Docosahexaenoic acid in plasma phosphatidylcholine may be a potential marker for in vivo phosphatidylethanolamine N-methyltransferase activity in humans1–3

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ABSTRACT
Background: Choline is an essential nutrient for humans, and part of this requirement is met by endogenous synthesis catalyzed by hepatic phosphatidylethanolamine N-methyltransferase (PEMT). PEMT activity is difficult to estimate in humans because it requires a liver biopsy. Previously, we showed that mice that lack functional PEMT have dramatically reduced concentrations of docosahexaenoic acid (DHA; 22:6n–3) in plasma and of liver phosphatidylcholine (PtdCho)—a phospholipid formed by PEMT.

Objective: The objective was to evaluate plasma PtdCho-DHA concentrations as a noninvasive marker of liver PEMT activity in humans.

Design: Plasma PtdCho-DHA concentrations were measured in 72 humans before and after they consumed a low-choline diet, and correlations were analyzed in relation to estrogen status, PEMT polymorphism rs12325817, the ratio of plasma S-adenosylmethionine (AdoMet) to S-adenosylhomocysteine (AdoHcy), and dietary choline intake; all of these factors are associated with changes in liver PEMT activity. PtdCho-DHA and PEMT activity were also measured in human liver specimens.

Results: At baseline, the portion of PtdCho species containing DHA (pmol PtdCho-DHA/nmol PtdCho) was higher in premenopausal women than in men and postmenopausal women (P < 0.01). This ratio was lower in premenopausal women with the rs12325817 polymorphism in the PEMT gene (P < 0.05), and PtdCho-DHA concentration and PEMT activity were lower in human liver samples from women who were homozygous for PEMT rs12325817 (P < 0.05). The ratio of DHA-PtdCho to PtdCho in plasma was directly correlated with the ratio of AdoMet to AdoHcy (P = 0.0001). The portion of PtdCho species containing DHA in plasma was altered in subjects who consumed a low-choline diet.

Conclusion: PtdCho-DHA may be useful as a surrogate marker for in vivo hepatic PEMT activity in humans. This trial was registered at clinicaltrials.gov as NCT00065546.

INTRODUCTION
Humans develop fatty liver, liver damage, and muscle damage when they consume diets low in choline (1, 2). This nutrient is obtained in the diet from a wide variety of foods (3–5) and also can be derived from de novo biosynthesis of phosphatidylcholine (PtdCho) in liver (6). This pathway is catalyzed by phosphatidylethanolamine N-methyltransferase (PEMT), which utilizes S-adenosylmethionine (AdoMet) to sequentially methylate phosphatidylethanolamine to form PtdCho. PtdCho is the most common phospholipid in eukaryotic cellular membranes and can be hydrolyzed by phospholipase D to release free choline (6). Most PEMT activity occurs in the liver (7).

Activity of hepatic PEMT is of significant interest because it can influence the amount of choline that must be obtained from the diet. We previously reported that individuals with a particular single nucleotide polymorphism (SNP) in the PEMT gene are at greater risk of developing liver and muscle dysfunction when they consume a diet low in choline (8). Moreover, because PEMT activity is regulated by estrogen in human and mouse hepatocytes (9–12), the choline requirement is increased for postmenopausal women (because they have lower estrogen concentrations than do premenopausal women) (13). Furthermore, knockout mice with no functional PEMT activity (PEMT−/−) cannot survive without adequate choline provided in the diet (14). Many other factors also can alter PEMT activity, including ethanol (15), certain hypolipemic drugs (16, 17), consumption of a choline-methionine deficient diet (18, 19), and other SNPs in the PEMT gene (20). In addition to its influence on dietary choline requirements, perturbations of hepatic PEMT activity also alter the normal secretion of VLDLs from liver (21–23) and are associated with the development of hepatosteatosis (20, 24).

