Fatty acid and fat-soluble antioxidant concentrations in milk from high- and low-input conventional and organic systems: seasonal variation

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Abstract

BACKGROUND: Previous studies showed differences in fatty acid (FA) and antioxidant profiles between organic and conventional milk. However, they did not (a) investigate seasonal differences, (b) include non-organic, low-input systems or (c) compare individual carotenoids, stereoisomers of α-tocopherol or isomers of conjugated linoleic acid. This survey-based study compares milk from three production systems: (i) high-input, conventional (10 farms); (ii) low-input, organic (10 farms); and (iii) low-input non-organic (5 farms). Samples were taken during the outdoor grazing (78 samples) and indoor periods (31 samples).

RESULTS: During the outdoor grazing period, on average, milk from the low-input systems had lower saturated FAs, but higher mono- and polyunsaturated FA concentrations compared with milk from the high-input system. Milk from both the low-input organic and non-organic systems had significantly higher concentrations of nutritionally desirable FAs and antioxidants – conjugated linoleic (60% and 99%, respectively) and α-linolenic (39% and 31%, respectively) acids, α-tocopherol (33% and 50%, respectively) and carotenoids (33% and 80%, respectively) – compared with milk from the high-input system. Milk composition differed significantly between the two low-input systems during the second half of the grazing period only; with milk from non-organic cows being higher in antioxidants, and conjugated linoleic acid, and that from organic cows in α-linolenic acid. In contrast, few significant differences in composition were detected between high-input and low-input organic systems when cows were housed.

CONCLUSIONS: Milk composition is affected by production systems by mechanisms likely to be linked to the stage and length of the grazing period, and diet composition, which will influence subsequent processing, and sensory and potential nutritional qualities of the milk.

Keywords: milk; low-input farming; organic farming; fatty acid profiles

INTRODUCTION

The fatty acid (FA) and fat-soluble antioxidant composition in milk fat is known to affect processing and sensory quality of dairy products,1,2 and may also affect their nutritional value.3,4 The degree of saturation in milk fat has a bearing on the hardness, texture and taste of manufactured dairy products, particularly butter and cheese.5 The presence of longer-chain saturated fatty acids (SFA) increases the hardness of butter, while milk with a high proportion of unsaturated FA content (typical range 275–400 g kg−1 fat) tends to give softer products (e.g., more spreadable butter). Unsaturated (especially polyunsaturated) FAs are also more prone to oxidation, which results in the development of off-flavour and reduced shelf-life in milk and dairy products.6 However, the sensory quality and shelf-life of milk and dairy products is determined by the balance of unsaturated FAs and fat-soluble antioxidants, which protect against oxidation and off-flavour development.6–8

High dietary intakes of SFA (which account for 60–70% of milk fat) is a risk factor for development of obesity, cardiovascular disease (CVD), impaired insulin sensitivity and the ‘metabolic syndrome’.9 In

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contrast, dietary intake of certain unsaturated fatty acids, in particular conjugated linoleic acid (CLA) and omega-3 fatty acids (n-3 FA), and fat-soluble antioxidants (e.g., α-tocopherol, carotenoids) has been linked to potential health benefits. CLA and n-3 FA have been shown to counteract the negative physiological effects of SFA, and CLA has also been linked to anticancer properties, reduced risk of type 2 diabetes, CVD and enhanced immune function. However, while CLA isomer C18:2 c9 t11 (CLA9) was only linked to beneficial health impacts, another CLA isomer, C18:2 t10 c12 (CLA10), was also associated with some negative health impacts in cell culture and animal models. In studies comparing the impact of different (e.g., organic and conventional) production systems on milk fat composition, it is therefore important to compare concentrations of both CLA isomers. Most previous comparative studies only reported concentrations of individual isomers or total CLA and also did not report concentrations of vaccinic acid (VA), the precursor for CLA. Milk contains significant concentrations of VA and, since a proportion can be readily converted to CLA9 in the human body, the total potential CLA9 supply can only be estimated if both VA and CLA9 levels are known.

Previous studies showed that the feeding regime has a major effect on the FA profiles of milk, but that other factors (including breed/genotype, stage and number of lactations) may also influence milk composition. Dietary unsaturated fatty acids are likely to undergo hydrogenation by rumen microorganisms and long-chain fatty acids may be subjected to desaturase activity in the mammary gland. The FA profile of milk, therefore, is primarily determined by: (i) the balance of fatty acids in the diet; (ii) the extent of rumen hydrogenation; and (iii) mammary desaturase activity. CLA levels are linked to dietary supply of α-linolenic acid (αLA) and linoleic acid. However, while 70–90% of CLA9 (which constitutes >70% of total CLA in milk) is generated from desaturation of VA in the mammary gland, all other CLA isomers (including CLA10) are generated as intermediates of rumen bihydrogenation and are therefore found at much lower concentrations than CLA9 in milk.

