Nutrient insult in early pregnancy

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Nutrient insults in early pregnancy, such as nutrient deprivation during famines, are often associated with an unfavourable outcome. Suboptimal nutrition in the early stage of gestation has been linked to a number of adverse effects on fetal growth and development. Historically, nausea and vomiting in pregnancy (NVP) was an important contributor to pregnancy-related mortality; indeed, Charlotte Bronte died from starvation and dehydration after suffering very severe NVP 4 months into her first pregnancy (Gaskell, 1858). Although NVP seldom now progresses to be life-threatening, it affects the majority of pregnant women, and potentially presents a challenge to nutrient intake in the most vulnerable period of development. Symptoms range from mild (nausea only) to severe (a level of vomiting that restricts nutrient intake and ultimately threatens metabolic and electrolyte balance). Although NVP has been documented for thousands of years, its cause has not yet been satisfactorily elucidated, but seems to be related to endocrinological changes. Pregnant women also frequently report dietary cravings and aversions during pregnancy which can be linked to both the incidence and severity of NVP. Paradoxically, NVP appears to be positively associated with a favourable outcome of pregnancy, including increased birth weight and gestational age. The mechanisms by which NVP favours the outcome of pregnancy are not known. They may be related to women increasing their nutrient intake to alleviate symptoms, improving the quality of their diet or reducing energy expenditure. Alternatively, adaptation to a reduced nutrient intake might stimulate the expression of growth factors and affect placentation or metabolism, thus favouring fetal growth when NVP resolves.

Nausea and vomiting in pregnancy: Nutrient deprivation: Fetal development:
Insulin-like growth factor: Maternal nutrition

Nausea and vomiting in pregnancy

Nausea and vomiting in pregnancy (NVP) affects 50–90% of pregnant women (Broussard & Richter, 1998). It usually presents as one of the first signs of pregnancy, within 2–4 weeks of fertilisation, peaks at the end of the first trimester and resolves by week 20 (Anderson, 1994). NVP is usually experienced as a complication of the first trimester of pregnancy, but a small proportion of women experience symptoms throughout pregnancy (Broussard & Richter, 1998). Although the onset of symptoms occurs typically in the morning, thus generating the term ‘morning sickness’, many women are affected episodically throughout the day (Gadsby et al. 1993). Our retrospective study in Guildford (Al-Rasasi et al. 2001) shows that 72.6% of pregnant women reported experiencing NVP, which is similar to the incidence seen in other studies. In this group of women 12.3% suffered only in the morning, 7.5% only at night, whilst the remaining 80.2% of the group either experienced symptoms continuously or episodically through the day; so NVP is clearly a more suitable term than morning sickness.

NVP is usually classified as mild (nausea only), moderate (nausea and vomiting) or severe (where the extent of vomiting is so severe and prolonged that it can induce maternal weight loss, electrolyte imbalance and dehydration; hyperemesis gravidarum). In the group of women

Abbreviations: hCG, human chorionic gonadotrophin; IGF, insulin-like growth factor; IGFBP, insulin-like growth factor-binding protein; NVP, nausea and vomiting in pregnancy.

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we studied, 57.5% could be classified as experiencing mild NVP, 37% moderate and 5.5% severe, including 14% diagnosed with hyperemesis gravidarum and hospitalised. However, within each of these classifications a range of symptoms resulting in quite different effects on dietary intakes might be experienced. For instance, a woman who experienced vomiting might only be affected at a certain period of the day, and although the NVP would be classified as moderate, her daily nutrient intake might have been less affected than that of a woman who experienced mild NVP continuously through the day.

NVP has been positively associated with a number of maternal characteristics. These characteristics include primiparity, younger maternal age and low educational achievement (Klebanoff et al. 1985), multiple gestation and intolerance of oral contraceptives (Jarnfelt-Samsioe et al. 1983), high maternal BMI (Klebanoff et al. 1985), high placental weight, non-smoking, family history and history of NVP in previous pregnancies (Gadsby et al. 1997). It is reported more often in Westernised countries (Broussaud & Richter, 1998) and is associated with poorer diets, irregular eating patterns and stressful pregnancies (Iatrakis et al. 1988). Women who report their occupation as housewife have also been found to have an increased incidence of NVP compared with those who are in paid employment (Weigel & Weigel, 1988).

