Gene–PUFA interactions and obesity risk

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Abstract

Although there are indications for modulatory effects of PUFA on associations between SNP and obesity risk, scientific evidence in human subjects is still scarce. The present analyses investigated interaction effects between SNP in candidate genes for obesity and PUFA in erythrocyte membranes on obesity risk. Within the second Bavarian Food Consumption Survey (cross-sectional, population-based), 568 adults provided blood samples. Fatty acid composition of erythrocyte membranes was analysed by means of GC. Genotyping was performed for twenty-one genes, including cytokines, adipokines, neurotransmitters and transcription factors. In addition, plasma IL-6 concentrations were analysed. For the statistical analysis, a logistic regression model assuming additive genetic effects was chosen. About 20 % of the study participants were classified as obese (BMI $\geq 30 \text{ kg/m}^2$). Several significant gene–PUFA interactions were found, indicating regulatory effects of PUFA by gene variants of IL-2, IL-6, IL-18, TNF receptor family member 1B and 21, leptin receptor and adiponectin on obesity risk. After stratification by genotype, the strongest effects were found for rs2069779 (IL-2) and all tested PUFA as well as for rs1800795 (IL-6) and linoleic or arachidonic acid. The obesity risk of minor allele carriers significantly decreased with increasing fatty acid content. The genetic PUFA–IL-6 interaction was also reflected in plasma IL-6 concentrations. If replicated in a prospective study with sufficient statistical power, the results would indicate a beneficial effect of high PUFA supply for a substantial proportion of the population with respect to obesity risk.

Key words: Obesity: Gene–PUFA interactions: Nutritional biomarkers: Bavarian Food Consumption Survey II

Obesity is a multifactorial disorder reflecting complex interactions of genes, environment and lifestyle(11). Industrialisation and modernisation encourage a sedentary lifestyle with concomitantly increased energy intake, resulting in an imbalance of energy intake and expenditure(22) and consequently in gaining surplus weight. About 40–70 % of the variance in the BMI is accounted for by genetic factors as several studies of twins, adoptees and families have shown(14–15). Screenings of candidate regions as well as genome-wide scans have helped to identify SNP that increase the risk of becoming overweight or obese(66). There is considerable evidence that not all fatty acids are obesogenic(71). Approximately 6–2–7–4 % of our average daily energy intake is accounted for by PUFA (n-3 PUFA 0·7–0·9%; n-6 PUFA 5·5–6·5 %)(68). PUFA exert their influence on cardiovascular function, insulin action, plasma lipid levels(69), neuronal development and the immune system inter alia through modulation of eicosanoid (PG and leukotrine) synthesis, activation of orphan nuclear receptor and T-lymphocyte signalling(9,109). They also regulate the transcription and activation of multiple genes(111–13). Hence, PUFA affect several metabolic pathways, and thus may have an impact on the

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Abbreviations: AA, arachidonic acid; LA, linoleic acid; LD, linkage disequilibrium; MET, metabolic equivalents; PA, physical activity.
development of a series of diseases, including obesity\textsuperscript{(13,14)} either directly or through interactive effects with the genetic background. This is supported by the results of different experimental studies\textsuperscript{(15–18)}.

So far, only a few studies have investigated gene–PUFA interactions with respect to obesity risk. We have mainly found human or animal cell studies or dietary intervention studies showing PUFA to have an increasing or decreasing effect on the expression of different candidate genes for obesity. Reseland et al.\textsuperscript{(17)} for example, found n-3 PUFA to decrease leptin gene expression in a dose- and time-dependent manner within a human trophoblast cell line. Alnajjar et al.\textsuperscript{(19)} described the effects of PUFA on the IL production on the basis of a dietary intervention study (Jordan population). A case–control study in the European Prospective Investigation into Cancer and Nutrition by Nieters et al.\textsuperscript{(20)} has obtained indications for possible interactive effects between dietary intake of PUFA and polymorphisms of different obesity candidate gene variants.

Obesity is the result of an interaction between genetic predisposition and the modern obesogenic environment\textsuperscript{(21)}. As it is spreading rather rapidly and presenting a major health problem, it is important to understand inter-individual variance and susceptibility. Thus, the aim of the present study is to investigate potential interactions between PUFA and selected SNP with respect to their impact on obesity risk.