PEMT also may have an important role in fatty acid distribution to tissues. The term PtdCho represents a family of phospholipids that differ in fatty acid composition but share a common glycerophosphocholine backbone. PtdCho derived from the PEMT pathway differs from that generated by the cytidine diphosphocholine (CDP)–choline pathway. The CDP-choline pathway forms PtdCho molecules primarily containing

Reference:
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medium-chain and saturated fatty acids, whereas PtdCho formed by the PEMT pathway primarily contains long-chain polyunsaturated fatty acids (PUFAs), such as arachidonic acid (20:4n-6) and docosahexaenoic acid (DHA; 22:6n-3) (25). The fact that the phosphatidylethanolamine used by PEMT is rich in DHA, whereas PtdCho synthesized from choline is not, makes PtdCho-DHA a good index of PEMT activity. In addition, we previously showed that PEMT⁻/⁻ mice have significantly lower DHA species in plasma and liver PtdCho than do wild-type mice—an effect not seen with any other fatty acids (26). In isolated rat hepatocytes, a rapid increase in PtdCho-DHA concentrations occurred after treatment with ethanolamine for 2 h, and this increase was prevented when PEMT was inhibited by 3-deazadenosine (27), which also suggests that the PtdCho-DHA concentration is a surrogate marker for PEMT activity.

Because liver tissue is not easily obtained from human subjects, it has been difficult to assess the activity of hepatic PEMT in humans. Considering the physiologic importance of PEMT and its ability to be modulated by many factors, a marker of in vivo hepatic PEMT activity in humans would be useful. Current measures of PEMT activity are quite often performed in vitro by using isolated tissues. In humans, this would require an invasive liver biopsy and would most likely yield inaccurate results because in vitro assays cannot replicate the fluctuating substrate and inhibitor concentrations found in vivo. Therefore, this investigation evaluated the usefulness of assessing plasma PtdCho-DHA concentrations as a surrogate marker of liver PEMT activity in humans.

**SUBJECTS AND METHODS**

**Subjects and diets**

Seventy-two healthy volunteers (20 men and 52 women) ranging in age from 18 to 70 y were recruited for this study. Subject enrollment began on 14 March 2001 for the first phase of the study and on 27 June 2007 for the second phase. Of these women, 25 were postmenopausal and 27 were premenopausal. Postmenopausal status was defined as having had the last spontaneous menstrual bleeding >12 mo previously and a follicle-stimulating hormone concentration of >30 IU/L. The details of subject recruitment and the clinical protocol were approved by the Institutional Review Board at the University of North Carolina at Chapel Hill. The subjects were admitted to the University of North Carolina at Chapel Hill Clinical and Translational Research Center where they received specified diets (28), and remained under the constant supervision of study staff for the entire duration of the study.

Initially, all participants received a conventional diet of normal foods containing 550 mg choline/70 kg daily (the current Adequate Intake level; 29), 50 mg betaine/70 kg daily, and 400 dietary folate equivalents (DFE) daily (baseline). After 10 d of this initial baseline diet, the subjects entered a choline-depletion phase and were fed a low-choline diet containing <50 mg choline/70 kg daily (with 6 mg betaine/70 kg daily), as confirmed by chemical analysis of a sample of duplicate food portions (3, 30). All diets were isocaloric, with 30% of energy from fat, 0.8 g high biological value protein/kg body weight, and the remaining energy from carbohydrate. These diets also met or exceeded the Estimated Average Requirement for methionine and cysteine and the dietary reference intake for all vitamins with the exception of folate (some of the subjects received 100 µg DFE/d, and others received 400 µg DFE/d; folate intake did not influence choline-deficiency organ dysfunction; 1, 8, 31). Periodic measurements of urinary choline and betaine concentrations (30) were used to confirm compliance with the dietary restrictions (data not shown). The subjects remained on the depletion diet until they developed organ dysfunction associated with choline deficiency or for 42 d if they did not.

The subjects were deemed to have organ dysfunction associated with choline deficiency if they had an increase in serum creatine phosphokinase activity >5-fold (2), an increase in aspartate aminotransferase >1.5 fold, or an increase in liver fat (measured by magnetic resonance imaging; 1) of >28% while consuming the choline-depletion diet. Aspartate aminotransferase and creatine phosphokinase analyses were performed by the McClendon Clinical Laboratory at University of North Carolina Hospitals (Clinical Laboratory Improvement Act and College of American Pathologists accredited) by using blood drawn by venipuncture. Additional samples of blood were collected. After centrifugation, plasma was removed and stored at –80°C until the day of analysis to measure plasma AdoMet, S-adenosylhomocysteine (AdoHcy), PtdCho, and PtdCho-DHA concentrations.