Although NVP has been recorded as a complication of pregnancy for over 4000 years (Fairweather, 1968), surprisingly little is known about its aetiology. Frustratingly, for many years most obstetricians attributed psychological factors, particularly negative personal relationships or undesired pregnancies, as the prime causes of NVP (Semmens, 1971). Now we know that a number of physiological factors are implicated. The symptoms of NVP follow a time-course that mirrors the secretion of placental human chorionic gonadotrophin (hCG; Soules et al. 1980), but there are no consistent patterns in levels of hCG and severity or duration of symptoms. Both oestrogen and progesterone levels increase during pregnancy. An increased sensitivity to oestrogen is implicated because women who report intolerable levels of nausea with oral contraceptives have a higher incidence of NVP (Jarnfelt-Samsioe et al. 1983). Both overweight women and those who do not smoke have elevated oestrogen levels and a higher incidence of NVP (Depue et al. 1987). However, again there is no correlation between hormone levels and symptoms.

The relationship between NVP and progesterone is related to the effects that progesterone has on smooth muscle tone of the gut that results in gastric hypotonicity, hyposecretion and hypoperistalsis, all of which promote gastric wave dysrhythmia and delayed gastric emptying. The increased progesterone levels of pregnancy reduce the patency of the lower oesophageal sphincter, thus increasing gastric reflex and affecting motility and emptying of the stomach. It has been found that protein-rich test meals (i.e. protein energy ratio (proportion of energy from protein) > 53 %), which seem to lessen symptoms of NVP, reduce gastric slow wave dysmotility (Jednak et al. 1999); however, this finding has been contested by Maes et al. (1999) who found that gastric emptying of solids was not significantly delayed in pregnant women.

The severity of symptoms of NVP has also been linked to altered thyroid function (Mori et al. 1988); a decrease in thyroid-stimulating hormone and an increase in thyroxine were measured. This change may occur because hCG, which has some structural similarities with thyroid-stimulating hormone, acts as a thyroid-stimulating factor. However, hyperthyroidism per se is not associated with nausea and vomiting outside pregnancy.

Other suggested causes of NVP include unidentified nutrient deficiency, vitamin B6 deficiency (Emeljanova et al. 1999), high prepregnancy intake of saturated fat (Signorello et al. 1998) and a pattern of food aversions before pregnancy (Crystal et al. 1999).

The effects of nutrient deprivation in early pregnancy

A woman’s reproductive capability is highly susceptible to nutrient insult or deficiencies during the period of ovum maturation and early embryonic development. During the Dutch Hunger Winter there was an increase in perinatal mortality and congenital abnormality in pregnancies where women were exposed to the famine early in gestation (Stein et al. 1975). Work from the Borough of Hackney in East London (Doyle et al. 1990) has provided convincing evidence of an association between maternal nutritional status and birth outcome, which can only have had its origin in early pregnancy. In these studies women are also likely to have been poorly nourished both before conception and throughout pregnancy rather than only in the early stages of gestation.

The early in utero period of growth and development of the human fetus represents a period of rapid cell division, occurring at different times in different tissues, resulting in the concept of fetal programming. During these critical periods of selective tissue growth, the nutritional and hormonal environment may exert influences on the underlying programmed changes, resulting in reduction in cell numbers, changes in the distribution of cell type and in the resetting of hormonal feedback (Godfrey & Robinson, 1998). The current evidence suggests that, although the fetal genome determines growth potential in utero, it plays only a minor role in the determination of growth that is actually achieved, although little is understood about the maternal influences (including nutrition) which programme the growth and development of the human fetus. The concept of fetal programming has important implications, because it is now well established that variations in fetal size and body proportions at birth have important connotations in terms of health outcomes and pathological changes in adult life.