Within the present study, twenty-one candidate genes for obesity, including cytokines, neurotransmitters, transcription factors and adipokines, were selected. Interaction effects with regard to obesity risk were calculated for the fatty acid composition of the erythrocyte membranes, a biomarker for dietary PUFA intake, and SNP located within these gene regions. The PUFA composition of the erythrocyte membranes reflects both PUFA intake and subsequent metabolism of the fatty acids over a period of weeks and months\textsuperscript{(22)}. In order to lend more credence to the statistical results of the genetic IL-6–PUFA analyses, plasma IL-6 concentrations were also considered.

Methods

Study design and population

The second Bavarian Food Consumption Survey is a cross-sectional study, representative of the Bavarian population and designed to investigate dietary and lifestyle habits. German-speaking subjects (n 1050) aged 13–80 years were recruited between September 2002 and June 2003 following a three-step random route sampling procedure. A total of forty-two communities served as sampling points and were stratified by county and community characteristics. With a given start address, a random walk (every third household) was conducted and one random household member who met the inclusion criteria was selected. Information on the subjects’ characteristics, lifestyle as well as health and socio-economic status were collected during a personal computer-assisted face-to-face interview at baseline. The participation rate was 71%. A non-responder analysis was performed. On average, non-responders had – among other characteristics – a higher BMI than the study participants. Within the following two weeks, data of the subjects’ dietary intake and physical activity (PA) were assessed by three 24 h dietary recalls (two weekdays and one weekend day), which were conducted via telephone and by trained interviewers. For the 24 h dietary recalls, the software EPIC-Soft (International Agency for Research on Cancer, Lyon, France) was used\textsuperscript{(23–25)}. The participants had to recall their dietary intake as well as their PA of the previous day. All adult subjects (≥ 18 years) who completed at least one 24 h recall (n 879) were invited to their nearest public health office for blood sampling and standardised anthropometric measurements within 6 weeks after recruitment. Of these subjects, 65% (n 568) accepted this invitation and represented the subgroup on which this evaluation is based on. The study was conducted according to the guidelines laid down in the Declaration of Helsinki, and all procedures involving human subjects were approved by the local ethics committee. Written informed consent was obtained from all study participants.

Anthropometric, dietary and physical activity assessment

Height was measured to the nearest 0·5 cm and weight to the closest 0·5 kg. BMI was calculated as weight/height\(^2\) (kg/m\(^2\)). Subjects were classified according to the definition of the WHO\textsuperscript{(26)} as obese (BMI ≥ 30 kg/m\(^2\)) and non-obese (BMI < 30 kg/m\(^2\)). Hip size was determined as the widest circumference measured over the buttocks and waist measurements were taken midway between the iliac crest and the margin of the lower rip.

The German food composition table BLS (Bundeslebensmittelschüllsel) (version II.3; BgVV, Berlin, Germany) was used to calculate nutrient intake. Data were weighted correspondingly to weekday or weekend day in order to calculate a mean daily intake per subject.

In the telephone interviews, participants were asked to recall their PA of the last 24 h. Standardised questions on type and duration of the PA in the categories of sports, occupation and other strenuous activities during leisure time as well as non-occupational television/personal computer use and duration of sleeping were part of the computer-based interview. Metabolic equivalents (MET) were matched to each activity and the energy expenditure (MET × h) of every individual was estimated\textsuperscript{(27)}.

Blood sampling

Venous blood was drawn, chilled at 4°C and further processed within 3 h. Plasma and buffy coat were separated from erythrocytes by centrifugation (2000 g for 15 min) before being divided into aliquots and stored at −80°C for further analyses.

Fatty acid composition of erythrocyte membranes

Membrane fatty acid analysis was conducted using an aliquot of 0·5 ml erythrocyte suspension. After cell lyses through addition of aqua destillata, the erythrocyte membranes were isolated via centrifugation (20 000 g for 20 min at 4°C) and
the pellet was resuspended with Tris-buffer (11 mM-Tris, 1 mM-Na-EDTA, pH 7.4); the washing procedure was repeated twice before adding 800 μl aqua destillata\(^{(28)}\). Fatty acid extraction was performed using a mixture of chloroform and methanol (2:1, v/v) according to a modification of the method described by Folch et al.\(^{(29)}\). The lipids were extracted twice using a chloroform-methanol mixture with the added antioxidant butylated hydroxytoluene (50 mg/l)\(^{(30)}\). The combination of these extracts was washed with a CaCl\(_2\) solution. The organic phase was collected and evaporated until dry. Resuspension of that extract was done using chloroform, and via trans-esterification with trimethylsulphonium hydroxide, the fatty acid methyl esters were obtained\(^{(31)}\).

