No significant difference existed between the prenatally diagnosed isolated versus prenatally diagnosed non-isolated subsets for median gestational age at delivery, prematurity rate, spontaneous delivery rate, median birth weight and fetal gender. Three neonatal deaths occurred in the non-isolated subset. In all cases delivery was premature at 29, 30 and 34 weeks of gestation. The first case was associated with a transposition of the great arteries and the second case revealed ventriculomegaly, Tetralogy of Fallot and a multicystic kidney. In the latter the anomalies were confirmed and additionally anal atresia and sacral vertebral anomalies were diagnosed, consistent with a VACTERL association. A duodenoduodenostomy was performed but neonatal death occurred at the age of 10 days. In the third case the fetus was known with a multicystic kidney and an apple peel jejunal atresia was diagnosed at surgery. The post operative period was complicated by encephalitis and meningitis and the infant died on day 19. The combined mortality (IUD and neonatal death) between the isolated (0/11) and non-isolated (6/17) subset did not reach statistical significance.

## **Prenatally versus postnatally diagnosed isolated DO (N=38)**

When comparing the prenatally (N=11; 29%) and postnatally detected subset of isolated DO (N=27; 71%) (Table 2), an overall significant difference for the type of obstruction (intrinsic versus extrinsic) was established. Intrinsic obstruction was significantly more common in the group with a postnatal diagnosis ( $p=0.017$ ). Due to the small numbers it was impossible to determine which subtype of obstruction led to this significant difference.

Data comparing delivery and outcome of infants between prenatally (N=11) and postnatally (N=27) diagnosed isolated DO are shown in Table 3. Whereas median gestational age at delivery and median birth weight were significantly lower, prematurity rate was significantly higher in the prenatal subset compared with the postnatal subset. The overall mode of delivery was similar in both subsets. The mode of delivery could not be retrieved in 3/27 cases in the postnatal subset. After delivery the detection of duodenal obstruction occurred significantly later ( $p<0.001$ ) in the postnatal subset with only 10/27 infants diagnosed before day 4. The method of diagnosis in the prenatal subset only required an abdominal x-ray, whereas in the postnatal subset several different and sometimes multiple methods were necessary to establish the diagnosis.

The nature of the surgical correction was significantly different ( $p=0.028$ ) between the prenatally diagnosed isolated versus postnatally diagnosed isolated subset. This could be contributed to the duodenoduodenostomy which was performed in all 11 prenatally detected cases as opposed to only 15/27 (55%) postnatally detected cases of DO. In the period after surgery no significant differences were seen between the pre- and postnatal subset for the time on parenteral nutrition, infants requiring more than one day artificial ventilation, the number of complications occurring and the length of hospital stay. Gastrointestinal complications required a total of 12 readmissions for eight infants in the postnatal subset as opposed to only one in the prenatal subset. All infants with an isolated DO survived.

## **Prenatally versus postnatally diagnosed non-isolated do (N=53)**

In the presence of non-isolated DO, 17 (32%) cases were detected prenatally and 36 (68%) cases postnatally. No significant difference was found for the type of chromosomal anomaly or congenital organ defect (Table 4) and no overall difference could be established for the type of obstruction between the two subsets (Table 5).

Data comparing delivery and outcome of liveborn infants between prenatally (N=12) and postnatally (N=36) detected non-isolated DO are given in Table 6. The prematurity rate was significantly higher and median birth weight significantly lower in the prenatal subset compared with the postnatal subset. Gestational age at delivery and overall mode of delivery did not reach a significant difference. For ten cases in the postnatal subset the delivery mode could not be retrieved. Both the prenatal and postnatal subset of non-isolated DO contained a high proportion of SGA infants (6/12 (50%) versus 14/36 (39%)). One case from the prenatal subset and one case from the postnatal subset was excluded from the analysis referring to treatment. This was due to demise within one week of delivery and no surgical intervention as a result of the associated anomalies. A significant difference ( $P<0.001$ ) was revealed for the age at diagnosis between the prenatal and postnatal subset of non-isolated DO. All cases of the prenatal subset were confirmed

before day 2. In the postnatal subset only 11/36 infants were diagnosed before day 4. Twelve infants were diagnosed between day 8 and day 814 (median day 116); 10 cases with a duodenal membrane obstruction and two cases with an annular pancreas. The methods required to reach a diagnosis and the nature of the surgical procedure were not significantly different between the prenatal and postnatal subset of non-isolated DO. In the period after surgery no difference was established between the two subsets for days on parenteral nutrition, complication rate, readmissions for gastrointestinal complications, and length of hospital stay. Significantly ( $p=0.03$ ) more infants required artificial ventilation for more than one day in the prenatal (5/12) versus the postnatal subset (3/36) of non-isolated DO. Neonatal death occurred in three prenatally and five postnatally diagnosed infants with DO. There were 29 cases of Down syndrome, seven of which were diagnosed in the prenatal subset of non-isolated DO and 22 cases in the postnatal subset of non-isolated DO. In one case the cause of obstruction was not established as autopsy was refused after intrauterine death had occurred. No overall difference could be established for the type of obstruction between the prenatal and postnatal subset with trisomy 21 and DO (Table 7).

## Discussion

This is a first report on the perinatal outcome of DO in which isolated or non-isolated cases diagnosed prenatally or after birth are analyzed separately. Both in the isolated and non-isolated form birth weight was significantly lower and prematurity rate significantly higher in the prenatal subsets. Additionally the gestational age at delivery was significantly lower in the prenatal isolated subset. Whereas the age at postnatal confirmation of the DO was significantly lower in both prenatal isolated and non-isolated subsets, perinatal outcome after surgery was similar between both pre- and postnatal isolated and non-isolated subsets. Intrauterine and neonatal death only occurred in the presence of non-isolated DO. Of the prenatal cases of DO presumed isolated, 25% revealed additional chromosomal or associated anomalies after delivery.

A recent analysis of 1047 liveborn infants with gastrointestinal malformations such as esophageal atresia, duodenal obstruction, omphalocele, diaphragmatic hernia or a combination of these anomalies from 14 population based registries of congenital malformations (EUROCAT)<sup>18</sup> pointed at the significantly shorter gestational age at delivery in the prenatally detected cases compared to those detected after birth. Although in this study the delivery mode was not registered, it was suggested that the shorter gestational age at birth in prenatally detected cases was due to clinicians planning the delivery under strictly controlled conditions. This is supported by a study on the effect of prenatal diagnosis on outcome of D-Transposition of the great arteries<sup>19</sup>, in which a reduced spontaneous

delivery rate and a lower birth weight was noted in the group with a prenatal diagnosis compared to the group with a diagnosis only at birth. In our study the mode of delivery was recorded in all prenatal cases, but data on delivery mode were missing in 3/27 postnatal isolated and 10/36 postnatal non-isolated cases. Even when the missing data on delivery mode were all replaced by a spontaneous vaginal delivery, the overall delivery mode would not show any significant difference between both the prenatal and postnatal subsets. The shorter gestational age at birth, the higher prematurity rate and lower birth weight in the prenatal isolated subset are more likely due to the difference in type of obstruction between these two isolated subsets, as demonstrated in Table 2. The postnatal isolated subset required no intervention during pregnancy. However, in the prenatal isolated subset, polyhydramnios gave rise to amniotic fluid drainage in 4/11 cases, presenting a risk factor for premature labor. No difference in type of associated anomaly or type of obstruction was recorded between the prenatal and postnatal non-isolated subset and only 1/17 prenatal non-isolated cases required amniotic fluid drainage. This does not explain the higher prematurity rate and lower birth weight in the prenatal non-isolated subset compared to the postnatal non-isolated subset. Other factors may have contributed to the high prematurity rate. Women who are informed that their offspring suffers a severe congenital anomaly experience increased psychological instability<sup>20</sup>. It has been well documented that stress and pregnancy related anxiety are associated with spontaneous preterm birth<sup>21,22</sup>.

Increased morbidity and mortality are described following preterm and near-term delivery<sup>23,24</sup>. The major morbidity from late prematurity may be respiratory. The prenatal non-isolated subset with DO showed an increased need for ventilatory support of more than one day. This subset contained significantly more premature infants although there was no difference in gestational age at birth compared to the postnatal non-isolated subset with DO. According to unit practice neonates will receive artificial ventilation for one day following upper abdominal surgery. Older infants, beyond the immediate postnatal age, do not need this support. At least 8 infants from the postnatal non-isolated subset underwent surgery beyond 3 months of age which explains this different requirement for artificial ventilation. Other post surgery parameters assessing outcome did not reveal a difference between the non-isolated prenatal and postnatal subsets. Despite increased prematurity and a lower gestational age at delivery in the prenatal isolated subset with DO the outcome following surgery was similar compared to the postnatally detected isolated subset. Earlier confirmation of the diagnosis and earlier surgery may have contributed to this favorable result. Similar to our findings, previous studies comparing pre- and postnatally established DO, describe earlier confirmation of the diagnosis and earlier surgery in prenatally detected DO with less metabolic complications before surgery<sup>9,10</sup> or earlier feeding transition<sup>11</sup> but no difference in length of hospital stay<sup>11</sup> and overall outcome<sup>4</sup>. Contrary to our investigation, gestational age at birth

was not taken into account in these studies. The need for artificial ventilation has not been assessed previously in this context.

In a study of prenatally and postnatally diagnosed small bowel atresia<sup>25</sup>, it was demonstrated that a prenatal diagnosis was associated with a lower birth weight and a longer period of parenteral feeding and hospital stay. It was presumed that small bowel atresia diagnosed before birth was associated with a more pronounced in utero distension of the bowel and hence a relative longer period of postnatal gut dysfunction, even after earlier repair. Alternatively, a more severe form of obstruction may have been the reason for the prenatal detection of DO when compared to cases detected after birth. In our series, polyhydramnios was present in 81% of the prenatal isolated and in 76% of the prenatal non-isolated subset. In two studies the incidence of polyhydramnios has been reported as high as 75% and 80% in the prenatal group, as opposed to 14% and 32% in the postnatal group<sup>4,10</sup>. Postnatal series have reported maternal polyhydramnios as a complication of DO in only 32-48%<sup>13,15,26</sup>.

Clinical symptoms that lead to the postnatal diagnosis of DO are bilious vomiting, distended abdomen and failure to pass meconium<sup>5</sup>. However, in the presence of complete obstruction non-bile stained vomit and normal passage of meconium have been documented in 39% and 30% of cases, respectively<sup>15</sup>. Membranous DO, in a number of cases containing a hole resulting in intermittent or less severe intestinal obstruction, has been reported to be associated with other anomalies in 75% of cases, mostly presenting trisomy 21<sup>27</sup>. Here, the obstruction is often associated with non-bilious vomiting and failure to thrive. A diagnosis of obstruction may be delayed as symptoms are frequently attributed to the other anomalies present<sup>9</sup>. In our study, 10/25 cases of membranous DO were of the isolated form, of which nine were diagnosed postnatally. The latter attributed to the majority of DO cases diagnosed between day 5 and day 39. In the postnatal non-isolated subset the delay in diagnosis was even more striking. Ten out of twelve cases of DO diagnosed between day 8 and day 814 represented the membranous form of DO, of which nine were associated with trisomy 21. A delayed diagnosis of DO is frequently reported<sup>11,15</sup> with in some cases a delay of up to 3, 4 or even 14 years<sup>14,27,28</sup>. Membranous obstruction can result in major diagnostic challenges as it results in intermittent feeding problems delaying an ultrasound examination or a contrast swallow examination. Especially the alternating periods of normal enteral feeding and vomiting is troublesome in many cases. Moreover in Down syndrome numerous reasons are present for recurrent refusal of enteral intake. The possibility of DO in these infants should be acknowledged.

Until January 2006 maternal age related detection of fetal chromosomal anomalies was only provided for women of 36 years and older. Screening tests such as the triple test or nuchal translucency measurement were not available for the general pregnant population. This may explain the high incidence of Down's syndrome cases (22/63, 35%) in the postnatal subset.



At the time of the study routine ultrasound to detect fetal anomalies was not a policy in the Netherlands. All pregnant women underwent an ultrasound scan in the first trimester to determine the gestational age but no further scans were performed unless an abnormal clinical finding would indicate this. This policy may explain why only 28 (30%) out of 91 cases were detected prenatally. Although this detection rate is double that of a study that had started in the seventies<sup>26</sup>, it is far from the 58% prenatal detection rate reported by Singh et al.<sup>14</sup> during the same period as the present investigation. Theoretically one may assume that in countries where 3rd trimester ultrasound scan is common practice more cases with DO will be detected prenatally. No such studies have been published so far.

In the entire prenatal group the non-isolated cases are diagnosed significantly earlier than the isolated cases, confirming data from the EUROSCAN study group<sup>3</sup>. Polyhydramnios was present in more than 75% in both isolated and non-isolated cases but only two non-isolated cases were detected before 24 weeks of gestation. One case was complicated by polyhydramnios and trisomy 21 and in the other esophageal atresia without a fistula was established after termination of pregnancy. In an earlier study<sup>2</sup>, four out of five cases of DO detected before 24 weeks of gestation were non-isolated with two presenting a chromosomal anomaly. The mechanism responsible for the early appearance of the 'double-bubble' sign in case of a chromosomal anomaly is not clear. Early diagnosis of DO has been reported in combination with esophageal atresia<sup>29,30</sup>. It has been suggested that accumulations of secretions in the closed loop of stomach and duodenum in case of DO with esophageal atresia without fistula<sup>31</sup> seem responsible for the early dilatation of stomach and duodenum.

Perinatal outcome between the prenatal isolated and non-isolated subset did not reveal any significant difference except for the high incidence of intrauterine and neonatal death in the non-isolated subset. This is in agreement with previous studies<sup>32,33</sup> in which intrauterine death is reported in non-isolated DO. Brantberg et al.<sup>7</sup>, however, described two cases of intrauterine death occurring in isolated DO.

Mortality in our study only involved the non-isolated cases with an incidence in the pre- and postnatal subset of 40% and 14%, respectively. The percentage in the prenatal non-isolated subset seems high compared to percentages between 17 and 24% reported in the literature<sup>2,7</sup>. These series, however, did not differentiate between isolated and non-isolated cases of DO.

For counselling purposes one must realize that not all associated or chromosomal anomalies are known at the time of the ultrasound detection of DO. Of the cases in our prenatal subset presumed isolated, 25% proved to be non-isolated after delivery with serious consequences for the outcome. This phenomenon is known in cases of prenatally suspected isolated talipes equinovarus<sup>34</sup>, but not yet described in the presence of DO. The nature of some of the anomalies, typically associated with DO such as esophageal

atresia with fistula, anal atresia, vertebral and rib anomalies are difficult to detect by antenatal ultrasound<sup>35,36</sup> and may therefore explain the relatively high percentage of missed anomalies.

It can be concluded that the overall outcome of liveborn infants after a pre- or postnatal diagnosis of isolated or non-isolated DO is similar despite more prematurity and a lower birth weight in the subsets with a prenatal diagnosis. The age at postnatal confirmation of the DO is significantly lower in both prenatal subsets and this may have contributed to the favorable outcome. Intrauterine and neonatal death only occurred in the presence of associated or chromosomal anomalies. Of the prenatal cases of DO assumed isolated, 25% revealed additional chromosomal or associated anomalies after delivery which influenced outcome.

**TABLE 1 Prenatal, delivery and outcome data from the prenatally detected isolated and non-isolated cases with duodenal obstruction.**

	Isolated N=11	Non-isolated N=17	P
GA at detection (weeks)	33.3 (27.8-37.3)	31.0 (20.3-36)	<0.05
Polyhydramnios <24 weeks	-	1	-
Polyhydramnios >24 weeks	9	12	1.00
Amniotic fluid drainage	4	1	0.15
Fetal growth retardation	1	2	1.00
Associated anomalies			
Cardiac	-	5	NT
Gastro-intestinal tract	-	5	NT
Renal tract	-	2	NT
Central nervous system	-	1	NT
Vertebral	-	4	NT
Other	-	3	NT
Karyotyping performed	10	17	0.39
Abnormal karyotype	-	7	NT
Intrauterine death (IUD)	0	3	0.26
Termination of pregnancy (TOP)	0	2	0.51
Delivery* <37w	8	9	1.00
Delivery mode overall			0.85
Spontaneous delivery *	6	8	0.68
Induced delivery*	2	1	0.57
Cesarean section*	3	3	1.00
GA at delivery (weeks) *	35.4 (30.1-39.3)	35.7 (29.3-39.1)	0.56
Birth weight (g) *	2320 (1455-3480)	1917 (1155-3390)	0.14
Birth weight <10 <sup>th</sup> centile *	3	6	0.40
Male/Female	7/4	9/8	0.70
Neonatal death * (NND)	0	3	0.22
Combined mortality (IUD + NND)	0	6	0.055

Data presented as median and range. GA, gestational age; g, gram; NT, not tested.

\* excluding cases of IUD or TOP.

**TABLE 2 Type of obstruction in the prenatal and postnatal subset with isolated duodenal obstruction.**

	Prenatal subset N=11	Postnatal subset N=27	P
Type of obstruction overall (intrinsic/extrinsic)	4/7	22/5	0. 017
Intrinsic obstruction			
Duodenal membranous obstruction	1	9	0. 23
Duodenal stenosis	0	2	1. 00
Duodenal atresia	3	11	0. 49
Extrinsic obstruction			
Annular pancreas	5	5	0. 12
Annular pancreas with intestinal malrotation (Ladd's bands)	2	0	0. 08

**TABLE 3 Delivery and outcome data of the prenatal and postnatal subset with isolated duodenal obstruction**

	Prenatal subset N=11	Postnatal subset N=27	P
GA at delivery (weeks)	35.4 (30.1-39.3)	40.0 (31.1-41.6)	0.011
Delivery <37 weeks	8	8	0.04
Mode of delivery overall‡			0.24
Spontaneous delivery	6	19	0.23
Induced delivery	2	1	0.23
Cesarean section	3	4	0.65
Birth weight (g)	2320 (1455-3480)	2945 (1670-4350)	0.04
Birth weight <10 <sup>th</sup> centile	3	2	0.3
Age at diagnosis (days)	0 (0-2)	4 (0-39)	<0.001
Method of diagnosis			
X-ray of abdomen	11	19	0.08
Contrast swallow examination	0	7	0.08
Ultrasound of abdomen	0	4	0.6
>1 method required for a diagnosis	0	4	0.3
Surgery overall			0.028
Duodenoduodenostomy	11	15	0.008
Duodenojejunostomy	0	4	0.3
Membrane resection	0	8	0.08
>1 day artificial ventilation	2	6	1.00
Days to first oral feeds	5 (4-10)	6 (3-23)	0.56
Days on parenteral nutrition	11 (4-16)	11 (5-33)	0.85
Infants with complications after surgery	2	5	1.00
Number of complications after surgery	2	6	1.00
Infants requiring readmission	1	8	0.24
Number of readmissions	1	12	0.23
Length of hospital stay (days)	23 (15-48)	22 (11-68)	0.67

Data presented as median and range. GA, gestational age; g, gram

‡ The mode of delivery could not be retrieved for 3/27 cases in the postnatal subset

**TABLE 4 Associated anomalies in the prenatal and postnatal subset with non-isolated duodenal obstruction and separated in cases with and without trisomy 21.**

	Prenatal subset N=17	Postnatal subset N=36	P
<b>Cases associated with trisomy 21</b>	7	22	0.24
Cardiac	1	8	0.63
Gastro-intestinal tract	0	5	0.55
Esophageal atresia	0	1	1.00
Renal tract	0	0	-
Central nervous system	0	1	1.00
Vertebral	0	1	1.00
Other	0	0	-
Total number of anomalies	1	16	0.25
<b>Cases not associated with trisomy 21</b>	10	14	0.24
Other abnormal karyotype	0	1	1.00
Cardiac	4	8	0.68
Gastro-intestinal tract	2	5	0.65
Esophageal atresia	3	1	0.27
Renal tract	2	2	1.00
Central nervous system	1	1	1.00
Vertebral	4	3	0.39
Other	3	5	1.00
Total number of anomalies	19	25	0.56

**TABLE 5 Type of obstruction in the prenatal and postnatal subset with non-isolated duodenal obstruction.**

	Prenatal subset N=17*	Postnatal subset N=36	P
Type of obstruction overall (intrinsic/extrinsic)	13/3	25/11	0.43
<b>Intrinsic obstruction</b>			
Duodenal membranous obstruction	3	12	0.34
Duodenal stenosis	1	2	1.00
Duodenal atresia	9	11	0.12
<b>Extrinsic obstruction</b>			
Annular pancreas	3	8	1.00
Annular pancreas with intestinal malrotation (Ladd's bands)	0	3	0.54

\* In one case of intrauterine death associated with trisomy 21 post mortem investigation was not permitted which prevented a final diagnosis of duodenal obstruction

**TABLE 6 Delivery and outcome of the liveborn infants in the prenatal and postnatal subset with non-isolated duodenal obstruction.**

	Prenatal subset N=12	Postnatal subset N=36	P
GA at delivery (weeks)	35.7 (29.3-39.1)	38 (30.1-42)	0.10
Delivery <37 weeks gestation	9	14	0.046
Mode of delivery overall ‡			1.00
Spontaneous delivery	8	16	1.00
Induced delivery	1	3	1.00
Cesarean section	3	7	1.00
Birth weight (g)	1918 (1155-3390)	2770 (1130-3780)	0.04
Birth weight <10 <sup>th</sup> centile	6	14	0.74
Age at diagnosis (days)*	0 (0-2)	5 (1-814)	<0.001
Method of diagnosis*			
X-ray of abdomen	10	21	0.13
Contrast swallow examination	2	13	0.45
Ultrasound of abdomen	0	5	0.56
>1 method required for a diagnosis	1	4	1.00
Surgery overall*			0.31
Duodenoduodenostomy	7	18	0.73
Duodenojejunostomy	3	4	0.34
Duodenogastrostomy	0	1	1.00
Membrane resection	1	12	0.14
>1 day artificial ventilation*	5	3	0.03
Days to first oral feeds*	5 (4-12)	5 (2-22)	0.53
Days on parenteral nutrition*	12.5 (5-18)	9 (3-31)	0.20
Infants with complications after surgery*	7	13	0.17
Number of complications after surgery*	15	24	0.16
Infants requiring readmission*	1	4	0.71
Number of readmissions*	1	5	0.55
Length of hospital stay (days)*	26 (12-46)	24 (9-332)	0.47
Neonatal/infant death	3	5	0.39

Data presented as median and range. GA, gestational age; g, gram

‡ The mode of delivery could not be retrieved for 10/36 cases in the postnatal subset.

\* Prenatal subset: one case of neonatal death after 6 days, no surgery, excluded from analysis. Postnatal subset: one case of neonatal death two days after delivery, not included in analysis

**TABLE 7 Causes of duodenal obstruction associated with trisomy 21**

	Prenatal subset N=7 *	Postnatal subset N=22	P
Type of obstruction overall (intrinsic/extrinsic)	5/1	16/6	0.83
Intrinsic obstruction			
Duodenal membranous obstruction	2	9	1.00
Duodenal stenosis	1	1	0.39
Duodenal atresia	2	6	1.00
Extrinsic obstruction			
Annular pancreas	1	6	1.00

\* In one case of intrauterine death associated with trisomy 21 post mortem investigation was not permitted, precluding a final diagnosis of duodenal obstruction





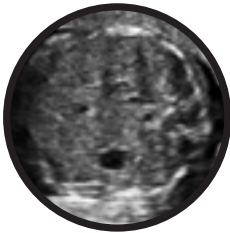
## REFERENCES

- 1 Garne E, Rasmussen L, Husby S. Gastrointestinal malformations in Funen county, Denmark-epidemiology, associated malformations, surgery and mortality. *Eur. J. Pediatr.* 2002; 12: 101-106.
- 2 Lawrence MJ, Ford WD, Furness ME, Hayward T, Wilson T. Congenital duodenal obstruction: early antenatal ultrasound diagnosis. *Pediatr. Surg. Int.* 2000; 16: 342-345.
- 3 Haeusler MC, Berghold A, Stoll C, Barisic I, Clementi M. Prenatal ultrasonographic detection of gastrointestinal obstruction: results from 18 European congenital anomaly registries. *Prenat. Diagn.* 2002; 22: 616-623.
- 4 Hancock BJ, Wiseman NE. Congenital duodenal obstruction: the impact of an antenatal diagnosis. *J. Pediatr. Surg.* 1989; 24: 1027-1031.
- 5 Shawis R, Antao B. Prenatal bowel dilatation and the subsequent postnatal management. *Early Hum. Dev.* 2006; 82: 297-303.
- 6 Tandler J. Zur Entwicklungsgeschichte des menschlichen Duodenums. *Morphol Jahrb* 1902; 29: 187-216.
- 7 Brantberg A, Blaas HG, Salvesen KA, Haugen SE, Mollerlokken G, Eik-Nes SH. Fetal duodenal obstructions: increased risk of prenatal sudden death. *Ultrasound Obstet. Gynecol.* 2002; 20: 439-446.
- 8 Grand RJ, Watkins JB, Torti FM. Development of the human gastrointestinal tract. A review. *Gastroenterology* 1976; 70: 790-810.
- 9 Romero R, Ghidini A, Costigan K, Touloukian R, Hobbins JC. Prenatal diagnosis of duodenal atresia: does it make any difference? *Obstet. Gynecol.* 1988; 71: 739-741.
- 10 Miro J, Bard H. Congenital atresia and stenosis of the duodenum: the impact of a prenatal diagnosis. *Am. J. Obstet. Gynecol.* 1988; 158: 555-559.
- 11 Bittencourt DG, Barini R, Marba S, Sbragia L. Congenital duodenal obstruction: does prenatal diagnosis improve the outcome? *Pediatr. Surg. Int.* 2004; 20: 582-585.
- 12 Dell'Agnola CA, Tomaselli V, Teruzzi E, Tadini B, Coran AG. Prenatal diagnosis of gastrointestinal obstruction: a correlation between prenatal ultrasonic findings and postnatal operative findings. *Prenat. Diagn.* 1993; 13: 629-632.
- 13 Grosfeld JL, Rescorla FJ. Duodenal atresia and stenosis: reassessment of treatment and outcome based on antenatal diagnosis, pathologic variance, and long-term follow-up. *World J. Surg.* 1993; 17: 301-309.
- 14 Singh MV, Richards C, Bowen JC. Does Down syndrome affect the outcome of congenital duodenal obstruction? *Pediatr. Surg. Int.* 2004; 20: 586-589.
- 15 Murshed R, Nicholls G, Spitz L. Intrinsic duodenal obstruction: trends in management and outcome over 45 years (1951-1995) with relevance to prenatal counselling. *Br. J. Obstet. Gynaecol.* 1999; 106: 1197-1199.
- 16 Nwosu EC, Welch CR, Manasse PR, Walkinshaw SA. Longitudinal assessment of amniotic fluid index. *Br. J. Obstet. Gynaecol.* 1993; 100: 816-819.
- 17 Niklasson A, Ericson A, Fryer JG, Karlberg J, Lawrence C, Karlberg P. An update of the Swedish reference standards for weight, length and head circumference at birth for given gestational age (1977-1981). *Acta Paediatr. Scand.* 1991; 80: 756-762.
- 18 Garne E, Loane M, Dolk H. Gastrointestinal malformations: impact of prenatal diagnosis on gestational age at birth. *Paediatr. Perinat. Epidemiol.* 2007; 21: 370-375.
- 19 Bartlett JM, Wypij D, Bellinger DC, Rappaport LA, Heffner LJ, Jonas RA, Newburger JW. Effect of prenatal diagnosis on outcomes in D-transposition of the great arteries. *Pediatrics* 2004; 113: e335-e340.
- 20 Hunfeld JA, Wladimiroff JW, Passchier J, Venema-Van Uden MU, Frets PG, Verhage F. Emotional reactions in women in late pregnancy (24 weeks or longer) following the ultrasound diagnosis of a severe or lethal fetal malformation. *Prenat. Diagn.* 1993; 13: 603-612.
- 21 Copper RL, Goldenberg RL, Das A, Elder N, Swain M, Norman G, Ramsey R, Cotroneo P, Collins BA, Johnson F, Jones P, Meier AM. The preterm prediction study: maternal stress is associated with spontaneous preterm birth at less than thirty-five weeks' gestation. National Institute of Child Health and Human Development Maternal-Fetal Medicine Units Network. *Am. J. Obstet. Gynecol.* 1996; 175: 1286-1292.
- 22 Orr ST, Reiter JP, Blazer DG, James SA. Maternal prenatal pregnancy-related anxiety and spontaneous preterm birth in Baltimore, Maryland. *Psychosom. Med.* 2007; 69: 566-570.
- 23 Wang ML, Dorer DJ, Fleming MP, Catlin EA. Clinical outcomes of near-term infants. *Pediatrics* 2004; 114: 372-376.
- 24 Kramer MS, Demissie K, Yang H, Platt RW, Sauve R, Liston R. The contribution of mild and moderate preterm birth to infant mortality. Fetal and Infant Health Study Group of the Canadian Perinatal Surveillance System. *JAMA* 2000; 284: 843-849.
- 25 Basu R, Burge DM. The effect of antenatal diagnosis on the management of small bowel atresia. *Pediatr. Surg. Int.* 2004; 20: 177-179.
- 26 la Vecchia LK, Grosfeld JL, West KW, Rescorla FJ, Scherer LR, Engum SA. Intestinal atresia and stenosis: a 25-year experience with 277 cases. *Arch. Surg.* 1998; 133: 490-496.
- 27 Mikaelsson C, Arnbjornsson E, Kullendorff CM. Membranous duodenal stenosis. *Acta Paediatr.* 1997; 86: 953-955.
- 28 Stanley P, Law BS, Young LW. Down syndrome, duodenal stenosis/annular pancreas, and a stack of coins. *Am. J. Dis. Child* 1988; 142: 459-460.

