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Detection of congenital heart defects in a non-selected population of 42,381 fetuses

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NTNU
Innovation and Creativity
CONTENT

ACKNOWLEDGMENTS 7
LIST OF PAPERS 9
ABBREVIATIONS 10
INTRODUCTION 11

Background 11
History of ultrasound 11
Early medical use of ultrasound 13
Ultrasound in cardiology 14
Ultrasound in obstetrics 15
Doppler ultrasound 18
Power Doppler ultrasound 20
Three- and four-dimensional ultrasound 21
Magnetic Resonance Imaging (MRI) 22

The fetal examination 23

Epidemiology of congenital heart defects 26
Incidence of CHDs 26
Fetal population versus postnatal population 27
Selected versus non-selected populations 27
Major and minor CHDs 28

The fetal heart examination 30

Teaching and training obstetric ultrasound 34

AIMS OF THE STUDIES 38

MATERIAL AND METHODS 40

Populations and study design 40

Methods 46
Classifications of congenital heart defects 46
Ultrasound equipment and computer software 47
3D imaging 47
Statistical analysis 49

RESULTS AND COMMENTS 50

Paper I 50
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Eva Tegnander
LIST OF PAPERS

The present thesis is based on the following papers:


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## Abbreviations

<table>
<thead>
<tr>
<th>Abbreviation</th>
<th>Description</th>
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<tbody>
<tr>
<td>2D</td>
<td>Two-dimensional</td>
</tr>
<tr>
<td>3D</td>
<td>Three-dimensional</td>
</tr>
<tr>
<td>A-mode</td>
<td>Display presentation of echo amplitude versus depth</td>
</tr>
<tr>
<td>ASD</td>
<td>Atrial septal defect</td>
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<tr>
<td>AVSD</td>
<td>Atrioventricular septal defect</td>
</tr>
<tr>
<td>BPD</td>
<td>Biparietal diameter</td>
</tr>
<tr>
<td>B-mode</td>
<td>Brightness mode (gray scale mode), display presentation of a spot brightness for each echo voltage delivered from the receiver</td>
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<tr>
<td>CHD</td>
<td>Congenital heart defect</td>
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<tr>
<td>CNS</td>
<td>Central nervous system</td>
</tr>
<tr>
<td>CRL</td>
<td>Crown-rump length</td>
</tr>
<tr>
<td>CW</td>
<td>Continuous wave</td>
</tr>
<tr>
<td>DIRV</td>
<td>Double inlet right ventricle</td>
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<tr>
<td>DORV</td>
<td>Double outlet right ventricle</td>
</tr>
<tr>
<td>HLHS</td>
<td>Hypoplastic left heart syndrome</td>
</tr>
<tr>
<td>IUFD</td>
<td>Intrauterine fetal death</td>
</tr>
<tr>
<td>IQ data</td>
<td>Complex demodulated radio frequency data</td>
</tr>
<tr>
<td>MHz</td>
<td>Megahertz</td>
</tr>
<tr>
<td>MRI</td>
<td>Magnetic resonance imaging</td>
</tr>
<tr>
<td>STIC</td>
<td>Spatio-temporal image correlation</td>
</tr>
<tr>
<td>TDOG</td>
<td>Tissue Doppler gated</td>
</tr>
<tr>
<td>TGA</td>
<td>Transposition of the great arteries</td>
</tr>
<tr>
<td>TOF</td>
<td>Tetralogy of Fallot</td>
</tr>
<tr>
<td>TOP</td>
<td>Termination of pregnancy</td>
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<tr>
<td>VSD</td>
<td>Ventricular septal defect</td>
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INTRODUCTION

Background

The aim of prenatal care has always been to prevent and alleviate maternal and fetal/newborn complications affecting the pregnancy, birth and the neonatal period. Traditionally, midwives and general practitioners have provided the basic prenatal care in Norway. Selection of high-risk pregnancies for referral to obstetric specialists has been an important part of the pregnancy care.

In 1982, a study analyzing stillbirths and neonatal deaths from five counties in Norway found that 30% of the deaths might have been avoided with provision of optimal care (Larssen et al. 1982). As a consequence, the Directorate of Health in Norway ordered guidelines to be developed in 1984 to ensure equal and complete prenatal care for all pregnant women in Norway (Norges offentlige utredninger, NOU 1984:17). The document outlined the organization of the pregnancy care in the future, but gave little attention to the potential of diagnostic ultrasound for future prenatal diagnosis and treatment. This laid the ground for an intense debate among obstetricians concerning the use of ultrasound in pregnancy in Norway in the following years. In 1986, a consensus conference concerning the use of ultrasound in pregnancy was organized. An implementation of one ultrasound examination at 17–18 weeks of gestation for all pregnant women in Norway was recommended (Backe and Buhaug 1986). Since then, the use of obstetric ultrasound has continuously expanded in Norway. It now represents the method of choice for diagnosis and treatment in pregnancy, and addresses the problems concerning the 30% deaths that might have been avoided with improved prenatal care (Larssen et al. 1982, Holt et al. 2000).

History of ultrasound

The Italian biologist Lazzaro Spallanzani demonstrated already in 1794 the ability of bats to navigate with the aid of echo reflections from high frequency inaudible sound. The scientific work towards defining the fundamental physics of sound vibrations,
transmission, propagation and refraction then followed in the centuries thereafter. This included the thesis by Lord Rayleigh from England in 1877, “The Theory of Sound”; this was the first description of a sound wave as a mathematical equation, and the foundation for future theoretical work in acoustics. For his work, Lord Rayleigh received the Nobel Prize in physics in 1904 (Woo 2005). Sir Francis Galton was able to generate high frequency sound waves in 1876 using the so-called “Galton Whistle” which he developed (Figure 1).

The real breakthrough for high frequency echo-sounding techniques came in 1880 when Pierre Curie and his brother Jacques Curie discovered the piezo-electric effect in certain crystals (Curie and Curie 1880) (Figure 2).

The Titanic tragedy in 1912 prompted research to detect underwater objects such as icebergs. The first working sonar system was designed in 1914, but was not particularly successful. World War I gave rise to new studies, as it became important to detect other underwater objects such as submarines. Among others, the French physicist Paul Langévin and the Russian scientist Constantin Chilowsky developed the hydrophone, a powerful high frequency ultrasonic echo sounding device involved in the sinking of a German submarine in the Atlantic in 1916 during World War I. Langévin’s hydrophones formed the basis of the future development of naval pulse-echo SONAR (Sound Navigation and Ranging) in the following years. Later the RADAR (Radio Detection and Ranging) system was developed, using electromagnetic waves rather than ultrasonic waves (Woo 2005).
The next important step was the development of pulse-echo ultrasonic metal flaw detectors in the 1930s (Figure 3). They were primarily constructed to check on the integrity of metal hulls of large ships and the armour plates of battle tanks (Figure 4). Together with the sonar and radar systems, the metal flaw detectors were precursors for the establishment of medical ultrasonic equipment (Woo 2005).

**Figure 3** Metal flaw detector. Note, camera supported by a wooden log! (Edler 1961).

**Figure 4** Interruptions in the passage of sound waves are used to locate and measure flaws and cracks in solid metal (Woo 2005).

**Early medical use of ultrasound**

Ultrasound is a relatively new modality within medicine and was initially used for therapy rather than diagnosis. In 1938, Raimar Pohlman, in Erlangen, Germany, demonstrated the therapeutic effects of ultrasonic waves in human tissue and introduced ultrasonic physiotherapy as a medical practice (Figures 5-7).

**Figure 5** Ultrasonic physiotherapy (Woo 2005).

**Figure 6** Ultrasonic therapy generator, the “Medi-Sonar” in the 1950s.
William Fry, University of Illinois, and Russell Meyers, University of Iowa, used ultrasound to destroy parts of the basal ganglia during craniotomies in patients with Parkinsonism. During the 1940s, ultrasound was regarded as a “cure-all” remedy, despite the lack of scientific evidence. The question of possible harmful tissue damaging started to be raised, and consequently curtailed the development of diagnostic ultrasound for a time (Woo 2005).

Figure 7 Treatment of ulcer (Woo 2005).

Early pioneers within diagnostic ultrasound were the Austrian neurologist Karl Theodore Dussik and his brother Friederich who were the first to use ultrasound for diagnostic purposes. They studied sound attenuation in brain tissue in the early 1940s, but their method was later described as worthless. George Ludwig, a physician from USA, investigated the use of ultrasound for the detection of gallstones (outside the human body) in the late 1940s (Ludwig and Struthers 1950). John Wild and John M. Reid built the first handheld B-mode scanner for clinical use in 1951 (Baker 2005, Woo 2005).

Ultrasound in cardiology
The development of ultrasound technology to be used in the field of cardiology started as a cooperation between cardiologist Inge Edler and physicist Carl Hellmuth Hertz in Lund, Sweden, in 1954 (Figure 8). In 2004 the 50th anniversary of this important scientific work in cardiology was celebrated (Olsson et al. 2004). This work has developed extensively and completely changed the diagnostic tools for adult cardiology. It is also the foundation for today’s fetal cardiology. Edler and Hertz utilized the metal flaw detectors as a basis for their research (Edler 1961). The need for improved diagnosis before heart surgery led Edler to look for non-invasive alternatives rather than cardiac catheterization and contrast x-rays of the heart that failed to give enough preoperative information, for example, about the mitral valve (Nilsson and Westling...
2004). For the first time in October 29, 1953, Inge Edler and Carl Hellmuth Hertz managed to record pictures from the heart (Edler and Hertz 1954). With a single sound beam aimed in a fixed direction through the heart, the various echoes in this sound beam could be recorded as a function of time, the so-called M-mode (motion) registration. Later on, Edler and his colleagues realized that some of the echoes came from the anterior leaflet of the mitral valve and the characteristic motion pattern for this valve was soon established (Figure 9). Edler and Hertz’s work made a decisive contribution to diagnostics in cardiology, neurosurgery and obstetrics and gynecology (Nilsson and Westling 2004).

Ultrasound in obstetrics

Ian Donald from Scotland was the first to introduce diagnostic ultrasound to obstetrics and gynecology. His 1958 publication in the Lancet, “Investigation of abdominal masses by pulsed ultrasound”, was the most important paper published on medical diagnostic ultrasound at that time and boosted the later clinical development (Donald et al. 1958). Ian Donald and his engineering group developed the Diasonograph produced by Nuclear Enterprise (Figure 10). He was the first to perform fetal head measurements and related them to age and weight (Willocks et al. 1964, Willocks et al. 1967), and the first to report sonographic evidence of a gestational sac (MacVicar and Donald 1963). Ian Donald also utilized ultrasound to study ovarian cysts, polyhydramnios and molar pregnancy (Donald et al. 1958, Donald 1962).
Inspired by Ian Donald, Bertil Sundén, at the University of Lund, took an early interest in obstetric and gynecological ultrasound and safety. Upon the publication of Ian Donald’s work on the echoscope generating a two-dimensional display, Sundén was sent to visit and work in Donald’s department in Glasgow for three weeks. As a consequence, Nuclear Enterprise sold their very first Diasonograph to the department of Obstetrics and Gynecology in Lund. Bertil Sundéns book titled “On the diagnostic value of ultrasound in obstetrics and gynaecology” from 1964 was the first thesis addressing the use of ultrasonography in obstetrics and gynecology (Sundén 1964). His thesis represented pioneer work with numerous first-time observations including an anencephalic fetus diagnosed at 31 weeks of gestation.

The pioneers in the development of transvaginal transducers were Hisaya Takeuchi from Japan and Alfred Kratochwil from Austria. Already in the late 1960s they reported on fetal heart pulsation in the early first trimester (Kratochwil and Eisenhut 1967, Takeuchi et al. 1968). Working with KretzTehnik AG in Austria, Kratochwil soon became an active user of the technique and from 1968 developed training courses in ultrasound (Woo 2005).

Until the mid-1970s, A- and B-mode scanners were used. In 1968, Stuart Campbell, in London, published his now classical paper “An improved method of fetal cephalometry by ultrasound” which immediately became a highly recognized work in which both A- and B-mode scans were used to measure the fetal biparietal diameter, today a standard measurement in fetal biometry (Campbell 1968). Utilizing a static B-scan, Campbell was the first to report on prenatal detection of an anencephalic fetus (Campbell et al.)
1972) and a fetus with spina bifida (Campbell 1977), resulting in a termination of pregnancy. In 1975, Hugh Robinson, in Glasgow, reported on the measurement of the fetal crown-rump length; his measurement curves are still in use today (Robinson 1973). By combining A- and B-mode ultrasound, Robinson successfully demonstrated fetal cardiac action from 7 weeks onwards (Robinson and Shaw-Dunn 1973). The advent of the analog scan converter in the early 1970s and the digital scan converter in the later 1970s were among the most important innovations in ultrasound imaging (Woo 2005).

The first mechanical real-time scanner was developed in Germany by Walter Krause and Richard Soldner in the 1960s. They used a rotating transducer and a parabolic mirror, sending the sound waves parallel into the tissue (Figure 11). A wide picture in real-time sequences of more than 10 frames per second could be obtained. This technique completely changed the practice of ultrasound scanning.

![Figure 11](image1.png) Videoson Siemens Medical Systems 1965, and its working mechanism.

Years of further development ensued. In 1973, one of the earliest commercial models of a linear-array real-time scanner was designed and produced by Martin H. Wilcox, the founder and engineer at the Advanced Diagnostic Research Corporation (Figure 12). In 1986, these electronic real-time scanners started to replace the static scanners.

![Figure 12](image2.png) The ADR real-time scanner.
Doppler ultrasound

The Doppler effect was first described by the Austrian physicist Christian Johann Doppler (1803–1853) in 1842. Echoes returned from moving objects have frequencies different from the frequencies sent into the body. This is called the Doppler effect. The change in frequency is called the Doppler shift, which might be used to detect and measure tissue motion, the presence or absence of blood flow and its direction, speed and character.

Figure 13 Satomura 1959.
Of importance for the use of ultrasound technology in the diagnosis of heart defects was the development of Doppler ultrasound technology (Satomura 1957, Satomura 1959 (Figure 13), Baker et al. 1964, Peronneau et al. 1969).

In Trondheim, research on the Doppler technique started in 1973. One instrument, PEDOF, Vingmed, (Figure 14) combined the use of continuous wave ultrasound for the measurement of high velocities, such as in a stenosis, with the pulsed wave ultrasound for blood flow measurements at a pre-selected depth. This laid the foundation for the future development of ultrasound in cardiology (Brubakk et al. 1977, Hatle et al. 1978). Later, the imaging ultrasound technique and Doppler ultrasound technique were combined in the same instrument, thereby making color-flow mapping and detailed registration of velocities in selected vessels possible (Hatle et al. 1980, Hatle et al. 1981).

In 1977, FitzGerald and Drumm managed to register crude Doppler ultrasound signals from the human fetus in utero (Fitzgerald and Drumm 1977). A year later McCallum et al. (1978) presented various types of waveforms from the umbilical cord in utero. In 1979, Gill and Kossoff managed to quantify the blood flow in the umbilical vein (Gill and Kossoff 1979). Their system, however, did not allow measurements of the high velocities present in fetal arteries. In 1978, Eik-Nes et al. started the work that culminated in the first prototype of a Duplex transducer combining a linear array transducer and a pulsed Doppler transducer for the qualitative and quantitative measurement blood flow in fetal arteries as well as veins (Eik-Nes et al. 1980, Eik-Nes et al. 1982, Kirkinen et al. 1983, Eik-Nes et al. 1984b, Maršál et al. 1984).
The use of Doppler ultrasound in obstetrics emerged late in the 1970s. Further development led to 2D color flow imaging (Angelsen et al. 1989). With rapid interfacing of 2D and Doppler ultrasound, color-coded presentations of Doppler information superimposed on gray-scale anatomic images were possible. Appropriate colors were assigned to the display pixels depending on whether the motion was toward or away from the transducer.

Color Doppler ultrasound is considered to be a second-line investigation for cardiac evaluation and completes the evaluation of the gray-scale information (Chaoui 2001). It allows for ascertainment of homogenous flow within the heart in the great vessels to rule out heart defects and a more detailed diagnosis once heart disease is suspected. Color Doppler may also highlight areas of turbulence or of important atrioventricular valve regurgitation and for assessment of valve opening and of symmetry in ventricular filling in cases where visualization using the gray scale alone is difficult (Chaoui 2001, Gardiner 2001).