A 100 m CP-Sil-88 capillary column (Varian-Chrompack, Darmstadt, Germany), which was installed in an HP 5890 series II gas chromatograph with a flame-ionisation detector (Hewlett Packard, Munich, Germany), was used to identify and separate the different fatty acid methyl esters. Authentic standards (Sigma-Aldrich, Steinheim, Germany) were applied to assure a correct identification and quantification of the fatty acid methyl ester peaks. As a result, the content of twenty-two types of fatty acids was measured and is expressed as a percentage of the total fatty acid methyl esters (\% fatty acid methyl esters) identified\(^{(32)}\). For each sample, data represent the mean of two injections.

Analysis of plasma IL-6

Plasma IL-6 was measured by means of a commercial ELISA kit (Biosource, Brussels, Belgium). The intra- and inter-assay CV were below 7 and 9\%, respectively.

SNP selection

The genetic dataset was developed in 2006 and consists of different candidate genes, which were chosen on the basis of an extensive literature research. SNP covering these genes as well as 100 kb of region 5' and 50 kb of region 3' were selected based on hapmap data (www.hapmap.org; phases 1 and 2) with a minimum minor allele frequency of 0.05 according to the data of dbSNP Build 125 (an SNP database).

For the present analyses, we carefully selected only genes for which an indication of a potential interaction with PUFA was provided by the literature. These include different cytokines and their receptors (IL-2, IL-6, IL-10, IL-18, TNF-α, TNFRSF1A, TNFRSF1B and TNFRSF21 (TNF receptor superfamily 1A, 1B and 21)), neurotransmitters and their receptors (NPY (neuropeptide Y), NPY1R, NPY5R (NPY receptors Y1 and Y5), MC4R (melanocortin 4 receptor), POMC (pro-opiomelanocortin), PPY (pancreatic polypeptide), and PYY (peptide YY)), transcription factors (PPAR\(_γ\) and PPAR\(_γ\)CA1) and adipokines (LEP (leptin), LEPR (leptin receptor), ADIPQ (adiponectin) and RETN (resistin)). For these genes, the genetic dataset holds a total number of 187 SNP, consisting of 180 tagging SNP, eighteen coding SNP and twelve candidate SNP previously reported to be associated with obesity. The median number of SNP per gene locus is 11 with a range of 2–16.

Genotyping and quality control

Genotyping was performed by GoldenGate Genotyping Assay (Illumina, Inc., San Diego, CA, USA) according to the standard protocol of the manufacturer. Additional inclusion criteria were a minor allele frequency of at least 5\% and genotyping call rate of not less than 95\%, leaving 157 SNP in the dataset.

Statistical analyses

The descriptive data are presented as median with 25 and 75\% quantiles for continuous parameters, as the majority of the variables were not normally distributed, or as percentage and absolute frequency for qualitative variables. Comparisons between the groups of obese and non-obese subjects were made by means of either the Mann–Whitney \(U\) test (continuous variables) or the Kruskal–Wallis test (qualitative parameters). Socio-economic status was categorised based on the values of three characteristics on a point scale including educational level, social position and the households’ net income\(^{(33)}\). PA data represent the estimated overall energy expenditure (MET \(\times\) h) of every individual\(^{(27)}\).

Departure from the Hardy–Weinberg equilibrium was tested by means of an exact test\(^{(34)}\). All selected SNP were in the Hardy–Weinberg equilibrium except for rs1061624 (TNFRSF1B), rs16475 (NPY), rs16480 (NPY region 5\_C1A), rs17366743 (ADIPOQ), applying a \(P\) value of 3·09\(10^{-4}\) (corrected for multiple testing) as the significance level.

