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A Uniform Management Approach to Optimize Outcome in Fetal Growth Restriction

Viola Seravalli, мд, Ahmet A. Baschat, мд*

KEYWORDS

- Fetal growth restriction Fetal acidemia Fetal Doppler Umbilical artery
- Middle cerebral artery Biophysical profile score Neonatal outcome Fetal testing

KEY POINTS

- A uniform approach to diagnosis and management of fetal growth restriction (FGR) produces better outcomes, prevents unanticipated stillbirth, and allows appropriate timing of delivery.
- An estimated fetal weight less than the tappercentile in association with either an elevated umbilical artery Doppler index, a decreased middle cerebral artery Doppler index, or a decreased cerebroplacental ratio should be considered evidence of FGR. Early-onset and late-onset FGR represent 2 distinct clinical phenotypes of placental dysfunction.
- Integration of different testing modalities allows adjustment of monitoring intervals based on Doppler parameters and a more precise prediction of acid-base status based on biophysical variables.
- Antenatal surveillance of the growth-restricted fetus requires adjustment of monitoring intervals based on signs of disease acceleration, when delivery is not yet indicated.
- Thresholds for interventions are defined by the balance of fetal risks of continuation of pregnancy versus the neonatal risks that follow delivery and depend on gestational age.

INTRODUCTION

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The main challenges in the management of pregnancies complicated by fetal growth restriction (FGR) are accurate identification of the small fetus at risk for adverse outcome, prevention of unanticipated stillbirth, and appropriate timing of delivery. A

- Authors declare no relationship with a commercial company that has a direct financial interest in the subject matter or materials discussed in the article or with a company making a competing product.
- Department of Gynecology and Obstetrics, The Johns Hopkins Center for Fetal Therapy, The
 - Johns Hopkins Hospital, 600 North Wolfe Street, Nelson 228, Baltimore, MD 21287, USA * Corresponding author.
- 48 E-mail addresses: aabaschat@hotmail.com; abascha1@jhmi.edu

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uniform management approach to diagnosis and management of FGR consistently
 produces better outcome than is reported in observational studies that rely on a range
 of diagnostic, surveillance, and delivery criteria.^{1–5} Once the diagnosis of FGR has
 been made, surveillance tests need to be applied at appropriate intervals until the rela tive risks of delivery outweigh the benefits of ongoing monitoring. These factors are
 determined by the clinical phenotype of FGR across gestational ages.

CLINICAL PHENOTYPE OF FETAL GROWTH RESTRICTION IN RELATION TO GESTATIONAL AGE

FGR evolves from a preclinical phase to clinically apparent growth delay and may eventually lead to fetal deterioration before the spontaneous onset of labor. Growth delay due to decreased nutrient delivery affects liver size and therefore the abdominal circumference (AC) first, and then growth of the head and entire body.⁶ Abnormal placental perfusion in the maternal compartment results in increased blood flow resis-tance in the uterine artery flow-velocity waveform.⁷ Abnormal perfusion of the fetal villous vascular tree is associated with decreased umbilical artery (UA) end-diastolic velocity proportional to the degree of flow impairment.⁸ Abnormal oxygen diffusion across the villous membrane leading to lower fetal arterial Pagars sociated with a decrease in middle cerebral artery (MCA) blood flow resistance, Hereas decreased CO₂ clearance additionally increases the MCA peak systolic velocity (Fig. 1).¹⁰ The relative predominance of these mechanisms determines the clinical picture of FGR.11-16

FGR that is established by the second trimester is associated with a greater degree of vascular abnormality in the maternal and fetal compartments of the placenta. In the mother, high-resistance uterine artery flow velocity waveforms and a 40% to 70% rate of associated pre-eclampsia are characteristic. In the fetal compartment, an elevation

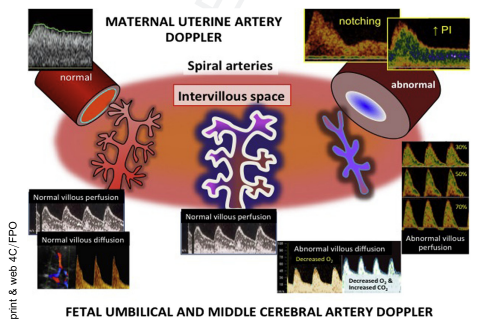


Fig. 1. Clinical correlates of maternal and fetal aspects of placental function.