The potential effect of different planes of maternal nutrition at different stages of pregnancy and the subsequent influence on the growth of the fetus is a complex subject. Barker (1995) hypothesised that undernutrition in early gestation would lead to a proportionately small baby predisposed to hypertension and haemorrhagic stroke, but not CHD, in adulthood. Lumey (1998a) contested this hypothesis after analysing data from the Dutch Hunger Winter and finding little effect on birth weight after undernutrition in the first and second trimester. There is compelling evidence from the Dutch Famine studies, when food availability was progressively reduced to < 5400 kJ/d
by April 1945, that exposure to famine in the first trimester of pregnancy had irreversible effects on the offspring which have become evident in adulthood (Stein et al. 1975). The effects of maternal undernutrition in early gestation include higher levels of obesity (Ravelli et al. 1999), more atherogenic lipid profiles and increased prevalence of CHD (Roseboom et al. 2000a), and altered hepatic function (Roseboom et al. 2000b) in adulthood. Interestingly, from a parallel study, there is evidence of compensatory placental growth after famine exposure in the first trimester of pregnancy (Lumey 1998b). It has been argued, however, that these historical observations do not represent the norm, and that thinness at birth is a result of suppressed placental development early in pregnancy (Godfrey et al. 1997).

More recent evidence of the influence of maternal nutrition in pregnancy in relation to placental and fetal growth has suggested that birth weight and the placental ratio (placental weight–birth weight) are affected early in gestation (Godfrey et al. 1996; see Table 1).

Women who consumed higher than normal amounts of food in early pregnancy (expressed by carbohydrate intake) had small placentas and infants with lower birth weights, especially if combined with low intakes of animal protein in late pregnancy (Godfrey et al. 1996). However, long-term effects following suboptimal nutrition in early gestation may result in altered body proportions or even occur in the absence of an effect observable at birth. Mild maternal dietary impairment in sheep demonstrates that when fetal and compensatory mechanisms are adequate, so that birth weight is not reduced, fetal cardiovascular development is altered (Hawkins et al. 2000). The effects of undernutrition in early pregnancy probably depend on both the magnitude of the deprivation and nutritional state before and afterwards.

It may seem paradoxical, therefore, that although NVP apparently presents a nutritional challenge early in pregnancy, a number of studies have reported that NVP in the first half of pregnancy is associated with a favourable outcome. The risks of miscarriage (Weigel & Weigel, 1989), perinatal death, low birth weight and premature delivery (Tiernan et al. 1986) and congenital heart defects (Boneva et al. 1999) are all reduced in women who experience NVP. However, very severe NVP or hyper-emesis gravidarum, especially if associated with loss of prepregnancy weight, has been associated with less favourable outcomes (Tsang et al. 1996). Although NVP is associated with a favourable outcome of pregnancy, it frequently has negative effects on quality of life during the pregnancy. It causes discomfort and is associated with irritability and tiredness, which may affect social and economic functioning, particularly employment.

**Possible mechanisms**

One possible mechanism by which NVP is associated with a favourable outcome of pregnancy is that despite causing discomfort and distress associated with food, NVP may not result in reduced nutrient intake, but may actually increase it or alter the macronutrient profile to favour the consumption of certain nutrients. We have found that some women (31 %) reported that their symptoms were alleviated by continually snacking, usually on carbohydrate-rich foods. Another possibility is that NVP stimulates a change in physical behaviour so that women change their level of activity, and hence energy expenditure will alter the balance in favour of maternal and fetal tissue growth.

Alternatively, quality of diet might be altered. Our studies have found a strong association between symptoms of NVP and reported dietary cravings and aversions (Al-Rasasi et al. 2001). Women reported experiencing cravings, particularly for fruit and sweet-tasting foods, dairy products and protein-rich foods, whereas the most commonly reported aversions were for drinks containing caffeine, strong tasting and smelling foods and fatty or greasy foods. These cravings and aversions may be linked to altered taste perceptions in pregnancy. Indeed, it has been shown that there are differences in bitter-taste perception in women with a history of severe vomiting during pregnancy (Sipiora et al. 2000).

Deutsch (cited in Flaxman & Sherman, 2000) proposed that NVP had evolved as a method of communication which alerted the pregnant woman’s partner and kin to her pregnancy. Although it was suggested that NVP might discourage intercourse or signal impending need for additional food or protection, it may simply constantly remind the woman of her own pregnancy and induce her to change behaviour. Women do appear to positively change their diet in pregnancy in response to public health messages (Anderson, 2001).