For comparison of the allele frequencies between obese and non-obese subjects, a Kruskal–Wallis test was used. The main effects of SNP and PUFA as well as their interaction effects are derived from a logistic regression model assuming additive genetic effects, and are presented as OR with the corresponding 95\% CI. For the genetic main-effect models, SNP were introduced as discrete parameters with either three categories (homozygous wild type, heterozygous or homozygous mutant type) or two categories (homozygous wild type and one or more mutant allele carriers), depending on the number of subjects in the third category (homozygous mutant type; minimum of ten subjects). As EPA and DHA were highly correlated, their sum was used in the present analyses. The fatty acid variables (presented as percentage of fatty acid methyl esters) were established as continuous parameters. For the interaction models, both SNP and PUFA were introduced as continuous parameters. A likelihood ratio test was used to compare the models with and without an interaction term and a \(P\) value of \(<0.05\) was regarded as nominally statistically significant. These \(P\) values are denoted as \(P_{\text{int}}\). The interaction models were adjusted for sex, age, PA and socio-economic status and significant SNP–PUFA interactions were analysed further. The effect of each PUFA with respect to obesity risk was estimated within the gene strata (homozygous wild type and one or more mutant allele carriers) of the SNP using logistic regression models adjusted for sex, age, PA and socio-economic status. As the analyses were done within the SNP strata and the number of people carrying two minor alleles was often quite small, two categories were used for sample size reasons. We tried different
procedures, *inter alia* by Bonferroni, Holm, Hochberg, Sidak or Benjamini and Hochberg, to correct the present results for multiple testing, but neither left us with any significant result. Therefore, all *P* values reported in the present study are uncorrected and only nominally significant at α 5%. The SNP were tested for pairwise linkage disequilibrium (LD). If SNP were in high LD (*r*² ≥ 0.7), one of them was selected as representative for the LD block. We chose rs1800795 to represent the IL-6 LD block with rs1800797 (*r*² 0.935) and IL-18 SNP rs3882891 to represent rs1946519 (*r*² 0.77).

As five values of plasma IL-6 were declared as outliers (greater than mean plus five times the standard deviation), they were excluded from the analyses. While plasma IL-6 levels ranged between 0.25 and 11.64 pg/ml, the levels of those five outliers were substantially higher (51.5, 61.8, 68.92, 93.62 and 620.6 pg/ml). The reason for those high values could neither be clarified nor was a re-analysis of the samples possible. To account for skewness, the parameter was log-transformed. Plasma IL-6 is presented as geometric means and 95% CI. All statistical analyses were performed with R software version 2.9.0 (R Development Core Team, 2009; http://www.r-project.org/).

**Results**

Table 1 summarises the characteristics of the study population. Obese subjects had a median BMI of 33.04 (25–75% quantiles 31.19–36.12) kg/m² and their median age exceeded that of the non-obese subjects (median BMI of 24.83 (25–75% quantiles 22.55–27.20) kg/m²) by 12 years. The median hip and waist circumference of obese and non-obese participants differed by 12.5 and 22.5 cm and the plasma IL-6 level of obese subjects was also elevated. There was no significant difference

| Table 1. Characteristics of obese (BMI ≥ 30 kg/m²) and non-obese (BMI < 30 kg/m²) subjects in a subsample of the Bavarian Food Consumption Survey II  
 (Medians, 25–75% quartiles, percentages or number of absolute frequencies) |
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<td>Non-obese subjects (79.93 %)</td>
<td><em>P</em></td>
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<td>Sex</td>
<td>%</td>
<td>Absolute n</td>
<td>%</td>
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<tr>
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<td>51</td>
<td>42.29</td>
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<tr>
<td>Female</td>
<td>55.26</td>
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<td>57.71</td>
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<tr>
<td>18 to &lt;30</td>
<td>5.26</td>
<td>6</td>
<td>11.45</td>
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<td>30 to &lt;40</td>
<td>12.28</td>
<td>14</td>
<td>25.55</td>
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<td>14.91</td>
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<td>23.79</td>
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<td>23.79</td>
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<td>Lower class</td>
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<td>33.26</td>
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<td>6</td>
<td>10.79</td>
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<tr>
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<td>23</td>
<td>12.33</td>
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<td>Age (years)</td>
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<td>Weight (kg)</td>
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<td>Height (cm)</td>
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<td>159.00–173.00</td>
<td>168.00</td>
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<td>BMI (kg/m²)</td>
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<td>Waist (cm)</td>
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<td>106.00–120.50</td>
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<td>Waist:hip ratio</td>
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<td>Physical activity (MET x h/d)</td>
<td>37.24</td>
<td>34.45–40.20</td>
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<td>Erythrocyte membranes</td>
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<tr>
<td>Linoleic acid (% FAME in erythrocytes)</td>
<td>10.98</td>
<td>9.40–12.12</td>
<td>11.50</td>
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<td>Arachidonic acid (% FAME in erythrocytes)</td>
<td>15.86</td>
<td>9.16–17.63</td>
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<tr>
<td>EPA + DHA (% FAME in erythrocytes)</td>
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<td>1.79–7.56</td>
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<td>EPA + DHA (% E)</td>
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<td>0.03–0.15</td>
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<tr>
<td>Plasma concentration‡</td>
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<tr>
<td>IL-6 (pg/ml)</td>
<td>2.37</td>
<td>2.12–2.65</td>
<td>1.38</td>
</tr>
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</table>

