A hypothesis linking low folate intake to neural tube defects due to failure of post-translation methylation of the cytoskeleton

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ABSTRACT Neural tube defects are serious congenital malformations which can be prevented by periconceptional folic acid supplementation. We hypothesize that folic acid provides the methyl group used for post-translational methylation of arginine and histidine in the highly conserved regulatory domains of the cytoskeleton and that these are required for neural tissue differentiation. Presumptive neural tissue has an unusually high need for folates due to the activity of phosphoethanolamine methyl transferase in producing neural tissue specific lipids at a time when the cytoskeleton is also competing for methylation. According to the cell state splitter hypothesis, the cytoskeleton is required to coordinate the spatial and temporal component of differentiation. When folate supply is low and the cytoskeleton is not methylated properly, the result is a neural tube defect due to failure of this coordination.

KEY WORDS: neural tube defect, folic acid, homocysteine, primary neural induction, cytoskeleton

Introduction

Except for heart defects, neural tube defects (NTDs) are the most common form of major congenital malformation with a birth prevalence of 1-5/1000 births worldwide (Frey and Hauser, 2003). NTDs result from a failure of the rising and fusing of the neural folds during the earliest stages of development. In humans, neural tube closure begins in the third week after conception, often before a woman knows that she is pregnant (Northrup and Volcik, 2000). Due to the high morbidity and mortality associated with NTDs, prenatal diagnosis and elective termination of an affected pregnancy has become the option chosen by 80% of families faced with an NTD pregnancy (Burton, 1986, Nicolaides, et al., 1986, Benacerraf, et al., 1989). Individuals with spina bifida will undergo multiple surgeries as they grow in order to treat a host of associated problems. Many affected individuals do not survive to adulthood (Bowman, et al., 2001, Rintoul, et al., 2002). While controversial experimental surgical closure of the defect in utero has shown some promise, it does not represent a cure and the surgery is high risk for both mother and fetus (Bannister, 2000, Holmes, et al., 2001).

History of research into folates and NTDs

The first clue that NTDs might be related to diet came from examining the Dutch famine winter of 1944, caused by Nazi occupiers. There was a doubling of the incidence of spina bifida among those conceived during the famine. The quality of nutrition was later investigated by gynecologist Hibbard (Hibbard and Smithells, 1965) in the 1960s and pediatrician Smithells in the early 1980s (Smithells, et al., 1985), among others. A series of studies soon pointed to multivitamins and folic acid in particular, as having the ability to prevent NTDs (Czeizel and Rode, 1984). The most widely cited study was published in 1991 (Medical Research Council, 1991). It found that supplementing with 0.4 mg of folic acid per day is capable of preventing 70% of NTDs in women with a past history of NTDs and 50% of all NTDs in the general population. Widespread periconceptional supplementation with folic acid has great promise for preventing the majority of isolated NTDs (Eskes, 1998a,b, 2000). Mandatory folic acid food fortification began in the USA on

Abbreviations used in this paper: NTD, neural tube defect; SAH, S-adenosylhomocysteine; SAM, S-adenosylmethionine.

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Homocysteine concentrations are normally inversely proportional (Rosenquist and Finnell, 2001).

Embryology of NTDs

The neural epithelium forms a thick flat plate, which then subdivides into multiple embryonic neural tissue types as it folds and seals (Gordon, et al., 1994). NTDs are most commonly thought to occur due to failure of the sides of the neural tube to rise or a failure of the neural tube margins to seal, or some combination thereof. There are multiple explanations of why this failure or failures might occur, but no one explanation is generally accepted (Gordon, 1985, Anon., 1992, Nieuwkoop, 1999, Candito, et al., 2003, McLone, 2003). The failure of any step of differentiation from the initial step of primary neural induction to the end of neural tube closure might result in an NTD. The relative timing of the different zones of differentiation is also critically important to development. For example, if the inner region of the neural plate has faster development compared to the outer edges of the plate, the edges won’t meet at a time when they are competent to bind to each other. Such localized differentiations are required in the multi-site closure hypothesis. This hypothesis suggests that there are separate gene(s) expressed at each closure site during neurulation (Van Allen, et al., 1993, Van Allen, 1996). Also, if one zone of the neural plate takes up too small or too large a portion of the whole plate, the mechanical properties of the entire plate might change such that the plate cannot fold properly and roll up to close. In general, earlier the failure of differentiation, the more severe the neural tube defect (Jones, 1997). Multifactorial NTDs, which represent about 70% of all NTDs, are thought to be due to a combination of environmental factors and genetic predisposition (Frey and Hauser, 2003), which would have to produce mechanical sequelae of failure of rising and/or sealing of the neural tube.

