Doppler ultrasound in pregnancy – quality requirements of DEGUM and clinical application (part 2)
Dopplersonografie in der Schwangerschaft – Qualitätsanforderungen der DEGUM und klinischer Einsatz (Teil 2)

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ABSTRACT
This second part on Doppler sonography in prenatal medicine and obstetrics reviews its clinical applications. While this has not become the initially anticipated screening tool, it is used for the diagnosis and surveillance of a variety of fetal pathologies. For example, the sonography-based determination of uterine artery blood flow indices is an important parameter for the first trimester multimodal preeclampsia risk assessment, increasing accuracy and providing indication for the prophylactic treatment with aspirin. It also has significant implications for the diagnosis and surveillance of growth-restricted fetuses in the second and third trimesters through Doppler-sonographic analysis of umbilical artery, middle cerebral artery and ductus venosus. Here, especially the hemodynamics of the ductus venosus provides a critical criterium for birth management of severe, early-onset FGR before 34 + 0 weeks of gestation. Further, determination of maximum blood flow velocity of the middle cerebral artery is a central parameter in fetal diagnosis of anemia which has been significantly improved by this analysis. However, it is important to note that the mentioned improvements can only be achieved through highest methodological quality. Importantly, all these analyses are also applied to twins and higher order multiples. Here, for the differential diagnosis of specific complications such as TTTS, TAPS and TRAP, the application of Doppler sonography has become indispensable. To conclude, the successful application of Doppler sonography requires both exact methodology and precise pathophysiological interpretation of the data.

ZUSAMMENFASSUNG
**Indications**

The technical principles and quality criteria for performing Doppler sonography in pregnancy have been recently defined [1].

Primary indications for the use of Doppler sonography are listed in the Maternity Guidelines (► Table 1). In addition, there are now further indications that are contained in the present recommendations (► Table 2).

**Screening**

Numerous studies in the 2nd and 3rd trimesters have demonstrated that increased uterine artery indices (► Fig. 1a, b) are associated with more frequent occurrence of preeclampsia, fetal growth restriction (FGR), premature placental abruption, stillbirth, etc. [2–4]. Measurement of the uterine arteries in "low-risk" cohorts between 18+0 and 24+0 gestational weeks resulted in sensitivities of 40–60 % with a false-positive rate around 10 %. However, the positive predictive values of 25 % for predicting FGR and/or preeclampsia were very low [5]. Nevertheless, the method is suitable for exclusion diagnostics due to the negative predictive value of 97–99 %. If the Doppler of the uterine artery is unremarkable, the risk of developing severe preeclampsia is very low, and further management of the pregnancy can be modified accordingly [6].

In "high-risk" cohorts, the results were better (sensitivity 70–80 %, PPV 50 %) [7]. However, no effect on perinatal outcome could be demonstrated [8].

At the time of first trimester screening (11+0–14+0 gestational weeks), the accuracy of predicting preeclampsia by Doppler ultrasonography of the uterine arteries alone is low; detection is 30–40 % with an FPR of 10 %. When using a multimodal algorithm (Fetal Medicine Foundation, London) that incorporates other anamnestic, biophysical (e.g., maternal BMI and blood pressure), and biochemical (e.g., PIGF, PAPP-A) parameters, the accuracy of predicting early preeclampsia (<34+0 gestational weeks) improves tremendously; detection rate 90–95 %, FPR 10 % [9–12]. This form of PE screening in first trimester screening is clinically relevant because prophylactic administration of ASA before 16+0 gestational weeks can significantly reduce the occurrence and severity of early preeclampsia [13]. Depending on the aspirin dose, the incidence of severe PE can be reduced to one-quarter, and the total number of PE can be reduced to one-half on average [14, 15]. The recommended dose is 150 mg ASA per day in the evening [15].

The technique of measuring the uterine arteries in the 1st trimester (► Fig. 2a, b) differs compared with the 2nd and 3rd trimesters and should meet certain quality criteria [16]. In particular, the measurement must be taken close to the cervix in the main branch of the uterine artery. Pulsatility at this time is significantly higher than in the 2nd trimester, and in the majority of pregnancies notching is still detectable, so this alone is not pathologically significant [17].