**Power Doppler ultrasound**

Power Doppler displays present two-dimensional Doppler information by color-encoding the strength of the Doppler shift. This method is free of aliasing, is not angle dependent, and is more sensitive than pulsed Doppler ultrasound to slow flow and flow in small or deep vessels (Kremkau 2006). On the other hand, power Doppler does not give information about direction, speed and flow character such as turbulence.

Power Doppler ultrasound was introduced in the early 1990s and has opened up the possibility of displaying flow independently of its velocity or direction. The technique is useful when applied to fetal cardiology; it facilitates the detection of small VSDs and enables spatial orientation of the great arteries (Chaoui 2001).
Three- and four-dimensional ultrasound

Three-dimensional ultrasound is a logical development of 2D imaging because the anatomy in reality is three-dimensional. Already in 1977, Brinkley et al. described a technique to generate a three-dimensional reconstruction of an organ, later applied to fetuses for weight estimation (Brinkley et al. 1982, Eik-Nes and Brinkley 1983). The technique described by Brinkley made possible crude 3D still-images of the fetal heart.

The heart is a moving object. In adult cardiology it is possible to synchronize (gate) the imaging of the heart movements in relation to the echocardiogram. Since this is not possible, or very difficult, in the fetus, other methods to cope with the temporal information had to be found. These methods are called “gating”. In 1996, Nelson et al. reported on possibilities examining the fetal heart by 3D and 4D ultrasound using a fast Fourier transform method, similar to what we today know as STIC. The same year, Deng et al. (1996) used real-time directed M-mode to gate the fetal heart rate and acquire and display 4D volume data sets of the fetal heart. The fetal heart rate acquired by Doppler ultrasonography (Deng et al. 2000, Meyer-Wittkopf et al. 2001, Herberg et al. 2005) or by cardiotocography (Herberg et al. 2003) has also been used to gate the spatial and temporal information. These described methods use a post processing technique. It would be of interest to find a technique that makes it possible to gate the images during the acquisition of data. In the future, it will be possible to obtain this by tissue Doppler gated technique and a 2D-array system.

The reported advantages of 3D over conventional 2D imaging are reduced scanning time, and reduced operator dependency and window dependency since the volumetric data are acquired within a few seconds from a single window. Another advantage is that access to volume data may provide information not obtainable with 2D ultrasound. In 3D ultrasound a volume of ultrasonographic data is acquired and stored, allowing the subsequent extraction of an infinite number of scan planes (Budorick and Millman 2000, Downey et al. 2000).
4D visualization of the fetal heart became a reality with the incorporation of STIC
(spatio-temporal imaging correlation) algorithms into commercially available
equipment (Gonçalves et al. 2005). The processing using STIC is done retrospectively.

Magnetic Resonance Imaging (MRI)

MRI has developed to become a powerful approach to imaging CHD in infants and
children (Chung 2000). The potential of MR for imaging the fetus was described in
1983 by Smith et al. (1983). MRI has proven to be a valuable complement to
ultrasonography when additional information is needed about fetal anatomy (Levine et
al. 1999a). Traditionally, MRI has been used to evaluate the maternal pelvic anatomy,
but the fetal anatomy has been difficult to image because of fetal motion during the
relatively long acquisition times (Weinreb et al. 1985, Powell et al. 1988). The
development of fast MRI techniques has revolutionized the ability to image both the
pregnant woman and the fetus (Levine et al. 1996).

Ultrasound evaluation of fetal anatomy may be limited by the nonspecific appearance of
some anomalies, by artifacts that reduce the image resolution and by advanced
gestational age. MRI uses no ionizing radiation, provides excellent soft tissue contrast
and has multiple planes for reconstruction and a large field of view. Thus, it is the
method of choice when additional information is needed following ultrasound diagnosis
(Levine 2001). In such cases MRI may give valuable information to add for the
counseling process (Levine et al. 1999a, Knox et al. 2005). MRI has proven to be
particularly beneficial in the evaluation of the fetal CNS system (Blaicher et al. 2005),
especially in ventriculomegaly and imaging of the corpus callosum (Levine et al.
1999b) and when therapeutic consequences might be expected, such as diaphragmatic
hernia, tumors (Knox et al. 2005) and fetal abdominal masses (Angtuaco et al. 1992).
Because of signal-to-noise limitations, small fetal structures may be difficult to image
by MRI (Blaicher et al. 2005), thus the fetal heart may be difficult to visualize and the
interventricular septum and the atroventricular valves may only occasionally be seen
(Trop and Levine 2001). The claustrophobic factor for those inside the magnet and the
vena cava syndrome often occurring in the third trimester of the pregnancy may be
factors that need to be addressed when imaging the fetus (Levine et al. 1999a). Increased experience with fast MRI techniques and 'real-time' cardiac MRI will most likely identify more conditions where this technique may add to the prenatal diagnosis (Fogel et al. 2005).

The fetal examination

Ultrasound technique made it possible to get information directly from the fetus in a non-invasive manner. As the development of this technique and equipment continued, the use of ultrasound in pregnancy rapidly expanded. It became obvious that ultrasound technology could reveal conditions not detectable by traditional ways of collecting information from the pregnant woman and the fetus.

The first screening program for systematic use of ultrasound in pregnancy was initiated in Malmö, Sweden, in 1974, starting with one ultrasound examination at 28 weeks of gestation. The primary intention was to detect twins. In the following years, the examination was successively moved to 17 weeks. From 1976 a second routine examination was included at 32 gestational weeks (Grennert et al. 1978). In 1980, Germany was the first country to officially introduce the two-stage screening program, adopted from Malmö, to their total pregnant population (Mutterschafts-Richtlinien 1980). In Norway, a consensus conference concerning the use of ultrasound in pregnancy was held in Oslo in 1986. The consensus panel recommended the offer of one ultrasound examination at approximately 18 weeks to all pregnant women (Backe and Buhaug 1986). The recommendation was based on the reported decrease in post-term pregnancies following introduction of routine ultrasound (Eik-Nes et al. 1984a) and the need to organize the already extensive use of ultrasound in pregnancy (Eik-Nes 1986), later confirmed by Backe (1994). In the years that followed, routine ultrasound scanning was officially introduced by the health authorities in a number of European countries. Iceland followed the Norwegian one-stage model in 1987 (Geirsson 1987), and Austria followed the German two-stage model in 1988. In 1995, Germany expanded their official offer with an early scan at 10 weeks (Änderung der
Mutterschafts-Richtlinien 1995), and in 1996, Switzerland introduced an early scan at 10 weeks followed by an 18 week scan. Although not all countries have introduced the practice officially, most countries offer ultrasound examinations to their pregnant population today (SBU-rapport nr. 139, 1998).

Randomized controlled trials were carried out during the 1980s with the purpose of showing any benefit of using ultrasound in every pregnancy (Bennet et al. 1982, Eik-Nes et al. 1984a, Bakketeig et al. 1984, Neilson et al. 1984, Waldenström et al. 1988, Saari-Kemppainen et al. 1990, Eik-Nes et al. 2000). One problem with such studies is that the window for carrying them out is narrow. A study evaluating a new technique too early, before it is fully developed, may not show the full benefit of the technique studied. If it is done too late, the technique may already be in use and the information needed for comparison is no longer available (Eik-Nes 1993a, Salvesen and Eik-Nes 1994). The early findings in the controlled trials included a more precise estimate of the gestational age, resulting in a 70% reduced frequency of induction for overdue pregnancies (Eik-Nes et al. 1984a, Waldenström et al. 1988, Neilson 2004) and a reduction in perinatal mortality mainly due to improved early detection of major anomalies (Saari-Kemppainen et al. 1990).

Other studies evaluated the effectiveness of the routine examination to detect fetal developmental disorders (Chitty et al. 1991, Crane et al. 1994, Papp et al. 1995, Grandjean et al. 1999, Vial et al. 2001, Garne et al. 2005, Dolk 2005). Due to differences in use of ultrasound in pregnancy and in experience of the operators, the results varied between 35–74%. Once detected, various conditions could be followed and information derived from prenatal observations and follow-up through delivery and into the neonatal period. With such information, follow-up of the fetus and child has been optimized to improve perinatal mortality and morbidity (Bonnet et al. 1999, Brantberg et al. 2004).

Traditionally, pregnant women at increased risk of having a fetus with congenital heart defect have been referred to specialist for an additional ultrasound examination (Allan
et al. 1986a). In the 1980s, pediatric cardiologists clearly saw the potential of ultrasound techniques such as M-mode, two-dimensional and Doppler echocardiography, techniques utilized for the fetal heart examination in high-risk pregnancies (Kleinman et al. 1980, Huhta 1986). In 1985, a French study first initiated the idea of prenatal cardiac screening (Fermont et al. abstract 1985), closely followed by Allan et al. (1986b). They suggested that 2/1000 major CHDs could be detected by including the four-chamber view into the second trimester fetal examination. It was soon realized that fetal heart defects were more difficult to detect than major defects in other fetal organs, particularly in non-selected populations (Brocks and Bang 1991, Crane et al. 1994, Rustico et al. 1995, Tegnander et al. 1995, Buskens et al. 1996, Bull 1999, Wong et al. 2003).

With the improvement in ultrasound technology, more and more details of the fetal heart could be imaged at an earlier gestation (Achiron et al. 1994, Allan et al. 1997). Data also showed that in the group of fetuses at increased risk for CHD, only a small percentage of the total CHD-population was found (Allan et al. 1994, Maher et al. 1994, Carvalho et al. 2002, Åmark et al. 2004). Most fetuses with CHD were born from mothers at no increased risk at all, thus the second trimester fetal examination became important for the prenatal detection of CHDs.

The poor detection rates of major CHDs from non-selected populations needed to be addressed. The equipment used, the gestational age at the time of the fetal examination and details obtained from the fetal heart were factors that had to be evaluated. The ultrasound machines used for second trimester screening purposes were often of poorer quality than the equipment used at referral centers. This was a paradox, since in order to make a referral for second opinion, the CHD had to be suspected when the routine scan was made; thus, good quality machines should be available in routine ultrasound settings. But, even with an ultrasound with poor quality, it is necessary to be able to use the potential of the machine to create optimized ultrasound images in order to detect developmental disorders in general. In addition, the gestational age seems important in imaging anatomical structures. Several studies have shown that examination of the fetal
heart prior to 18 gestational weeks is too early to obtain details from the heart; thus, the fetal heart examination should be done after 18 weeks, preferably after 20 weeks (Schwärzler et al. 1999b, Allan 2003). The various ways of organizing the use of ultrasound in pregnancy and the upper limit for termination of the pregnancy are decisive factors for setting the optimal time for the second trimester fetal examination. During the last years, the 11–13+6 week ultrasound examination has been introduced (Nicolaides et al. 1992). Increased nuchal translucency has proven to be a marker for chromosome aberrations, congenital heart defects and other defects such as diaphragmatic hernia and genetic syndromes (Nicolaides et al. 1992, Hyett et al. 1997, Hafner et al. 1998a, Schwärzler et al. 1999a, Souka et al. 2001, Haak et al. 2002, Hyett 2002). Due to the fact that the fetal heart is tiny at that time of gestation, that an increased nuchal translucency may not be present in all CHDs, and because of the progression of CHDs during the time of the pregnancy, some heart defects may not be detected at such an early gestation (Maeno et al. 1999, Gardiner 2001, Mavrides et al. 2001, Trines and Hornberger 2004).

**Epidemiology of congenital heart defects**

Congenital heart defect is one of the most common serious congenital anomalies found in live births with a reported incidence of 8/1000 live births (Mitchell et al. 1971). The use of ultrasound in non-selected populations has resulted in new information about the incidence, range and natural history of congenital heart defects seen prenatally (Allan 1989).

**Incidence of CHDs**

Traditionally, the incidence of CHDs in large non-selected populations has been reported to be 8/1000 live births, of these, 4/1000 as major CHDs and 4/1000 as minor CHDs. A large variety of incidences are found from published data, between 3–12/1000 live births (Carlgren 1969, Mitchell et al. 1971, Hoffman and Christianson 1978, Dickinson et al. 1981, Ferencz et al. 1985, Meberg et al. 1990). The lower incidences
were reported before there were well-trained pediatric cardiologists and before cardiac surgery required a detailed preoperative diagnosis (Hoffman 1995a). The introduction of the ultrasound technique, including Doppler ultrasound and Color Flow Mapping, has made it possible to diagnose minor CHDs and even asymptomatic CHDs postnatally (Martin et al. 1989, Meberg et al. 1994), and has thus resulted in an increase in the reported incidence of CHDs.

Fetal population versus postnatal population

Most studies from large populations evaluating the incidence of CHD have been based on postnatal populations (Mitchell et al. 1971, Dickinson et al. 1981, Ferencz et al. 1985, Samánek et al. 1989). In postnatal populations, the ascertainment of cases might be biased due to asymptomatic CHDs that are missed at the ordinary clinical exam (mild pulmonary stenosis, small ASDs or VSDs), incorrectly diagnosed cause of death, deaths prior to recognized CHD and intrauterine deaths and postnatal deaths without autopsy (Abu-Harb et al. 1994, Hoffman 1995a).

With the introduction of prenatal ultrasound, the natural history of CHDs could be studied. Some forms of CHD were seen more frequently prenatally than postnatally; some conditions, like HLHS, AVSD and critical aortic stenosis, were often presented with intrauterine heart failure, and a particularly high rate of chromosome aberrations was found among fetuses with CHD resulting in a high rate of intrauterine fetal deaths (Allan 1989). A high incidence of CHD and chromosome aberrations among spontaneously aborted fetuses has also been found (Gerlis 1985, Chinn et al. 1989). This suggests that the true incidence of CHD in early fetal life may be higher than what our knowledge from born populations would lead us to believe.

Selected versus non-selected populations

It has been a well-established routine to offer one extra ultrasound examination at 20–24 weeks of gestation when there is an increased risk for CHD (Callan et al. 1991, Allan et al. 1994, Oberhänsli et al. 2000). Maternal factors that might increase the risk are
family history of CHD; previous child affected; parental CHD; maternal diabetes; exposure to known cardiac teratogens (such as alcohol, amphetamine, anticonvulsants), intrauterine infections (rubella, parvovirus, coxackievirus, cytomegalovirus) or high doses of ionizing radiation. Fetal factors are findings of nonimmune fetal hydrops, extra cardiac anomalies or arrhythmia (Allan 1989, Allan et al. 1994, Gembruch and Geipel 2003). Because of the increased risk of CHD among these fetuses, the ultrasound examinations have been done by obstetric ultrasound specialists or pediatric cardiologists utilizing advanced ultrasound equipment. The detection rates of CHDs from selected populations have thus been reported as high as 92% (Copel et al. 1987, Stewart et al. 1987). Despite the high detection rates, the CHDs detected constituted only a small percentage of the total CHD population, as most fetuses with CHD are found in non-selected populations (Allan et al. 1994, Maher et al. 1994, Carvalho et al. 2002, Åmark et al. 2004).

As routine ultrasound became an integrated part of prenatal care, an increasing number of women at no risk for disease were being referred to tertiary centers because of abnormal fetal hearts seen at ultrasound examinations (Allan et al. 1994, Sharland 2004). With new knowledge about where to find the CHDs, the focus changed towards fetal routine ultrasound in non-selected populations. The ultrasound operators were not as skilled, and the ultrasound machines not as advanced among those working with non-selected populations as among those working with selected populations at tertiary centers. In addition, the low incidence of malformations in non-selected populations made defects harder to find. CHDs seemed particularly difficult to detect, as the reported detection rates were lower than expected, and lower than the results from selected populations (Crane et al. 1994, Buskens et al. 1996, Bull 1999). Thus, investigating ways to increase the detection rate of CHDs in non-selected populations became important.

Major and minor CHDs

There does not seem to be a universally accepted definition of major CHD (Allan et al. 2001). As a result, studies define major CHDs in various ways depending on the study
design, the population defined and the experience of the authors (Ferencz et al. 1990, Wren et al. 2000, Allan et al. 2001). These differences are reflected in the reported incidences of CHDs. Most investigators define major CHDs as defined by Mitchell et al. (1971) as a gross structural abnormality of the heart that is actually or potentially of functional significance (Hoffman 1995b, Allan et al. 2001).

For many years, an incidence of CHD of 8/1000 live births has been accepted in non-selected populations (Achiron et al. 1992, Boldt et al. 2002, Wong et al. 2003), whereas 4/1000 as major CHDs. New data show that the incidence of major CHDs has remained unchanged (Carvalho et al. 2002). One may conclude that the majority of the major CHDs have always been recognized, thus an incidence of approximately 4/1000 reflects proper ascertainment of the major CHD cases.