Fat-soluble antioxidants/vitamins present in milk are derived from dietary sources, either from (i) natural constituents in feedstuff (especially the forage component of the diet) or (ii) synthetic compounds added as supplements to the diet of lactating cows. Carotenoids derived from fresh forage are dominated by β-carotene, but also include lutein, zeaxanthin, cryptoxanthin, lycopene and α-carotene. The main vitamin E activity in fresh forage is associated with the RRR isomer of α-tocopherol (the only isomer synthesized by plants), with some activity being associated with β-, γ- and δ-tocotrienol and α-, β-, γ- and δ-tocopherol.

Most high-input conventional dairy production systems supplement diets with proprietary mineral and/or vitamin products containing A vitamins, vitamin D3 and E vitamins (in particular α-tocopherol); such supplements are prohibited in organic production. The naturally occurring RRR isomer of α-tocopherol has a higher vitamin E activity (1.49 IU mg⁻¹) than synthetic vitamin E (1.0 IU mg⁻¹), which contains equal proportions of the eight different stereoisomers of α-tocopherol. Synthetic α-tocopherol products are referred to as ‘all rac’ α-tocopherol and consist mainly of 2R stereoisomers. Synthetic α-tocopherol is absorbed with the same efficiency as the RRR stereoisomer of α-tocopherol, but levels of uptake into key tissues (e.g. the brain) are lower. Also, a recent study with dairy cows found higher α-tocopherol concentrations in blood and milk following supplementation of RRR compared with ‘all rac’ α-tocopherol and reported preferential transfer of RRR isomers into milk by cows receiving the synthetic isomer mix.

Milk and dairy products from certified organic dairy production systems have been reported to contain higher concentrations of polyunsaturated fatty acids (PUFA), αLA (the main n-3 FA in milk), and/or CLA, and fat-soluble antioxidants than those from high-input conventional production. These studies did not include non-organic, low-input systems in comparisons. However, an increasing number of dairy farms in Europe, New Zealand/Australia and North America are adapting ‘lower-input’ production methods similar to those used in organic farming, but do not comply with all input restrictions prescribed by organic farming standards. Most importantly, these systems use mineral NPK fertilizers, but often at reduced levels compared with conventional high-input systems. It is unclear whether such non-organic, low-input systems can provide similar benefits in milk composition to certified, organic dairy production systems.

Milk composition is known to change when switching from outdoor grazing to indoor forage-based diets in winter; however, little is known about whether this dietary change affects the differential in milk quality between organic and conventional systems reported previously. There is also limited information on differences in the composition of fat-soluble antioxidants in milk from high- and low-input dairy systems and the few studies available show contradictory results. Such information would, however, be essential to assess (i) the overall nutritional value of milk from low-input systems and (ii) whether the higher unsaturated fat content of organic milk (and associated risk of oxidation and off-flavour development) is compensated for by higher concentrations of antioxidants.
The objectives of this study were therefore to: (i) compare the fatty acid and fat-soluble antioxidant composition of milk from three UK production systems – certified-organic ‘low-input’ (O-LI), non-organic certified ‘low-input’ (NO-LI) and standard ‘high-input’ (HI) conventional production systems, during the outdoor grazing period; (ii) quantify differences in fatty acid and fat-soluble antioxidant content of milk between O-LI and HI systems, during the winter indoor (conserved forage-based) feeding period; and (iii) identify whether there are differences in milk composition between certified-organic ‘low-input’ (O-LI) and non-certified ‘low-input’ (NO-LI) systems that use spring block calving systems and graze cows outdoors throughout lactation.