Folate metabolism and NTDs

Studying the place of folates in embryological functioning has provided important clues about the etiology of NTDs. Folic acid is an artificial form of folate that is readily converted into the metabolically active derivatives of folate such as S-adenosylmethionine (SAM). This methylating intermediate is required in enormous quantities for proper construction and functioning of DNA, proteins and lipids. As developing embryos are rapidly creating new cells, they are especially sensitive to reduced levels of folates. Ectoderm and early neural epithelium tissue may be the most sensitive of all embryonic tissues. This idea is supported by the observation that neural plate epithelium expresses extremely high levels of messenger RNA for folate receptors when compared to other embryonic tissues such as the developing heart (Rosenquist, et al., 1996, Rosenquist and Finnell, 2001).

Neural epithelium is also extremely sensitive to homocysteine. Homocysteine concentrations are normally inversely proportional to the concentration of folate derivatives (Fig. 1). Homocysteine is converted to methionine. The enzyme S-adenosylmethionine synthase (also known as methionine adenosyltransferase), joins a methionine to an adenosine triphosphate by linking the sulfur of the methionine to the 5’ carbon of the ribose unit. This forms the high energy compound SAM (Fig. 2). SAM is the single most important methylating reagent in the cell whose action includes, but is not limited to, the methylation of lysine, arginine and histidine in post-translational modifications of proteins. Critical methylations of DNA, RNA and lipids also require SAM (Miner, et al., 1997).

SAM is converted back to S-adenosylhomocysteine (SAH) when the methylation reaction occurs. Several different enzymes catalyze this methylating reaction depending on which compound in the cell requires a methyl group. The product, SAH, is a feedback inhibitor of all the enzymes that create SAM and can therefore be regarded as a potentially toxic waste product. The ratio of SAM to SAH in the cell is tightly regulated in each cell type. In normal adult rat liver, SAH exists at 13 µmol/g while SAM is normally at 60 to 90 µmol/g (Skovby, 1989). In the rat embryo, the difference is even more profound with SAM concentration being 42 times higher than SAH in some tissues (Vanaerts, et al., 1994). After the methylation reaction has occurred, the adenosyl group is normally stripped off, leaving homocysteine. Homocysteine must be quickly condensed to cysteine via the pathway called transsulfuration, or recycled back to methionine (remethylation), or exported from the cell, to avoid causing elevated levels of SAH and the resulting inhibiting activity of SAH on the generation of SAM (see Figs. 1 and 2).

In cultured rat embryos, exposure to a 3 millimol/L concentration of homocysteine in the culture media leads to a drop of the SAM/SAH ratio from 42 to 3.6 (Vanaerts, et al., 1994). Any buildup of homocysteine in the cell results in reversal of the hydrolase reaction that changes SAH to homocysteine. This causes an increase in SAH levels. This will in turn inhibit the production of SAM, reducing methylation reactions throughout the cell and impeding normal cellular functioning. Amounts of four µmol/L of homocysteine in culture media can cause NTDs in chick embryos (Rosenquist, et al., 1996).

There are differences between human males and females in terms of metabolism of methionine (Blom, et al., 1988). Females

![Fig. 1. The structural and enzymatic relationships between methionine, homocysteine and cysteine and the cofactors involved. Arrows represent a single enzymatic reaction. Adapted from Ueland and Refsum, 1989.](image-url)
are more likely to shunt off a portion of the available methionine to 4-methyl-2-oxobutyrate in the presence of increased homocysteine. This is because the enzymes in this pathway appear to be enhanced by estrogen. It is unlikely there is any gender based effect at the time of neural tube closure because the embryo is not yet producing significant amounts of its own estrogen and testosterone at that early stage. However, at later stages of pregnancy, there may be survival advantages for one gender over another. The enhancement the 4-methyl-2-oxobutyrate pathway offered by estrogen may explain the epidemiological differences in prevalence of males versus females among fetuses with NTDs by allowing differential survival of one gender over the other. Females reach term with anencephaly three times more often than males, see (Elwood and Elwood, 1980).