of the UA pulsatility index (PI) is typical.^{11,12} In FGR that is not established until 31 to 100 34 weeks (late-onset FGR), villous diffusion and perfusion defects coexist in various 101 proportions,¹⁷⁻²¹ leading to cerebral or UA Doppler abnormalities that may be present 102 103 independent of each other (Fig. 2).²²⁻²⁴ Because of this variable association between 104 small fetal size and abnormal Doppler velocimetry, distinction between growth restric-105 tion and constitutional smallness can be challenging. Accordingly, management chal-106 lenges in early-onset FGR revolve around prematurity and coexisting maternal 107 hypertensive disease, whereas in late-onset disease, failure of diagnosis or surveil-108 lance leading to unanticipated stillbirth is the primary issue.^{25,26}

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110 DIAGNOSIS OF FETAL GROWTH RESTRICTION

The diagnosis of fetal growth delay can be based on fetal biometry alone or by also 112 taking umbilical or cerebral artery Doppler indices into consideration. An AC less 113 than the tenth percentile has the highest sensitivity for the diagnosis of FGR, whereas 114 a sonographically estimated fetal weight (SEFW) less than the tenth percentile has 115 greater specificity.¹¹ Most national societies agree on the tenth percentile for the 116 SEFW as a diagnostic cutoff for small for gestational age (SGA). The disadvantage of 117 this cutoff is the inclusion of a variable number of normal constitutionally small fetuses 118 that do not require surveillance. Using an SEFW less than the third percentile or a 119 decreased AC growth rate is more likely to identify "true FGR,"27 but has the disadvan-120 tage that less severe forms of FGR at risk for deterioration are missed and therefore their 121 risk for stillbirth remains. Combining an SEFW less than the tenth percentile with either 122 an abnormal UA, MCA, or cerebroplacental ratio (CPR, defined as UA/MCA index), in-123 creases the identification of the small fetus at risk for adverse outcome. Although UA 124 Doppler velocimetry is sufficient for the diagnosis of FGR before 32 weeks gestation, 125 thereafter MCA Doppler is also required to represent the whole clinical spectrum found 126 in early-onset and late-onset placental disease. 12,14,16,24 Because the CPR mathemat-127 ically amplifies mild abnormalities in the umbilical and middle cerebral arteries, it is 128



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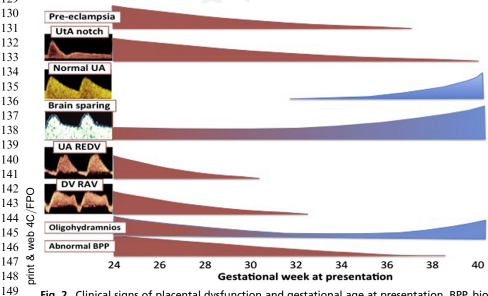


Fig. 2. Clinical signs of placental dysfunction and gestational age at presentation. BPP, biophysical profile; RAV, reversed a-wave velocity; UtA, uterine artery.

151 the most sensitive Doppler parameter, especially after 28 weeks of gestation, and its 152 decrease should alert the clinician to the possibility of evolving brain sparing. Here, 153 an SEFW less than the tenth percentile in association with either an elevated UA Doppler 154 index, a decreased MCA Doppler index, or a decreased CPR should be considered evidence of FGR (Table 1).^{11,12,14,16,24} The proportion of growth-restricted fetuses 155 156 with normal UA blood-flow resistance but isolated MCA brain sparing is higher toward 157 the late third trimester. Accordingly, MCA Doppler better identifies FGR after 34 weeks of gestation, when the predictive accuracy of CPR decreases.¹² 158

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160 ASSESSMENT OF THE DEGREE OF FETAL DETERIORATION 161

Fetal surveillance tests are applied to pregnancies with suspected FGR to estimate the 162 163 risk for hypoxemia, prelabor acidemia or stillbirth, as well as the rate of clinical deterioration. The required accuracy of this assessment is highest at early gestational ages 164 165 wherein prematurity-related risks are high and each additional day gained in utero can 166 significantly increase chance of neonatal survival. An accurate estimation of pH is 167 important to predict fetal compromise that precedes stillbirth and therefore critical 168 to time delivery.