The incidence of minor CHDs, on the other hand, has steadily increased during the last years (Martin et al. 1989, Meberg et al. 1994, Roguin et al. 1995). Better postnatal clinical diagnosis and technique utilizing diagnostic ultrasound have resulted in an increased detection of minor CHDs previously not diagnosed. These are defects with little long-term consequences for the child and a vast majority close spontaneously within the first year after birth (Evans et al. 1960, Du et al. 1998, Helgason 1999, Meberg et al. 2000). There is reason to believe that the closure of these small defects towards the end of the pregnancy and during the first months of life may be a delay of a normal process, thus it might be questioned whether these defects are true congenital heart defects (Ben-Shachar et al. 1985, Hiraishi et al. 1992). As long as they are defined as CHDs, however, they will affect the incidence of CHDs in the population.

Obviously we have to reconsider the incidence of CHDs and the spectrum of CHDs as new data emerge.
The fetal heart examination

Real-time 2D ultrasound still is the best ultrasound technique for structural evaluation of the fetal heart. The technical development has resulted in ultrasound machines with advanced features such as high image resolution and high frame rate, making it possible to obtain a detailed image of the tiny structures in the moving fetal heart (Chaoui 2001). For routine use in the second trimester, the ultrasound examinations are most often performed transabdominally. As the attenuation factor sometimes limits imaging depth (Kremkau 2006), especially in obese women, alternative routes for examination may be preferred. Kratochwil and Eisenhut (1967), Takeuchi et al. (1968) and Jouppila (1971) were the first to propose the transvaginal route to identify fetal heart pulsations. With real-time transvaginal transducers, the fetal heart could be imaged in detail, resulting in the first publication, in 1990, of a CHD detected at 11 weeks of gestation (Gembruch et al. 1990).

The first studies based on real-time images of the fetal heart were published in 1980 by Lindsey Allan et al. (1980), Kleinman et al. (1980) and Lange et al. (1980). Lindsey Allan’s work from the early 1980s laid the basis for subsequent studies showing that fetal cardiac anomalies could be detected by real-time ultrasound in the second trimester of the pregnancy (Allan et al. 1986).

The fetal heart appears somewhat different than an adult heart. As a consequence of the normal large liver in fetal life, the heart is positioned more horizontally and the apex is displaced more cranially than in the adult. This results in a slightly different orientation of the heart with the right side positioned anteriorly in the fetal chest and the left side posteriorly. The two ventricles are equal in size in the fetus, and there is normally a patent foramen ovale between the two atria and a patent ductus arteriosus between the main pulmonary artery and the aortic isthmus. Fetal lungs are filled with water, which makes the heart visualized in planes not obtainable after birth (Allan et al. 1980, Allan 1986).
Since most fetuses with CHD are found in non-selected populations, the ultrasound examination for screening purposes is the only opportunity to detect the majority of CHDs. For this reason, an easy way of examining the fetal heart during the routine examination was introduced in the mid 1980s (Fermont et al. 1985, Allan et al. 1986b). Among the eight scan planes Allan et al. (1980) investigated, the four-chamber view was one of the easiest views to obtain, thus the four-chamber view was the plane chosen to be obtained in every fetus at the routine scan.

Table 1 Sonographic details from the four-chamber view.

<table>
<thead>
<tr>
<th>Details obtained</th>
</tr>
</thead>
<tbody>
<tr>
<td>Transverse abdomen</td>
</tr>
<tr>
<td>Abdominal aorta and inferior vena cava</td>
</tr>
<tr>
<td>Stomach is on the left side of the abdomen</td>
</tr>
<tr>
<td>Aorta is on the left side of the spine</td>
</tr>
<tr>
<td>Inferior vena cava is on the right side of the spine, ventral to the aorta</td>
</tr>
<tr>
<td>The atrioventricular junction (The four-chamber view)</td>
</tr>
<tr>
<td>Position</td>
</tr>
<tr>
<td>Apical view</td>
</tr>
<tr>
<td>Subcostal view</td>
</tr>
<tr>
<td>The heart occupies 1/3 of the thoracic area</td>
</tr>
<tr>
<td>The position is in the middle of the thorax with the apex pointing towards left</td>
</tr>
<tr>
<td>The rhythm is regular between 120–160 beats/minute. Short periods of bradycardia are normal in the second trimester of the pregnancy.</td>
</tr>
<tr>
<td>There is equal contractility on the left and the right side of the heart</td>
</tr>
<tr>
<td>There are two atria of equal size</td>
</tr>
<tr>
<td>There are two ventricles of equal size</td>
</tr>
<tr>
<td>The left ventricle forms the apex</td>
</tr>
<tr>
<td>The moderator band is found in the right ventricle</td>
</tr>
<tr>
<td>There are two atrioventricular valves</td>
</tr>
<tr>
<td>The tricuspid valve is closer to the apex than the mitral valve</td>
</tr>
<tr>
<td>The two septa and the atrioventricular valves form an “offset cross” in the middle of the heart</td>
</tr>
<tr>
<td>There is an intact ventricle septum</td>
</tr>
<tr>
<td>There is a patent foramen ovale in the atrial septum</td>
</tr>
<tr>
<td>Two pulmonary veins connect to the left atrium</td>
</tr>
</tbody>
</table>

The four-chamber view seemed to be an easy view to obtain, as the reported success rates in viewing the four chambers in all fetuses were more than 95% (Copel et al. 1987, Achiron et al. 1992, Todros et al. 1997). Despite the easy access to this view, it
seemed to increase the detection rate of major CHDs to only a small degree (Achiron et al. 1992, Ott 1995, Tegnander et al. 1995, Todros et al. 1997).

The importance of evaluating all the details in the four-chamber view, as shown in Table 1, has been emphasized by several authors (Allan 1986, Chaoui 2003, Sharland 2004).

To be able to detect CHDs with a normal four-chamber view the great arteries need to be included into the fetal heart examination. Educational sites and organizations recommend examination of both the four-chamber view and the great arteries when the fetal heart is evaluated. Details to evaluate when the great arteries are obtained are shown in Table 2. In 1997, Yoo et al. introduced the three-vessel view as an easy way to examine the great arteries (Table 2). The three-vessel view has proven to be an important view for detection of defects affecting the outflow tracts and the great arteries (Yoo et al. 1999, Gardiner 2001).

**Table 2** Sonographic details from the great arteries.

<table>
<thead>
<tr>
<th>Details obtained</th>
<th>The ventriculo-arterial junction</th>
<th>Ascending aorta</th>
<th>Pulmonary artery</th>
<th>Three-vessel view</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>The ascending aorta arises from the left ventricle</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>There is continuity between the anterior wall of the ascending aorta and between the ventricle septum and the posterior wall and the mitral valve</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>The main pulmonary artery arises from the right ventricle and branches into the right pulmonary artery and the ductus arteriosus</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>The main pulmonary artery crosses over with the ascending aorta</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>The two arteries are approximately equal in diameter (the main pulmonary artery slightly larger)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Three-vessel view</td>
<td>The main pulmonary artery is seen in a longitudinal view from the right ventricle to the ductus arteriosus</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>The ascending aorta is seen in a transverse view on the right side of the main pulmonary artery</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>The superior vena cava is seen in a transverse view on the right side of the ascending aorta</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>The vessels are arranged in a straight line and sized in decreasing order</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
When the great arteries were introduced, the heart examination also included the aortic arch and the ductal arch (Bromley et al. 1992). Details of importance to notice when the arches are examined are shown in Table 3. Later, a five-step approach to the examinations of the fetal heart was introduced including only transverse views from the fetus (Yagel et al. 2001). The authors concluded that a proper evaluation of all the five views, also shown in Tables 1 and 2, make the view of the arches unnecessary. As the arches might be difficult to obtain due to the fetal position, the three-vessel view has been recommended as an optional view (Chaoui 2003a). Prenatal detection of ductal dependent CHDs may also improve from the effort of obtaining the three-vessel view (Viñals et al. 2002).

In a screening situation, the time allocated for the complete scan is limited. With an awkward fetal position or fetal movements, planes expected to be identified during the scan may not be obtainable. A higher detection rate of cardiac structures may be achieved if unlimited time is given for the examination (Allan et al. 1980) or if a repeat scan is offered when the fetal heart examination at the routine scan is difficult (Allan et al. 1980, Rustico et al. 1995, Kirk et al. 1997). For other fetal anatomical structures, the specific focus and standardization of the examination procedure, for instance nuchal translucency measurement at an early gestation, has resulted in an increased detection of certain developmental disorders (Nicolaides et al. 1992, Hafner et al. 1998a, Nicolaides 1999, Schwärzler et al. 1999a, Souka et al. 2001, Hyett 2002). Had the same high quality standard used in the nuchal translucency measurements been applied to the

<table>
<thead>
<tr>
<th>Table 3 Sonographic detail of the aortic arch and ductal arch.</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Details obtained</strong></td>
</tr>
<tr>
<td>The arches</td>
</tr>
<tr>
<td>Aortic arch</td>
</tr>
<tr>
<td>- The aortic arch is seen as a tight hook (“candy cane shape”) tucking into the center of the heart</td>
</tr>
<tr>
<td>- The aortic arch gives rise to the head and neck vessels</td>
</tr>
<tr>
<td>Ductal arch</td>
</tr>
<tr>
<td>- The ductal arch arises from the right ventricle close to the anterior chest wall, is directed straight backwards (“hockey stick shape”) and connects to the descending aorta</td>
</tr>
<tr>
<td>- The ductal arch have no head and neck vessels</td>
</tr>
</tbody>
</table>

33
evaluation of the four-chamber view and the great arteries, our detection rate of CHDs most likely would have been improved.

Fetal echocardiography is defined as a specific fetal heart examination where the four-chamber view, the great arteries and the venous return to the heart are evaluated (Huhta and Rotondo 1991, Stümpflen et al. 1996). To optimize the image of the heart, additional techniques such as spectral Doppler, color Doppler (Chaoui 2001) and even 3D ultrasound (Budorick and Millman 2000, DeVore et al. 2003) may be used. Advanced techniques may well be of diagnostic value for the selected group of fetuses where a heart defect is suspected. For routine use in a screening situation, the use of advanced techniques is time consuming and requires skilled personnel. Until the views of the four chambers and the great arteries are properly learned and interpreted among the personnel performing routine fetal examinations, the advanced technology belongs in a fetal echocardiography setting.

**Teaching and training obstetric ultrasound**

In the early days of diagnostic ultrasound development, the static scanners were huge, bulky machines and required extremely skilful operators. As a consequence the average physician did not give much attention to the ultrasound technique (Eik-Nes 1993, Woo 2005). The obstetrician and gynecologist Alfred Kratochwil, an enthusiast in obstetric ultrasound development, saw the need for training and offered courses in ultrasound in Vienna, starting in 1968. This was after the introduction of the articulated-arm design, which made the ultrasound machines easier to manipulate (Woo 2005). The same year Kratochwil published his first book “Ultraschalldiagnostik in Geburtshilfe und Gynäkologie”, one of the earliest textbooks in ultrasonography in this field (Kratochwil 1968). Kratochwil has followed up his teaching tradition and, after his retirement in 1992, started regular courses for 3D ultrasound.

In the mid-1970s, the machines became easier to use with improved image quality. Since then, the ultrasound technique has gone through extensive improvement, both
when it comes to quality, size and cost of the machines. Ultrasound equipment is now available to all physicians. However, because modern ultrasound machines are highly developed technology with detailed resolution of the images and applications for different examination procedures, an understanding of the basic principles of ultrasound physics is a prerequisite for a proper and safe use of the equipment and an optimal ultrasound examination.

The educational and training aspect regarding the use of obstetric ultrasound has become increasingly more important with the improvement in ultrasound technology. Still, the requirements regarding ultrasound education and training vary greatly. With the introduction of routinely performed ultrasound scans in pregnancy, the need for ultrasound operators suddenly increased. It was emphasized that the ultrasound operators needed ultrasound training and experience, but there were no requirements for such training and thus no quality control of the examinations was carried out (Backe and Buhaug 1986).

The need for professions other than physicians to perform ultrasound examinations soon became obvious as the indications for the use of ultrasound in obstetrics increased. Already in the mid-1960s, nurses and radiographers assisted physicians in performing ultrasound examinations (Baker 2005). Approved by the American Institute of Ultrasound in Medicine, the American Society of Ultrasound Technical Specialists was established in 1970. After 10 years of discussions and struggling to survive as a profession, the first accreditation of educational programs occurred in 1982 (Baker 2005). The term ‘sonographer’ came to be used for this profession. Today, postgraduate ultrasound programs for sonographers are well established in several countries, such as the USA, Canada, Australia and the United Kingdom.

In the department of Obstetrics and Gynecology in Malmö, Sweden, where the routine scans were originally introduced in 1976, the scans were performed by nurse/midwives (Grennert et al. 1978). In the Scandinavian system, nurse/midwives have a strong tradition in pregnancy care, and it was therefore natural to base the scanning on this
profession. This was immediately adopted in Norway. In Ålesund, a number of midwives were systematically trained in obstetric ultrasound by Eik-Nes in the early 1980s. In Denmark, registered nurses initially took an interest in obstetric ultrasound, while nurse/midwives became involved at a later stage. For many years, obstetric ultrasound has been performed by personnel with varied ultrasound experience. In 1997, postgraduate education in obstetric ultrasound for nurse/midwives was established at the National Center for Fetal Medicine in Trondheim and is now part of the general offer of education at the Norwegian University of Science and Technology in Trondheim. This one-year part-time education includes both academic and clinical education and is at present the only education of this kind in Scandinavia.

Ultrasound imaging is not a passive activity but rather an interactive process involving the sonographer, patient, transducer and the ultrasound instrument (Kremkau 2006). The ultrasound operator has to interpret the images while the examination is being performed, placing strong demands on the ultrasound operator’s knowledge and experience in ultrasound technique and interpretation of ultrasound images. The importance of experience has clearly been demonstrated in the RADIUS study where the poor detection rate of developmental disorders most likely was a result of poor training and not a result of the diagnostic limitations of the ultrasound technique (Ewigman et al. 1993, Gonçalves and Romero 1993, Crane et al. 1994). Studies also show that tertiary centers have higher detection rates of fetal malformations than do non-tertiary centers. This is most likely due to the academic environment and a higher number of examinations resulting in increased ultrasound skills (Saari-Kemppainen et al. 1990, Crane et al. 1994, Wong et al. 2003). Other studies show the success in implementing teaching programs for the detection of fetal malformations (Sharland and Allan 1992, Hunter et al. 2000, Carvalho et al. 2002, Cuneo et al. 2004), emphasizing that the basic training has not been sufficient. Ultrasound societies and educational sites have guidelines for the proper way of examining the fetus (Meyer et al. 1990, ISUOG education committee 1996, Burr and Johanson 1998, Abuhamad et al. 2004, Allan et al. 2004, ISUOG guidelines 2006). These are of great value for the harmonization of the
examinations between countries, but may not reach their expectations as long as the basic ultrasound education insufficient.

With the rapid development of ultrasound technology, new techniques and examination procedures have been introduced in obstetric ultrasound. This implies that the appropriate way of performing ultrasound examinations a few years ago has changed and may be different and improved today. Regular updates are necessary to keep up with the development, and each ultrasound operator carries this responsibility. Interest and motivation have been reported as important factors when it comes to applying new ultrasound techniques (Roberts et al. 1995, Nicolaides 2004).

As long as governments and professional societies do not understand the necessity of ultrasound education, we have to accept that the quality of the examinations will vary, resulting in a variable quality of the scan offered to the pregnant population. Thus, the intention stated 20 years ago in the Consensus Statement from 1986 (Backe and Buhaug 1986) about improvement of the quality of the examinations conducted, has not yet been fulfilled for the total pregnant population in Norway.
AIMS OF THE STUDIES

One of the purposes of the second trimester routine ultrasound examination is to detect fetal developmental disorders that will benefit from prenatal diagnosis. Already in the late 80s it was clear that CHDs were detected far less frequently than defects in other fetal organs (Lys et al. 1989, Rosendahl and Kivinen 1989, Brocks and Bang 1991). To increase the prenatal detection rate of congenital heart defects, the present study has evaluated different methods to examine the fetal heart during the second trimester routine examination. An important factor was to find easy ways to examine the heart, as the fetal heart examination is only a small part of the complete routine scan. At the same time, the aim was to get as much information as possible from the heart to correctly differentiate normality from pathology. In addition to evaluating the sonographic appearance of the fetal heart, this study also addressed the sonographer/midwives’ scanning skills to see if their amount of experience affected their ability to obtain fetal heart structures, and thus affected their ability to detect heart defects. It was also of interest to follow the outcome of the CHDs, detected and not detected prenatally, to measure the impact of prenatal diagnosis of CHDs.

