Impact of n-3 Docosapentaenoic Acid Supplementation on Fatty Acid Composition in Rat Differs Depending upon Tissues and Is Influenced by the Presence of Dairy Lipids in the Diet

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Impact of n-3 Docosapentaenoic Acid Supplementation on Fatty Acid Composition in Rat Differs Depending upon Tissues and Is Influenced by the Presence of Dairy Lipids in the Diet

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Supporting Information

ABSTRACT: The n-3 docosapentaenoic acid (n-3 DPA) could be a novel source of n-3 long-chain polyunsaturated fatty acids (LCPUFA) with beneficial physiological effects. Following the supplementation of 0.5% purified n-3 DPA for 3 weeks from weaning, the n-3 DPA content increased in one-half of the 18 studied tissues (from +50% to +110%, p < 0.05) and mostly affected the spleen, lung, heart, liver, and bone marrow. The n-3 DPA was slightly converted into DHA (+20% in affected tissues, p < 0.05) and mostly retroconverted into EPA (35–46% of n-3 DPA intake in liver and kidney) showing an increased content of these LCPUFA in specific tissues. The partial incorporation of dairy lipids in the diet for 6 weeks increased overall n-3 PUFA status and brain DHA status. Furthermore, the n-3 DPA supplementation and dairy lipids had an additive effect on the increase of n-3 PUFA tissue contents. Moreover, n-3 DPA supplementation decreased plasma cholesterol.

KEYWORDS: polyunsaturated fatty acid (PUFA) metabolism, dairy lipids, cholesterol, eicosapentaenoic acid (EPA), tissue composition

1. INTRODUCTION

The n-3 docosapentaenoic acid (n-3 DPA) is a member of the n-3 long-chain polyunsaturated fatty acid (PUFA) essential family. It is also an intermediate between eicosapentaenoic acid (EPA) and docosahexaenoic acid (DHA) in the n-3 PUFA conversion pathway from α-linolenic acid (ALA).

Many studies have shown the beneficial role of n-3 PUFA in the reduction of cardiovascular risk, metabolic syndrome, cancer, and inflammation. Unfortunately, the n-3 PUFA conversion pathway is limited in humans as in rodents. Indeed, the conversion from ALA occurs through a sequence of desaturations and elongations including two enzymes considered as limiting steps: the Δ-6 desaturase and the elongase-2. Moreover, the main part of ALA intake is lower than the dietary recommendations in humans and is deposited in adipose tissues or catabolized by mitochondrial β-oxidation.

Furthermore, direct supplementation with the different precursors of DHA, α-linolenic acid (C18:3 n-3, ALA), stearidonic acid (C18:4 n-3), or EPA showed an increase in EPA and n-3 DPA, whereas the DHA status remained stable or was only slightly increased.

n-3 DPA was described as a potential reservoir of n-3 PUFA. Indeed, n-3 DPA could be interesting both for its potential capacity to be converted into DHA or retroconverted into EPA and for its potential physiological effects. This fatty acid (FA) has been poorly studied especially in vivo as compared to EPA and DHA, probably due to its relative lack of commercial availability in high quantity and purity. Consequently, the whole-body assimilation and FA metabolism of n-3 DPA has not been addressed yet. Nutritional sources of n-3 DPA are mainly sea products, meat including giblets, n-3 enriched food, and dairy products. Thereby, the n-3 DPA intake is mainly in the form of phospholipids and triglycerides in the food or in ethyl ester form in food supplement. Moreover, n-3 PUFA bioavailability seems better when they are consumed in complex matrix than as supplement form. This raises the question of the lipid mixture in which n-3 DPA is ingested to modulate its tissue delivery and metabolism after digestion and absorption.

Indeed, including dairy lipids in the diet increased the n-3 PUFA content, as compared to a lipid mixture of vegetable oils, in different nutritional and physiological conditions. More specifically, the DHA level was increased in the brain and the retina, while n-3 DPA was increased in some tissues like the heart, the liver, and red blood cells (RBC). These authors also showed that n-3 DPA content in RBC, which increased with dairy lipids diet, could act as a predictor of brain DHA status. Moreover, dairy products provide 9% of the n-3 DPA intake in Canadian pregnant and lactating women and 6% in children. However, the absolute intake remains low in Europe and is about 12–80 mg/day in human adults and about 30–40 mg/day in children. Nevertheless, dairy lipids could be a good lipid mixture for the assimilation of dietary n-3 DPA and for increasing the n-3 PUFA status.

The present study contains two experiments with the aim of showing whether the dietary n-3 DPA is assimilated within the...
tissues, and whether n-3 DPA is converted into DHA or retroconverted into EPA and could increase the overall n-3 PUFAs tissue status depending on the lipid mixture inside which it was included. In experiment 1, we studied the whole body assimilation and the overall FA metabolism of n-3 DPA, and we investigated which tissues were the most impacted by the n-3 DPA dietary supplementation for 3 weeks as ethyl ester. In experiment 2, we studied if n-3 DPA supplementation and the addition of dairy lipids in the diet for 6 weeks could have a complementary effect to increase n-3 PUFAs tissue status.

2. MATERIALS AND METHODS

2.1. Chemicals. Solvents and chemicals were obtained from Thermo Scientific (Elancourt, France), VWR (Fontenay-sous-Bois, France), or Sigma (Saint-Quentin-Fallavier, France). Polaris (Quimper, France) graciously provided high n-3 PUFAs marine oil (Omegavie 4020 EE Qualitysilver). Anhydrous milk fat and oleic sunflower oil were provided by Lactalis group (Retiers, France).

2.2. Animals. This study is composed of two experiments (Figure 1). In experiment 1, 32 rats (n = 8/group) were fed from weaning for 3 weeks with VO diet supplemented or not supplemented with 0.5% n-3 DPA (VO+DPA). In experiment 2, 32 rats (n = 8/group) were fed from weaning for 6 weeks with VO or DL diets not supplemented or supplemented with 0.5% n-3 DPA (VO+DPA and DL+DPA). For both experiments, Sprague–Dawley male rats (mean body weight 50 ± 1 g, 3 weeks old at beginning of the experiment) (Janvier Laboratories, Le Genest-Saint-Isle, France) were randomly assigned into the groups and were housed four animals by cage with free access to water and food on a 12 h light–dark cycle, maintained at a temperature of 21 ± 2 °C and at a hygrometry of 50% ± 10%. Body weight, food, and water intakes were measured two times a week. Sprague–Dawley rats were chosen considering their FA metabolism close to human, the nonconsanguinity of Sprague–Dawley rats ensures heterogeneity of the populations, the animal is large enough to obtain large enough samples for analysis purposes, and it is small enough to consume enough DPA to allow nutritional study. Only male animals were chosen to overcome the influence of estrogen and the menstrual cycle on fatty acid and lipid metabolisms in female rats.

