The term placental insufficiency has been increasingly used in recent years in Britain to describe a condition where the fetus is particularly at risk, or dies, antepartum, intrapartum or sometimes postpartum, without trauma. Postmortem examination of the child shows that it is entirely healthy and the only changes demonstrable are those of anoxia, namely multiple petechial hemorrhages, and the presence of meconium and liquor in the lungs. The placenta may show disturbance, for example infarction or abruptio, or it may be unduly small. On the other hand it may seem perfectly normal. Such silent intra-uterine death is common in pre-eclamptic toxemia or in hypertension, and it is occasionally seen when the only abnormality is that the pregnancy is prolonged beyond the normal span.

**Placental Ischaemia**

On the basis of animal experiments Page (1939), Bastiaanse and Mastboom (1949) and Beker (1949) all suggested that placental ischaemia was the causative factor. Evidence of the occurrence of placental ischaemia in the human has however only comparatively recently been presented. Greenfield, Shephard and Whelan (1951) calculated that to supply the oxygen requirements of a fetus weighing 3,000 g., 350 ml. of maternal blood must flow through the placental bed each minute. Villee (1954) reached almost the same conclusion. Assali, Douglas, Baird, Nicholson and Suyemoto (1953) inserted a catheter in a uterine vein in the human and found, using nitrous oxide and the Fick principle, that at 38 weeks the amount of maternal blood flowing through the uterus was 750 ml. per minute. Metcalfe, Romney, Ramsey and Reid (1953), by a similar method found a slightly lower figure. Our own work (Browne and Veall, 1953) showed that under normal conditions at 38 weeks 600 ml. of maternal blood were supplied to the placenta each minute, so that it seemed that the major portion of blood entering the uterus went to the placenta. Details of our technique have been published elsewhere, but it may be of interest to recall it in general terms.

The situation of the placenta within the uterus is first determined by outlining the area of the uterus which shows greatest radio-activity after an intravenous injection of 10 microcuries of radio-sodium $^{24}$Na. If the placenta or part of it is found to be situated on the anterior wall of the uterus, it is then possible to investigate further. After the lapse of a few days to allow elimination of all radio-activity due to the placental localization, 5 microcuries of $^{24}$Na are injected into the intervillous space, and the time taken for the injected sodium to be carried away by the circulation is recorded. In a normal healthy woman at 38 weeks half of the injected sodium is removed in about 20 seconds. If hypertension or toxemia is present the rate of clearance is substantially reduced, so that half of the injected dose may require as much as 60 seconds for its removal. This shows very clearly that in the presence of hypertension or of pre-eclamptic toxemia there is a measurable degree of placental ischaemia. Morris, Osborn and Wright (1955), using a modification of our technique, showed that a similar reduction in blood flow occurred in the uterine muscle in toxemia, and we ourselves showed that the same occurred in the presence of hypertension without toxemia. Cox and Chalmers (1953) and Johnson and Clayton (1955) have also shown that in these conditions transfer of sodium across the placenta to the fetus is considerably slowed, and Clemetson and Churchman (1954) found that the transfer of amino-acids was similarly impaired. On the other side of the placenta Walker and Turnbull (1953) showed that in these conditions the fetus is short of oxygen. It is clear therefore that in toxemia and hypertension there is placental impairment which may amount to insufficiency.

In our own series, although the amount of maternal blood (normally 600 ml. per minute), was reduced in cases of toxemia or hypertension to as little as 200 ml. per minute, no babies were lost. It is apparent therefore that under normal conditions the placenta has a very considerable functional reserve, and moreover that the fetus has various means of counteracting the adverse effects of placental insufficiency. Only when the
placental function is very seriously impaired, and the fœtus has used to the full all its powers of compensation, will fetal death occur.

**Diminished Placental Permeability**

Impairment of the maternal circulation to the placenta is one cause of placental insufficiency. Another cause is diminished placental permeability. Flexner, Cowie, Hellman, Wilde and Vosburgh (1948) showed that the rate of transfer of sodium from mother to fœtus is impaired in late pregnancy, and the nearer to term the greater the degree of impairment. This seems to be a normal ageing process in every placenta, and is aggravated if the pregnancy goes beyond term.