169 The association between the abnormalities in Doppler parameters and the deterioration of fetal acid-base status has been demonstrated in several studies,²⁸⁻³¹ pre-170 dominantly in the preterm fetus. Abnormal umbilical flow patterns indicate an 171 172 increased risk of hypoxemia and acidemia proportional to the severity of Doppler abnormality. Although Doppler findings in each of the examined vascular beds correlate 173 with fetal acid-base status, there is a wide variation in fetal pH with abnormal results. 174 175 Among Doppler parameters, the elevation of the precordial venous Doppler indices provides the best prediction of acidemia in fetuses with FGR.^{31,32} Therefore, fetal 176 177 Doppler assessment that is based on the UA indices alone is no longer appropriate 178 in early-onset FGR, and the incorporation of venous Doppler is necessary to assess 179 the rate and degree of fetal compromise. In preterm growth-restricted fetuses, MCA 180 Doppler study has limited accuracy to predict acidemia and adverse outcome and 181 should not be used to time delivery. Beyond 34 weeks, the UA waveform may be 182 normal, and therefore, the best predictor of fetal adaptation to hypoxemia is considered the MCA PI. However, studies on fetal brain circulation in late-onset FGR^{33,34} pri-183 184 marily evaluated the relationship of MCA Doppler with intrapartum fetal distress or neonatal acidosis rather than prelabor acid-base status. Accordingly, conclusions 185 relating MCA Doppler to fetal pH are generally extrapolated. 186

188 189 Table 1 190 Implications of diagnostic cutoffs for management of fetal growth restriction 191 **Diagnostic Cutoff** Advantage Disadvantage 192 193 AC <10th percentile Highest sensitivity for FGR Lowest specificity for FGR 194 SEFW <10th percentile Acceptable sensitivity for FGR Unnecessary monitoring of normal fetuses 195 SEFW <3rd percentile Less severe FGR is missed 196 Greater specificity for FGR 197 SEFW <10th percentile Greatest specificity for FGR at Misses term FGR with normal & abnormal UA Doppler risk for adverse outcome **UA** Doppler 198 199 SEFW <10th percentile with Greatest specificity for FGR at Requires interpretation of abnormal UA or MCA umbilical and cerebral risk for adverse outcome 200 across all gestational ages Doppler studies

202 The 5-component biophysical profile scoring (BPS) shows a reliable and 203 reproducible relationship with the fetal pH, irrespective of gestational age.^{35,36} An 204 abnormal BPS of 4 or less is associated with a mean pH of less than 7.20 and a score 205 of less than 2 has a sensitivity of 100% for acidemia.³⁶ When the relationship between 206 the various testing modalities and fetal acid-base status is compared, biophysical 207 parameters show a closer relationship with the pH, whereas there is a wide variation 208 in fetal pH with abnormal Doppler results. On the other hand, the BPS alone has limited utility in the prediction of longitudinal deterioration,^{37,38} which is better assessed with 209 210 multi-vessel Doppler studies.

211 Fetal heart rate is one of the 5 components of the BPS. A nonreactive cardiotoco-212 gram (CTG) has been correlated with fetal hypoxemia and acidemia,^{39,40} but it is associated with a wide range of pH values,³⁹ and as for the other components of the BPS, it 213 214 does not anticipate the rate of deterioration. Computerized heart rate monitoring 215 (cCTG) has been introduced to improve the interpretation of fetal heart rate traces, 216 by determining quantitative parameters, such as the short-term variation, that cannot 217 be visually assessed. In fetuses with intrauterine growth restriction, a short-term vari-218 ation less than 3.5 ms appears the best predictor of an UA pH of less than 7.20.41 219 However, cCTG as a stand-alone test in FGR offers limited accuracy, and it performs 220 best when combined with venous Doppler or as a substitute for the traditional NST in 221 the BPS.42 222

223 224 SELECTION OF MONITORING INTERVALS

The goal of fetal surveillance is to prevent stillbirth and irreversible fetal deterioration;
 this requires adjustment of monitoring intervals based on signs of disease accelera tion, when delivery is not yet indicated.

With standardization of antenatal surveillance, a reduction in antenatal mortality might be achieved without worsening neonatal outcome.³ The optimal surveillance pattern and timing of delivery remain the objects of much debate and research. There is no general consensus between national guidelines on the appropriate frequency of testing, and they are based on expert opinion of key authors because there is no highquality evidence to guide practice.

