Higher efficacy of dietary DHA provided as a phospholipid than as a triglyceride for brain DHA accretion in neonatal piglets

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Running Title: Higher brain DHA accretion with PL-DHA

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Abstract

Long chain polyunsaturated fatty acids occur in foods primarily in the natural lipid classes triacylglycerols (TAG) or phospholipids (PL). We studied the relative efficacy of the neural omega-3 docosahexaenoic acid (DHA) provided in formula to the growing piglet as a dose of 13C-DHA bound to either TAG or phosphatidylcholine (PC). Piglets were assigned to identical formula-based diets from early life and provided with TAG-13C-DHA or PC-13C-DHA orally at 16 days. Days later, piglet organs were analyzed for 13C-DHA and other fatty acid metabolites. PC-13C-DHA was 1.9-fold more efficacious for brain gray matter DHA accretion than TAG-13C-DHA, and was similarly more efficacious in gray matter synaptosomes, retina, liver, and RBC. Liver labeling was greatest implying initial processing in that organ followed by export to other organs, and suggesting that transfer from gut to bloodstream to liver in part drove the difference in relative efficacy for tissue accretion. Apparent retroconversion to 22:5n-3 was more than double for PC-13C-DHA and was more prominent in neural tissue than in liver or RBC. These data directly support greater efficacy for PC as a carrier for LCPUFA compared to TAG, consistent with previous studies of arachidonic acid and DHA measured in other species.

Keywords. docosahexaenoic acid, triacylglycerol, phosphatidylcholine, piglets, brain, nutrition
Docosahexaenoic acid (DHA) is the most abundant omega-3 fatty acid in the mammalian brain (1), accounting for 8-14% of fatty acids perinatally in primates (2) and humans (3, 4). In humans, DHA accumulates at an accelerating rate from mid gestation reaching an inflection point during the first months after birth (5) and continuing well past the plateau of brain weight to reach a plateau around 18 years of age and remain stable to the end of life (6).

DHA can be synthesized de novo by term and preterm human infants from precursor omega-3 fatty acids, α-linolenic acid (ALA) or eicosapentaenoic acid (EPA) (7-9) but the process is generally considered to be inefficient, also challenged by current dietary intakes of omega-6 linoleic acid at relatively high levels (10, 11). Direct measurement of percent incorporation of DHA into the primate brain from precursor ALA indicate that it is 7-fold less efficient than for preformed DHA. This finding is in contrast to that for arachidonic acid (ARA), present in the brain at similar/comparable levels but not strongly affected by dietary preformed ARA (12). Numerous human studies have shown efficacy of dietary preformed DHA to support brain and visual development (13), and follow-up studies suggest that preformed DHA intake in infancy provides long term benefits on specific tests of higher cognitive function, although it is widely agreed that no effects can be seen using the Bayley scales, a standard test for normalcy (14).

While all preformed DHA is structurally identical, in human foods and food supplements DHA is normally esterified to one of three lipid classes as molecular
carriers, triacylglycerol (TAG), phospholipid (PL), or ethyl ester. In primate milk, including human breastmilk, DHA content is higher in PL than in TAG, but the bulk of DHA is still delivered via TAG because of the overwhelming prevalence of TAG over PL (15). TAG-DHA is a common molecular form of DHA used as an ingredient in infant formula. Ongoing development of alternative sources of DHA has identified components potentially suitable for use in infant formulas that use PL-DHA. PL-DHA and -ARA supplemented infant formula has been studied since the 1990s (16, 17), and infant formula products were also widely available on the European market before this market expanded and changed for single-cell and fish oil based sources of TAG-bound ARA and DHA.

Once hydrolyzed from its esterified molecular carrier, DHA and ARA may be handled identically by normal metabolic processes. In vivo and in vitro studies show pancreatic lipase hydrolyses TAG in the sn-1 and sn-3 positions consistent with the survival of the predominantly saturated sn-2 fatty acid of breastmilk post-absorption (18, 19), while intestinal phospholipases in the rat hydrolyze the sn-2 position leaving the predominantly saturated sn-1 position intact (20). Positional specificity of intestinal glycerolipid hydrolysis insures that the post-absorptive reassembled TAG and PL will have a non-randomized distribution of fatty acids, though sn-1/3 fatty acids hydrolyzed by intestinal lipases and phospholipases in the intestine are re-esterified in a random fashion onto TAG and PL post-absorption.
Positional specificity may be metabolically relevant for subsequent metabolism as fatty acids are incorporated into membranes, oxidized, or excreted on the skin. Previously we showed by $^{13}$C labeling that ARA-PL, specifically as phosphatidylcholine (ARA-PC), when added to artificial formula is about twice as efficacious as a substrate for supplying ARA to the brain as ARA-TAG in neonatal baboons (21). We report here a similar study in neonatal piglets using $^{13}$C-DHA, specifically investigating efficacy for supplying the gray matter of the cerebral cortex with DHA.
MATERIALS AND METHODS