The human embryo/fetus has both less efficient transsulfuration and remethylation abilities when compared to the adult (Sturman, et al., 1970, Skovby, 1989) (Fig. 3). Yet, the human embryo/fetus nevertheless maintains a much higher SAM/SAH ratio in spite of this dual “impairment”. Maintaining the high SAM/SAH ratio in the embryo/fetus can only be accomplished by lowering maternal homocysteine levels. Increasing both transsulfuration and remethylation in the mother, which in fact occurs, can do this. As well, estrogen related compounds enhance removal of methionine from the cycle, which enhances remethylation of homocysteine. Perhaps the dramatic physiologic drop in homocysteine during normal pregnancy of 50% or more (Andersson, et al., 1992, Adams, et al., 1995) evolved as a protective mechanism for nurturing a hyperhomocysteinemia susceptible embryo by allowing it to export excess homocysteine.

Even though high levels of homocysteine are toxic to the embryo, the embryo still needs to import some homocysteine as an essential amino acid. This may explain why women who have enzyme variants that increase homocysteine later in pregnancy have been shown to have fewer cases of intrauterine growth restriction (IUGR) among their babies (Infante-Rivard, et al., 2003). The embryo may also control how much homocysteine it imports from the mother and so only suffer problems if both the mother’s levels are too high to allow homocysteine export and the embryo is metabolically impaired, so it cannot maintain low levels for itself. Women with homocystinuria due to cystathionine β-synthase deficiency appear to have normal pregnancy outcomes (Vargas, et al., 1999, Levy, et al., 2002) indicating that high homocysteine levels in the mother are not necessarily a problem for the embryo.

**Homocysteine and experimentally induced NTDs**

Cow serum must be supplemented with methionine in order to prevent NTDs in rat embryos grown *in vitro* on cow serum during the period of neural tube closure (Coelho, et al., 1989). Cow serum is low in methionine and causes low SAM levels in rat embryos. This indicates a critical period of increased requirement for methionine during neural tube closure in rats. This critical period corresponds to the previously noted increased need for folates in neural plate tissues. It would appear that homocysteine is embryotoxic in a general sense. However, it does not directly cause NTDs. Rather, it is the loss of folate derivatives and reduced SAM that cause NTDs. This hypothesis is supported by recent research (Greene, et al., 2003).

Rat embryos grown on low methionine cow serum and with NTDs, have been shown to have reduced amounts of two post-translational methylations of specific proteins in the neural tissue. Methylation of histidine to form a trimethyl compound was reduced by up to 56%. Methylation of arginine to form a dimethyl compound was reduced by 42%. This effect was not seen in normally developing, but methionine deprived, heart tissue from the same embryo. Methylation of histidine and arginine was normal in embryos that were methionine supplemented and that subsequently developed normally without NTDs (Coelho and Klein, 1990). Why is this specific decrease in methylation re-
restricted to rat embryo neural tissue? NTDs may represent a borderline case of SAM deficiency. Too great a deficiency results in death of the embryo. A borderline deficiency affects only embryonic neural tissue because this is the only tissue with elevated SAM demands during the critical period of neural tube closure, at least in rats.

Polyunsaturated fatty acid synthesis may contribute to NTDs

In adult rats, SAM provides the methyl group for a brain specific phosphoethanolamine methyl transferase (PEMT) enzyme, which preferentially methylates polyunsaturated fatty acids. Most polyunsaturated fatty acids used in the adult are produced in the liver and transported through the blood stream to wherever they are needed (Vance, et al., 1997). PEMT2 is the version of PEMT normally expressed only in adult liver. PEMT2 knockout mice develop normally, from a behavioral perspective, if given choline supplements, because extra phosphotidylcholine is produced via the alternate pathway of phosphorylation of choline. They do have some anatomical brain abnormalities even though their behaviour is normal (Merouani, et al., 2001).

PEMT activity also occurs at very low levels in some other tissue types. There is a brain specific PEMT enzyme required which must exist because polyunsaturated fatty acids cannot pass the blood brain barrier, yet they accumulate in brain tissue (Vance and Ridgway, 1988). This brain specific PEMT has been isolated in newborn rat brain tissue. The PEMT enzyme of neonatal rat brain is different from the adult rat version in the liver. The neonatal PEMT has a high affinity for SAM and methylates polyunsaturated fatty acids four times faster than the adult liver version. Its concentration also increases 2-fold from neonatal days 5 to 20, a period of rapid rat brain growth (Blusztajn, et al., 1985).

Recent research has shown that PEMT activity in the adult liver regulates homocysteine levels. Thus it is not implausible to suggest that there may be a form of PEMT in developing neural tissue that creates polyunsaturated fatty acids in embryonic neural tissue and that this enzyme may also regulate homocysteine in the neural tissue. Certainly changes in homocysteine levels result in changes in brain membrane composition in chick embryos. Increases in homocysteine levels result in decreases in the levels of polyunsaturated fatty acids. The likely mechanism is through inhibition of PEMT by elevations in SAH levels (Noga, et al., 2003).

