Circulating fatty acids as biomarkers of dairy fat intake: data from the lifelines biobank and cohort study

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ABSTRACT

Background: C14:0, C15:0, C17:0 and trans-C16:1(n-7) are often used as biomarkers for dairy fat intake. Trans-C18:1(n-7) and CLA, two fatty acids which are also present in dairy, have hardly been explored. We investigated whether trans-C18:1(n-7) and CLA can enrich the existing biomarker portfolio.

Methods: Data were obtained from Lifelines (n = 769). Dairy fat intake was determined by FFQ. Fatty acids were measured in fasting plasma triglycerides (TG), phospholipids (PL) and cholesterol esters (CE).

Results: Median (25th–75th percentile) intakes of dairy and dairy fat were 322(209–447) and 12.3(8.4–17.4) g/d respectively. A pilot study showed that trans-C18:1(n-7) and CLA were only detectable in TG and PL. Of the established markers, TG C15:0 was most strongly associated with dairy fat intake (standardized β (std.β) = 0.286, R² = 0.111). Of the less established markers, TG trans-C18:1(n-7) was most strongly associated with dairy fat intake (std.β = 0.292, R² = 0.115), followed by PL CLA (std.β = 0.272, R² = 0.103) and PL trans-C18:1(n-7) (std.β = 0.269, R² = 0.099). In TG, a combination of C15:0 and trans-C18:1(n-7) performed best (R² = 0.128). In PL, a combination of C14:0, C15:0, trans-C18:1(n-7) and CLA performed best (R² = 0.143).

Conclusion: Trans-C18:1(n-7) and CLA can be used as biomarkers of dairy fat intake. Additionally, combining established with less established markers allowed even stronger predictions for dairy fat intake.

Introduction

Dairy consumption is part of the food culture in many countries. Worldwide intake of dairy products is high, particularly in developed countries such as Canada, North America and parts of the European Union. For instance, in the European Union and in the USA, the average per capita milk consumption is 1.01 and 0.91 servings per day respectively (Dougkas et al. 2011). While dietary recommendations tend to set an upper limit to certain types of dairy products as a result of the high saturated fat or sodium content, dairy intake is generally considered healthy, particularly because of the numerous essential nutrients it contains, including calcium, proteins, phosphorus, magnesium, and several vitamins and minerals (Haug et al. 2007). It has been observed that high dairy intake is associated with a reduced risk for hypertension (Soedamah-Muthu et al. 2012). Furthermore, dairy has been suggested to have a positive influence on bone health by increasing bone mineral density (Heaney 2009, Sahni et al. 2013). There is also increasing evidence that fat from dairy products (as reflected by circulating dairy fat biomarkers) has a positive effect on various cardiovascular risk factors such as BMI, fasting glucose and insulin sensitivity (Smedman et al. 1999, Mozaffarian et al. 2010, 2013). Moreover, prospective studies have reported lower risk of stroke and coronary heart disease with higher levels of circulating dairy fat biomarkers (Waren sj et al. 2009, de Oliveira Otto et al. 2013). Additionally, high circulating concentrations of dairy fat biomarkers were found to be associated with reduced risk of development of type 2 diabetes as was evidenced in several longitudinal observational studies (Krachler et al. 2008, Forouhi et al. 2014, Santaren et al. 2014, Yakoob et al. 2016).

Dairy fat consists of more than 400 different fatty acids (Mansson 2008). The fatty acids in dairy fat derive either directly from the animal’s diet, or indirectly from ruminal fermentation. The fatty acid composition of plasma, serum,
Fatty acids, such as trans-C18:1(n-7), are known to partially reflect the fatty acid composition of the diet. Several fatty acids are to a large extent unique for dairy fat intake. They are present in measurable amounts in the circulation and tissue of humans (Smedman et al. 1999). Myristic acid (C14:0), pentadecanoic acid (C15:0), heptadecanoic acid (C17:0) and trans-palmitoleic acid (Trans-C16:1(n-7)) are commonly used dairy fat biomarkers (Wolk et al., 2001, Rosell et al. 2004, Biong et al., 2006, Sun et al., 2007, Yakoob et al., 2014, Warenosj et al., 2015, Lund-Blix et al., 2016, Yakoob et al. 2016). Particularly the odd-chain fatty acids C15:0 and C17:0 (Brevik et al., 2005, Biong et al., 2006, Sun et al., 2007, Yakoob et al., 2014, Warenosj et al., 2015, Yakoob et al., 2016, Lund-Blix et al., 2016), and the natural ruminant trans fat trans-C16:1(n-7) are often considered as fatty acids that mainly originate from dairy fat (Sun et al., 2007, Yakoob et al., 2014, Yakoob et al., 2016), and because these fatty acids are synthesized by the bacterial flora in the rumen of the animal and cannot be synthesized in the human body (Jenkins et al., 2015). Correlations of C15:0, C17:0 and trans-C16:1(n-7) with dairy fat intake are variable. Papers that reported on the correlation between circulating dairy fat biomarkers in plasma and dairy fat intake showed correlations of 0.10–0.53 with C15:0 (Smedman et al., 1999, Wolk et al., 2001, Rosell et al., 2004, Sun et al., 2007, Yakoob et al., 2014, Warenosj et al., 2015, Lund-Blix et al., 2016, Yakoob et al., 2016), of 0.16–0.36 with C17:0 (Wolk et al., 2001, Rosell et al., 2004, Sun et al., 2007, Yakoob et al., 2014, Warenosj et al., 2015, Yakoob et al., 2016), and of 0.13–0.30 with Trans-C16:1(n-7) (Sun et al., 2007, Yakoob et al., 2014, Yakoob et al. 2016). Despite the specific origin of these fatty acids, the correlations are relatively low and interest in other potential dairy fat biomarkers is therefore growing.