Chemical

Uniformly labeled DHA (perlabeled, [U-\(^{13}\)C]-DHA) was prepared in our labs using published methods (22) and esterified into lipid classes by Avanti Polar Lipids (Alabaster, AL, US). The TAG tracer (TAG-\(^{13}\)C-DHA) had \(^{13}\)C-DHA in the sn-2 position with unlabeled 16:0 in the sn-1 and sn-3 positions. The PL tracer used was specifically phosphatidylcholine (PC) with \(^{13}\)C-DHA in the sn-2 position and unlabeled 16:0 in the sn-1 position (PC-\(^{13}\)C-DHA). Solvents for tissue lipid extraction were HPLC grade from Sigma-Aldrich (St. Louis, MO, US) or Burdick & Jackson (Muskegon, MI, US).

Animals and diet

The overall study and all procedures involving live piglets were approved by the Institutional Animal Care and Use Committee (IACUC) at Cornell University. Sows (Yorkshire and Landrace cross) were bred with Hampshire boars. After birth, piglets were left to nurse with the sow for at least 48 hours. Twenty piglets, 2 to 4 days of age, were chosen for the study based on the following criteria: weight (about 2.0 kg), gender (10 males and 10 females), health (active and apparently normal, and not the smallest animal in the litter) and sow. Piglets were given a unique number/color combination identifier and then were transported to Large Animal Research and Teaching Unit (LARTU), Cornell University.

At the study site, piglets were housed in individual stainless steel cages and were
placed on a single, fully balanced, commercially available young farm animal milk replacer formula immediately upon arrival.

The fatty acid profile of the milk replacer (piglet milk formula) is presented in Table 1. DHA is present in low levels in form of TAG (0.02 %, w/w) and PL (0.11 %, w/w). TAG, PL, and related DHA measurements are presented in Table 2, when appropriate tabulated for a 500 mL feeding used with the dose. As with human milk and human infant formulas, total energy from fat was about 47% and mostly in the form of TAG (94%). Despite the low absolute DHA concentration in TAG, the predominant mass of TAG over PL provided 73% of total DHA compared to 27% from PL, in line with human milk and other mammalian milks.

Fresh formula was provided every 6 hours for the first week and 8 hours thereafter in individual troughs as in our previous studies (23, 24). All piglets had free access to fresh water. Formula consumption for each group was recorded daily for the first week to insure the piglets thrived, and monitored every third day thereafter. Piglet body weight was recorded every other day in the first week and every third day thereafter. Home cage enrichment, included bedding, contact with other piglets, positive human contact and rubber toys.

Dose and sampling

On day 16 of life, 16 robust piglets out of a total set of 20 piglets were assigned to two
dosing groups, PC and TAG, balanced with respect to gender, weight, and age but otherwise distributed randomly. Group PC was orally dosed with PC-$^{13}$C-DHA as described below. Group TAG was dosed with TAG-$^{13}$C-DHA. The four smallest piglets were used as natural abundance isotopic controls and did not receive a dose but were otherwise treated identically to the other animals.

Doses were prepared by dissolving CHCl$_3$ solutions of PC-$^{13}$C-DHA or TAG-$^{13}$C-DHA in about 1 mL of refined olive oil and driving off the CHCl$_3$ by gentle heating under flowing dry N$_2$ over 4 hours. The resulting olive oils were carefully placed on the surface of a few mL of reconstituted piglet formula and sonicated until homogeneous. The resulting labeled formula was split gravimetrically into 1 to 2 mL aliquots and administered orally to piglets via syringe. Labeled formula was withdrawn by syringe from the vials which were subsequently washed several times with unlabeled formula which was then also administered to the piglets at the beginning of a normal feeding time. Piglets were therefore hungry but could be considered as fasted insofar as their normal meal interval was concerned. Care was taken to insure all of the labeled formula and the washes, consisting of a total of up to 10 mL, were swallowed by the piglets. Piglets were then provided with their normal meal of 500 mL milk replacer which they routinely consume completely. Doses to piglets corresponded to about 20 mg $^{13}$C-DHA delivered in PC and 86 mg of $^{13}$C-DHA delivered in TAG, and the difference was only due to the amount of each labeled tracer that was available.
Six days after the receiving the labeled DHA dose and in total 20 days of formula and 22-24 days of life, all piglets were euthanized by exsanguination under anesthesia on a single day. Blood for fatty acid analysis was collected in tubes with EDTA and spun to prepare red blood cells. The surface few mm of the gray matter on the cerebral cortex superior surface was rapidly collected at necropsy. A piece approximately 5 × 10 × 3 mm was immediately used for preparation of synaptosomes, and the remaining gray matter was snap frozen in liquid nitrogen for fatty acid analysis. Retina, heart, liver, biceps and femoris muscle were also harvested, and all tissues were snap frozen and stored at -80°C until prepared for analysis.