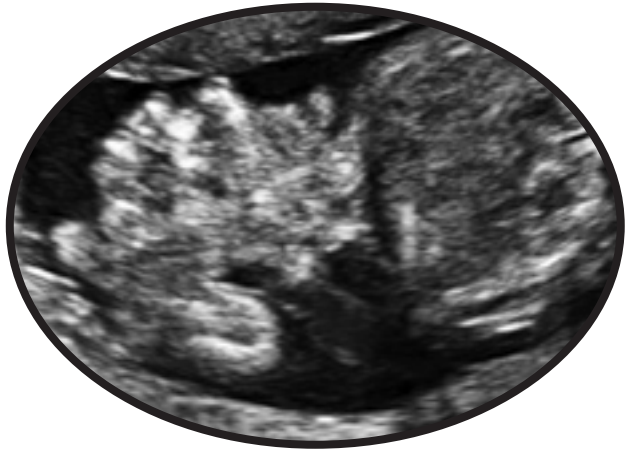
- 29 Chitty LS, Goodman J, Seller MJ, Maxwell D. Esophageal and duodenal atresia in a fetus with Down syndrome: prenatal sonographic features. *Ultrasound Obstet. Gynecol.* 1996; 7: 450-452.
- 30 Dundas KC, Walker J, Laing IA. Esophageal and duodenal atresia suspected at the 12 week booking scan. *BJOG.* 2001; 108: 225-226.
- 31 Estroff JA, Parad RB, Share JC, Benacerraf BR. Second trimester prenatal findings in duodenal and esophageal atresia without tracheoesophageal fistula. *J. Ultrasound Med.* 1994; 13: 375-379.
- 32 Heydanus R, Spaargaren MC, Wladimiroff JW. Prenatal ultrasonic diagnosis of obstructive bowel disease: a retrospective analysis. *Prenat. Diagn.* 1994; 14: 1035-1041.
- 33 Nicolaides KH, Snijders RJ, Cheng HH, Gosden C. Fetal gastro-intestinal and abdominal wall defects: associated malformations and chromosomal abnormalities. *Fetal Diagn. Ther.* 1992; 7: 102-115.
- 34 Bakalis S, Sairam S, Homfray T, Harrington K, Nicolaides K, Thilaganathan B. Outcome of antenatally diagnosed talipes equinovarus in an unselected obstetric population. *Ultrasound Obstet Gynecol* 2002; 20: 226-229.
- 35 Stoll C, Alembik Y, Dott B, Roth MP. Evaluation of prenatal diagnosis of congenital gastro-intestinal atresias. *Eur. J. Epidemiol.* 1996; 12: 611-616.
- 36 Goldstein I, Makhoul IR, Weissman A, Drugan A. Hemivertebra: prenatal diagnosis, incidence and characteristics. *Fetal Diagn. Ther.* 2005; 20: 121-126.



3



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2

- 1 Transverse plane of section through the normal fetal abdomen with the spine at 3 o'clock and stomach at 6 o'clock.
- 2 Cross sectional plane of the fetal abdomen showing free floating intestines in the amniotic fluid.
- 3 Newborn infant with gastroschisis; the intestines are herniated through an abdominal wall defect at the right of the umbilical cord insertion.

# 3.2

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The outcome of gastroschisis after a prenatal diagnosis or a diagnosis only at birth. Recommendations for prenatal surveillance.

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Titia E. Cohen-Overbeek (1), Titi R. Hatzmann (1), Eric A.P. Steegers (1),  
Wim C.J. Hop (2), Juriy W. Wladimiroff (1), Dick Tibboel (3)

1 Department of Obstetrics and Gynecology, Division of Obstetrics and Prenatal Medicine,  
Erasmus MC, Rotterdam, the Netherlands

2 Department of Epidemiology & Biostatistics, Erasmus MC, Rotterdam, the Netherlands

3 Department of Pediatric Surgery, Erasmus MC, Rotterdam, the Netherlands

# Abstract

**Objectives** To establish in infants with gastroschisis whether outcome is different when comparing a prenatal diagnosis with a diagnosis only at birth with the intention to develop a prenatal surveillance protocol. Intestinal atresia established after birth and preterm versus term delivery were studied as risk factors.

**Study design** All 24 fetuses and 9 infants diagnosed with gastroschisis and referred to our tertiary center between January 1991 and June 2003 were studied retrospectively.

**Results** The infants of the prenatal subset delivered at our tertiary center and 18 survived. There were two pregnancy terminations, three intrauterine deaths at 19,33 and 36 weeks respectively and one neonatal death. All nine infants in the postnatal subset survived. Eight were out born and one was delivered at our tertiary center. Prenatal bowel dilatation did not correlate with outcome. Between the prenatal and postnatal subset no significant difference in outcome of liveborn infants was established. For four infants with intestinal atresia a significant difference was demonstrated for induction of preterm labor ( $P<0.05$ ), duration of parenteral nutrition ( $P<0.01$ ), number of additional surgical procedures ( $P<0.001$ ) and length of hospital stay ( $P<0.01$ ). The fifteen infants born prior to 37 weeks of gestation spent a significantly longer period in hospital compared to those delivered at term. When the cases with bowel atresia were excluded this difference was no longer present. Five of the 33 cases were diagnosed with associated anomalies which mainly involved the urinary tract.

**Conclusion** Neonatal outcome of liveborn infants following a prenatal diagnosis of gastroschisis is not different from a diagnosis at birth. The presence of intestinal atresia is the most important prognostic factor for morbidity. The supplemental value of prenatal diagnosis to the outcome of infants with gastroschisis may be in the prevention of unnecessary intrauterine death and detection of intestinal complications. A proposed surveillance protocol for fetuses with gastroschisis focused on intrauterine signs of pending distress such as a dilated stomach, intra abdominal bowel dilatation with peristalsis, notches in the umbilical artery Doppler signal, development of polyhydramnios and an abnormal CTG registration may improve outcome.

# Introduction

Gastroschisis is a congenital defect of the abdominal wall characterized by the evisceration of abdominal organs without a covering membranous sac. The anomaly can easily be detected by prenatal ultrasound. The detection rate has been reported to be as high as 83%, with the majority being diagnosed before the third trimester<sup>1</sup>.

Infants with gastroschisis have a high overall survival rate, however, the rate of intrauterine fetal death and morbidity due to gastrointestinal complications is considerable<sup>2</sup>. Bowel dilatation<sup>3</sup> and polyhydramnios<sup>4</sup> have been suggested as prenatal predictors of adverse postnatal outcome, although for the size of the dilated bowel<sup>5</sup> this may be questioned. Preterm delivery and particularly pre-labor cesarean section may result in a reduction in the formation of a fibrous coating on the bowel as well as reduce complications such as atresia, stenosis, necrosis and perforations of the bowel<sup>6</sup>. However there is still controversy regarding the use of cesarean section and preterm instead of term delivery<sup>7</sup>. Several studies have come to the conclusion that term vaginal delivery will improve outcome<sup>8,9</sup>. The positive effect of prenatal diagnosis on the outcome of gastroschisis has not been established, although these studies were conducted a decade ago<sup>10-12</sup>. Routine ultrasound to detect congenital anomalies was not part of recent policy in the Netherlands therefore some cases of gastroschisis remained undiagnosed in the antenatal period. This provided us with the opportunity to assess the value of prenatal diagnosis for fetuses with gastroschisis and to propose a surveillance protocol according to outcome parameters and data from the literature. We studied (i) the influence of prenatal diagnosis on the outcome of fetuses with gastroschisis compared with a group of infants where this diagnosis was only established at birth and (ii) the influence of bowel atresia on outcome of infants with gastroschisis and (iii) the outcome of infants with gastroschisis born before or after 37 weeks of gestation.

## Material and Methods

The University Medical Center Rotterdam serves as the referral center for both fetal anomaly scanning and pediatric surgery in the South West of the Netherlands. We performed a retrospective analysis of all infants from singleton pregnancies diagnosed with a gastroschisis either pre- or postnatally between January 1991 and June 2003. Cases were identified from our ultrasound database and from the patient database of the Department of Pediatric Surgery.

Following prenatal detection of a gastroschisis, regular ultrasound scans were performed during pregnancy to evaluate fetal growth, the presence of bowel dilatation and amniotic fluid volume. During the study period no structured protocol was applied to monitor the fetal condition. Fetal growth

restriction (FGR) was defined as a fetal upper abdominal circumference <10<sup>th</sup> centile. Bowel dilatation was determined by the inner to inner bowel wall distance. Maximum bowel dilatation prior to delivery was stratified employing a cut-off level of 10 and 17 mm and correlated with outcome. Amniotic fluid volume was considered abnormal for measurements below the 5<sup>th</sup> and above 95<sup>th</sup> percentile<sup>13</sup>. Counselling by a pediatric surgeon was offered to all couples during the prenatal period. According to the hospital protocol all women with a prenatal diagnosis of gastroschisis were delivered in our center by vaginal delivery unless obstetric reasons required otherwise. Prematurity was defined as a delivery prior to 37 weeks of gestation. Infants with a birth weight below 10<sup>th</sup> percentile, adjusted for sex<sup>14</sup>, were defined as small for gestational age (SGA).

After delivery the presence or absence of a fibrous coat on the bowel (bowel peel) was documented. Management of the gastroschisis consisted of closure of the defect within hours following birth, by means of primary closure or Silastic Silo depending on the size of the defect and ventilatory pressures during the procedure. Due to motility disorders of the bowel infants were given primarily parenteral nutrition (TPN) and enteral feeding was instituted as soon as stomach retentions diminished.

Adverse neonatal outcome was defined as neonatal death or complications resulting from the abnormality itself (intestinal atresia, necrotising enterocolitis), the subsequent surgical procedure (anatomic or functional short bowel syndrome, TPN associated cholestasis jaundice, venous line sepsis), or other significant morbidity not directly related to the gastroschisis.

Distinction was made between: (i) a diagnosis prenatally and a diagnosis only postnatally, (ii) presence and absence of intestinal atresia and (iii) delivery before and after 37 weeks of gestation. Charts were reviewed for maternal demographic data, fetal ultrasound surveillance data, fetal outcome, neonatal surgical and subsequent outcome data.

Statistical analysis of comparing groups was performed using the Mann-Whitney test or Fisher's exact test in case of continuous or categorical data, respectively.  $P=0.05$  (two-sided) was considered as the limit of statistical significance.

## Results

We reviewed the data of 24 prenatally diagnosed and 9 postnatally diagnosed cases of gastroschisis. Three prenatal cases have been described previously<sup>15</sup>. In the prenatal subset, mean maternal age ( $\pm$  SD) at diagnosis was  $25.9 \pm 6$  years. Mean gestational age at diagnosis was  $23.3 \pm 6.6$  weeks. In six out of 24 women (25%) the gastroschisis was detected after 24 weeks gestation. Two pregnancies were terminated before 24 weeks gestation. Three women were diagnosed with FGR. Five pregnancies were complicated by oligohydramnios and two by polyhydramnios. Associated fetal anomalies were detected in



4/22 (17%) pregnancies, unilateral hydronephrosis in three cases and bilateral talipes equinovarus in one. Three fetuses (12.5%) died in utero at 19, 33 and 36 weeks of gestation, respectively. One intrauterine death represented a combination of gastroschisis, bilateral talipes equinovarus and early FGR. The two intrauterine deaths in the third trimester revealed normal amniotic fluid volume, no altered aspect of the stomach and intestines and no evidence of FGR or abnormal Doppler measurements within two weeks of fetal demise. Apart from the gastroschisis neither intestinal atresia nor other anomalies were revealed in both cases at post mortem investigation.

Delivery data and outcome variables of the liveborn prenatal subset (N=19) and postnatal subset (N=9) are provided in Tables 1 and 2. The postnatal subset consisted of one infant born at home, seven infants delivered in a regional general hospital and one infant delivered in our tertiary care facility at a gestational age of 36 week. No statistically significant difference was found between the two groups. In the liveborn prenatal subset labor was induced before 37 weeks in 3/7 cases: at 32 week due to ruptured membranes and developing fever; at 36 weeks because of FGR and dilated bowel loops; and at 36 weeks due to FGR, oligohydramnios and dilated bowel loops. In the latter two infants, atresia of the bowel was suspected prenatally, but only confirmed in one. One cesarian section was performed because of fetal distress noticed during spontaneous labor at 36 weeks of gestation. Postnatal evaluation of the prenatal subset revealed seven infants presenting solely with bowel in the evisceration, eight infants with evisceration of the bowel and stomach, one with evisceration of the bowel and ovaries and three with evisceration of bowel, stomach and ovaries (50% of females). In the subset in which gastroschisis was first diagnosed after delivery (postnatal subset), two infants presented with bowel evisceration, six with bowel and stomach evisceration, and in one infant the evisceration included the urinary bladder.

All infants were transferred to the pediatric surgical ICU on the day of delivery. In the prenatal subset all three cases of suspected unilateral hydronephrosis were confirmed. They were the result of a pelvic ureteric junction obstruction, a primary mega ureter and a vesicoureteric reflux, respectively. In addition, one other case of vesico-urethral reflux was diagnosed. Complications developing in the postnatal period included a frontal temporal hypoxia causing a convulsion and a venous infarct in the corticospinal tract resulting in mild hemiplegia in one patient. In the postnatal subset no other anomalies or complications other than those related to the gastroschisis were diagnosed.

Intestinal atresia was suspected prenatally in 3/19 liveborn infants, twice due to a herniated stomach together with dilated bowel loops of 11 and 33 mm, respectively. The latter case was also complicated by polyhydramnios and intra abdominal dilated bowel loops with active peristalsis; the third case revealed dilated echogenic bowel loops with a diameter of 23 mm. Bowel atresia was confirmed at birth in the first two cases. The first infant

was delivered at 36 weeks and its bowel was covered by peel. Closure of the defect required a Siliastic Silo. Definitive closure of the defect was undertaken on day 14. At this time the presence of bowel peel was not documented. Additional surgery necessary for persistent stomach retentions was performed at the age of 3 months and a jejunal atresia was corrected. The second infant required almost total bowel resection due to complete bowel necroses established at birth, resulting in withdrawal of treatment and neonatal death one day after delivery. This fetus presented at 35 weeks gestation with a polyhydramnios, dilated intra-abdominal bowel loops. He was born at 37 weeks gestation with a birth weight of 2745 grams. One infant was delivered after premature rupture of membranes and onset of labor at 31 weeks. There was no ultrasound evidence of polyhydramnios or dilated bowel loops at 29 weeks of gestation, however atresia was established.

In none of the nine infants diagnosed with gastroschisis only at birth, an atresia was noted at initial surgery. In one infant re-evaluation due to persistent stomach retention showed a jejunal atresia at the age of 8 weeks, leading to two additional surgical procedures.

Extra-abdominal bowel dilatation was seen in 14/21 fetuses (67%) of the prenatal subset which reached the third trimester. The mean bowel dilatation measured was  $18.3 \pm 6$  mm with a minimum of 11 and a maximum of 33 mm. Based on reported cut-off levels of 10 and 17 mm the number of fetuses with a bowel diameter of <10 mm was seven, between 10 and 17 mm six and  $\geq 17$  mm eight. One case of atresia was diagnosed postnatally in each of these subgroups.

To evaluate the effect of atresia on outcome, the 4 infants with atresia were compared with the 24 infants without this complication. A significant difference was established for induction of labor prior to 37 weeks, 2/4 compared to 1/24 ( $P < 0.05$ ). Excluding the neonatal death, the median total duration of parenteral nutrition (range) 148 days (148-149) versus 32 days (19-96) ( $P < 0.01$ ), the number of additional surgical procedures 3 versus 0 ( $P < 0.001$ ) and the median length of hospital stay (range) 145 days (68-149) versus 33 days (12-100) ( $P < 0.01$ ) demonstrated a significant difference.

Liveborn delivery occurred in 15 of 28 infants prior to 37 weeks. Bowel peel was documented in 7/15 cases of the premature infants and 3/13 infants of the subset born after 37 weeks of gestation and did not show a significant difference between these two subsets. Intestinal atresia in combination with bowel peel was present in only one of the premature infants. There was a significant difference for mean birth weight ( $\pm$  SD) 2350 grams ( $\pm$  310) versus 2717 grams ( $\pm$  305) ( $P < 0.01$ ) and median length of hospital stay (range) 58 days (28-149) versus 31 days (12-74) ( $P < 0.05$ ) in infants born prior to 37 weeks of gestation compared to infants born after this period. The prolonged duration of artificial ventilation for premature infants just did not reach statistical significance ( $P = 0.053$ ). When the cases with bowel atresia were excluded the significant difference for length of hospital stay (range) 44 days (28-100) versus 31 days (12-74) was no longer present.

## Comment

The majority of infants born with gastroschisis are detected before birth. This provides an opportunity to influence management with the aim to improve outcome. In order to diminish insults to the exposed intestines, elective preterm delivery<sup>6</sup> or pre-labor cesarean section has been recommended<sup>16</sup>. In a meta-analysis of studies investigating delivery modes in fetuses with gastroschisis, no advantage of cesarean section over vaginal delivery could be established<sup>17</sup>. Furthermore, no advantage could be demonstrated for preterm delivery in a first randomized controlled trial studying elective preterm delivery versus spontaneous delivery<sup>18</sup>. In this study the median duration of hospital stay in the elective preterm delivery group versus the spontaneous group was not significantly different (47.5 versus 53 days). These findings remained unaltered if data from infants with intestinal atresia were excluded.

Our study confirms previous data<sup>10-12</sup> where a significant difference in outcome could not be demonstrated between liveborn infants following a prenatal diagnosis or only a diagnosis at birth. The Netherlands is a small country with easily available tertiary level resources and out born infants with severe anomalies can be transported within 15-30 minutes to one of these facilities. This may have biased our results compared to remote areas where infants are born distant from tertiary care resources. A recent study from Australia<sup>19</sup>, however, analyzing data from 181 liveborn infants with gastroschisis revealed that place and mode of delivery, distance from tertiary centers, time to surgery and type of closure did not influence neonatal outcome.

In our preterm delivery group a significantly longer length of hospital stay was demonstrated compared with infants born at term. When cases with bowel atresia were excluded, this difference was no longer present but tended towards significance ( $P=0.10$ ). Calculation of the 95% confidence interval for the difference (subset 1 minus subset 2) of mean values resulted in a wide interval ranging from -0.6 up to 32.6 days. The power to demonstrate a difference was apparently too low with the studied number of patients and a larger study would be valuable to address this question. However, similar findings were reported by Huang et al.<sup>20</sup> who studied 57 infants with gastroschisis. In their study all cases of atresia were delivered before 37 weeks. Prematurity together with atresia may contribute to the unfavorable outcome.

Infants with gastroschisis can be divided in two categories; those with a simple defect comprising approximately 80% of cases and those where the gastroschisis is complicated by bowel pathology such as atresia, stenosis, perforation or volvulus<sup>21</sup>. Morbidity in the latter group is considerable and expressed by a significantly prolonged hospital stay and longer periods of mechanical ventilation and enteral feeding<sup>22</sup>. Although our study consists of small numbers the unfavorable outcome for infants with atresia could

be demonstrated. Prenatal selection of these cases proved difficult and measurement of extra abdominal bowel dilatation was not useful. One of the two cases of polyhydramnios was diagnosed with atresia, in agreement with data from Japaraj et al.<sup>4</sup> who found a clear correlation between polyhydramnios and severe bowel complications. Intra abdominal bowel dilatation together with active peristalsis was in our study associated with bowel atresia and necrosis. This combination has only been previously described in 19 cases<sup>16,23-26</sup> all with confirmed atresia.

Amniotic fluid of fetuses with gastroschis contains an inflammatory exudate and this may have a deleterious effect on the exposed bowel<sup>27</sup>.

It has been suggested that amnioexchange using warm saline<sup>28</sup> or amnioinfusion in case of oligohydramnios<sup>29</sup> may prevent or diminish intestinal compromise. On the other hand a higher incidence of premature delivery has been reported as a result of this intervention<sup>30</sup>. In the present study 7/15 premature infants presented with bowel peel compared to 3/13 infants delivered after 37 weeks of gestation. Only one of the 7 premature infants revealed bowel peel together with intestinal atresia. This is in agreement with a recent large neonatal study<sup>8</sup> containing 75 liveborn infants with gastroschisis where infants born before or after 37 weeks showed evidence of bowel peel in 50% and 57%, respectively. Only 5 cases in this study suffered from an intestinal atresia, whether or not associated with bowel peel was not documented. Until a prospective randomized trial with sufficient numbers can demonstrate an improved outcome after prenatal intervention, this treatment should not generally be instituted in the management of fetuses with gastroschisis.

Intrauterine death in our study occurred in 3/22 (13.6%) prenatal cases. This is comparable to previous investigations<sup>1,2</sup>, before the commencement of intense fetal monitoring, where figures of 10.6% and 12%, respectively, are quoted. The two cases with intrauterine death in our study did not reveal an explanation for fetal demise at post mortem investigation which was also previously described in the literature<sup>2</sup>. The contribution of prenatal diagnosis to the management of gastroschisis may be in the prevention of intrauterine deaths and adverse neurological outcome<sup>2</sup>. Intrauterine deaths usually occur in the third trimester as was the case in the present study despite normal Doppler studies of the umbilical artery and no evidence of FGR within 2 weeks of fetal demise. Regular fetal cardiotocography (CTG) in third trimester management has been proposed in order to avert these intrauterine deaths. In one study intense third trimester CTG monitoring was not commenced unless there was evidence of FGR or abnormal umbilical artery Doppler studies and no intrauterine death occurred in 45 fetuses<sup>4</sup>. Others suggest weekly CTG from 34 weeks<sup>31</sup> onwards or daily or every second day CTG starting at 34 weeks<sup>16</sup>. Salomon et al.<sup>29</sup> performed daily CTG examinations from 27 until 35 weeks, but despite intensive monitoring both latter studies<sup>16,29</sup> could not prevent the occurrence of an intrauterine death. In the first study the intrauterine death occurred at 35 weeks but

only a CTG 2 weeks prior to this event was registered. Pathologic CTG's are documented in a high percentage of fetuses with gastroschisis who after delivery do not demonstrate evidence of hypoxia<sup>16</sup>. It is suggested that fetal heart rate variability may be affected by alterations in vagal output caused by stomach dilatation<sup>32</sup> or by the mechanical effect of gut herniation<sup>31</sup>. Other signs of fetal compromise such as a notch in the umbilical artery Doppler waveform<sup>23</sup>, possible caused by compression of the cord due to dilated bowel, and the development of polyhydramnios<sup>4</sup> may be equally important to detect those fetuses at risk of fetal demise or a complicated neonatal road to recovery. A new surveillance protocol summarizing the most recent information from the literature integrated with our own data is presented in Box 1. We speculate that this protocol may improve the detection of fetus at risk and in the future an analysis of outcome data after the commencement of this regimen should be performed.

In the present study one intrauterine death occurred at a gestational age of nineteen weeks and besides gastroschisis, bilateral talipes equinovarus had been diagnosed. In five out of our 33 cases (15%), associated extra gastrointestinal anomalies were diagnosed, the majority of which concerned the urinary tract. Two previous studies<sup>33,34</sup> demonstrated similar numbers of associated urinary tract anomalies. Some cases of urinary tract obstruction may be the result of evisceration of the bladder<sup>35,36</sup> which over time returns to normal after correction of the abdominal wall defect. In our study only one case of bladder evisceration was documented and this was not associated with urinary tract obstruction. Surgical correction of the urinary tract anomalies was not required but antibiotic prophylaxis was commenced to prevent infection. The relative high frequency with which urinary tract obstructions are encountered would suggest that in all infants with gastroschisis ultrasound investigation of the urinary tract should be recommended after the first week of life.

It can be concluded, that for liveborn infants there is no difference in perinatal outcome between a prenatal diagnosis of gastroschisis and a diagnosis only at birth. Preterm delivery results in prolonged hospital stay which is mainly associated with bowel atresia. In the presence of gastroschisis, infants with bowel atresia have a prolonged total duration of parenteral nutrition and hospital stay, and require more additional surgical procedures, compared with infants without additional gastrointestinal complications. The added value of prenatal diagnosis to the outcome of infants with gastroschisis may be in the prevention of unnecessary intrauterine death and detection of intestinal complications. Protocols to monitor fetuses with gastroschisis focused on intrauterine signs of pending distress such as a dilated stomach, intra abdominal bowel dilatation with peristalsis, notches in the umbilical artery Doppler signal, development of polyhydramnios and an abnormal CTG registration may improve outcome.

**TABLE 1 Delivery data of the prenatal liveborn subset and postnatal subset with gastroschisis**

	Prenatal subset N=19	Postnatal subset N=9	P
Male/female ratio	11/8	4/5	NS
GA at delivery (weeks)*	36.2 ± 1.7	37 ± 1.3	NS
Delivery <37 weeks (%)	11 (58%)	4 (44%)	NS
Induced delivery <37 weeks (%)	3 (16%)	0 (0%)	NS
Vaginal delivery (%)	17 (90%)	9 (100%)	NS
Mean birth weight (g)*	2461 ± 408	2582 ± 302	NS
Birth weight <P10 (%)	4 (21%)	4 (44%)	NS
Apgar score at 1 min#	8 (1-10)	7 (6-9)	NS
Apgar score at 5 min#	9 (3-10)	9 (8-10)	NS

\* Data presented as mean ± SD. # Data presented as median (range). GA, gestational age; g, gram; NS, not significant.

**TABLE 2 Outcome data of the prenatal liveborn subset and postnatal subset with gastroschisis**

	Prenatal subset N=19	Postnatal subset N=9	P
Bowel peel	8 (42%)	2 (22%)	NS
Primary closure (%)	13 (68%)	9 (100%)	NS
Silo closure	6 (32%)	0 (0%)	NS
Atresia	3 (16%)	1 (11%)	NS
Days to first oral feeds# §	13.5 (6-126)	18 (7-68)	NS
Days on parenteral nutrition# §	32 (21-149)	45 (19-148)	NS
Days on antibiotics# §	6 (1-34)	8 (2-15)	NS
Days on ventilation# §	5 (1-36)	4 (1-17)	NS
Additional surgery §	2 (11%)	1 (11%)	NS
Short bowel syndrome §	1 (5%)	0 (0%)	NS
Sepsis §	8 (44%)	2 (22%)	NS
Cholestatic jaundice §	1 (5%)	1 (11%)	NS
Necrotizing enterocolitis §	0	1 (11%)	NS
Length of hospital stay (d)# §	33 (12-149)	40 (22-145)	NS
Neonatal death	1 (5%)	0 (0%)	NS
Other complications*§	2 (11%)	0 (0%)	NS
Associated anomalies	4 (21%)	0 (0%)	NS

# Data presented as median (range). § excluded data of infant with neonatal death at day 1.

\*\* Includes a frontal temporal hypoxia and a venous infarct in the corticospinal tract. d, days; ns, not significant.