Table 1. Differences in management and production system parameters between high-input conventional (HI), organically certified (O-LI) and non-organic (NO-LI) low-input farms (mean values over all samples, with standard deviation in parentheses)

<table>
<thead>
<tr>
<th>Parameters recorded</th>
<th>HI</th>
<th>O-LI</th>
<th>NO-LI</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Herd characteristics</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Herd size (milking cows)*</td>
<td>252 (125)</td>
<td>160 (93)</td>
<td>322 (141)</td>
</tr>
<tr>
<td>Breed Indexa</td>
<td>0.0 (0)</td>
<td>0.2 (0.3)</td>
<td>0.3 (0.1)</td>
</tr>
<tr>
<td>% primiparous cows*</td>
<td>25 (7)</td>
<td>27 (12)</td>
<td>30 (8)</td>
</tr>
<tr>
<td>Live weight of cows (kg)b</td>
<td>650 (0)</td>
<td>610 (34)</td>
<td>588 (21)</td>
</tr>
<tr>
<td>Dry matter intake (kg d⁻¹)c</td>
<td>19.5 (0.5)</td>
<td>17.6 (1.0)</td>
<td>16.9 (0.7)</td>
</tr>
<tr>
<td><strong>Diet composition</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>1. Outdoor period</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Fresh forage (proportion DMI)</td>
<td>0.37 (0.24)</td>
<td>0.84 (0.23)</td>
<td>0.95 (0.07)</td>
</tr>
<tr>
<td>Conserved forage (proportion DMI)</td>
<td>0.29 (0.15)</td>
<td>0.08 (0.16)</td>
<td>0 (0)</td>
</tr>
<tr>
<td>- Grass silageæ*</td>
<td>0.73 (0.28)</td>
<td>0.72 (0.40)</td>
<td>0.0 (0)</td>
</tr>
<tr>
<td>- Maize silageæ*</td>
<td>0.10 (0.20)</td>
<td>0 (0)</td>
<td>0 (0)</td>
</tr>
<tr>
<td>- Other silageæ,æ*</td>
<td>0.13 (0.18)</td>
<td>0 (0)</td>
<td>0 (0)</td>
</tr>
<tr>
<td>- Straw/hayeæ</td>
<td>0.04 (0.09)</td>
<td>0.28 (0.40)</td>
<td></td>
</tr>
<tr>
<td>Concentrate (proportion of DMI)</td>
<td>0.34 (0.13)</td>
<td>0.08 (0.09)</td>
<td>0.05 (0.07)</td>
</tr>
<tr>
<td>- Cerealsæ</td>
<td>0.31 (0.24)</td>
<td>0.23 (0.40)</td>
<td>0.05 (0.14)</td>
</tr>
<tr>
<td>- By-productsæ γ</td>
<td>0.30 (0.23)</td>
<td>0.20 (0.40)</td>
<td>0.52 (0.50)</td>
</tr>
<tr>
<td>- Other concentratesæ,æ</td>
<td>0.40 (0.31)</td>
<td>0.57 (0.49)</td>
<td>0.43 (0.53)</td>
</tr>
<tr>
<td>Mineral supplementsæ (g cow⁻¹ day⁻¹)</td>
<td>142 (75)</td>
<td>8 (17)</td>
<td>3 (13)</td>
</tr>
<tr>
<td>Vitamin E supplementæ (IU cow⁻¹ day⁻¹)</td>
<td>450–750</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>2. Indoor period</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Fresh forage (proportion DMI)f</td>
<td>0 (0)</td>
<td>0.24 (0.38)</td>
<td>NA</td>
</tr>
<tr>
<td>Conserved forage (proportion of DMI)</td>
<td>0.56 (0.08)</td>
<td>0.54 (0.30)</td>
<td>NA</td>
</tr>
<tr>
<td>- Grass silageæ</td>
<td>0.69 (0.29)</td>
<td>0.80 (0.19)</td>
<td>NA</td>
</tr>
<tr>
<td>- Maize silageæ</td>
<td>0.05 (0.12)</td>
<td>0 (0)</td>
<td>NA</td>
</tr>
<tr>
<td>- Other silageæ,æ</td>
<td>0.24 (0.28)</td>
<td>0.20 (0.19)</td>
<td>NA</td>
</tr>
<tr>
<td>- Straw/hayeæ</td>
<td>0.02 (0.04)</td>
<td>0 (0)</td>
<td>NA</td>
</tr>
<tr>
<td>Concentrate (proportion of DMI)</td>
<td>0.44 (0.08)</td>
<td>0.23 (0.10)</td>
<td>NA</td>
</tr>
<tr>
<td>- Cerealsæ</td>
<td>0.31 (0.17)</td>
<td>0.42 (0.16)</td>
<td>NA</td>
</tr>
<tr>
<td>- By-productsæ γ</td>
<td>0.24 (0.16)</td>
<td>0.07 (0.11)</td>
<td>NA</td>
</tr>
<tr>
<td>- Other concentratesæ,æ</td>
<td>0.45 (0.24)</td>
<td>0.51 (0.23)</td>
<td>NA</td>
</tr>
<tr>
<td>Mineral supplementsæ (g cow⁻¹ day⁻¹)</td>
<td>150 (53)</td>
<td>22 (31)</td>
<td>NA</td>
</tr>
<tr>
<td>Vitamin E supplementæ (IU cow⁻¹ day⁻¹)</td>
<td>250–674</td>
<td>0</td>
<td>0</td>
</tr>
</tbody>
</table>