*MET, metabolic equivalents; % FAME in erythrocytes, fatty acids in percentage of total fatty acid methyl esters in erythrocyte membranes; % E, percentage of energy.
*Kruskal–Wallis test.
†Mann–Whitney U test.
‡Values are geometric means and 95% CI.
in sex distribution, PA or dietary fatty acid intake between the
two groups. The linoleic acid (LA), arachidonic acid (AA), and
EPA+DHA compositions of erythrocyte membranes were
higher for non-obese (r. obese) participants but only the
difference in LA content reached statistical significance.

The following results are not corrected for multiple testing
and therefore are only nominally significant at α 5 %. Table
S1 of the supplementary material (available online at http://
www.journals.cambridge.org/bjn) shows the distribution of
the alleles within the two groups of obese and non-obese sub-
jects, the P-values of the Kruskal–Wallis test and the main
effects of all SNP analysed on the risk of obesity. Risk esti-
mates were calculated for models with either three categories
(homozygous wild type, heterozygous and homozygous
mutant type) or two categories (homozygous wild type and
one or more mutant allele carriers). The P value of the con-
tinuous model is also given as a P trend. A nominal P value
<0·05 was reached by rs4779714 and rs12700386 (IL-6
region 5'), rs2069849 (IL-6), rs1061628 (TNFRSF1B) and
rs1116656 (LEP region 3'). The crude main effects for LA
(OR 0·90, 95 % CI 0·82, 0·99, AA (OR 0·97, 95 % CI 0·93,
1·01) and EPA+DHA (OR 0·96, 95 % CI 0·89, 1·05) in erythro-
cyes showed an indication of an inverse association with
obesity, but only for LA, the statistical significance was
reached.

SNP for which the interaction term with PUFA reached nomi-
nal statistical significance (α 5 %) were stratified by genotype
(two categories: homozygote wild type and one or more
mutant allele carriers), and risk estimates were calculated for
the corresponding PUFA within these strata (Tables 2–4).

Concerning cytokine genes, we found several SNP–PUFA
interactions in relation to obesity risk. For IL-2, the interaction
term between rs2069779 and all three PUFA as well as
between rs2069772, rs4833248 (IL-2 region 5') and LA were
statistically significant. Each PUFA associated with rs2069779
had a decreasing effect on obesity risk for minor allele carriers
(Fig. 1). In this gene stratum, the risk decreased for each mol %
increase in erythrocyte membrane-bound PUFA with OR of
0·63 (LA), 0·83 (AA) and 0·66 (EPA+DHA). For rs4833248 as
well as rs2069762, homozygous wild-type carriers benefited
from increased LA content in erythrocyte membranes. Ana-
lyses for IL-6 revealed three SNP: rs1800795; rs10242595;
rs2069861. The first two showed a significant interaction
effect with LA with P_int = 0·0341 and 0·0315, respectively.
For each percentage increase in erythrocyte membrane-bound
LA, obesity risk decreased in carriers of at least one minor
allele of rs1800795 with an OR of 0·86 (95 % CI 0·76, 0·96); a
similar effect was found for the rs1800795–AA interaction
(Fig. 2). In the case of rs10242595, the relative risk decreased
for carriers of the homozygous wild-type alleles with increas-
ing LA content (OR 0·81, 95 % CI 0·71, 0·93). The interaction
effect of rs2069861 and EPA+DHA was significant as well.
Here, the obesity risk decreased with each mol % increase
of erythrocyte membrane-bound EPA+DHA for minor allele
carriers (OR 0·74, 95 % CI 0·57, 0·94). For IL-18, one SNP
(rss882891) interacted significantly with LA when analysing
obesity risk. Minor allele carriers of this SNP had a reduced
obesity risk with increasing LA content in erythrocyte mem-
branes (OR 0·84, 95 % CI 0·75, 0·94, P_int = 0·0203).