**BOX 1     Prenatal diagnosis and surveillance of a fetus with gastroschisis.**

<b>1</b>	<b>2nd trimester diagnosis of gastroschisis</b>
	Ultrasound assessment of
	a. Biometry
	b. Presence of associated anomalies
	c. Intra-abdominal bowel dilatation and peristalsis
	d. Amniotic fluid volume
	Arrange counselling by a pediatric surgeon
<b>2</b>	<b>From 30 weeks gestational age</b>
	Once a week ultrasound assessment of
	a. Biometry
	b. Position and size of stomach
	c. Intra abdominal bowel dilatation and peristalsis
	d. Extra abdominal bowel dilatation
	e. Position and size of bladder
	f. Amniotic fluid volume
	g. Doppler of the umbilical artery at the abdominal insertion
	CTG monitoring twice weekly
<b>3</b>	<b>Arrange for vaginal delivery from 37 weeks onwards</b>
	In case of detection of alterations or signs of fetal distress during surveillance consult with fetal maternal medicine specialist, neonatologist and pediatric surgeon for possible adaptation of management

## REFERENCES

- 1 Barisic I, Clementi M, Hausler M, Gjergja R, Kern J, Stoll C. Evaluation of prenatal ultrasound diagnosis of fetal abdominal wall defects by 19 European registries. *Ultrasound Obstet. Gynecol.* 2001; 18: 309-316.
- 2 Burge DM, Ade-Ajayi N. Adverse outcome after prenatal diagnosis of gastroschisis: the role of fetal monitoring. *J. Pediatr. Surg.* 1997; 32: 441-444.
- 3 Pryde PG, Bardicef M, Treadwell MC, Klein M, Isada NB, Evans MI. Gastroschisis: can antenatal ultrasound predict infant outcomes? *Obstet. Gynecol.* 1994; 84: 505-510.
- 4 Japaraj RP, Hockey R, Chan FY. Gastroschisis: can prenatal sonography predict neonatal outcome? *Ultrasound Obstet. Gynecol.* 2003; 21: 329-333.
- 5 Alsulyman OM, Monteiro H, Ouzounian JG, Barton L, Songster GS, Kovacs BW. Clinical significance of prenatal ultrasonographic intestinal dilatation in fetuses with gastroschisis. *Am. J. Obstet. Gynecol.* 1996; 175: 982-984.
- 6 Moir CR, Ramsey PS, Ogburn PL, Johnson RV, Ramin KD. A prospective trial of elective preterm delivery for fetal gastroschisis. *Am. J. Perinatol.* 2004; 21: 289-294.
- 7 Salvesen KA. Fetal abdominal wall defects--easy to diagnose--and then what? *Ultrasound Obstet. Gynecol.* 2001; 18: 301-304.
- 8 Ergun O, Barksdale E, Ergun FS, Prosen T, Qureshi FG, Reblook KR, Ford H, Hackam DJ. The timing of delivery of infants with gastroschisis influences outcome. *J. Pediatr. Surg.* 2005; 40: 424-428.
- 9 Puligandla PS, Janvier A, Flageole H, Bouchard S, Laberge JM. Routine cesarean delivery does not improve the outcome of infants with gastroschisis. *J. Pediatr. Surg.* 2004; 39: 742-745.
- 10 Haddock G, Davis CF, Raine PA. Gastroschisis in the decade of prenatal diagnosis: 1983-1993. *Eur. J. Pediatr. Surg.* 1996; 6: 18-22.
- 11 Rinehart BK, Terrone DA, Isler CM, Larmon JE, Perry KG, Jr., Roberts WE. Modern obstetric management and outcome of infants with gastroschisis. *Obstet. Gynecol.* 1999; 94: 112-116.
- 12 Adra AM, Landy HJ, Nahmias J, Gomez-Marin O. The fetus with gastroschisis: impact of route of delivery and prenatal ultrasonography. *Am. J. Obstet. Gynecol.* 1996; 174: 540-546.
- 13 Nwosu EC, Welch CR, Manasse PR, Walkinshaw SA. Longitudinal assessment of amniotic fluid index. *Br. J. Obstet. Gynaecol.* 1993; 100: 816-819.
- 14 Niklasson A, Ericson A, Fryer JG, Karlberg J, Lawrence C, Karlberg P. An update of the Swedish reference standards for weight, length and head circumference at birth for given gestational age (1977-1981). *Acta Paediatr. Scand.* 1991; 80: 756-762.
- 15 Heydanus R, Raats MA, Tibboel D, Los FJ, Wladimiroff JW. Prenatal diagnosis of fetal abdominal wall defects: a retrospective analysis of 44 cases. *Prenat. Diagn.* 1996; 16: 411-417.
- 16 Brantberg A, Blaas HG, Salvesen KA, Haugen SE, Eik-Nes SH. Surveillance and outcome of fetuses with gastroschisis. *Ultrasound Obstet. Gynecol.* 2004; 23: 4-13.
- 17 Segel SY, Marder SJ, Parry S, Macones GA. Fetal abdominal wall defects and mode of delivery: a systematic review. *Obstet. Gynecol.* 2001; 98: 867-873.
- 18 Logghe HL, Mason GC, Thornton JG, Stringer MD. A randomized controlled trial of elective preterm delivery of fetuses with gastroschisis. *J. Pediatr. Surg.* 2005; 40: 1726-1731.
- 19 Singh SJ, Fraser A, Leditschke JF, Spence K, Kimble R, Dalby-Payne J, Baskaranathan S, Barr P, Halliday R, Badawi N, Peat JK, Glasson M, Cass D. Gastroschisis: determinants of neonatal outcome. *Pediatr. Surg. Int.* 2003; 19: 260-265.
- 20 Huang J, Kurkchubasche AG, Carr SR, Wesselhoeft CW, Jr., Tracy TF, Jr., Luks FL. Benefits of term delivery in infants with antenatally diagnosed gastroschisis. *Obstet. Gynecol.* 2002; 100: 695-699.
- 21 Snyder CL, Miller KA, Sharp RJ, Murphy JP, Andrews WA, Holcomb GW, III, Gittes GK, Ashcraft KW. Management of intestinal atresia in patients with gastroschisis. *J. Pediatr. Surg.* 2001; 36: 1542-1545.
- 22 Molik KA, Gingalewski CA, West KW, Rescorla FJ, Scherer LR, Engum SA, Grosfeld JL. Gastroschisis: a plea for risk categorization. *J. Pediatr. Surg.* 2001; 36: 51-55.
- 23 Kalache KD, Bieriich A, Hammer H, Bollmann R. Is unexplained third trimester intrauterine death of fetuses with gastroschisis caused by umbilical cord compression due to acute extra-abdominal bowel dilatation? *Prenat. Diagn.* 2002; 22: 715-717.
- 24 McMahon MJ, Kuller JA, Chescheir NC. Prenatal ultrasonographic findings associated with short bowel syndrome in two fetuses with gastroschisis. *Obstet. Gynecol.* 1996; 88: 676-678.
- 25 Brun M, Grignon A, Guibaud L, Garel L, Saint-Vit D. Gastroschisis: are prenatal ultrasonographic findings useful for assessing the prognosis? *Pediatr. Radiol.* 1996; 26: 723-726.
- 26 Nick AM, Bruner JP, Moses R, Yang EY, Scott TA. Second-trimester intra-abdominal bowel dilation in fetuses with gastroschisis predicts neonatal bowel atresia. *Ultrasound Obstet. Gynecol.* 2006; 28: 821-825.
- 27 Morrison JJ, Klein N, Chitty LS, Kocjan G, Walshe D, Goulding M, Geary MP, Pierro A, Rodeck CH. Intra-amniotic inflammation in human gastroschisis: possible aetiology of postnatal bowel dysfunction. *Br. J. Obstet. Gynaecol.* 1998; 105: 1200-1204.
- 28 Luton D, De Lagausie P, Guibourdenche J, Oury J, Sibony O, Vuillard E, Boissinot C, Aigrain Y, Beaufils F, Navarro J, Blot P. Effect of amnioinfusion on the outcome of



- prenatally diagnosed gastroschisis. *Fetal Diagn. Ther.* 1999; 14: 152-155.
- 29 Salomon LJ, Mahieu-Caputo D, Jouvett P, Jouannic JM, Benachi A, Grebille AG, Dumez Y, Dommergues M. Fetal home monitoring for the prenatal management of gastroschisis. *Acta Obstet. Gynecol. Scand.* 2004; 83: 1061-1064.
  - 30 Mahieu-Caputo D, Muller F, Jouvett P, Thalabard JC, Jouannic JM, Nihoul-Fekete C, Dumez Y, Dommergues M. Amniotic fluid beta-endorphin: a prognostic marker for gastroschisis? *J. Pediatr. Surg.* 2002; 37: 1602-1606.
  - 31 Ingamells S, Saunders NJ, Burge D. Gastroschisis and reduced fetal heart-rate variability. *Lancet* 1995; 345: 1024-1025.
  - 32 Aina-Mumuney AJ, Fischer AC, Blakemore KJ, Crino JP, Costigan K, Swenson K, Chisholm CA. A dilated fetal stomach predicts a complicated postnatal course in cases of prenatally diagnosed gastroschisis. *Am. J. Obstet. Gynecol.* 2004; 190: 1326-1330.
  - 33 Gibbin C, Touch S, Broth RE, Berghella V. Abdominal wall defects and congenital heart disease. *Ultrasound Obstet. Gynecol.* 2003; 21: 334-337.
  - 34 Puligandla PS, Janvier A, Flageole H, Bouchard S, Mok E, Laberge JM. The significance of intrauterine growth restriction is different from prematurity for the outcome of infants with gastroschisis. *J. Pediatr. Surg.* 2004; 39: 1200-1204.
  - 35 Reiss RE, Landon MB, Jayanthi VR, Caniano DA, Mutabagani K, O'Shaughnessy RW. Functional urinary tract obstruction developing in fetuses with isolated gastroschisis. *Ultrasound Obstet. Gynecol.* 2000; 15: 194-198.
  - 36 Ikkena SE, de Chazal RC, Konje JC. Gastroschisis associated with bladder evisceration complicated by hydronephrosis presenting antenatally. *Ultrasound Obstet. Gynecol.* 1999; 13: 370-372.

Due to an embargo

*Chapter 3.3. Omphalocele; comparison of the perinatal outcome following a prenatal diagnosis or a diagnosis at birth*

is temporarily removed.

# 4

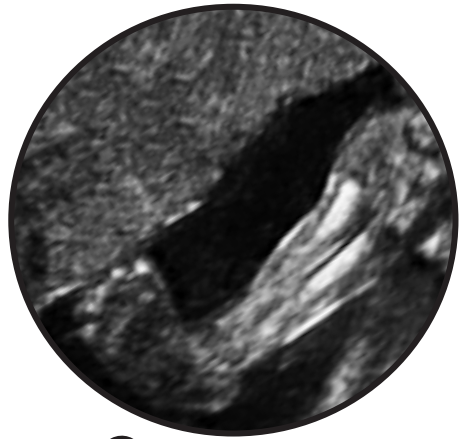
## Diagnosis of Talipes equinovarus

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The diagnosis of fetal talipes equinovarus is based on the tibia, fibula and sole of the foot being presented in the same plane on an ultrasound scan. Approximately half the prenatally detected cases, are associated with other anomalies or an abnormal karyotype, contrary to postnatal detected cases where only a minority is associated with these abnormalities. The prenatal diagnosis of talipes equinovarus appears to be associated with easier and less complicated postnatal surgical treatment providing the anomaly is isolated and the treatment is arranged at a Pediatric Orthopedic Center



1



2



3

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- 1 Normal position of fetal knee, lower leg and foot.
  - 2 Lower leg of a 20 week fetus with talipes equinovarus. The tibia and the fibula are visible in the same plane as the sole of the foot.
  - 3 3-months old infant with bilateral talipes equinovarus. The right foot shows the abnormal position whereas the left foot is in a plaster cast.

# 4.1

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## Congenital talipes equinovarus; comparison of outcome between a prenatal diagnosis and a diagnosis after delivery

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T. E. Cohen-Overbeek (1), E.W.M. Grijseels (1), E.A.G. Lammerink(1),  
W.C.J. Hop (2), J.W. Wladimiroff (1), A.F.M. Diepstraten (3)

1. Department of Obstetrics and Gynecology, Erasmus MC, Rotterdam, the Netherlands
2. Department of Epidemiology & Biostatistics, Erasmus MC, Rotterdam, the Netherlands
3. Department of Pediatric Orthopedics, Erasmus MC, Rotterdam, the Netherlands

# Abstract

**Objectives** To establish the impact on outcome of prenatally versus postnatally detected talipes equinovarus (TEV).

**Methods** The prenatal group was represented by pregnancies with sonographically detected TEV of which 18 were isolated and 39 were complex. The postnatal group contained 64 infants with an isolated and 10 infants with a complex TEV detected at birth. Treatment consisted of redressement followed by surgical postero-lateral or postero-medial release in the University Pediatric Orthopedic Center. The postnatal isolated TEV group underwent redressement treatment in the University Center (subset A, N=39) or in a regional general hospital (subset B, N=25).

**Results** For isolated TEV statistically significant difference existed for the surgical procedure ( $p < 0.001$ ), age at surgery ( $p < 0.01$ ) and admission time ( $p < 0.001$ ) between the prenatal and postnatal subset B and between the postnatal subsets A and B. For the complex TEV no significant difference was found for these variables between the six surviving infants of the prenatal group and the postnatal group.

**Conclusion** Prenatal detection of isolated TEV results in earlier and less complicated postnatal surgery and a shorter admission time providing treatment is arranged in a Pediatric Orthopedic Center. After prenatal detection of a complex TEV survival is low and determined by associated anomalies.

# Introduction

In the last 2 decennia, reports have appeared on the prenatal detection of talipes equinovarus (TEV). The first publications concentrated on the occurrence of TEV with other anomalies (complex cases) and the relative high frequency of an associated abnormal karyotype<sup>1-3</sup>. The prenatal detection rate demonstrated an increase from 11% in 1991 to 64% in 1999<sup>4</sup>. Apart from complex cases, a substantial number of isolated cases were revealed. During pregnancy or after delivery it may become evident, however, that a proportion of the isolated cases was in fact complex<sup>5-7</sup>. Furthermore, the false positive rate for isolated TEV varies between 6.4 and 35.3%<sup>8,9</sup>. In the presence of associated anomalies the prognosis is poor with only 10% normal outcome in contrast to a 94% normal outcome for isolated cases<sup>5</sup>. The need for surgical treatment in isolated cases varies between 18 and 72%<sup>8-10</sup>. This may be the result of different treatment modalities and the fact that prenatal ultrasound cannot distinguish between positional and structural TEV<sup>9</sup>. Particularly for the group of isolated TEV, counselling by an orthopedic surgeon will help future parents to understand the condition and the availability of treatment<sup>11</sup>.

It may be assumed that the prognostic advantage of prenatal diagnosis of TEV is the detection of associated anomalies<sup>5,9</sup>. Depending on the severity of the associated anomalies, future parents have the opportunity to choose between different management options such as termination of pregnancy or continuation of pregnancy with non-intervention or standard management. In case of an isolated TEV they can prepare for the birth of an affected child that will require intensive treatment, particularly in the first year. In the Netherlands, consensus was reached for the treatment of congenital TEV in 1992. Treatment should start early and initially consist of repeated redressements followed by plaster cast immobilisation. In the more severe cases surgical treatment is subsequently needed. This should preferably be performed between the age of 4 and 9 months and be completed before the child starts walking in order to facilitate normal motor development<sup>12</sup> without the need of wearing modified shoes. The aim of the present study was to retrospectively determine whether, for congenital TEV requiring surgical treatment, prenatal detection makes a difference in infant outcome compared to detection only at birth.

## Patients and Methods

A diagnosis of TEV was made when both the tibia and the fibula were seen in the same plane as the sole of the foot. TEV was considered complex if other anomalies whether or not combined with an abnormal karyotype were also present.

Two groups of women were included in the study: a prenatal group in which the diagnosis of TEV was made by ultrasound in our own tertiary center and a postnatal group in which the diagnosis of TEV was only made at birth. In this latter group a prenatal ultrasound examination, if any, had always been carried out elsewhere. Inclusion criteria for both groups were as follows: a singleton pregnancy between 2000 and 2005 and redressement followed by surgical treatment; the latter always performed in the department of Pediatric Orthopedics of the Erasmus Medical Center (EMC). Adopted children were excluded from the study, since they always had started treatment much later.

In the postnatal group two subsets of infants could be distinguished. In the first subset (subset A) redressement treatment was commenced soon after birth followed by surgery in the department of Pediatric Orthopedics of the EMC. The second subset (subset B) received redressement treatment primarily in another hospital in the region followed by surgery at the department of Pediatric Orthopedics of the EMC.

In case of a complex TEV, treatment was only provided in the department of pediatric orthopedics of the EMC.

Redressement treatment consisted of application of a snugly fitted plaster cast well padded above the knee bent at 90 degrees to prevent the cast from sliding totally. Surgery was not considered necessary if at the end of the redressement treatment period radiographic examination demonstrated a completely corrected position of the foot. If the foot only showed a residual equinus deformity surgery primarily involved a simple postero-lateral release. In case the foot on clinical examination could not be put in valgus and abduction and on radiographic examination showed an abnormal relation between talus and os calcis, a more extensive postero-medial release was indicated.

On the basis of the retrospective nature of the study, the couples of the postnatal group were sent a questionnaire only following completion of the surgical treatment of the TEV. The purpose of this questionnaire was to obtain data concerning family and obstetric history by telephone. Notes were reviewed for maternal demographic data, fetal sonographic data and neonatal and surgical outcome with a minimal follow up of 12 months. A family history was considered positive if a family member up to the third degree or less of relation was known with a TEV. Age at surgery, type of surgery, number of surgical procedures, duration of hospital admission and recurrence rate of TEV were compared between the pre- and postnatal cohorts.

The Pearson chi-square test or Fisher's exact test was used to analyse categorical variables and the unpaired t-test or the Mann-Whitney test was applied to analyse continuous variables. All calculations were performed using the SPSS software package (release 10.1, SPSS Inc, Chicago, IL, USA). A p-value of <0.05 was considered statistically significant.



# Results

The total study group consisted of 151 cases from singleton pregnancies. The prenatal group was represented by 65 cases, of which 2 with an isolated TEV only underwent redressment therapy and a further 4 were treated elsewhere resulting in 59 cases for further analysis. The postnatal group consisted of 86 infants, of which 3 were adopted and 5 for which there was no response to the questionnaire resulting in 74 infants for further analysis. TEV was isolated in 84 cases and complex in 49 cases.

## Isolated TEV (N=84)

Isolated TEV was diagnosed in 20/59 (34%) fetuses in the prenatal group at a mean gestational age of 23 (19-36) weeks. Referral for a prenatal ultrasound examination was 14 (70%) times for a suspected abnormality and 6 (30%) times for an *a priori* risk of a fetal anomaly amongst those who had a positive family history of TEV (N=3). Unilateral TEV was established in 6 cases. There were 2 false positive cases associated with oligohydramnios due to premature rupture of the membranes or fetal growth restriction, yielding a 5.8% false-positive rate. One pregnancy ended in intrauterine death as a result of placental abruption at 31 weeks. In this case, a bilateral instead of the suspected unilateral TEV was diagnosed after delivery. Bilateral TEV was diagnosed prenatally and confirmed after delivery in 14 (70%) cases. Altogether 17 newborn from the prenatal group were eligible for treatment, representing 13 males and 4 females.

The postnatal group consisted of 64 infants with an isolated TEV, of which 43 were males and 21 females. A positive family history was present in 15/64 cases (23%). Bilateral TEV was established in 28/64 cases (44%). A total of 39 infants was referred directly after delivery for redressment treatment followed by surgery at a later stage (subset A) while in the remaining 25 infants redressment therapy was started at a regional hospital followed by surgery in the University Center (subset B). Delivery and postnatal treatment data for both the prenatal and total postnatal group are presented in Table 1. No significant difference was found for gestational age at delivery, birth weight and gender ratio between the prenatal and total postnatal group. The ratio of unilateral/bilateral TEV in the prenatal group (3/14) was significantly different ( $p < 0.01$ ) from the total postnatal group (36/28). No significant difference existed for age at surgery, type of surgery (postero-lateral or postero-medial release) and duration of admission between the prenatal group and postnatal subset A. However, a significant difference was found for all three variables ( $p < 0.01$ ;  $p < 0.001$ ;  $p < 0.001$ ) when comparing (1) the prenatal group and postnatal subset B and (2) postnatal subsets A and B. In postnatal subset A, three infants suffered a recurrence following surgery: one following a postero-lateral release at 9 months and two following a postero-medial release at 16 and 18 months, respectively.

## Complex TEV (N=49)

Complex TEV was diagnosed prenatally in 39/59 (66%) fetuses at a mean gestational age of 24 (14-36) weeks. Reason for referral for a prenatal ultrasound examination was a suspected fetal anomaly in every instance; in one case, there was also a family history of TEV. There were 19 males and 20 females. Twelve pregnancies were complicated by oligo/anhydramnios (N=5) or polyhydramnios (N=7). Fetal chromosome analysis was carried out in 34/39 (87%) cases, resulting in six abnormal karyotypes: triploidy (N=2), trisomy 18 (N=2), trisomy 21, marker chromosome. Termination of pregnancy was opted by the patient in 19 cases and intrauterine death occurred in 5 cases resulting in 15 liveborn infants with a normal karyotype in which 2 demonstrated bilateral instead of unilateral complex TEV and 1 turned out to be a pes equinus resulting in a 2.5% false positive rate. A further nine infants died during the neonatal period. The remaining six survivors underwent postero-medial surgical release.

The postnatal group consisted of 10 infants, of which nine were male and one was female. None had a positive family history. Bilateral TEV was diagnosed in eight cases.

All infants underwent both redressement and surgical treatment in the University Center, nine of which a postero-medial release and one a postero-lateral release.

A detailed list of the associated anomalies and outcome in complex TEV is given in Table 2. Delivery and treatment data are presented in Table 3. No significant difference was found for gestational age at delivery, birth weight, and distribution of unilateral and bilateral TEV between the prenatal and postnatal group. The same applied to the age at surgery, type of surgery and duration of admission. In the prenatal group, one infant with spina bifida suffered a recurrence of the TEV six months after a postero-medial release. When considering the confirmed isolated and complex cases of TEV together, a significant difference ( $p < 0.001$ ) in the proportion of isolated and complex TEV cases was established between the prenatal group (18 isolated and 39 complex TEV cases) and postnatal group (64 isolated and 10 complex cases)

## Discussion

To our knowledge this is the first study that analyses the outcome of treatment between a group of neonates with a diagnosis of TEV before birth and a group in which this anomaly was diagnosed only after delivery. In the presence of isolated TEV, there is no difference in outcome after a prenatal or postnatal diagnosis if redressement treatment followed by surgery is performed at a specialised Pediatric Orthopedic Center. However, if after a postnatal diagnosis of isolated TEV redressement treatment was commenced outside such a specialised center, surgery took place later, was of a more

complicated nature (postero-medial instead of postero-lateral release) and required a longer admission period. Counselling of women by a pediatric orthopedic surgeon during the prenatal period will, in most cases, result in postnatal treatment in the corresponding department. The majority of infants with isolated TEV that require surgical treatment are referred to the pediatric orthopedic department. Our study may be slightly biased as a small proportion of infants will remain for treatment in a regional general hospital. This mainly concerns the positional TEV where surgery is often not indicated. There is only one other study which compared the outcome of a prenatal and postnatal group with isolated TEV<sup>4</sup>. They only assessed the percentage of cases requiring surgery in both groups, which was 55 and 60%, respectively. Detailed analyses of age at surgery, type of surgery and admission period have not been previously reported.

During embryologic limb development the foot passes through three different positions. At first the foot is in a straight line with the leg. Then at the 30-mm stage the foot passes to a marked equinovarus-adductus position which is eventually followed at the end of the embryologic phase to only a slight equinovarus-adductus position<sup>13</sup>. It is believed that the arrest of bone growth between the 30 and 50 mm stage of the embryo will result in congenital TEV<sup>14</sup>. Therefore the majority of congenital TEV should be detectable at the 18-22 week anomaly scan. Tillet et al.<sup>10</sup> assumed that the reason why orthopedic surgeons still see cases of isolated TEV at birth that have not been diagnosed prenatally, is the possibility of a transient TEV in pregnancy<sup>15</sup>. However, studies investigating the detection rate of TEV<sup>4,11</sup> reached a maximum detection rate at the end of their investigation period of 35% and 64%, respectively. Although an obvious learning curve was demonstrated and experience would most likely continue to rise, even tertiary centers present false negative figures between 3 and 29% for bilateral TEV where, in the prenatal period, a unilateral TEV was assumed<sup>9,16,17</sup>. It is therefore highly plausible that still a number of cases of isolated TEV may only be detected at birth.

The male preponderance in isolated TEV present in both our prenatal and combined postnatal group confirms data from neonatal studies where the male female ratio is 2:1<sup>18</sup>.

Although the present study concentrated on TEV requiring surgery, it is remarkable that only two fetuses (10%) with a prenatal diagnosis of isolated TEV received redressement treatment only, whereas in the other 90% this was followed by surgery. This is a much higher percentage than the 48-72% requiring surgery reported in other studies of prenatally detected isolated TEV<sup>4,6,9,10</sup>. In the study by Woodrow et al.<sup>8</sup> only 18% of isolated cases needed surgery which was performed well beyond the first year of life. This discrepancy is most likely the result of different treatment modalities. Furthermore, in our clinic, the necessity for surgery is defined by radiographic and clinical examination at the end of the redressement treatment period. Only a completely corrected foot will not require surgery. To stimulate normal

motor development in neonates with a TEV, it is policy in the Netherlands to obtain a stable plantigrade foot by the time the infants start to walk. This approach has led to the present management with surgery being carried out between the age of 4 and 9 months<sup>12</sup>. Also, in the present study, only three patients (4%) in the isolated group suffered a recurrence which is comparable to other studies with surgically treated isolated TEV<sup>19,20</sup>.

In the prenatal group, 66% of the fetuses were diagnosed with a complex TEV whereas this was only 16% of the postnatal group. The significant distribution difference in proportion of isolated and complex TEV cases between the prenatal and postnatal group appears to be caused by the severity of the associated anomalies in the former group, resulting in a high incidence of termination of pregnancy, intrauterine and neonatal death. Previous prenatal studies<sup>3,7,17,21</sup> have shown similarly high figures but reports from countries with a longstanding experience in ultrasound screening demonstrate a more equal proportion of isolated and complex TEV cases<sup>5,9</sup>. It is likely that this is the result of more experience in the detection of congenital anomalies and as a consequence an increase in those isolated anomalies which are more difficult to detect.

In our prenatal complex TEV group, there were six cases (15.4%) with a chromosome anomaly. For complex TEV cases similar percentages were reported in previous studies with no chromosomal anomalies in isolated cases<sup>5,9,17</sup>. The majority of chromosomal anomalies represented trisomy 18 and triploidy. In the present study a case of hydrops associated with trisomy 21 was documented. Trisomy 21 is not typically related to TEV, as opposed to TEV hyperflexibility of the joints is a feature of this chromosomal anomaly<sup>22</sup>. The combination of complex TEV and trisomy 21 has been reported in a number of prenatal studies<sup>5,16,17,21</sup>. When a TEV is established prenatally, the present study clearly demonstrates that the ultrasonographer should be aware of the fact that in more than 50% of cases associated anomalies may be present. If the resolution of the ultrasound scan and the experience of the sonographer allow exclusion of additional anomalies in case of isolated TEV, we agree with Malone et al.<sup>23</sup> that there is no indication for karyotyping. Of the survivors with a complex TEV, the associated anomalies mainly represented musculoskeletal abnormalities and neural tube defects. These anomalies generally cause a stiff TEV which explains why the postero-medial release operation was required in all but one case to obtain a plantigrade foot. For both the prenatal and the postnatal group of complex TEV, the treatment was commenced in our Pediatric Orthopedic department and no difference was observed in timing and nature of surgery, period of admission and recurrence rate.