Trans fatty acids which are also thought to mainly originate from rumen derived products are vaccenic acid (trans-C18:1(n-7)) and rumenic acid, a conjugated linoleic acid with the lipid name cis-9, trans-11 conjugated linoleic acid (CLA). The trans fatty acids are synthesized via microbial fermentation of C18 unsaturated fatty acid in the rumen of the animal (Lock and Bauman 2004, Gebauer et al. 2011a). They can be found in several ruminant derived foods (Gebauer et al., 2006, Sun et al., 2007, Thiebaut et al. 2009, Gebauer et al. 2011b, Aslibekyan et al., 2012, de Oliveira Otto et al., 2013, Yakoob et al. 2014, Yakoob et al. 2016). Despite their occurrence in dairy products, vaccenic acid and CLA are hardly explored as potential dairy fat biomarkers. Current studies on the association of trans fats with dairy intake use a cluster of trans-C18:1, which is actually a combination of mainly rumen-derived trans-C18:1(n-7) and industrially (trans-C18:1(n-9) produced trans fats, possibly leading to biased associations (Yu et al., 2012, Zong et al. 2014). It is therefore of interest to determine the association of dairy fat intake and the genuine rumen derived Trans-C18:1(n-7).

The main objective of this study was to investigate the fatty acids trans-C18:1(n-7) and CLA as potential biomarkers of dairy fat intake and to compare them with more commonly used dairy fat biomarkers. Secondly, we investigated whether a combination of biomarkers provides a better predictor for dairy fat intake compared to single biomarkers. For this we developed and validated a model for the prediction of dairy fat intake in the general population. We used data from the LifeLines Biobank and Cohort Study, a large observational cohort study of initially healthy participants living in the Northern part of the Netherlands.

Clinical significance

- Despite their occurrence in dairy products, trans-C18:1(n-7) and CLA are hardly explored as potential dairy fat biomarkers in the general population.
- The present study confirmed the commonly used biomarkers C14:0, C15:0, C17:0 and trans-C16:1(n-7) and identified the biomarkers trans-C18:1(n-7) and CLA as additional dairy fat biomarkers. A combination of the commonly used biomarkers with less established biomarkers allowed even stronger predictions for dairy fat intake.

Materials and methods

Study design and subjects

The LifeLines Cohort Study is a large observational population-based cohort study and Biobank that examines the health and health-related behaviours of more than 167,000 persons (Stolk et al., 2008). The participants were recruited from the three Northern provinces of the Netherlands between 2006 and 2013. The first group of 25–50 years old participants was recruited via general practitioners. Participants could indicate whether family members were interested as well. In addition, individuals who were interested in the study had the possibility to register via an online self-registration. Individuals with insufficient knowledge of the Dutch language, with severe psychiatric or physical illness, and those with limited life expectancy (<5 years) were excluded from the study. Participants (>18 years old) completed several questionnaires, including topics such as occurrence of diseases, general health, medication use, diet, physical activity and personality. Participants were invited to the Lifelines Research sites for a comprehensive health assessment. A more detailed description of the Lifelines Cohort study can be found elsewhere (Klijs et al., 2015, Scholtens et al., 2015). For the present study, we used a subset of the cross-sectional data. 864 participants were randomly selected from the Lifelines biobank. Additionally, 776 of the 864 participants had filled in a food frequency questionnaire (FFQ). Cases with missing data on either FFQ or circulating fatty acids were removed before analysis leaving 769 participants in the lifelines cohort with complete data (Figure 1). All participants provided written consent. The Lifelines Cohort Study was conducted according to the principles of the Declaration of Helsinki and approved by the Medical ethical committee of the University Medical Center Groningen, The Netherlands.
**Dietary assessment**

To assess dietary intake in the Lifelines Cohort, a 110-item semi-quantitative baseline food frequency questionnaire (FFQ) assessing food items over the previous month was developed and validated by the Wageningen University using the Dutch FFQTOOL™, in which food items were selected based on the Dutch National Food Consumption Survey of 1997/1998 (Voedingscentrum 1998). Seven answer categories were used to assess consumption frequency,
ranging from ‘not this month’ to ‘6–7 days a week’. Portion size was estimated by fixed portion sizes (e.g. slices of bread, pieces of fruit) and commonly used household measures (e.g. cups, spoons). Energy and macronutrient intake, including dairy consumption, was estimated from the FFQ data by using the Dutch food composition database of 2011 (NEVO) (Nederlands Voedingsstoffenbestand 2011). For this study, ‘total dairy intake’ and ‘total dairy fat intake’ (cheese, milk, buttermilk, yoghurt, sweetened yoghurt drinks, custard, curd cheese, ice cream, whipped cream and porridge) were calculated. Butter was not included in the ‘total dairy intake’ and ‘total dairy fat intake’ calculation.

Clinical measurements

Anthropometric measurements (weight, height, and waist circumference) and blood pressure were measured by well-trained staff. The anthropometric measurements were measured without shoes. Body weight was measured to the nearest 0.1 kg. Height was measured to the nearest 0.5 cm with a stadiometer placing their heels against the rod and the head in Frankfort Plane position. Waist circumference was measured to the nearest 0.5 cm in standing position with a tape measure all around the body, at the level midway between the lower rib margin and the iliac crest. BMI was calculated as weight (kg) divided by height squared (m²). Systolic and diastolic blood pressures were measured 10 times during a period of 10 min, using an automated Dinamap Monitor (GE Healthcare, Freiburg, Germany). The average of the final three readings was used for each blood pressure parameter.