Another intriguing explanation may be that NVP, which coincides with the most sensitive periods of embryonic organogenesis, protects the developing embryo by causing women to reject or avoid foods containing potentially-harmful teratogens or abortifacients (Hook, 1980; Profet, 1988). Foods implicated include strong-tasting vegetables, beverages containing alcohol and caffeine, and meat, fish, poultry and eggs. This ‘Stone Age theory’ has been strongly contested by Brown et al. (1997). However, Flaxman & Sherman (2000) investigated the concept further and hypothesised that NVP causes women to avoid foods that are more likely to be contaminated with parasites and pathogens at a time when pregnancy causes immunosuppression, leaving women more vulnerable to infection. This concept was supported by the findings that societies (e.g. Bhil, Mbundu, Omaha, Papago, Siriono, Tarahumara and Woleai)

![Table 1. Mean birth weight (adjusted for gender; g) in 538 pregnancies according to the mother’s daily intake of carbohydrate in early pregnancy and meat protein intake in late pregnancy (intakes calculated by food-frequency questionnaire; Godfrey et al. 1996)](https://doi.org/10.1079/PNS2001136)

<table>
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<th>Carbohydrate intake (g/d)†</th>
<th>&lt; 265</th>
<th>265–340</th>
<th>&gt; 340</th>
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<td>3419</td>
<td>3321</td>
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<tr>
<td>All</td>
<td>3501</td>
<td>3444</td>
<td>3381</td>
<td>3442</td>
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* Birth weight fell by 3·1 g for each 1 g decrease in meat-protein intake in late pregnancy.
† Birth weight fell by 165 g (P = 0·005) for each log 1 g increase in carbohydrate intake in early pregnancy.
where NVP had never been observed were more likely to have only plants as their dietary staple. Flaxman & Sherman (2000) analysed the results of twenty studies and found that the most common aversions were to meat and non-alcoholic beverages.

Placental development

Much of what we know about the effects of maternal nutrition on fetal and placental growth and development is based on animal models. In experimental animals effects of undernutrition on placental weight are particularly variable (Harding & Johnston, 1995). Fetal and placental responses to undernutrition in ewes depend on the magnitude, timing and duration of nutrient restriction (Symonds et al. 2001). Maternal nutrient restriction in ewes in early gestation tends to increase placental weight at term. In early gestation, during the period of rapid placental growth, maternal nutrient restriction initially restricts placental growth (Clarke et al. 1998), but when the mother is subsequently fed to requirements the placental size increases and fetal growth is restored (Heasman et al. 1999). These experiments performed in sheep have demonstrated that severe undernutrition reduces placental mass, but lesser reductions in maternal intake have the opposite effect (Robinson et al. 1995; Gadd et al. 2000). In contrast to the discoid structure of the human placenta, sheep have cotyledonary placentas in which villus contact is established in a number (forty to 100) of small areas called placentomes. Each placentome is formed by the projection of interdigitating fetal villi into specialised predetermined regions of the maternal stroma called caruncles. In the nutrient-restricted group of sheep both placental weight and the number of placentomes were significantly increased ($P < 0.05$; Heasman et al. 1999). The practice of ‘flushing’ in sheep husbandry, in which sheep are transferred from good pasture to poorer pasture just for the early period of gestation, appears to promote fetal growth by stimulating placental development (McCrabb et al. 1992).

Other studies have shown that overfeeding adolescent (still growing) sheep early in gestation promotes rapid maternal growth, but results in a restriction of placental mass and a significant ($P < 0.001$) decrease in birth weight which is related to nutritional status rather then gynaecological immaturity (Wallace, 2000). In these overnourished adolescent sheep an altered maternal hormonal profile apparently promoted maternal tissue anabolism at the expense of fetal growth (Wallace et al. 1997). In contrast, adolescent ewes fed moderately during the first trimester had significantly higher fetal cotyledon numbers than those fed on a higher nutritional plane ($P < 0.007$; Wallace et al. 1999). From studies in sheep, it seems that the placenta undergoes an adaptive compensatory response to mild maternal undernutrition during this period of rapid placental growth, thus optimising transplacental exchange efficiency. It is possible that NVP is associated with a favourable outcome of pregnancy because maternal nutrient restriction early in gestation favours placental development. Can we extrapolate these findings to man?