Synaptosomes were prepared using Syn-PER Synaptic Protein Extraction Reagent (Thermo Scientific, Waltham, MA, US) according to manufacturer’s instructions. Briefly, 1 ml of Syn-PER Reagent was added to ~100 mg brain tissue sample. Samples were homogenized and the homogenate centrifuged at 1,200 g for 10 mins. The supernatant was collected and further centrifuged at 15,000g for 20 mins. Synaptosomes were recovered from the pellet and lipids were immediately extracted as described below.

Lipid extraction and analysis

Total lipids were extracted from samples of brain gray matter, the synaptosome pellet, liver, heart, and retina. Samples were simultaneously digested and fatty acid methyl esters (FAME) prepared using a one step method as described in detail previously.
(25). For plasma and erythrocytes, the Bligh and Dyer method (26) was employed to extract total lipids and FAME were prepared using 14% BF$_3$ in methanol. A known quantity of freshly prepared heptadecanoic acid in chloroform (99% pure, Sigma Chemical) was added as an internal standard to tissue samples just before extraction. FAME were dissolved in heptane and stored at -20°C until analysis.

FAME were analyzed using a Hewlett Packard 5890 series II GC-FID with a BPX 70 column (60 m; 0.32 mm inner diameter; 0.25 µm film; Hewlett Packard, Palo Alto, CA, U.S.A.) and H$_2$ as carrier gas. Quantitative profiles were calculated using the internal standard and an equal weight FAME mixture to derive response factors for each fatty acid. GC-FID conditions and calibration details have been reported (21).

$^{13}$C-DHA analysis

Tracer analysis for $^{13}$C-DHA is performed on FAME mixtures using similar GC column conditions as for the quantitative analysis, as has been described in detail previously (27). Instrumentation for tracer analysis is an Agilent 6890 gas chromatograph coupled to a combustion furnace interface, and to a Thermo Scientific 253 isotope ratio mass spectrometer (IRMS). FAME eluting from the GC are combusted to CO$_2$, dried, and admitted to the IRMS. Data processing is as described previously (21). Isotope ratios in the conventional high precision notation, δ$^{13}$C, defined previously (27), is converted to fraction of $^{13}$C. For each fatty acid, the mean isotope ratio of the control group was subtracted from the isotope ratio of the means for the enriched groups to
yield an atom fraction enrichment, which was subsequently converted to %Dose, which reflects the appearance of tracer in the specific pool. The primary outcome is a relative comparison of the %Dose appearing in the brain gray matter for TAG and PL, respectively. Total %Dose found in liver and retina was calculated directly from their respective weights. The total labeled DHA in cerebral gray matter and RBC was estimated based on brain weight, and using the relative amount of gray matter in the brain as 60%, estimated from human imaging data (28). For RBC, the blood volume was estimated as 8.5% of body weight (29) and hematocrit was about 35%. For gray matter synaptosomes, we did not attempt to estimate the total amount and normalized the %Dose to the highest value found. In all cases, estimated masses apply to both experimental groups and cancel in the primary and secondary outcome calculations, and thus do not affect the final results. Detailed calculations have been previously presented (21).

Statistics.

Primary Outcome, relative DHA %Doses from PC and TAG. The primary outcome is the relative %Dose of $^{13}$C-DHA found in the gray matter of the cerebral cortex in the PC-$^{13}$C-DHA vs the TAG-$^{13}$C-DHA dosed groups. The %Dose in the two dosing groups were tested for equivalence by one way analysis of variance with $p<0.05$ considered significant.

Secondary outcomes. Total unlabeled fatty acids in the various pools were compared
in a pairwise manner in the two dosing groups and were not significantly different, and were therefore pooled. Since these two groups were fed the same formulas and treated identically except for a few mg doses, no differences were expected based on treatment.

Relative $^{13}$C-DHA %Doses for synaptosomes, retina, liver, and RBC were compared for similarity to the primary outcome. Relative meal wise amounts of total DHA delivered in TAG and PL were calculated from the determination of relative amounts of unlabeled DHA in formula in a 500 ml meal.
Results

Piglets in both dosing groups grew at comparable rates and attained non-significantly different final weights of 9.4±0.3 kg and 9.1±0.4 kg for the PC and TAG dosed groups, respectively. The isotopic control groups were purposely chosen as the smallest animals but grew in parallel to the other groups, starting at about 6% lower body weight and finishing at about 12% lower weight. These animals were expected to yield accurate estimates of baseline isotope ratios, which were applied identically to both experimental groups.