**Liver tissue procurement**

Human liver samples from premenopausal women were obtained from the Liver Procurement and Distribution System (University of Minnesota, Minneapolis MN; funded by NIH contract N01 DK92310). Of the 8 donors, 7 had fatty liver (2 of which were cirrhotic as well). Synopses of the tissue donors’ medical histories, including the pathologists’ impression diagnoses concerning liver fat content, were obtained. The specimens were snap-frozen after removal from the organ donors, delivered on dry ice, and stored at −80°C until analyzed.

**Analysis of plasma PtdCho fatty acids**

Lipids were extracted from duplicate plasma samples, and the livers were pulverized in liquid nitrogen by using the method of Bligh and Dyer (32) in the presence of an internal standard (1,2-diheptadecanoyl PtdCho; Avanti Polar Lipids, Alabaster, AL) and 0.02% butylated hydroxytoluene. Lipid classes were separated by thin-layer chromatography (CHCl₃; Chromatograph; Agilent Technologies, Santa Clara, CA), with fatty acid methyl esters (FAMEs) being separated by gas chromatography by using a BPX-70 capillary column (SGE, Austin, TX) and a split injection system (Agilent Technologies, Santa Clara, CA), with helium as the carrier gas. The injector temperature was 240°C, and the flame ionization detector 280°C. Data were analyzed by using ChemStation Firmware A.01.09 (Agilent). DHA was identified and quantified against authentic standards (NuCheck Prep, Elysian, MN). Data were expressed as pmol PtdCho-DHA/nmol PtdCho.
Determination of plasma PtdCho, AdoMet, and AdoHcy

PtdCho was extracted from plasma by the method of Bligh and Dyer (32). Aqueous and organic compounds were separated, analyzed, and quantified directly by using liquid chromatography/electrospray ionization-isotope dilution mass spectrometry after the addition of internal standards labeled with stable isotopes to correct for recovery (30). AdoMet and AdoHcy were measured in a subset of plasma samples from subjects enrolled in 2001–2004 by using an HPLC method (34). Briefly, 500 μL plasma (in duplicate) was deproteinized with trichloroacetic acid, and AdoMet and AdoHcy were separated by HPLC with the use of a C8 column. Fractions were collected and derivatized by using naphthalenedialdehyde and cyanide. The resulting fluorescent isoindoles were separated by HPLC with the use of a C18 column and a fluorescence detector.

PEMT activity assay

Liver homogenates were assayed for PEMT activity by using a modified method of Ridgway and Vance (35). Briefly, activity was assessed by using 50 μg protein in 125 mmol Tris-HCl/L (pH 9.2) and 5 mmol DTT buffer/L (Sigma, St Louis, MO) in the presence of 200 μmol S-adenosyl-L-methionine/L containing 0.5 μCi S-adenosyl-L-[methyl-3H] methionine (55.70 Ci/mmol; GE Health Care, Piscataway, NJ) and 0.4 mmol exogenous phosphatidyl-dimethylethanolamine/L (P2; Avanti Polar Lipids, Alabaster, AL). The reaction was carried out for 30 min at 37°C and stopped by adding CHCl3:MeOH:1 N HCl (100:50:1, vol:vol:vol). An aliquot of the chloroform phase was applied to a silica gel thin-layer chromatography plate [Si250-PA (19C)-Silica Gel, Baker Inc, Phillipsburg, NJ] and developed in CHCl3:MeOH:acetic acid:H2O (50:30:5:2, vol:vol:vol:vol). Disintegrations per minute from [3H]-PtdCho were determined in bands that comigrated with a standard and expressed as the Pearson correlation coefficient. Data are reported as means ± SEs, and all results were considered significant at P < 0.05.