Infants from both the prenatal and postnatal complex TEV group showed 75 and 80% bilateral TEV, respectively. This confirms data by Bakalis et al.<sup>5</sup> and Treadwell et al.<sup>21</sup> who also detected bilateral TEV in the majority of prenatal complex TEV cases. Similarly to a previous report<sup>24</sup>, isolated TEV in our postnatal group was bilateral in approximately 50% of cases. However,

in the prenatal group of isolated TEV, only 17% revealed the unilateral form and a significant difference was established in the proportion of unilateral and bilateral cases between the prenatal and postnatal isolated TEV cases. Detection of a relative minority of isolated unilateral TEV cases was also shown in other studies<sup>4,16</sup>. This low detection rate of isolated unilateral TEV is probable the result of inexperience in recognising congenital anomalies. Maffulli<sup>25</sup> considered prenatal detection of TEV only of prognostic relevance in complex cases. This is at variance with the present study, which has demonstrated that there is a prognostic advantage in the prenatal detection of isolated TEV.

It can be concluded that prenatal detection of isolated TEV results in earlier and less complicated postnatal surgery and a shorter admission time, provided treatment is arranged in a pediatric orthopedic center. After prenatal detection of a complex TEV survival is low and determined by associated anomalies: outcome of TEV treatment is not dependent on timing of diagnosis.

**TABLE 1** The delivery and treatment data for isolated TEV of the prenatal group and the postnatal group A, referred directly for redressement and surgery and group B, referred for first surgery after redressement treatment is started elsewhere.

	Prenatal group N=17	Postnatal group A N=39	Postnatal group B N=25	P
GA at delivery (weeks)	39 ± 2	39 ± 2	39 ± 3	NS
Birth weight (g)	3193 ± 430	3359 ± 656	3211 ± 789	NS
Male/female	13/4	27/12	16/9	NS
Unilateral TEV	3	24	12	* < 0.01
Bilateral TEV	14	15	13	
Age at surgery (month)	6.3 ± 2.4	6.7 ± 2.9	8.6 ± 2.4	** < 0.01
				*** < 0.01
Type of treatment				
Postero-lateral release	10	20	1	** < 0.001
Postero-medial release	7	19	24	*** < 0.001
Duration of admission (days)	2.4 ± 1.7	2.8 ± 1.9	5.0 ± .4	** < 0.001
				*** < 0.001
Recurrence	0	3	0	

Values represent means ± SD. TEV, talipes equinovarus; GA, gestational age; ns, not significant; g, gram

\* p value for significant difference between the prenatal group and the combined postnatal groups A and B

\*\* p value for significant difference between the prenatal group and the postnatal group B

\*\*\* p value for significant difference between the postnatal group A and the postnatal group B

**TABLE 2 Associated anomalies and outcome in the prenatal and postnatal group with complex TEV**

Associated anomalies	TOP	IUD	NND	Alive	Alive
	Prenatal group			Postnatal group	
Central nervous system					
Exencephaly	2	-	-	-	-
Spina bifida	2	-	2	3	3
Encephalocele	1	-	-	-	-
Walker Warburg	1	-	-	-	-
Chromosomal					
Triploid	1	1	-	-	-
Trisomy 18		2	-	-	-
Trisomy 21	1	-	-	-	-
Marker chromosome with ventriculomegaly	-	1	-	-	-
Abdominal wall defect					
Canthrell pentalogy	-	-	1	-	-
Limb bodywall complex	1	-	-	-	-
Skeletal dysplasia					
Chondrodysplasia punctata	1	-	-	-	-
Achondrogenesis type II	1	-	-	-	-
Atelo-osteogenesis type II	1	-	-	-	-
Urogenital anomaly with oligohydramnios					
Obstructive uropathie	-	-	1	-	-
Polycystic renal dysplasia	-	1	-	-	-
Meckel Gruber syndrome	2	-	-	-	-
MCA					
Multiple vertebral anomalies with tethered cord, rib agenesis, anal atresia and dextrocardia	-	-	-	1	-
Hypoplastic right heart syndrome with pulmonary stenosis and transposition of the great arteries, multiple vertebral anomalies, horseshoe kidney	-	-	-	-	1
Arthrogryposis	1	-	1	1	4
Myotonic dystrophy	-	-	2	-	-
Hydrops	4	-	1	-	-
Larsen syndrome	-	-	-	-	2
Diaphragmatic hernia	-	-	1	-	-
Cleft lip/palate	-	-	-	1	-
Totaal	19	5	9	6	10

TEV, talipes equinovarus; TOP, termination of pregnancy; IUD, intrauterine death; NND, neonatal death; MCA, multiple congenital anomalies

**TABLE 3 The delivery and treatment data for complex TEV of the survivors of the prenatal and the postnatal group.**

	Prenatal group N=6	Postnatal group N=10	P
GA at delivery (weeks)	39 ± 1	39 ± 1	NS
Mean birth weight (g)	3440 ± 431	3031 ± 510	NS
Unilateral TEV	2	2	NS
Bilateral TEV	4	8	
Number with surgery	6	10	NS
Age at surgery (month)	11.6 ± 5.6	14.6 ± 9.4	NS
Type of treatment			
Postero-lateral release	0	1	NS
Postero-medial release	6	9	NS
Mean duration of admission (days)	4.5 ± 1.6	5.2 ± 1	NS
Recurrence	1	0	

Values represent means ± SD. TEV, talipes equinovarus; GA, gestational age; g, gram.



## REFERENCES

- 1 Jeanty P, Romero R, d'Alton M, Venus I, Hobbins JC. In utero sonographic detection of hand and foot deformities. *J. Ultrasound Med.* 1985; 4: 595-601.
- 2 Benacerraf BR. Antenatal sonographic diagnosis of congenital clubfoot: a possible indication for amniocentesis. *J Clin. Ultrasound* 1986; 14: 703-706.
- 3 Pagnotta G, Maffulli N, Aureli S, Maggi E, Mariani M, Yip KM. Antenatal sonographic diagnosis of clubfoot: a six-year experience. *J. Foot Ankle Surg.* 1996; 35: 67-71.
- 4 Keret D, Ezra E, Lokiec F, Hayek S, Segev E, Wientroub S. Efficacy of prenatal ultrasonography in confirmed club foot. *J Bone Joint Surg Br* 2002; 84: 1015-1019.
- 5 Bakalis S, Sairam S, Homfray T, Harrington K, Nicolaides K, Thilaganathan B. Outcome of antenatally diagnosed talipes equinovarus in an unselected obstetric population. *Ultrasound Obstet Gynecol* 2002; 20: 226-229.
- 6 Bar-On E, Mashlach R, Inbar O, Weigl D, Katz K, Meizner I. Prenatal ultrasound diagnosis of club foot: outcome and recommendations for counselling and follow-up. *J. Bone Joint Surg. Br.* 2005; 87: 990-993.
- 7 Rijhsinghani A, Yankowitz J, Kanis AB, Mueller GM, Yankowitz DK, Williamson RA. Antenatal sonographic diagnosis of club foot with particular attention to the implications and outcomes of isolated club foot. *Ultrasound Obstet Gynecol* 1998; 12: 103-106.
- 8 Woodrow N, Tran T, Umstad M, Graham HK, Robinson H, de Crespigny L. Mid-trimester ultrasound diagnosis of isolated talipes equinovarus: accuracy and outcome for infants. *Aust. N. Z. J Obstet Gynaecol* 1998; 38: 301-305.
- 9 Carroll SG, Lockyer H, Andrews H, Abdel-Fattah S, McMillan D, Kyle PM, Soothill PW. Outcome of fetal talipes following in utero sonographic diagnosis. *Ultrasound Obstet Gynecol* 2001; 18: 437-440.
- 10 Tillett RL, Fisk NM, Murphy K, Hunt DM. Clinical outcome of congenital talipes equinovarus diagnosed antenatally by ultrasound. *J Bone Joint Surg Br* 2000; 82: 876-880.
- 11 Burgan HE, Furness ME, Foster BK. Prenatal ultrasound diagnosis of clubfoot. *J Pediatr. Orthop.* 1999; 19: 11-13.
- 12 Keesen W, van der Eijken JW, Diepstraten AFM. De behandeling van de congenitale klompvoet in Nederland; resultaten van een enquête onder orthopedisch chirurgen. *Ned Tijdschr Geneesk* 2002; 136: 1710-1712.
- 13 Victoria-Diaz A, Victoria-Diaz J. Pathogenesis of idiopathic clubfoot. *Clin. Orthop. Relat Res.* 1984; 14-24.
- 14 Miedzybrodzka Z. Congenital talipes equinovarus (clubfoot): a disorder of the foot but not the hand. *J. Anat.* 2003; 202: 37-42.
- 15 Bar-Hava I, Bronshtein M, Orvieto R, Shalev Y, Stal S, Ben-Rafael Z. Caution: prenatal clubfoot can be both a transient and a late-onset phenomenon. *Prenat. Diagn.* 1997; 17: 457-460.
- 16 Shipp TD, Benacerraf BR. The significance of prenatally identified isolated clubfoot: is amniocentesis indicated? *Am J Obstet Gynecol* 1998; 178: 600-602.
- 17 Mammen L, Benson CB. Outcome of fetuses with clubfeet diagnosed by prenatal sonography. *J Ultrasound Med* 2004; 23: 497-500.
- 18 Wynne-Davies R. Family studies and the cause of congenital club foot. Talipes equinovarus, talipes calcaneo-valgus and metatarsus varus. *J. Bone Joint Surg. Br.* 1964; 46: 445-463.
- 19 Diepstraten AF. Congenital clubfoot. *Acta Orthop. Scand.* 1996; 67: 305-312.
- 20 Tschopp O, Rombouts JJ, Rossillon R. Comparison of posteromedial and subtalar release in surgical treatment of resistant clubfoot. *Orthopedics* 2002; 25: 527-529.
- 21 Treadwell MC, Stanitski CL, King M. Prenatal sonographic diagnosis of clubfoot: implications for patient counseling. *J Pediatr. Orthop.* 1999; 19: 8-10.
- 22 Jones, K. L. Smith's recognizable patterns of human malformation. In Saunders Staff WB (eds). W. B. Saunders Company: Philadelphia, 1988; 10-13.
- 23 Malone FD, Marino T, Bianchi DW, Johnston K, D'Alton ME. Isolated clubfoot diagnosed prenatally: is karyotyping indicated? *Obstet Gynecol* 2000; 95: 437-440.
- 24 Wainwright AM, Auld T, Benson MK, Theologis TN. The classification of congenital talipes equinovarus. *J Bone Joint Surg Br* 2002; 84: 1020-1024.
- 25 Maffulli N. Prenatal ultrasonographic diagnosis of talipes equinovarus: does it give the full picture? *Ultrasound Obstet Gynecol* 2002; 20: 217-218.



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## Diagnosis of urinary tract anomalies

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Fetal kidneys and urinary bladder filling can be visualised as early as 12-14 weeks of gestation. Renal tract abnormalities whether at urethral or at urethral level will provide well-defined images by medical ultrasound.

Whereas assessment of renal function during fetal life has been limited, grading of urinary tract dilation appears to be helpful in determining which infant will need postnatal evaluations (chapter 5. 1).

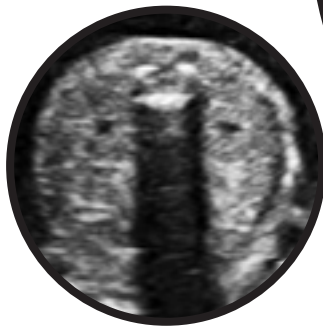
Abnormal renal morphology may not only be the result of urinary tract obstruction, it may also reflect abnormal development of the kidney itself, as is the case in multicystic dysplastic kidney. The postnatal outcome of this renal abnormality is determined by associated renal or non-renal structural pathology rather than the size or location of the renal anomaly itself (chapter 5. 2)



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- 1 Transverse image of the normal fetal abdomen with the spine at 12 o'clock. Right and left adjacent to the spine the kidney's are visible with a normal size pyelum.
  - 2 Transverse image of the fetal abdomen with the spine at 12 o'clock. Right and left adjacent to the spine the kidney's are visible with a dilated pyelum of the left kidney.
  - 3 Intra-venous pyelogram in infant showing the left kidney with pelviureteric junction obstruction.

# 5. 1

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## Mild renal pyelectasis in the second trimester; determination of cut-of levels for postnatal referral.

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T.E. Cohen-Overbeek (1), P. Wijngaard-Boom (1), N.T.C. Ursem (1),  
W.C.J. Hop (2), J.W. Wladimiroff (1), K.P. Wolffenbuttel (3)

1 Department of Obstetrics and Gynecology, Erasmus MC, Rotterdam, the Netherlands

2 Department of Epidemiology & Biostatistics, Erasmus MC, Rotterdam, the Netherlands

3 Department of Pediatric Urology, Erasmus MC, Rotterdam, the Netherlands

# Abstract

**Objective** To establish guidelines for postnatal referral of fetuses presenting with mild pyelectasy in the second trimester of pregnancy.

**Methods:** In a retrospective study 87 fetusus with a renal pelvis anterior posterior (RPAP) diameter of  $\geq 4$  mm and  $\leq 10$  mm before 28 weeks of gestation were included. All patients had a third trimester scan and fetuses with an RPAP diameter of  $\geq 10$  mm were referred for postnatal assessment. The family practitioner of all infants with an RPAP of  $< 10$  mm in the third trimester was contacted for follow up information. The RPAP diameter most predictive of renal pathology was determined with receiver-operating characteristics (ROC) curve analysis for both the first and second scans.

**Results:** In 36 of 87 infants, 49 abnormal kidneys were diagnosed. Seven infants required surgery on 8 renal tracts. The ROC curves of the first scan, second scan and differences between scans resulted in an area under the curve of 0.60, 0.87 and 0.85, respectively. The sensitivities and specificities for a cut-off level of 8, 9 and 10 mm at the second scan are 80%, 71% and 61% and 79%, 90% and 93% respectively. At a cut-off level of 10 mm, only cases of insignificant dilatation and a case of vesicoureteric reflux (VUR) requiring surgery were not detected.

**Conclusion:** After establishing a diagnosis of mild pyelectasy before 28 weeks a second scan is mandatory to determine which infants need postnatal evaluation. A cut-off level of 8 mm has a low specificity but includes most cases of pathology. A cut-off level at 10 mm detects most significant pathology, however VUR may be missed.

# Introduction

The fetal kidneys and bladder are readily visualised during prenatal scanning due to the collection of urine. In the second trimester, many consider the renal pelvis dilated when the renal pelvic anteroposterior (RPAP) diameter is  $\geq 4$  mm<sup>1-5</sup>. Mild pyelectasis is associated with an increased risk of associated pathology or chromosomal defects<sup>6,7</sup>. The Society of Fetal Urology introduced a scheme to grade excessive accumulation of fluid within the fetal kidneys<sup>8</sup>. From grade III upwards, when the renal pelvis is dilated with at least almost all calyces visible, there appears to be no doubt that infants should be referred for postnatal assessment and treatment in order to minimize renal injury and scarring<sup>9</sup>. In the case of mild pyelectasis, when only the renal pelvis with one or two calyces at most is visible, the need for postnatal assessment is less clear. RPAP cut-off levels varying from  $\geq 5$  mm at any stage in pregnancy<sup>10,11</sup> to  $\geq 8$  mm<sup>1</sup> in the third trimester are recommended for referral for postnatal assessment. This results in sensitivities ranging from 70%<sup>4</sup> to 100%<sup>11,12</sup>. Langer et al.<sup>13</sup> and Sairam et al.<sup>14</sup> have recommend postnatal referral only when the renal pelvis is  $\geq 10$  mm in the third trimester. These varying recommendations are confusing for doctors in the prenatal assessment of potential pathology and the advice they should give to parents, and they may lead to an unnecessarily large burden for pediatric urological clinics. Previous studies have included patients presenting in both the second and third trimester<sup>12,13</sup>, or included moderate as well as mild pyelectasis in the analysis<sup>10,11</sup>. Although Sairam et al.<sup>14</sup> evaluated the natural history of fetal hydronephrosis in a large prospective study, no follow up of infants with an antenatal RPAP  $< 10$  mm in the 3<sup>rd</sup> trimester was obtained. In the present study, fetuses presenting with mild second trimester pyelectasis and available third trimester and postnatal follow-up were investigated in order to establish clear guidelines for postnatal referral to a pediatric urology unit and to determine the risk of uropathy at different cut-off levels for renal pelvic size.

## Methods

From the 1<sup>st</sup> of January 1996 until the 31<sup>st</sup> of December 1999 all women with a singleton pregnancy and an RPAP diameter of  $\geq 4$  mm and  $\leq 10$  mm before 28 weeks of gestation were selected from the database of our tertiary referral center. Only kidneys with renal pelvis dilatation without calyceal dilatation were included. A third-trimester ultrasound scan was mandatory for inclusion. Fetuses with obvious obstructive uropathy as presented by a dilated urinary bladder and keyhole urethra at the first scan were excluded, as well as fetuses with associated or chromosomal anomalies. All fetuses with an RPAP diameter of  $\geq 10$  mm in the third-trimester scan were referred to a pediatric urologist for assessment. Postnatal ultrasound scans

were performed at the end of the first week. The postnatal cut-off level for a normal renal pelvis is <10 mm. However, caliectasis or ureterectasis were considered abnormal regardless of the size of the renal pelvis. Generally, when the renal pelvis exceeded 15 mm or calyceal dilatation was present, a Tc-99 mercaptoacetyltriglycine (MAG3) scan and micturating cystourethrogram (MCU) were performed.

The family practitioner of all infants with an RPAP of <10mm in the third trimester was contacted for follow-up information on possible renal tract anomalies and urinary tract infections. If referral to the pediatrician was indicated, this person was approached for further information on follow-up. Statistical comparison of percentages was carried out using the Chi-Square test. Continuous data were compared with the Mann-Whitney *U*-test. Sensitivities and specificities for various cut-off levels were calculated and graphically displayed using receiver-operating characteristics (ROC) curves. Logistic regression analysis was applied to determine the predictive chance of pathology for each RPAP diameter at the first and second scan. In case of bilateral pathology, both renal pelvises were analyzed separately.

## Results

Of the 6087 fetuses in the database 101 (1.7%) were found to have a RPAP diameter of  $\geq 4$  mm and  $\leq 10$  mm before 28 weeks of gestation. Three of these fetuses had a dilated bladder and keyhole urethra due to posterior urethral valves and were therefore excluded. In six cases associated anomalies were seen including hydrops ( $N=2$ ), omphalocele ( $N=1$ ), ventriculomegaly ( $N=2$ ), and a cardiac anomaly ( $N=1$ ). In two of these six fetuses a genetic syndrome was diagnosed at post mortem or after delivery. Trisomy 18 was diagnosed in two fetuses with multiple anomalies. The mean maternal age of the study population was 31.2 (range, 19-44) years ( $SD$  4.8). Three women were lost to follow up. Eighty-seven newborns with 174 renal units were available for the final analysis. Due to the retrospective design of the study, follow-up ranged from at least 1 year to 5 years.

The mean gestational age at the second and third trimester scan was 21.6 weeks ( $SD$  2.5) and 32.5 weeks ( $SD$  2.3). There were 65 males and 22 females, of which 26 (40%) and 10 (45%) showed anomalies in 37 and 12 renal units, respectively. There was no significant difference in the number of abnormal kidneys between males and females ( $P=0.65$ ). In Table 1 the postnatal diagnosis is classified according to cases and number of abnormal kidneys. A distinction is made between fetuses with both RPAP diameters <10 mm or at least one RPAP diameter  $\geq 10$  mm after 28 weeks of gestation. In the group where both RPAP diameters were <10 mm after 28 weeks' gestation, 43 (81%) showed no evidence of any renal tract anomaly. In 8 infants and 12



renal units a mild idiopathic dilatation, defined as a RPAP  $\geq 10$  mm and  $< 15$  mm, was established. In one infant, bilateral vesicoureteric reflux (VUR) was diagnosed. A scan at 32 weeks' gestation had shown both RPAP diameters to be 8 mm. In another infant, a unilateral dysplastic kidney was diagnosed at the age of 1 year. The antenatal RPAP diameter was 9 mm at 34 weeks' gestation. A postnatal ultrasound scan had been performed and no anomaly was established.

In the group where at least one RPAP diameter was  $\geq 10$  mm after 28 weeks, isolated pelviureteric junction obstruction (PUJO) was diagnosed in 17 renal units in 14 infants (41%). Isolated unilateral VUR existed in two infants. Unilateral VUR was diagnosed once in combination with PUJO and twice with renal duplication. A unilateral kidney cyst was diagnosed in one infant. Mild idiopathic dilatation was seen in six infants in 11 renal units. The RPAP diameter after 28 weeks in these 11 renal units varied between 4 and 18 mm. In 8 infants and 34 renal units there was no evidence of any renal tract pathology. In the total study group, 7 of 87 neonates (8%) underwent surgery, the nature of which is described in Table 2. The RPAP diameters at the first and second scan of all infants requiring surgery are provided in Table 2. In five infants, surgery was performed for PUJO, which involved bilateral pyeloplasty for one of them. In another infant, a partial nephrectomy was performed for renal duplication and VUR. One infant required bilateral endoscopic antireflux surgery and this was the only infant requiring surgery where both RPAP diameters after 28 weeks were  $< 10$  mm.

Antibiotic prophylaxis was prescribed to 21 newborn on the first day of life. This involved 18 of 33 newborns with a RPAP diameter  $\geq 10$  mm and 3 of 54 newborn with a RPAP diameter  $< 10$  mm. Despite this treatment six infants developed urinary tract infections, including two with VUR. In two infants with RPAP diameters of  $< 7$  mm at the second scan, a urinary tract infection was diagnosed. Mild idiopathic dilatation was subsequently diagnosed in one case, whereas in the other, no urogenital pathology was established. The predictive chance of all uropathology, described in Table 1, is calculated for each RPAP diameter at the first and second scan, and the results are displayed in Figures 1. For the first measurement the risk shows a moderate increase for greater diameters, whereas for the second measurement the risk demonstrates a much steeper rise. In Figure 2, the ROC curves for the RPAP diameter at the first and second scan, and for the RPAP diameter difference between the two scans, are displayed. The areas under the curve were 0.60, 0.87 and 0.85 respectively. When considering the second scan as the best performer (ROC area of 0.87), the sensitivities for a cut-off level of 8, 9 and 10 mm were 80%, 71% and 61%, respectively. The specificities for these cut-off values were 79%, 90% and 93%, respectively. At a cut-off level of 10 mm, only cases of insignificant minimal dilatation and a case of VUR requiring surgery were not detected.

## Discussion

The prevalence of 1.7% for mild renal pyelectasis in this study is comparable to that of other studies of low- and high-risk populations<sup>10,14,15</sup>. In our high-risk population associated anomalies were seen in 8% of cases. In other studies<sup>16-18</sup> this percentage varied between 3.9 and 31.6%. The relative high percentage of associated anomalies in the present study is likely to be due to the fact that a high-risk population was investigated. Mild renal pyelectasis has an increased risk of aneuploidy, particularly trisomy 21 when another risk factor such as advanced maternal age ( $\geq 36$  years) or associated anomalies are involved<sup>7</sup>. No case of trisomy 21 was encountered in the present study, but in two cases trisomy 18 was diagnosed. Both fetuses showed multiple associated anomalies at a maternal age of 25 and 32 years, respectively. Encountering mild renal pyelectasis should trigger the investigator to exclude associated anomalies and instigate invasive diagnosis if other risk factors are involved.

Our study confirms that isolated mild renal pyelectasis is significant more common in males than in females. This predominance of affected males has also been observed by others<sup>1,3,16,19</sup> and is in accordance with the postnatal incidence of hydronephrosis where PUJO is the most common diagnosis<sup>20</sup>. In our study, it is evident that from the 174 kidneys with mild renal pyelectasis in the second trimester, 148 kidneys did not reveal significant pathology in the postnatal period. There were 23 kidneys with a mild idiopathic dilatation that did not require surgery and in which there was no evidence of renal impairment. PUJO was diagnosed in 18 of the 174 kidneys; five infants with six renal units needed pyeloplasty. PUJO was the most common diagnosis in the pathology group followed by VUR, as has also been evident in other studies<sup>2,5</sup>. Apart from one case with bilateral VUR and one case with a unilateral dysplastic kidney diagnosed at the age of 1 year, all infants in whom significant uropathology was diagnosed had displayed RPAP diameters of  $\geq 10$  mm after 28 weeks of gestation.

VUR was diagnosed in six infants. Two infants required antireflux surgery, one of them showed bilateral RPAP diameters in the third trimester of only 8 mm. Reported percentages of VUR established after prenatal diagnosis of mild renal pyelectasis vary from 4.1 to 22%<sup>10,16</sup> but many cases of VUR are not detected<sup>16,21,22</sup>. To prevent this, all cases with a prenatal RPAP of  $\geq 5$  mm at any stage of pregnancy, would require an MCU<sup>10</sup>. Yerkes et al.<sup>23</sup> demonstrated that adopting such a policy would establish a high prevalence of VUR, whereas most of these cases would resolve without intervention and only require conservative management within 1-4 years. This policy would cause a high burden on medical resources without significantly improving morbidity in infants with mild degrees of VUR. Furthermore, there appears to be a high rate of spontaneous resolution of VUR within the first 2 years<sup>24,25</sup>. The issue of antibiotic prophylaxis in the presence of mild pyelectasis remains unclear in the literature. While Misra et al.<sup>26</sup> advise only to give

prophylaxis in case of bilateral hydronephrosis, others<sup>27</sup> recommend antibiotic prophylaxis when renal calyces are dilated. In the presence of VUR there is consensus about prophylactic therapy. Despite antibiotic prophylaxis, six infants in our study developed a urinary tract infection. In two of these cases VUR had been diagnosed. These figures are higher than those reported by Blachar et al.<sup>27</sup> and only emphasize the importance of prophylactic treatment in order to prevent urinary tract infection and subsequent renal scarring<sup>28</sup>. In two infants with RPAP diameters of <7 mm at the second scan, a urinary tract infection was diagnosed. Only in one case renal pathology in the form of mild idiopathic dilatation was subsequently established. These results are consistent with other studies<sup>2</sup>. Considering these findings we would support the advice from Kent et al.<sup>4</sup> that, if ultrasound examination in the third trimester is considered normal, no further investigations are required. However, the parents should be advised that if the infant develops a fever of unknown origin, a urine culture is necessary and infants with a proven urinary tract infection should be referred for treatment and radiologic work-up, including ultrasound and MCU. Despite the retrospective study design, contrary to other studies, we obtained a complete follow-up of all cases that entered the study, including those with a RPAP diameter in the third trimester of <10 mm. Furthermore, only 3% of cases were lost to follow up. This enabled us to provide a true picture of the incidence of postnatal pathology. The areas under the ROC curves for the first scan, the difference between the two scans, and the second scan indicate that the difference between measurements and the anteroposterior measurement of the pelvis after 28 weeks are both tests with a good prediction. It confirms that a renal pelvis which remains stable in size throughout pregnancy, especially with a RPAP diameter of ≤9 mm, is of little clinical significance<sup>5,13</sup>. In 60% of the patients of our study the RPAP diameter did not exceed 10 mm in the third trimester and in only a few of these cases a minimal dilatation was detected in the postnatal period. However, one case of VUR requiring surgery also belonged to this group. The graph for the predictive chance of pathology for each RPAP diameter at the first scan shows a linear increase with a maximum prediction of 55% at an RPAP diameter of 10 mm. The graph for the predictive chance of pathology for each RPAP diameter at the second scan shows a steeper increase particularly from 8 mm onwards. This information may be of use to both care providers and parents when confronted with a mildly dilated renal pelvis in the second trimester. As observed by others, a second scan in the third trimester is required to distinguish between normal and potential pathologic cases<sup>29,30</sup>. When counselling parents on the ability of ultrasound to detect renal anomalies one should bear in mind that not all cases of urogenital anomalies are identified at a second-trimester scan. Several authors have reported that cases of VUR and some cases of obstructive uropathy may be associated with a normal scan in the second trimester, with a pathological dilatation of the renal pelvis only being revealed in the third trimester or the postnatal

period<sup>21,31</sup>. All parents experience anxiety when confronted with any kind of fetal anomaly and serial investigations may only aggravate this<sup>32</sup> where, in case of mild renal pyelectasis no additional information is obtained<sup>33</sup>. Only, when a significant reduction of the amniotic fluid is detected in the presence of bilateral urinary tract pathology should repeat scans be recommended. In conclusion, after establishing mild renal pyelectasis before 28 weeks' gestation, a second scan is mandatory to determine which infants need postnatal evaluation. A cut-off level of 8 mm will include most cases of pathology, however, with a low specificity. At a cut-off level of 10 mm most significant pathology will be detected. In the presence of a RPAP diameter of <10 mm in the third trimester of pregnancy, postnatal investigations are not recommended. However, cases of VUR may not be detected. Our recommendation is that in case of a proven urinary tract infection further investigations are indicated.

