

Beyond folic acid: can optimizing maternal status of other methyl donors contribute to further reducing the risk of neural tube defects?

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1 Beyond folic acid: can optimizing maternal status of other methyl donors contribute to

2 further reducing the risk of NTD?

- 3 Helene McNulty
- 4 Author Affiliation: Nutrition Innovation Centre for Food and Health, School of Biomedical
- 5 Sciences, Ulster University, Coleraine, Northern Ireland, United Kingdom
- 6 Author's last name: McNulty
- 7 Corresponding Author: Prof Helene McNulty, Nutrition Innovation Centre for Food and
- 8 Health (NICHE), School of Biomedical Sciences, Ulster University, Cromore Road,
- 9 Coleraine, Northern Ireland BT52 1SA. Telephone: +44 2870124583 Email:
- 10 h.mcnulty@ulster.ac.uk
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In this issue of The American Journal of Clinical Nutrition, Petersen et al (1) present important 16 findings from a multicentered, population-based, case-control study of >40,000 US 17 pregnancies, suggesting that higher intakes of multiple micronutrients involved in one-carbon 18 metabolism, individually or in combination, are associated with a reduced risk of neural tube 19 defects (NTDs) in the offspring of women meeting current folic acid recommendations. NTDs 20 21 are major congenital malformations of the central nervous system occurring as a result of failure in early pregnancy of the neural tube to close properly to form the brain and spinal cord, 22 resulting in death of the fetus or newborn or lifelong disability. The most common forms of 23 NTDs are an encephaly (a brain defect) and spina bifida (a spinal cord defect), depending on the 24 portion of the neural tube that fails to close. Most children with NTDs who survive beyond birth 25 will have serious lifelong disabilities. There are profound impacts for the individual, their 26 families and society. 27

The finding of a beneficial effect of periconceptional folic acid supplementation of 28 mothers in preventing NTD in their offspring ranks as one of the most important discoveries in 29 30 human nutrition, and indeed in birth defects, research. Despite the availability of conclusive evidence for this effect for over 30 years (2, 3), the precise mechanism whereby folic acid (the 31 synthetic vitamin form) protects against NTD remains to be fully elucidated. It must however 32 involve one-carbon metabolism, a network of folate-dependent pathways requiring close 33 34 interaction with other methyl donors. Thus, the latest finding by Petersen et al, suggesting that 35 lower intakes of methyl donor nutrients apart from folate are implicated in NTD risk, is biologically plausible and consistent with certain previous reports, including an earlier study 36 by the current authors (4, 5). In particular, there is considerable evidence showing that low 37 maternal vitamin B12 status is an independent risk factor for having an NTD-affected 38 39 pregnancy (6); likewise, lower amniotic fluid B12 concentrations and lower B12 binding capacity were reported in NTD-affected pregnancies (7). 40

One-carbon metabolism involves the transfer and utilization of one-carbon units (i.e 41 methyl, formyl or formimino groups) in a network of reactions required for DNA and RNA 42 biosynthesis, serine and glycine metabolism, histidine catabolism, methionine synthesis and 43 44 methylation processes (8). For effective functioning of this network, folate needs to interact closely with vitamins B12, B6 and riboflavin. Reduced folates enter the one-carbon cycle as 45 tetrahydrofolate (THF), which then acquires a carbon unit from serine in a vitamin B6-46 dependent reaction to form 5,10 methyleneTHF. Once formed, this folate co-factor is converted 47 48 to 5 methylTHF, or serves as the one-carbon donor in the synthesis of nucleic acids, where it is

required by thymidylate synthetase in the conversion of deoyxuridine to deoxythymidine for 49 pyrimidine biosynthesis, or is converted to other folate co-factor forms required for purine 50 biosynthesis. Methylenetetrahydrofolate reductase (MTHFR) is the riboflavin-dependent 51 enzyme that catalyzes the reduction of 5,10 methyleneTHF to 5 methylTHF, the folate form 52 used by methionine synthase for the vitamin B12-dependent conversion of homocysteine to 53 methionine and the formation of THF. Methionine in turn is activated by ATP to form S-54 adenosylmethionine, the 'universal methyl donor' which donates its methyl group to more than 55 100 methyltransferases for a wide range of substrates such as DNA, hormones, proteins, 56 neurotransmitters and membrane phospholipids (8). Thus, effective folate functioning requires 57 essential metabolic interaction with vitamins B12, B6 and riboflavin, and therefore, inadequate 58 59 intake of one or more of these B vitamins, or polymorphisms in folate genes, can impair onecarbon metabolism, even if folate intake is adequate (9-13). Given that other methyl donors 60 play such critical roles in folate recycling, it is unsurprising, but not so widely reported, that 61 their status would be important in preventing the range adverse health outcomes associated with 62 impaired one-carbon metabolism including NTD. Petersen et al adds the most comprehensive 63 evidence to date that optimizing maternal status of other methyl donors could contribute to 64 further reducing the risk of NTD. 65