Data on smoking and general health

Information about smoking and general health was collected from the self-administered questionnaire. Subjects were classified as nonsmokers, former smokers or current smokers. Information on history and prevalence of hypertension, hypercholesterolaemia and diabetes type 2 were collected in the same questionnaire.

Biochemical measurements

Blood samples were collected in the fasting state, between 8.00 and 10.00 a.m, and transported on ice to the Central Lifelines Laboratory in the University Medical Center Groningen. Serum levels of total and HDL cholesterol were measured using an enzymatic colorimetric method, triglycerides using a colorimetric UV method, and LDL cholesterol using an enzymatic method, all on a Roche Modular P chemistry analyzer (Roche, Basel, Switzerland). Fasting blood glucose was measured using a hexokinase method. HbA1c was determined in whole blood (EDTA-anticoagulated) by means of turbid metric inhibition immunoassay on a Cobas Integra 800 CTS analyzer (Roche Diagnostics Nederland BV, Almere, The Netherlands). The hs-CRP was determined by nephelometry (BN II system Siemens, Marburg, Germany). Serum creatinine was measured on a Roche Modular P chemistry analyzer (Roche, Basel, Switzerland).

Fatty acid analyses

EDTA-plasma samples were collected at baseline and stored frozen at −80°C until use for assessment of fatty acid profiles. Analyses of fatty acids were performed in the Department of Laboratory Medicine of the University Medical Center Groningen, The Netherlands using the methodology as described by Hoving et al. (1988). In short, total lipids were extracted by the method of Folch et al., using 6 ml of chloroform-methanol (2:1) and a 200 µl EDTA-plasma sample (Folch et al. 1957). After that, a shortened version of the method of Kaluzny et al. was used to isolate plasma cholesterol esters (CE), triglycerides (TG) and phospholipids (PL), using aminopropyl SPE columns (Isolute, Biotage) (Kaluzny et al. 1985). Fatty acids were transmethylated with methanolic-HCl into fatty acid methyl esters (FAME). The samples were extracted with hexane and eventually redissolved into 100 µl hexane. Internal standards for the quantification of fatty acids in CE (100 µl of a solution of 50.1 mg C17:0/100 ml chloroform-methanol, 2:1 v/v) and TG (100 µl of a solution of 19.9 mg of C19:0/100 ml chloroform-methanol, 2:1 v/v), both obtained from Sigma-Aldrich (Zwijndrecht, The Netherlands), were added before isolation of lipid classes. An internal standard for the quantification of fatty acids in PL (100 µl of a solution of 50.0 mg free fatty acid 19:0/100 ml methanol), obtained from Larodan (Solna, Sweden), was added after isolation of lipid classes. 100 µl Butylated Hydroxytoluene (1 g/100 ml methanol) from Sigma-Aldrich (Zwijndrecht, The Netherlands) was added to prevent fatty acid oxidation.

Aliquots of 2 µl were injected into an Agilent model 6890 gas chromatography equipped with a 200 m × 0.25 mm polar column (CP Select for FAME) and detected with an Agilent 7683 series flame ionization detector. The 200 m polar column is specifically developed to allow for better separation between trans C18:1 fatty acids (Peene 2011). FAMEs were identified by comparing retention times with those of known standards (Supelco 37 component FAME mix (Sigma-Aldrich)). Fatty acid compositions were expressed into mol%.

Firstly, circulating fatty acid compositions were analyzed in 96 samples to investigate whether the potential dairy fat biomarkers, and especially trans-C18:1(n-7) and CLA, could be detected in the three fractions (plasma CE, TG and PL). Subsequently, we proceeded with analyses in those fractions in which all potential dairy fat biomarkers could be detected. The precision of the potential dairy fat biomarkers was calculated by measuring 10 quality-control samples. Potential dairy fat biomarkers had an average variation coefficient of ≤10%. In TG, the CVs for the potential new biomarkers Trans-C18:1(n-7) and CLA were 8.5% and 7.9%, respectively, and in PL, the CVs for Trans-C18:1(n-7) and CLA were 9.3% and 9.6% respectively.