The fatty acid profile of neural tissue is presented in Table 3. Retina was richest in DHA (22:6) at 12%, followed by gray matter (8.3%) and white matter (5.2%). Docosapentaenoic acid (22:5n-6), the DHA analogue among omega-6 fatty acids, mean value was 9.7% in gray matter, and thus slightly greater than DHA. Domestic piglets have long been considered to require small amounts of linoleic acid and that n-3 incidental to practical diets is sufficient to support rapid growth (30). These levels are apparently insufficient to enable DHA levels similar to wild animals (1) though they are apparently developing normally.

Results for the primary outcome in gray matter and gray matter synaptosomes are presented in Figure 1. $^{13}$C label was detected only for DHA and 22:5n-3 (DPAn-3) among all fatty acids. The %Dose detected in cerebral cortex gray matter in the $^{13}$C-DHA-PC dosed animals was about 0.4%, greater than that for the TAG dose.
$^{13}$C-DHA from the TAG dose was about half this value, thus dietary PC was 1.9-fold more efficacious to supply DHA to the developing piglet brain compared to TAG. Results in gray matter synaptosomes, also shown in Figure 1, were consistent with the relative values. The ratio of relative accretion of labeled DHA in the synaptosomes was 1.7-fold greater for PC than for TAG.

The retroconversion product DPAn-3 was labeled below 0.05% Dose for the PC dose group and less than half of that for the TAG dose group. The relative efficacy of PC over TAG for the retroconversion of DHA to DPAn-3 was about 2.8. For gray matter synaptosomes, the superiority of PC over TAG was the same with a relative efficacy of 2.8. Gray matter and synaptosome $^{13}$C-DHA was approximately 10-fold greater than $^{13}$C-DPA.

Figure 2 presents the results for liver and RBC. Liver retained 6.8% of the PC $^{13}$C-dose compared to 3.5% of the TAG dose at 6 days post-dose. The relative efficacy was 1.9, similar to gray matter and synaptosomes. The ratio of $^{13}$C-DHA to $^{13}$C-DPA was about 50, substantially greater than found for gray matter. RBC relative labeling paralleled liver levels, with PC-derived $^{13}$C-DHA 1.6-fold that of TAG-derived $^{13}$C-DHA, and a similar 50-fold dominance of $^{13}$C-DHA over $^{13}$C-DPA.

Figure 3 presents the results for retina tissue. Relative efficacy of PC over TAG was 2.2-fold for DHA, and about 2.4-fold for DPAn-3, all similar and consistent with results
in other tissues examined. Consistent with the brain gray matter but in contrast to the liver, $^{13}$C-DHA was 6.3-fold greater in the retina than the retroconversion product $^{13}$C-DPA-3, again suggesting that retroconversion took place outside the liver and possibly in the retina. As seen before, TAG predominated as the source of retina DHA.
Discussion

Our study investigated the relative efficacy of dietary DHA carried by PC compared to DHA carried by TAG. The results show that dietary DHA carried as PC is more efficiently used for brain gray matter DHA accretion than when carried as TAG. The data in isolated white matter synaptosomes are consistent with those for whole gray matter, and match results obtained in other tissues. Moreover, the data is quantitatively consistent with our previous results using a similar approach to study ARA membrane incorporation in neonatal baboons (21). The similarity of these findings with respect to specific LCPUFA (ARA or DHA), various species, and different laboratories, supports the conclusion that LCPUFA carried as PL are utilized differently and more efficiently deliver LC-PUFA as source of membrane components than when provided as TAG.

Despite the well recognized importance of brain DHA accretion, transport forms of DHA into the brain are not fully understood. Deletion of CD36, an unesterified fatty acid binding protein, in mice does not reduce brain DHA(31). Similar results were obtained for VLDL receptor null mice (32). Consistent with the present data, albumin-bound sn-1-lyso-2-DHA-PC is more efficiently incorporated into brain PL in the rat compared to unesterified DHA (33, 34). In our study, 1-lyso-2-\(^{13}\text{C}\)-DHA-PC can be synthesized by a phospholipase A1 action on \(^{13}\text{C}\)-DHA-PC, recently proposed to be mediated by an endothelial lipase (35), analogous to results suggesting extensive phospholipase A1 activity mediated by hepatic lipase (36). This pathway would not be
available to DHA-TAG, and could at least in part mediate some of our whole body results. A very recent report shows that mice expressing human apoE4 take up less unesterified DHA and have lower brain DHA compared to mice expressing apoE2 or apoE3, suggesting a mechanism involving free DHA as well (37). A DHA-specific lysophosphatidic acid acyl transferase that may mediate specific transfer of unesterified DHA into phospholipids was also recently reported (38).