The aims of the five studies were:

I  To assess if incorporation of the four-chamber view of the fetal heart was achievable within the 30 minutes allocated to the second trimester routine examination at 18 weeks of gestation, and to evaluate any fetal, maternal or technical reasons for not being able to obtain this view.

II To evaluate prospectively whether the implementation of the four-chamber view to the second trimester routine examination in a large non-selected population would improve the detection rate of congenital heart defects compared to a period when the heart was not systematically evaluated, and to evaluate the CHDs detected and not detected to find ways to further improve the prenatal detection rate.
III To evaluate the detection rate of congenital heart defects in a non-selected population following the introduction of the four-chamber view and the great arteries to the second trimester routine scan, to understand the pattern of prenatal recognition of heart defects to further improve the detection rate, and to follow the fetuses/children with CHDs into the postnatal period to get a better understanding of the outcome, in order to improve the accuracy of counseling.

IV To compare the experienced and the less experienced sonographer/midwives’ ability to obtain fetal heart structures leading to detection of CHDs at the second trimester routine scan in a large non-selected population with the purpose of evaluating whether basic education and experience, as opposed to advanced examination procedures, was of importance in detecting CHDs, and to suggest improvements to the current ultrasound training program.

V To introduce a new three-dimensional technique based on tissue Doppler gating (TDOG) to create reconstructed, dynamic 3D ultrasound images of the fetal heart and to evaluate the accuracy of the signals and the reconstruction of the most common two-dimensional views used during a fetal heart examination.
MATERIAL AND METHODS

Populations and study design

The non-selected population came from a geographically well-defined area including the city of Trondheim and eight surrounding communities with approximately 200,000 inhabitants (Figure 15).

Figure 15 Communitites included in the study:
1 Klæbu
2 Malvik
3 Melhus
4 Midtre Gauldal
5 Rissa
6 Selbu
7 Trondheim
8 Tydal
9 Åfjord

This area is served solely by Trondheim University Hospital for all health care services. To ensure a true non-selected population, a community was included only when more than 97% of the pregnant women residing in the selected community received a prenatal ultrasound at the National Center for Fetal Medicine and more than 97% delivered at the University Hospital. The University Hospital also includes the only Neonatal Intensive Care Unit and the only Pediatric Cardiology Unit for the same population.

During the study period from August 1986 to December 2001, this non-selected population constituted the total population of 42,381 fetuses, all scheduled to be born at Trondheim University Hospital. Of those, 41,354 (98%) had an ultrasound examination performed at the National Center for Fetal Medicine, and thus formed the study population (screened population).
The second trimester routine ultrasound examinations were scheduled at 18 completed weeks based on the last menstrual period and/or early clinical assessment. The examinations were performed mainly by nurse/midwives with basic training in obstetric ultrasound. Following the official university-based education in obstetric ultrasound they are referred to as sonographer/midwives. Ultrasound examinations prior to the routine scan were done by physicians with ultrasound experience, while normal follow-up ultrasound examinations after the routine scan were mainly performed by sonographer/midwives. Except for the routine scan, the ultrasound examinations were performed on clinical indications, only.

The routine fetal examinations were scheduled at 30-minute intervals, and included a basic obstetric history as well as general information to the parents. The ultrasound examination comprised assessment of the number of fetuses, regular biometric measurements, placental location, a detailed survey of the fetal anatomy and an overview of the adnexae uteri. The final gestational age was based on the fetal BPD between 16+1 and 22+5 weeks (BPD outer-outter 38–60 mm) or the BPD and/or CRL for those detected prior to the routine examination. All data from the ultrasound examinations were prospectively registered in a computer database.

When a heart defect was diagnosed or suspected, a physician at the center carried out a detailed survey of the heart and the fetal anatomy, in most cases in the presence of a pediatric cardiologist.

A perinatal team consisting of an obstetrician, a sonographer/midwife, a pediatric cardiologist and a social worker supported the parents and the fetus/newborn from the time of diagnosis until after the delivery. If the pregnancy was continued, a specialist pediatric cardiac nurse became a part of the perinatal team. When immediate intervention after birth was expected, the planned delivery took place at the National Hospital in Oslo where neonatal cardiac surgery service is centralized in Norway. If a pregnancy was terminated, the parents were supported during that process and during
the time thereafter. In Norway, termination of the pregnancy may be approved until approximately 22 weeks.

Within the first 24 hours after birth, all newborns had a general clinical examination by a pediatrician. If a heart defect was suspected, the child was referred to a pediatric cardiologist for an echocardiogram. CHDs detected after discharge from the hospital were referred to the Pediatric Cardiology Unit. The health care program in Norway ensures all children regular physical examinations at child health care centers until school age. For the first part of the study these examinations took place at 3, 6 and 12 months, and at 2, 4 and 7 years of age (Paper II), the examinations were later changed to 6 weeks, 6 and 12 months, and at 2, 4 and 6 years of age (Paper III).

**Paper I**

**Incorporating the four-chamber view into the second trimester routine examination**

The prospective study included 7,322 consecutive fetuses at the second trimester routine examination and was divided into two periods: Period I, from June 1988 to May 1989, included 2,993 fetuses; Period II, from June 1989 to September 1990, included 4,329 fetuses. The criteria for a proper four-chamber view were: two ventricles of equal size; two atria of equal size; the heart occupying one-third of the thorax area; the heart located in the middle of the thorax with the apex pointing to the left anterior chest wall.

The five sonographer/midwives’ ability to include the four-chamber view into the second trimester routine protocol was assessed during Period II. Every fetus for which a four-chamber view could not be obtained was registered on a separate form with a comment about any reason why a proper image of this view failed.
Paper II
Prenatal detection of CHDs at the second trimester routine examination including the four-chamber view of the fetal heart
The prospective study included a total population of 12,232 fetuses, of which 11,894 (97%) fetuses had a second trimester routine ultrasound examination and constituted the study population (screened population). The study was divided into two phases: Phase I, from August 1986 to May 1988, included 4,435 fetuses; Phase II, from June 1988 to January 1991, included 7,459 fetuses. The criteria for a proper four-chamber view were: two ventricles of equal size; two atria of equal size; the heart occupying one-third of the thorax area; the heart located in the middle of the thorax with the apex pointing to the left anterior chest wall.

The detection rate of CHDs at the second trimester routine examination and a total prenatal detection rate was compared between Phase I, when no specific attention was paid to the fetal heart, and Phase II, when the four-chamber view was obtained in every fetus. The longest and shortest postnatal follow-up times for Phase I were 77 months and 58 months, for Phase II, 57 and 26 months, respectively.

Paper III
Prenatal detection of CHDs including the four-chamber view and the great arteries of the fetal heart – detection rates and outcome
The prospective study included a total population of 30,149 fetuses, of which 29,460 (98%) fetuses had prenatal ultrasound and constituted the study population (screened population). The study period lasted from February 1991 to December 2001. The criteria for obtaining the four-chamber view and the great arteries properly were: two atria of equal size; two ventricles of equal size; moderator band in the anterior right ventricle; tricuspid valve closer to apex than the mitral valve; intact ventricle septum; crossing of the ascending aorta with main pulmonary artery; aortic arch; ductal arch. From January 1995, a new scan was offered at approximately 20 weeks of gestation if a proper four-chamber view had not been seen at the routine scan. The examination at
20 weeks included the four-chamber view, the great arteries and the venous return to the fetal heart.

The postnatal follow-up of the children born with CHD included a description of their clinical condition. A classification system has been developed to critically evaluate the use of diagnostic procedures and therapies in the management of prevention of disease states (American Heart Association, 1994). The functional capacity is based on subjective symptoms as well as objective assessment (Bennett et al. 2002). According to the New York Heart Association classification system, “healthy” was defined as a person with cardiac disease without limitations of physical activity or objective evidence of cardiovascular disease (Class I).

The detection rate of CHDs was assessed after the four-chamber view and the great arteries of the fetal heart were included in the fetal examination protocol; the pattern of prenatal recognition was sought to further improve the detection rate, and CHDs were followed-up into the postnatal period to improve the accuracy of counseling. The children were followed for a minimum of two years and a maximum of 13 years.

**Paper IV**

**Comparison between experienced and less experienced sonographer/midwives in obtaining fetal heart structures and detecting CHDs prenatally**

All sonographer/midwives assessed during the study period were specially trained in obstetric ultrasound. The basic training included a minimum of 200 routine examinations supervised by experienced sonographer/midwives in addition to a Level I ultrasound course. For this study, “experienced” was defined as a sonographer/midwife who had performed more than 2000 routine examinations after completing the basic ultrasound training. “Less experienced” was defined as a sonographer/midwife who had performed between 200 and 2000 routine ultrasound examinations after completing the basic ultrasound training. The ultrasound experience following the basic training varied from 3 months to 17 years.
The prospective study included a population of 29,460 fetuses, of which 29,035 (99%) had had a second trimester routine examination performed by a sonographer,midwife. Those who did not receive a second trimester routine scan and those who were scanned by a physician were excluded. During the study period from February 1991 to December 2001, the four-chamber view and the great arteries were obtained from the fetal heart. During the first five years of the study period, heart structures obtained from every fetus were registered in detail on a separate form: four chambers; intact ventricular septum; tricuspid valve closer to the apex than mitral valve; ascending aorta; main pulmonary artery; aortic arch; ductal arch. From January 1995, a new scan was offered at approximately 20 weeks if a proper four-chamber view had not been seen at the routine scan. The examination at 20 weeks included the four-chamber view, the great arteries and the venous return to the fetal heart. Minor CHDs were excluded from this study.

A comparison between the experienced and the less experienced sonographer/midwives was carried out regarding their success rate in obtaining the fetal heart structures and prenatal detection of major CHDs.

**Paper V**

**Introduction of three-dimensional ultrasound imaging of the fetal heart (TDOG)**

The study included 8 selected pregnant women who were scheduled for a regular cardiac scan between 20–24 gestational weeks following a second trimester routine examination. The scanning time was 30 minutes for each fetus. Tissue Doppler data from the fetal heart were used to calculate a gating signal to create dynamic 3D reconstruction of the heart. The classical 2D views for reconstruction were: four-chamber view; ascending aorta; main pulmonary artery; three-vessel view; short-axis view of the great arteries; aortic arch; ductal arch; long-axis view of inferior and superior venae cavae.

A new method to image the fetal heart was introduced. Tissue Doppler gated (TDOG) dynamic 3D ultrasound images of the fetal heart were created, the accuracy of the
TDOG signals and the reconstruction of the most common views through the fetal heart were evaluated.

**Methods**

Classifications of congenital heart defects

In Paper II the heart defects were classified as critical or non-critical, in Papers III and IV as major or minor, which were synonymous to critical and non-critical. The difference in terminology was due to improved postnatal treatment over time, leading to survival of children with critical CHDs that in the early study period had had no treatment options. The classification was done retrospectively after a final diagnosis was achieved either by postnatal echocardiogram or by autopsy. The majority of the spontaneous losses after 22 weeks were autopsied. A perinatal pathologist at the National Center for Fetal Medicine performed the postmortem examinations (Isaksen et al. 1999). When postnatal echocardiogram or autopsy was not available, the prenatal video recordings and hard copies were scrutinized to ensure the correct heart diagnosis.

The defects were classified as *major* when surgical repair was most likely to be required because of gross structural complexity having functional significance (Mitchell et al. 1971), e.g. TGA, HLHS, AVSD, coarctation of aorta, large VSD (Papers II–IV). Heart defects were classified as *minor* when no intervention was likely to be required, e.g. mild pulmonary stenosis, mild aortic stenosis, small VSD, small ASD (Papers II and III). If there was doubt as to whether a heart defect was major or minor, the defect was classified as major if intervention was actually needed during the first month of life. Excluded from this study were arrhythmias with a structurally normal heart, patent ductus arteriosus, atrial septal defects < 3 mm (Radzik et al. 1993) and absent inferior vena cava with azygos continuation. Rhabdomyomas were categorized as tumors and thus excluded as heart defects.

Fetal karyotyping was offered in all prenatally detected cases. During the first 10 years of the study it was mainly performed by fetal blood sampling; later, by amniocentesis
and fluorescence in situ hybridization analysis (FISH) (Wakui et al. 1999). The 22q11 microdeletion was not routinely checked during this study. When a karyotype was not obtained, it was considered as probably normal.

The major CHDs were grouped as *isolated* in fetuses with a normal karyotype and absence of extra-cardiac malformations; as *abnormal karyotype* in fetuses with a chromosome aberration with or without extra-cardiac malformations; as *associated malformations* in fetuses with extra-cardiac malformations but a normal karyotype (Papers III and IV). The most common major CHDs (AVSD, TGA, HLHS, coarctation of aorta) were grouped as *simple* when additional cardiac defects were absent, e.g. TGA or AVSD; as *complex* when additional heart defects were present, e.g. TGA with associated single ventricle, HLHS with additional coarctation of aorta (Paper III).

**Ultrasound equipment and computer software**

The ultrasound scanners used during the total study period were replaced over time. For Papers I and II, a Hitachi EUB 40 ultrasound scanner with 3.5 or 5 MHz transducers and a Hitachi EUB 410 with a 5 MHz transducer (Hitachi, Tokyo, Japan) were used. For Papers III and IV the ultrasound scanners used were Hitachi EUB 415 and EUB 6000 with 5 MHz and 3.5 MHZ transducers (Hitachi, Tokyo, Japan), VingMed CFM 800 with a 5 MHz mechanical sector transducer and VingMed System Five with a 3.5 MHz curvilinear transducer (Vingmed Sound, Horten, Norway). For Paper V, a Vivid 7 scanner (GE Medical Systems) with a 1.5D probe with a frequency range of 2–5 MHz (Vingmed Sound, Horten, Norway) was used and the reconstruction was performed with the EchoPAC-3D software product (GE Medical Systems).

**3D imaging**

**Data acquisition**

3D data were acquired by collecting several datasets in one sweep covering the complete heart for each fetus. Using a free-hand technique, the transabdominal scan plane was slowly tilted manually over a period of 8–20 seconds (20–50 cardiac cycles).
The total angle of the sweep was estimated by recording a separate loop through the center of the heart, in the elevation direction of the sweep. Each dataset consisted of a duplex sector scan with a visible B-mode image and 2D tissue Doppler not visible during acquisition. Based on subjective evaluation of the clinical data, a second harmonic setup with a transmitting frequency of 2.5 MHz and a receiving frequency of 5.1 MHz, an inter-beam distance of 0.3˚ and a data rate of 96 frames/second was chosen as an adequate trade-off between frame-rate and image quality. The tissue Doppler settings were adjusted to provide a small bandwidth, but the frame size and frame rate were the same, with transmitting and receiving frequencies of 2.4 MHz and an inter-beam distance of 3.8˚. There were three pulse firings in each beam direction of the tissue Doppler.

The gating signal was derived from clutter-filtered IQ data. Following calculation of autocorrelation of lag 1, the imaginary part of this was averaged over each frame, and then the local minima were extracted.

Reconstruction
The EchoPAC-3D software program converted the B-mode scan-lines into a sequence of voxel datasets, yielding a 4D dataset with temporal resolution equal to the B-mode frame rate and duration of one cardiac cycle. The 4D data were then visualized by two standard examination techniques for 3D medical imaging: 2D slicing and volume projection.

Gating signal accuracy
CW spectral Doppler from the umbilical artery was used to evaluate the accuracy of the TDOG signal, and was recorded simultaneously with the B-mode and tissue Doppler IQ data, but with a separate ultrasound system. The recordings were synchronized for offline analysis by simultaneously recording an independent signal on both ultrasound systems. The cycle lengths from the two methods were then compared and the variation in time delay between upstroke readings from the spectrum and the TDOG signal was examined.
The use of the tissue Doppler signal for synchronization was evaluated visually by running side-by-side consecutive cineloops.

Statistical analysis

The observed frequencies were tested for significance using the $X^2$ (Chi-square) statistical test (Papers I and II). Groups were compared in 2x2 contingency tables using Pearson Chi-Square or Fisher’s exact test (Papers III and IV). The level of significance was set at $p < 0.05$. 
RESULTS AND COMMENTS

Paper I

Incorporating the four-chamber view into the second trimester routine examination

In period I, a four-chamber view could not be obtained in 220 (7%) of the 2993 fetuses scanned using a Hitachi EUB 40 ultrasound scanner. A significant difference was found in the number of four-chamber views not obtained when the 3.5 MHz and the 5 MHz transducers were compared, 35/131 (26.7%) and 185/2862 (6.5%), respectively (p < 0.001).