All experiments were performed in accordance with the European Union Guidelines for Animal Care and Use (2010/63/CPE). The experimental procedures (no. APAFIS#1389-2015080411586889 v4 and no. APAFIS#S139-201605121200996 v1) were approved by the French Animal Care Committee (Rennes, approval number A3523838) and the Ministry of Higher Education, Research and Innovation, in compliance with recommendations of the 2013-118 French directive for animal experimentation. The number of animal by group was chosen considering a power calculation based on tissue FA composition fold-change in PUFAs after a dietary supplementation with 0.5% DHA in similar experimental conditions.24

At the end of the experiments, fasted rats were anesthetized with an intraperitoneal injection of sodium pentobarbital (140 mg/kg of body weight) (Euthasol Vet, France). Blood samples were collected by cardiac punctures into lithium heparin-treated vacutainers (Dominique Dutcher, Brumath, France), and plasma was separated from blood cells by centrifugation (3000g, 20 min, room temp) and stored at −80 °C. After removing the peripheral blood mononucleated cells layer, red blood cells (RBC) were mixed, weighted, and stored at +4 °C before lipid extraction. Whole eyes were removed by section of the optic nerve and then stored in 4% formaldehyde in PBS at 4 °C for at least 1 week; the retina was dissected by removing muscle membrane, optic nerve, cornea, lens, and vitreous body, and then washed in PBS. Tissues were removed, washed with PBS, weighted, and around 200 mg of each tissue was stored in dimethyloxime/methanol (4/1, v/v) at −20 °C. The rest of the organs were snap-frozen in liquid nitrogen and stored at −80 °C. In experiment 1, 18 tissues were collected and analyzed to realize the whole-body impact of n-3 DPA on fatty acid composition in tissue: bone marrow, brain, epididymal adipose tissue, heart, jejunum, kidney, liver, lung, muscle, pancreas, plasma, red blood cells, retina, subcutaneous adipose tissue, skin, spleen, stomach, and testis. In experiment 2, the tissues most impacted by n-3 DPA supplementation were collected as well as the brain. In experiment 2, we also studied the FA composition of the RBC and the brain, particularly interesting for the potential impact of dairy lipids on these tissues.

2.3. Diets. Four different lipid blends were prepared from a combination of natural fat sources (Table 1). Lipid blends were composed of a blend of vegetable oils (VO) or a blend of vegetable oils and dairy lipids (DL), not supplemented or supplemented with 0.5% n-3 DPA (w/w, 0.1% of total energy) (VO+DPA and DL+DPA). Lipid blends were made with particular attention to ALA and linoleic acid (LA) levels, with an LA/ALA ratio around 5 as nutritionally recommended in humans in France.22 The addition of short- and medium-chain fatty acids (SMCFA) with the introduction of dairy lipids was adjusted with oleic acid. FA composition of lipid blends composing the diets was analyzed by GC–MS after Folch extraction of diets and derivation as described above in section 2.5. Total sterols were measured in DL and VL blends by Liebermann method to estimate the cholesterol intake of animals as described previously.24

All of the diet contained 10% (w/w, 21% of total energy) of lipid blend and 90% (w/w) of a lipid-free base. The lipid-free base was prepared at the Unité de Production d’Aliments Expérimentaux (UPAE, INRA, Jouy-en-Josas, France). The lipid-free base contained for 1 kg of diet: 220.0 g of HCl-casein, 1.5 g of rhamnose, 402.3 g of corn starch, 201.2 g of sucrose, 20.0 g of cellulose, 45.0 g of mineral mix 102, and 10.0 g of vitamin mix 102. Next, 100.0 g of fat/kg (21% of total energy) of diet from the lipid blends was added in our laboratory.

To prepare the rat diet, the lipid-free base and the lipid blend were mixed then sifted twice, 13% water was added (w/w), and then the mixture was homogenized and shaped as croquettes. The rat diets were sealed in an airtight package and frozen at −20 °C, as all diets were prepared at the beginning of the study. During the experiments, rat diets were thawed at +4 °C 24 h before being given to animals. The n-3 DPA–EE possible peroxidation was controlled by TBA/R test on diets,24 and malondialdehyde was not detected in diets at the
The isocratic mode of methanol/water (98/2, v/v) (step 7). The granulometry of Interchrom preparative column (Uptisphere strategy C18-HQ, diameter 50 mm, length 250 mm, Interchim) with an isocratic mode of methanol/water (99/1, v/v) with a constant flow of 120 mL/min. Mass spectra were recorded with an Agilent Technologies 5975C inert MSD with triple axis detector. The mass spectrometer was operated under electron impact ionization conditions (electron energy 70 eV, source temperature 230 °C). Data were obtained in the full scan mode with a mass range of m/z 50–550 atomic mass units (amu), using the MSD Chemstation software E.02.02.14.31. Identification of the FA methyl esters was based upon retention times obtained for methyl ester of authentic standards, when available. The National Institute of Standards and Technology database v2.0 was also sometimes used to identify unknown FA. All identified FA with a signal/noise >10 were considered in the analysis of the proportions, and FA identified with a signal/noise between 3 and 10 was marked as trace.

The apparent retroconversion of n-3 DPA into EPA in the liver was estimated as percentage using the following formula:

\[
\frac{\Delta \text{EPA}_{(\text{supplemented-control})}}{\Delta \text{EPA + DPA}_{(\text{supplemented-control})}} \times 100
\]

2.4. n-3 Ethyl Docosapentaenoate (DPA-EE) Purification

The n-3 DPA-EE with purity >99% of total detected FA was obtained from a high n-3 PUFA marine oil (Omegavie 4020EE Qualisylvers, Polaris). The purification was performed by liquid chromatography in seven steps with a puriFlash4250 system (Interchrom, Montluçon, France) with a Chromabond flash column (RS 330 C18 ec, porosity 45 μm, diameter 60 mm, length 200 mm, Macherey Nagel, Hoerdt, France) followed by a Puriflash flash column (PF-15C18 HP-F0330, granulometry 15 μm, diameter 60 mm, length 226 mm, Interchim) and coupled with an UV detector. For the first two steps, the elution was performed at room temperature in an isocratic mode of methanol/water (99/1, v/v) with a constant flow of 120 mL/min. The fraction corresponding to n-3 DPA-EE was collected and was evaporated with a rotatory evaporator, then analyzed by gas chromatography—mass spectrometry (GC–MS) to control purity and efficiency. The collected fraction of step 2 then was reinjected with an elution of methanol/water (98/2, v/v) in the same conditions (step 3). This purification cycle was repeated three more times (steps 4–6) to gradually increase the purity in n-3 DPA-EE of the fraction of interest until 95%.