The majority of dietary DHA was carried as TAG because of its greater proportion of DHA compared to PL in the formula (piglet milk replacer) (Figure 1C,D, 2C,D, 3B). Qualitatively, this is comparable with the composition of human/mammalian milks of other species. However, the amount of DHA in our piglet formula was very low, probably only being a minor component of bovine milk protein as base of the formula. Variation in human milk DHA is over at least an order of magnitude, from 0.1% to over 1% of fatty acids (39) in apparently well nourished mothers and is strongly related to the mother’s DHA intake (13). The lowest DHA content of human milk of which we are aware has been reported in resource-limited northern Sudanese mothers with carbohydrate rich diets (0.06%, w/w) (40). Because of the low dietary DHA, our study is more similar to previous studies conducted with LCPUFA-free control formulas reported until the early years of the present century compared to recent studies comparing formulas with various DHA. However, the relative advantage of about 1.9-fold for PC-DHA compared to TAG-DHA applies closely to current formulas because it is relatively unaffected by overall formula DHA levels and reflects the
handling of a single physiological dose of DHA; it should therefore apply also to
greater formula DHA levels as have been investigated in primates and humans (12, 41).

Liver $^{13}$C-DHA was about 20-fold greater overall than the $^{13}$C-DHA in the gray matter,
in contrast to the results reported previously for the baboon neonate (21). One
explanation for this effect is likely to be a species difference in the relative mass of the
organs, which in the baboon is near 10% of body weight, while in our piglets was only
0.57±0.04% of body weight. In absolute terms, relative sizes of the different organs
are relevant: the piglet gray matter represented about 29 g while the liver was about
321 g. Even with these considerations, the relative efficacy of $^{13}$C-DHA from PC and
TAG was similar to that reported for 20:4n-6, arachidonic acid, in the baboon (21).

The retroconversion product DPAn-3, though also almost 3-fold greater in the liver for
PC compared to TAG, was a smaller proportion of the liver $^{13}$C-DHA than found in gray
matter. In gray matter, $^{13}$C-DHA was about 8-fold greater than $^{13}$C-DPA while in liver
the $^{13}$C-DHA was 40-fold greater. If DPAn-3 had been synthesized in the liver and
transported to the brain, similar ratios would be expected. These observations
suggest that retro-conversion takes place in organs other than the liver and may well
be more active in the brain than in liver.

The piglet has long been used as a model for infant nutrition because of its similar
metabolism to humans, and in numerous studies it has been used to investigate the aspects of LCPUFA delivery from TAG vs PL (e.g. (42, 43)). Piglet growth is several-fold more rapid than for human infants, thus also reflecting several fold greater intake and exposure to test substances than human infants (44). Others have shown that piglet lipoprotein characteristics depend on the source of DHA, whether TAG oil or egg PL (45, 46), a mechanism that may be operating in our piglets to influence the relative amounts captured by the liver. Notably, the major difference in $^{13}$C-DHA among the measured pools was in total labeling, with liver enrichment much greater than RBC or neural tissue (%Dose in all figures). For instance, liver PC-$^{13}$C-DHA was about 7%Dose while cerebral gray matter %Dose was about 0.4%. These results are consistent with the liver’s central role as a processor/repackager of PUFA including DHA and that the form of DHA emerging from the lymph was of key importance. Within an organ, the relative efficacy is expected to accurately reflect human metabolism.

PL as a carrier of LCPUFA became of interest for infant nutrition in the 1990s with the introduction of sources that provided LCPUFA in PL form, and the recognition that, although PL is richer in LCPUFA than TAG, the vast bulk of LCPUFA in breast milk is carried by TAG because of its relative abundance(21). More recently, interest in the potential efficacy of PL to deliver DHA has become of renewed interest also for use in adults with the widespread availability of krill oil. Krill oil carries greater than one-third of its high level of EPA and DHA as PL (47), though notably, a recent report notes that
about 22% of the EPA and DHA in one krill oil product is in the form of free fatty acids, which is at least several fold greater than normally found in foods or supplements (48). In the first day post-dose, the timing and peak of blood levels is generally greater when ingested as free fatty acids, and similar to that of TAG (49, 50) and/or as PL compared to ethyl esters. However, these short term kinetics may not correspond to long term differences in DHA status as measured in human blood pools (48). Notably, however, blood FA status reflects fatty acids in circulation and does not address the uptake and incorporation of specific FA’s into target tissues, which can be measured directly with isotopically labeled precursors in animal models. Greater efficacy of PC compared to TAG carrier for ARA found with $^{13}$C-precursors in previous studies (21, 51) led to speculation that the same would be the case for DHA. Others have considered the relative efficacy of labeled DHA as carried by TAG and PC in the rat (52). In a previous study with radiolabeled ($^{14}$C) version of the same lipids we considered here, PC-DHA was twice as efficacious for brain DHA accretion as TAG-DHA at 10 weeks but not earlier. The peak of the rat brain growth spurt is maximal between 1 and 2 weeks (53), and accordingly the rat is long past the developmental time when the brain is growing. It is possible, however, that rat brain DHA may continue past the time when the brain ceases to increase in weight, similar to humans (6). The brains of our piglets were certainly increasing in weight during the period of our study and indicate that the advantage of PC is found perinatally. The pig is generally considered a good model for human brain development because it is among the few experimental animals with a brain growth spurt similar to humans (53), and it is a large non-ruminant omnivore.
Our data also show apparent retroconversion of DHA to $^{13}$C-22:5n-3, and a parallel advantage for PC. No labeled appeared in 20:5n-3 indicating chain shortening to 20:5n-3 was found in contrast to results in humans (54). Retroconversion in this context refers to the flow of $^{13}$C label from its original molecular form, 22:6n-3, to the more saturated 22:5n-3. Our measurements do not indicate the location of the labeling, whether all 22 carbons are labeled or possibly that 22:6n-3 might be β-oxidized to acetate and used for new synthesis of 22:5n-3 from 18:3n-3 or 20:5n-3. We did not find labeling in 20:5n-3 which would have been expected if 18:3n-3 was elongated with labeled acetate, and no evidence of label in other fatty acids, suggesting that the acetate pool was not a factor. In humans, 22:6n-3 β-oxidation is low compared to other fatty acids (55). These considerations all suggest that the label in 22:5n-3 did not proceed via acetate.