* Based on farm records and collected by questionnaire; a estimated proportion of non-Holstein–Friesian genetics in the herd; b estimated based on breed index; c estimated based on live weight and milk yield; d whole-crop wheat, barley and/or oats, dry matter; e proportion of total conserved forage intake; f when weather permitted, most organic herds were grazed in the day; g brewing and distillers’ waste and/or sugar beet pulp; h bought in or farm produced compound/mixed concentrate feeds; i no oilseed or fat supplementation was recorded by farmers; NA, not applicable (NO-LI cows were grazed throughout the lactation).
and assumed live weight (DMI = 0.025 LW + 0.125 milk yield). Grazing was calculated at the herd level by difference: DMI (fresh grass) = total DMI – DMI (conserved forage + concentrate; recorded by producers). Since cow live weight varied between farming systems, recorded levels of dietary components were used to calculate proportions of total intake, to allow a more relevant comparison between systems. Tables 1 and 2 list diet composition for each production system during grazing and the housed periods of this study.

Conventional ‘high-input’ (HI) farms
Ten farms were selected representing common conventional production and feeding systems in the UK. HI farms used predominantly pure ryegrass swards during the grazing period, winter diets based on grass silage and higher concentrate:conserved forage ratio diets during the indoor feeding period than LI farms (see Table 1 for the diets used during the outdoor grazing and indoor feeding periods). The HI group did not include farms with extremely high-input/output systems (e.g., farms which use more than 50% of the diet coming from concentrates, regularly milk three times per day and/or those that house animals throughout their lactation). All farms were all-year round-calving and had similar proportions of cows in early lactation at all sampling dates.

Organically certified ‘low-input’ (O-LI) farms
Ten farms were selected representing two principal organic dairy systems found in the UK: (a) an all-year-round calving system (five farms) in which lactating cows are grazed when conditions allow (spring to autumn), but fed on conserved forage-based diets during the winter indoor period (see Table 1); and (b) a spring block calving system in which cows are grazed throughout lactation (March to October) and were only indoors when not lactating between November to February. All-year-round calving farms had similar proportions of cows in early lactation at all sampling dates. Diets used in both organic systems were similar during the outdoor grazing period (Table 1); all O-LI farms used mixed grass–clover swards and did not apply mineral N or water-soluble P fertilizers. Where appropriate, on the basis of soil analyses, finely ground rock phosphate fertilizers were applied.

Non-organically certified ‘low-input’ (NO-LI) farms
Five farms representing the main non-organic, ‘low-input’ system found in the UK were selected. All farms used a New Zealand-type production system26 with spring block calving, in which cows were grazed throughout the lactation and no, or low levels of concentrate and/or other feed supplements included in the diet (see Table 1). As with the organic spring block calving herds, cows were only housed when not lactating between November and February. NO-LI farms selected used mixed grass–clover swards, but applied up to 120 kg N ha⁻¹ per year of mineral N and water-soluble P fertilizer at levels determined from soil analyses.

Samples were taken in August and October in 2004 and in January, March and May in 2005 from all farms. In January 2005 samples could only be collected from O-LI and HI farms that used an all-year-round calving system. Samples of milk were taken from the stirred bulk tank after two milkings (representing a 24 h production period), at each participating farm and frozen immediately after sampling and kept at −20 °C until dispatched for analysis.

<table>
<thead>
<tr>
<th>Sampling date</th>
<th>Dietary components (proportion of DMI)</th>
<th>O-LI</th>
<th>NO-LI</th>
</tr>
</thead>
<tbody>
<tr>
<td>August</td>
<td>Fresh forage 0.96 (0.04) 0.92 (0.08)</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Conserved forage 0 (0) 0 (0)</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Concentrate 0.04 (0.04) 0.08 (0.08)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>October</td>
<td>Fresh forage 0.88 (0.11) 0.96 (0.08)</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Conserved forage 0.04 (0.06) 0 (0)</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Concentrate 0.08 (0.08) 0.05 (0.08)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>March</td>
<td>Fresh forage 0.86 (0.20) 0.95 (0.07)</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Conserved forage 0.11 (0.15) 0 (0)</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Concentrate 0.03 (0.06) 0.05 (0.07)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>May</td>
<td>Fresh forage 0.96 (0.06) 1.00 (0)</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Conserved forage 0 (0) 0 (0)</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Concentrate 0.04 (0.06) 0 (0)</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

DMI, dry matter intake.