Transferring the multimodal screening concept to the 2nd and 3rd trimesters also achieves a high detection rate of 85 % with a false-positive rate of 10 % for early preeclampsia [18]. However, it should be noted that this means that the preventive effect of ASA treatment is significantly lower and that the screening benefit is the adaptation of the monitoring intervals and the time of delivery.

The umbilical artery (► Fig. 3) is not suitable for screening for FGR, preeclampsia, or other pathologies [19].

**Monitoring of high-risk pregnancies**

The importance of the use of Doppler sonography in obstetrics must ultimately be measured by the prevention of complications and the reduction of perinatal or neonatal morbidity and mortality.

In principle, a distinction must be made between pregnancies with pathological Doppler values without vs. with clinical fetal (FGR) and/or maternal symptoms (preeclampsia). Pathological uterine Doppler findings (► Fig. 1, 2) are almost always measurable long before the development of clinical symptoms (early warning effect), while pathological fetal Doppler findings (► Fig. 3) are practically always detectable shortly before or with the onset of complications (e.g., FGR).

If it is already assumed that measurement of the uterine arteries is becoming increasingly important, especially in the context of first trimester screening for risk assessment of various complications, the question arises whether monitoring of pathological findings is useful in the further course of pregnancy. There is

**Table 1** Indications for the use of Doppler sonography according to the Maternity Guidelines.

<table>
<thead>
<tr>
<th>Indication</th>
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<tbody>
<tr>
<td>suspicion of fetal growth restriction (FGR)</td>
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<tr>
<td>pregnancy-induced hypertension, preeclampsia</td>
</tr>
<tr>
<td>status post FGR or IUFT</td>
</tr>
<tr>
<td>status post preeclampsia/eclampsia</td>
</tr>
<tr>
<td>abnormalities of the fetal heart rate (CTG)</td>
</tr>
<tr>
<td>suspicion of malformation/fetal disorder</td>
</tr>
<tr>
<td>multiple pregnancy with discordant growth</td>
</tr>
<tr>
<td>suspicion of heart defects or fetal arrhythmias</td>
</tr>
</tbody>
</table>

**Table 2** Additional indications for the use of Doppler sonography according to the DEGUM (German Society for Ultrasound in Medicine) recommendations.

<table>
<thead>
<tr>
<th>Indication</th>
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<tbody>
<tr>
<td>pre-existing maternal vascular diseases (hypertension, nephropathy, D. mellitus, auto-immune diseases)</td>
</tr>
<tr>
<td>diagnosis and monitoring of fetal growth restriction (FGR)</td>
</tr>
<tr>
<td>suspicion of fetal anemia</td>
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<tr>
<td>suspicion of fetal infection</td>
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<tr>
<td>fetal hydrops</td>
</tr>
<tr>
<td>monitoring of monochorionic multiple pregnancies (FFTS (TOPS, TAPS), TRAP, sFGR)</td>
</tr>
<tr>
<td>diagnosis of pathological umbilical cord insertions and vasa previa</td>
</tr>
<tr>
<td>diagnosis of placental disorders (e.g. placenta increta)</td>
</tr>
</tbody>
</table>

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clear evidence that the adaptation of uterine perfusion to the needs of pregnancy continues well beyond the 1st half of pregnancy [17, 20]. That is, pathological findings in first trimester screening (▶ Fig. 2) may normalize and formerly normal findings in this screening may become pathological. Since this has prognostic significance for the individual pregnancy [17], we recommend monitoring uterine perfusion between 20+0 and 24+0 weeks.

In case of persistence of pathological uterine Doppler findings according to a “Uterine Artery Score” (UAS; PI of both uterine arteries > 1.4 or 1.2 or notch) [21], a check-up between 26\(^{+0}\) and 28\(^{+0}\) weeks (biometry, Doppler, placenta) is recommended to detect early-onset deterioration of fetal hemodynamics in a timely manner [21, 22]. The perinatal risk profile increases significantly if more than 2 of the above parameters (Uterine Artery Score = UAS) are pathological in the sum of both sides [23]. At a UAS of 4, weekly clinical and Doppler sonographic checks are recommended. In this case, supplemental serum screening by sFlt-1/PlGF ratio is also discussed, but generally accepted management protocols are still pending [24].

Employment of fetal Doppler sonography usually begins with the onset of complications to assess fetal status. This can be as early as the first trimester (e.g., ductus venosus in FFTS in multiples), but usually becomes necessary after 20\(^{+0}\) weeks (FGR, PE, anomaly, anemia).