The pathway by which this transition occurs is not clear. The last few steps of the widely accepted coupled mitochondrial-peroxisomal pathway of 22:6n-3 synthesis proceed as (20:5 → 22:5 → 24:5 → 24:6) → 22:6, where all but the last step occurs on the endoplasmic reticulum, and 24:6 is transported to the peroxisomes for one round of β-oxidation (56), the latter of which is a four step FAD and NAD dependent process. Retroconversion in the sense of reversal of this pathway would require entry of 22:6n-3 into peroxisomes for an energetically and entropically expensive reversal of β-oxidation to yield 24:6n-3, followed by transport to the ER for a reversal of 6-desaturation and elongation, another four step process. Conventional elongation of
22:6n-3 to 24:6n-3 in the ER is a known process and is therefore the simplest explanation for a first (set of) steps, but it would then require reversal of desaturation and presumably β-oxidation of the resulting product, as $24:6 \rightarrow 24:5 \rightarrow 22:5$. Direct conversion of 22:6n-3 to 22:5n-3 would require the equivalent of a reversal of Δ4-desaturation in one step and is the most parsimonious explanation. Further studies are required to establish the molecular details of this pathway which have been hypothesized (57), and which operates in numerous species including vertebrate fish (58).

Gray matter DPA was greater than in reported for other species, and DHA lower (Table 3). Gray matter DHA in normal neonatal baboons has recently been found to be around 12-14%, while DPA was between 1-2% (2). DPA is well known to rise in neural tissue when omega-3 fatty acids are limited in the diet and with omega-6 being more abundant. It has been used as an index of relative omega-3 deficiency (59). In well nourished normally growing piglets, the relatively high amount of DPA indicates active demand for highly unsaturated fatty acids and presumably a maximally upregulated pathway for their biosynthesis. The sum of DHA and DPA in gray matter was 18% (w/w), similar to that for normal primates of about 16% (w/w). White matter consisting of a large proportion of mostly saturated myelin has 5.2% DHA and 3.7% DPA, suggesting that the DHA supply more adequately satisfies the demand in white matter. Although retinal DHA concentration is high, the total retina mass is small and the total DHA required is correspondingly small. We previously showed in primates
that small neural structures normalize DHA levels at lower dietary concentrations of DHA than the very large cerebral cortex (2); these data are thus consistent.

In summary, stable isotope tracer doses of DHA bound in the sn-2 position of PC was 1.9-fold more efficacious for supply of cerebral cortex gray matter than DHA bound to the sn-2 position of TAG. These data are generally consistent with numerous previous measurements on other LCPUFA and on DHA studied in other species. Together with previous work, the results indicate that PC is a highly efficacious source of both DHA and ARA, the two major LCPUFA of human breast milk (60).
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Figure legends

Figure 1. Results for primary outcome, cerebral cortex gray matter. (A) DHA* ($^{13}$C-22:6n-3) and DPAn-3* ($^{13}$C-22:5n-3) expressed as %Dose found in gray matter. PC was 1.9-fold more efficacious for supplying DHA to the neonatal piglet cortex than TAG. The ratio of efficacy was 2.8 for the retroconversion product DPAn-3. (B) $^{13}$C fatty acids found in gray matter synaptosomes normalized to the $^{13}$C-DHA-PC value. The ratio of efficacies for DHA and DPAn-3 in synaptosomes was 1.7 and 2.8, respectively. (C) The mass of unlabeled (tracee) DHA and DPAn-3 from a 500 mL meal from PC and TAG for gray matter and (D) synaptosomes. TAG is favored because it predominates in formula; no difference was found for DPAn-3. Unlabeled material is calculated from the tracer percentages and concentrations of tracee DHA in formula.