Statistical analyses

All analyses were performed using IBM SPSS Statistics, version 22.0 for Windows software (IBM, Armonk, NY, USA) and STATATA/MP version 13.1 for Windows software (Statacorp, College Station, Texas, USA). In total, circulating fatty acids in
plasma TG and PL were measured in 864 participants, whereof 776 participants had also completed an FFQ. Cases with missing data on either FFQ or circulating fatty acids were removed before analysis leaving 769 participants in the lifelines cohort with complete data. Baseline data are presented for the total population and separately for males and females to check for potential differences between males and females. Data are presented as N (%), mean ± SD or as in case of non-normally distributed data as median (25th–75th percentile). Differences in baseline data between males and females were tested with an independent t-test (continuous data), Mann-Whitney U test (non-normally distributed data) or Chi-square test (categorical data). For the main analyses, we used energy-adjusted total dairy fat intake. Energy-adjusted intakes were generated by adjustment for total energy intake using the residual method (Willett et al. 1997). Because men usually have a higher energy and macronutrient intake compared to women, adjustment for energy intake was performed for men and women separately. Energy-adjusted intakes are expressed in grams/day. Linear regression models were used to examine the association between circulating fatty acids and total dairy fat intake. We also examined the association of circulating fatty acids with (semi-) skimmed dairy products, full fat dairy products, fermented dairy products, milk and cheese. As sensitivity analyses, we tested whether removal of outliers for energy intake (participants with energy intakes lower or higher than 3 SDs from the mean value of energy intake) influenced the associations between dairy fat biomarkers and energy-adjusted dairy fat intake. We also tested for potential effect modification of the associations of circulating fatty acid biomarkers with total dairy fat intake by sex. Furthermore, in subsidiary analyses, the associations were adjusted for age, sex, BMI, total meat intake, and the industrialized trans fatty acid trans-C18:1(n-9) (Elaicid acid). Skewed data were transformed before analyses (square root or log-transformed). Multivariate backwards regression analyses were carried out with all potential dairy fat biomarkers in TG, PL and a combination of both to identify the best combination of markers to predict energy-adjusted dairy fat intake in our population (cut-off for removal $\rho < 0.10$). We compared the performance of the models against each other by comparing the explained variance ($R^2$). Bland-Altman plots were created to indicate the agreement of observed (FFQ-derived data) compared to the estimated models (based on circulating fatty acids). The mean of the observed and estimated models were plotted against the difference between the observed and estimated models. To be able to compare both models with each other the observed measurement as well as the estimated measurement were standardized before analyses. Validity of the models was evaluated by ranking ability (cross-classification between the observed (FFQ-derived) intake data and the estimated intake data (based on circulating fatty acids)) (Sluik et al. 2016). For the ranking ability, both observed (FFQ-derived) intake data and estimated intake data (based on circulating fatty acids) were ranked into quintiles. To validate the models, we repeated the multivariate backwards regression analyses in STATA, by implementing the svrboot command which created a table with the number of times each variable is selected after 1000 bootstrap samples (De Boer et al. 2012). Variables were included in the bootstrap models when these were present in the majority of the bootstrap attempts (>50%). In secondary analyses, variables were included in the bootstrap models when these were present in >70% of the bootstrap attempts. As additional analyses, we repeated the multivariate backwards regression analyses for unadjusted dairy fat intake (rather than energy-adjusted dairy fat intake which was used for the primary analyses), and also assessed predictive capacity for energy-adjusted dairy intake. All reported probability values are two-tailed, and $p \leq 0.05$ was considered statistically significant.

Results
The baseline characteristics of the 769 participants overall and for males and females separately are shown in Table 1. The total population had a mean age of 53.0 ± 15.5 years and a BMI of 26.0 ± 4.0 kg/m². Blood pressure was slightly higher in males (130/77 mmHg) compared to females (123/71 mmHg). Total and HDL cholesterol were higher in females, whereas total triglycerides was higher in males. Energy intake of the population was 1975 ± 624 kcal/d (8274 ± 2613 KJ/d). Of the total intake, 46 En% was derived from carbohydrates, 15 En% from proteins, and 35 En% from fats. The Lifelines dataset confirms that males overall have a higher dietary intake compared to females. Total meat intake of the population was 73.5 g/d, which contributed for 13% to the daily fat intake. Median dairy intake of the population was 322 (209–447) g/d and consisted for 12.8% of cheese, for 35.1% of milk, for 10.0% of yoghurt, and for 42.1% of other milk (based) products. Of the total dairy intake, 34% was from fermented dairy products. Median dairy fat intake of the population was 12.3 (8.4–17.4) g/d. Skimmed dairy products contributed for 2% to total dairy fat intake, while semi-skimmed and full-fat products contributed for 20% and 78% respectively. Median dairy fat intake, rather than median dairy intake overall, was significantly higher in males compared to females (median intakes males: 13.5 (9.0–18.3) g/d vs females: 11.3 (8.0–15.6) g/d; $p < 0.001$). Males consumed more semi-skimmed and full-fat dairy products, while females consumed more skimmed dairy products.

Circulating fatty acids were measured in the first 96 samples (Table S1). Trans-C18:1(n-7) appeared not detectable in plasma CE, since it was below the detection limit. All potential dairy fat biomarkers, including trans-C18:1(n-7) and CLA, were detectable in plasma TG and PL. Since we were especially interested in the potentially less established biomarkers, we decided to further only analyze plasma TG and PL. Data on potential circulating dairy fat biomarkers in plasma TG and PL of all 769 participants are shown in Table 1. Dairy fatty acids in plasma TG were higher compared to plasma PL. The quantitatively most abundant dairy fat marker was C14:0 (Total TG: 1.87 mol%, PL: 0.49 mol%), followed by C17:0 and C15:0 in both fractions. Trans-C18:1(n-7) was the highest trans fat in both fractions (TG: 0.10 mol%, PL: 0.09 mol%).
In the univariate main analyses, all potential dairy fat biomarkers in plasma TG, except C17:0, were associated with energy-adjusted dairy fat intake (Table 2, Model 1). The highest associations were found for the commonly used biomarker C15:0 (standardized $\beta$ (std $\beta$) = 0.286, $p < 0.001$) and the less established biomarker trans-C18:1(n-7) (std $\beta$ = 0.292, $p < 0.001$). All potential dairy fat biomarkers in plasma PL were also moderately associated with energy-adjusted dairy fat intake (Table 2, Model 1). Slightly higher, but still moderate associations were found with the less established biomarkers trans-C18:1(n-7) (std $\beta$ = 0.269, $p < 0.001$) and CLA (std $\beta$ = 0.272, $p < 0.001$) compared to the commonly used biomarkers. The highest associations with more commonly used biomarkers were found for C14:0 (std $\beta$ = 0.214, $p < 0.001$) and C15:0 (std $\beta$ = 0.215, $p < 0.001$), but these were still moderate. In sensitivity analyses, removal of outliers for energy intake did not materially change the association between dairy fat biomarkers and energy-adjusted dairy fat intake. In multivariate regression analyses, associations between fatty acids in plasma TG and PL with energy-adjusted dairy fat intake remained after adjustments for age, sex, BMI, total meat intake, and the industrial trans fatty acid Trans-C18:1(n-9) (Table S2(A)). The association between the biomarkers and energy-adjusted dairy fat intake did not differ between males and females, except for PL C14:0 ($p = 0.03$ for interaction, association in males: std $\beta$ = 0.264, $p < 0.001$, association in females: std $\beta$ = 0.173, $p < 0.001$ (Table S2(B), Model 1).