The most important vessel is the umbilical artery (UA), for which there is clear evidence regarding its benefit for perinatal outcome [25]. It is the first vessel in the temporal cascade to assess fetal status [26], which is complemented by measurement of the middle cerebral artery (MCA) and ductus venosus (DV) when the umbilical artery becomes pathological (▶ Fig. 3).

In recent years, it has become clear that early severe FGR differs from late FGR in the cascade of Doppler sonographic changes [26–28]. In the early form, as the pathology increases,
Doppler indices in the umbilical artery (▶ Fig. 3–5), later in the middle cerebral artery (▶ Fig. 6), and just before fetal decompensation, those of the venous duct (▶ Fig. 7) become conspicuous. According to the TRUFFLE study, the decision criterion for delivery before 34+0 weeks is considered to be negative or reversal of the a-wave in the ductus venosus (▶ Fig. 7) in addition to the STV in the cCTG (short-time variability in computer-supported CTG) [29]. After the appearance of pathological Doppler indices in the umbilical artery (▶ Fig. 3), individualized monitoring begins, which should now be performed by a specialist at shorter intervals after 1–2 weeks. Many different fetal parameters (biometry, amniotic fluid volume, middle cerebral artery, DV, cCTG, biophysical profile, gestational weeks) as well as maternal conditions (history, symptoms) play a modifying role, but in particular the development of the individual Doppler parameters over time (no, slow or rapid deterioration) [26], which also determine the need for inpatient monitoring and the administration of antenatal steroids for lung maturation prophylaxis [30]. Using a cerebro-placental ratio (CPR) could further improve the prediction of the outcome of perinatal disorders [31].

With the more common late fetal growth discrepancy in the third trimester, a distinction must be made between SGA (small for gestational-age) and true FGR (fetal growth restriction). This is only possible with multimodal approaches in which Doppler findings play an important role [31, 32].

In this late period, the end of adaptive mechanisms is reached with fetal circulatory centralization in the middle cerebral artery, which is best documented by the cerebro-placental ratio (CPR) [31, 33, 34]. Prediction of perinatal outcome disorders by CPR is good for individual parameters in FGR [31], perhaps successful multimodally in SGA fetuses [31, 35], and inadequate in unselected pregnancies [36, 37]. This could perhaps also be related to methodological problems and a lack of quality criteria for measur-
ing CPR to date [38]. The umbilical artery and venous duct may also show elevated indices, but are usually not highly pathological (e.g., zero flow, reverse flow) [30, 34].

Doppler sonography in fetal anemia

Direct and definitive diagnosis of fetal anemia is made by cordocentesis, an invasive procedure that results in fetal loss due to the procedure in 1% of cases, depending on gestational age. Doppler sonographic detection of elevated peak systolic velocity (PSV) in the middle cerebral artery is currently the best method to non-invasively diagnose fetal anemia and to serially monitor the fetus if the risk is appropriate (▶ Fig. 8) [39]. In anemia, there is an increase in blood flow rates in all fetal vessels—arteries and veins—as a result of increased cardiac output and decrease in blood viscosity. Primarily for technical reasons, peak systolic velocities (PSV) in the MCA are best suited to detect the presence and extent of anemia, regardless of its etiology. The decrease in hemoglobin and pCO₂ correlate with MCA-PSV, but not the pO₂ [40]; fetal anemia is not a hypoxemia-triggered blood redistribution, because pO₂, pCO₂ and pH are normal, as are the Doppler indices (PI, RI) or the Doppler spectrum in the arterial and venous vessels.

At this point, it should be mentioned with respect to differential diagnosis that even with severe and early onset of fetal growth restriction, not only is the pulsatility in the middle cerebral artery reduced, but PSV can also be raised [41]. In late pregnancy, fetal hypercapnia may occur in the setting of progressive placental insufficiency with prolongation of the villous diffusion distance, which may also lead to flow acceleration in the middle cerebral artery [40]. In this case, the diagnostic value is therefore limited.

Other less sensitive signs of fetal anemia may include: cardiomegaly with biventricular dilatation, hepatomegaly, splenomegaly, increased echogenicity of the bowel, polyhydramnios, placentomegaly, and hydrops fetalis [42, 43].