In the authors' opinion, the best approach consists of a longitudinal surveillance
starting at 24 to 26 weeks with integrated fetal testing, including multivessel Doppler
examination, fetal heart rate analysis, and assessment of fetal activity through BPS,
because the combination of tests improves the prediction of acidemia and stillbirth
compared with single tests.^{37,42-44}

239 Monitoring interval choice depends on gestational age at onset and signs of dete-240 rioration at Doppler study. When new features indicating disease acceleration or fetal 241 deterioration develop, monitoring frequency needs to be increased until the delivery 242 threshold is reached. Because early-onset and late-onset FGR represent 2 distinct 243 clinical phenotypes of placental dysfunction, they show different signs of disease pro-244 gression. In early-onset FGR, fetal deterioration typically evolves from abnormal UA 245 Doppler studies, to brain-sparing, abnormal venous Doppler parameters, abnormal 246 computerized CTG, and finally, an abnormal 5-component BPS.^{38,45-52} The rate of 247 progression is determined by the interval between diagnosis to loss of UA end-248 diastolic velocity^{49-51,53} and typically takes 4 to 6 weeks.⁵¹ Once forward velocities 249 in the ductus venosus (DV) become absent or reversed, fetal survival of longer than 250 1 week is unlikely.⁵⁴ Late-onset FGRs are characterized by a slower progression 251 (up to 9 weeks), with predominant cerebral or UA Doppler abnormalities. There are 252 no evident Doppler changes in the precordial veins and brain sparing may be the 6

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only observed Doppler sign of hypoxemia (see Fig. 2).^{16,55} Importantly, however, ter-253 minal deterioration resulting in stillbirth occurs more rapidly and unanticipated in term 254 255 FGR.⁵⁶ Therefore, a closer surveillance is required after 34 weeks, and new onset of 256 Doppler abnormalities at this age should raise consideration for delivery. 257

The observed progression of Doppler abnormalities should determine the interval of monitoring as follows, until the threshold for delivery is reached. 259

Early-onset fetal growth restriction

- Elevated UA Doppler flew PI (>2 SDs greater than the mean for gestational age), no other abnormality: every 2 weeks Doppler, weekly BPS
- Low MCA or CPR: weekly Doppler + BPS
- UA absent end-diastolic velocity (AEDV): consider admission, 2 times per week Doppler + BPS
 - UA reversed end-diastolic velocity (REDV), increased DV Doppler indices, and/or oligohydramnios (maximum vertical pocket of fluid <2 cm): admission, 3 times per week Doppler + BPS, daily CTG
- Absent/reversed DV a-wave: admission, daily Doppler + BPS, prepare for delivery

Late-onset fetal growth restriction (>34 weeks)

- Elevated UA Doppler flow PI (>2 SDs greater than mean for gestational age), no other abnormality: weekly Doppler + BPP
- Low MCA or abnormal CPR: 2 to 3 times per week Doppler + BPS

PLANNING DELIVERY: GESTATIONAL AGE AS A DETERMINANT OF INTERVENTION THRESHOLDS

278 In pregnancies complicated by FGR, the thresholds for interventions are defined by the balance of fetal risks of continuation of pregnancy versus the neonatal risks that 280 follow delivery. The principle neonatal risks are neonatal mortality, major neonatal morbidity, which is associated with long-term impacts on health, and adverse 282 neonatal development. These risks change in specific gestation age epoch (Fig. 3, 283 Table 2), and the outcome is comparable to that of appropriate for gestational age in-284 fants born at a 2-week shorter gestational age.⁵⁷ Accordingly, the threshold for deliv-285 ery needs to be higher at earlier gestational age. 286

The neurodevelopmental outcome of growth-restricted babies has received growing attention in recent years, given the impact on quality of life.^{4,58,59} In earlyonset FGR, gestational age has been found to be one of the major determinants of neurodevelopment. However, it remains to be determined if interventions other than modulating disease course might improve neurodevelopment.