In plasma TG, multivariate backwards regression analyses identified a combination of C15:0 and trans-C18:1(n-7) as the best combination to predict energy-adjusted dairy fat intake (Table 2, Model 3). The explained variance of this
Table 2. Univariate regression analyses and multivariate models to predict dairy fat with circulating fatty acids in plasma TG, plasma PL and a combination of the two in the Lifelines Cohort.

| Model | Plasma TG | Plasma PL | Plasma TG + PL |
|-------|-----------|-----------|----------------|
|       | Std β     | $R^2$     | p-value        | Std β     | p-value | Std β     | p-value | Std β     | p-value |
| TG C14:0 | 0.204 | 0.071 | <0.001 | 0.049 | 0.27 |
| TG C15:0 | 0.286 | 0.111 | <0.001 | 0.133 | 0.01 | 0.162 | 0.001 |
| TG C17:0 | 0.040 | 0.032 | 0.19 | 0.016 | 0.67 | 0.002 | 0.181 | 0.001 |
| TG T-C16:1(n-7) | 0.233 | 0.072 | <0.001 | 0.031 | 0.54 | 0.000 | 0.94 |
| TG T-C18:1(n-7) | 0.292 | 0.115 | <0.001 | 0.162 | 0.002 | 0.181 | 0.001 |
| TG CLA | 0.215 | 0.076 | <0.001 | -0.004 | 0.94 | 0.130 | 0.128 |

Model 1: Univariate linear regression analysis of the fatty acid with dairy fat intake; Model 2: Multivariate model with all fatty acids from our population in one model; Model 3: Multivariate model with the best combination of fatty acids in our population (backwards regression).

C14:0, C15:0, trans-C16:1(n-7), trans-C18:1(n-7) and CLA in plasma TG and Trans-C16:1(n-7), Trans-C18:1(n-7) and CLA in plasma PL were transformed before analyses. Dairy fat intake was transformed and adjusted for energy intake according to the residual method. β’s are standardized beta’s.

TG: Triglycerides; PL: Phospholipids; C14:0: Myristic acid; C15:0: Pentadecanoic acid; C17:0: Heptadecanoic acid; T-C16:1(n-7): Trans-Palmitoleic acid; T-C18:1(n-7): Trans-Palmitoleic acid; CLA: Conjugated linoleic acid.

Agreement and ranking ability

To indicate the agreement between the observed intake (FFQ-derived) and estimated intake (based on circulating fatty acids) Bland-Altman plots were made. Plots were made for the best single markers (from a set of markers) which were found by backward regression analyses (Figures 2 and 3). In addition, validity of the models was evaluated by ranking ability (cross-classification between the observed intake data (FFQ-derived) and the estimated intake data (TG, PL and TG + PL model based on circulating fatty acids)). Both the observed and estimated intake data were ranked into quintiles and compared. For all three estimated models, ≥67% of the participants were classified in the same or adjacent quintile of the observed model, whereas 2–3% of the participants were classified in extreme quintiles. Overall, both methods showed acceptable agreement between the observed and estimated models.

Internal consistency of the models

To judge the internal consistency of the models, we performed backwards multivariate analyses using a bootstrap combination was 0.128, which was higher than the explained variance of the single markers (TG model: $R^2 = 0.128$ vs C15:0: $R^2 = 0.111$ and trans-C18:1(n-7): $R^2 = 0.115$). In plasma PL, the fatty acids C14:0, C15:0, trans-C18:1(n-7) and CLA were identified as the best combination to predict energy-adjusted dairy fat intake in the general population. This model had an explained variance of 0.143, which is again higher than the explained variance of the single markers (PL model: $R^2 = 0.143$ vs C14:0: $R^2 = 0.074$, C15:0: $R^2 = 0.075$, trans-C18:1(n-7): $R^2 = 0.099$ and CLA: $R^2 = 0.103$). A combination of fatty acids from TG (C15:0, trans-C18:1(n-7), CLA) and PL (C15:0, trans-C16:1(n-7), trans-C18:1(n-7), CLA) explained energy-adjusted dairy fat intake in the general population with an $R^2$ of 0.154. Based on these data, a combination of biomarkers can be seen as a better predictor for energy-adjusted dairy fat intake compared to single biomarkers.

Total dairy intake, total dairy fat intake and dairy product groups

For all, but one, circulating potential dairy fatty acid biomarkers in TG and PL, associations with total dairy fat intake were present and strongest compared to associations with total dairy intake and dairy product groups (Table S3). C17:0 is a single exception, with absence of an association with total dairy fat intake for TG and a stronger association with total dairy intake than with total dairy fat intake for PL. All fatty acid biomarkers, except C17:0 in TG and PL and Trans-C16:1(n-7) in PL were relatively strongly associated with intake of cheese and full fat dairy.