In period II, there was no difference in the number of four-chamber views not obtained with a 5 MHz transducer when the Hitachi EUB 40 ultrasound scanner used through period I was compared with a new Hitachi EUB 410 ultrasound scanner, 3.8% versus 4.3%, respectively. When the results for the Hitachi EUB 40 scanner with a 5 MHz transducer were compared between Period I (6.5%) and Period II (3.8%), significant improvement was found (p < 0.001).

The reasons why a four-chamber view could not be obtained were; unfavorable position of the fetus, poor imaging and maternal obesity. The main reason for not obtaining a four-chamber view utilizing the 3.5 MHz transducer was poor imaging (52%); for the 5 MHz transducer, the main reason was unfavorable fetal position (64%).

Figure 16 Four-chamber views not obtained for each gestational week at the second trimester routine scan.
Figure 16 shows the percentage of four-chamber views not obtained for each gestational week between 16 and 21 weeks.

The five sonographer/midwives active in both periods significantly improved their ability to obtain a four-chamber view from period I to period II when a 5 MHz transducer was used. The individual results for not obtaining a four-chamber view varied from 3-10% in period I to 2-6% in period II (p < 0.001).

Comments:
The four-chamber view was introduced in the mid-1980s (Fermont et al. 1985, Allan et al. 1986b) with the purpose of increasing the prenatal detection rate of CHDs. At that time, most studies concerning the fetal heart came from selected populations. In one such study (Copel et al. 1987), it was suggested that a four-chamber view of the fetal heart should be obtained in more than 95% of the fetuses between 18 and 40 weeks. As these results were based on ultrasound examinations throughout the pregnancy in a selected population it was not obvious that the same results would be reached in a non-selected population at an earlier gestational age. At our center, 98% of the pregnant population received a second trimester routine scan between 16 and 22 gestational weeks and the 30 minutes allocated to the complete consultation were already exhausted to serve other tasks.

Our study showed that a four-chamber view could easily be incorporated into the second trimester routine scan at approximately 18 weeks, as this view was successfully obtained in 96% of the fetuses. The result was similar to findings published from other non-selected populations at that time (Achiron et al. 1992, Vergani et al. 1992) and could also be compared with results from selected populations (Copel et al. 1987). It is assumed that 50% of the major CHDs can be detected by evaluating the four-chamber view (Allan et al. 1986b, Cullen et al. 1992), thus, an incorporation of this view may increase the prenatal detection rate of heart defects. During this study period, 87% of the pregnant women in the total country of Norway accepted the second trimester scan, 97% in the region served by Trondheim University Hospital (Nafstad and Backe 1989).
Incorporation of the four-chamber view to the routine scan protocol would thus cover most of the pregnant population with the possibility to detect 2/1000 major CHDs.

During the two periods, there were significant improvements in obtaining the four-chamber view. In Period I, the 5 MHz transducer was superior to the 3.5 MHz transducer in obtaining this view. This could be explained by the improvement in resolution when a higher frequency was chosen. The different reasons why a four-chamber view could not be obtained may support this explanation. The main reason for not obtaining this view utilizing the 3.5 MHz transducer was poor imaging, which most likely was caused by poor resolution. In both Periods I and II, the main reason for not being able to obtain the four-chamber view with the 5 MHz transducer was fetal position. Thus, the reason was not dependent on the quality of the transducer. Obviously, the awareness of the ultrasound equipment and the choice of transducer are of significant importance when small fetal structures are to be imaged. As a consequence of the obvious difference between the transducers, the 5 MHz transducer was preferred for the total second trimester routine scan, as better resolution due to an increase in frequency was expected to affect the image of the fetal anatomy in general, and thus improve the overall quality of the routine scan (DeVore 1985, Kremkau 2006).

A four-chamber view could not be obtained in 7% of the fetuses in Period I when a Hitachi EUB 40 ultrasound scanner with a 5 MHz transducer was utilized. With a new Hitachi EUB 410 scanner with the same transducer in Period II, 4% could not be obtained, an improvement that could be explained by a new and better ultrasound machine. Interestingly, the results for the Hitachi EUB 40 improved from 7% in Period I to 4% in Period II. Although the same machine and transducer were used, the results improved significantly. Also when evaluating the results for the 5 sonographer/midwives active in both periods, significant improvements were found. The improvements may thus also be explained by a learning effect. When particular focus was set on the four-chamber view, the technique to obtain this view and the familiarity of the details in the four-chamber view would most likely improve over time (Sharland and Allan 1992). The relatively long learning curve to obtain the four-chamber view,
also found by Sharland and Allan (1992), may suggest that the examination of the fetal heart is difficult to learn in a second trimester routine setting.

The gestational age seemed to be of significant importance for the success in obtaining the four-chamber view. Prior to 18 weeks, the number of four-chamber views not obtained were higher than at 18–21 weeks. Sharland and Allan (1992) also found that four-chamber views were obtained less frequently prior to 18 weeks than after 18 weeks of gestation. Several authors have shown a high ability to obtain the four-chamber view after 18 weeks, and some report the results up to 40 gestational weeks, but without specification for each week (Copel et al. 1987, Achiron et al. 1992). As the routine fetal examination preferably is done in the second trimester, an ultrasound scan at 18–20 weeks performed with a 5 MHz transducer seems to be the optimal choices for the best results according to our data. The examiner’s ultrasound experience also needs to be considered when evaluating the success in obtaining the four-chamber view.

**Paper II**

**Prenatal detection of CHDs at the second trimester routine examination including the four-chamber view of the fetal heart**

**Detection of CHDs**

In Phase I, when no particular attention was paid to the fetal heart, 49 fetuses in the population of the 4435 scanned had a CHD. Of the 17 major CHDs, three (18%) were detected at 18 weeks of gestation. None of the 32 minor CHDs were prenatally detected.

In Phase II, when the four-chamber view of the fetal heart was obtained, 90 within the population of the 7459 fetuses scanned had a CHD. Nine (39%) of the 23 major CHDs were detected prenatally: 6/23 (26%) at the second trimester routine scan and 3/23 (13%) at later scans.

Table 4 shows the major CHDs missed at the 18-week routine scan with respect to the expected four-chamber view presentation, including the three detected later in the
pregnancy. In three of the 17 cases missed at 18 weeks (VSD, overriding aorta; TOF; double aortic arch), an adequate four-chamber view could not be obtained due to poor imaging or unfavorable position of the fetus. In 7/17 (41%) cases, the routine scan was performed prior to 18 weeks of gestation.

Table 4  Major CHDs not detected prenatally (including three detected later in the pregnancy) at the 18 week examination with four-chamber view presentation in Phase II.

<table>
<thead>
<tr>
<th>CHD diagnosis</th>
<th>n</th>
<th>Normal four-chamber view expected</th>
</tr>
</thead>
<tbody>
<tr>
<td>AVSD (twin, trisomy 21)</td>
<td>1</td>
<td>no</td>
</tr>
<tr>
<td>Coarctation of the aorta</td>
<td>2</td>
<td>yes</td>
</tr>
<tr>
<td>Coarctation of the aorta, ASD</td>
<td>1</td>
<td>yes</td>
</tr>
<tr>
<td>Common arterial trunk, large VSD*</td>
<td>1</td>
<td>no</td>
</tr>
<tr>
<td>Coarctation of the aorta, DIRV, VSD, mitral stenosis*</td>
<td>1</td>
<td>no</td>
</tr>
<tr>
<td>Double aortic arch</td>
<td>1</td>
<td>yes</td>
</tr>
<tr>
<td>TOF</td>
<td>2</td>
<td>yes</td>
</tr>
<tr>
<td>TGA</td>
<td>3</td>
<td>yes</td>
</tr>
<tr>
<td>Pulmonary atresia, tricuspid atresia, VSD*</td>
<td>1</td>
<td>no</td>
</tr>
<tr>
<td>Tricuspid atresia, pulmonary stenosis, VSD</td>
<td>1</td>
<td>no</td>
</tr>
<tr>
<td>VSD, large</td>
<td>2</td>
<td>no</td>
</tr>
<tr>
<td>VSD, overriding aorta (trisomy 18)</td>
<td>1</td>
<td>yes</td>
</tr>
<tr>
<td>Total</td>
<td>17</td>
<td></td>
</tr>
</tbody>
</table>

*CHD detected in the third trimester

Among the 4% in which a four-chamber view could not be obtained at the second trimester routine examination, 3 fetuses with major CHDs were found (incidence 10/1000 fetuses).

Epidemiology
To ensure complete epidemiological data for the total population, 2 newborns with minor CHDs from Phase I (muscular VSD; secundum ASD) and 2 from Phase II (muscular VSD; pulmonary artery stenosis), where the mothers had not attended the ultrasound program at all, were included. The incidences of congenital heart defects for the total population from Phases I and II were: 11/1000 fetuses and 12/1000 fetuses, respectively; for major CHDs 4/1000 and 3/1000 fetuses; for minor CHDs 7/1000 and 9/1000 fetuses, respectively.
Isolated VSDs constituted 18/34 (53%) of the minor CHDs in the total population in Phase I, 50/69 (73%) in Phase II. The same numbers for the isolated secundum ASDs were 5/34 (14%) and 7/69 (10%), respectively. Within the first year of life, 50% of the isolated VSDs from Phase I and 70% from Phase II closed spontaneously.

Chromosome aberrations and associated malformations
Chromosome aberrations were found in 7/40 (18%) of the major CHDs in the total population from Phase I and II, in 6/103 (6%) among the minor CHDs. The corresponding numbers for CHDs with associated malformations and normal karyotype were 6/40 (15%) and 11/103 (11%), respectively. Chromosome aberrations and/or associated malformations were present in 7/12 (58%) of the major CHDs prenatally detected, and in 6/28 (21%) of those not detected (p = 0.03).

Outcome
Of the 12 fetuses with a major CHD detected prenatally during Phases I and II, 9 (75%) were born alive and 5 (42%) were alive after 2 years. Nine CHDs were detected prior to 22 weeks when TOP was an option, and 3/9 (33%) chose to terminate the pregnancy (Trisomy 18, VSD, overriding aorta; Trisomy 21, AVSD; pulmonary atresia, single ventricle complete situs inversus). Of the 28 fetuses with CHD detected postnatally, 26 (93%) were born alive, and 22 (79%) were alive after 2 years. One (4%) of the major CHDs from Phase II not detected prenatally was terminated due to prenatal findings of Trisomy 18. The heart defect, VSD and overriding aorta, was found during autopsy. One intrauterine death was due to associated malformations (hydrocephaly, bilateral pes equino varus, single umbilical artery) and the postductal coarctation of aorta was found during autopsy.

Comments:
Detailed structures from the fetal heart imaged by real-time two-dimensional ultrasound were reported as early as in 1980 (Allan et al. 1980, Kleinman et al. 1980). At that time, pregnancies at increased risk for CHD were offered fetal echocardiography (Callan et al. 1991, Allan et al. 1994). Since then we have learned that most fetuses and children
with CHD are born to mothers at no increased risk at all (Davis et al. 1990, Sharland et al. 1990, Callan et al. 1991, Allan et al. 1994). The second trimester routine examination, offered to most pregnant women, therefore seems like the best opportunity to detect most of the CHDs in the population. Subsequent studies have shown that the four-chamber view of the fetal heart can easily be obtained at this routine scan (Achiron et al. 1992, Vergani et al. 1992, Sharland and Allan 1992).

Our study showed that the prenatal detection rate of major CHDs at the second trimester examination improved from 18 to 26% when the four-chamber view was obtained in nearly every fetus. Among the 4% where a four-chamber view could not be obtained due to poor imaging or unfavorable position of the fetus (Paper I), three major CHDs were found. At that time a rescan was not a part of the protocol when a four-chamber view was not obtained, unless there was a suspicion of a heart defect. Retrospectively we may conclude that these fetuses represented a high-risk group, as the incidence of major CHD among those 4% was 10/1000 fetuses compared to 4/1000 in the general population. Those CHDs would most likely have been detected with a follow-up, detailed scan; this would have increased the detection rate at the routine scan from 26% to 39%.

In the early nineties, few studies reported detection rates of CHDs from non-selected populations. In two studies from low-risk populations (Achiron et al. 1992, Vergani et al. 1992) the prenatal detection rates of CHDs after implementation of the four-chamber view varied from 48–81%. Those studies differed with the present study in population selection, gestational age and duration of the routine scan, a rescan if the first failed and shorter postnatal observation time affecting the ascertainment of cases. These differences make the results difficult to compare. One study from a selected population reported a 92% sensitivity of the four-chamber view (Copel et al. 1987). However, one should be cautious comparing results from selected populations to non-selected populations, since results from selected populations are influenced by the high incidence of the disease. Despite the differences, a common finding for all the studies was the increase in prenatal detection of CHDs after the four-chamber view was
implemented to the fetal heart examination – which was the desired effect. One may conclude that a detailed evaluation of the four-chamber view is necessary to detect CHDs prenatally.

Of the major CHDs missed at the routine scan (including the three CHDs detected later), 35% would be expected to have an abnormal four-chamber view at that time in the pregnancy (Table 4), thus a four-chamber view was interpreted to be normal when it obviously was not. Several studies have documented that the four-chamber view is difficult to obtain prior to 18 weeks in the pregnancy (Sharland and Allan 1992). In the present study, 41% of those missed had a routine scan at a gestation prior to 18 weeks, which might have affected the ability to interpret the four-chamber view correctly. Two of those scanned prior to 18 weeks (tricuspid atresia; coarctation of the aorta, DIRV) were detected at a later third trimester scan, which supports the experience that the four-chamber view is easier to examine at a later gestation (Copel et al. 1987, Achiron et al. 1992, Tegnander et al. 1994). As the routine fetal examination is a compromise accommodating the various reasons for performing such a scan we rather need to look at ways of improving the scan qualities than at changing the time of its performance. Also, the 22 week upper limit for termination of a pregnancy in Norway requires an anomaly scan prior to 20 gestational weeks.

Evaluation of the four-chamber view can possibly only detect heart defects with an abnormal four-chamber view. CHDs such as TGA, TOF, DORV, mild pulmonary stenosis, mild aortic stenosis, and small VSDs will most likely appear with a normal four-chamber view and thus be missed. For this reason, the four-chamber view is expected to detect 50% of the major CHDs (Allan et al. 1986b, Cullen et al. 1992). Sharland and Allan (1992) excluded the CHDs with a normal four-chamber view and evaluated the detection rate of CHDs with an abnormal four-chamber view, only. They implemented a teaching program where ultrasonographers from ten obstetric ultrasound units were taught to obtain the four-chamber view. As a result of the teaching program, 77% of the CHDs with an abnormal four-chamber view were prenatally detected. The percentage of four-chamber views not obtained decreased dramatically from 15-20%
and ended with less than 5% in most units. Finally, an increase was found in the number of CHDs detected prenatally because of suspected abnormal four-chamber views at the routine scan. Obviously, the teaching program had an impact on the way the four-chamber view was evaluated. Still, 23% of the CHDs with an abnormal four-chamber view were not detected, emphasizing the difficulties in prenatal evaluation of this view.

Our criteria for a proper four-chamber view were two ventricles of equal size; two atria of equal size; the heart occupying one-third of the thorax area and located in the middle of the thorax with the apex pointing toward the left anterior chest wall. In addition to those criteria, for their teaching program, Sharland and Allan (1992) also evaluated the two atrioventricular valves which meet the atrial and ventricular septa at the crux of the heart in an offset cross-appearance, the presence of the foramen ovale defect in the atrial septum and the appearance of an intact ventricle septum. As the present study reported the detection rate among all major CHDs in the population and Sharland and Allan (1992) among major CHDs with an abnormal four-chamber view, these results are difficult to compare. Still, one may speculate that requiring more details from the four-chamber view forces the examiner to obtain a better image and thus makes it easier to detect CHDs.

Chromosome aberrations and/or associated malformations were found in 32% of the major and in 16% of the minor CHDs. A high percentage of extra-cardiac malformations in CHDs has been reported by others (Berg et al. 1988, Copel et al. 1988), and the importance of cytogenetic evaluation when a CHD is detected has been emphasized. The presence or absence of extra-cardiac malformations is important for the outcome and thus the counseling process.
Paper III

Prenatal detection of CHDs including the four-chamber view and the great arteries of the fetal heart – detection rates and outcome

Detection of CHDs

Of the 29,460 fetuses scanned, 97 were found to have a major CHD, 333 a minor CHD. The incidences of major and minor CHDs were 3.3 and 11.3/1000, respectively, making an incidence of 14.6/1000 fetuses in the study population. Figure 17 shows the incidences of major and minor CHDs in the total population for the Phases I, II and III.