Fetal anemia is present when hemoglobin concentration and/or hematocrit are below two standard deviations of the gestational age-related reference range. Since both increase with increasing gestational age (mean fetal hemoglobin concentration is 10.6 g/dL at 18<sup>+</sup>0 gestational weeks, 12.0 g/dL at 24<sup>+</sup>0 weeks, 12.8 g/dL at 30<sup>+</sup>0 weeks, and 13.8 g/dL at 40<sup>+</sup>0 weeks) [39], hemoglobin concentration is calculated as the multiple of median (MoM) relative to gestational age, so that the extent of anemia can be specified independently of gestational age (mild anemia: MoM 0.83–0.65; moderate anemia: MoM: 0.64–0.55; severe anemia: MoM <0.55) [39].

Since MCA-PSV increases during pregnancy from 20–30 cm/s at the end of the first trimester to 50–70 cm/s in the third trimester, the PSV (cm/s) is also converted to MoM to express the extent of deviation of PSV from the mean value independent of gestational age [39].

To realize the diagnostic potential of Doppler sonography of MCA-PSV, the measurement must be performed extremely carefully and accurately by an experienced examiner as described in detail in the section on the method [1].

Immunologically induced fetal anemia

In a prospective study of 111 pregnancies with Rh incompatibility, the sensitivity with respect to moderate and severe fetal anemia was 100% and the false-positive rate was 12% when the MCA-PSV exceeded the cutoff of 1.5 MoM [39], thus reducing invasive procedures by about 70% (serial amniocentesis) that would otherwise be required. The choice of intervals between measurements depends on the respective measured velocities in the case of blood group incompatibility [39]. After 35<sup>+</sup>0 gestational weeks, the proportion of false-positive results increases. Other studies showed slightly worse results [44, 45].

In case of blood group incompatibility confirmed by detection of maternal alloimmune antibodies (critical titer usually ≥ 1:16, in case of Kell incompatibility ≥ 1:8), paternal zygosity is determined...
first; if necessary, direct genotyping of the fetus from amniotic fluid or from free fetal DNA in maternal blood [46] or from amniotic fluid is subsequently performed. If the fetus has the appropriate antigen, fetal monitoring by serial measurements of MCA-PSV starts at 16+0–20+0 weeks and is then performed every 1–2 weeks, or at shorter intervals in the case of borderline findings and/or other indications [47, 48].

To determine the timing of repeat intrauterine blood transfusion, velocity measurement in the middle cerebral artery is also suitable, but with limitations. Due to the smaller size and viscosity of the previously transfused adult erythrocytes, the MCA-PSV is higher for the same hemoglobin concentration than for fetuses that have not yet been transfused; the optimal borderline after a transfusion was 1.69 MoM, not 1.5 MoM [49]. Because of the mixture of fetal and adult erythrocytes, velocity measurements in the MCA alone are not adequate to fully estimate the extent of anemia, and a daily hemoglobin drop of 0.3 g/dL should be assumed for timing further transfusions [49–51].

The key advantages of measuring PSV are noninvasive serial monitoring and avoidance of amniocentesis or cordocentesis. Furthermore, early detection of fetal anemia with subsequent therapy prevents the occurrence of fetal hydrops. The number of transfusions required cannot be reduced [51, 52].

Non-immunologically-induced fetal anemia

Measurement of MCA-PSV is also applicable in fetal anemias of other causes (Table 3), and in almost all hemolytic and nonhemolytic anemias, such as parvovirus B19, Kasabach-Merritt sequence, fetomaternal transfusion, and TAPS. An exception regarding the strict inverse correlation between MCA-PSV and fetal Hb is fetal α-thalassemia [53, 54]. In this case, the method of choice is the measurement of the cardiothoracic ratio (CTR). Even in the first trimester, a CTR >0.5 is the best diagnostic parameter (sensitivity 97 % with a false-positive rate of 9 %) [54].

In the case of sonographic evidence of non-immunological fetal hydrops, measurement of MCA-PSV allows simple, rapid, and reliable detection or exclusion of anemia, in the presence of which intrauterine blood transfusion can immediately improve the condition of these usually severely anemic fetuses.