To judge the internal consistency of the models, we performed backwards multivariate analyses using a bootstrap
Figure 2. Bland-Altman plots for the agreement between the observed intake (FFQ-derived) and the estimated intake (based on the best circulating fatty acids in plasma TG) for the measurement of dairy fat intake in the general population. Plots of the differences between the observed intake (FFQ-derived) and the estimated intake (based on the best circulating fatty acids in plasma TG) vs the mean of the observed intake (FFQ-derived) and the estimated intake (based on the best circulating fatty acids in plasma TG) method, with (a) representing C15:0 and (b) Trans-C18:1(n-7) as the estimated intake. The red line displays the mean of the difference. The black dotted lines represents the limits of agreement (lower limit: mean - SD*1.96; upper limit: mean + SD*1.96). C15:0: Pentadecanoic acid; FFQ: food frequency questionnaire; SD: standard deviation; TG: triglycerides; Trans-C18:1(n-7): Vaccenic acid.

Figure 3. Bland-Altman plots for the agreement between the observed intake (FFQ-derived) and the estimated intake (based on the best circulating fatty acids in plasma PL) for the measurement of dairy fat intake in the general population. Plots of the differences between the observed intake (FFQ-derived) and the estimated intake (based on the best circulating fatty acids in plasma PL) vs the mean of the observed intake (FFQ-derived) and the estimated intake (based on the best circulating fatty acids in plasma PL) method, with (a) representing C14:0, (b) C15:0, (c) Trans-C18:1(n-7) and (d) CLA as the estimated intake. The red line displays the mean of the difference. The black dotted lines represents the limits of agreement (lower limit: mean - SD*1.96; upper limit: mean + SD*1.96). C14:0: Myristic acid; C15:0: Pentadecanoic acid; CLA: Conjugated Linoleic; FFQ: food frequency questionnaire; PL: phospholipids; SD: standard deviation; Trans-C18:1(n-7): Vaccenic acid.
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In the TG model, C15:0 and trans-C18:1(n-7) were present in more than half of the bootstrapped attempts, whereas in the PL model this was the case for C14:0, C15:0, trans-C18:1(n-7) and CLA. When fatty acids from TG and PL were combined, C15:0, trans-C18:1(n-7) and CLA from both plasma TG and PL were found in the majority of the bootstrap attempts. Trans-C16:1(n-7) in plasma PL did not remain in the prediction model. Complete bootstrap models of TG, PL and a combination of the two can be found in Table 3. Overall, the internal consistency for the fatty acids in plasma TG and PL was good, with exception for trans-C16:1(n-7) measured in plasma PL.

In secondary analyses, the bootstrap selection method was repeated with selection of fatty acids that were present in more than 70% of the 1000 bootstrap attempts. In the TG model, C15:0 and trans-C18:1(n-7) were again present, whereas the PL model included C15:0, trans-C18:1(n-7) and CLA, but not C14:0. The stricter selection criteria slightly worsened the internal consistency of the PL model. Fatty acids from TG and PL were also combined for these analyses. Interestingly, only the fatty acids trans-C18:1(n-7) and CLA from plasma PL remained in the model.

## Discussion

The main objective of the current study was to investigate whether the circulating fatty acids trans-C18:1(n-7) and CLA are an addition to the already existing dairy fat biomarkers in the general population. First of all, we found that the less established biomarkers for dairy fat intake were best quantifiable in plasma TG and PL. Additionally, we showed that trans-C18:1(n-7) and CLA, in both plasma TG and PL, were moderately associated with dairy fat intake. In plasma PL, these biomarkers were slightly higher compared to the commonly used biomarkers. Lastly, we developed and validated a model to predict dairy fat intake in the general population.

Compared to the single markers, we found that a combination of less established and commonly used biomarkers, including the C15:0, trans-C18:1(n-7) and CLA, showed a slightly higher association with dairy fat intake in the general population.

Significant associations of C15:0 and C17:0 with dairy fat intake were reported in the literature before (Wolk et al. 1998, Wolk et al. 2001, Brevik et al. 2005, Rosell et al. 2005, Biong et al. 2006, Sun et al. 2007, Golley and Hendrie 2014, Yakoob et al. 2014, Warensjo et al. 2015, Albani et al. 2016, Lund-Blix et al. 2016, Yakoob et al. 2016). In the Lifelines Cohort, C15:0 was also associated with dairy fat intake, while there was no consistent independent association of C17:0 with dairy fat intake. Plasma TG C17:0 was not associated with total dairy fat intake. Plasma PL C17:0 was associated with total dairy fat intake, however in backwards regression analyses taking all potential dairy fat biomarkers from plasma PL into account, plasma PL C17:0 did not remain in the PL model. Additionally, the bootstrapping with stepwise backwards algorithm showed that C17:0 remained in the minority of the 1000 bootstrap attempts (TG: 91x, PL: 149x) (Table S4).

A Norwegian study with 110 healthy men (age 21–55 years), published in 2005, also found that C15:0 from serum and adipose tissue were significantly correlated with dairy fat intake (both $r = 0.28$), while this was not the case for C17:0.

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### Additional analyses

As additional analyses, we repeated the multivariate backwards regression analyses for unadjusted dairy fat intake (rather than energy-adjusted dairy fat intake), and also assessed predictive capacity for energy-adjusted total dairy intake (Tables S5 and S6). The associations of the fatty acids with unadjusted dairy fat intake were weaker compared to the associations of the fatty acids with energy-adjusted dairy fat intake. However, C15:0 and trans-C18:1(n-7) remained the best predictive fatty acids in plasma TG, whereas trans-C18:1(n-7) and CLA remained the best predictive fatty acids in plasma PL.