**Liquor Volume**

Recent work by Elliott and Inman (1961) in my own Department has shown that the volume of the liquor amnii reflects the functional capacity of the placenta. Thus they found that in a healthy woman the volume of the liquor reached a peak (1,100 ml.) at 37 weeks, diminishing steadily thereafter until at 42 weeks there was only about 300 ml. altogether. Every clinician knows from his own experience that in some cases of prolonged pregnancy when the membranes rupture there is virtually no liquor, and what little there is is thick and stained with meconium. In pre-eclamptic toxæmia and in hypertension Elliott and Inman found that the volume of liquor at 37 weeks was less than in the normal woman (approximately 800 ml.) and the reduction in volume proceeded more rapidly than normal, so that volumes of 100 to 200 ml. were found between 40 and 41 weeks. Elliott’s findings suggest that the volume of the liquor amnii reflects accurately the functional state of the placenta, and volumes below 300 ml. suggest that the fœtus is in grave danger from placental insufficiency.

**Twin Pregnancy**

Though I have not yet been able to prove it I strongly suspect that placental insufficiency occurs particularly in twin pregnancy, and that in this case it is of dual origin, for Morris, Osborn, Wright and Hart (1956) found that uterine blood flow was somewhat slower in twin pregnancy, and undoubtedly they combined weights of the fœtuses being greater than that of a single fœtus there is a greater demand for oxygen to cross the placenta. I consider therefore that in twin pregnancy the optimum duration is 38 weeks and not 40 weeks. In other words that if "term" for a single pregnancy is 40 weeks it is 38 weeks for a twin pregnancy.

Because both hypertension and postmaturity impair the efficiency of the placenta it will be evident why these two conditions when combined carry a considerable risk for the fœtus.

**Other Factors**

Several other factors impair the efficiency of the placenta:

1. **Age.** There is good evidence that the blood pressure rises steadily with age. Thus in a survey of the blood pressure of 7,000 primigravidae (Browne 1961) the mean blood pressure at the age of 20 was 124/73, rising steadily to 130/76 at the age of 40. It is also well known clinically that the older woman runs a greater risk of losing her baby from no obvious cause. It seems reasonable, when we remember the effect of hypertension on the maternal placental circulation, to suppose that the less resilient circulation of the older woman should give rise to placental insufficiency.

2. **Gravidity.** There seems to be no doubt that the primigravida and grand multipara are more likely to have a silent intrauterine fœtal death than women having their second or third children. It may be that in the primigravida the circulatory responses to the demands of pregnancy are not fully met as pregnancy is a new experience for her. The grand multipara on the other hand, though knowing what is demanded of her, is unable because of a degenerating vascular system to respond fully to these demands.

3. **Exercise.** The beneficial effects of rest in the management of toxæmia and of hypertension in pregnancy have been known for generations. Morris and others (1956) showed that exercise resulted in considerable slowing of the uterine circulation. He first determined the rate of clearance of $^{42}$K from the uterine muscle of women at rest, then during exercise, and finally at rest again. In every case the slowing of uterine circulation produced by exercise was very marked, and moreover the return to normal on cessation of exercise was slow. It will readily be seen that the effect of physical activity on a uterine and placental circulation already impaired by hypertension or prolonged pregnancy may be lethal for the fœtus.

4. **Labour.** Wright, Morris, Osborn and Hart (1958) also showed very clearly that during labour the uterine circulation was impaired and that the degree of impairment increased the longer the labour went on. Caldeyro Barcia (1959), using our own method, showed that during a uterine contraction the maternal placental blood flow was diminished. This may well account for many of the cases of fœtal distress and even fœtal death during labour in women suffering from pre-eclampsia and hypertension.

Thus we can say that placental insufficiency may occur in the following conditions: age over 30, primigravida, toxæmia, hypertension, chronic
nephritis, diabetes, prolonged pregnancy, physical exercise, labour, and in twin pregnancy after 38 weeks.

Diagnosis

How can placental insufficiency be detected? Scott Russell, Payne and Coyle (1957) have shown that when serial determinations of pregnanediol excretion are made during pregnancy there is a steady rise as the months go by until the last two weeks or so before delivery, when it falls. In cases of placental insufficiency, however, pregnanediol excretion, which may initially rise quite normally, never reaches the normal peak, and some two weeks or so before the child dies in utero pregnanediol excretion falls. Zondek and Goldberg (1957) have shown that urinary oestriol elimination can be used in late pregnancy as a qualitative test of placental function. Elliott and Inman (1961) as mentioned above have shown that the volume of the liquor amnii reflects the functional efficiency of the placenta. Dawkins, MacGregor and McLean (1959) suggested that the placental enzyme iso-citric dehydrogenase could be used as an index of placental function. This work, however, has not yet been confirmed, and there is great need for a satisfactory laboratory test to determine placental function.