The fraction of step 6 was injected in the same conditions in an Interchrom preparative column (Uptisphere strategy C18-HQ, granulometry 5 μm, diameter 50 mm, length 250 mm, Interchim) with an isocratic mode of methanol/water (98/2, v/v) (step 7). The fraction corresponding to the n-3 DPA-EE peak was collected and pooled to obtain n-3 DPA-EE >99% purity of total detected FA (see Supporting Information A, Figure S1). The n-3 DPA-EE possible peroxidation was controlled by a TBARS test on n-3 DPA-EE oil. Malondialdehyde (TBARS) was not detected in the n-3 DPA-EE final purified fraction.

2.5. Lipid Extraction and Fatty Acid Composition

Lipids were extracted twice from tissues with 20 volumes dimethoxy-methane/methanol (4/1, v/v) after homogenization with an UltraTurrax. RBC lipid membranes and plasma were extracted twice with hexane/isopropanol (3/2, v/v) after acidification with HCl 3 M, as previously described. Brain phospholipids were separated from total lipids on preparative thin-layer chromatography (TLC) using silica gel H (Merck Millipore, Darmstadt, Germany) plates impregnated with a mixture of hexane/diethyether/acetich acid (85/15/1, v/v/v). Total lipids and phospholipids were saponified by 1 mL of 0.5 M NaOH in methanol at 70 °C for 30 min and then methylated with 1 mL of BF3 (12% in methanol) at 70 °C for 30 min. Fatty acid methyl esters were extracted twice with pentane, washed twice with NaCl 0.9%, and were taken up in hexane. Fatty acid methyl esters from brain phospholipids were purified by another step of TLC with a mixture of hexane/diethyether ethanol (80/20, v/v), extracted a first time by a mixture of methanol/hexane/NaCl 0.9% (3/4/3, v/v/v), then extracted a second time with 4 volumes of hexane, washed with NaCl 0.9%, and were taken up in hexane.

GC–MS analysis was performed using an Agilent 7890A (Agilent technologies, Santa Clara, CA) with a bonded silica capillary column (BPX 70, 60 m × 0.25 mm; SGE, Melbourne, Australia) containing a polar stationary phase of 70% cyanopropyl polysilphenylene–siloxy (0.25 µm film thickness). Helium was used as the carrier gas (average velocity 36 cm/s). The column temperature program started at 150 °C and gradually increased at 4 °C/min to 250 °C, and held at 250 °C for 10 min. Mass spectra were recorded with an Agilent Technologies 5975C inert MSD with triple axis detector. The mass spectrometer was operated under electron impact ionization conditions (electron energy 70 eV, source temperature 230 °C). Data were obtained in the full scan mode with a mass range of m/z 50–550 atomic mass units (amu), using the MSD Chemstation software E.02.02.14.31. Identification of the FA methyl esters was based upon retention times obtained for methyl ester of authentic standards, when available. The National Institute of Standards and Technology database v2.0 was also sometimes used to identify unknown FA. All identified FA with a signal/noise >10 were considered in the analysis of the proportions, and FA identified with a signal/noise between 3 and 10 was marked as trace.

The apparent retroconversion of n-3 DPA into EPA in the liver was estimated as percentage using the following formula:
variable coefficient of the first component calculated for each tissue or for each FA.

The aim of experiment 2 was to compare the combined effect of n-3 DPA supplementation with two lipid mixtures after 6 weeks of diet. FA composition of tissues, plasma assays, and organs weights of VO, VO+DPA, DL, and DL+DPA groups were compared with two-way ANOVA. The major aims were to study the effects of DPA supplementation (VO and DL vs VO+DPA and DL+DPA) and lipid mixture (VO and VO+DPA vs DL and DL+DPA) on FA composition of tissues, and the interaction between both effects. Type-II ANOVA was used when the interaction was nonsignificant, and type-III ANOVA was used when the interaction was significant. Body weight and food consumption were analyzed with the same model for repeated measures; whenever the model was significant, multiple comparisons according to Tukey procedure adjusted with fdr method were performed. Prior to analysis, normality of the data of the residuals was graphically assessed and homoscedasticity was supposed verified. For all procedures, a p-value <0.05 was considered as significant. All data were reported as mean ± SEM.

Redundancy analysis (RDA) was used in experiment 2 to study the global interaction between the n-3 DPA supplementation and the lipid mixture. RDA corresponds to ANOVA in a multivariate framework. Constraints (i.e., factors) were the n-3 DPA supplementation effect, the lipid mixture effect, and the interaction between both factors. RDA was performed on FA composition of the four most impacted tissues by the n-3 DPA supplementation in experiment 1. As FA compositions were presented as the percentage of total detected FA, and that the FA proportion took place in a large range of values, the data were transformed by centering log-ratio, then by centering-scaling. A permutation test was used to assess the significance of all constraints. This test was followed by pairwise comparisons using factor fitting to an ordination adjusted with fdr method to assess the significance between the four diets. FA explaining most of the variability between the two effects was chosen by considering the test of Pearson correlation coefficient between FA data and individual scores, and a Pearson correlation coefficient $R^2 > 0.5$.

Univariate analysis and redundancy analysis were realized using R software v3.4.2 and R packages car v2.1.5, RVAideMemoire v0.9.68, vegan v2.4.4, and chemometrics v0.1. OPLS-DA was performed with Simca software v14.1 (Umetrics, Umeå, Sweden).

3. RESULTS

The aim of experiment 1 was to study the whole-body assimilation and FA metabolism of the n-3 DPA after 3 weeks of dietary n-3 DPA supplementation, and it focused on 18 tissues. The complete FA compositions of tissues from experiment 1 are presented in Supporting Information B, Tables S1–S6. nd: not detected.

Figure 2. Fatty acid composition of tissues in n-3 docosapentaenoic acid (DPA), docosahexaenoic acid (DHA), and eicosapentaenoic acid (EPA) of rats fed with a control VO diet (white) or a 0.5% n-3 DPA-supplemented VO diet (black) during 3 weeks. Results were expressed as mean ± SEM. Significance between groups was evaluated by multiple Student t test ($^* p < 0.05; ^** p < 0.01; ^*** p < 0.001; td 0.1 < p < 0.05$). Complete fatty acids profiles of all tissues are available in Supporting Information B, Tables 1–6. nd: not detected.
of Diet. At the end of experiment 1, body weight gain, organ weight, and food intake did not differ significantly between groups (see Supporting Information A, Figure S2 and Table S1).

3.1.1. n-3 DPA Supplementation Increased n-3 PUFA and Decreased n-6 PUFA. Following the n-3 DPA supplementation, the n-3 DPA content ranked from 0.1% in epididymal adipose tissue to 2.6% in red blood cells, and it was not detected in testis and subcutaneous adipose tissue (Figure 2).