Figure 17  Incidences of major and minor CHDs for each phase in the total population of 42,381 fetuses.

Of the major CHDs, 55/97 (57%) were detected prenatally: 9/55 (16%) prior to, 36/55 (66%) at, and 10/55 (18%) after the routine scan, respectively. Of all major CHDs in the study population, 45/97 (46%) were detected prior to 22 weeks.

Figure 18  Major CHDs in a non-selected population of 41,354 fetuses, detection rates and time of detection.
Of the major CHDs classified as complex, 11/14 (79%) were detected prenatally compared to 22/40 (55%) of those classified as simple (p = 0.12). Figure 18 shows the percentages of major CHDs prenatally detected for the Phases I, II and III and the time of detection.

Of the 9 fetuses with major CHD detected prior to the routine scan, 5 (56%) had an increased nuchal translucency or cystic hygroma.

Six of the 10 major CHDs detected late in the pregnancy were missed at the routine scan, while four had a first scan at our institution after 29 weeks. Four (67%) of the 6 missed would have had a normal four-chamber view at the 18-weeks scan (2 TOF; TGA; pulmonary valve stenosis).

Isolated CHDs constituted 43/97 (44%) of all major CHDs in the study population, and 19/43 (44%) were detected prenatally. Of the isolated CHDs, 23/43 (54%) were ductal dependent and 11/23 (48%) of those were detected. Of the 24 isolated CHDs missed prenatally, 14 (58%) would be expected to have an abnormal four-chamber view at the second trimester routine scan.

Major heart defects with associated malformations or abnormal karyotype totaled 54/97 (56%). Thirty-six (67%) of them were detected prenatally, a significantly better result compared to the number of major isolated CHDs detected prenatally (p = 0.03).

<table>
<thead>
<tr>
<th>Study phase</th>
<th>Population N</th>
<th>Isolated VSDs n</th>
<th>Incidence</th>
<th>Secundum ASD n</th>
<th>Incidence</th>
</tr>
</thead>
<tbody>
<tr>
<td>I</td>
<td>4,615</td>
<td>18</td>
<td>0.6</td>
<td>5</td>
<td>0.2</td>
</tr>
<tr>
<td>II</td>
<td>7,617</td>
<td>51</td>
<td>1.6</td>
<td>7</td>
<td>0.2</td>
</tr>
<tr>
<td>III</td>
<td>30,149</td>
<td>192</td>
<td>6.3</td>
<td>39</td>
<td>1.3</td>
</tr>
<tr>
<td>Total</td>
<td>42,381</td>
<td>261</td>
<td>6.2</td>
<td>51</td>
<td>1.2</td>
</tr>
</tbody>
</table>

Twelve (4%) of the 333 minor CHDs in the study population were detected prenatally. Of all minor CHDs, 188 (57%) were isolated VSDs and 38 (11%) were isolated
secundum ASDs. Table 5 shows the incidences of isolated VSDs and ASDs in the total population for Phases I, II and III.

Abnormal karyotype and associated malformations
Abnormal karyotype was found in 37 (38%) of the 97 major CHDs. Of the CHDs grouped as complex and simple, 2/14 (14%) and 21/40 (53%) had abnormal karyotype, respectively (p = 0.02). Among the 55 CHDs detected, 26 (47%) had an abnormal karyotype and 10 (18%) had associated malformations and normal karyotype. The corresponding numbers for the 42 major CHDs not detected prenatally were 11 (26%) and 7 (17%), respectively. A significant difference was found between the prenatally and postnatally detected major CHDs with associated malformations including abnormal karyotype (p = 0.03). Forty-five (82%) major CHDs were detected prior to 22 weeks when termination of the pregnancy was an option. Among the 24/45 (53%) who opted for a termination, 21 (84%) had associated anomalies including abnormal karyotype, compared to 10/21 (48%) of those who continued their pregnancy (p = 0.01). Five (63%) of the 8 intrauterine fetal deaths had abnormal karyotype.

Of the 333 minor CHDs, 27 (8%) had abnormal karyotype and 32 (10%) associated malformations. Of the 12 detected prenatally, all had abnormal karyotype or associated malformations.

Outcome
The outcomes of the major CHDs detected and not detected prenatally are shown in Figure 19. Intrauterine fetal death occurred in 8/55 (15%) of the fetuses with a major defect detected, but in none of the 42 fetuses with a major defect that was not detected (p = 0.01). When the terminated pregnancies were excluded, 23/31 (74%) of the fetuses with a major CHD detected prenatally were born alive compared to 41/41 (100%) of those not detected prenatally (p = 0.001). Of the children who survived until two years of age, all were still alive at a median follow-up of six (range 2-13) years.
Figure 19  Outcome of major CHDs in a non-selected population of 29,460 fetuses.

Major CHDs  N = 97

Detected prenatally  n = 55

TOP  n = 24
IUFD  n = 8

Chromosome aberration  n = 14
Serious anomalies  n = 7
Serious CHD  n = 3

Detected postnatally  n = 42

IUFD  n = 0
TOP  n = 1

Serious associated anomalies

Born alive  n = 23/55 (42%)*

Born alive  n = 41/42 (98%)*

Died postnatally  n = 8

Died postnatally  n = 7

Chromosome aberration  n = 4
Serious CHD  n = 4

Chromosome aberration  n = 1
Serious CHD  n = 6

Alive after 2 years  n = 15/55 (27%)*

Alive after 2 years  n = 34/42 (81%)*

Morbidity  n = 9/15 (60%)

Healthy  n = 6/15 (40%)

Healthy  n = 17/34 (50%)

Morbidity  n = 17/34 (50%)

Chromosome aberration  10
Cerebral  2
Cardiac  4

*p < 0.001

Chromosome aberration  3
Cerebral  3
Cardiac  3
One of the 12 fetuses with a minor CHD detected prenatally was born alive with a possible genetic defect and a syndrome.

Isolated ventricular septal defects without extra-cardiac malformations totaled 57% (188/333) of the minor heart defects. Of those, 162 (86%) were muscular and 26 (14%) were perimembranous. During the first year of life, 77/162 (48%) of the isolated muscular and 3/26 (12%) of the isolated perimembranous VSDs closed spontaneously.

Comments:
This study showed that inclusion of the great arteries to the fetal heart examination, in addition to the four-chamber view, increased the detection rate of major CHDs at the second trimester routine scan from 26% (Paper II) to 37%, and the total prenatal detection rate from 39% to 57%.

The classification of CHDs seems arbitrary when comparing papers evaluating the prenatal detection of CHDs, making both detection rates and outcome difficult to compare (Cullen et al. 1992, Fernandez et al. 1998, Bull 1999, Jaeggi et al. 2001, Carvalho et al. 2002, Wong et al. 2003). Even among experts the definition of major CHDs seems to differ, although the majority seem to classify major CHDs as those that will require surgical or interventional treatment during the first year of life (Allan et al. 2001). In addition, the evaluation of the severity of major CHDs has changed over time and should be seen in relation to the improvement in postnatal treatment and surgery. In the late-1980s, HLHS was considered a critical and lethal diagnosis and most parents chose to terminate the pregnancy after being counseled about treatment options (Allan et al. 1991). With improved surgical techniques (Jacobs et al. 1994) HLHS is no longer a CHD without option of postnatal treatment and is thus not considered lethal or critical (Andrews et al. 2001, Tworetzky et al. 2001, Wilkins-Haug et al. 2005). In line with the therapeutic improvements we renamed the two groups of CHDs from critical and non-critical to major and minor, although the definition of the CHDs remained the same through the complete study period.
The study period extended over 11 years. During that time, advanced imaging techniques and diagnostic tools were introduced, making it easier to detect fetal developmental disorders at an early gestation (Blaas et al. 1999, Devine and Malone 1999, Schwärzler et al. 1999a). Thus, the different use of prenatal tests makes studies evaluating the second trimester routine examination difficult to compare. The 11–13+6 week nuchal translucency screening (Nicolaides et al. 1992) and the development of OSCAR-clinics (One-Stop Clinic for Assessment of Risk) (Nicolaides et al. 2002) result in early detection of major CHDs and consequently a lower incidence of CHDs at the time of the second trimester examination. In one study from a non-selected population, a fetal heart training program was carried out, resulting in a 38% detection rate of major CHDs at the second trimester routine scan (Carvalho et al. 2002). Although the result seems similar to ours, 80% of the women in their population had an early scan detecting 20% of the major CHDs which would affect the number of CHDs present at the second trimester scan. Our data also show the same trend. Although nuchal translucency screening is not an offer to the pregnant population in Norway, 9% of the major CHDs were detected prior to the routine scan, changing our previous pattern of prenatal detection of CHDs (Figure 18).

The introduction of the four-chamber view to the fetal heart examination raised high expectations for the detection rate of major CHDs (Copel et al. 1987). As the detection rate from second trimester routine scans started to emerge, the results clearly did not meet the expectation of 50% detection (Allan et al. 1986b, Sharland et al. 1992) but rather between 4.5–48% from non-selected populations (Achiron et al. 1992, Ott 1995, Buskens et al. 1996, Todros et al. 1997). When these studies revealed that the four-chamber view provided poorer results than anticipated, more details of the heart were included rather than question the methods of four-chamber view acquisition. Retrospectively, one may wonder if the potential of this view had been fully utilized (Chaoui 2003b).

With both the four-chamber view and the great arteries included into the fetal heart examination, studies showed that total prenatal detection rates of major CHDs increased
to 40–83% (Achiron et al. 1992, Bromley et al. 1992, Hafner et al. 1998b, Carvalho et al. 2002, Wong et al. 2003). Our total detection rate of 57% was in accordance with the published data, but the detection rate at the second trimester routine scan, 37%, was still low. The details of the fetal heart are better imaged after 20 weeks (Schwärzler et al. 1999b, Allan 2003), thus a scan after 20 weeks would probably make it easier to detect CHDs. With an early scan for nuchal translucency measurements, including dating of the pregnancy, an anomaly scan can be performed at a later gestation (Carvalho et al. 2002). Since the 18-week routine scan still is the only ultrasound examination offered to all pregnant women in Norway, efforts have to be made to increase the detection rate of major CHDs at this scan. Also, the upper limit for termination of the pregnancy is 22 weeks in Norway, thus CHDs need to be detected prior to 22 weeks to give the parents this option.

Figure 18 shows data from a period ranging more than 15 years from our large non-selected screened population of 41,354 fetuses and summarizes the prenatal detection of major CHDs through all the three Phases. Since 1986, all pregnant women in Norway have been offered one routine ultrasound examination in the second trimester (Backe and Buhaug 1986), and since then every second trimester scan at our center has been prospectively registered in our database. The data from this database clearly showed that the detection of major CHDs through 1986 and 1987 was poor, only 18% (Paper II). This was the time that the four-chamber view of the fetal heart was introduced, and during Phase II our target was to obtain a four-chamber view in all fetuses scanned. As we started to pay more attention to the fetal heart, the results improved and, as Figure 18 shows, CHDs were even detected in the third trimester when ultrasound scans were done on indications only. An increase in the detection rate to 26% at the routine scan (Phase II) still left most of the major CHDs undetected. Encouraged by the improvement after implementation of the four-chamber view, we added the great arteries to the fetal heart examination to further increase the detection rate. In Figure 18, the improved detection rate as more details of the fetal heart were added to the ultrasound examination is clearly shown. The impact of the second trimester scan is emphasized here, since this is the time that most of the major CHDs were detected, and
the changing pattern in prenatal detection of the defects was observed. The focus on the early scan has resulted in detection of CHDs prior to the second trimester scan, a change that now has to be addressed. Although the focus on the fetal heart examination through many years clearly has resulted in increased detection rates, Figure 18 shows that too many CHDs still are left undetected, clearly challenging our skill to organize and perform the fetal heart examination by ultrasound.

To help in further improving the skills in fetal heart scanning, characteristic findings of the defects detected as opposed to those not detected were sought. When looking at our data, we found that we detected significantly more of the CHDs with abnormal karyotype and/or associated malformations than we detected isolated CHDs. This shows that extra-cardiac malformations seem easier to detect than defects in the heart itself. This has also been found by other investigators (Stoll et al. 2002). Only 44% of all isolated CHDs and 48% of the isolated ductal dependent CHDs were detected prenatally, leaving more than one half of the cases undetected. Since the heart is the only affected organ in those cases, we can conclude that the examination of the fetal heart most likely has not been sufficient. Of all the isolated CHDs, we detected 47% with a normal four-chamber view, but, 58% of those we missed were expected to have an abnormal four-chamber view. Thus, a true abnormal four-chamber view was obviously not always recognized. When detection of the complex and the simple CHDs were compared, we detected more of the complex CHDs where the defect was severe than the simple CHDs. This pattern seemed to favor the detection of the more serious defects. Convincing data today show that the outcome seems better for isolated CHDs compared to those with extra-cardiac malformations, especially when prenatally detected (Bonnet et al. 1999), thus, efforts have to be made to increase the detection of isolated CHDs. Overall, we may conclude that the fetal heart examination in general is not sufficiently mastered, and more focus has to be put on both the examination of the four-chamber view and the great arteries.

Minor CHDs have been difficult to detect prenatally, especially as isolated defects (Allan et al. 1989). In our series, we did not detect minor CHDs in Phase I or II (Paper
II), but 12 (4%) minor CHDs were detected in the present study (Phase III). One may speculate whether improved ultrasound skills and knowledge over time have resulted in detection of those minor defects, or if it was the extra-cardiac malformations rather than the heart itself that was the reason for the detection (Hafner et al. 1998b). The real importance of detecting those minor CHDs may be questioned, unless they are associated with a chromosome aberration.

Published data show a strong association between CHDs and extra-cardiac malformations (Copel et al. 1988, Schwanitz et al. 1990, Allan et al. 1994, Carvalho et al. 2002, Stoll et al. 2002), emphasizing the importance of cytogenetic evaluation when CHDs are prenatally detected (Berg et al. 1988, Paladini et al. 1993). In the present study, as much as 38% of the major CHDs had abnormal karyotype and 17% associated malformations and normal karyotype. One large population-based study, the Baltimore-Washington Infant Study, found only 12% chromosome aberrations among live born infants (Ferencz et al. 1989). As we know that fetal populations include stillbirths and terminations of pregnancies where the incidence of CHDs associated with extra-cardiac malformations is high (Chinn et al. 1989, Tennstedt et al. 1999), a comparison between a fetal and an infant population will not be correct (Allan et al. 1994). When the results from the Baltimore-Washington Infant Study were adjusted for the effect of intrauterine lethality, the risk of aneuploidy for a fetus detected with CHD at the routine scan was found to be close to 40% (Berg et al. 1988), which is in accordance with the findings from the present study. Our data also showed a significantly higher association of chromosome aberrations among simple CHDs compared to complex CHDs, emphasizing the importance of detecting those simple CHDs. The results challenge our ultrasound skills as we detected significantly more of the complex than the simple CHDs.

The tendency to detect the most severe CHDs prenatally also reflects the outcome (Montaña et al. 1996, Boldt et al. 2002). As shown in Figure 19, the outcome was significantly better for the children where the heart defect was detected after birth, which supports previous findings (Allan et al. 1994, Bull 1999). The significantly
higher number of associated malformations and/or abnormal karyotype among the prenatally detected CHDs emphasizes the seriousness of those defects. The risk for spontaneous intrauterine death has been shown to increase in chromosomally abnormal fetuses (Hoffman 1995b). The fact that intrauterine fetal death occurred in 15% of the fetuses with a CHD detected prenatally (of which 63% had a chromosome abnormality), compared to zero fetal deaths among the postnatally detected, underlines the difference in seriousness between these two groups also in the present study. The same tendency was found among the terminated pregnancies where 58% of the fetuses had associated lethal chromosome aberrations or had developed a life-threatening condition at an early gestational age, and the rest, except for the Trisomy 21 cases, were severe anomalies requiring major surgery.

During recent years the prenatal detection of CHDs has improved and a trend towards more survivors has been found (Copel et al. 1997, Bonnet et al. 1999, Tworetzky et al. 2001, Brick and Allan 2002, Franklin et al. 2002). Those findings are encouraging in our effort to further increase the number of prenatally detected CHDs. With new surgical options and results, the long-term follow up will be of importance to give the best possible counseling when a major CHD is detected at the routine scan (Allan and Huggon 2004).