We also obtained evidence for SNP–EPA+DHA interaction
effects of the TNFRSF1B gene. Subjects carrying at least one
minor allele of rs3766730 (OR 0·86, 95 % CI 0·75, 0·98,
P_int = 0·0225) or rs2275416 (OR 0·88, 95 % CI 0·78, 0·99,
P_int = 0·0455) had a lower obesity risk than homozygous
wild-type carriers. In relation to obesity risk, two SNP
of TNFRSF21 were shown to interact with EPA+DHA;
s9381530 (region 3') with an OR of 0·79 (95 % CI 0·68, 0·91,
P_int = 0·0065) and rs2236039 with an OR of 0·87 (95 % CI
0·78, 0·96, P_int = 0·0482). In each case, obesity risk decreased
with increasing PUFA content in subjects carrying two
major alleles.

Among the selected adipokine genes, few significant results
were obtained. The interaction terms of two ADIPOQ SNP
(rs1063539 and rs2241766) and membrane EPA+DHA
content were significantly associated with obesity risk. In
homozygous wild-type carriers, obesity risk decreased with

<table>
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<th>Genes</th>
<th>SNP</th>
<th>Allele</th>
<th>Non-obese (%)</th>
<th>Obese (%)</th>
<th>OR</th>
<th>95 % CI*</th>
<th>P_int‡</th>
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<tr>
<td>IL-2</td>
<td>rs2069779</td>
<td>CC</td>
<td>85·8</td>
<td>86·7</td>
<td>0·93</td>
<td>0·85, 1·03</td>
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<td>≥ 1 T</td>
<td>14·2</td>
<td>13·3</td>
<td>0·63</td>
<td>0·42, 0·87</td>
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* Adjusted for age, sex, physical activity and socio-economic status.
† Main effect: OR 0·90, 95 % CI 0·82, 0·99.
‡ P value of the likelihood-ratio test (adjusted continuous interaction model).
increasing EPA+DHA content (Table 4). Concerning a leptin receptor gene polymorphism (rs1805096), an inverse association with obesity risk existed in carriers of the minor allele, with an increasing erythrocyte membrane content of LA (OR \( \text{OR} = 0.95, 95\% \text{CI} 0.89, 1.02 \); Table 2) or AA (OR \( \text{OR} = 0.97, 95\% \text{CI} 0.94, 1.01 \)).

Table 5 gives the geometric means and the 95% CI of plasma IL-6 stratified by IL-6 SNP rs1800795, rs10242595 and rs2069861 (homozygote wild type and one or more mutant alleles) and tertiles of LA, AA and EPA+DHA for obese and non-obese subjects. The plasma IL-6 concentrations were generally higher in obese subjects compared with non-obese subjects (see also Table 1) and decreased with increasing PUFA content in erythrocyte membranes (tertiles). These results lend credit to the identified IL-6 SNP–PUFA interactions and obesity risk derived from the statistical models.

**Discussion**

The present study aimed to investigate the additional effect on obesity brought on by the interaction of selected genetic variants and PUFA content of erythrocyte membranes. Out of the four different groups of genes, including cytokines, adipokines, neurotransmitters and transcription factors, we obtained significant interaction effects between the SNP of IL-2, IL-6, IL-18, TNFRSF1B, TNFRSF21, LEPR or ADIPOQ and PUFA content in erythrocyte membranes. We found a reduced obesity risk for minor allele carriers of most variants with high PUFA content in erythrocyte membranes, except for the SNP of TNFRSF21, ADIPOQ, rs2069762 (IL-2), rs4833248 (IL-2 region 5′) and rs10242595 (IL-6 region 3′). With the latter genes, subjects, homozygote for the major allele, benefited from an increased PUFA content of erythrocyte membranes. In the case of IL-6, the analysed plasma IL-6 protein concentration supports the statistical findings.