From the clinical point of view the presence of any one of the conditions listed above should make the obstetrician aware of the possibility of fetal death occurring. Certain clinical observations give an indication of impending fetal death.

1. Weight loss. The average weight gain throughout pregnancy is of the order of 24 pounds, and in the last four weeks of pregnancy weight is gained at the rate of about 1 pound per week. For a long time we have been aware of the significance of a sudden excessive weight gain as an indication of fluid retention and impending toxemia. In placental insufficiency it may be noted that weight gain is not maintained, and indeed the overall gain of weight in pregnancy may be much less than normal. In placental insufficiency it may be noted that weight gain ceases and there may even be an actual loss of weight of 1 or 2 pounds. When this occurs and there is no other explanation for it such as vomiting, fetal death is likely to occur within the next ten days or so. In cases where placental insufficiency seems likely it is desirable therefore to weigh the woman daily, so that on any one day her weight may be compared with that of the week before. A cessation of weight gain or a sustained weight loss may indicate the necessity for early delivery of the child by whatever means is appropriate.

2. Abdominal girth. The girth of the abdomen at term is on the average 40 in., and at 36 weeks 36 in., though due allowance must be made of course for any obesity. If the abdominal girth, which has been increasing steadily, begins to diminish, this again is an indication that the child should be delivered within a week or so.

3. Diminution in the amount of liquor. Wrigley (1939) in discussing postmaturity pointed out that the same observer palpating the uterus daily may detect a diminution in the amount of liquor. This may indicate the necessity for delivery. Elliott's method of determination of the liquor volume is not at present universally applicable.

Management

When placental insufficiency is suspected and the fetus is still premature, it is advisable to keep the woman at rest in bed, so as to favour the placental circulation. On the whole diuretics are best avoided as their use may mimic weight changes due to placental deterioration and so lead the obstetrician to effect delivery before it is necessary. The chances of survival of an infant born prematurely depend on the stage of gestation rather than on its birth weight, which in any case will not increase if the placenta cannot keep pace with fetal demands. Decision as to the best time to effect delivery demands a nice obstetric judgement, weighing on the one hand the risks of prematurity against those of placental insufficiency on the other. Taking these all into account, the time comes when delivery seems imperative if a live child is to be secured.

In many cases surgical induction of labour is the method of choice. When this is performed by artificial rupture of the membranes, careful note should be taken of the volume, consistency and appearance of the liquor so obtained. If the liquor escapes freely and is colourless, and there is plenty of it, placental insufficiency is not likely and delivery can be awaited calmly. On the other hand when the liquor is scanty, thick and stained with meconium the fetus is already at risk because of placental insufficiency, and a special watch should be maintained on the fetal heart until the child is safely delivered. It should be remembered that the contractions of labour themselves impair placental function, and may be the last straw for a fetus already embarrassed by placental insufficiency from some other cause such as toxemia or prolonged pregnancy. For the same reason a 'pitocin drip' should be avoided. When the infant is an especially precious one, e.g. in the elderly primigravida, it may be deemed desirable to proceed straight away to Cesarean section in the interests of the child.

Finally, let me emphasize that the combination of two or more of the conditions which produce placental insufficiency is particularly hazardous.
for the fœtus, and the obstetrician should be ready to intervene by whatever means are appropriate in order to secure a living child.

Summary
1. There is a considerable amount of clinical and experimental evidence to suggest that placental insufficiency is a common cause of intra-uterine fœtal death.
2. Placental insufficiency may occur in a variety of clinical conditions, of which toxaemia and prolonged pregnancy are the commonest.
3. Age, parity, physical exercise, and labour also affect placental function.
4. Diagnosis is mainly based on clinical observation. There may be failure of the woman to gain weight (or loss of weight), or failure of increase of abdominal girth (or decrease), or diminution in the amount of liquor amnii. All three often are detected together. Laboratory determination of the rate of pregnanediol excretion or aœstriol excretion may help, as may the calculation of the volume of liquor amnii.
5. Induction of labour, or in some cases Cesarean section, is indicated, taking into account the risks of prematurity.
6. The combination of two or more conditions, each of which in itself conduces to placental insufficiency, may be lethal for the fœtus unless delivered without delay.

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