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### Table 3. Multivariate models to predict dairy fat with circulating fatty acids in plasma TG, plasma PL and a combination of the two in the Lifelines Cohort based on 1000 bootstraps.

| Variable     | Beta ± SE | Std (beta) | p-value | 95%CI     |
|--------------|-----------|------------|---------|-----------|
| Plasma TG    |           |            |         |           |
| TG C15:0     | 1.752 ± 0.55 | 0.162  | 0.001  | 0.74; 2.86 |
| TG T-C18:1(n-7) | 1.798 ± 0.48 | 0.181  | 0.001  | 0.85; 2.71 |
| Plasma PL    |           |            |         |           |
| PL C14:0     | 0.538 ± 0.27 | 0.074  | 0.04   | 0.02; 1.07 |
| PL C15:0     | 1.102 ± 0.47 | 0.085  | 0.02   | 0.19; 2.02 |
| PL T-C18:1(n-7) | 2.032 ± 0.52 | 0.148  | 0.001  | 0.94; 3.05 |
| PL CLA       | 2.844 ± 0.74 | 0.151  | 0.001  | 1.41; 4.31 |
| Plasma TG + PL|         |            |         |           |
| TG C15:0     | 1.083 ± 0.63 | 0.100  | 0.08   | −0.20; 2.31|
| TG T-C18:1(n-7) | 1.144 ± 0.59 | 0.115  | 0.05   | −0.10; 2.56|
| TG CLA       | −1.131 ± 0.67 | −0.091 | 0.09   | −2.53; 0.15|
| PL C15:0     | 0.912 ± 0.52 | 0.070  | 0.08   | −0.11; 1.96|
| PL T-C18:1(n-7) | 1.155 ± 0.59 | 0.084  | 0.04   | 0.04; 2.28 |
| PL CLA       | 3.112 ± 0.78 | 0.166  | 0.001  | 1.56; 4.63 |

Fatty acids are included in the multivariate model when the circulating fatty acids were present in more than 50% of the models during the bootstrap process based on the backwards stepwise algorithm.

C14:0, C17:0, Trans-C16:1(n-7), Trans-C18:1(n-7) and CLA in plasma TG and Trans-C16:1(n-7), Trans-C18:1(n-7) and CLA in plasma PL were transformed before analyses. Dairy fat intake was transformed and adjusted for energy intake according to the residual method. β’s are standardized beta’s.

TG: Triglycerides; PL: Phospholipids; C14:0: Myristic acid; C15:0: Pentadecanoic acid; C17:0: Heptadecanoic acid; T-C16:1(n-7): T-Palmitoleic acid; T-C18:1(n-7): Vaccenic acid; CLA, Cis-9, Trans-11 Conjugated linoleic acid.
may be the best combination to predict the dairy fat intake and in the bootstrap models. Therefore, a combination of C15:0 and trans-C18:1 as the best biomarkers from the set of investigated biomarkers for the prediction of dairy fat intake. Every prediction model in the lifelines cohort included C15:0 as an important marker of dairy fat intake with standardized beta’s for the association with dairy fat intake of 0.29 in plasma TG and 0.22 in plasma PL. These results are in line with the current literature. For instance, in a study with more than 3000 subjects, published in 2016, the correlation for total plasma C15:0 with dairy fat intake was 0.29 (Yakoob et al. 2016). In addition, in a study with 279 subjects, published in 2015, the correlation of plasma PL C15:0 with dairy fat intake was 0.32 (Warenajo et al. 2015). This is one of the first studies investigating the association between trans-C18:1(n-7), measured in plasma TG and PL, and dairy fat intake. A single study measured trans-C18:1(n-7) in total plasma and erythrocytes, but did not find a significant correlation \( r = 0.06 \) and \( r = -0.01 \) respectively (Sun et al. 2007). Most other studies investigated a cluster of trans-C18:1 fatty acids, which is a combination of the rumen (trans-C18:1(n-7)) and industrial derived (trans-C18:1(n-9)) fatty acids, rather than trans-C18:1(n-7) alone (Yu et al. 2012, Zong et al. 2014). One of these studies is a Chinese study of 2091 adult participants aged 50–70 years, published in 2014, which found an association of 0.37 between the cluster of trans-C18:1 fatty acids assessed in erythrocytes and total dairy consumption (Zong et al. 2014). The current study showed a lower, but significant association between the ruminant derived trans fatty acid trans-C18:1(n-7) and dairy fat intake with a standardized beta of 0.29 for TG and of 0.27 for PL.

CLA in plasma PL was also found to be associated with dairy fat intake in the Lifelines population, both as a single marker and in combination with C14:0, C15:0 and trans-C18:1(n-7), while CLA in plasma TG was less strongly associated with dairy fat intake. CLA in plasma TG as a single marker was significantly associated with dairy fat intake (std.\( \beta = 0.22 \)), however the association of CLA with dairy fat intake did not remain in the backwards regression model when only plasma TG fatty acids were included. Current papers report mainly on the correlation between dairy fat intake and CLA in adipose tissue. For example, in a study of 3630 Costa Rican adults, published in 2012, the correlation of CLA with dairy fat intake was 0.24 (Aslibekyan et al. 2012). The correlation in adipose tissue is in line with the correlation found with CLA in plasma PL.

In the current cohort, we found that a combination of biomarkers is a slightly better predictor for dairy fat intake in the general population when compared to single markers. C15:0 and trans-C18:1(n-7) were found in all three models and in the bootstrap models. Therefore, a combination of dairy fat biomarkers, particularly models including these two, may be the best combination to predict the dairy fat intake in the general population.