Obesity has been described as a state of chronic low-grade inflammation\(^{35,36}\). Thus, polymorphisms in different cytokines were included in the analyses of gene–PUFA interactions. Cytokines are a group of modulatory proteins which respond to various stimuli, thereby activating second messengers and signal transduction pathways within the cells.\(^{37}\) Several cell studies and also dietary intervention studies have shown a reduced production of IL-2 in response to PUFA\(^{16,18,19,38}\); however, the exact mechanism behind this phenomenon still remains unclear\(^{39,40}\). In the present study, obesity risk decreased with increasing PUFA content

<table>
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<th>Genes</th>
<th>SNP</th>
<th>Allele</th>
<th>Non-obese (%)</th>
<th>Obese (%)</th>
<th>OR</th>
<th>95% CI*</th>
<th>( P_{\text{int}} )‡</th>
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<td>0·95</td>
<td>0·91, 1·00</td>
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* Adjusted for age, sex, physical activity and socio-economic status.
‡ \( P \) value of the likelihood-ratio test (adjusted continuous interaction model).

**Table 3.** Significant SNP–arachidonic acid interactions on obesity risk, showing the adjusted* relative risk of obesity per 1 mol% increase of arachidonic acid† in erythrocyte membranes by allelic variants

(Odds ratios and 95% confidence intervals)

<table>
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<th>Genes</th>
<th>SNP</th>
<th>Allele</th>
<th>Non-obese (%)</th>
<th>Obese (%)</th>
<th>OR</th>
<th>95% CI*</th>
<th>( P_{\text{int}} )‡</th>
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<td>IL-2</td>
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<td>( \geq 1 \text{T} )</td>
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* Adjusted for age, sex, physical activity and socio-economic status.
‡ \( P \) value of the likelihood-ratio test (adjusted continuous interaction model).

**Table 4.** Significant SNP–EPA+DHA interactions on obesity risk, showing the adjusted* relative risk of obesity per 1 mol% increase of EPA+DHA† in erythrocyte membranes by allelic variants

(Odds ratios and 95% confidence intervals)

<table>
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<th>Genes</th>
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<th>Non-obese (%)</th>
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* Adjusted for age, sex, physical activity and socio-economic status.
‡ \( P \) value of the likelihood-ratio test (adjusted continuous interaction model).
Gene–PUFA interactions and obesity risk

been associated with excess adiposity (51). Our IL-18 variant higher levels of IL-18 than lean subjects (50), and IL-18 has

the present study is indicative towards the importance of this SNP for the development of obesity. Variations in the IL-18 gene have been associated with IL-18 plasma concentrations and measures of obesity (49). Obese subjects show higher levels of IL-18 than lean subjects (50), and IL-18 has been associated with excess adiposity (51). Our IL-18 variant is in complete LD with rs5744292, an IL-18 SNP whose minor allele has been reported to be associated with lower circulating IL-18 levels and lower mRNA expression in immortalised lymphocytes (52,53). Furthermore, a suppressing effect of PGE₂ on the expression of IL-18 has been shown in cell studies (54). The finding of an IL-18–PUFA interaction in the present study fits well with these data. Even though the

of the erythrocyte membranes, and thus confirms the expected direction. PUFA eicosanoid derivates, such as PGE₂, are involved in the modulation of the intensity and duration of inflammatory processes and suppress the production of IL-6 (42). Adipose tissue in human subjects releases IL-6 and serum levels are inversely correlated with body fat mass (43). Himmerich et al. (42) confirmed this relation for the present population. IL-6 gene transcription was found to be influenced in vitro by the rs1800795 polymorphism within the promoter region (43). The G allele of this SNP was described as to be more common in lean subjects (44); additionally, a lower BMR was measured in subjects with the CC genotype (45), which might eventually predispose to weight gain; this hypothesis has been supported by some studies but not confirmed in two meta-analyses of the association of this SNP with BMI (46,47). Our findings on SNP–PUFA interactions confirmed in two meta-analyses of the association of this SNP with BMI (46,47). Estimates are adjusted for age, sex, physical activity and socio-economic status. Padj, P value of the likelihood-ratio test comparing models with and without an interaction term.

The finding of an IL-18–PUFA interaction in the present study fits well with these data. Even though the production of TNF-α by monocytes and macrophages is also suppressed through PGE₂ (10), no significant interactions could be determined within the present study. However, we found evidence for significant interaction effects for its receptors, TNFRSF1B and TNFRSF21.