RESULTS

At baseline, plasma PtdCho-DHA/PtdCho ratios in plasma were greater in premenopausal women than in men or postmenopausal women (Figure 1; P < 0.01). This pattern was maintained when subjects were placed on a low-choline diet (data not shown). Because PtdCho concentrations were expected to decrease with choline deficiency, we expressed the data as a concentration ratio so that PtdCho-DHA could be represented by the actual amount of PtdCho present. However, we found that expressing the PtdCho-DHA data as nmol/mL plasma gave similar results (see Supplemental Figure 1 under “Supplemental data” in the online issue). The PEMT gene polymorphism rs12325817 was previously reported to greatly increase the likelihood of developing organ dysfunction when subjects with the variant allele consumed a low-choline diet (8), and we suggested that this indicated diminished capacity to form choline via PEMT activity. In the current study, premenopausal women homozygous for this SNP had a lower plasma PtdCho-DHA/PtdCho ratio in plasma than did premenopausal women without the SNP at baseline (Figure 1; P < 0.05 by t test) and after the low-choline diet (data not shown). The premenopausal women with one allele of the SNP had intermediate values (47 ± 6 nmol PtdCho-DHA/nmol PtdCho; n = 14), which were not statistically different from those with 2 alleles of the SNP. The plasma PtdCho-DHA/PtdCho ratio did not differ significantly between postmenopausal women and men by genotype, including in those with one allele of the SNP.

Statistical analysis

The effect of sex and menopausal status on the plasma PtdCho-DHA/PtdCho ratio was determined by mixed-model analysis of variance by using JMP software (V2; SAS Institute, Cary, NC), followed by pairwise comparisons of individuals with and without the PEMT rs12325817 genotype and by dietary treatment after grouping by sex and menopausal status. For the human liver data, statistical differences in the plasma PtdCho-DHA/PtdCho ratio and PEMT activity were determined by using Student’s t test after grouping by genotype. The relation of plasma AdoMet and AdoHcy concentrations to the plasma PtdCho-DHA/PtdCho ratio was determined by linear regression and expressed as the Pearson correlation coefficient. Data are reported as means ± SEs, and all results were considered significant at P < 0.05.

Genotyping

SNP analysis was performed as previously described (8, 13). Briefly, genomic DNA was extracted from liver tissues by using TriZol (Invitrogen, Carlsbad, CA) and from peripheral lymphocytes isolated from blood by using PureGene (Gentra Systems, Minneapolis, MN) according to the manufacturer’s instructions. DNA sequencing was performed on double-stranded DNA templates obtained from genomic DNA by polymerase chain reaction (PCR) amplification. Sequencing reactions were performed by the University of North Carolina at Chapel Hill Genome Analysis Facility by using a capillary sequencing machine (model 3100; Applied Biosystems, Foster City, CA). Sequence results were interpreted by using the program Sequencher (Gene Code Corp, Ann Arbor, MI). Successful amplification of the 1896-base pair DNA fragment of the PEMT gene was performed by using Takara Ex Taq polymerase (Fisher Scientific, Fair Lawn, NJ) with forward and reverse primers: 5’GAGCACGTGAGCTGT-CAGTGCCCTTTTG3’ and 5’CACAACCCTCTTATACACAC-AGGTCC3’, respectively. A 3-step PCR was performed on an Applied Biosystems 2720 Thermal Cycler under the following conditions: 96°C for 2 min, 30 cycles (94°C for 30 s, 60°C for 1 min, and 72°C for 2 min), extend 72°C for 7 min, and soak at 4°C. For the sequence determination of PEMT rs12325817, we used an additional primer, 5’TAGATTTGTCATGGGAGGCTT3’, to verify the sequence in a region containing Alu repeats and a poly-A tail. We also designed forward primers specific to the rs12325817 allele using GeneFisher (http://bibiserv.techfak.uni-bielefeld.de/genefisher) so that the SNP would be located at the 3’-end of the priming sequence, which allows for specific PCR products to be synthesized only if the primer is 100% complementary to its template DNA. We purchased them from Qiagen (Huntsville, AL). The PCR reactions were optimized for each pair of primers, and the products were visualized on a 1.5% agarose gel to determine the genotype.