Parvovirus B19 infection

The first and early 2nd trimesters appear to be the critical period of fetal parvovirus B19 infection. Placental transmission of the virus to the fetus results in consecutive apoptosis of erythroid progenitor cells and transient aplastic anemia. Placentalitis, hepatitis, and myocarditis may develop with generalized infection [55, 56]. Myocarditis appears to be responsible for more frequent signs of cardiac dysfunction in contrast to fetuses with blood group incompatibility.

Severe anemia is developed almost exclusively in fetuses almost exclusively during the first and second trimesters; this can lead to fetal hydrops in about 5–10 % and death of the fetus in about 3–5 % around 2–6 weeks after infection; severe fetal anemia occurs less frequently in infections after 16+0 weeks of gestation [55, 56].

If maternal PV B19 infection is detected, regular Doppler sonographic measurements of MCA-PSV up to 12 weeks after the time of maternal infection are recommended [56, 57].

Highly-elevated PSV is found in PV B19-induced fetal anemia in the first and early second trimesters (Fig. 8). This may be associated with thickened nuchal translucency, cardiomegaly, skin edema (“space suit” phenomenon), polyhydramnios, placental-megaly, and hyperreflective bowel [43]. Since intrauterine transfusions before 20+0 gestational weeks are difficult and have a higher rate of complications, it seems reasonable to perform intrauterine transfusion not as soon as MCA-PSV is elevated, but only when there are discrete signs of hydrops [42, 43].

Feto-maternal transfusion

The majority of feto-maternal transfusions with subsequent fetal anemia occur in the third trimester, rarely result in hydrops, but may end up as IUFT. Decreased fetal movements and pathologic fetal heart rate tracings on CTG are an indication of this. Causes may include placental disorders, in combination with fetal growth restriction and preeclampsia, as well as maternal abdominal trauma or external turning. Anemia due to feto-maternal transfusion should be rapidly detected or excluded by Doppler ultrasonography of the middle cerebral artery in all of these constellations [58, 59].

Other fetal anemias

Other uncommon fetal anemias also result in increased flow velocities in the middle cerebral artery (Table 3). Extremely rare aplastic and hemolytic anemias requiring transfusion in the fetal

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**Table 3** Causes of fetal anemia.

<table>
<thead>
<tr>
<th>Cause</th>
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<tbody>
<tr>
<td>aplastic anemia</td>
</tr>
<tr>
<td>Blackfan-Diamond, Fanconi</td>
</tr>
<tr>
<td>Parvovirus B19</td>
</tr>
<tr>
<td>transitory myeloproliferative disorder (TMD)</td>
</tr>
<tr>
<td>hemolytic anemia</td>
</tr>
<tr>
<td>Alloimmunization (Rh-, Kell incompatibility, etc.)</td>
</tr>
<tr>
<td>α-Thalassemia</td>
</tr>
<tr>
<td>erythrocyte membrane defects</td>
</tr>
<tr>
<td>G6PD deficiency</td>
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<tr>
<td>Kasabach-Merritt sequence</td>
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<tr>
<td>cytomegaly</td>
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<tr>
<td>syphilis</td>
</tr>
<tr>
<td>bleeding anemia</td>
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<tr>
<td>feto-maternal transfusion (FMT)</td>
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<tr>
<td>hemorrhage (sacrococygeal teratoma, intestinal perforation)</td>
</tr>
<tr>
<td>fetal vessel damage</td>
</tr>
<tr>
<td>hemorrhage after death of monochorionic twin</td>
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<tr>
<td>blood shift as a consequence of TAPS</td>
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period occur in Blackfan-Diamond syndrome, erythrocyte membrane defects, glucose 6-phosphate dehydrogenase deficiency, type VII mucopolysaccharidosis, or congenital dyserythropoietic anemia (CDA I and II). Not quite as rare are anemias in the context of Kasabach-Merritt sequence in chorangiomas, hepatic hemangiomas, or diffuse hemangiomatosis, as well as in coccogyal teratomas. Fetal cytomegaly infection or syphilis are not infrequently associated with anemia and thrombocytopenia of the fetus, although the anemia is rather mild or moderate, even if fetal hydrops is present. Limited to fetuses with trisomy 21, transient myeloproliferation (transient myeloproliferative disorder, TMD) may be responsible for fetal anemia [60]. Finally, in a few cases of in utero anemia requiring transfusion, the cause is not found antenatally or postnatally [58].