More precisely, the proportion of n-3 DPA increased significantly in the heart (2.1-fold), the lung (1.8-fold), the gut (1.8-fold), the spleen (1.8-fold), the bone marrow (1.5-fold), the RBC (1.4-fold), the kidney (1.3-fold), and the skin (1.5-fold) in the DPA-supplemented group. The n-3 DPA content tended to increase in the stomach (2.0-fold, \( p = 0.072 \)) and the pancreas (1.2-fold, \( p = 0.083 \)). In the epididymal adipose tissue, the n-3 DPA was detected only in the n-3 DPA supplemented group. By contrast, n-3 DPA levels were not

Figure 3. Ratio n-6 LCPUFA/LA from fatty acid composition of tissues of rats fed with a control VO diet (white) or a 0.5% n-3 DPA-supplemented VO diet (black) during 3 weeks. Results were expressed as mean ± SEM. Significance between groups was evaluated by multiple Student’s t test (\( *p < 0.05 \); \( **p < 0.01 \); \( ***p < 0.001 \)). Complete fatty acids profiles of all tissues are available in Supporting Information B, Tables 1–6.

Figure 4. OPLS-DA model of all fatty acids of the studied tissues of rats fed with a control diet (Ctrl) or a 0.5% n-3 DPA-supplemented diet (DPA) for 3 weeks. (A) OPLS-DA score plot. Each rat is represented by a point (i.e., 16 points), falling into either CTRL or DPA. A significant discrimination between both groups is obtained along with the horizontal axis (CV-ANOVA, \( p < 0.05 \)). \( R^2 \) (explaining class variance) and \( Q^2 \) (explaining class prediction) values were 0.986 and 0.776, respectively. (B) OPLS-DA loading plot. Each fatty acid (or variable) from one tissue is represented by a point. The horizontal axis is significantly characterized by colored fatty acids, according to their variable importance in projection coefficient (>1) and their jack-knife CI errors bars. In straight relationship with the score plot, rats fed the n-3 DPA diet are associated with higher n-3 PUFA status and lower n-6 PUFA status as compared to Ctrl. The converse occurs for the rats fed the Ctrl diet. To make it clearer, some nonsignificant variables were randomly removed from the loading plot. Supporting Information B, Tables 1–6 gathers all fatty acids considered in the OPLS-DA model. Bmw, bone marrow; Brn, brain; Epf, epididymal adipose tissue; Hrt, heart; Kdy, kidney; Liv, liver; Lng, lung; Mus, muscle; Pan, pancreas; Plm, plasma; Rbc, red blood cells; Ret, retina; Scf, subcutaneous adipose tissue; Skn, skin; Spn, spleen; Stc, stomach.
Table 2. n-3 PUFA Tissue Contents

<table>
<thead>
<tr>
<th></th>
<th>FA %</th>
<th>VO</th>
<th>VO+DPA</th>
<th>DL</th>
<th>DL+DPA</th>
<th>DPA</th>
<th>LM</th>
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<tr>
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<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
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</tr>
<tr>
<td>ALA</td>
<td>0.48 a</td>
<td>±0.07</td>
<td>0.50 a</td>
<td>±0.07</td>
<td>0.63 ab</td>
<td>±0.06</td>
<td>0.69 b</td>
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<td>EPA</td>
<td>0.14 A</td>
<td>±0.01</td>
<td>0.16 A</td>
<td>±0.02</td>
<td>0.33 B</td>
<td>±0.04</td>
<td>0.67 C</td>
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<tr>
<td>n-3 DPA</td>
<td>0.37 A</td>
<td>±0.02</td>
<td>0.44 A</td>
<td>±0.02</td>
<td>0.47 A</td>
<td>±0.02</td>
<td>0.96 B</td>
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<tr>
<td>DHA</td>
<td>6.11</td>
<td>±0.29</td>
<td>6.69</td>
<td>±0.52</td>
<td>5.86</td>
<td>±0.21</td>
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<td>±0.54</td>
<td>7.29 a</td>
<td>±0.22</td>
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<tr>
<td>ALA</td>
<td>0.25</td>
<td>±0.03</td>
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<td>±0.07</td>
<td>0.35</td>
<td>±0.04</td>
<td>0.33</td>
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<tr>
<td>EPA</td>
<td>0.10 A</td>
<td>±0.02</td>
<td>0.13 AB</td>
<td>±0.01</td>
<td>0.19 C</td>
<td>±0.01</td>
<td>0.16 BC</td>
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<td>n-3 DPA</td>
<td>1.38 a</td>
<td>±0.15</td>
<td>2.36 c</td>
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<td>±0.08</td>
<td>2.51 c</td>
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<tr>
<td>DHA</td>
<td>11.9 a</td>
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<td>13.1 b</td>
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<td>±0.50</td>
<td>15.9 bc</td>
<td>±0.26</td>
<td>14.8 ab</td>
<td>±0.47</td>
<td>16.2 c</td>
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<tr>
<td>ALA</td>
<td>0.34</td>
<td>±0.03</td>
<td>0.26</td>
<td>±0.04</td>
<td>0.32</td>
<td>±0.03</td>
<td>0.32</td>
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<td>EPA</td>
<td>0.30 a</td>
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<td>0.34 ab</td>
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<td>0.42 b</td>
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<td>±0.12</td>
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<td>±0.04</td>
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<td>2.02 B</td>
<td>±0.18</td>
<td>2.16 B</td>
<td>±0.07</td>
<td>2.19 B</td>
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<td>3.28 b</td>
<td>±0.34</td>
<td>4.27 bc</td>
<td>±0.09</td>
<td>4.68 c</td>
</tr>
<tr>
<td>spleen</td>
<td></td>
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<td></td>
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<td></td>
<td></td>
</tr>
<tr>
<td>ALA</td>
<td>0.27</td>
<td>±0.05</td>
<td>0.64</td>
<td>±0.22</td>
<td>0.33</td>
<td>±0.02</td>
<td>0.49</td>
</tr>
<tr>
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<td>±0.03</td>
<td>0.13 a</td>
<td>±0.02</td>
<td>0.19 ab</td>
<td>±0.02</td>
<td>0.22 b</td>
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<tr>
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<td>±0.03</td>
<td>1.00 b</td>
<td>±0.12</td>
<td>1.09 b</td>
<td>±0.06</td>
<td>1.39 c</td>
</tr>
<tr>
<td>DHA</td>
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<td>±0.10</td>
<td>1.95</td>
<td>±0.20</td>
<td>1.93</td>
<td>±0.11</td>
<td>1.98</td>
</tr>
<tr>
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<td>±0.26</td>
<td>3.62 ab</td>
<td>±0.35</td>
<td>3.55 ab</td>
<td>±0.17</td>
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<td></td>
</tr>
<tr>
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<td>0.68 AB</td>
<td>±0.23</td>
<td>1.06 B</td>
<td>±0.09</td>
<td>0.42 A</td>
</tr>
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<td>±0.11</td>
<td>0.47 ab</td>
<td>±0.04</td>
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<td>±0.03</td>
<td>1.05 b</td>
<td>±0.05</td>
<td>1.46 d</td>
</tr>
<tr>
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<td>4.23 a</td>
<td>±0.17</td>
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<td>±0.14</td>
<td>4.18 a</td>
</tr>
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<td>±0.26</td>
<td>6.58 b</td>
<td>±0.29</td>
<td>5.96 b</td>
<td>±0.23</td>
<td>6.70 b</td>
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</table>