Statistical analysis

The effect of sex and menopausal status on the plasma PtdCho-DHA/PtdCho ratio was determined by mixed-model analysis of variance by using JMP software (V2; SAS Institute, Cary, NC), followed by pairwise comparisons of individuals with and without the PEMT rs12325817 genotype and by dietary treatment after grouping by sex and menopausal status. For the human liver data, statistical differences in the plasma PtdCho-DHA/PtdCho ratio and PEMT activity were determined by using Student’s t test after grouping by genotype. The relation of plasma AdoMet and AdoHcy concentrations to the plasma PtdCho-DHA/PtdCho ratio was determined by linear regression and expressed as the Pearson correlation coefficient. Data are reported as means ± SEs, and all results were considered significant at P < 0.05.
In human livers (n = 8) and plasma (n = 72), \( \approx 3\% \) of PtdCho was present as the DHA species (liver: 2.7 \( \pm \) 0.2 g/100 g fatty acids; plasma: 3.0 \( \pm \) 0.1 g/100 g fatty acids). Because cirrhotic livers have fewer hepatocytes and more fibroblasts, we excluded such livers (n = 2) from our analyses. It is interesting to note that the cirrhotic livers (both GC genotype) had a lower PtdCho-DHA content (1.6 \( \pm \) 0.1 g/100 g fatty acids) than did the other GG/GC samples. Samples from premenopausal women homozygous for the variant allele (CC genotype) had significantly less PtdCho-DHA than did samples from women with the other genotypes (Figure 2A; \( P = 0.01 \)). We measured PEMT activity in the human livers to determine whether there was a correlation with PtdCho-DHA concentrations. We found that women homozygous for the SNP had less PEMT activity than did the other women (Figure 2B; \( P = 0.008 \)). There was an insufficient sample size to analyze the GG and GC genotype groups separately.

We previously reported that some subjects, when placed on a low-choline diet, developed signs of organ dysfunction, whereas others did not (1). Plasma PtdCho concentration decreased in premenopausal women and men fed a low-choline diet (Table 1; \( P < 0.05 \) by paired \( t \) test). However, only the men who developed organ dysfunction (n = 14) had significantly lower plasma PtdCho concentrations (1615 \( \pm \) 70 nmol/mL) than their baseline values (1950 \( \pm \) 68 nmol/mL; \( P = 0.0003 \) by paired \( t \) test). The postmenopausal group did not differ regardless of diet or depletion status.

A low-choline diet lowered plasma PtdCho-DHA/PtdCho ratios in premenopausal women (Figure 3). Of these women, those who developed organ dysfunction with the low-choline diet had a 24% decrease in the plasma PtdCho-DHA/total PtdCho ratio from their basal concentration (-14 pmol DHA/nmol PtdCho) compared with 3% (-2 pmol DHA/nmol PtdCho) in those who did not develop organ dysfunction with the deficient diet (Figure 3; \( P = 0.023 \), \( t \) test). Men and postmenopausal women did not experience a change in the plasma PtdCho-DHA/total PtdCho ratio when they consumed the deficient diet.

AdoMet is a substrate for PEMT, and AdoHcy (a product of PEMT) inhibits the enzyme. While consuming the low-choline diet, subjects with the lowest AdoMet/AdoHcy ratio had the lowest plasma PtdCho-DHA/PtdCho as determined by linear regression (Figure 4; \( r = 0.716 \), \( P = 0.0011 \)).

**DISCUSSION**

We found that plasma PtdCho-DHA/PtdCho ratios in humans varied in a manner that was consistent with this measure being a good surrogate marker for in vivo PEMT activity in liver. PEMT activity is induced by estrogen (11, 12) in premenopausal women, and it is expected to be lower in men and postmenopausal women (who have low estrogen concentrations). We observed that premenopausal women had the highest plasma PtdCho-DHA/PtdCho ratios compared with men and postmenopausal women.
**TABLE 1**

Plasma phosphatidylcholine concentrations in subjects after consumption of a low-choline diet