Overall, all identified (significant) interactions between cytokine SNP and PUFA indicate an inverse association with obesity risk for minor allele carriers, with increasing PUFA content in erythrocyte membranes, except for SNP of TNFRSF21, rs4833248 (IL-2 region 5') and rs10242595 (IL-6 region 3').

Besides its role for lipid storage, adipose tissue functions as an endocrine organ, regulating metabolism and different vital functions related, among others, to inflammation (55,56). Thus, different adipokines have been included in the present analyses. Adiponectin is exclusively secreted by adipose tissue and serum levels are inversely correlated with body fat mass (57). The mRNA expression is reduced in obese individuals (58). Serum adiponectin levels are highly heritable (approximately 50%) and are linked to the ADIPOQ gene locus (59–61). Different cell and dietary intervention studies found EPA + DHA to stimulate the expression of ADIPOQ and to increase plasma adiponectin levels (62–64); however, findings differ (65). EPA + DHA might possibly up-regulate ADIPOQ by acting through PPARy, affecting the ADIPOQ promoter (15). The present results are in line with these findings and show a significantly decreased obesity risk for carriers of two major alleles of rs2241766 or rs1063539, with increasing EPA + DHA concentrations in erythrocyte membranes. Different animal, human and cell studies have shown an inverse effect of PUFA on the LEP mRNA expression (17,66). The present analyses resulted in one significant interaction for a variant of the leptin receptor gene, which is in line with these findings.

We see two major mechanisms of how PUFA may in conjunction with genetic variants affect obesity risk: either via
Plasma IL-6 concentrations (pg/ml) by tertiles of linoleic acid, arachidonic acid and EPA þ DHA in erythrocyte membranes by allelic variants of IL-6 SNP, rs1800795, rs10242595 and rs2069861 (data not shown). A high erythrocyte membrane content of LA or AA or EPA þ DHA thereby did either show no association or an inverse association with obesity risk in each of the SNP strata. Therefore, we conclude that PUFA exert their effects rather via modification of gene transcription than through metabolites derived during eicosanoid synthesis, since the latter would have led to differential effects of n-3 and n-6 PUFA. In mutually adjusted analyses, we have also not received any indication for changes of effects of n-3 PUFA-adjusted n-6 PUFA and vice versa. It is important to mention that the frequency of these SNP for which we observed interactive effects with PUFA is fairly high, except for IL-2 SNP (rs2069779). This implicates that a substantial part of the population would benefit from a high PUFA intake with respect to obesity risk.

A major limitation of the present study is obviously the small sample size. Studies of genetic associations with complex diseases need thousands of cases and controls (67); however, this requirement is not easily fulfilled with respect to the costly fatty acid analyses. The small number of cases and controls and consequently the limited statistical power strongly argue for a careful interpretation of the results and a replication in a second, larger and independent study. Because of the small sample size and the resulting limited power, we did not correct for multiple testing. No correction method for multiple testing left us with any significant result. Therefore, all reported P values are not corrected and are only nominally significant at α 5 %. The observational nature of the study does not allow for interpreting causal associations, and we cannot rule out the possibility of reverse causation. However, we controlled for potential confounding by adjusting for sex, age, PA and socio-economic status.

The major strength of the present study is the use of erythrocyte membranes to assess biologically available PUFA at the cellular level and its association with genetic variants influencing the risk of obesity. With the utilisation of biomarkers as an objective metabolic correlate of dietary PUFA intake, misclassifications can be largely avoided. A further strength of the present study is its population-based design aiming at representativeness for the adult Bavarian population and the strict quality control in the analyses.

Acknowledgements

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contributions were as follows: C. J. performed the statistical analyses and drafted the manuscript. S. K., A. N., H. H. and M. A. K. were responsible for SNP selection and genotyping; S. K. also wrote these sections of the manuscript. H. S. contributed to the data collection. H.-E. W., G. W. and S. L. provided critical revision. C. G. gave statistical advice. J. L. was responsible for the study design, fatty acid data acquisition as well as supervision and together with G. W. for funding. All authors contributed to the interpretation and discussion of the results and read and approved the final version of the manuscript. The present study represents original work that has not been published previously and the authors declare that there are no conflicting interests.

References