The insulin-like growth factor axis

The mechanisms by which maternal nutrient intake affect fetal and placental growth seem to be mediated by the insulin-like growth factor (IGF) axis. The IGF are low-molecular-weight polypeptides (7 kDa), with structural homology to proinsulin, which promote mitosis and differentiation. They play an important role in determining fetal and placental growth (van Kleffens et al. 1998). The IGF and their binding proteins (IGFBP), which are produced by the placenta, act as autocrine or paracrine factors at or close to their site of synthesis. Several components of the IGF system, including IGF and IGFBP, have been demonstrated to be affected by nutrition. The six homologous IGFBP have high binding affinity for IGF-I and IGF-II (Baxter, 2000) which may be altered by modification of IGFBP such as phosphorylation and proteolysis. Binding of IGF to IGFBP can result in either inhibition or enhancement of IGF action. The actions of IGF-I and IGF-II are influenced both by levels of the specific IGFBP and by the expression of IGF receptors on the target tissues. Studies using different types of knockout mice have led to the identification of the roles of various components of the IGF axis and their effects on fetal growth in utero.

Both maternal and fetal IGF-I levels affect fetal growth rate (Gluckman, 1995). IGF-I predominantly influences growth in late gestation and postnatally, whereas IGF-II has a stronger influence on embryonic growth in early development (Fig. 1). Both human fetal and neonatal size correlate with circulating IGF-I concentrations in cord blood at term (Gluckman, 1995), and IGF-I levels are sensitive to maternal nutrient intake. Placental glucose transfer controls fetal insulin release, which regulates IGF-I release and, therefore, fetal growth; IGF-I also affects placental metabolism and controls placental substrate delivery to the fetus. There is a switch during development from IGF-II to IGF-I, but the actions of both IGF on fetal growth are primarily mediated by the type-1 receptor (Gluckman & Harding, 1997). IGF-II seems to require much greater changes in nutritional and hormonal factors than IGF-I before plasma and tissue levels are affected (Straus & Takemoto, 1990). This situation means that in early development, when IGF-II is the dominant growth factor, the fetus is less sensitive to environmental influences such as nutrient deprivation. Later in gestation maternal fasting causes a decrease in IGF-I and increases fetal liver expression of IGFBP-1, which contributes to fetal growth retardation (Straus et al. 1991). Thus, when IGF-I is the predominant influence the fetus responds to nutrient deprivation by promptly decreasing growth rate.

All six IGFBP are produced by human decidualised endometrium. IGFBP-1 is the main secretory product of the human decidualised endometrium and is the predominant IGFBP in the amniotic fluid and fetal plasma (Drop et al. 1984). Concentrations of IGFBP-1 increase during the first trimester and plateau at mid-gestation, and are higher in intrauterine growth retardation (Giudice et al. 1997). Both IGF-I and IGF-II regulate IGFBP-1 in a biphasic manner; low concentrations stimulate IGFBP-1, while high concentrations are inhibitory (Westwood, 1999). Production of
IGFBP-1 is inhibited by insulin and stimulated by progesterone and relaxin. The affinity of IGFBP-1 is affected by its phosphorylation state; phosphorylated IGFBP-1 has a higher affinity for IGF-I and so can inhibit IGF-I activity (Westwood, 1999). The IGFBP-1 regulated by nutritional state to different extents; IGFBP-1 is regulated acutely by nutrient intake, with high levels in the fasted state and rapid postprandial decreases. IGFBP-3 is regulated acutely by nutrient intake, with high levels in the fasted state and rapid postprandial decreases. IGFBP-3 is relatively stable, but is depressed after prolonged periods of severe malnutrition, and IGFBP-2 is highly dependent on dietary protein intake (Ketelslegers et al. 1996).

The effects of the IGF are mediated via two specific receptors, type-1 and type-2, which are expressed in high density on most fetal and placental cells. The type-1 and type-2 receptors co-localise with IGF-II, which suggests the receptors compete for IGF-II (Zhou & Bondy, 1992). IGF-II interacts with type-I and type-2 IGF receptors; the type-2 receptor binds IGF-II with high affinity and IGF-I with an affinity about 100-fold lower (Han & Carter, 2000). While the type-1 receptor promotes growth, the competitive non-signalling type-2 receptor limits growth (Czech, 1989); it is also implicated in IGF-II degradation (O’Dell & Day, 1998). A soluble circulating form of the type-2 receptor inhibits IGF-II-mediated DNA synthesis and therefore probably constrains fetal growth (Ong et al. 2000). Size at birth correlates with circulating IGF-II:soluble type-2 receptor levels (Ong et al. 2000).