<table>
<thead>
<tr>
<th>Group</th>
<th>Baseline diet</th>
<th>Low-choline diet</th>
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<tr>
<td></td>
<td>nmol/mL</td>
<td>nmol/mL</td>
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<tr>
<td>Premenopausal women (n = 27)</td>
<td>1742 ± 47</td>
<td>1629 ± 52&lt;sup&gt;1&lt;/sup&gt;</td>
</tr>
<tr>
<td>Men (n = 20)</td>
<td>1842 ± 63</td>
<td>1574 ± 58&lt;sup&gt;2&lt;/sup&gt;</td>
</tr>
<tr>
<td>Postmenopausal women (n = 25)</td>
<td>1997 ± 82</td>
<td>1906 ± 69</td>
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<sup>1</sup> All values are means ± SEs. Healthy volunteers consumed a standardized diet that delivered 550 mg choline/70 kg body weight daily for 10 d (baseline), followed by a diet containing <50 mg choline/70 kg body weight daily (low choline) until they developed liver or muscle dysfunction (determined as described in Subjects and Methods) or for 42 d. Blood was collected at the end of each diet phase, and plasma phosphatidylycerol concentrations were measured by mass spectrometry as described in Subjects and Methods.

<sup>2</sup> Significantly different from the baseline diet, P < 0.05 (paired t test).

PEMT uses AdoMet as a methyl donor and is maximally active at ~200 μmol AdoMet/L. PEMT is inhibited by AdoHcy (36, 37); thus, PEMT activity should be proportional to the AdoMet/AdoHcy ratio. We found that the AdoMet/AdoHcy ratio measured in plasma was correlated with PEMT activity in vivo. The correlation was assessed by using linear regression analysis (P = 0.0001, r = 0.716). n = 41.

**FIGURE 3.** Mean (±SE) ratios of phosphatidylcholine docosahexaenoic acid to phosphatidylcholine (PtdCho-DHA/PtdCho) concentrations in plasma in healthy premenopausal women (F), men (M), and postmenopausal women (P) after consumption of a standardized diet that delivered 550 mg choline/70 kg body weight daily for 10 d and after consumption of a diet containing <50 mg choline/70 kg body weight daily for ≤42 d or until they developed liver or muscle dysfunction (D) as described in Subjects and Methods. Subjects who developed signs of organ dysfunction while consuming the low-choline diet had decreased plasma PtdCho-DHA/PtdCho ratios. Some subjects did not develop organ dysfunction within 42 d of consuming the low-choline diet (ND). Blood was collected at the end of each diet phase, and the portion of PtdCho species containing DHA was measured in plasma by using gas chromatography as described in Subjects and Methods. Plasma AdoMet and AdoHcy concentrations were measured at the end of the low-choline phase of the study as described in Subjects and Methods. The correlation was assessed by using linear regression analysis (P = 0.0001, r = 0.716). n = 41.

**FIGURE 4.** Ratios of phosphatidylcholine docosahexaenoic acid to phosphatidylcholine (PtdCho-DHA/PtdCho) concentrations in plasma correlated with ratios of plasma S-adenosylmethionine (AdoMet) to S-adenosylhomocysteine (AdoHcy). Healthy volunteers consumed a standardized diet that delivered 550 mg choline/70 kg body weight daily for 10 d and then consumed a diet containing <50 mg choline/70 kg body weight daily for ≤42 d or until they developed liver or muscle dysfunction as described in Subjects and Methods. At the end of this period, blood was collected and the portion of PtdCho species containing DHA was measured in plasma by using gas chromatography as described in Subjects and Methods. Plasma AdoMet and AdoHcy concentrations were measured at the end of the low-choline phase of the study as described in Subjects and Methods. The correlation was assessed by using linear regression analysis (P = 0.0001, r = 0.716). n = 41.
Nutritional and the Egg Nutrition Research Center for studies other than
analyses and data interpretation, provided major input in the writing of the
and statistical analyses and data interpretation, provided major input in the writing of the
and was responsible for the conceptualization, implementation, and
design of the human study. None of the authors had a financial conflict of
and the Egg Nutrition Research Center for studies other than
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