“Total fatty acids proportions of a-linolenic acid (ALA, C18:3n-3), eicosapentaenoic acid (EPA, C20:5n-3), n-3 docosapentaenoic acid (n-3 DPA, C22:5n-3), docosahexaenoic acid (DHA, C22:6n-3), and total n-3 long-chain polyunsaturated fatty acids (∑n-3) in tissues of rats fed the vegetable oils blend diet (VO) or the vegetable oil and dairy lipid blend diet (DL) not supplemented or supplemented with 0.5% DPA (VO+DPA and DL+DPA). Values are mean ± SEM. DPA supplementation effect (DPA) compared both not supplemented groups with both supplemented groups; lipid mixture effect (LM) compared both VO groups with both DL groups; *p < 0.05, **p < 0.01, ***p < 0.001. Different letters indicate significant differences for multiple comparisons (Tukey, p < 0.05). Uppercase letters indicate a significant interaction between DPA and LM effects. nd, nondetected; FA, fatty acids; RBC, red blood cells.

affected by DPA supplementation in liver, plasma, brain, retina, and muscle.

Following the n-3 DPA supplementation, the DHA content in studied tissues ranked from 0.2% in skin to 15.5% in brain and was not detected in adipose tissues. More precisely, the proportion of DHA increased significantly in the spleen (1.2-fold) and the lung (1.2-fold). The DHA content tended to increase in the bone marrow (1.6-fold, p = 0.099) and the stomach (1.5-fold, p = 0.069). DHA levels were similar in both groups in the other tissues.

Following the n-3 DPA supplementation, the EPA content in studied tissues ranked from 0.1% in the bone marrow to 1.4% in the pancreas. More precisely, the proportion of EPA increased significantly in the liver (2.0-fold), the plasma (2.0-fold), the spleen (1.5-fold), the lung (1.3-fold), and the stomach (2.6-fold). The EPA content tended to increase in the bone marrow (1.1-fold, p = 0.074) and the pancreas (1.4-fold, p = 0.096). The estimated retroconversion of DPA into EPA in the liver was 46.6% ± 7.9%. EPA was not detected in the brain, the retina, the skin, and adipose tissues.

Concerning n-6 PUFA, the ratio of the n-6 PUFA (sum of C18:3 n-6, C20:3 n-6, C20:4 n-6, C22:4 n-6, and C22:5 n-6) divided by the n-6 PUFA precursor (C18:2 n-6) was used to estimate the n-6 PUFA conversion pathway rate (Figure 3). Following the n-3 DPA supplementation, the ratio decreased significantly in the RBC (1.3-fold), the pancreas (1.3-fold), the liver (1.2-fold), the kidney (1.2-fold), and the brain (1.1-fold).

3.1.2. Global and Comparative Impact of n-3 DPA Supplementation on Tissues. To further underline the differences in tissue FA composition induced by n-3 DPA supplementation, an OPLS-DA was performed (Figure 4). All identified FA were taken into account in the model, which expressed 40.7% of the total variance R² (CV-ANOVA, p = 0.047) (see Supporting Information B for complete tissue FA profiles). A significant discrimination between both groups is obtained along with the horizontal axis (CV-ANOVA, p < 0.05) (Figure 4A), showing that the n-3 DPA dietary supplementation affected the overall FA tissue compositions. R² (explaining class variance) and Q² (explaining class prediction) values were 0.986 and 0.776, respectively. The n-3 DPA supplemented group was discriminated to the right of the horizontal axis of the score plot (Figure 4A). Thus, the n-3 DPA group was positively associated with variables at the right of the loading plot (Figure 4B) and negatively associated with variables at the left of the loading plot. The converse occurs for the rats fed the control group. Moreover, the association is stronger for variables close to the origin of the vertical axis than distant variables.
First, we hierarchized the different FA contents affected by n-3 DPA supplementation (Figure 4B). As a general result, n-3 DPA supplementation significantly increased only n-3 PUFA, decreased only n-6 PUFA, and did not impact saturated n-7 and n-9 FA. More precisely, among the n-3 PUFA, n-3 DPA > EPA > DHA > ALA were positively impacted by n-3 DPA supplementation. Among the n-6 PUFA, n-6 DPA > C22:4 n-6 > C20:4 n-6 > C18:2 n-6 were negatively impacted by n-3 DPA supplementation.

We then were interested in the comparison between tissues (Figure 4B). When considering all FA contents, the tissues the most impacted by the n-3 DPA supplementation were ranked as follows: spleen > lung > heart > liver > skin > gut > kidney > pancreas > bone marrow > epididymal adipose tissue > RBC > brain. When considering n-3 PUFA content, the tissues most impacted by the n-3 DPA supplementation were ranked as follows: spleen > lung > heart > liver > bone marrow > epididymal fat > skin > kidney > RBC ≫ brain. When considering n-6 PUFA content, the tissues most impacted by the n-3 DPA supplementation were ranked as follows: heart > lung > spleen > liver > pancreas > skin > kidney > gut > brain.

3.2. Experiment 2: Combined Effects of Dairy Lipid Mixture (DL) and n-3 DPA Supplementation after 6 weeks of Diet As Compared to a Vegetable Oils Mixture (VO). The major aim of experiment 2 was to study the potential impact of two lipid mixtures (lipid mixture effect: VO and VO+DPA vs DL and DL+DPA) not supplemented or supplemented with DPA (DPA effect: VO and DL vs VO +DPA and DL+DPA) on PUFA metabolism. Moreover, we studied if the impact of n-3 DPA supplementation was affected by the lipid mixture (interaction between DPA effect and lipid mixture effect).

We focused on the FA composition of the liver as the major metabolic organ, and the heart, the spleen, and the lung as the most impacted tissues by the n-3 DPA supplementation in experiment 1. We also studied the FA composition of the RBC and the brain, particularly interesting for the potential impact of fatty acids on these tissues, as previously described.25 The tables of tissues FA composition of all identified FA are presented in Supporting Information C, Tables S1–S6. At the end of experiment 2, body weight gain, organ weight, and food intake did not differ significantly between groups (see Supporting Information A, Figure S2 and Table S1).

3.2.1. Dairy Lipids Improved n-3 PUFA Status in Tissues. We first considered the effect of the lipid mixture on n-3 PUFA metabolism (Table 2). The n-3 DPA content was higher in liver, heart, lung, spleen, and RBC of rats fed DL diets than in rats fed VO diets. The DHA content of rats fed DL diets was higher in the lung only, as well as the total n-3 PUFA content.