It has been suggested by Haig & Graham (1991) that, as the gene for the type-2 receptor is maternally imprinted and IGF-II is paternally imprinted (DeChiara et al. 1991), this situation presents an example of genetic conflict. Paternal genome expression promotes growth via the expression of IGF-II, but the maternally-expressed type-2 receptor can act to mop up IGF-II, preventing it from binding to the growth-promoting type-1 receptor. Thus, an excess of type-2 receptor expression will limit growth of placenta and, therefore, constrain fetal growth, thus limiting the metabolic demands placed on the mother.

**Nutrient regulation of growth**

The preimplantation embryo bathes in fluid that is rich in IGF and IGFBP (van Kleffens et al. 1998). IGF-II is also expressed very early after fertilisation and has been demonstrated in the two-cell-stage embryo (Heyner et al. 1989). Fetal tissues overexpress IGF-II compared with postnatal tissues. Successful implantation, placental development and fetal growth depend on migration of the IGF-II-producing trophoblast into the maternal decidua (Minniti et al. 1992). IGF-II is produced in abundance by the trophoblast cells of the placenta (van Kleffens et al. 1998), and is particularly prevalent in sites of cell differentiation; its concentration is highest in the trophoblastic columns of anchoring villi, especially at the leading edge, implying it plays a pivotal role in trophoblastic invasion (Westwood, 1999). The maternal decidua and uterine vessels express IGFBP-1, which is involved in orchestrating trophoblast migration (Westwood, 1999). This interaction between IGF-II from the cytotrophoblast and decidua IGFBP at the maternal–fetal interface of the primate placenta is thought to be critical during trophoblastic invasion and decidualisation (Han & Carter, 2000). IGFBP-2 is not detectable early in gestation (Reynolds et al. 1997). There appears to be a functional relationship between IGF-II and IGFBP-2; during the early invasive period when the placenta is establishing, IGF-II expression is high and unopposed by IGFBP-2, whereas later in gestation the ratio is reversed (Zhou & Bondy, 1992).

IGF-II-knockout mice have significant growth retardation, especially in the early stages of gestation; IGF-II gene disruption is associated with severe placental growth retardation (DeChiara et al. 1990). IGF-II appears to be
involved in the regulation of body composition in the mouse; it controls fluid uptake by direct action on the maternal capillaries (Gardner et al. 1999), increasing vascular endothelial growth factor, which increases capillary permeability and promotes angiogenesis. Vascular endothelial growth factor increases NO production, which is positively correlated with birth weight and placental weight (Hata et al. 1998). Other processes mediated by IGF-II are thought to include increased cell mass (Zaina & Squire, 1998) and cell survival (Christofori et al. 1994) and rate of progression through the cell cycle (Eggenschwiler et al. 1998). IGF-II also regulates the cell number in the placenta and plays a role in the differentiation of placental glycanogen-producing cells (Lopez et al. 1996).

In summary, the mechanism by which NVP is associated with a favourable outcome of pregnancy, particularly increased birth weight, may be mediated through adaptation to reduced nutrient intake in early gestation, favouring compensatory placental development, which subsequently optimises fetal development. During the early stages of development, when trophoblastic invasion is occurring and the placenta is establishing, the predominant growth factor involved in promoting growth via the type-1 receptor is IGF-II, which is fairly resistant to fluctuations in maternal nutrient intake, thus placental growth is maintained. At this time IGFBP-2 is not detectable in the fetal circulation, thus potential competition with the type-1 receptor is minimal. It is necessary to be cautious when drawing conclusions from mechanisms elucidated in different species. The structures of the primate discoid placenta and the cotyledonous placenta of ruminants are very different. None of the species studied share identical expressions of IGF and IGFBP. Also placental weight per se is a gross measurement; weight alone does not necessarily indicate placental efficiency of nutrient transfer, which is likely to be affected more by depth of insertion or differential expression of transport mechanisms rather than by weight.

