How good are we at first-trimester prevention and prediction of early fetal growth restriction?

A series of recent publications have shown that early preeclampsia can be successfully predicted in the first trimester using a combination of maternal, biophysical and biochemical factors [1–4], and administration of low-dose aspirin to women identified as high-risk can reduce the development of preterm preeclampsia by about 60% [5]. Given that abnormal placentation and placental insufficiency are common pathways to many cases of preeclampsia and fetal growth restriction (FGR), we intuitively use the same strategies to predict and prevent FGR. However, are we as successful in FGR as we are in preeclampsia?

A variety of definitions

In contrast to preeclampsia, which is a clearly defined condition, multiple terms describe suboptimal fetal growth, and two of these, i.e. small for gestational age (SGA) and FGR, are often used interchangeably. This happens despite the two being two different conditions. SGA merely signifies a fetus (or a neonate) that is smaller than a given centile (usually the 10th), whereas FGR involves the failure of the fetus to reach its developmental potential, and it normally refers to a small fetus with some evidence of hypoxia. Therefore, one might say that FGR is a subset of SGA, and this is the case most of the time, but not always. A growth-restricted fetus can be non-SGA (i.e. > 10th centile for estimated weight or abdominal circumference) and still be FGR, if its growth potential was meant to be higher. This reality has been acknowledged by the recent consensus definition for FGR, which introduces the option of a fetus crossing centiles, even if its eventual centile is > 10th [6]. Despite this development, different definitions have traditionally been used in the existing studies, even within the terms SGA and FGR (different cutoffs, Doppler parameters, pre- or postnatal weight etc), adding to the heterogeneity of the literature.

Targeting small fetuses

Prediction of FGR (or SGA) has been pursued along that of preeclampsia, and it was early acknowledged that the yield of screening models is higher when FGR coexists with preeclampsia rather than when it is an isolated condition. Four years ago, the Fetal Medicine Foundation (FMF) developed a dedicated screening algorithm targeting SGA, and they reported that, by combining maternal, biochemical and biophysical factors, about 50% of SGA can be predicted for a 10% false positive rate [7]. However, is this the case?

The yield of screening models is significantly higher when FGR coexists with preeclampsia, in the absence of preeclampsia, this screening strategy can predict about 40% of those requiring delivery before 37 or 32 weeks [1]. So, in practice, screening for preeclampsia with a 10% screen-positive rate (which is where 1:100 corresponds to) can predict about 35–40% of the small and very small fetuses requiring early delivery.

Prevention of small fetuses through preeclampsia-oriented screening

Administration of prophylactic aspirin to these screen-positive (for preeclampsia) cases, will significantly reduce the risk for small neonates with preeclampsia, but will only reduce the risk for the subgroup of such neonates without preeclampsia that require delivery before 32 weeks, by about 60%. Of course this is quite an important group in terms of morbidity, but it only comprises about 0.25% of the population [1].

A synopsis...

It appears thus that combined first-trimester screening for preeclampsia can predict about 40% of the subgroup of small fetuses who will need a preterm delivery. Prophylactic aspirin in screen-positive cases will reduce the risk for the small but clinically important group of very preterm (<32 weeks) small fetuses by about 60%.
...and some more unanswered questions

There are yet more unanswered questions, including the possible role of maternal hemodynamics as part of a combined screening, or the ability of the fetal fraction of cell-free DNA in the maternal blood to act as an independent predictor. And, of course, there is the issue with twins. There is an ongoing debate whether twin-specific growth charts should be used, or charts of singleton are applicable as well, so problems like this need to be solved before aiming at screening.

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References


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