The EPA content of rats fed DL diets increased in the liver and the RBC as compared to VO diets. In brain phospholipids, the DHA content increased in the DL group as compared to both VO and VO+DPA groups (Figure 5). The n-3 DPA supplementation did not increase the n-3 PUFA status in the brain (see Supporting Information C, Table S6 for brain complete FA profile).

We then considered the effect of the lipid mixture on n-6 PUFA metabolism (Table 3). The total n-6 PUFA increased in the heart, the lung, and the RBC. The n-6 DPA content of rats fed DL diets decreased in the heart and the spleen. The arachidonic acid (C20:4 n-6) content of rats fed DL diets decreased in the RBC, while it increased in the lung. The arachidonic acid (C20:4 n-6) content of rats fed DL diets decreased in the liver. When considering n-3 PUFA content, the tissues most impacted by the n-3 DPA supplementation were ranked as follows: spleen > lung > heart > liver > pancreas > bone marrow > epididymal adipose tissue > RBC > brain.

3.2.2. n-3 DPA Supplementation Increased n-3 PUFA and Decreased n-6 PUFA. We first considered the effect of the n-3 DPA supplementation on n-3 PUFA metabolism (Table 2). The n-3 DPA content as well as the total n-3 PUFA increased in all studied tissues after the n-3 DPA supplementation. The DHA content increased in the heart, the lung, and the RBC. The EPA content increased in the liver only. The estimated retroconversion of DPA into EPA in the liver was 35.1 ± 6.4% and did not differ between the four groups. The ALA content increased in the spleen and the RBC.

We then considered the effect of the n-3 DPA supplementation on n-6 PUFA metabolism (Table 3). The total n-6 PUFA content decreased only in the heart after the n-3 DPA supplementation. The n-6 DPA content decreased in the liver, the heart, and the spleen. The arachidonic acid (C22:4n-6) content decreased in the heart and the spleen. The arachidonic acid content decreased only in the spleen. The LA content decreased in the liver and the heart of n-3 DPA supplemented rats.

3.2.3. n-3 DPA Supplementation Differently Affected Tissue FA Composition in Combination with Dairy Lipids or Vegetable Oils Mixtures. We used redundancy analysis (RDA) to observe if the n-3 DPA supplementation affected the tissues FA composition differently, depending on the lipid mixture where n-3 DPA was included (VO or DL). We chose to study the four tissues most impacted by the n-3 DPA supplementation reported in experiment 1: the liver, the heart, the lung, and the spleen.

We first included all identified FA in the model to observe the potential interaction between n-3 DPA supplementation and the lipid mixture on the global fatty acid metabolism (Figure 6A). The model expressed 37.9% of constrained variance. The first component (horizontal axis) corresponded to the lipid mixture effect (VO and VO+DPA vs DL and DL+DPA, p = 0.001). The second component (vertical axis)
corresponded to the n-3 DPA supplementation (VO and DL vs VO+DPA and DL+DPA, p = 0.007). Differences between FA profiles were more enhanced by the lipid mixture effect (73.9% of constrained variance) than by the n-3 DPA supplementation (17.6% of constrained variance). The significant interaction between the lipid mixture and the n-3 DPA supplementation (p < 0.020) demonstrated that the n-3 DPA supplementation differently affected rats depending of the lipid mixture. Therefore, if we then compared each group, the n-3 DPA supplementation did not affect the total FA profiles when rats were fed with DL diets (DL vs DL+DPA, p = 0.3710). Conversely, the total FA profiles of rats fed VO diets were impacted by the n-3 DPA supplementation (VO vs VO+DPA, p = 0.0012). Moreover, both DL and DL+DPA groups were significantly different from both VO and VO+DPA groups (p = 0.0012). The C14:0, C15:0, and the n-3 DPA to a lesser extent were the most associated FA with both DL diets (see Supporting Information D, Figure S1 for loading plots).

As DL diets were mostly discriminated by saturated fatty acids (C14:0 and C15:0) and n-3 DPA, we then considered all of the identified PUFA to observe only the potential interaction of the n-3 DPA supplementation and the lipid mixture on the PUFA acid metabolism (Figure 6B). The model explained 31.4% of constrained variance. The first component expressed 68.2% of constrained variance and significantly discriminated all groups (p < 0.05). The first component was positively characterized by the n-3 PUFA, and negatively by the n-6 PUFA (Figure 6B). Thus, the VO group was associated with the highest n-6 PUFA status, while the DL+DPA group was associated with the highest n-3 PUFA status. The VO+DPA group had an intermediate PUFA status between the VO and the DL group. The DL group had an intermediate PUFA status between the VO+DPA and the DL+DPA group. In conclusion, the n-3 PUFA global status corresponded to VO < VO+DPA < DL < DL+DPA (see Supporting Information D, Figure S2 for loading plots).

3.2.4. Plasma Cholesterol Decreased with n-3 DPA Supplementation and Increased with DL Diet.

DIET. DL diets contained 6.6 mg of cholesterol/100 g of diet, whereas cholesterol was absent in VO diets. Plasma total cholesterol increased in the DL group only (Figure 7). This increase in total cholesterol was associated with an increase in cholesterol esters and an increase of the non-HDL cholesterol fraction and so the cholesterol/HDL cholesterol ratio. The supplementation of DL diet with n-3 DPA maintained all of these amounts at the same level as both VO and VO+DPA groups.

Table 3. n-6 PUFA Tissue Contents

<table>
<thead>
<tr>
<th>FA %</th>
<th>VO</th>
<th>VO+DPA</th>
<th>DL</th>
<th>DL+DPA</th>
<th>DPA</th>
<th>LM</th>
</tr>
</thead>
<tbody>
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<td>liver</td>
<td>LA</td>
<td>±0.35</td>
<td>6.83 b</td>
<td>±0.27</td>
<td>9.38 c</td>
<td>±0.27</td>
</tr>
<tr>
<td></td>
<td>ARA</td>
<td>±0.63</td>
<td>14.3</td>
<td>±0.92</td>
<td>14.1</td>
<td>±0.54</td>
</tr>
<tr>
<td></td>
<td>AdA</td>
<td>±0.02</td>
<td>0.21 ab</td>
<td>±0.02</td>
<td>0.19 b</td>
<td>±0.01</td>
</tr>
<tr>
<td>n-6 DPA</td>
<td>±0.02</td>
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<td>±0.02</td>
<td>0.19 a</td>
<td>±0.01</td>
<td>0.11 b</td>
</tr>
<tr>
<td></td>
<td>∑n-6</td>
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<td>2.22</td>
<td>±1.10</td>
<td>24.8</td>
<td>±0.73</td>
</tr>
<tr>
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<td>LA</td>
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<td>±0.36</td>
<td>15.0 C</td>
<td>±0.34</td>
</tr>
<tr>
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<td>±0.27</td>
<td>20.9</td>
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</tr>
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<td>±0.77</td>
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<td>0.23 b</td>
<td>±0.03</td>
<td>0.24 ab</td>
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<td>0.20 b</td>
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<tr>
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<td>34.9 A</td>
<td>±0.97</td>
<td>32.0 B</td>
<td>±0.86</td>
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*Proportions of linoleic acid (LA, C18:2n-6), arachidonic acid (ARA, C20:4n-6), adrenic acid (AdA, C22:4n-6), n-6 docosapentaenoic acid (n-6 DPA, C22:5n-6), and total n-6 long-chain polyunsaturated fatty acids (∑n-6) in tissues of rats fed the vegetable oils blend diet (VO) or the vegetable oil and dairy lipid blend diet (DL) not supplemented or supplemented with 0.5% DPA (VO+DPA and DL+DPA). Values are mean ± SEM. DPA supplementation effect (DPA) compared both not supplemented groups with both supplemented groups; lipid mixture effect (LM) compared both VO groups with both DL groups (*p < 0.05, **p < 0.01, ***p < 0.001). Different letters indicate significant differences for multiple comparisons (Tukey, p < 0.05). Uppercase letters indicate a significant interaction between DPA and LM effects. FA, fatty acids; RBC, red blood cells.
4. DISCUSSION