There is an extremely high demand for neural lipids during early development. The proportion of the neural plate that is made up of cell membrane increases dramatically during neural plate formation and neural tube closure. Individual neural plate cells are both elongating and rapidly dividing (Jacobson and Gordon, 1976, Jacobson and Tam, 1982, Zhu, et al., 2004). This indicates there would be a high demand on any embryonic version of PEMT that might be present in early neural tissue. Normally some of the choline stored in yolk platelets is used for producing phosphorylated choline derivatives such as sphingomyelin. Some of the choline is also converted to betaine, which then acts as a methyl donor for methylation of homocysteine to methionine. This then increases available SAM. In addition, some limited amounts of phosphotidylcholine are normally produced by PEMT during gastrulation and neurulation. However, the betaine pathway is not active until the end of neurulation (Fisher, et al., 2001, 2002). This could then explain why neural tissue is so sensitive to reduced availability of folate derived SAM. During neurulation SAM is only available via the folate pathway. SAM is perhaps also being used up by high affinity PEMT for production of phosphotidylcholine first. Other required methylations such as methylation of histidine and arginine in proteins, cannot be completed because of the lack of methyl groups if there is a shortage of folate.

There are three components of the cytoskeleton that are normally methylated using SAM. These are actin, α and β tubulin and neurofilament L. The active site of actin and myosin binding has a highly conserved 3-methylhistidine residue. This residue is actively methylated by SAM during neural tube closure. Reduced methylation results in failure of these cytoskeletal elements to localize in the basal and apical ends of the cells (Moephuli, et al., 1997). Failure of appropriate localization of the cytoskeletal elements of neural tissue would result in mechanical failures of cell contractions and movements and thereby cause NTDs. Therefore, reduced methylation of cytoskeletal elements is an ideal candidate for reduced methylation causing changes in differentiation that affect the developing neural plate, which then leads to an NTD.

Cell state splitter and NTDs

This hypothesis suggests a direct connection between the cytoskeleton and differentiation. While no one model of differentiation is accepted universally, there is a model of neural tissue differentiation that is based on cytoskeletal functioning. This model is called the “cell state splitter model” (Gordon and Brodland, 1987, Gordon, 1999).

Development is generally considered to be a series of bifurcating choices to produce each new cell type. For example, the ectoderm is changed to two new types of tissue, presumptive
neural epithelium and presumptive epidermal cells. In classical embryological terms, the ectoderm is considered to be on a pathway to forming epidermis unless “induced” to form neural tissue. A signal that induces this alternate pathway of development is presumed to come from the dorsal lip of the blastopore during early gastrulation while mesoderm and endoderm is being tucked inside the embryo and ectoderm is expanding to cover the whole exterior. The primary neural induction signal spreads over half of the ectoderm. The ectoderm changes in response to the induction signal. It begins expressing new gene products unique to neural tissue. The other half of the ectoderm does not receive the induction signal. It, therefore, follows the default program, which is to change gene expression to form the epidermis. Thus, there are two new tissues arising out of one earlier tissue. By continually repeating this process throughout the embryo and in many different tissue types, the entire embryo forms (Schoenwolf and Smith, 1990).

The cell state splitter model of differentiation postulates that the combination of actin, microtubules and intermediate filaments, seen at the apical end of the neural plate cells using electron microscopy (Martin and Gordon, 1997), forms a bistable organelle (the cell state splitter) that is responsible for sensing and then propagating a contraction or expansion wave across presumptive neural epithelium (Fig. 4). The bistability of the device is presumed to account for the binary nature of differentiation, with one nuance: the previous state is not necessarily continued, but one of two new cell states (cell types) is produced when the cell state splitter changes to one of two new stable forms (Gordon, 1985, Gordon and Brodland, 1987, Gordon, 1999).

Many waves of contraction have been observed in the open neural plate of the developing axolotl (Ambystoma mexicanum) (Björklund and Gordon, 1994). The axolotl ectoderm also has its own unique contraction wave and a corresponding expansion wave. The ectoderm contraction wave begins at the dorsal lip of the blastopore at the same time and place that primary neural induction is known to start. Its trajectory is limited to the same region of the ectoderm that is affected by classical primary neural induction, i.e. the presumptive neural plate. After this contraction wave has ended, an expansion wave begins at the bottom of the embryo and travels upward over the region of ectoderm that forms presumptive epidermis. This expansion wave ends at the border of the newly forming neural plate. Both these observed contraction and expansion waves are hypothesized to begin propagating due to a specific mechanical signal (Gordon and Brodland, 1987).