Despite years of persistent attempts to increase the prenatal detection rate at the second trimester routine examination, the fetal heart remains the most difficult organ to examine. Once prenatally diagnosed, the chance to survive until 2 years of age was only 27%, compared to 81% for those who were not detected prior to birth. The difficulties in showing improved outcome in prenatally detected CHDs may reflect the nature of the fetal cardiac disease that often is severe and with associated malformations (Yates 2004), as demonstrated in our study. Prenatal studies also tend to include those heart defects that have not been included in postnatal series because of death prior to surgery and a post-mortem diagnosis, influencing the outcome data (Abu-Harb et al. 1994, Yates 2004). In our effort to detect more of the major CHDs prenatally, we also need to focus on detecting these serious defects as early in the pregnancy as possible to allow
parents the choice of termination of the pregnancy (Allan and Huggon 2004). Also, when the pregnancy is continued, prenatal diagnosis will ensure optimal perinatal management and may influence the long-term morbidity (Eapen et al. 1998, Mahle et al. 2000, Yates 2004).


In fetal populations, the incidence of CHD is found to be higher than in live born populations (Rosendahl and Kivinen 1989, Hafner et al. 1998b). This may be due to the higher incidence of malformations in general in spontaneous abortions and stillbirths (Mitchell et al. 1971, Chinn et al. 1989). Figure 17 shows the increase in the overall incidence of CHDs in the present study from 11.1/1000 in Phase I to 14.7/1000 in Phase III. Interestingly, the incidences of major CHDs, 3–4/1000, were consistent and in accordance with the reported incidence of major CHDs (Ferencz et al. 1985). This emphasizes that the overall high incidences are not due to selection biases. The development of advanced ultrasound techniques has made it possible to detect minor CHDs today not detectable some years ago, either prenatally or postnatally. This is clearly demonstrated in Table 5 as both the incidence of isolated VSDs and ASDs have increased over time and are probably responsible for the increase in the incidence of minor CHDs during the last few years (Meberg et al. 1994, Roguin et al. 1995, Wren et al. 2000). It may be questioned whether these minor defects are true heart defects and not just a delay of a normal closure process towards the end of the pregnancy (Ben-Shachar et al. 1985, Hiraishi et al. 1992). The fact that none of these children needed treatment and that approximately 50% of the isolated VSDs closed spontaneously after
birth, indicate that we are observing a physiological process rather than pathology (Radzik et al. 1993, Meberg et al. 1994, Roguin et al. 1995, Helgason and Jonsdottir 1999).

Paper IV

Comparison between experienced and less experienced sonographer/midwives in obtaining fetal heart structures and detecting CHDs prenatally

During the study period, 29,035 fetuses had a second trimester routine examination performed by a sonographer/midwife. There was no difference in the experienced and less experienced midwives’ ability to obtain the four-chamber view at the routine scan (97%). Significant differences were found when the ability to obtain the ascending aorta (85% and 66%, respectively), the pulmonary artery (78% and 36%, respectively) or both arteries (75% and 36%, respectively) was evaluated (p < 0.001). Figure 20 shows the ability to obtain both the four-chamber view and the great arteries over time for the five experienced sonographer/midwives and the four less experienced sonographer/midwives who were new in this work in 1991, 1993 and 1995. A significant increase in the ability to obtain both views were found among the new, less experienced sonographer/midwives during the period from 1991 – 1995.

Figure 20 The ability to obtain the four-chamber view and the great arteries in 2994 fetuses, over time, for the experienced sonographer/midwives and the less experienced sonographer/midwives who started in 1991, 1993 and 1995, respectively.

* p < 0.0001
Of the 82 major CHDs in the study population, 35 (43%) were detected at the routine examination. The experienced sonographer/midwives detected 22/42 (52%) and the less experienced 13/40 (33%) (p = 0.07). Of the CHDs grouped as isolated, the detection rates were 8/18 (44%) and 6/22 (27%), respectively (p = 0.26), and for the CHDs with associated malformations 14/24 (58%) and 7/18 (39%), respectively (p = 0.21).

Of the 82 fetuses with a major CHD, 36 (44%) had a routine scan prior to 18 weeks. Of those, the experienced sonographer/midwives detected 11/22 (50%) of the CHDs prenatally, the less experienced 6/14 (43%). Of the 46 fetuses with a major CHD scanned after 18 completed weeks, the corresponding numbers were 11/20 (55%) and 7/26 (27%), respectively (p = 0.05).

Fifty-eight (71%) of the 82 major CHDs would be expected to have an abnormal four-chamber view at the time of the routine scan. Of these, 34/58 fetuses were examined by experienced sonographer/midwives and 28 (88%) of the abnormal views were recognized. The corresponding number for the less experienced sonographer/midwives was 13/26 (50%) (p = 0.002).

Of the isolated CHDs missed at the routine scan, 16/26 (62%) would be expected to have an abnormal four-chamber view: the experienced sonographer/midwives missed 5/10 (50%) and the less experienced 11/16 (69%). In 3/47 (6%) of the major CHDs missed, the sonographer/midwives had suspected a malformation at the routine scan, but the finding was not interpreted as pathology by the consulted physician. Nineteen (40%) of the 47 CHDs missed had a routine scan prior to 18 completed weeks.

Among the 19,237 fetuses scanned between 1995 and 2001, the percentage of fetuses with a four-chamber view not obtained at the second trimester routine examination decreased per year from 2.4% in 1995 to 1% in 2001. Unfavorable fetal position (41%) and poor imaging (40%) were the main reasons why the four-chamber view could not be obtained.
Comments:
The recent advances in ultrasound technology have placed great responsibility on the ultrasound operator. The different ultrasound techniques allow the operator to examine the fetus in more detail and to survey the fetus in better ways than only a few years ago (Eik-Nes et al. 1982, Eik-Nes 1993b, Blaas 1999, Hofstaetter et al. 2002, Brantberg et al. 2004). Proper training is necessary to utilize the potential of the ultrasound machine and to keep up with the new information, techniques and the development in the field of obstetric ultrasound.

Post-graduate ultrasound education and ultrasound courses are available in many countries, but are not mandatory for those who work with diagnostic ultrasound. With no requirements of education, it is the ultrasound operators’ responsibility to realize their own need for training. This has resulted in a large variation of background knowledge and experience among those working with obstetric ultrasound. Each sonographer/midwife in the present study received three months of basic ultrasound training, including a one-week basic theoretical course in addition to a minimum of 200 ultrasound scans supervised by an experienced sonographer/midwife.

The analysis of our results showed that 62% of the isolated major CHDs missed prenatally were expected to have an abnormal four-chamber view. Other studies also show that too many CHDs with an abnormal four-chamber view were missed (Wong et al. 2003). Although the detection rates increased when the great arteries were obtained, the detection rates did not reach the expected numbers for the four-chamber view alone (Stoll et al. 1998, Hunter et al. 2000). The lack of increase in the detection rate when more views were included into the fetal heart examination may indicate that other factors ought to be addressed.

Our study showed that experience played a significant role when it came to obtaining the fetal heart structures. In the mid-eighties, when the four-chamber view was introduced (Fermont et al. 1985, Allan et al. 1986b), studies demonstrated that this view could easily be incorporated into the fetal heart examination at the routine scan as it was
obtained in more than 95% of the fetuses (Copel et al. 1987, Achiron et al. 1992, Vergani et al. 1992, Tegnander et al. 1994, Todros et al. 1997). The same tendency was found in the present study where the four-chamber view was obtained in 97% of the fetuses. No difference was observed between the experienced and the less experienced sonographer/midwives, thus the four-chamber view seemed like an easy view to obtain no matter the degree of experience.

On the other hand, when we look at the isolated CHDs missed prenatally, 62% were expected to have an abnormal four-chamber view at the time of the routine scan. One may speculate if the four-chamber view had actually been properly obtained, and if so, whether it then had been correctly interpreted. Both the technique to obtain a correct four-chamber view and the way it is interpreted are a consequence of basic learning and experience. Ultrasound experience did seem to be of importance when it came to interpreting the view, as the experienced sonographer/midwives recognized significantly more of the expected abnormal four-chamber views than did the less experienced sonographer/midwives. But, the experienced sonographer/midwives also missed CHDs with abnormal four-chamber views, emphasizing the need for extra focus on this basic view (Chaoui 2003b).

The success in obtaining the sonographic views of the great arteries seemed to be more dependent on experience than on the ability to obtain the four-chamber view. The differences between the experienced and the less experienced sonographer/midwives were significant, and the main pulmonary artery seemed like the most difficult vessel to obtain. Since this registration was carried out, the three-vessel view has been introduced as an easy view to obtain the great arteries (Yoo et al. 1997), which may affect the way the great arteries are imaged today. Still, the significant difference between the sonographer/midwives clearly demonstrated the importance of experience. Even the experienced sonographer/midwives showed improvement of their results, most likely due to the extra focus on the four-chamber view and the great arteries (Figure 20). Thus, intensive focus and training on the examination of the fetal heart seem to be needed for all categories of ultrasound personnel. The poor detection rates of CHDs reported from
non-selected populations (Crane et al. 1994, Buskens et al. 1996, Bull 1999, Garne et al. 2001) may indicate that the basic training has not paid enough attention to the teaching of the four-chamber view and the great arteries of the fetal heart, and that there is potential for improvement.

Statistically, the differences found between the experienced and the less experienced sonographer/midwives were not all significant, but the consistent trend between them does suggest a difference. A comparison of the experienced sonographer/midwives’ ability to detect major CHDs to the less experienced sonographer/midwives’ results showed that the experienced personnel did better. When breaking the CHDs down to isolated CHDs, ductal dependent CHDs and CHDs with associated malformations including abnormal karyotype, we found the same tendency, i.e. that the experienced sonographer/midwives did better than the less experienced sonographer/midwives. Because heart defects are low in number, a background population of three times the size of our population, approximately 100,000 fetuses, would be needed to reach a level of significance given the same detection rates. However, the experienced sonographer/midwives did significantly better in obtaining both the four-chamber view and the great arteries – details necessary for the detection of CHDs. In addition, the experienced sonographer/midwives’ ability to recognize an abnormal four-chamber view was significantly better than that of the less experienced sonographer/midwives. The significant improvement in the ability to obtain fetal heart structures over time also emphasizes the difference between the experienced and less experienced sonographer/midwives.

The gestational age may be of importance for the detection of CHDs. Several studies have shown that the fetal heart structures are more difficult to image prior to 18 weeks than after 18 weeks (Schwärzler et al. 1999b). As much as 40% of the fetuses with a major CHD missed prenatally in the present study had a routine scan performed prior to 18 completed weeks. According to published data, some of these might have been detected if the ultrasound examination had been carried out at a later gestation (Allan 2003). The results from Paper I demonstrating a higher success in obtaining the four-
chamber view after 18 weeks, still seem to be valid. Despite the advances in technology, it is interesting to notice that those offering both an early and second trimester ultrasound examination prefer to do the second trimester anomaly scan between 20-24 weeks (Gembruch et al. 1993, Zosmer et al. 1999, Rustico et al. 2000). It is worth noting, though, that the gestational age most likely is only one of several factors involved in the poor detection of CHDs. There were no differences between the experienced and the less experienced sonographer/midwives when it came to detecting CHDs prior to 18 weeks of gestation, but the difference was significantly better for the experienced sonographer/midwives after 18 completed weeks. It may be that the less experienced sonographer/midwives had problems detecting the CHDs no matter what the gestation, and that experience still is important even at later gestations.

To further improve the results, it has been suggested that additional views of the fetal heart and new modalities be incorporated to the fetal heart examination (Yoo et al. 1997, Chaoui and McEwing 2003). Adding the great arteries to the four-chamber view clearly increased the detection rate in the present study. The present study also showed that the ability to obtain the four-chamber view at the routine scan increased to 99%, but still too many CHDs with an abnormal four-chamber view were missed. Clearly, the four-chamber view was evaluated as normal when it was not. Also, too many CHDs with a normal four-chamber view were missed, underscoring that the examination of the great arteries was not sufficiently mastered. It may be that the potential of including the great arteries can not be fully exhausted as long as the four-chamber view is not properly mastered. The stepwise introduction to the fetal heart examination, suggested by Allan in 1986, still seems to be fundamental to the success in fetal heart scanning. The first step, the four-chamber view, should be mastered before the next step is considered (Allan 1986, Sharland 2004). Adding even more views to the fetal heart examination may not give the expected results as long as the basic views have not been properly learned.

The present study emphasizes the importance of dedicated personnel performing the obstetric ultrasound examinations, as experience seems crucial for the detection of
defects, specifically of the fetal heart. In most non-selected populations, sonographers with basic ultrasound training are performing the routine examinations. After years working with ultrasound, sonographers develop an awareness to pathology gained from intensive experience (Cullen et al. 1992). This was also shown in the present study as three true CHDs were suspected by the sonographer/midwives, but later evaluated as normal by the consultant physician. It takes years to develop this experience; thus, fetal ultrasound examinations should be performed by personnel specifically trained in obstetric ultrasound (Cullen et al. 1992). Non-dedicated personnel will probably not get enough experience and will, over time, remain less experienced ultrasound personnel.

The long learning curve to become experienced in fetal heart scanning may be shortened by a more focused education on the four-chamber view and the great arteries. The educational and training systems today expect the sonographers to build up their experience through a high number of ultrasound examinations following their training. The busy schedule most sonographers are faced with does not give much room for developing scanning skills; this is demonstrated by the long learning curve. Since studies have shown that the fetal heart is more difficult to examine than other fetal organs (Stoll et al. 2002a, Levi et al. 2003), the educational sites have to accept that the heart needs more focus during the time of education. With a better basic learning process, the learning curve may increase faster.

What have we learned during the 20 years since the sonographic appearance of the fetal heart and the subsequent demand for increasing the detection rates of CHDs? During all these years the ultrasound equipment has improved significantly, making it easier to obtain the details of the fetal heart, but also requiring more technological knowledge from the ultrasound operator. Already in the late eighties (Paper II, Phase II) we demonstrated the importance of utilizing the correct equipment when examining the fetal heart: using a 5 MHz transducer gave a significantly better image of the four-chamber view than the 3.5 MHz transducer. Although most modern ultrasound machines are equipped with multifrequency transducers, a proper image of the fetal heart still requires the ultrasound operator’s knowledge and awareness of correct use.
and adjustments of the ultrasound machine; these need to be optimized for the imaging of the fetal heart.

In continuation of the discussion on equipment, the poor detection rates of CHDs may also be a result of old and poor ultrasound equipment. The fact that most of the fetuses with CHD are found in non-selected populations (Allan et al. 1994, Maher et al. 1994, Carvalho et al. 2002, Åmark et al. 2004) means that the detection of fetuses with CHD needs to be done in the routine setting. This scan is often the only opportunity to detect CHDs prenatally, as ultrasound examinations later in the pregnancy often are done on indications only. If the ultrasound machines used during the routine scan are of poor quality, this may affect the detection rate of defects. This is an organizational issue that should be addressed by each ultrasound department.

Experience is of importance for the detection of CHD (Cullen et al. 1992, Hunter et al. 2000, Carvalho et al. 2002, Wong et al. 2003). When the four-chamber view was introduced and evaluated (Paper II, Phase II), a significant improvement in the ability to obtain this view was observed over time. The same results were found in the present study with the introduction of the great arteries. For both periods, the learning curve was relatively long, despite better ultrasound equipment during the present study. Thus, good equipment is not the only answer to better imaging of the heart; experience must also be considered. The experienced sonographer/midwives were superior to the less experienced sonographer/midwives in obtaining details and detecting defects of the fetal heart. Thus, if only experienced sonographer/midwives performed the routine examinations, one would expect a further increase in the prenatal detection rate of major CHDs. The opposite situation is often the reality in ultrasound departments. Experienced sonographer/midwives tend to end up with administrative work while less experienced sonographer/midwives are left on their own to do the majority of the scans to gain experience. Since there always will be a need for recruitment of new personnel, the basic education must focus on the need for more experience in fetal heart scanning.
To provide the best possible fetal heart scans, we must apply quality assurance to the work we do. This has already been addressed by the Fetal Medicine Foundation in London. The Fetal Medicine Foundation requires minimum standard for the ultrasound examination, and has shown the potential of nuchal translucency measurement in detecting developmental disorders when this measurement is performed with a high degree of accuracy and quality assessment (Nicolaides et al. 1999). The importance of complying with a detailed procedure when measuring the nuchal translucency has been encouraged in numerous courses around the world to ensure that this accurate and standardized method is followed. Multiple series have shown that focusing on nuchal translucency measurements as taught by the Fetal Medicine Foundation has good results (Hafner et al. 1998, Hyett et al. 1997, Schwärzler et al. 1999). This way of standardization and assessment of nuchal translucency measurements probably also applies to other aspects of the fetal examination. Based on results from the present study there is reason to believe that a standardized procedure for the fetal heart examination should be given increased focus. Images of the nuchal translucency that do not follow the standardized method are not accepted by the Fetal Medicine Foundation quality assessment system. There is no such quality check when it comes to images from the fetal heart. If the examination of the four-chamber view with all its details described 20 years ago (Allan 1986) had been followed up by the same rules and quality assessment as for the nuchal translucency measurement, this would most likely have made a great impact on the examination of the fetal heart and thus the detection of CHDs.