This study first addressed the impact of the nutritional n-3 DPA supplementation on tissues FA composition after 3 or 6 weeks of diet.

Nowadays, only few studies have examined the nutritional and biochemical effects of n-3 DPA in vivo. This could be mainly due to the high cost and the limited availability of highly purified DPA these days, not allowing wide intervention studies in humans for the moment. In this study, we proposed an innovative approach to purify n-3 DPA from fish oil enriched in n-3 PUFA by liquid chromatography (see section 2.2). This methodology, accessible in research laboratory, could help.

First, we showed that n-3 DPA supplementation increased n-3 DPA content only in some tissues: the spleen, the lung, the heart, the liver, the RBC, the bone marrow, and the kidney. This specificity could come from the fact that the tissues most impacted by the n-3 DPA supplementation are (i) either tissues in which the metabolism of fatty acids is important and the enzymes of the PUFA conversion pathways are strongly

![Figure 6. Redundancy analysis of fatty acids (FA) composition of the liver, heart, lung, and spleen for (A) all identified FA, and (B) long-chain polyunsaturated FA, of rats fed the vegetable oils blend diet (VO, ○) or the vegetable oil and dairy lipid blend diet (DL, □) not supplemented or supplemented with 0.5% DPA (VO+DPA, ●; and DL+DPA, ■). The represented barycenter of each group of rats. A permutation test was performed to consider the significance of the n-3 DPA supplementation effect (VO and DL vs VO+DPA and DL-DPA) and the lipid mixture effect (VO and VO+DPA vs DL and DL+DPA) with the two components, or considering their interaction (n-3 DPA supplementation:lipid mixture effects). This test was followed by pairwise comparisons using fitting factor into an ordination to assess the significance between the four groups (p < 0.05), indicated by different superscript characters on the group name. Both loadings plot are presented in Supporting Information D, Figures 1 and 2.]

![Figure 7. Plasma cholesterol concentrations of rats fed the vegetable oils blend diet (VO) or the vegetable oil and dairy lipid blend diet (DL) not supplemented or supplemented with 0.5% DPA (VO+DPA and DL+DPA). Values are mean ± SEM. Different superscript characters indicate significant differences for Tukey posthoc test for multiple comparisons (p < 0.05), following the one-way ANOVA if significant (p < 0.05).]
expressed (liver, kidney, bone marrow),\textsuperscript{36} (ii) either tissues rich in mitochondria and energy intensive, the energetic role being the first quantitative use of fatty acids (heart, lung),\textsuperscript{37} (iii) either marker tissues of food consumption and fatty acid metabolism of animals (RBC, spleen),\textsuperscript{38} or (iv) either tissues exhibiting the highest proportions of n-3 DPA.\textsuperscript{39}

Despite the relative lack of literature on the subject, our results are of interest when considering the physiological effects reported in these tissues. Concerning the spleen, a major organ involved in inflammation regulation, some experiments showed that n-3 DPA reduced inflammation in in vitro models,\textsuperscript{40} associated with the action of some hydroxy-metabolites directly derived from n-3 DPA like resolvins.\textsuperscript{41} Concerning the lung, one epidemiological study showed that n-3 DPA is the most potent fatty acid associated with better forced expiratory volume and could prevent chronic obstructive pulmonary disease.\textsuperscript{42} Indeed, in addition to its anti-inflammatory properties, n-3 DPA have antiproteolytic and antioxidative abilities by decreasing nuclear factor-kappa B (NF-kB) and p38 mitogen-activated protein kinases (p38 MAPK) activation, leading to a decrease in vascular endothelial growth factor (VEGF), and matrix metalloproteinase-2 and -9 (MMP-2 and MMP-9) expression levels in lung in a model of rat pulmonary hypertension.\textsuperscript{43} Concerning the heart, n-3 DPA seems to exert similar cardiometabolic protection effects as DHA in high-fat diet fed mice,\textsuperscript{44} and a high n-3 DPA RBC level was associated with a lower risk of metabolic syndrome in human adults.\textsuperscript{3,5} These effects could arise from the capacity of dietary DPA to decrease plasma lipid parameters\textsuperscript{25} by inhibiting the expression of genes involved in the lipogenesis like Sterol Regulating Element Binding Protein (SREBP), 3-Hydroxy-3-Methyl-Glutaryl-Coenzyme A reductase (HMG-CoA-R), Acetyl Coenzyme A Carboxylase (ACC), and Fatty Acid Synthase (FAS).\textsuperscript{46} Moreover, some studies showed that n-3 DPA inhibited platelet aggregation stimulated by collagen or arachidonic acid by accelerating the lipoxygenase pathway and by interfering with the cyclooxygenase pathway dose-dependently.\textsuperscript{47} These suggested effects linked to n-3 DPA could arise from the recently described bioactive metabolites derived from n-3 DPA, which could require a special attention in further studies.\textsuperscript{48,49} Surprisingly, while n-3 DPA has been described to exert a neuroprotective effect, we did not found a major change in brain FA composition following the n-3 DPA supplementation, but brain n-3 DPA content increased following the n-3 DPA dietary supplementation.\textsuperscript{4} We now wonder if the effects of n-3 DPA are independent of the other n-3 PUFA or shared with them.\textsuperscript{14}