**TABLE 1 Post partum diagnosis of renal anomalies in fetuses with a renal pelvic anteroposterior (RPAP) diameter of  $\geq 4$  mm and  $\leq 10$  mm before 28 weeks' of gestation and  $< 10$  mm or  $\geq 10$  mm after 28 weeks of gestation**

Diagnosis	Bilateral RPAP diameters $< 10$ mm after 28 wks gestation (N(%))		At least one RPAP diameter $\geq 10$ mm after 28 wks gestation (N(%))	
	Fetuses	Kidneys	Fetuses	Kidneys
PUJO	0	0	14 (41)	17
VUR	1 (2)	2	2 (6)	2
PUJO and VUR	0	0	1 (3)	1
Renal duplication and VUR	0	0	2 (6)	2
Dysplastic kidney	1 (2)	1	0	0
Solitary kidney cyst	0	0	1 (3)	1
Mild idiopathic dilatation	8 (15)	12	6 (18)	11
No pathology	43 (81)	91	8 (23)	34
Total	53	106	34	68

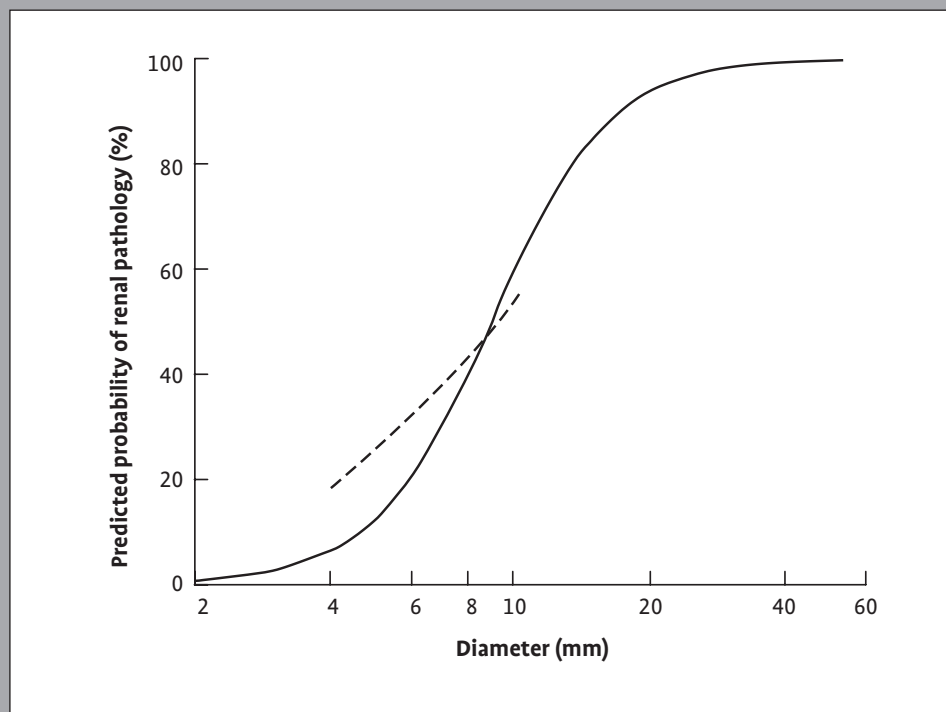
PUJO, pelviureteric junction obstruction; VUR, vesicoureteric reflux.

**TABLE 2 Measurements of the renal pelvic anteroposterior (RPAP) diameters in the first and second scan in all cases where surgery was performed.**

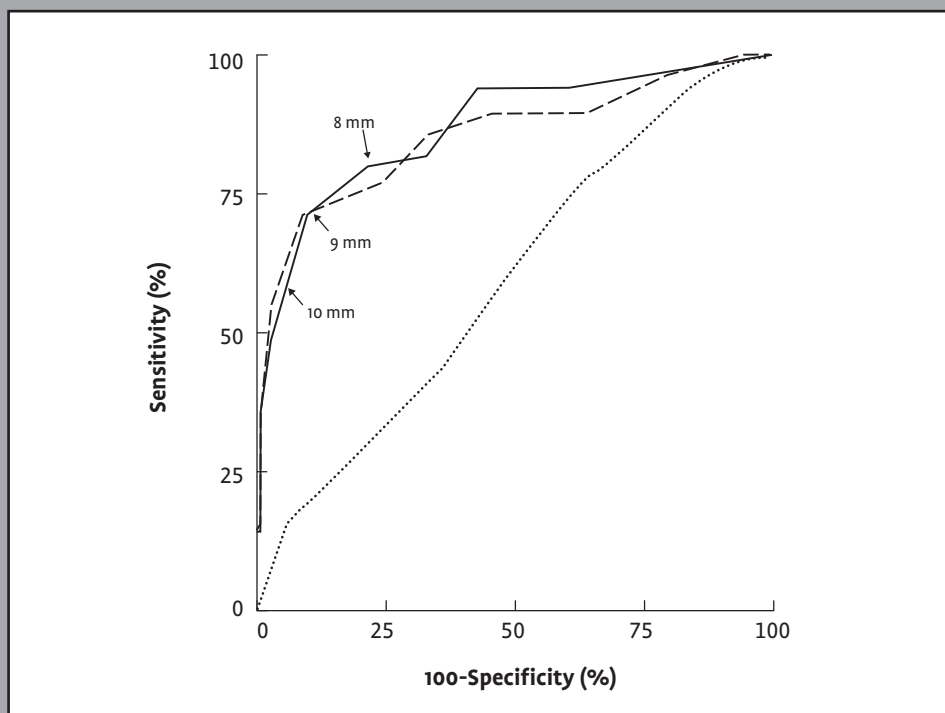
Gender	RPAP diameter at ultrasound scan $< 28$ weeks' gestation (mm)		RPAP diameter at ultrasound scan $\geq 28$ weeks' gestation (mm)		Postnatal diagnosis	Surgery
	L	R	L	R		
F	3	6	3	23	R PUJO	R pyeloplasty
M		8	4	25	R PUJO	R pyeloplasty
M	6	5	8	8	VUR L gr IV, VUR R gr II	Endoscopic antireflux surgery
F	6	8	54	27	Bilateral PUJO	Bilateral pyeloplasty
M	4		31	3	L PUJO and VUR gr II	L pyeloplasty
M	7	3	10	6	Renal duplication and VUR L lower pole gr V	L partial nephrectomy
M	3	5	6	16	R PUJO	R pyeloplasty

F, female; gr, grade; L, left; M, male; PUJO, pelviureteric junction obstruction; R, right; VUR, vesicoureteric reflux.

**FIG. 1** The predictive chance of pathology for each renal pelvic anteroposterior (RPAP) diameter of the first (dashed line) and second (solid line) scan. Equation of the dashed line:  $\log_e(\text{odds}) = -3.99 + 4.2 \times \log_2(\text{diameter}(\text{mm}))$ , with odds representing the odds ( $P/(100-P)$ ) of the predicted probability (%). Equation of the solid line:  $\log_e(\text{odds}) = -7.32 + 2.31 \times \log_2(\text{diameter}(\text{mm}))$ , with odds representing the odds ( $P/(100-P)$ ) of the predicted probability  $P$  (%).



**Figure 2** Receiver-operating characteristics curves for detection of renal pathology according to the renal pelvic anteroposterior diameter measurement at the first scan (dotted line) and the second scan (solid line) and the difference (dashed line) between the two measurements at the first and second scan.



## REFERENCES

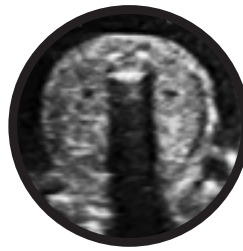
- Adra AM, Mejides AA, Dennaoui MS, Beydoun SN. Fetal pyelectasis: is it always "physiologic"? *Am. J. Obstet. Gynecol.* 1995; 173: 1263-1266.
- Morin L, Cendron M, Crombleholme TM, Garmel SH, Klauber GT, D'Alton ME. Minimal hydronephrosis in the fetus: clinical significance and implications for management. *J. Urol.* 1996; 155: 2047-2049.
- Stocks A, Richards D, Frentzen B, Richard G. Correlation of prenatal renal pelvic anteroposterior diameter with outcome in infancy. *J. Urol.* 1996; 155: 1050-1052.
- Kent A, Cox D, Downey P, James SL. A study of mild fetal pyelectasia-outcome and proposed strategy of management. *Prenat. Diagn.* 2000; 20: 206-209.
- Aviram R, Pomeran A, Sharony R, Beyth Y, Rathaus V, Tepper R. The increase of renal pelvis dilatation in the fetus and its significance. *Ultrasound Obstet. Gynecol.* 2000; 16: 60-62.
- Nicolaides KH, Cheng HH, Abbas A, Snijders RJ, Gosden C. Fetal renal defects: associated malformations and chromosomal defects. *Fetal Diagn. Ther* 1992; 7: 1-11.
- Chudleigh PM, Chitty LS, Pembrey M, Campbell S. The association of aneuploidy and mild fetal pyelectasis in an unselected population: the results of a multicenter study. *Ultrasound Obstet. Gynecol.* 2001; 17: 197-202.
- Fernbach SK, Maizels M, Conway JJ. Ultrasound grading of hydronephrosis: introduction to the system used by the Society for Fetal Urology. *Pediatr. Radiol.* 1993; 23: 478-480.
- Newell SJ, Morgan ME, McHugo JM, White RH, Taylor CM, Chapman S, Shah KJ, Gornall P, Corkery JJ. Clinical significance of antenatal calyceal dilatation detected by ultrasound. *Lancet* 1990; 336: 372.
- Jaswon MS, Dibble L, Puri S, Davis J, Young J, Dave R, Morgan H. Prospective study of outcome in antenatally diagnosed renal pelvis dilatation. *Arch. Dis. Child Fetal Neonatal* Ed 1999; 80: 135-138.
- Ouzounian JG, Castro MA, Fresquez M, al Sulyman OM, Kovacs BW. Prognostic significance of antenatally detected fetal pyelectasis. *Ultrasound Obstet. Gynecol.* 1996; 7: 424-428.
- Corteville JE, Gray DL, Crane JP. **Congenital hydronephrosis: correlation of fetal ultrasonographic findings with infant outcome.** *Am. J. Obstet. Gynecol.* 1991; 165: 384-388.
- Langer B, Simeoni U, Montoya Y, Casanova R, Schlaeder G. Antenatal diagnosis of upper urinary tract dilation by ultrasonography. *Fetal Diagn. Ther* 1996; 11: 191-198.
- Sairam S, Al Habib A, Sasson S, Thilaganathan B. Natural history of fetal hydronephrosis diagnosed on mid-trimester ultrasound. *Ultrasound Obstet. Gynecol.* 2001; 17: 191-196.
- Wickstrom EA, Thangavelu M, Parilla BV, Tamura RK, Sabbagha RE. A prospective study of the association between isolated fetal pyelectasis and chromosomal abnormality. *Obstet. Gynecol.* 1996; 88: 379-382.
- Persutte WH, Koyle M, Lenke RR, Klas J, Ryan C, Hobbins JC. Mild pyelectasis ascertained with prenatal ultrasonography is pediatrically significant. *Ultrasound Obstet. Gynecol.* 1997; 10: 12-18.
- Corteville JE, Dicke JM, Crane JP. Fetal pyelectasis and Down syndrome: is genetic amniocentesis warranted? *Obstet. Gynecol.* 1992; 79: 770-772.
- Snijders RJ, Sebire NJ, Faria M, Patel F, Nicolaides KH. Fetal mild hydronephrosis and chromosomal defects: relation to maternal age and gestation. *Fetal Diagn. Ther* 1995; 10: 349-355.
- Wilson RD, Lynch S, Lessoway VA. Fetal pyelectasis: comparison of postnatal renal pathology with unilateral and bilateral pyelectasis. *Prenat. Diagn.* 1997; 17: 451-455.
- Walsh PC, Retic AB, Stamey TA, Vaughan ED Jr. (eds). *Campbell's urology* (6th edn), W. B. Saunders: Philadelphia, PA, 1992;897-900.
- Economou G, Egginton JA, Brookfield DS. The importance of late pregnancy scans for renal tract abnormalities. *Prenat. Diagn.* 1994; 14: 177-180.
- Lepercq J, Beaudoin S, Bary F. Outcome of 116 moderate renal pelvis dilatations at prenatal ultrasonography. *Fetal Diagn. Ther* 1998; 13: 79-81.
- Yerkes EB, Adams MC, Pope JC, Brock JW, III. Does every patient with prenatal hydronephrosis need voiding cystourethrography? *J. Urol.* 1999; 162: 1218-1220.
- Marra G, Barbieri G, Moiolì C, Assael BM, Grumieri G, Caccamo ML. Mild fetal hydronephrosis indicating vesicoureteric reflux. *Arch. Dis. Child Fetal Neonatal* j112 Ed 1994; 70: 147-149.
- Yeung CK, Godley ML, Dhillon HK, Gordon I, Duffy PG, Ransley PG. The characteristics of primary vesicoureteric reflux in male and female infants with prenatal hydronephrosis. *Br. J. Urol.* 1997; 80: 319-327.
- Misra D, Kempey ST, Hird MF. Are patients with antenatally diagnosed hydronephrosis being over-investigated and overtreated? *Eur. J. Pediatr. Surg.* 1999; 9: 303-306.
- Blachar A, Blachar Y, Livne PM, Zurkowski L, Pelet D, Mogilner B. Clinical outcome and follow-up of prenatal hydronephrosis. *Pediatr. Nephrol.* 1994; 8: 30-35.
- Smellie JM, Poulton A, Prescod NP. Retrospective study of children with renal scarring associated with reflux and urinary infection. *BMJ* 1994; 308: 1193-1196.
- Ismaili K, Hall M, Donner C, Thomas D, Vermeylen D, Avni FE. Results of systematic screening for minor degrees of fetal renal pelvis dilatation in an unselected population. *Am. J. Obstet. Gynecol.* 2003; 188: 242-246.
- Langer B. Fetal pyelectasis. *Ultrasound Obstet. Gynecol.* 2000; 16: 1-5.
- Fugelseth D, Lindemann R, Sande HA, Refsum S, Nordshus T. Prenatal diagnosis of urinary tract



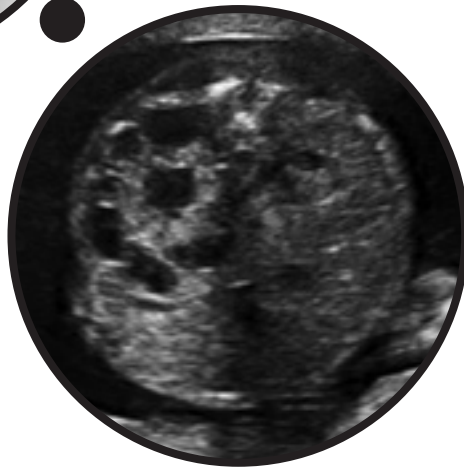
- anomalies. The value of two ultrasound examinations.  
Acta Obstet. Gynecol. Scand. 1994; 73: 290-293.
- 32 Harding LJ, Malone PS, Wellesley DG. Antenatal  
minimal hydronephrosis: is its follow-up an  
unnecessary cause of concern? Prenat. Diagn. 1999; 19:  
701-705.
- 33 Sherer DM. Is fetal hydronephrosis overdiagnosed?  
Ultrasound Obstet. Gynecol. 2000; 16: 601-606.



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- 1 Transverse image of the normal fetal abdomen with the spine at 12 o'clock. Right and left adjacent to the spine the kidney's are visible with a normal size pyelum.
- 2 Transverse image of the fetal abdomen with the spine at 12 o'clock. The left kidney is normally developed whereas the right kidney shows multiple cysts, consistent with a multicystic dysplastic kidney.
- 3 Multicystic dysplastic kidney after surgical excision.

# 5. 2

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## Unilateral multicystic dysplastic kidney: a combined pre- and postnatal assessment

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L. van Eijk (1), T.E. Cohen-Overbeek (1), N.S. den Hollander (1), J.M. Nijman (2), J.W. Wladimiroff(1)

1 Department of Obstetrics and Gynecology, University Hospital Rotterdam Dijkzigt and Sophia Children's Hospital, Rotterdam, the Netherlands

2 Department of Pediatric Urology, University Hospital Rotterdam Dijkzigt and Sophia Children's Hospital, Rotterdam, the Netherlands

# Abstract

**Objectives** To review the prenatal assessment of associated renal pathology, non-renal pathology and renal biometry, fetal outcome and postnatal urological management in the presence of unilateral fetal multicystic dysplastic kidney.

**Methods** A total of 38 singleton pregnancies with fetal unilateral multicystic dysplastic kidney was studied over a 13-year period. Prenatally, fetal biometry, including head and abdominal circumferences and largest longitudinal diameter of the affected and contralateral kidneys, was performed. The amount of amniotic fluid was assessed. Fetal karyotyping was offered in cases of contralateral renal or non-renal pathology. A MAG3 scan and voiding cystogram was performed approximately 4 weeks after delivery to establish renal function and to exclude urinary reflux.

**Results** Unilateral fetal multicystic dysplastic kidney was left-sided in 53% and right-sided in 47% of cases. The fetus was male in 63% and female in 37% of cases. Associated renal and non-renal pathology existed in 21% and 5% of cases, respectively. The fetal karyotype in these subsets was always normal. The longitudinal diameter of the multicystic dysplastic kidney was above the 95<sup>th</sup> centile in 87%. There was polyhydramnios in three cases and oligohydramnios in two cases. The prematurity rate was 16%. Postnatal examination revealed a non-functional multicystic kidney in 87% (33/38) of cases. Following surgical removal of the affected kidney, these infants progressed normally. Of the remaining five infants, four died because of associated anomalies and one infant developed normally without surgery.

**Conclusions:** Fetal outcome is determined by associated renal and/or non-renal structural pathology and not by the size/location of the unilateral multicystic dysplastic kidney or amniotic fluid volume.

# Introduction

The multicystic dysplastic kidney (MCDK) is a relatively common form of renal pathology. The MCDK can be unilateral, bilateral or segmental and probably results from atresia of the urethral bud system during embryogenesis<sup>1</sup>. The incidence of MCDK is approximately 1:4300 live births<sup>2,3</sup>. In approximately 20–50%, there are also abnormalities of the contralateral kidney which should also be evaluated. These abnormalities are mostly bilateral multicystic dysplastic kidneys, vesicoureteric reflux, ureteropelvic junction obstruction or renal agenesis<sup>3,4</sup>. Prenatal detection of MCDK allows the pediatric urologist to verify the diagnosis soon after delivery, to monitor the infant for urinary tract infection and to decide on further diagnostic procedures and treatment modalities. We review (i) the prenatal assessment of associated renal pathology, non-renal pathology and renal biometry; (ii) outcome and (iii) postnatal urological management in 38 cases of unilateral fetal MCDK seen in our department over a 13-year period.

## Methods

During a period of 13 years, a total of 38 singleton pregnancies was diagnosed with unilateral fetal MCDK. All women were referred to our prenatal center from regional community hospitals for further sonographic analysis. Thirty-four women were referred because of suspected fetal structural pathology at the referring hospital, four women were seen because of a previously affected infant. Details are provided in Table 1. Gestational age at referral was 18–39 weeks (median, 28 weeks). Maternal age ranged between 18 and 41 years (median, 26 years). All ultrasound examinations were carried out on a Toshiba SSA 270 (Toshiba Medical Systems Europe, Zoetermeer, the Netherlands) with 3.5- and 5.0-MHz curvilinear transducers.

Ultrasound evaluation included a detailed anomaly scan of the entire fetus. The diagnosis of MCDK was made from a sagittal and transverse cross-section of the abdomen at the level of the fetal kidneys. MCDK is characterized by normal size, hypoplastic or grossly enlarged kidneys, in which the normal renal parenchyma is replaced by multiple non-communicating cysts that may vary from only a few millimeters to several centimeters in diameter, giving the kidney a distinct hypoechoic appearance<sup>5,6</sup>. The largest MCDK and contralateral longitudinal kidney diameters were measured and data were plotted on the normal reference chart of Romero et al.<sup>7</sup>.

Oligohydramnios was defined as a largest fluid pocket of 2 cm or less; polyhydramnios was established when a largest fluid pocket of 10 cm or more was measured. Fetal karyotyping was offered to 12 out of 38 women by means of amniocentesis or cordocentesis because of contralateral renal or non-renal pathology. Fetal biometry included measurement of the biparietal diameter,

head circumference and abdominal circumference. For retrospective data analysis, normal reference charts by Snijders and Nicolaides<sup>8</sup> were used. On the second day following delivery, the infant underwent a confirmatory ultrasound scan of the renal area. A renal scan and voiding cystogram was performed approximately 4 weeks later to establish ipsi- and contralateral renal function and to exclude urinary reflux.

## Results

Of our series of 38 patients with unilateral fetal MCDK, 20 (53%) were left-sided and 18 (47%) were right-sided. Twenty-four (63%) fetuses were male and the remaining 14 (37%) were female. Contralateral renal pathology existed in five (13%) out of 38 cases, including three cases of hydronephrosis, one case of ureteral dilatation and one case of renal agenesis. Ipsilateral dilatation of the ureter was established in three (8%) cases. Extrarenal pathology consisted of a case of bilateral cleft lip and a case of combined microcephaly, micrognathia and complicated congenital heart disease. Fetal karyotyping was carried out through amniocentesis or cordocentesis in 12 cases (31%) because of contralateral renal or non-renal pathology. The chromosome pattern was always normal.

Fetal measurement of longitudinal MCDK diameter was available in 30 cases, with measurements above the 95<sup>th</sup> centile of the normal reference charts<sup>7</sup> in 26 cases (87%) and within the normal range in four cases (13%). Measurements of the contralateral kidney length were carried out in 20 cases, of which six (30%) were above the 95<sup>th</sup> centile, 12 (60%) were within the normal range and two (10%) were below the 5<sup>th</sup> centile, one because of renal dysplasia.

Fetal abdominal circumference data were available in 32 out of 38 patients. The fetal abdominal circumference was situated above the 95<sup>th</sup> centile of the normal reference chart<sup>8</sup> in seven cases (22%), was within the normal range in 21 cases (65%) and was below the 5<sup>th</sup> centile in the remaining four cases (13%).

The amniotic fluid volume was normal in 32 cases (84%). Oligoanhydramnios was established in two cases and polyhydramnios was observed in three cases although the increased amount of amniotic fluid was only temporary in two of these. Amniotic fluid volume was not estimated in one case.

Premature delivery (<37 weeks) took place in six out of 38 cases (16%). Instrumental delivery was performed in nine cases (24%), six of which were cesarean sections because of prolonged labor, fetal distress or breech position. Birth weight was normal in 30 out of 34 cases (88%), below the 2.5<sup>th</sup> centile in two cases and above the 97.5<sup>th</sup> centile in two cases. Data on birth weight could not be traced in the remaining four cases.

Postnatal sonographic examination confirmed the presence and location of the MCDK in every case. The infant with multiple congenital anomalies

died soon after delivery. The three cases of prenatally established contralateral hydronephrosis displayed vesicoureteric reflux in two cases and posterior urethral valves in the remaining case. Four infants (11%) developed a urinary tract infection, which was successfully treated. In cases of pre- or postnatal dilatation of the ureter or contralateral kidney, a voiding cystogram was performed 2–4 days following delivery. Renal scintigraphy (diethylenetriaminepenta-acetic acid and recently MAG 3) and voiding cystogram was routinely performed approximately 4 weeks after delivery to determine ipsi- and contralateral renal function and possible renal reflux in the remaining 37 infants. The MCDK was non-functional in 32 cases (86%) but demonstrated some activity varying between 7% and 18% of total renal function in the remaining five cases. Contralateral renal function was normal in all but two infants. In the infant with posterior urethral valves, the contralateral kidney showed dysplasia with diminished function and, in one infant, the contralateral kidney was absent. Vesicoureteric reflux was found in four infants.

Two infants died before surgery was performed (renal failure, epileptic insult). A further two infants were treated expectantly: one infant displayed agenesis of the contralateral kidney and died before the age of 1 year; in the other infant, the MCDK turned out to be so dysplastic that it could not be visualized by ultrasound. This infant subsequently developed normally. In the remaining 33 cases, the MCDK was surgically removed at the age of 3–6 months; in two cases, this included the ipsilateral ureter because of severe vesicourethral reflux. Pathological examination confirmed the diagnosis of MCDK in each case. All infants subsequently progressed normally.

## Discussion

MCDK seems to be related to urethral bud abnormalities<sup>9</sup> that lead to abnormal metanephric induction as well as defects involving the genitals, the hindgut and the cloacal derivatives.

The multicystic kidney lacks a discernable pelvis and calices and the abnormality is nearly always associated with atresia at the ureteropelvic junction<sup>10</sup>. Prenatally, the diagnosis of MCDK is only feasible using diagnostic ultrasound.

There is evidence that urological malformations are increasing. This rise may be due to the application of prenatal ultrasound revealing a previously unrecognized group of patients, or reflect the existence of a true increase in the incidence of this entity<sup>11</sup>. Since MCDK is nearly always non-functional, the prognosis depends entirely on the contralateral kidney. In the presence of bilateral MCDK or contralateral renal agenesis, these malformations are nearly always lethal<sup>12</sup>.

When both kidneys are involved, the amount of amniotic fluid may be severely reduced because of the virtual absence of fetal urine production.

In the present study, the amount of amniotic fluid was abnormal in five cases. In three cases, there was polyhydramnios which disappeared in two cases during the pregnancy. In the case where polyhydramnios did not disappear, an infant was born with a MCDK, hydroureter and reflux. In one of the two cases of anhydramnios, there was contralateral renal agenesis. The other case was characterized by posterior urethral valves. Kleiner et al.<sup>12</sup>, in a series of 27 cases of MCDK, established oligohydramnios in nine and polyhydramnios in two cases. In eight cases, oligohydramnios was associated with bilateral non-functional kidneys; five with bilateral MCDK and three with contralateral renal agenesis. The underlying mechanism for the development of polyhydramnios in the presence of MCDK is unclear. In the present study, MCDK was left-sided in 53% and right-sided in 47% of cases. These percentages are approximately the same as reported in other studies, where the MCDK tends to be slightly more left-sided<sup>13-15</sup>. In a recent study by Gough et al.<sup>11</sup>, left-sided MCDK was established in 63% of cases. In the present study, approximately 90% of the multicystic kidneys were enlarged. The contralateral kidney was enlarged in 30% of cases, 10% of which were a result of hydronephrosis. The remaining 20% represented enlargement of the normal kidney, which most likely reflects a compensatory effect in the presence of a non-functional MCDK. In two cases, the size of the contralateral kidney was below the 5<sup>th</sup> centile; one of which was the result of a dysplastic kidney. The size of the MCDK was not related to fetal outcome. The contralateral kidney was affected by pathology other than MCDK in 21% of cases, which is similar to a figure of 25% recently established by Lazebnik et al.<sup>16</sup>. However, the percentage of extrarenal abnormalities was five-fold (25%) higher in the latter study compared to our own data (5%), indicating the different populations presented in these studies. In the present study, contralateral renal and extrarenal pathology was associated with a normal fetal karyotype. Both numerical (trisomy 13 and 18) and structural chromosomal abnormalities have been documented by Lazebnik et al.<sup>16</sup> in association with extrarenal pathology. The incidence of extrarenal structural pathology in our study was low (N=2). Whereas, the clinical relevance of fetal chromosome analysis in the presence of contralateral renal pathology is questionable, we advocate fetal karyotyping when extrarenal pathology is established. In the presence of associated renal and/or non-renal structural pathology, the prenatal outcome was poor. Four out of eight infants died because of associated anomalies.