66 Petersen and co-authors set out to investigate whether intakes of the folate-related B vitamins B12, B6, riboflavin, and other methyl donors methionine, choline, betaine, along with 67 68 thiamine, and zinc, individually or in combination, were associated with NTD risk reduction in offspring of women meeting the folic acid recommendations. Data were drawn from the 69 National Birth Defects Prevention Study, a population-based, case-control cohort in the US, 70 during the period 1999 and 2011. Cases (n=1227) were live births, stillbirths, or terminations 71 72 affected by NTD. Controls were live births without a major birth defect (n=7095). Intakes of each micronutrient were categorized as 'higher' or 'lower' based on a combination of diet and 73 74 periconceptional vitamin supplementation. The results show although that NTD associations with each individual micronutrient were weak to modest, much greater NTD reductions were 75 observed with concurrent higher-level intakes of multiple micronutrients, with the strongest 76 association - equating to ~75% lower NTD risk - observed with concurrent consumption of 77 higher B6, B12, choline, betaine, and methionine, compared with intake of only one or no 78 methyl donors in the higher range. The authors conclude that NTD prevention, in the context 79 80 of folic acid fortification, could be augmented with intakes of methyl donors and other micronutrients involved in folate metabolism. The strengths of the Petersen study include its 81

population-based design, the relatively large number of cases, the rigor of case classification,and adjustment for sociodemographic factors.

84 Despite this latest evidence suggesting benefits of other methyl donors, readers should note that the proven protective effect of periconceptional folic acid against NTD established 85 86 over 30 years ago (2, 3) provides the evidence to support policy for women of reproductive age globally and has led to clear recommendations to take 0.4 mg/d folic acid before conceiving 87 and in early pregnancy. But translating these recommendations into effective policy and 88 practice has proved problematic over the years and, for huge proportions of women, we have 89 not reached the stage where the primary goal of optimizing maternal folate status to prevent 90 NTD has been achieved (14). On the one hand, in the US, along with 90 other countries 91 92 worldwide, mandatory folic acid fortification has proven to be highly effective in optimizing folate status and reducing NTDs. In marked contrast, effective folic acid policy to prevent NTD 93 94 has not been implemented in Ireland, the UK or other European country, and as a result, there has been no change in the incidence of NTDs over the 25-year period that the current strategy, 95 96 recommending periconceptional folic acid supplements to women, has been in place. Concerns regarding potential adverse effects of folic acid have delayed the implementation of effective 97 folic acid policy in Europe. Although the balance of available scientific evidence suggests that 98 the proven benefits of mandatory folic acid fortification would more than outweigh any risks, 99 100 folic acid is biologically highly potent, and dose remains an important consideration in any 101 emerging policy, whether folic acid fortification or supplementation (15). Notably, recent studies from North America have addressed concerns regarding high-dose folic acid usage and 102 concluded that higher-than-recommended folic acid doses are unwarranted for the prevention 103 of first occurrence of NTDs (16). The implementation of mandatory fortification, wherever it 104 105 is introduced, must therefore be accompanied by rigorous monitoring to ensure that the target folic acid levels for beneficial effects are reached, whilst avoiding any risk of overexposure at 106 107 a population level.

The Petersen et al paper provides important new information with potential impacts for policy and practice in relation to NTD prevention. It provides convincing evidence that optimizing the status of methyl donors other than folate will have additional benefits in NTD prevention. As the authors point out, a randomized trial to confirm the roles of each of these nutrients in protecting against NTD is unlikely to ever be performed. In the absence of such conclusive evidence, it is important to recognize that the findings of the current paper, together with other lines of evidence, can make a meaningful contribution to emerging policy aimed at

preventing NTD. Nonetheless, intervention with folic acid in early pregnancy remains the only 115 proven measure to reduce NTDs and must remain the driver of evidence-based policy in the 116 area globally. Thus, efforts worldwide should focus first and foremost on implementing 117 mandatory folic acid fortification in the many countries worldwide (including throughout 118 Europe) who have yet to introduce this effective measure and where current policy (educating 119 women to take folic acid supplements) has proven to be ineffective in reducing NTDs for the 120 past 25 years, and thus preventable NTDs are not being prevented. Urgent action is needed on 121 implementation of mandatory food fortification with folic acid particularly in Ireland, where 122 123 NTD rates are among the highest in the world, so that mothers and their babies can benefit.

124 Conflict of Interest Statement:

125 The author has no conflict of interest to declare

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