291 Taking in account the data on neonatal survival derived from 2 large observational 292 studies (see Fig. 3),^{3,5} the following delivery indications per gestational epoch are 293 suggested. 294

24 to 26 Weeks Gestation

296 The survival rate of FGR neonates averages less than 50%.⁵ In surviving babies, the 297 risks for major neonatal complications are as high as 80%. With these neonatal mor-298 bidities, especially higher grades of intraventricular hemorrhage, the motor neurode-299 velopmental adverse outcomes are equally high. These risks gradually decrease 300 and there is an improvement in survival by an average of 2% per gestational day 301 that is gained in utero. The survival rates exceed 50% once the estimate of fetal weight 302 exceeds 500 g or 26 weeks are reached. Because of these significant neonatal mor-303 bidities, delivery for fetal deterioration may not be considered in certain health care

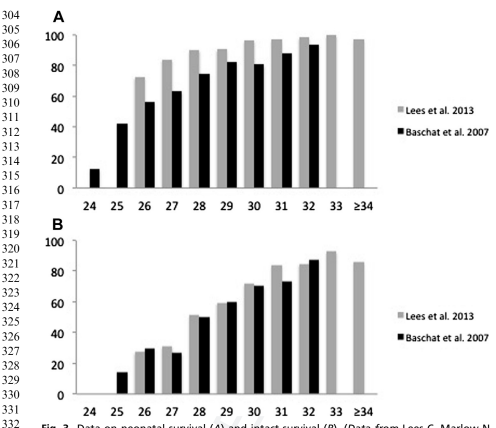


Fig. 3. Data on neonatal survival (A) and intact survival (B). (Data from Lees C, Marlow N,
Arabin B, et al. Perinatal morbidity and mortality in early-onset fetal growth restriction:
cohort outcomes of the trial of randomized umbilical and fetal flow in Europe (TRUFFLE).
Ultrasound Obstet Gynecol 2013;42(4):400–8; and Baschat AA, Cosmi E, Bilardo CM, et al.
Predictors of neonatal outcome in early-onset placental dysfunction. Obstet Gynecol 2007;109(2 Pt 1):253–61.)

- settings. Maternal indications such as severe pre-eclampsia are the primary indica-tions for delivery.
- 341 342 **26 to 28 Weeks Gestation**

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343 Neonatal survival exceeds 50%. However, intact survival at 26 to 27 weeks remains 344 around 30% (see Fig. 3).^{3,5} Because neonatal morbidity rates are high, additional fetal 345 deterioration before delivery does not appear to produce a statistical impact on survival. 346 Although maternal disease remains an absolute delivery indication, fetal status may not 347 qualify until acidemia is certain. Although an abnormal 5-component BPS (<6/10) is an 348 indication to delivery from 26 weeks of gestation, because of its strong association with 349 fetal acidemia, the evidence of venous Doppler abnormalities is not considered an indi-350 cation to intervention until 28 weeks. The observed median time interval between the 351 detection of abnormal venous Doppler indices and the deterioration of the BPS is 352 1 week,⁵² which could potentially increase neonatal survival by 14% (see Table 2). Indi-353 vidualization of care in these pregnancies needs to be discussed with the patient, 354 including the option of nonintervention.

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Each day in utero

of 1%

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Reversed DV a-wave

associated with

lower neonatal survival

Reversed DV a-wave

before delivery is

SGA fetuses

receiving prenatal

lower rate of RDS,

steroids have

BPD, IVH, and

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UA REDV

SGA neonates

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38 wk have a

higher rate of

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Evidence

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Delay not justified	
Risks of surveillance	Ŀ.
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decline in growth,	
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delivery at 38 wk	U

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Delivery threshold Maternal conditions Abr Abbreviation: NICU, neonatal intensive care unit.

Birth-weight <500 g

associated with

>50% mortality

& gestational age

<26 wk at delivery



Each day in utero

Fetal deterioration

Abnormal BPS (<6)

impact on neonatal outcome

has no statistical

of 2%

increases neonatal

survival by median

406 28 to 32 Weeks Gestation

407 Neonatal survival exceeds 70% at 28 weeks and increases to more than 90% at 408 32 weeks (see Fig. 3). Survival gain per day in utero now averages 1% and neonatal 409 mortality and morbidity progressively decrease. Fetal deterioration of venous Doppler 410 parameters may be tolerated as long as DV a-wave velocities are antegrade. Reversal of 411 the DV a-wave before delivery has an independent additional impact on neonatal mor-412 bidities, and persistence of this abnormality beyond 1 week carries significant risk for 413 stillbirth. For this reason, the presence of a DV reversed a-wave is generally considered 414 an indication to intervention from 28 weeks. However, delivery before 30 weeks gesta-415 tion still carries a significantly higher risk for adverse neurodevelopment at age 2 416 because of neonatal complications and their impact on motor development.⁴