Most cases of MCDK are sporadic malformations. A positive family history is rare, since there is only a small risk of recurrence. However, MCDK may occur as part of the syndrome of hereditary renal A dysplasia (HRA), a predominantly autosomal-dominant abnormality with variable expression<sup>17</sup>. Other inheritance patterns of HRA have been suggested, such as recessive inheritance and X-linked inheritance. MCDK has been described in other syndromes, including the VATER association<sup>18</sup>, Williams syndrome<sup>19,49</sup>, XXXXX syndrome<sup>20</sup> and branchio-oto-renal syndrome<sup>21</sup>. Since nonrenal



abnormalities also are involved in these syndromes, it is essential to scan the entire fetus when MCDK is diagnosed. Accurate genetic counselling of couples with a family history of MCDK is mandatory.

The most common urological problem after MCDK seen prenatally was dilatation of the ureter (13%) of which 8% was contralateral. Postnatal examination showed that vesicoureteric reflux was the most common urological problem (13%) after MCDK, of which 8% was contralateral. This is somewhat lower than the percentages of contralateral vesicoureteric reflux reported in literature which range between 11% and 28%<sup>16,22,23</sup>.

Non-functional MCDK may be removed to avoid the risk of hypertension or malignancy<sup>24</sup>, although there is no consensus at present. An expectant non-intervention policy, on the other hand, makes long-term follow-up necessary as the risk of hypertension and malignancy is still uncertain. Cost-benefit studies are presently being carried out, but the outcome depends greatly on the various differences in healthcare systems. Certainly, in those cases with associated pathology of the contralateral kidney, a poorly functioning MCDK does not necessarily have to be removed, although with time most of these kidneys ultimately lose whatever remaining function they initially have. Most pediatric urologists do not feel comfortable to leave a severely dysplastic kidney *in situ*. In these cases, an individual approach is warranted. Because these children only have one functional kidney, which is at risk due to an increased incidence of contralateral abnormalities, a full postnatal evaluation of the urogenital tract has to be carried out and, until these studies are performed, the child should be given antibiotic prophylaxis. In one infant, the MCDK became so dysplastic that it could no longer be visualized. Earlier follow-up ultrasound studies<sup>16,25</sup> have documented incomplete or even complete involution of the antenatal MCDK.

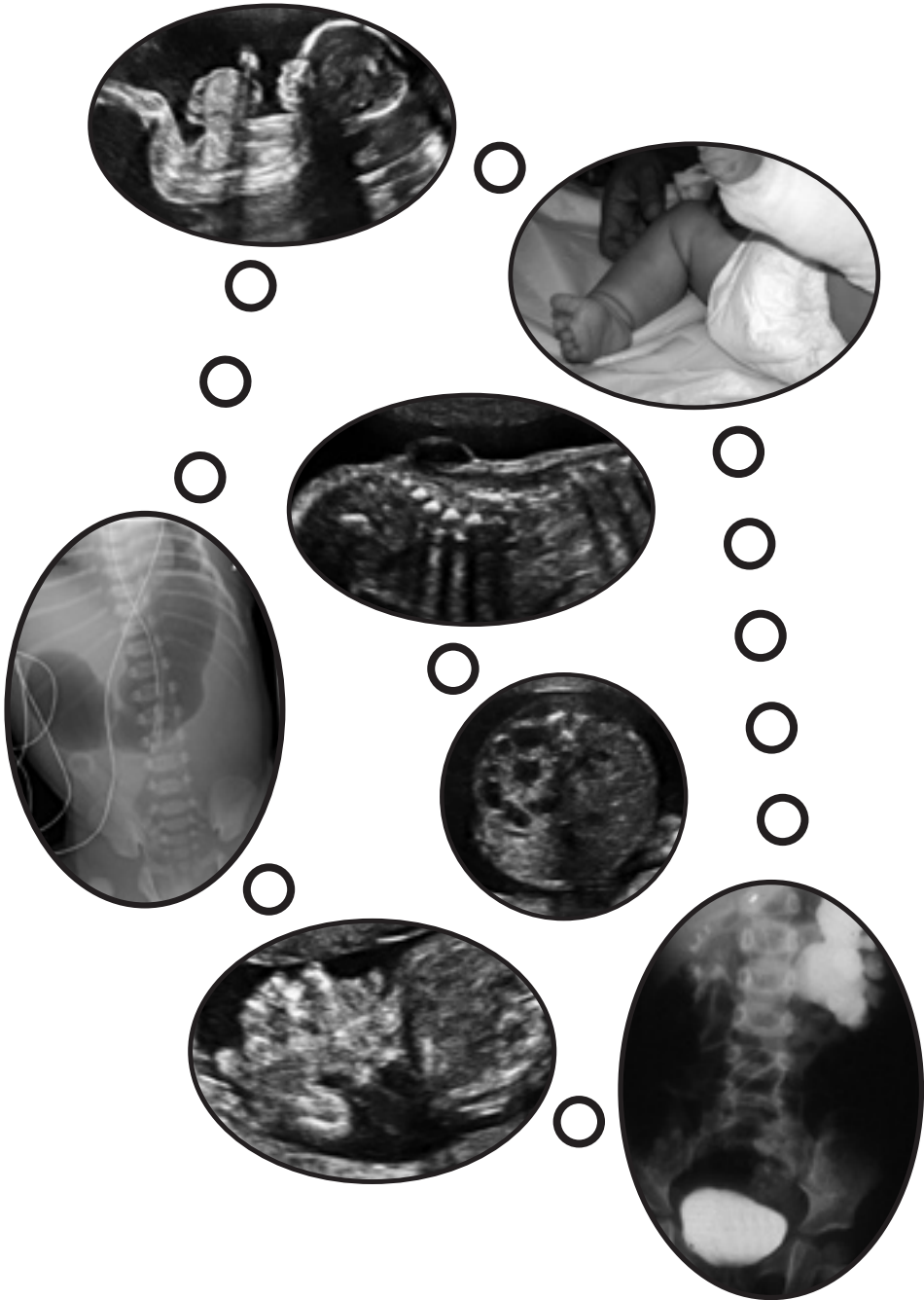
It can be concluded that, in the presence of unilateral MCDK, fetal outcome is determined by associated renal and/or non-renal structural pathology and not by the size/location of the unilateral MCDK or amniotic fluid volume. Prenatal knowledge of fetal unilateral MCDK allows optimization of diagnosis and treatment by the pediatric urologist after delivery. Chromosome studies are not needed in cases of isolated fetal unilateral MCDK.

**TABLE 1 Referral reasons for fetal anomaly scan**

	Number of patients
Suspected renal pathology	30
Dilated bowel loops	1
Fetal cardiac arrhythmia	1
'Double-bubble' phenomenon	1
Suspected urethral stenosis	1
Previous child with	
Dandy Walker syndrome	1
Potter syndrome	1
Congenital heart disease	1
Cleft lip	1

## REFERENCES

- 1 Felson B, Cussen LJ. The hydronephrotic type of unilateral congenital multicystic disease of the kidney. *Semin. Roentgenol.* 1975; 10: 113-123.
- 2 Gordon AC, Thomas DF, Arthur RJ, Irving HC. Multicystic dysplastic kidney: is nephrectomy still appropriate? *J. Urol.* 1988; 140: 1231-1234.
- 3 Orejas G, Malaga S, Santos F, Rey C, Lopez MV, Merten A. Multicystic dysplastic kidney: absence of complications in patients treated conservatively. *Child Nephrol. Urol.* 1992; 12: 35-39.
- 4 Glassberg KI and Filmer RB. Renal dysplasia, renal hypoplasia and cystic disease of the kidney. In *Clinical Pediatric Urology*, Kelalis PP, King LR, Bleman AB (eds). Saunders: Philadelphia, 1985; 922-971.
- 5 Stuck KJ, Koff SA, Silver TM. Ultrasonic features of multicystic dysplastic kidney: expanded diagnostic criteria. *Radiology* 1982; 143: 217-221.
- 6 Mahony BS, Filly RA, Callen PW, Hricak H, Golbus MS, Harrison MR. Fetal renal dysplasia: sonographic evaluation. *Radiology* 1984; 152: 143-146.
- 7 Romero R, Pilu G, Jeanty P, Ghidini A, and Hobbins JC. The urinary tract and adrenal glands: normal anatomy of the urinary tract. In *Prenatal diagnosis of congenital anomalies*, Delauter DL (eds). Appleton & Lange: Connecticut, 1988; 256.
- 8 Snijders RJM, Nicolaides KH. Fetal biometrie at 14-40 weeks' gestation. *Ultrasound Obstet. Gynecol.* 1994; 4: 34-38.
- 9 Squiers EC, Morden RS, Bernstein J. Renal multicystic dysplasia: an occasional manifestation of the hereditary renal adysplasia syndrome. *Am. J. Med. Genet. Suppl* 1987; 3: 279-284.
- 10 Bernstein J. The multicystic kidney and hereditary renal adysplasia. *Am. J. Kidney Dis.* 1991; 18: 495-496.
- 11 Gough DC, Postlethwaite RJ, Lewis MA, Bruce J. Multicystic renal dysplasia diagnosed in the antenatal period: a note of caution. *Br. J. Urol.* 1995; 76: 244-248.
- 12 Kleiner B, Filly RA, Mack L, Callen PW. Multicystic dysplastic kidney: observations of contralateral disease in the fetal population. *Radiology* 1986; 161: 27-29.
- 13 Wacksman J, Phipps L. Report of the Multicystic Kidney Registry: preliminary findings. *J. Urol.* 1993; 150: 1870-1872.
- 14 Mandell J, Paltiel HJ, Peters CA, Benacerraf BR. Prenatal findings associated with a unilateral nonfunctioning or absent kidney. *J. Urol.* 1994; 152: 176-178.
- 15 Selzman AA, Elder JS. Contralateral vesicourethral reflux in children with a multicystic kidney. *J. Urol.* 1995; 153: 1252-1254.
- 16 Lazebnik N, Bellinger MF, Ferguson JE, Hogge JS, Hogge WA. Insights into the pathogenesis and natural history of fetuses with multicystic dysplastic kidney disease. *Prenat. Diagn.* 1999; 19: 418-423.
- 17 Buchta RM, Viseskul C, Gilbert EF, Sarto GE, Opitz JM. Familial bilateral renal agenesis and hereditary renal adysplasia. *Z. Kinderheilkd.* 1973; 115: 111-129.
- 18 Lloyd DJ, McKenzie J, Kaye HH, Russell G. Vater syndrome: hypothesis and report of two further cases. *Teratology* 1977; 15: 43-46.
- 19 Fischbach M, Lutz JD, Tongio J, Sauvage P, Geisert J, Levi JM. Syndrome de Williams et Beuren avec hypertension et anomalies rénales associées. *Semin Hop Paris* 1979; 55: 13-14.
- 20 Toussi T, Halal F, Lesage R, Delorme F, Bergeron A. Brief clinical report: renal hypodysplasia and unilateral ovarian agenesis in the penta-X syndrome. *Am. J. Med. Genet.* 1980; 6: 153-162.
- 21 Widdershoven J, Monnens L, Assmann K, Cremers C. Renal disorders in the branchio-oto-renal syndrome. *Helv. Paediatr. Acta* 1983; 38: 513-522.
- 22 Atiyeh B, Husmann D, Baum M. Contralateral renal abnormalities in multicystic-dysplastic kidney disease. *J. Pediatr.* 1992; 121: 65-67.
- 23 Flack CE, Bellinger MF. The multicystic dysplastic kidney and contralateral vesicourethral reflux: protection of the solitary kidney. *J. Urol.* 1993; 150: 1873-1874.
- 24 Webb NJ, Lewis MA, Bruce J, Gough DC, Ladusans EJ, Thomson AP, Postlethwaite RJ. Unilateral multicystic dysplastic kidney: the case for nephrectomy. *Arch. Dis. Child* 1997; 76: 31-34.
- 25 Mesrobian HG, Rushton HG, Bulas D. Unilateral renal agenesis may result from in utero regression of multicystic renal dysplasia. *J. Urol.* 1993; 150: 793-794.



# 6

## General discussion

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# Introduction

Medical ultrasound is invaluable for the diagnosis of fetal anomalies. Since its introduction in prenatal diagnosis in the 1980's<sup>1</sup>, technical progress in ultrasound imaging enabled detection of fetal anomalies with increasing accuracy at earlier stages in pregnancy<sup>2</sup>. Obviously, fetal anomaly scanning would not have been given its prominent position in antenatal care, if it had not resulted in improvement of fetal and maternal welfare. Although a 50% reduction in perinatal mortality was established as a result of ultrasound detection of lethal anomalies, it proved more difficult to demonstrate changes in neonatal morbidity<sup>3</sup>. We therefore approached this subject by looking at the perinatal outcome of a number of different anomalies that were chosen to reflect both lethal and non-lethal anomalies in the context of pre- or postnatal detection. These different situations allow the estimation of the qualitative effect of prenatal detection on mortality and morbidity. Termination of pregnancy for a condition that is lethal postnatally, will not change mortality, when the mortality is not artificially considered separately before or after birth. Morbidity remains unchanged if infants with serious disabilities are allowed to survive. But in these situations mortality will increase if the pregnancy is terminated with as a result a reduction in morbidity. We will now consider the different conditions studied in this thesis separately within this context.

## Spina bifida

The spectrum of open isolated spina bifida involves on the one hand a complete rachischisis, with an invariably lethal outcome and on the other hand a minor meningocele without any physical or mental handicap following treatment. The majority of liveborn infants will present with a myelomeningocele and therefore with a certain degree of handicap depending on the spinal level of the defect<sup>4</sup>. Parents faced with this anomaly at a time when management options are still open will in the majority of cases request a termination of pregnancy<sup>5-7</sup>. Not all terminated cases would have resulted in death at birth as a number would have been eligible to treatment. Although improvement in the degree of disabilities or quality of life of affected individuals has been demonstrated, long term morbidity will be unavoidable for the majority of these individuals<sup>8</sup>. One may conclude that prenatal diagnosis of isolated spina bifida prior to 24 weeks of gestation will result in an increased mortality with a decreased morbidity and prevalence of the disease<sup>9</sup>. The audit of prenatal and postnatal diagnosis of isolated open spina bifida in the Netherlands proved that an ultrasound screening policy based on indications will only detect 22% of all cases before 24 weeks of gestation in contrast to the 66% detection rate in the second trimester reported by the Eurocat study<sup>10</sup>. The ultrasound screening policy in the

Netherlands was recently changed to a population based screening policy<sup>11</sup> and it remains to be seen whether as a result of this, the birth prevalence of spina bifida will reduce as much as is reported in other countries<sup>12</sup>. Preconceptional use of folic acid may additionally contribute to a reduction in the prevalence of neural tube defects<sup>13</sup>.

## Gastrointestinal and abdominal wall defects

Gastro-intestinal and abdominal wall defects represent another group of anomalies for which perinatal outcome was investigated following a prenatal diagnosis or a diagnosis at birth. Gastroschisis and omphalocele are both abdominal wall defects that are easily recognized with prenatal ultrasound.

### Gastroschisis

Contrary to omphalocele, gastroschisis is not associated with a high incidence of associated or karyotype anomalies. During pregnancy there is a 10% risk of intrauterine death and another 10% of cases is associated with small bowel atresia which may cause long term morbidity. The timing of the development of associated bowel atresia is not well understood but a recent study reported the development of dilated intra-abdominal bowel loops before 25 weeks of gestation in 8/10 of cases with bowel atresia, suggesting this complication may occur already early in the second trimester<sup>14</sup>. Contrary to spina bifida the majority of parents with prenatal detected cases will continue with their pregnancy. Our study did not reveal a difference in perinatal outcome following a pre- or postnatal diagnosis of liveborn infants, in spite of the fact that 90% of postnatal detected cases was delivered outside our tertiary center. The contribution of prenatal diagnosis to perinatal outcome lies in surveillance protocols with the purpose to detect those cases at risk of intrauterine death or developing complications from intestinal atresia such as perforations. Prenatal diagnosis may thus contribute to a decrease in mortality of this anomaly whereas morbidity will probably remain unaltered.

### Omphalocele

As the majority of omphaloceles are either associated with other anomalies or an abnormal karyotype (non-isolated omphalocele) parents often opt for a termination of pregnancy when confronted with this type of anomaly in the second trimester of pregnancy, even if the associated anomalies are not presumed lethal. Isolated omphaloceles form the minority of prenatal detected cases and survival is excellent although again approximately 20% of parents will opt for a termination of pregnancy<sup>15,16</sup>. Surprisingly, in the study presented in this thesis the perinatal outcome of liveborn

infants with prenatally diagnosed isolated omphalocele was considerably worse compared to infants where the anomaly was established only at birth. This was due to larger defects and more liver containing defects in the prenatal subset. Perinatal outcome for liveborn infants with non-isolated omphaloceles did not demonstrate a major difference between the prenatally and postnatally diagnosed infants. Three infants in the postnatal subset died as a result of a concomitant lethal karyotype, however. An indication based ultrasound screening policy will detect those isolated omphaloceles with a more complicated route to recovery. Karyotype anomalies of non-isolated postnatal cases may be missed, with as a result admission to a neonatal intensive care unit and use of medical resources until the aneuploidy is established. From the above data one may conclude that prenatal diagnosis of omphaloceles will contribute to an increase in mortality, at the same time morbidity and the prevalence will be reduced. In our studies of abdominal wall defects only 70% of both omphaloceles and gastroschisis were detected prenatally and a substantial number not before 24 weeks of gestation. Countries with a population based screening policy for congenital anomalies report much higher detection rates<sup>17,18</sup> of these anomalies and as a consequence prenatal diagnosis is likely to have a greater impact on mortality as well as morbidity<sup>19</sup>.

## Duodenal obstruction

In contrast to the prenatally diagnosed abdominal wall defects, duodenal obstruction (DO) is a gastrointestinal anomaly that presents only with symptoms late in the second or third trimester and in many cases clinical symptoms will only develop after birth. The obstruction can be complete or partial. As is the case with omphaloceles, associated anomalies complicate DO in 50% of cases and another 30% is associated with an abnormal karyotype, particularly trisomy 21. In our study only 30% of cases was detected prenatally of which two prior to 24 weeks of gestation. Both these latter cases were associated with another anomaly or trisomy 21 and the parents requested a termination of pregnancy. Countries with a population based screening policy report a 60% detection rate<sup>20</sup>, with the majority of the non-isolated cases detected before 24 weeks of gestation<sup>21</sup>. Both in the isolated and non-isolated form of DO birth weight was significantly lower and prematurity rate significantly higher in the prenatal subsets. For isolated DO this could be explained by the different types of obstruction between the pre- and postnatal subset. Non-isolated DO revealed similar associated anomalies and similar obstructions between both pre- and postnatal subset. The delivery mode was similar for both subsets and other factors such as for instance maternal stress could be responsible for the high prematurity rate of prenatally diagnosed DO. There was no difference in perinatal outcome after surgery between both pre- and postnatal isolated and non-isolated subsets despite the higher prematurity rate in both



prenatal subsets. Adjustment of obstetric management in the prenatal subsets could have contributed to this favorable result. Intrauterine and neonatal death occurred only in the presence of non-isolated DO. In view of the above data one may assume that an ultrasound screening policy based on indications will have a minimal influence on the mortality of the disease whereas the morbidity may improve slightly. If uptake of first trimester screening would be high, a substantial number of Down syndrome cases could be detected. Furthermore a population based ultrasound screening policy could detect approximately 60% of DO cases. Future evaluations of DO are necessary to investigate whether the latter two screening policies have an impact on the mortality, morbidity and prevalence of the anomaly.

## Talipes equinovarus (clubfoot)

Compared to the previously discussed abnormalities, talipes equinovarus (TEV) is a more frequently occurring anomaly. Recent prenatal series report an equal occurrence of isolated and non-isolated cases<sup>22,23</sup>, the latter in 15% associated with an abnormal, mostly lethal, karyotype. The perinatal outcome of isolated and non-isolated TEV detected prenatally or at birth was investigated. During a 5 year period 131 cases of TEV were studied. Fifty-nine cases (44%) were detected prenatally of which 39 (66%) were non-isolated and only six infants in this latter subset survived. This proportion was different in the postnatal detected subset where only 14% were non-isolated and all these infants survived. Following a prenatal diagnosis of isolated TEV, parents were counseled by a pediatric orthopedic surgeon and after delivery immediate redressement treatment was commenced in the Pediatric Orthopedic Center. Prenatally detected isolated TEV cases received therefore more often a simple type of surgery at an earlier stage with a shorter length of hospital stay compared to isolated cases detected after birth where redressement treatment was commenced in a district general hospital prior to surgery in the Pediatric Orthopedic Center. An indication based ultrasound screening policy would allow less than half of the TEV cases to be detected prenatally, of which the greater part would present with associated anomalies and a poor prognosis. Regions with a population based screening policy for congenital anomalies report higher detection rates<sup>24</sup> with equal proportions of isolated and non-isolated cases<sup>25</sup>. Our data indicated that as long as all infants with isolated TEV are treated in a Pediatric Orthopedic Center prenatal diagnosis for these cases would not be required to reduce morbidity. Generalisation of this finding is, however, critically dependent upon the accurate diagnosis of isolated TEV. Detection of non-isolated cases offers the possibility of additional investigations and adjustment of obstetric management. The mortality and morbidity of non-isolated TEV could thus alter as parents request a termination of pregnancy when confronted with

these anomalies. Although the effect is not estimated to be large, prenatal detection of TEV will lead to an increase in mortality. However, there will be a reduction in morbidity, due to termination of pregnancies in which the TEV is a marker for chromosomal or other associated anomalies.

## Urinary tract anomalies

### **Pyelectasis**

Prenatal detection of soft markers such as mild dilatation of the renal collecting system (pyelectasis) has a different impact on the efficacy of prenatal diagnosis. Soft markers are defined as subtle morphological changes that are often transient and have little or no pathological significance. The majority of cases with pyelectasis will resolve during pregnancy or shortly thereafter whereas a minority will develop pathology in need of treatment. When confronted with this knowledge half way through pregnancy, it may create uncertainty for both the pregnant woman and the provider of care but is unlikely to lead to any actions to be taken. Repeated ultrasound scans, to reassure the prospective mother will only increase her anxiety<sup>26</sup>. Boyd et al.<sup>17</sup> reported that soft markers will only marginally increase the detection of anomalies with a substantial increase in the false positive rate. In the working environment, it is impossible to ignore the finding of mild pyelectasis. A management strategy was required in order to determine which fetuses should be referred after delivery for postnatal assessment of the renal system while not unnecessarily increasing the level of anxiety of future parents. The number of infants with a pathologic and normal outcome was determined for different cut-off levels in the third trimester. Our study led to a new protocol for pre- and postnatal surveillance of fetuses with mild pyelectasis. Whilst this approach does not change mortality or morbidity it may change anxiety levels and hence morbidity in the parents.

### **Multicystic dysplastic kidney**

Another relatively easily recognized anomaly is the unilateral multicystic dysplastic kidney (MCDK). The kidney undergoes abnormal development as the connection between the ureteric bud and the metanephros is not properly established. The nephrons produce urine which can not be removed resulting in the formations of many unconnected cysts. This dilated multicystic structure is readily seen by ultrasound and may already be recognized early in the second trimester of gestation. The kidney may shrink in due course and ultimately disappear altogether, sometimes already during pregnancy, but most often later in life. The unilateral nature of the anomaly precludes loss of renal function which is completely taken over by the contralateral kidney. Associated anomalies including contralateral renal malformations determine the prognosis. Our study revealed that vesico-

ureteric reflux is the most common contralateral complication. This may not be recognisable with prenatal ultrasound and a combined pre- and postnatal assessment is therefore indicated. Unilateral MCDK seldom gives rise to clinical symptoms. An indication based ultrasound screening policy will recognize substantially less cases than a population based screening policy where 97% will be detected prenatally<sup>27</sup>. Mortality will not be affected but morbidity will be reduced as contra-lateral renal pathology is recognized and associated complications prevented.

## Conclusions

The impact on morbidity and mortality of a population based screening policy can be summarized as follows. In the presence of a lethal anomaly the mortality will not alter but will shift toward an earlier period in gestation. The morbidity associated with an abnormality may be reduced, at least assuming that even with a lethal anomaly infants will survive for a certain period of time. Prenatal termination of non-lethal major anomalies will increase mortality and reduce morbidity. Prenatal management can be adjusted following the detection and continuation of non-lethal major anomalies. Depending on the anomaly, like in gastroschisis, mortality may be reduced, but morbidity will not be altered. Prenatal detection at 18-22 weeks of gestation of major anomalies such as spina bifida, omphalocele and to a lesser extent DO will result in increased mortality and decreased morbidity. For other anomalies such as TEV and MCDK, morbidity may be reduced by early commencement of optimal treatment in a pediatric center. Knowledge of the postnatal development of prenatally detected minor anomalies such as mild pyelectasis will ensure selection of those infants at risk of developing pathology. The subsequent prevention of complications carries the capacity to reduce morbidity.

Several studies presented in this thesis demonstrate missed fetal diagnoses in spite of ultrasound investigations. For isolated spina bifida cases established at birth, 12% had been overlooked at a third trimester ultrasound scan in district general hospitals. Furthermore, 39% of prenatal isolated omphaloceles revealed associated anomalies following delivery with consequences for outcome. Of the prenatal cases of duodenal obstruction presumed isolated, 25% revealed additional chromosomal or associated anomalies after delivery. The prenatal isolated subset with TEV contained more infants with bilateral TEV compared to both postnatal subsets. Bilateral TEV is apparently more easily detected by ultrasound than unilateral TEV. The last three studies were conducted in our tertiary center. The first study underlies the fact that a screening programme can only be successful if carried out by competent sonographers equipped with modern ultrasound machines. Several learning points can be picked up from the false negative diagnoses in a tertiary center. Firstly, rapid postnatal follow up of prenatally

diagnosed anomalies and continuing reporting of postnatal diagnosis obtained from neonatology, pediatric surgery, genetic and post mortem investigations should increase knowledge of the variability in presentation of prenatal anomalies. The increasing number of anomalies detected by prenatal ultrasound will also contribute to the body of knowledge on the natural history of these anomalies. Secondly, continuing education on these subjects for both sonographers in primary care and doctors in prenatal medicine is important for obtaining a high standard upon which future parents can rely. Despite this, it is inevitable that some anomalies will be missed during pregnancy. Health care practitioners should be aware of this inadequacy of prenatal diagnosis when counselling parents.

## Suggestions for future research

Relatively few studies<sup>3,24,28-42</sup> have reported on the outcome of congenital anomalies with respect to the timing of diagnosis, the majority of which concerned cardiac anomalies<sup>36,38,40,41</sup>. We propose that the timing of diagnosis of congenital anomalies, whether before or after birth, or early or late in gestation, is further investigated for other organ systems, in order to evaluate the contribution of different prenatal ultrasound screening policies on fetal and maternal welfare. Follow up periods should be longer to evaluate the influence of the timing of detection, and possible change in obstetric management on child development.

Work in this thesis gave rise to a surveillance protocol for fetuses with gastroschisis and a management protocol for mild pyelectasis detected in the second trimester of gestation. Prospective evaluations of these protocols should be performed.