Experience shows that it takes time to learn the ultrasound technique in general. As a consequence, most post-graduate ultrasound educational programs last for a minimum of one year. During that year, the basic ultrasound technique has to be learned and the ultrasound images must be understood and correctly interpreted. Obtaining correct images is dependent on the operator’s understanding and mastering of the ultrasound technique. The poor detection rates reported on CHDs may suggest that the basic technique has not been properly understood and learned. The educational sites’ future focus should be on each ultrasound operator’s skills to obtain and interpret sections through the fetal heart; time must be allowed for necessary maturation of these
ultrasound skills. The technique to obtain the correct four-chamber view and the interpretation of all the details from this view have to be mastered before the great arteries are added. As the basic ultrasound education includes a range of different subjects to be learned, the great arteries may be introduced through additional ultrasound courses following the first year of training, perhaps as an obligatory extension of the basic education.

Paper V

Introduction of three-dimensional ultrasound imaging of the fetal heart (TDOG)

A new 3D technique utilizing tissue Doppler as a gating signal was introduced. The purpose of this technique was to see if structures in the fetal heart could be more easily obtained and imaged than was possible with the conventional 2D ultrasound technique. The images were of adequate quality for four of the eight fetuses examined. Two fetuses showed several motion artifacts and for two fetuses there was no adequate access to the four-chamber view during the scanning time. To cover the fetal heart with maximal diameter of 2 cm at a depth of 5–7 cm, a scanning angle of approximately 25° showed good agreement between the B-mode and a planar slice.

The best image quality of the 3D reconstruction was achieved by a compromise between high B-mode quality and high frame rate. Second harmonic imaging gave the highest lateral resolution and was used on all recordings. As for the gating signal, cardiac motion could positively be used, even when the motion was not visible within the B-mode image. The basic signal used for the tissue Doppler synchronization emphasized strong echoes and massive motion of tissue in the ultrasound beam direction.

The variation in the quality of the reconstructions between different recordings were mainly due to fetal position not suitable for high-quality B-mode imaging of the fetal heart, fetal movements during the sweep and uneven probe motion during the sweep.
Figure 21 demonstrates reconstructed volume projections of the heart from a 24-week fetus. All images were retrospectively produced from one cineloop of 3D data, derived by apical insonation.

**Figure 21** Volume projection of the heart of a 24 weeks fetus. a) Apical four-chamber view. b) Subcostal four-chamber view. c) Ascending aorta. d) Three-vessel view. e) Short-axis view of the great arteries. f) Aortic arch. g) Ductal arch. h) Long-axis view of inferior and superior vena cavae.

Comments:
Dynamic 3D imaging of the fetus has received increasing interest during the past few years (Hata et al. 1998, Pöhls and Rempen 1998). This may be a natural result of the rapid development of technology and imaging modalities to meet the demand of increased image quality and thus the information derived from those images. The interest for dynamic 3D imaging of the fetal heart may also be a result of poor detection rates of major CHDs and the recognition of the fetal heart as a difficult organ to examine with 2D ultrasound imaging (Chaoui 2003b, Sharland 2004).

The acquisition techniques with 3D ultrasound imaging have been difficult, and different approaches have been investigated, such as using both gated and non-gated B-mode reconstruction techniques (Merz et al. 1995, Sklansky et al. 1998) or real-time 3D acquisition using a 2D array transducer (Deng et al. 2002a). Different gating techniques
have also been studied, including operator input based on an M-mode display (Deng et al. 1996), temporal Fourier transforms of the volume data itself (Sklansky et al. 1998) and Doppler spectrum acquisition and analysis performed by an independent ultrasound system (Deng et al. 2000, Deng et al. 2002b). The spatio-temporal image correlation (STIC) technique has been used to produce 3D gray scale images with the gating signal automatically detected from the underlying B-mode sequence (Devore et al. 2003, Viñals et al. 2003), and to reconstruct 3D color Doppler images (Chaoui et al. 2004, Gonçalves et al. 2004). 3D images of the fetal heart have also been obtained by free-hand scanning gated with cardiotocography (CTG) (Herberg et al. 2003).

The current study looked into another approach, using 2D tissue Doppler from the fetal heart to generate the gating signal (TDOG). Tissue Doppler has been used for clinical application since 1989, first described by Isaaz et al. (1989) to measure the low velocity motion of the left ventricular posterior wall. The higher velocity in blood than in other tissue during normal circumstances results in a lower Doppler shift from the myocardial wall than in blood. The tissue signal will have a stronger amplitude because of more scatter returning from tissue than from blood. Thus, the tissue Doppler signal can be recorded either by low pass filtering or gain damping, or with a combination of both (Brodin 2004). For the TDOG method, those strong tissue Doppler signals could positively contribute to the gating signal.

Spatial and temporal tracking, or gating, is usually preferred when 3D images are to be acquired in-utero, during which fetal movements, maternal respiration or probe-induced movements often occur (Deng 2003). Nongated cardiac imaging generates a single volume that mixes together several parts of the cardiac cycle, resulting in suboptimal accuracy and resolution; the gated techniques have shown to improve the resolution of image planes generated from volumetric data (Sklansky et al. 1998). Electrocardiogram used for gating in 3D echocardiography in infants, children and adults has been difficult to achieve in fetuses, although gating by cardiotocography in the fetus has been attempted (Herberg et al. 2003).
The disadvantages of using two different systems are the inconvenience of two devices, in addition to possible artifacts caused by acoustic interference between the transducers (Deng et al. 2000, Downey et al. 2005). Such acquisition artifacts were noted during the present study when Doppler CW data were recorded simultaneously with the tissue Doppler for quality assessment. To ease the use for clinical setting, the gating signals should be easy to obtain, accurate and technically reliable. With the TDOG-method, tissue Doppler data can be acquired simultaneously and with the same transducer as the B-mode data, allowing the recording of tissue Doppler data as part of a gated 3D scan.

Handheld sweeps allow the sonographer to make fine adjustments during the sweep to minimize acoustic shadowing, but the image quality depends largely on how smoothly and steadily the transducer is moved (Downey et al. 2000, Sklansky 2004). A long sweep yields a high scan plane density as a high number of consecutive cineloops become neighboring planes in the 3D dataset. However, too long a sweep makes it difficult to perform the sweep with constant tilt velocity. A method in which the transducer is kept stationary and electronic scanning with a 2D array processes the echoes to generate 3D information may be an alternative to freehand systems (Downey et al. 2000), used for the STIC technique (DeVore et al. 2003, Viñals et al. 2003). However, the STIC-technique is a post-processing technique and does not allow 3D in real-time scanning, i.e. 4D scanning.

Acquiring volumetric data within a few seconds from a single window, with the possibility of a retrospective visualization of the heart in a multiplanar display is of interest. Such a procedure reduces the scanning time, is independent of the experience of the ultrasound operator, allows reconstruction of standardized fetal echocardiographic views without the pregnant woman present, and allows 2D reconstruction of views from the fetal heart not accessible by regular 2D B-mode imaging (Budorick and Millman 2000, Sklansky 2004). On the other hand, there are also limitations to 3D ultrasound. The present study showed that important factors for an optimal and useful recording were: no bone shadows or other disturbances in front of the fetal heart, the heart positioned within a certain distance from the ultrasound.
transducer (ideally less than 6 cm), and no fetal or respiratory motion during the sweep time. These limitations well known from 2D ultrasound (DeVore et al. 1993, Hendler et al. 2005), also recognized from 3D imaging (Budorick and Millman 2000, Hull et al. 2000, Abuhamad 2005), should be expected, as long as the 3D reconstructions are based on 2D recordings.

Performance of the 3D scan by less experienced ultrasound personnel may be questioned. Difficult fetal position, which can affect the acquisition of the data, can be dealt with by using an alternative insonation angle gained through experience. Data from Paper IV show that it takes time and experience to obtain adequate images of the fetal heart needed for 3D reconstruction. In addition, the acquisition, display and manipulation of 3D volumes require experience (Abuhamad 2005). Thus, as the utilization of the multiplanar review may be time-consuming, the timesaving factor may not be obvious (Budorick and Millman 2000). Standardization in the acquisition and display of 3D volumes may ease the training process (Abuhamad 2005).

There are advantages and disadvantages with both 2D imaging systems and 3D imaging systems. Some will remain the same for both imaging modalities as long as 2D images are acquired to reconstruct 3D images. The present paper demonstrated the weakness of the ultrasound system due to poor resolution when the perimembranous part of the ventricle septum was to be imaged from an apical view of the heart. The nature of ultrasound physics cannot be overruled, thus one single 3D ultrasound insonation angle will not give an optimal image quality of all structures, regardless of resolution. A typical example would be the imaging of the ventricular septum. In the apical view using 3D ultrasound, it still would appear very thin (Figure 21), whereas in the long-axis view, true imaging of the texture and thickness would be appropriate.

Today most 3D ultrasound images represent a compromise between the optimal diagnostic perspective (large data set, higher-resolution 3D images) and what the computers actually are capable of processing (Downey et al. 2000). With the increased
clinical use of 3D ultrasound and further advances in computer technology, 3D ultrasound may play an important role in obstetric ultrasound.
SUMMARY

Background
The second trimester fetal ultrasound examination has become an integrated and important part of prenatal care. One of the aims of this scan is to detect developmental disorders which might have consequences for the management of the pregnancy and the neonatal period. The fetal heart has received particular attention, as heart defects seem difficult to detect prenatally and data show when CHDs are prenatally detected, that the outcome improved. As a teaching institution, our focus has been on improvement in the fetal heart examination. The knowledge about what kind of CHDs we do and do not detect is of importance in order to understand why we fail to detect CHDs and to learn about the next steps needed to improve the results.

Purpose
The objectives of the studies were to implement and prospectively evaluate new examination procedures of the fetal heart at the second trimester routine examination, in order to increase the prenatal detection rate of CHDs (Papers I, II, III and V). The training and experience of the ultrasound personnel were evaluated to see if educational aspects were factors that would have influence on the prenatal detection of CHDs and to evaluate current ultrasound-training programs (Paper IV). Finally, the CHD cases were followed into the postnatal period to get a better understanding of the outcome, in order to improve the basis and the accuracy for our counseling once the defects are detected (Papers II and III).

Material and methods
A well-defined non-selected population of 42,381 fetuses was prospectively studied from August 1986–December 2001. Of those, 41,354 (98%) received an ultrasound examination at our hospital and formed the study population (Papers I, II, III and V). The second trimester routine ultrasound examinations were scheduled at 18 completed weeks and were mainly performed by specially trained sonographer/midwives. In Paper
IV, these sonographer/midwives were grouped as experienced or less experienced and constituted the study population.

During the regular second trimester routine scan, the four-chamber view was incorporated into the fetal heart examination (Papers I and II), and later the views of the great arteries were added (Paper III). Starting in 1995, a new scan was offered at approximately 20 weeks of gestation if a proper four-chamber view had not been seen at the routine examination (Papers III and IV). In 8 of these fetuses, a new method to image the fetal heart, Tissue Doppler gated (TDOG) 3D ultrasound, was evaluated (Paper V).

A heart defect was retrospectively classified as major or minor after a final diagnosis was achieved. The major CHDs were grouped as isolated; abnormal karyotype; associated malformations (Papers III and IV). Fetal karyotyping was offered in all prenatally detected cases. Every fetus/child was followed-up to ensure outcome data. The longest period of follow-up for the live-born children was 13 years (Paper III).

Results
A four-chamber view of the fetal heart could be obtained in 96% of the fetuses at 18–21 weeks of gestation, the time of the second trimester routine scan. Prior to 18 weeks, the results were poorer. The results with a 5 MHz transducer were significantly better than with a 3.5 MHz transducer. A significant improvement in the ability to obtain the four-chamber view was noticed over time for the sonographer/midwives performing the ultrasound examinations (Paper I).

Introducing the four-chamber view into the routine examination increased the detection rate of major CHDs at that scan from 18 to 26% (Paper II). By adding the views of the great arteries, the detection rate further increased to 39% (Paper III). During this last period (Paper III), 9% of the major CHDs in the study population were detected prior to the routine scan.
An abnormal karyotype was found in 38% of the 97 major CHDs, and 18% had associated malformations with normal karyotype (Paper III). Major CHDs with associated malformations including abnormal karyotype were significantly more often detected than were isolated major CHDs. Of the isolated ductal dependent major CHDs, 48% were detected prenatally. Of the isolated major CHDs missed prenatally, 58% would be expected to have an abnormal four-chamber view.

The incidence of major CHDs was constant between 3.0–3.7/1000 through the complete study period. The incidence of minor CHDs increased from 7.4–11.3/1000 through the same period (Papers II and III).

There were significant differences in the outcomes between the fetuses with a CHD detected prenatally compared to postnatally (Paper III). Of the prenatally detected cases, 44% of the pregnancies were terminated. Of those, 58% had associated lethal chromosome aberrations, and the rest, except for the trisomy 21 cases, were severe anomalies. Intrauterine fetal death occurred in 15% of the fetuses with a detected CHD, but in none of those not detected prenatally. Of the prenatally detected cases, 42% were born alive. At the age of 2 years, only 27% of the children with prenatally detected CHDs were alive. The corresponding numbers for those detected postnatally were 98% and 81%, respectively.

Compared to the less experienced sonographer/midwives, the experienced sonographer/midwives showed significantly better results in obtaining a view of the great arteries and in recognizing an abnormal four-chamber view (Paper IV). The experienced sonographer/midwives did better in the overall detection of major CHDs, as well as in the detection of isolated CHDs and CHDs with associated malformations. The less experienced sonographer/midwives showed a significant improvement over time in their ability to obtain the heart structures and reach the level of the experienced sonographer/midwives. CHDs with an abnormal four-chamber view were missed in both groups.
With the new 3D technique utilizing tissue Doppler as a gating signal, it was possible to obtain images of adequate quality in four out of eight fetuses and reconstructed volume projections of the heart from one cineloop of 3D data could be demonstrated (Paper V). Two fetuses showed several motion artifacts; in two fetuses, a four-chamber view could not be obtained due to awkward fetal position.

Conclusions
The fetal heart examination, and thus the prenatal detection of CHDs, is still a challenge in non-selected populations. Incorporation of the four-chamber view and the great arteries increases the detection rate to some extent, but not as much as expected. The fetuses with the most severe CHDs and associated malformations seem easier to detect than fetuses with isolated CHDs.

Education and training seem crucial for the ability to obtain fetal heart structures. Obtaining and interpreting basic views of the fetal heart failed for all sonographer/midwives, emphasizing that the basic education on the fetal heart examination needs to be strengthened. Both the views of the four chambers and the great arteries must be mastered before new views and advanced technology are added.

Future aspects
Recent data show that results for newborns with a heart defect detected prenatally are improving. To detect the major CHDs, the sonographer/midwife must master the 2D ultrasound technique as well as the evaluation of the four-chamber view and the great arteries. Thus, the fetal heart examination must be given special attention in the education and training of sonographer/midwives. With a thorough basic understanding and experience, advanced ultrasound techniques may be more easily adopted.

Recent studies also indicate that the 11–13+6 week scan might be of importance for the overall detection of heart defects during pregnancy.
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severe cardiac malformations by 2D echocardiography. Presented at the 2nd World congress of Paediatric Cardiology New York, p. 10.


CORRECTIONS

Paper II  Page 376, second column, line six:
“Eighteen percent of the critical and 11%...” should be “Fifteen percent of the critical and 11%...”.

Page 377, Table 10, first line under “Outcome”:
“Intrauterine fetal death” should be “Alive, died 2 days old, Trisomy 18”.
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2003
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2004
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