In the case of the axolotl’s ectoderm contraction wave, the invaginating pharyngeal endoderm touches the underside of the ectoderm, probably inducing the contraction wave. This particular wave travels over that entire presumptive neural developmental field, moving from cell to cell and thereby propagating the induction signal. The newly forming neural plate cells narrow apically and elongate perpendicularly to the apical surface. The neural plate thereby exerts mechanical pull on the remaining ectoderm tissue in the lower half of the ectoderm. This is hypothesized to start an expansion wave over the remainder of the ectoderm carrying a different signal in this region (Björklund and Gordon, 1994, Gordon, et al., 1994).

The cell state splitter model proposes that while each individual cell is contracting or expanding, a simultaneous chemical signal is sent to the nucleus indicating that the cell has participated in a wave and what kind (Björklund and Gordon, 1993). The nucleus responds to the expansion or contraction wave signal with changes in gene expression that correspond to differentiation into the next cell type (Fig. 5). The trajectory of each wave is shaped and limited by simple mechanical forces in cell sheets. The final stage of differentiation of an individual cell is the preparation of a new cell state splitter capable of responding to and propagating the next wave (except for terminally differentiated cells).

**Nuclear state splitter and NTDs**

Contraction and expansion waves provide the spatial and physical component that is missing from most other embryological models (Gordon, 1985, Schoenwolf and Smith, 1990). The classical primary neural induction model is unchanged except that the inducer is a mechanical signal sensed and propagated by cytoskeleton to adjacent cells. The cytoskeleton is specified in this model as that which transduces the mechanical induction signal into a chemical signal. The nucleus senses and responds to the transduced chemical signal, changing its state of gene expression. We therefore speak of a “nuclear state” and “nuclear state splitting”. Contrary to the classical model, there is no default state for a tissue type that is not induced. Instead, there is an active change from a previous tissue type to one of two new cell types. The change occurs as a response to which of two types of waves the individual cell actively participates in. Although there is direct correlation between the timing and spatial components of the waves for all the inductions of all presumptive tissue types that appear during axolotl gastrulation, a direct cause and effect relationship remains to be proven (Björklund and Gordon, 1994).

If the cell state splitter model is correct, any malfunction of microfilament function would cause NTDs by preventing contraction waves and thereby preventing localized differentiations in the open neural plate, which would probably alter the mechanics of neural tube closure (Brodland and Clausi, 1994). The reduced
methyllations of critical regulatory sites may be the underlying mechanism behind multifactorial NTDs. Failures of differentiation may occur because malfunctioning cytoskeletal elements cannot propagate the contraction wave. These cytoskeleton malfunctions could be due to the reduced level of posttranslational methylation in apical cytoskeleton that has been observed in rat embryos with NTDs (Moephuli, et al., 1997). In the embryo affected by an NTD, this particular failure is confined to embryonic neural tissue. The neural tissue is most affected because of its higher SAM requirement relative to other tissues. The higher requirement is due to probable PEMT activity creating new membranes during neural plate formation and closure. This failure (or combination of failures) is circumvented by folic acid supplementation because increasing folate increases SAM levels and perhaps allows proper methylation of the apical cytoskeletal elements of neural epithelium. Any variant enzyme that affects SAM availability is a candidate for the genetic predisposition component of multifactorial NTDs. Reduced folate intake or increased homocysteine become the environmental triggers for NTDs.

Mutations or syndromic conditions that result in the failure of expression of any of the proteins required for cell to cell signaling, cytoskeleton to nucleus signaling, or the subsequent differentiations would also cause NTDs. These types of NTDs would not be folate sensitive (for example, Haigo, et al., 2003). Such mutations or syndromes could account for the 30% of NTDs that are not prevented by folic acid supplementation. Thus this model interrelates the genetics, epidemiology, morphogenesis and cytoskeletal cell physiology of NTDs.

Summary

NTDs are serious, common congenital malformations that can be prevented by periconceptional folic acid supplementation. We hypothesize that folic acid provides the methyl group used for posttranslational methylation of arginine and histidine in the highly conserved regulatory domains of the cytoskeleton that is required for neural tissue differentiation (Björklund, 2005). Presumptive neural tissue has an unusually high need for folates due to the activity of phosphoethanolamine methyl transferase in producing neural tissue specific lipids at a time when the cytoskeleton is also competing for methylation. When folate supply is low and the cytoskeleton is not methylated properly, the model predicts that the result is a neural tube defect.

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