This thesis has concentrated on the medical aspects of pre- or postnatal diagnosis of congenital anomalies. It is generally assumed that prenatal diagnosis of congenital anomalies gives parents and health-care providers the opportunity to prepare for the birth of an infant requiring special attention. Additional information can be acquired and management options can be discussed without the medical or emotional pressure brought about when faced with a critically ill newborn<sup>43</sup>. Parents may experience a different reaction and suffer different degrees of stress when confronted with the fact that their child may not be healthy. It is likely that the use of prenatal diagnosis will only expand. Knowledge about the perception and coping mechanisms of parents, when confronted with the possibility that their child will suffer from a congenital anomaly, is important in order to be able to provide the best possible support during this stressful period. Although a number of studies concerning these matters have been presented<sup>44-47</sup> this field requires more research.

Finally, as stated before, prenatal ultrasound diagnosis has the capability to reduce suffering in infants and parents, although at a price of earlier

mortality. This can only be done when the quality of ultrasound diagnosis is high. It may therefore be questioned if potentially premature release of the procedure in the hands of relatively untrained health professionals like ultrasound technicians or midwives is effective<sup>48</sup>. This also requires structured training as well as continuing assessment of the role of these professionals. Economic or managerial considerations should not play a role in this before sufficient evaluations have been done.

## REFERENCES

- Campbell S, Pearce JM. The prenatal diagnosis of fetal structural anomalies by ultrasound. *Clin. Obstet. Gynaecol.* 1983; 10: 475-506.
- Souka AP, Nicolaides KH. Diagnosis of fetal abnormalities at the 10-14-week scan. *Ultrasound Obstet. Gynecol.* 1997; 10: 429-442.
- Bucher HC, Schmidt JG. Does routine ultrasound scanning improve outcome in pregnancy? Meta-analysis of various outcome measures. *BMJ* 1993; 307: 13-17.
- Cochrane DD, Wilson RD, Steinbok P, Farquharson DF, Irwin B, Irvine B, Chambers K. Prenatal spinal evaluation and functional outcome of patients born with myelomeningocele: information for improved prenatal counselling and outcome prediction. *Fetal Diagn. Ther.* 1996; 11: 159-168.
- Stoll C, Alembik Y, Dott B, Roth MP. Impact of prenatal diagnosis on livebirth prevalence of children with congenital anomalies. *Ann. Genet.* 2002; 45: 115-121.
- Morris JK, Wald NJ. Prevalence of neural tube defect pregnancies in England and Wales from 1964 to 2004. *J. Med. Screen.* 2007; 14: 55-59.
- Olde Scholtenhuis MA, Cohen-Overbeek TE, Offringa M, Barth PG, Stoutenbeek P, Gooskens RH, Wladimiroff JW, Bilardo CM. Audit of prenatal and postnatal diagnosis of isolated open spina bifida in three university hospitals in the Netherlands. *Ultrasound Obstet. Gynecol.* 2003; 21: 48-52.
- Bruner JP, Tulipan N. Tell the truth about spina bifida. *Ultrasound Obstet. Gynecol.* 2004; 24: 595-596.
- Busby A, Abramsky L, Dolk H, Armstrong B, Addor MC, Anneren G, Armstrong N, Bagueette A, Barisic I, Berghold A, Bianca S, Braz P, Calzolari E, Christiansen M, Cocchi G, Daltveit AK, De WH, Edwards G, Gatt M, Gener B, Gillerot Y, Gjergja R, Goujard J, Haeusler M, Latos-Bielenska A, McDonnell R, Neville A, Olars B, Portillo I, Ritvanen A, Robert-Gnansia E, Rosch C, Scarano G, Steinbicker V. Preventing neural tube defects in Europe: a missed opportunity. *Reprod. Toxicol.* 2005; 20: 393-402.
- Boyd PA, Wellesley DG, De Walle HE, Tenconi R, Garcia-Minaur S, Zandwijken GR, Stoll C, Clementi M. Evaluation of the prenatal diagnosis of neural tube defects by fetal ultrasonographic examination in different centers across Europe. *J. Med. Screen.* 2000; 7: 169-174.
- Hoogervorst J. F. Reply to questions from parlement concerning the changes in the set of measures regarding the new health insurance. 2005; DBO-K-U-2629961.
- Vinchon M. Deliberate termination of life of newborns with spina bifida. *Childs Nerv. Syst.* 2008; 24: 39-41.
- Prevalence of congenital malformations in the Northern Netherlands 1981-2006. 2007; <http://www.rug.nl/umcg/faculteit/disciplinegroepen/MedischeGenetica/Eurocat/professionals/tabellen>
- Nick AM, Bruner JP, Moses R, Yang EY, Scott TA. Second-trimester intra-abdominal bowel dilation in fetuses with gastroschisis predicts neonatal bowel atresia. *Ultrasound Obstet. Gynecol.* 2006; 28: 821-825.
- Rankin J, Dillon E, Wright C. Congenital anterior abdominal wall defects in the north of England, 1986-1996: occurrence and outcome. *Prenat. Diagn.* 1999; 19: 662-668.
- Barisic I, Clementi M, Hausler M, Gjergja R, Kern J, Stoll C. Evaluation of prenatal ultrasound diagnosis of fetal abdominal wall defects by 19 European registries. *Ultrasound Obstet. Gynecol.* 2001; 18: 309-316.
- Boyd PA, Chamberlain P, Hicks NR. 6-year experience of prenatal diagnosis in an unselected population in Oxford, UK. *Lancet* 1998; 352: 1577-1581.
- Brantberg A, Blaas HG, Haugen SE, Eik-Nes SH. Characteristics and outcome of 90 cases of fetal omphalocele. *Ultrasound Obstet. Gynecol.* 2005; 26: 527-537.
- Lakasing L, Cicero S, Davenport M, Patel S, Nicolaides KH. Current outcome of antenatally diagnosed exomphalos: an 11 year review. *J. Pediatr. Surg.* 2006; 41: 1403-1406.
- Singh MV, Richards C, Bowen JC. Does Down syndrome affect the outcome of congenital duodenal obstruction? *Pediatr. Surg. Int.* 2004; 20: 586-589.
- Haeusler MC, Berghold A, Stoll C, Barisic I, Clementi M. Prenatal ultrasonographic detection of gastrointestinal obstruction: results from 18 European congenital anomaly registries. *Prenat. Diagn.* 2002; 22: 616-623.
- Carroll SG, Lockyer H, Andrews H, Abdel-Fattah S, McMillan D, Kyle PM, Soothill PW. Outcome of fetal talipes following in utero sonographic diagnosis. *Ultrasound Obstet Gynecol* 2001; 18: 437-440.
- Bakalis S, Sairam S, Homfray T, Harrington K, Nicolaides K, Thilaganathan B. Outcome of antenatally diagnosed talipes equinovarus in an unselected obstetric population. *Ultrasound Obstet Gynecol* 2002; 20: 226-229.
- Keret D, Ezra E, Lokiec F, Hayek S, Segev E, Wientroub S. Efficacy of prenatal ultrasonography in confirmed club foot. *J Bone Joint Surg Br* 2002; 84: 1015-1019.
- Offerdal K, Jebens N, Blaas HG, Eik-Nes SH. Prenatal ultrasound detection of talipes equinovarus in a non-selected population of 49 314 deliveries in Norway. *Ultrasound Obstet. Gynecol.* 2007; 30: 838-844.
- Harding LJ, Malone PS, Wellesley DG. Antenatal minimal hydronephrosis: is its follow-up an unnecessary cause of concern? *Prenat. Diagn.* 1999; 19: 701-705.
- Wiesel A, Queisser-Luft A, Clementi M, Bianca S, Stoll C. Prenatal detection of congenital renal malformations by fetal ultrasonographic examination: an analysis of 709, 030 births in 12 European countries. *Eur. J. Med. Genet.* 2005; 48: 131-144.

- 28 Romero R, Ghidini A, Costigan K, Touloukian R, Hobbins JC. Prenatal diagnosis of duodenal atresia: does it make any difference? *Obstet. Gynecol.* 1988; 71: 739-741.
- 29 Miro J, Bard H. Congenital atresia and stenosis of the duodenum: the impact of a prenatal diagnosis. *Am. J. Obstet. Gynecol.* 1988; 158: 555-559.
- 30 Sipes SL, Weiner CP, Sipes DR, Grant SS, Williamson RA. Gastroschisis and omphalocele: does either antenatal diagnosis or route of delivery make a difference in perinatal outcome? *Obstet. Gynecol.* 1990; 76: 195-199.
- 31 Dillon E, Renwick M. The antenatal diagnosis and management of abdominal wall defects: the northern region experience. *Clin. Radiol.* 1995; 50: 855-859.
- 32 Adra AM, Landy HJ, Nahmias J, Gomez-Marín O. The fetus with gastroschisis: impact of route of delivery and prenatal ultrasonography. *Am. J. Obstet. Gynecol.* 1996; 174: 540-546.
- 33 Haddock G, Davis CF, Raine PA. Gastroschisis in the decade of prenatal diagnosis: 1983-1993. *Eur. J. Pediatr. Surg.* 1996; 6: 18-22.
- 34 St Vil D, Shaw KS, Lallier M, Yazbeck S, Di Lorenzo M, Grignon A, Blanchard H. Chromosomal anomalies in newborns with omphalocele. *J. Pediatr. Surg.* 1996; 31: 831-834.
- 35 Skari H, Bjørnland K, Bjørnstad-Ostensen A, Haugen G, Emblem R. Consequences of prenatal ultrasound diagnosis: a preliminary report on neonates with congenital malformations. *Acta Obstet. Gynecol. Scand.* 1998; 77: 635-642.
- 36 Bonnet D, Coltri A, Butera G, Ferment L, Le BJ, Kachaner J, Sidi D. Detection of transposition of the great arteries in fetuses reduces neonatal morbidity and mortality. *Circulation* 1999; 99: 916-918.
- 37 Marshall KW, Blane CE, Teitelbaum DH, van LK. Congenital cystic adenomatoid malformation: impact of prenatal diagnosis and changing strategies in the treatment of the asymptomatic patient. *AJR Am. J. Roentgenol.* 2000; 175: 1551-1554.
- 38 Verheijen PM, Lisowski LA, Stoutenbeek P, Hitchcock JF, Brenner JJ, Copel JA, Kleinman CS, Meijboom EJ, Bennink GB. Prenatal diagnosis of congenital heart disease affects preoperative acidosis in the newborn patient. *J. Thorac. Cardiovasc. Surg.* 2001; 121: 798-803.
- 39 Bittencourt DG, Barini R, Marba S, Sbragia L. Congenital duodenal obstruction: does prenatal diagnosis improve the outcome? *Pediatr. Surg. Int.* 2004; 20: 582-585.
- 40 Nikkila A, Bjørkhem G, Kallen B. Prenatal diagnosis of congenital heart defects-a population based study. *Acta Paediatr.* 2007; 96: 49-52.
- 41 Fuchs IB, Müller H, Abdul-Khalik H, Harder T, Dudenhausen JW, Henrich W. Immediate and long-term outcomes in children with prenatal diagnosis of selected isolated congenital heart defects. *Ultrasound Obstet. Gynecol.* 2007; 29: 38-43.
- 42 Anumba DO, Scott JE, Plant ND, Robson SC. Diagnosis and outcome of fetal lower urinary tract obstruction in the northern region of England. *Prenat. Diagn.* 2005; 25: 7-13.
- 43 Steinhorn RH. Prenatal ultrasonography: first do no harm? *Lancet* 1998; 352: 1568-1569.
- 44 Hunfeld JA, Wladimiroff JW, Passchier J, Venema-Van Uden MU, Frets PG, Verhage F. Emotional reactions in women in late pregnancy (24 weeks or longer) following the ultrasound diagnosis of a severe or lethal fetal malformation. *Prenat. Diagn.* 1993; 13: 603-612.
- 45 Skari H, Malt UF, Bjørnland K, Egeland T, Haugen G, Skreden M, Dalholt BM, Bjørnstad OA, Emblem R. Prenatal diagnosis of congenital malformations and parental psychological distress--a prospective longitudinal cohort study. *Prenat. Diagn.* 2006; 26: 1001-1009.
- 46 Brosig CL, Whitstone BN, Frommelt MA, Frisbee SJ, Leuthner SR. Psychological distress in parents of children with severe congenital heart disease: the impact of prenatal versus postnatal diagnosis. *J. Perinatol.* 2007; 27: 687-692.
- 47 Brisch KH, Munz D, Bemmerer-Mayer K, Terinde R, Kreienberg R, Kachele H. Coping styles of pregnant women after prenatal ultrasound screening for fetal malformation. *J. Psychosom. Res.* 2003; 55: 91-97.
- 48 Taipale P, Ammala M, Salonen R, Hiilesmaa V. Two-stage ultrasonography in screening for fetal anomalies at 13-14 and 18-22 weeks of gestation. *Acta Obstet. Gynecol. Scand.* 2004; 83: 1141-1146.





## Summary / samenvatting

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# Summary

Ultrasound has made it possible to observe in ever more detail the growth and development of the embryo and fetus. Unfortunately not all embryo's develop normally. In many instances this will result in a miscarriage but if this does not come about the congenital anomalies may be recognized by ultrasound. Minor and major anomalies occur with a frequency of 7 and 2.3%, respectively. Depending on the type of anomaly detection is possible from early, middle or late gestation. However, for a proportion of congenital anomalies such as congenital myopathies or metabolic storage diseases detection by ultrasound is not feasible. In the Netherlands universal population based screening for congenital anomalies was only introduced at the beginning of 2006, which was the reason why prior to this period many congenital anomalies were detected either late in pregnancy or sometimes only at birth. This situation set the scene to investigate whether the outcome of minor and major anomalies is altered by a prenatal detection versus a detection at birth. **Chapter 1** reviews the role of diagnostic ultrasound in Obstetrics and Gynecology and informs on the different types of ultrasound screening policies used to detect congenital anomalies. Targeted screening versus universal population based screening of congenital anomalies using these different methods are discussed. Additionally, the timing of detection is addressed as knowledge of the anomaly before 24 weeks of gestation provides the opportunity for parents to request a termination. Preparing for the birth of a possible critically ill infant in a controlled environment with the presence of a multidisciplinary team is equally important. At the end of the chapter the specific research objective are presented.

*Spina bifida* is one of the major congenital abnormalities that can be detected by ultrasound. We asked the question what proportion of isolated spina bifida cases an indication based system could detect. The study presented in **chapter 2** revealed that of the isolated spina bifida cases observed over a 4 year period in 3 university hospitals less than 25% was detected before 24 weeks of gestation. When faced with the diagnosis at a time that a choice of management in pregnancy is still possible, most parents requested a termination of pregnancy.

The outcome of three major gastrointestinal anomalies diagnosed prenatally or at birth are introduced in **chapter 3**. Two separate abdominal wall defects, gastroschisis and omphalocele, each present a different picture. The association of *gastroschisis* with other anomalies is low but intestinal atresia occurs in 10%. Additionally, approximately 10-15% of all cases die during pregnancy. Perinatal outcome did not reveal a difference for liveborn infants between the pre- and postnatal subset despite the fact that 8/9 postnatal infants were not delivered in our tertiary center, this in contrast to all prenatal diagnosed cases. Intestinal atresia was present in four infants,

three from the prenatal subset and one from the postnatal subset. These four cases experienced more morbidity and a longer length of hospital stay compared to all infants without atresia. The supplementary value of prenatal diagnosis for cases with gastroschisis lies in the recognition of fetuses at risk of a compromising outcome and a surveillance protocol was introduced in order to provide for an optimal outcome.

*Omphaloceles* are associated with other structural anomalies in 40% of cases and another 50% present with an abnormal karyotype. The defect is classified as isolated (without associated structural anomalies or an abnormal karyotype) or non-isolated. Non-isolated omphaloceles formed 80% of the prenatal cases and the majority (86%) did not survive as a result of termination of pregnancy, intrauterine or neonatal death contrary to the isolated cases where 62% survived. Surprisingly, liveborn infants with a prenatally detected isolated omphalocele displayed a worse outcome compared to infants with a postnatally detected isolated omphalocele. Both the larger sized and more liver containing omphaloceles in the prenatal subset could explain this difference. The outcome for liveborn infants with non-isolated omphaloceles did not reveal a major difference. However, the postnatal non-isolated subset contained three infants who died as a result of a lethal karyotype. An indication based system for prenatal recognition of congenital anomalies could only detect 70% of the abdominal wall defects. Particularly for omphaloceles and to a lesser extent for gastroschisis this policy has consequences for the mortality and morbidity of the disease. Comparable to omphalocele, *obstruction of the duodenum* (DO) is in 50% associated with other structural anomalies and in 30% with an abnormal karyotype, particularly trisomy 21. DO was considered isolated in the absence of other structural anomalies or an abnormal karyotype, whereas in all other cases DO was classified as non-isolated. The DO may be partial or complete and contrary to abdominal wall defects the anomaly is usually only detected by ultrasound late in the second or third trimester. In our study only 28/91 cases (30%) were detected before birth and only two before 24 weeks of gestation. The detection rate is approximately doubled in countries with a population based screening policy where the majority of the non-isolated cases are discovered prior to 24 weeks gestational age. Although liveborn infants from the prenatal isolated and non-isolated subsets had a higher prematurity rate and lower birth weight compared to the postnatal subsets, outcome was similar. The mechanisms responsible for the higher prematurity rate in the prenatal subsets are discussed. With an indication based system for prenatal recognition of congenital anomalies less than half of DO cases are detected before birth and only a minority before 24 weeks of gestation, even in the presence of a substantial number of associated anomalies.

*Talipes equinovarus* (TEV) is a frequently occurring anomaly and may be isolated or non-isolated. The perinatal outcome of TEV detected prenatally

or at birth is discussed in **chapter 4**. Of the 59 cases detected prenatally, 66% was associated with other structural or chromosome anomalies (non-isolated TEV) and only six infants survived. TEV was detected after birth in 74 infants of which only 13% were non-isolated and all survived. Following the prenatal diagnosis of TEV all infants were treated in a Pediatric Orthopedic Center. The prenatal diagnosis of isolated TEV was associated with less complicated postnatal surgical treatment compared to infants with a postnatal diagnosis where treatment was commenced in a district general hospital. Optimal outcome for infants with isolated TEV is ensured when treatment starts directly after birth in a Pediatric Orthopedic Center.

Anomalies of the urinary tract are presented in **chapter 5**. *Mild dilatation of the renal collecting system* (pyelectasis) is a common finding in second trimester ultrasound. The majority resolves during pregnancy or shortly thereafter. To determine which fetuses should be referred after delivery for postnatal assessment of the renal system, a retrospective study was conducted of 87 fetuses with mild pyelectasis in the second trimester. All fetuses underwent an ultrasound examination in the third trimester and follow up information after birth was arranged. The number of infants with a pathologic and normal outcome was determined for different cut-off levels in the third trimester. This resulted in a new protocol for pre- and postnatal surveillance of fetuses with mild pyelectasis.

The combined pre- and postnatal assessment of 38 cases with a unilateral *multicystic dysplastic kidney* (MCDK) is presented in chapter 5.2. Contralateral renal pathology was revealed in 13% of cases of which the majority demonstrated vesicoureteric reflux. Extrarenal pathology was present in 8%. Outcome was determined by the association of contralateral renal and non-renal pathology and not by the size or location of the unilateral MCDK. Prenatal knowledge of fetal unilateral MCDK allows optimization of diagnosis and treatment by the pediatric urologist after delivery.

**Chapter 6** of this thesis is dedicated to the analysis of the different studies and the consequences for perinatal mortality and morbidity. In the presence of a lethal anomaly a population based screening policy will not alter the mortality but shift this toward an earlier period in gestation. The morbidity associated with the abnormality may be reduced. Termination of non-lethal major anomalies will increase mortality and reduce morbidity. Prenatal management can be adjusted following the detection and continuation of non-lethal major anomalies. Depending on the anomaly, for instance in case of gastroschisis, mortality may be reduced, but morbidity will not be altered. Prenatal detection at 18-22 weeks of gestation of major anomalies such as spina bifida, omphalocele and to a lesser extent DO will result in an increase in mortality and a decrease in morbidity of the anomaly. For other anomalies such as TEV and MCDK morbidity may be reduced by early commencement of optimal treatment in a pediatric center. Knowledge of

the postnatal development of prenatally detected minor anomalies such as mild pyelectasis will ensure selection of those infants at risk of developing pathology. The subsequent prevention of complications carries the capacity to reduce morbidity.

# Samenvatting

Echoscopisch onderzoek heeft het mogelijk gemaakt om de groei en ontwikkeling van het embryo en de foetus met steeds meer detail waar te nemen. Helaas ontwikkelen niet alle embryo's zich normaal. Veelal resulteert dit in een miskraam, maar bij het uitblijven hiervan kunnen eventuele structurele aangeboren afwijkingen door middel van echoscopisch onderzoek worden herkend. Men maakt bij de aangeboren afwijkingen een onderscheid tussen ernstige ('major') en geringe ('minor') afwijkingen. Major aangeboren afwijkingen zijn levensbedreigend, vereisen uitgebreide chirurgie of hebben een ernstige cosmetisch effect. Alle andere afwijkingen worden als minor beschouwd. De frequentie van major en minor aangeboren afwijkingen bij de levend- en doodgeborenen is respectievelijk 2.3% en 7%. Afhankelijk van het soort afwijking is het mogelijk deze vroeg, in het midden of pas laat in de zwangerschap te ontdekken. Een deel van de aangeboren afwijkingen, zoals bij voorbeeld de congenitale spierziekten en stofwisselingsziekten, kan echter niet door middel van echoscopisch onderzoek worden opgespoord. Prenatale screening van de bevolking op structurele aangeboren afwijkingen werd in Nederland begin 2006 geïntroduceerd. Hiervoor werden structurele aangeboren afwijkingen veelal pas laat in de zwangerschap of na de geboorte ontdekt. Deze situatie bood de gelegenheid om te onderzoeken of de uitkomst van minor en major aangeboren afwijkingen verandert na een prenatale diagnose ten opzichte van een diagnose pas bij de geboorte. In **hoofdstuk 1** wordt de rol van het echoscopisch onderzoek in de zwangerschap besproken en wordt informatie gegeven over de verschillende echoscopische screeningsmethoden om structurele aangeboren afwijkingen op te sporen. Het verschil tussen de toepassing van echoscopie om aangeboren afwijkingen op te sporen bij gericht prenataal onderzoek vanwege een verhoogd risico op aangeboren afwijkingen ten opzichte van prenataal screeningsonderzoek van de totale bevolking wordt belicht. Daarenboven komt het tijdstip van detectie van een aangeboren afwijking aan de orde, aangezien bij een diagnose voor een zwangerschapsduur van 24 weken de ouders de mogelijkheid hebben om een verzoek te doen om de zwangerschap te beëindigen. Ook wordt besproken dat een belangrijk aspect van prenatale screening de mogelijkheid is om voorbereid te zijn op de geboorte een eventueel ernstig ziek kind in een gecontroleerde omgeving en in het bijzijn van een multidisciplinair team. Aan het eind van het hoofdstuk worden de afzonderlijke onderzoeksvragen gepresenteerd.

Een *spina bifida* of open rug is een major aangeboren afwijking die door middel van echoscopisch onderzoek kan worden aangetoond. Wij onderzochten welk percentage van de foetussen met een geïsoleerde spina bifida wordt gedetecteerd wanneer het echoscopisch onderzoek op indicatie wordt verricht. De studie in **hoofdstuk 2** laat zien dat van alle casussen met een geïsoleerde spina bifida die over een periode van 3 jaar in drie universiteitsklinieken wer-

den waargenomen minder dan 25% werd ontdekt voor een zwangerschapsduur van 24 weken. Ouders die met deze diagnose waren geconfronteerd op een termijn wanneer nog over het beleid in de zwangerschap mocht worden beslist, kozen voor het merendeel voor een afbreking van de zwangerschap en slechts 2/38 foetussen (5%) bleef in leven. Van de groep die ontdekt werd tussen 24 weken en de geboorte, werden 8 van de 50 zwangerschappen ondanks de late termijn getermineerd omdat de afwijking als lethaal werd beschouwd en 23 kinderen (46%) bleven uiteindelijk in leven. Wanneer de spina bifida bij de geboorte werd vastgesteld (N=65) bleef 79% in leven. De studie toonde aan dat ouders veelal een terminering van de zwangerschap verkozen wanneer deze afwijking voor 24 weken werd aangetoond. Bovendien liet de studie zien dat spina bifida voor de geboorte veelal een ernstiger aspect en somberder prognose heeft dan wanneer deze afwijking pas bij de geboorte aan het licht komt.

In **hoofdstuk 3** wordt de uitkomst besproken van drie major afwijkingen van de buikwand en het maagdarm kanaal die prenataal of bij de geboorte zijn vastgesteld. Twee verschillende buikwanddefecten, gastroschisis en omphalocèle laten ieder een andere uitkomst zien.

Een *gastroschisis* is een paramediaan buikwanddefect, meestal rechts, waardoor de darmen zich vrij in de amnionholte bevinden zonder omgeven te zijn door een vlies. De associatie van gastroschisis met andere structurele afwijkingen is laag maar een darmafsluiting komt bij 10% voor. Intra-uteriene sterfte komt voor bij 10-15%. De perinatale uitkomst toont geen verschil tussen de levendgeboren kinderen met een pre- of postnatale diagnose, ondanks het feit dat 8/9 kinderen met een postnatale diagnose niet in een tertiair centrum zijn geboren in tegenstelling tot alle kinderen met een prenatale diagnose. De toegevoegde waarde van prenatale diagnostiek voor de foetus met een gastroschisis ligt in de mogelijkheid om die foetussen te herkennen die een verhoogde kans hebben op een ongunstige uitkomst. Een bewakingsprotocol werd geïntroduceerd om hiermee een optimale uitkomst te bewerkstelligen.

Een *omphalocèle* is een defect van de buikwand in de umbilicale ring, waarbij abdominale structuren zich buiten de buikwand bevinden, omgeven door een membraan. De afwijking gaat in 40% gepaard met andere afwijkingen en is in 50% geassocieerd met een chromosoomafwijking. De afwijking was geïsoleerd (zonder andere afwijkingen of een chromosoomafwijking) of geassocieerd. De prenatale groep bestond voor 80% uit geassocieerde omphalocèles waarvan het merendeel (86%) niet overleefde als gevolg van een zwangerschapsafbreking, intra-uteriene vruchtdood of neonatale sterfte. Dit in tegenstelling tot de geïsoleerde omphalocèles waarbij 62% in leven bleef. Tot onze verrassing was de uitkomst voor levendgeboren kinderen met een geïsoleerde omphalocèle slechter na een prenatale diagnose in vergelijking tot de kinderen waarbij deze afwijking pas bij de geboorte werd ontdekt. Dit verschil in uitkomst kon verklaard worden doordat de omphalocèles van

de prenatale groep groter bleken en vaker lever bevatten. De uitkomst van levend geboren kinderen met een geassocieerde omphalocèle liet niet een duidelijk verschil zien tussen de pre- en postnataal gediagnosticeerde groep. De postnatale geassocieerde omphalocèle groep bevatte wel 3 pasgeborenen die overleden als gevolg van een letale chromosoomafwijking. Wanneer voor de detectie van aangeboren afwijkingen het echoscopisch onderzoek op indicatie wordt verricht, wordt slechts 70% van de buikwanddefecten prenataal ontdekt. Dit indicatie beleid heeft met name voor de omphalocèle en in mindere mate voor de gastroschisis gevolgen voor de mortaliteit en morbiditeit. Analooq aan de omphalocèle is een obstructie van het *duodenum* (de twaalfvingerige darm) in 50% geassocieerd met andere afwijkingen en in 30% met een chromosoomafwijking, met name trisomie 21. De afwijking werd gerubriceerd als geïsoleerd (zonder andere afwijkingen of een chromosoomafwijking) of geassocieerd. Een obstructie in het duodenum kan partieel of compleet zijn en in tegenstelling tot de buikwanddefecten wordt deze afwijking meestal pas laat in het tweede trimester of in het derde trimester van de zwangerschap met echoscopisch onderzoek ontdekt. In onze studie werden slechts 28/91 (30%) casussen voor de geboorte gedetecteerd waarbij niet meer dan 2 casussen voor de 24<sup>ste</sup> week van de zwangerschap werden opgespoord. De kans op detectie is verdubbeld in landen met screenings onderzoek op aangeboren afwijkingen van de totale bevolking, waarbij het merendeel van de geassocieerde duodenum obstructies wordt ontdekt voor de 24<sup>ste</sup> week van de zwangerschap. Hoewel de levendgeboren kinderen met een prenataal bekende geïsoleerde of geassocieerde duodenum obstructie een kortere zwangerschapsduur en lager geboorte gewicht hadden in vergelijking tot de groepen levendgeborenen en dezelfde afwijking, die pas na de geboorte aan het licht kwamen, maakte dit niet uit voor de uiteindelijke uitkomst. De oorzaken voor de hogere prematuriteit bij de prenatale groepen worden besproken. Bij een beleid gericht op indicatie om aangeboren afwijkingen voor de geboorte door middel van echoscopisch onderzoek op te sporen, zal de helft van de casussen met een duodenum obstructie voor de geboorte worden ontdekt en slechts enkelen voor een zwangerschaps termijn van 24 weken, zelfs als er sprake is van geassocieerde afwijkingen.