Figure 2. Liver and RBC tracer and tracee. (A) $^{13}$C-DHA in liver from PC and TAG. Liver retained about 7% of the DHA from the PC dose at 6 days compared to about 3.8% for the TAG dose, corresponding to a relative efficacy of about 1.9; PC-derived DPAn-3 was significantly greater than TAG-derived DPAn-3, with a relative efficacy of 2.6. (B) Similar relative efficacy was found for RBC DHA (1.6); RBC DPAn-3 was not significantly different. (C) and (D) show the mass (in mg) of DHA and DPAn-3 derived from unlabeled (tracee) TAG in the 500 mL meal predominated in whole liver and all RBC due to the relative amounts of tracee in the respective pools. Liver weight was as
measured at necropsy and RBC mass was estimated as described in the Methods section.

Figure 3. Retina. (A) Relative efficacy of PC compared to TAG was about 2.2 for DHA and 2.4 for DPA-3. (B) More retina DHA was derived from diet TAG than PC.
Table 1. Fatty acid profile of piglet milk formula (replacer), as % (w/w). *indicative of ruminant fat.

<table>
<thead>
<tr>
<th>FA</th>
<th>TAG</th>
<th>PL</th>
</tr>
</thead>
<tbody>
<tr>
<td>Saturated FA</td>
<td>42.32</td>
<td>55.84</td>
</tr>
<tr>
<td>Branched/CLA*</td>
<td>0.67</td>
<td>2.90</td>
</tr>
<tr>
<td>Monoene FA</td>
<td></td>
<td></td>
</tr>
<tr>
<td>18:2n-6</td>
<td>14.12</td>
<td>7.35</td>
</tr>
<tr>
<td>20:2n-6</td>
<td>0.57</td>
<td>0.34</td>
</tr>
<tr>
<td>20:3n-6</td>
<td>0.10</td>
<td>0.95</td>
</tr>
<tr>
<td>20:4n-6</td>
<td>0.25</td>
<td>2.63</td>
</tr>
<tr>
<td>22:4n-6</td>
<td>0.10</td>
<td>0.12</td>
</tr>
<tr>
<td>18:3n-3</td>
<td>0.53</td>
<td>0.69</td>
</tr>
<tr>
<td>20:3n-3</td>
<td>0.12</td>
<td>0.00</td>
</tr>
<tr>
<td>22:5n-3</td>
<td>0.04</td>
<td>0.38</td>
</tr>
<tr>
<td>22:6n-3</td>
<td>0.02</td>
<td>0.11</td>
</tr>
</tbody>
</table>
Table 2. Lipid and DHA distribution in piglet milk replacer.

<table>
<thead>
<tr>
<th></th>
<th>TAG</th>
<th>PL</th>
</tr>
</thead>
<tbody>
<tr>
<td>g FA per 500 mL</td>
<td>23.9</td>
<td>1.6</td>
</tr>
<tr>
<td>Approximate energy from FA (%)</td>
<td>43%</td>
<td>4%</td>
</tr>
<tr>
<td>% FA</td>
<td>94%</td>
<td>6%</td>
</tr>
<tr>
<td>mg DHA per 500 mL</td>
<td>4.8</td>
<td>1.8</td>
</tr>
<tr>
<td>% DHA</td>
<td>73%</td>
<td>27%</td>
</tr>
</tbody>
</table>
Table 3. Fatty acid composition of neonatal piglet tissues (pooled)