On the other hand, the n-3 DPA could be interesting to lead to an increase in the total n-3 PUFA status. It is well-known that the n-3 PUFA conversion pathway was described as limited by the Δ-6 desaturase enzyme, the elongase-2 enzyme, and the final step of β-peroxisomal oxidation of the C24:6 n-3 to DHA.\textsuperscript{50} The elongase-2 acts twice in PUFA conversion pathway, one time in association with the elongase-5 to convert EPA to DPA (or arachidonic acid to C22:4n-6) and another time to convert EPA to C24:5 n-3 (or C22:4n-6 to C24:4n-6), for the n-3 (and n-6) conversion pathway, respectively.\textsuperscript{51} A few years ago, some authors showed that the elongase-2 enzyme could be the more limiting step to the conversion of n-3 DPA to DHA in brain astrocyte\textsuperscript{52} and in rat liver.\textsuperscript{53} As the direct supplementation with n-3 DPA shunted the contribution of the elongase-2 for converting EPA to n-3 DPA, one could hypothesize that n-3 DPA could be a good precursor for DHA or act as an in vivo stock of DHA.\textsuperscript{24,25} We showed a significant but slight increase in DHA content in the spleen and in the lung after 3 weeks of n-3 DPA supplementation and of the lung, the heart, and the RBC after 6 weeks. The n-3 DPA could therefore be considered for its ability to increase the DHA status. However, we must keep in mind that the increase in DHA remained slight. These results are in good agreement with previous reports describing a low level of the saturation plateau in the conversion pathway to DHA.\textsuperscript{6,7,9,10}

We also showed in both experiments that a large part of the n-3 DPA content in the liver was retroconverted into EPA (46% and 35% of apparent retroconversion). Indeed, the EPA content increased in the liver, the plasma, the spleen, and the lung in our study. The n-3 DPA apparent retroconversion, as well as the DHA apparent retroconversion to EPA, were previously described in vitro\textsuperscript{52} and in vivo,\textsuperscript{53} indicating that both n-3 DPA and DHA retroconversion involved the peroxisomal acyl-CoA oxidase.\textsuperscript{54}

Another point of our study is the decrease in n-6 PUFA content in tissues following the n-3 DPA supplementation, and mainly n-6 DPA and arachidonic acid (C22:4 n-6). We also showed by OPLS-DA that n-3 DPA supplementation only impacted n-3 and n-6 PUFA tissue compositions, and not saturated n-7 and n-9 FA. These results are surely due to the similarity of both n-3 and n-6 conversion pathways as compared to each others and by the competition effect of n-3 and n-6 PUFA on the Δ-6, Δ-5 desaturases and elongase-2 enzymes, which act in the synthesis of n-6 DPA from linoleic acid.\textsuperscript{35}

Furthermore, our results are interesting in terms of human nutrition because n-3 DPA is the only n-3 PUFA between EPA and DHA that is present in diet source in non-negligible amounts.\textsuperscript{17} Thus, we showed for the first time that DPA intake should be considered as a novel source of EPA as well as DHA for assessing the n-3 LC-PUFA nutritional status in rats, and to a larger extent in humans because the DPA content of products from animal husbandry or agriculture is not taken into account nowadays.\textsuperscript{55} Moreover, n-3 DPA decreased n-6 PUFA status and could have interesting and poorly known beneficial physiological effects that need more investigations.

This study second addressed the impact of the lipid mixture to modulate the effects of the n-3 DPA supplementation on the FA composition of the tissues most impacted by the n-3 DPA supplementation. Indeed, the second main finding of our study was that a partial incorporation of dairy lipids in the diet increased the PUFA content in tissues, as compared to a diet composed of a vegetable oil blend. Interestingly, this increase in PUFA status was higher in the nonsupplemented DL group than in the VO+DPA group, suggesting a more important impact of the lipid mixture than the direct n-3 DPA supplementation. Moreover, the n-3 PUFA status increasing effect was higher in the DL+DPA group than in the DL group in all tissues except in the brain, which was only affected by the lipid mixture. This showed an additive effect of dairy lipids and n-3 DPA supplementation on PUFA status. It has been previously shown that dairy lipids could improve the n-3 PUFA tissue content, principally up to n-3 DPA in most organs and up to DHA in the brain and retina in specific conditions.\textsuperscript{31,23,25,56}

Three mechanistic hypotheses could explain the increase of n-3 PUFA with DL diets. First, it has been reported that myristic acid (C14:0) present in dairy lipids improved the

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activity and the mRNA expression of the elongase-2 and the Δ-6 desaturase, the limiting enzymes of the n-3 and n-6 PUFA conversion pathway, resulting in an increase in PUFA tissues content. Second, dairy lipids contain short- and medium-chain FA, which have a higher β-mitochondrial oxidation rate than ALA, and could so prevent part of the β-oxidation of PUFA precursors. In our study, ALA content was higher in the liver of the DL group, strengthening this hypothesis. Third, DL diets contained a lower level of oleic acid as compared to VO diet. As oleic acid is also a substrate of Δ-6 desaturase, it could also compete with this enzyme more for VO groups than for DL groups. These results open new perspectives for the improvement of n-3 LC-PUFA status in early life in humans, through the incorporation of dairy lipids instead of high amounts of DHA. Indeed, the lipid compositions of our diets were based on commercial infant formulas, and most infant formula contains a mixture of vegetable oils and no dairy lipids, although closer to the composition of breastmilk.

To conclude, the 0.5% n-3 DPA supplementation increased some EPA and DHA tissue content and mainly affected the spleen, the heart, the lung, and the liver. A partial incorporation of dairy lipids in the diet increased n-3 PUFA status. Moreover, the n-3 DPA supplementation combined with a partial incorporation of dairy lipids had a positive and complementary effect on the n-3 PUFA status in all studied tissues except in the brain. Finally, the DHA status increased in brain phospholipids with dairy lipid mixture only.

**ASSOCIATED CONTENT**

Supporting Information The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acs.jafc.8b03069.

Supporting Information A, diets and animals; Figure S1, purification of n-3 DPA by liquid chromatography; Figure S2, rats cumulated body weight gain (A) and total weight gain (B); Table S1, organs weight at the end of the experimentation; Supporting Information B, fatty acid composition of tissues of experiment 1; Table S1, liver, red blood cells, and plasma total fatty acid composition; Table S2, gut, stomach, and heart fatty acid composition of total lipids; Table S3, kidney, brain, and retina fatty acid composition of total lipids; Table S4, spleen, lung, and bone marrow fatty acid composition of total lipids; Table S5, pancreas, epididymal adipose tissue (TAE), and skin fatty acid composition of total lipids; Table S6, subcutaneous adipose tissue (SCAT), muscle, and testis fatty acid composition of total lipids; Supporting Information C, fatty acid composition of tissues of experiment 2; Table S1, liver fatty acids composition of total lipids; Table S2, heart fatty acids composition of total lipids; Table S3, lung fatty acids composition of total lipids; Table S4, spleen fatty acids composition of total lipids; Table S5, red blood cells fatty acids composition of total lipids; Table S6, brain fatty acids composition of phospholipids; Supporting Information D, loadings plot of Figure 3; Figure S1, loadings plot of Figure 3A; Figure S2, loadings plot of Figure 3B (PDF).
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