Een *klompvoet* is een frequent voorkomende afwijking die zich zowel geïsoleerd als geassocieerd presenteert. De perinatale uitkomst van kinderen met een of twee klompvoet(en) ontdekt voor of bij de geboorte wordt besproken in **hoofdstuk 4**. Van de 59 prenataal ontdekte casussen met een klompvoet waren 66% geassocieerd met bijkomende afwijkingen of een chromosoomafwijking (geassocieerde klompvoet). Slechts 6 kinderen van deze groep bleven in leven. Bij de geboorte werd een klompvoet bij 74 kinderen ontdekt. Er was sprake van een geassocieerde klompvoet bij 13% en allen bleven in leven. Nadat de diagnose klompvoet prenataal was vastgesteld, werden alle kinderen behandeld in een orthopedische kinderkliniek. Kinderen met een prenataal vastgestelde klompvoet kregen een eenvoudiger operatie gevolgd



door een kortere opnameduur ten opzichte van kinderen met een klompvoet vastgesteld na de geboorte bij wie de behandeling gestart werd in een algemeen ziekenhuis. De beste uitkomst voor kinderen met een geïsoleerde klompvoet wordt bewerkstelligd als de behandeling direct na de geboorte start in een orthopedische kinderkliniek.

Twee nierafwijkingen worden besproken in **hoofdstuk 5**. Een *milde dilatatie van het nierbekken* wordt frequent gezien in het tweede trimester van de zwangerschap. Het merendeel hiervan verdwijnt vanzelf gedurende de zwangerschap of kort daarna en bij slechts enkele kinderen zal een operatie noodzakelijk zijn. Om te bepalen welke kinderen na de geboorte verwezen moeten worden vanwege een grote kans op afwijkingen aan nieren en urinewegen werd retrospectief een onderzoek gedaan bij 87 foetussen met een milde dilatatie van het nierbekken in het tweede trimester van de zwangerschap. Bij alle foetussen werd het echoscopisch onderzoek van de nieren herhaald in het derde trimester en voor allen was follow-up tot minimaal 1 jaar na de geboorte beschikbaar. Voor de verschillende metingen van het nierbekken in het derde trimester van de zwangerschap kon het aantal kinderen met een normale en pathologische uitkomst worden bepaald. Met deze informatie kon een nieuw protocol voor pre- en postnataal beleid bij foetussen met een milde dilatatie van het nierbekken worden opgesteld.

De gecombineerde pre- en postnatale beoordeling van 38 casussen met een unilaterale *multicysteuze nierdyplasie* komt vervolgens aan de orde. Een multicysteuze nierdyplasie is een nier waarbij het grootste deel van het weefsel is vervangen door cystes waardoor de nier niet functioneert. Contralaterale nierpathologie werd bij 13% waargenomen waarvan het merendeel bestond uit vesico-urethrale reflux. Andere dan nierafwijkingen werd bij 8% aangetroffen. De uitkomst werd bepaald door de contralaterale nierpathologie en de geassocieerde andere afwijkingen en niet door de grootte of plaats van de unilaterale multicysteuze nierdyplasie. De diagnose en behandeling door de kinderuroloog wordt gunstig beïnvloed wanneer deze voor de geboorte reeds op de hoogte is van het bestaan van een unilaterale multicysteuze nierdyplasie.

In **hoofdstuk 6** van dit proefschrift wordt de invloed van de verschillende studies met betrekking tot perinatale morbiditeit en mortaliteit besproken. Bij een letale afwijking zal screeningsonderzoek van de totale bevolking naar aangeboren afwijkingen de mortaliteit niet veranderen, maar de termijn hiervan naar een vroeger tijdstip in de zwangerschap verplaatsen. De morbiditeit kan hierdoor wel afnemen. Het afbreken van de zwangerschap bij een niet letale afwijking zal de mortaliteit doen toenemen maar daarmee de morbiditeit verminderen. Het obstetrisch beleid kan worden aangepast nadat een niet letale afwijking is aangetoond en de zwangerschap wordt voortgezet. Afhankelijk van het soort afwijking, zoals bij voorbeeld bij een gastroschisis, kan de mortaliteit worden verlaagd terwijl de morbiditeit nauwelijks

verandert. Het prenataal vaststellen bij een zwangerschapsduur tussen 18 en 22 weken van major aangeboren afwijkingen, zoals spina bifida, omphalocele en in mindere mate een duodenum obstructie, zal de mortaliteit doen toenemen, waarbij de morbiditeit afneemt. Voor andere afwijkingen, zoals geïsoleerde klompvoeten en unilaterale multicysteuze nierdyplasie, kan de morbiditeit afnemen door na de geboorte vroeg te starten met de behandeling in een centrum voor kindergeneeskunde. Wanneer men op de hoogte is van het postnatale beloop van prenataal vastgestelde minor afwijkingen, zoals een dilatatie van het nierbekken, is het mogelijk alleen die kinderen na de geboorte te verwijzen waarbij een grote kans bestaat op het ontwikkelen van pathologie. Door het voorkomen van complicaties kan de morbiditeit afnemen.

# Curriculum Vitae

After secondary school at the Werkplaats Kindergemeenschap in Bilthoven, Titia Cohen-Overbeek (1954) started her medical studies at Leiden University in 1972. During her studies she did practical work in the Tel Hashomer Hospital, Tel Aviv, Israel and in the Department of Pediatrics at the Hammersmith Hospital in London, England. She obtained her medical degree in 1980. From March 1980 until November 1981 she worked as a resident at the department of Obstetrics and Gynecology in the Bronovo Hospital, The Hague. She moved to London in 1981 and worked in family planning clinics for the Lewisham Area Health Authority and Brook Advisory Centers in the South London area. In the meantime she commenced research in the field of Doppler ultrasound and prenatal diagnosis at Kings College Hospital, London, supervised by professor Stuart Campbell, where she remained as a part-time investigator until May 1987. From November 1987 she continued her work in prenatal diagnosis at the Department of Clinical Genetics (head Professor Dr. H. Galjaard) and the Department of Obstetrics and Gynecology, Division of Obstetrics and Prenatal Medicine (Professor Dr. J.W. Wladimiroff, Professor Dr. E.A.P. Steegers) of the Erasmus University Rotterdam. For her work she obtained three awards. The prize for the best clinical paper in volume 11 of *Ultrasound in Medicine and Biology* in 1985, the Dutch Organon ultrasound prize in 2000 for her contribution to prenatal ultrasound between 1998 and 2000 and the 2nd prize for all posters contributed at the 13<sup>th</sup> World Congress on Ultrasound in Obstetrics and Gynecology in Paris 2003 for the poster entitled 'Mild renal pyelectasis in the second trimester; determination of cut-off levels for postnatal referral'. She is married to Adam and the mother of Danielle, Sophie and Ben.

# PUBLICATION LIST

- Van De Putte LB, Hegt VN, **Overbeek TE**. Activators and inhibitors of fibrinolysis in rheumatoid and nonrheumatoid synovial membranes. A histochemical study. *Arthritis Rheum*. 1977; 20: 671-678.
- Campbell S, Diaz-Recasens J, Griffin DR, **Cohen-Overbeek TE**, Pearce JM, Willson K, Teague MJ. New doppler technique for assessing uteroplacental blood flow. *Lancet* 1983; 1: 675-677.
- Griffin D, **Cohen-Overbeek T**, Campbell S. Fetal and utero-placental blood flow. *Clin. Obstet. Gynaecol*. 1983;10:565-602.
- Cohen-Overbeek T**, Pearce JM, Campbell S. The antenatal assessment of utero-placental and feto-placental blood flow using Doppler ultrasound. *Ultrasound Med. Biol.* 1985; 11: 329-339.
- Campbell S, Pearce JM, Hackett G, **Cohen-Overbeek T**, Hernandez C. Qualitative assessment of uteroplacental blood flow: early screening test for high-risk pregnancies. *Obstet. Gynecol*. 1986; 68: 649-653.
- Hackett GA, Campbell S, Gamsu H, **Cohen-Overbeek T**, Pearce JM. Doppler studies in the growth retarded fetus and prediction of neonatal necrotising enterocolitis, haemorrhage, and neonatal morbidity. *Br. Med. J. (Clin. Res. Ed)* 1987; 294: 13-16.
- Campbell S, Bewley S, **Cohen-Overbeek T**. Investigation of the uteroplacental circulation by Doppler ultrasound. *Semin. Perinatol*. 1987; 11: 362-368.
- Pearce JM, Campbell S, **Cohen-Overbeek T**, Hackett G, Hernandez J, Royston JP. References ranges and sources of variation for indices of pulsed Doppler flow velocity waveforms from the uteroplacental and fetal circulation. *Br. J. Obstet. Gynaecol*. 1988; 95: 248-256.
- Cohen-Overbeek TE**, Jahoda MG, Wladimiroff JW. Uterine bloodflow velocity waveforms before and after transcervical chorionic villus sampling. *Ultrasound Med. Biol.* 1990; 16: 129-132.
- den Ouden M., **Cohen-Overbeek TE**, Wladimiroff JW. Uterine and fetal umbilical artery flow velocity waveforms in normal first trimester pregnancies. *Br. J. Obstet. Gynaecol*. 1990; 97: 716-719.
- Jahoda MG, Pijpers L, Reuss A, Brandenburg H, **Cohen-Overbeek TE**, Los FJ, Sachs ES, Wladimiroff JW. Transabdominal villus sampling in early second trimester: a safe sampling method for women of advanced age. *Prenat. Diagn*. 1990; 10: 307-311.
- Cohen-Overbeek TE**, Hop WC, den OM, Pijpers L, Jahoda MG, Wladimiroff JW. Spontaneous abortion rate and advanced maternal age: consequences for prenatal diagnosis. *Lancet* 1990; 336: 27-29.
- Jahoda MG, Brandenburg H, Reuss A, **Cohen-Overbeek TE**, Wladimiroff JW, Los FJ, Sachs ES. Transcervical (TC) and transabdominal (TA) CVS for prenatal diagnosis in Rotterdam: experience with 3611 cases. *Prenat. Diagn*. 1991; 11: 559-561.
- van den Anker JN, **Cohen-Overbeek TE**, Wladimiroff JW, Sauer PJ. Prenatal diagnosis of limb-reduction defects due to maternal cocaine use. *Lancet* 1991; 338: 1332.
- Hackett GA, **Cohen-Overbeek T**, Campbell S. The effect of exercise on uteroplacental Doppler waveforms in normal and complicated pregnancies. *Obstet. Gynecol*. 1992; 79: 919-923.
- Los FJ, Hagenaars AM, Marrink J, **Cohen-Overbeek TE**, Gaillard JL, Brandenburg H. Maternal serum alpha-fetoprotein levels and fetal outcome in early second-trimester oligohydramnios. *Prenat. Diagn*. 1992; 12: 285-292.
- Den Hollander NS, Stewart PA, **Cohen-Overbeek TE**, Heydanus R, Brandenburg H, Jahoda MGJ, Wladimiroff JW. Cordocentese en structureke afwijkingen bij de foetus. *Ned. Tijdschr. voor Obstetrie en Gynaecologie* 1992; 105: 343-345.
- van den Anker JN, van Vught EE, Zandwijken GR, **Cohen-Overbeek TE**, Lindhout D. Severe limb abnormalities: analysis of a cluster of five cases born during a period of 45 days. *Am. J. Med. Genet*. 1993; 45: 659-667.
- Jahoda MG, Brandenburg H, **Cohen-Overbeek T**, Los FJ, Sachs ES, Wladimiroff JW. Terminal transverse limb defects and early chorionic villus sampling: evaluation of 4, 300 cases with completed follow-up. *Am. J. Med. Genet*. 1993; 46: 483-485.
- Wladimiroff JW, Heydanus R, Stewart PA, **Cohen-Overbeek TE**, Brezinka C. Fetal renal artery flow velocity waveforms in the presence of congenital renal tract anomalies. *Prenat. Diagn*. 1993; 13: 545-549.
- van den Elzen HJ, **Cohen-Overbeek TE**, Grobbee DE, Wladimiroff JW. The predictive value of uterine artery flow velocity waveforms in miscarriage in older women. *Br. J. Obstet. Gynaecol*. 1993; 100: 762-764.
- Los FJ, Hagenaars AM, **Cohen-Overbeek TE**, Quartero HW. Maternal serum markers in second-trimester oligohydramnios. *Prenat. Diagn*. 1994; 14: 565-568.
- Den Hollander NS, **Cohen-Overbeek TE**, Heydanus R, Stewart PA, Brandenburg H, Los FL, Jahoda MG, Wladimiroff JW. Cordocentesis for rapid karyotyping in fetuses with congenital anomalies or severe IUGR. *Eur. J. Obstet. Gynecol. Reprod. Biol*. 1994; 53: 183-187.
- van den Elzen HJ, **Cohen-Overbeek TE**, Grobbee DE, Quartero RW, Wladimiroff JW. Early uterine artery Doppler velocimetry and the outcome of pregnancy in women aged 35 years and older. *Ultrasound Obstet. Gynecol*. 1995; 5: 328-333.
- van den Elzen HJ, Wladimiroff JW, **Cohen-Overbeek TE**, Morris CD, Grobbee DE. Calcium metabolism, calcium supplementation and hypertensive disorders of pregnancy. *Eur. J. Obstet. Gynecol. Reprod. Biol*. 1995; 59: 5-16.
- Naudin ten CL, Vermeij-Keers C, Smit DA, **Cohen-Overbeek TE**, Gerssen-Schoor KB, Dijkhuizen T. Intracranial

- teratoma with multiple fetuses: pre- and post-natal appearance. *Hum. Pathol.* 1995; 26: 804-807.
- van der Ham LI, **Cohen-Overbeek TE**, Geuze HD, Vermeij-Keers C. The ultrasonic detection of an isolated craniosynostosis. *Prenat. Diagn.* 1995; 15: 1189-1192.
- Brandenburg H, **Cohen-Overbeek TE**, Los FJ, Jahoda MJ. Transabdominal chorionic villus sampling and bowel loops as a restricting factor. *Prenat. Diagn.* 1995; 15: 387-388.
- Wladimiroff JW, Bhaggoo WR, Kristelijan M, **Cohen-Overbeek TE**, Den Hollander NS, Brandenburg H, Los FJ. Sonographically determined anomalies and outcome in 170 chromosomally abnormal fetuses. *Prenat. Diagn.* 1995; 15: 431-438.
- Brandenburg H, Los FJ, **Cohen-Overbeek TE**. A case of early intrauterine parvovirus B19 infection. *Prenat. Diagn.* 1996; 16: 75-77.
- van den Elzen HJ, Wladimiroff JW, **Cohen-Overbeek TE**, de Bruijn AJ, Grobbee DE. Serum lipids in early pregnancy and risk of pre-eclampsia. *Br. J. Obstet. Gynaecol.* 1996; 103: 117-122.
- Cha'ban FK, **Cohen-Overbeek TE**, Frohn-Mulder IM, Wladimiroff JW. Multiple intracardiac tumors: spontaneous prenatal recovery of fetal bradyarrhythmia. *Ultrasound Obstet. Gynecol.* 1996; 8: 120-122.
- van Opstal D, van den BC, Deelen WH, Brandenburg H, **Cohen-Overbeek TE**, Halley DJ, Van den Ouweland AM, In 't V, Los FJ. Prospective prenatal investigations on potential uniparental disomy in cases of confined placental trisomy. *Prenat. Diagn.* 1998; 18: 35-44.
- Mul TF, van Herwerden LA, **Cohen-Overbeek TE**, Catsman-Berrevoets CE, Lotgering FK. Hypoxic-ischemic fetal insult resulting from maternal aortic root replacement, with normal fetal heart rate at term. *Am. J. Obstet. Gynecol.* 1998; 179: 825-827.
- Cohen-Overbeek TE**, Wladimiroff JW. Standaard echoscopisch onderzoek: kijken, meten en zien. *Ned. Tijdschr. voor Obstetrie en Gynaecologie* 1998; 111: 5-8.
- Los FJ, van den BC, van OD, Noomen P, Braat AP, Galjaard RJ, Pijpers L, **Cohen-Overbeek TE**, Wildschut HI, Brandenburg H. Abnormal karyotypes in semi-direct chorionic villus preparations of women with different cytogenetic risks. *Prenat. Diagn.* 1998; 18: 1023-1040.
- van Gessel PH, Wildschut HI, **Cohen-Overbeek TE**, Vermeij-Keers C. Parvovirus B19 infection in pregnancy: a cause of non-immune hydrops fetalis. *Ned. Tijdschr. Geneesk.* 1999; 143: 3-7.
- van Eijk L, **Cohen-Overbeek TE**, Den Hollander NS, Nijman JM, Wladimiroff JW. Unilateral multicystic dysplastic kidney: a combined pre- and postnatal assessment. *Ultrasound Obstet. Gynecol.* 2002; 19: 180-183.
- Wessels MW, Den Hollander NS, **Cohen-Overbeek TE**, Lesnik Oberstein MS, Nash RM, Wladimiroff JW, Niermeijer MF, Willems PJ. Prenatal diagnosis and confirmation of the acrofacial dysostosis syndrome type Rodriguez. *Am. J. Med. Genet.* 2002; 113: 97-100.
- Olde Scholtenhuis MA, **Cohen-Overbeek TE**, Offringa M, Barth PG, Stoutenbeek P, Gooskens RH, Wladimiroff JW, Bilardo CM. Audit of prenatal and postnatal diagnosis of isolated open spina bifida in three university hospitals in the Netherlands. *Ultrasound Obstet. Gynecol.* 2003; 21: 48-52.
- Wladimiroff JW, **Cohen-Overbeek TE**, Ursem NT, Bijma H, Los FJ. Twenty years of experience in advanced ultrasound scanning for fetal anomalies in Rotterdam. *Ned. Tijdschr. Geneesk.* 2003; 147: 2106-2110.
- de Weerd S, Polder JJ, **Cohen-Overbeek TE**, Zimmermann LJ, Steegers EA. Preconception care: preliminary estimates of costs and effects of smoking cessation and folic acid supplementation. *J. Reprod. Med.* 2004; 49: 338-344.
- Cohen-Overbeek TE**, Wijngaard-Boom P, Ursem NT, Hop WC, Wladimiroff JW, Wolffenbuttel KP. Mild renal pyelectasis in the second trimester: determination of cut-off levels for postnatal referral. *Ultrasound Obstet. Gynecol.* 2005; 25: 378-383.
- van de Laar I, van Lange I, **Cohen-Overbeek TE**, Wilde A., Govaert L, ten Harkel A. Het Jervell en Lange Nielsen syndroom: klinische presentatie bij een zuigeling. *Tijdschr. Kindergeneesk.* 2006; 74: 114-118.
- Cohen-Overbeek TE**, Grijzeels EW, Lammerink EA, Hop WC, Wladimiroff JW, Diepstraten AF. Congenital talipes equinovarus: comparison of outcome between a prenatal diagnosis and a diagnosis after delivery. *Prenat. Diagn.* 2006; 26: 1248-1253.
- Klaassens M, Galjaard RJ, Scott DA, Bruggenwirth HT, van OD, Fox MV, Higgins RR, **Cohen-Overbeek TE**, Schoonderwaldt EM, Lee B, Tibboel D, de Klein A. Prenatal detection and outcome of congenital diaphragmatic hernia (CDH) associated with deletion of chromosome 15q26: two patients and review of the literature. *Am. J. Med. Genet. A* 2007; 143: 2204-2212.
- Wessels MW, De Graaf BM, **Cohen-Overbeek TE**, Spitaels SE, de Groot-de Laat LE, Ten Cate FJ, Frohn-Mulder IF, de KR, Bartelings MM, Essed N, Wladimiroff JW, Niermeijer MF, Heutink P, Oostra BA, Dooijes D, Bertoli-Avella AM, Willems PJ. A new syndrome with noncompaction cardiomyopathy, bradycardia, pulmonary stenosis, atrial septal defect and heterotaxy with suggestive linkage to chromosome 6p. *Hum. Genet.* 2008; 122: 595-603.
- Cohen-Overbeek TE**, Hatzmann TR, Steegers EA, Hop WC, Wladimiroff JW, Tibboel D. The outcome of gastroschisis after a prenatal diagnosis or a diagnosis

- only at birth Recommendations for prenatal surveillance. *Eur. J. Obstet. Gynecol. Reprod. Biol.* 2008; 139: 21-27.
- Dam L, **Cohen-Overbeek TE**, Poll-The BT, van Zalen-Sprock RM, Bilardo CM, Pajkrt E. OP03.06: Prenatally diagnosed ventriculomegaly: Associations and outcome. *Ultrasound Obstet. Gynecol.* 2008;32:318-319.
- Grijseels EWM, **Cohen-Overbeek TE**, Adama van Scheltema PN, Groenenberg IAL, Schoonderwaldt EM, Steegers EAP, Wildschut HJ. Kenmerken van de geïsoleerde sonomarker als nevenbevinding van het structureel echoscopisch onderzoek tijdens het tweede trimester van de zwangerschap. *Ned. Tijdschr. Geneesk.* accepted for publication.
- Cohen-Overbeek TE**, Grijseels EWM, Niemeijer N, Hop WCJ, Wladimiroff JW, Tibboel D. Isolated or non-isolated duodenal obstruction: perinatal outcome following a prenatal diagnosis or a diagnosis only at birth. *Ultrasound Obstet. Gynecol.* accepted for publication.
- Dorleijn DMJ, **Cohen-Overbeek TE**, Groenendaal F, Bruinse HW, Stoutenbeek Ph. Idiopathic polyhydramnios and postnatal findings. Submitted.
- Cohen-Overbeek TE**, Tong WH, Hatzmann TR, Wilms JF, Govaerts LCP, Steegers EAP, Hop WCJ, Wladimiroff JW, Tibboel D. Omphalocele; comparison of the perinatal outcome following a prenatal diagnosis or a diagnosis at birth. Submitted.
- Campbell S and **Cohen-Overbeek TE** The clinical value of bloodflow measurement in pregnancy. In *The fetus as a patient*, Kurjak A (eds). Excerpta Medica: Amsterdam, 1985; 300-311.
- Hackett GA, **Cohen-Overbeek TE** and Campbell S. The clinical value of bloodflow studies in obstetrics. In *Doppler techniques in obstetrics*, Jung H, Fendel H (eds). Georg Thieme Verlag: Stuttgart, 1986; 58-63.
- Cohen-Overbeek TE** and Campbell S. Doppler ultrasound techniques for the measurement of uterine and umbilical bloodflow. In *The uterine circulation*, Rosenfeld C. R. (eds). Perinatology Press: New York, 1989; 75-112.
- Wildschut HJ and **Cohen-Overbeek TE**. Afwijkingen van de tractus digestivus. In *Echoscopie in de gynaecologie en obstetrie*, Stoutenbeek PH, van Vught JMG, Wladimiroff JW (eds). Bunge: 1997; 113-117.
- Wladimiroff JW and **Cohen-Overbeek TE**. Afwijkingen van het centraal zenuwstelsel. In *Echoscopie in de verloskunde en gynecology*, van Vught J. M. G., Stoutenbeek Ph, Emanuel M. H., Wladimiroff JW (eds). Elsevier: Maarsen, 2003; 113-120.
- Wladimiroff, JW, **Cohen-Overbeek TE** and Laudy JAM. Ultrasound evaluation of the fetal thorax. In *Ultrasonography in Obstetrics and Gynecology*, Callen PW (eds). Saunders Elsevier: Philadelphia, 2007; 493-510.
- Cohen-Overbeek TE**. Het Structureel Echoscopisch Onderzoek in de zwangerschap. In *Ontwikkelingen in de geneeskunde 2007*, van der Meer LW, van der Poel BN M, de Vries AD, Weber RFA (eds). Erasmus MC, Het Congresbureau Rotterdam: Rotterdam, 2007; 87-91.

# Dankwoord

Toen in Nederland steeds maar niet het besluit werd genomen om structureel echoscopisch onderzoek bij een termijn van 18-22 weken aan de zwangere aan te bieden, leek het tijd een studie te doen naar de uitkomst van verschillende aangeboren afwijkingen waarbij een diagnose voor of pas bij de geboorte werd gesteld. De Gezondheidsraad had een onderzoek naar de uitkomst bij foetussen en kinderen met spina bifida geïnitieerd, hetgeen geleid werd door Katia Bilardo. Katia, ik ben je zeer dankbaar dat je direct je medewerking toezegde toen ik lang geleden suggereerde het onderzoek om te zetten in een publicatie zodat dit als startpunt voor mijn proefschrift kon dienen. Dat je in mijn promotiecommissie plaats neemt, hadden we niet kunnen fantaseren toen onze vriendschap lang geleden startte in King's College Hospital. De 23 overige co-auteurs ben ik zeer erkentelijk voor hun bijdrage en enkele hiervan wil ik in het bijzonder noemen.

In de eerste plaats mijn promotor, Juriy Wladimiroff. Als hoofd van de afdeling verloskunde en prenatale diagnostiek ondersteunde je aan het begin van de 90-er jaren onze lang strijd om gebruik te kunnen maken van een eigen database voor de registratie van het echoscopisch onderzoek. Dat was aanvankelijk tegen alle wensen van de Informatie Technologie afdeling in, maar uiteindelijk konden we in 1996 aan de slag. Dit geeft ons nu het grote bestand waardoor we zoveel studies kunnen doen met betrekking tot de klinische uitkomst na echoscopische diagnostiek, waarvan een deel in dit proefschrift is beschreven. Niet in de laatste plaats zorgde je directe bereikbaarheid voor snel overleg over de geschreven stukken en daarmee voor een gestage voortgang van dit project.

De prenatale diagnostiek van aangeboren afwijkingen vereist regelmatig overleg met de kindergeneeskunde. Het was daardoor niet meer dan logisch dat Dick Tibboel mijn tweede promotor zou zijn. Door het wekelijkse perinataal overleg en de samenwerking in publicaties hebben we in de loop der jaren steeds meer begrip gekregen van en voor elkaars patiënten populatie. Beste Dick, je hebt me veel kunnen uitleggen over de ziekteprocessen bij neonaten met aangeboren afwijkingen en daarbij een belangrijke bijdrage aan het tot stand komen van dit proefschrift geleverd. Dat de wetenschappelijke samenwerking continueert zegt iets over wederzijdse bereidheid om de foetus als neonat en de neonat als foetus te onderzoeken.

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Nicolette Ursem en Els Grijseels hebben ieder hun eigen bijdrage geleverd en stonden altijd klaar met een gewillig oor, vaak gevolgd door nuttige actie, waarvoor ik hen zeer erkentelijk ben.

De Nederlandse opleiding tot arts heeft in het curriculum gelukkig nog steeds een verplichte wetenschappelijke stage. Dankbaar heb ik gebruik kunnen maken van de inzet van de medisch studenten, Leon van Eijk, Pauline Wijngaard-Boom, Ellen Lammerink, Titi Hatzmann, Nienke Niemeijer en Wing Tong.

Het manuscript werd beoordeeld door professor Bax, professor Bonsel en professor Deprest. Hartelijk dank voor uw tijd. Professor Jan Deprest ben ik tevens erkentelijk voor zijn bijdrage aan het symposium. Meerdere studies zouden niet volledig zijn zonder de medewerking van de ouders van de patiënten, waarvoor ik hen zeer dankbaar ben.

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