<table>
<thead>
<tr>
<th>Fatty acid</th>
<th>Gray matter</th>
<th>White matter</th>
<th>GM Synaptosomes</th>
<th>Retina</th>
<th>Liver</th>
<th>RBC</th>
</tr>
</thead>
<tbody>
<tr>
<td>16:0</td>
<td>20.45 ± 0.67</td>
<td>19.04 ± 1.03</td>
<td>22.49 ± 0.68</td>
<td>20.42 ± 0.78</td>
<td>17.00 ± 0.65</td>
<td>23.07 ± 1.47</td>
</tr>
<tr>
<td>18:0</td>
<td>20.05 ± 0.40</td>
<td>18.48 ± 0.76</td>
<td>18.06 ± 0.27</td>
<td>21.85 ± 0.72</td>
<td>24.04 ± 0.84</td>
<td>15.97 ± 0.75</td>
</tr>
<tr>
<td>∑SFA</td>
<td>41.95 ± 1.00</td>
<td>39.29 ± 1.53</td>
<td>42.83 ± 1.06</td>
<td>44.67 ± 1.05</td>
<td>42.22 ± 0.44</td>
<td>41.42 ± 2.10</td>
</tr>
<tr>
<td>MUFA</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>16:1</td>
<td>1.18 ± 0.15</td>
<td>2.85 ± 0.48</td>
<td>2.26 ± 0.22</td>
<td>1.64 ± 0.13</td>
<td>1.10 ± 0.13</td>
<td>1.02 ± 0.19</td>
</tr>
<tr>
<td>18:1</td>
<td>15.51 ± 0.75</td>
<td>25.77 ± 3.25</td>
<td>15.35 ± 0.55</td>
<td>16.74 ± 0.67</td>
<td>12.67 ± 0.68</td>
<td>34.36 ± 1.75</td>
</tr>
<tr>
<td>20:1</td>
<td>0.47 ± 0.08</td>
<td>1.50 ± 0.45</td>
<td>0.41 ± 0.09</td>
<td>0.30 ± 0.07</td>
<td>0.65 ± 0.19</td>
<td>0.34 ± 0.28</td>
</tr>
<tr>
<td>∑MUFA</td>
<td>18.12 ± 0.83</td>
<td>30.66 ± 4.01</td>
<td>18.28 ± 0.68</td>
<td>19.00 ± 0.73</td>
<td>14.50 ± 0.78</td>
<td>36.28 ± 1.70</td>
</tr>
<tr>
<td>n-6 PUFA</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>'18:2n-6</td>
<td>1.15 ± 0.07</td>
<td>1.23 ± 0.14</td>
<td>1.09 ± 0.11</td>
<td>2.29 ± 0.25</td>
<td>14.05 ± 0.79</td>
<td>15.40 ± 1.45</td>
</tr>
<tr>
<td>18:3n-6</td>
<td>0.09 ± 0.14</td>
<td>0.11 ± 0.03</td>
<td>0.11 ± 0.06</td>
<td>0.23 ± 0.10</td>
<td>0.29 ± 0.11</td>
<td>0.15 ± 0.07</td>
</tr>
<tr>
<td>'20:4n-6</td>
<td>11.15 ± 0.37</td>
<td>9.57 ± 0.74</td>
<td>9.77 ± 0.34</td>
<td>9.37 ± 0.47</td>
<td>17.19 ± 0.96</td>
<td>3.23 ± 1.23</td>
</tr>
<tr>
<td>'22:4n-6</td>
<td>6.64 ± 0.61</td>
<td>6.88 ± 1.00</td>
<td>6.97 ± 0.55</td>
<td>4.33 ± 0.28</td>
<td>2.16 ± 0.23</td>
<td>0.75 ± 0.29</td>
</tr>
<tr>
<td>'22:5n-6</td>
<td>9.69 ± 0.73</td>
<td>3.68 ± 0.89</td>
<td>9.47 ± 0.59</td>
<td>5.67 ± 0.72</td>
<td>3.11 ± 0.57</td>
<td>0.45 ± 0.17</td>
</tr>
<tr>
<td>∑n-6 PUFA</td>
<td>29.92 ± 0.97</td>
<td>22.78 ± 2.28</td>
<td>28.65 ± 1.02</td>
<td>22.91 ± 1.16</td>
<td>38.75 ± 0.60</td>
<td>21.00 ± 2.65</td>
</tr>
<tr>
<td>n-3 PUFA</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>18:3n-3</td>
<td>0.05 ± 0.03</td>
<td>0.08 ± 0.04</td>
<td>0.13 ± 0.08</td>
<td>0.10 ± 0.09</td>
<td>0.09 ± 0.07</td>
<td>0.12 ± 0.05</td>
</tr>
<tr>
<td>'22:5n-3</td>
<td>0.54 ± 0.17</td>
<td>0.35 ± 0.07</td>
<td>0.50 ± 0.07</td>
<td>0.74 ± 0.06</td>
<td>1.51 ± 0.17</td>
<td>0.48 ± 0.24</td>
</tr>
<tr>
<td>'22:6n-3</td>
<td>8.32 ± 0.68</td>
<td>5.21 ± 1.01</td>
<td>8.16 ± 0.65</td>
<td>12.02 ± 0.83</td>
<td>2.44 ± 0.23</td>
<td>0.47 ± 0.17</td>
</tr>
<tr>
<td>∑n-3 PUFA</td>
<td>8.93 ± 0.77</td>
<td>5.82 ± 1.03</td>
<td>8.86 ± 0.65</td>
<td>12.87 ± 0.82</td>
<td>4.04 ± 0.32</td>
<td>1.06 ± 0.39</td>
</tr>
</tbody>
</table>
Figure 1
Liu et al.
Figure 3

**Retina DHA (ng)**

- Retina DHA (%Dose)

**22:5n3**

- TAG
- PC

**22:6n3**

- TAG
- PC

* Values indicate significance.