For an accurate diagnosis of rare fetal anemia, it is critical that extensive blood tests be performed at the time of the first fetal blood draw to narrow down the cause of the anemia. This is hardly possible in already transfused fetuses; the same applies to the detection of fetal HbF cells in maternal blood in case of feto-maternal transfusion [58].

Doppler sonography for twin pregnancies

Doppler sonography of the fetal vessels is a key monitoring modality in the care of multiple pregnancies. Staging and management are largely determined by Doppler findings. Determination of chorionicity in the first trimester is essential for counseling and differentiated further care [61]. Doppler sonography of the uterine arteries is of secondary importance in multiples [62].

In dichorionic pregnancies, Doppler sonography is essentially used analogously to singleton pregnancies. Here, the more frequent incidence of complications to be monitored, such as FGR and PE, should be noted, also the use in the case of anomalies.

The exceptions arise in complications of monochorionic multiple pregnancies. Classic twin transfusion syndrome (feto-fetal transfusion syndrome, FFTS) is caused by vascular anastomoses between the placental portions of the multiples [63]. Sonographically, the donor twin has oligohydramnios up to a “stuck twin” and a slightly filled to empty bladder, while the recipient twin has polyhydramnios and a full urinary bladder. Further staging includes Doppler criteria that reflect the hemodynamic changes in the fetus [64]. The donor typically shows an increase in pulsatility in the umbilical artery, while the recipient has a cardiac load due to the increase in volume in the form of an increase in pulsatility in the ductus venous up to a negative or reverse a-wave and pulsations in the umbilical vein. The gold standard of treatment for FFTS is laser coagulation of the anastomoses at the placental portions in the umbilical vein. Here, the flow rate in the umbilical Doppler sonogram must be set to slow to record sufficient cycles to monitor the flow pattern over an extended period of time. Type 3 is at high risk of sudden IUFT or neurological damage.

Anemia of the surviving twin

After death of a monochorionic twin, spontaneously or after selective fetocide, related anemia of the surviving fetus not infrequently occurs in a few hours to days. This effect is caused by a more or less rapid and large blood loss into the placental vascular bed, rarely also into the other twin [75]. This complication can be detected by close-meshed assessment of MCA-PSV and cardiac contractility and filling, initially every hour immediately after the
Summary

Doppler sonography of the maternal, uterine and fetal vessels is a method that is now widely used in prenatal and obstetric medicine. The great advantage lies in the fact that the method is non-invasive and can be used repeatedly if employed correctly.

Doppler sonography of the uterine and fetal vessels alone is not a suitable single procedure for screening. However, incorporated into multimodal algorithms, Doppler findings of uterine arteries are an important pillar of prediction accuracy. In particular, screening for early PE in the first trimester has high accuracy. This is all the more significant because ASA provides an effective prophylaxis for PE.

The Doppler findings of the fetal arteries play a major role in the diagnosis of the condition and monitoring of high-risk pregnancies. The changes in the Doppler findings in the UA, MCA and venous duct in their chronological order are important for monitoring the early FGR. The venous duct can be used to optimize the birth management of the fetuses with FGR. The cerebro-placental ratio (CPR) appears to be the most suitable for predicting and monitoring late FGR; a final assessment is still pending, however.

The measurement of the maximum systolic velocity in the middle cerebral artery is of significant importance for the diagnosis of fetal anemia. Thus, former invasive diagnostics with a high complication rate could be replaced by a non-invasive method. Both the indication and the monitoring of fetuses requiring transfusions are very successful.

The advantages of using Doppler sonography seem obvious. This has been proven in a large number of studies for various aspects of the child's outcome after a complicated progression. Ultimately, however, the critical question remains whether perinatal, neonatal, or early childhood morbidity or mortality can be improved. So far, this has only been confirmed for the use of the umbilical artery in high-risk pregnancies.

The clinical utility of Doppler sonographic findings depends crucially on the quality of the examination. It is therefore essential that the specified quality criteria are applied in every examination. Accurate methodological application coupled with correct physiological or pathophysiologica interpretation ensure the success of the use of Doppler sonography. Parts 1 and 2 of this "Best Practice Guideline for Doppler Sonography" are intended to provide the foundation for this.

Conflict of Interest

The authors declare that they have no conflict of interest.

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