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419 <u>32 to 34 Weeks Gestation</u>

420 Thirty-two to 34 weeks gestation is a time in fetal development whereby the cerebral circulation gains an additional structural layer, and, accordingly, there is a significant 421 422 reduction in the rates of intraventricular hemorrhage. This reduction has measurable 423 impact on motor development at age 3. Now, up until 34 weeks gestational age espe-424 cially, the administration of antenatal steroids has an added benefit in reducing respi-425 ratory neonatal morbidity as well as intraventricular hemorrhage rates, and babies who 426 have received steroids have improved survival. Moreover, recent evidence suggests that neurodevelopment is also improved by the administration of steroids⁶⁰; this is 427 428 most likely due to the beneficial impact on the respiratory performance and the 429 decrease of ventilation related intraventricular bleeding.

430 Evidence of reversed UA end-diastolic velocity is generally considered a delivery indi-431 cation from 32 weeks onward, whereas an AEDV is an indication from 34 weeks onward.

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433 **34 to 38 Weeks Gestation**

434 At this gestational age, the gain in survival as well as neonatal morbidity is minimal; 435 however, up to 38 weeks gestation, the rate of neonatal admissions to the intensive 436 care nursery is still significantly greater for FGR infants, and the overall neonatal 437 adverse outcome scores are higher. Accordingly, delivery thresholds should be based 438 on clear maternal or fetal indications. The absence of UA end-diastolic velocity at 439 Doppler study is considered an indication to delivery from 34 weeks onward. In 440 late-onset FGR, the MCA Doppler is considered the best predictor of fetal adaptation 441 to hypoxemia, and some national guidelines recommend the use of this parameter to 442 time delivery in fetuses with normal UA Doppler.^{61,62} 443

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445 After 38 Weeks Gestation

446 Neonatal adverse events in SGA infants are negligible and, accordingly, ongoing preg-447 nancy must be weighed carefully against the risks of unanticipated stillbirth if the pa-448 tient remains undelivered. Risks of surveillance failure, risks for progressive decline in 449 head growth, and low neonatal risks favor delivery. The Disproportionate Intrauterine 450 Growth Intervention Study at Term (DIGITAT)² showed that among women with sus-451 pected intrauterine growth restriction at 36 to 41 weeks, a policy of labor induction 452 affects neither the rate of adverse neonatal outcomes nor the rates of instrumental 453 vaginal delivery or caesarean section, indicating that both approaches are acceptable. 454 The consensus view from the DIGITAT is that the optimum time for induction in SGA 455 with normal Doppler study is at around 38 weeks, because it is associated with the 456 lowest neonatal morbidity⁶³ and seems to minimize the risk of stillbirth.⁶⁴

457 Between 24 and 34 weeks, a single course of should be administered over a period or 458 of 48 hours for fetal lung maturity if delivery is being considered. At this age, delivery 459 should be planned at a center with a neonatal intensive care unit. The route of delivery 460 depends on the severity of fetal compromise, along with maternal condition and other 461 obstetric factors. If prelabor acidemia is suspected, cesarean section is recommen-462 ded. In FGR cases with abnormal UA Doppler, induction of labor can be offered, 463 but rates of emergency caesarean section are increased. The use of prostaglandin 464 for cervical preparation is usually discouraged. Because of the increased risk of intra-465 partum asphyxia in growth-restricted fetuses, continuous fetal heart rate monitoring is 466 recommended from the onset of uterine contractions.

469 SUMMARY

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470 Detection of FGR must be accompanied by uniform approaches to management to 471 improve perinatal outcomes. The understanding of the clinical phenotype of early-472 onset and late-onset FGR is actively evolving. A decreased estimated fetal weight 473 coupled with abnormal umbilical, MCA, or CPR studies provides the best identification 474 of fetuses requiring surveillance. Doppler abnormalities precede biophysical deterio-475 ration and therefore allow adjustment of monitoring frequency. Concurrent deteriora-476 tion of Doppler and biophysical variables best predict prelabor acidemia and therefore 477 allow timing of delivery. The threshold for delivery is determined by the neonatal risks 478 at each gestational epoch and decreases with advancing gestational age. 479

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