**Nausea and vomiting in pregnancy**

An alternative explanation of the mechanisms is that NVP affects nutrient partitioning via altered metabolism. During the first half of pregnancy increased sensitivity to insulin promotes maternal anabolism. Subsequently, in the latter part of pregnancy increasing insulin resistance stimulates maternal catabolism and maintains higher substrate levels, favouring placental transport when fetal growth is high. Altering intake in early gestation could moderate the extent of deposition of maternal stores. A high nutritional intake in adolescent sheep early in gestation resulted in suppressed placental and fetal growth (Wallace et al. 1997). This phenomenon was suggested to be mediated by elevated insulin levels and maternal IGF-I, which promoted maternal tissue deposition in early gestation at the expense of placental development and caused down regulation of placental IGF-I levels. It has been hypothesised that, by causing reduced energy intake in early gestation, NVP would lower maternal insulin and IGF-I levels, thus abrogating the anabolic drive and ensuring that nutrient partitioning favoured placental, and ultimately fetal, growth (Huxley, 2000). As insulin potentially inhibits hCG production (Barnea et al. 1993), reducing maternal insulin levels would optimise hCG production and subsequent effects of thyroxine on placental development. Huxley (2000) also suggests that the positive effect of NVP on placental development could be potentially augmented by placental leptin suppressing appetite. Placental production of leptin peaks in the first trimester (Masuzaki et al. 1997); thus, it could either augment the effect of NVP or provide an alternative mechanism for suppressing maternal energy intake during placental development. Maternal leptin concentration is negatively correlated with placental size (Schubring et al. 1997). Maternal dietary intake is also inversely related to peripheral progesterone concentration, so that a lower nutrient intake in early gestation facilitates progesterone production, which has positive effects on growth of the embryonic inner cell mass (Wallace, 2000).

**Summary**

Although mild-to-moderate NVP is associated with a positive outcome of pregnancy, favouring both fetal growth and gestational length, this positive effect depends on nutritional status before conception and after the symptoms of NVP have resolved. Providing a woman enters pregnancy with adequate nutrient stores, the fetal growth trajectory will be set optimally and there will be less competition between maternal nutrient requirements and those of the developing conceptus. Indeed, it has been suggested that women of below normal body weight are likely to experience NVP to a lesser extent (Huxley, 2000). Certainly, women who are less well nourished before and during conception tend to gain more weight during the course of pregnancy and to produce lighter babies (Doyle et al. 1990). Similarly, adolescent women, who presumably have a stronger anabolic drive, also have a poorer outcome of pregnancy (Scholl et al. 1995) in a similar pattern to that of pregnant adolescent sheep (Wallace, 2000).

The other requirement for enhanced fetal development is that dietary intake is adequate to match the transfer capacity of the placenta after symptoms of NVP have abated, during which time fetal growth is constrained by nutrient availability in the second half of pregnancy (Bauer et al. 1998). At this time in gestation IGF-I takes over as the predominant promoter of fetal growth and stimulates maturation of fetal organs. Optimal early development of the placenta could affect subsequent levels of hormones, growth factors and transporter proteins which are produced later to control distribution of substrates and regulate fetal growth. As well as IGF-I, glucose-transporter proteins (Illsley, 2000), placental growth hormone (Alsat et al. 1998) and placental lactogen (O’Dell & Day, 1998) have been shown to be important in regulating both placental substrate availability and uptake, and maternal appetite and metabolism.

As well as causing decreased nutrient intake which may affect either placental development or nutrient partitioning, NVP results in an increased nutrient intake in some women (Fig. 2). It can be hypothesised that this situation may be related to rates of maternal growth before conception. NVP can also potentially result in dietary changes either by the effects of changing intake on severity of symptoms, or because it acts to continuously remind the woman she is...
Fig. 2. Maternal diet, size and maturity affect the severity of nausea and vomiting in pregnancy (NVP). The mechanisms by which NVP is positively associated with the outcome of pregnancy could include (1) increasing or decreasing energy expenditure, (2) reducing intake, which could either alter maternal metabolism and nutrient partitioning in favour of the conceptus and/or stimulate placental adaptation and/or (3) altering the quality of the diet.

Acknowledgements

B.A.-R. is in receipt of a University of Surrey Research scholarship that we gratefully acknowledge. We thank Dr John Nicholls for his support and helpful discussions.

References