The combined fatty acids assessed in PL provided a slightly better prediction of dairy fat intake than the combined fatty acids in TG (\( R^2 \) PL: 0.143 vs \( R^2 \) TG: 0.128). Consistent with this, when the selection criterion for inclusion in the model was set at present in >70% of the models, it was found that only plasma PL trans-C18:1(n-7) and plasma PL CLA remained in the model, if both TG and PL fatty acids were combined. This suggests that fatty acids in PL are stronger predictors than fatty acids in TG. This might be due to the fact that plasma TG is influenced by recent dietary intake (intake in the last few hours), whereas this is less so for plasma PL (Hodge et al. 2007). Recent meals, including large amounts of carbohydrates, can for instance turn on de novo lipogenesis which changes the fatty acid balance in plasma triglycerides and may therefore change the association with dairy fat intake. Researchers of the FORCE consortium, therefore, recommend to measure biomarkers of dietary intake in plasma CE and PL over plasma TG (Hodson et al. 2008, Del Gobbo et al. 2016, Wu et al. 2017). Indeed, most of the biomarker studies are carried out in plasma PL (Smedman et al. 1999, Wolk et al. 2001, Rosell et al. 2005, Biong et al. 2006, Warenajo et al. 2015, Lund-Blix et al. 2016), while only one study reported on biomarkers in serum TG (Biong et al. 2006). Importantly, however, one of the most promising markers, i.e. ruminant trans-C18:1(n-7), was only detectable in PL and TG, and not in CE. Furthermore, by measuring the fatty acids in both plasma TG and PL, it was possible to identify differences between both fractions.

Prediction of energy-adjusted dairy fat intake was the primary outcome for this paper. In additional analyses, we repeated the analyses for unadjusted dairy fat intake, and also assessed predictive capacity for energy-adjusted dairy intake. Both C15:0 and trans-C18:1(n-7) in plasma TG and both trans-C18:1(n-7) and CLA in plasma PL were associated with unadjusted dairy fat intake, albeit with slightly lower effect sizes than with energy-adjusted dairy fat intake. The associations between dairy biomarkers and energy-adjusted dairy intake were weaker than for energy-adjusted dairy fat intake. Obviously, dairy fatty acid biomarkers serve best to predict dairy fat intake.

This study has several strong points. Firstly, we are one of the first studies to identify trans-C18:1(n-7) and CLA in both plasma TG and PL as biomarkers of dairy fat intake. Another strength of the present study is its large population size. In total, complete data of 769 participants were used for analyses. Additionally, we were able to measure a large amount of dairy fat biomarkers in both plasma TG and PL in one cohort.

We must acknowledge that the current study has some limitations. First of all, the \( R \)-squares of the models for prediction of dairy fat intake were relatively low. This means that the dairy fatty acid biomarkers are not strongly related to the dairy fat intake from the FFQ. However, it cannot be discerned whether the weak associations are intrinsic to the dairy fatty acid biomarkers, or that the FFQ as a reference is not optimal. Concerning the dairy fatty acid biomarkers, it has been shown that some of the fatty acids in plasma are
not only influenced by dairy fat intake, but can also be affected by metabolic processes (e.g. trans-C16:1(n-7) can be influenced by β-oxidation of trans-C18:1(n-7)), or by the intake of dietary fibres (e.g. C17:0 can be influenced by insuln), meat and coconut oil (e.g. C14:0) (Zock et al. 1994, Valsta et al. 2005, Jaudszus et al. 2014, Weitkunat et al. 2017). A limitation of this study is that we have no specific data on types of oils and types of meat, which precluded adjustment of these potential confounders. However, we had data on total meat intake. In our cohort adjustment for total meat intake did not alter the association between fatty acid biomarkers and energy-adjusted dairy fat intake (Table S2(A)). Another limitation is that circulatiny fatty acids as biomarkers for dietary intake were expressed as proportions rather than as absolute concentrations, which makes them relative and therefore dependent on the presence of other major circulating fatty acids. The reason that we report on proportions of dairy fatty acids is to align with existing literature and make results comparable. It should also be noted that the fatty acid composition of dairy fat can vary depending on factors such as cow genetics, forage content, production level, time of lactation, physiological status of the animal and seasonal variation (Jensen 2002, Mansson 2008, Hanus et al. 2018; German and Dillard 2006). Nevertheless, fatty acid biomarkers can provide an objective estimation of dietary fat intake, while this is intrinsically not the case with FFQ, which may be subject to under- and over reporting by participants (Kristal et al. 2005, Sluik et al. 2016). Overall, there is no gold standard for measuring dietary intakes and the FFQ can therefore be seen as the best method so far.

This study is the first observational study investigating commonly used biomarkers in combination with less investigated trans fatty acid biomarkers. A first step towards validation would be performance of similar studies in other populations. Furthermore, as the applied assessment method is labour intensive, the results of this study should first be replicated in other studies to show that incorporation is worthwhile. Concluding, this paper showed associations between circulating fatty acids and dairy fat intake in the general population, which can be seen as a first step towards better measurement techniques for dairy fat intake.

Conclusion

In this study, models were developed and validated for the prediction of dairy fat intake in the general population. The study confirmed the utility of commonly used biomarkers and identified the biomarkers trans-C18:1(n-7) and CLA as potential dairy fat biomarkers, which are one of the best biomarkers to predict dairy fat intake in the Lifelines population. Moreover, a combination of the biomarkers may be even a stronger predictor for dairy fat intake. A first step towards validation would be performance of similar studies in other populations.

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The manuscript is based on data from the LifeLines Cohort Study. LifeLines adheres to standards for open data availability. The data catalog of LifeLines is publicly accessible on www.LifeLines.net. All international researchers can apply for data at the LifeLines research office (LLscience@umcg.nl). The Lifelines system allows access for reproducibility of the study results.

Disclosure statement

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