Extraction of fat from milk
The extraction of fat from the milk was carried out as described by Havemose et al.,23 with minor modifications. Milk fat was extracted from milk (2 mL) by adding methanol (2 mL) and chloroform (4 mL). The mixture was shaken vigorously for 1 min, then centrifuged for 10 min at 3000 × g at 4 °C. The lower phase containing the lipid fraction was isolated and evaporated to dryness under nitrogen.

Methylation of fatty acids from milk
The methylation of fatty acids extracted from the milk was carried out as described by Havemose et al.,23 with minor modifications. Fat (approx. 10 mg) was dissolved in sodium methylate solution (2 g L⁻¹ methanol) in sealed glass tubes filled with argon, incubated at 60 °C for 30 min, and then cooled on ice. Saturated sodium chloride solution (4 mL) and pentane (1 mL) were added. The samples were mixed on a vortex mixer for 1 min and centrifuged at 1700 × g for 10 min. The upper pentane phase was collected and used for gas chromatographic analysis.
Analysis of fatty acid composition by gas chromatography
Separation and quantification of the fatty acids isolated from milk was carried out as described by Havemose et al., with modifications. Samples (1 μL) of the pentane phase containing the fatty acid methyl esters were analysed by gas chromatography (HP6890 GC system, Hewlett Packard Co., Palo Alto, CA, USA) with a flame ionization detector and a Supelco SI 2560 column (100 m × 0.25 mm × 0.20 μm, Supelco, Bellafonte, PA, USA). The inlet temperature was 275 °C with a split ratio of 40:1, and the carrier gas was helium with a constant flow of 1.5 mL min⁻¹. The starting temperature of 140 °C was held for 5 min and increased by 4 °C min⁻¹ to an end temperature of 240 °C. The detector temperature was 300 °C.

The concentrations of saturated (SFA), monounsaturated (MUFA) and polyunsaturated (PUFA) fatty acids and the ratio of n-3 and n-6 isomers of linoleic acid (C18:3) were then calculated as a proportion of total fatty acids recovered, based on the use of external standards. To calculate the n-3:n-6 FA ratio, the concentration of the main n-3 FA (α-LA) was divided by the sum of the concentrations of the following n-6 FA isomers: 18:2 t9 t12, 18:2 t10 t12, 18:2 c9 c12, 18:3 c6 c9 c12 and 20:4 c5 c8 c11 c14.

**Analysis of fat-soluble antioxidant composition**
Fat-soluble antioxidants (α-tocopherol, β-carotene, lutein and zeaxanthin) were analysed using the high-performance liquid chromatographic method described by Havemose et al. Isomers of α-tocopherol were analysed using the methods described by Meglia et al.

**Statistical analysis**
Linear mixed-effects models were used to investigate differences in milk quality parameters under the different systems (HI, O-LI and NO-LI). These models use two types of explanatory variables: fixed effects, which affect the mean of the response; and random effects, which affect the variance of the response. In these analyses, farm identifier was used as a random effect. Three sets of analyses were undertaken: (i) comparison of milk samples from all three systems (HI, O-LI and NO-LI) taken during the outdoor grazing period (samples from the spring block and all-year calving organic farms were pooled, because no major differences could be detected in preliminary analyses; results not shown); (ii) comparison of samples taken from HI and all-year calving O-LI farms during the indoor period when cows were on conserved forage-based diets; and (iii) comparison of samples taken from spring block calving O-LI and NO-LI herds at four different sampling dates using a two-factorial model (system and date), adapted to account for repeated measures from the four dates, to identify (a) whether at any time during the grazing period milk quality differed between the two LI systems and (b) interactions between the two factors for any of the milk quality parameters assessed. All proportion data were arcsine transformed prior to statistical analysis, but means presented were calculated from non-transformed data. Pairwise comparisons of means were carried out, where appropriate, using Tukey’s honest significant difference tests.

All statistical analyses were carried out using the R statistical environment.

**RESULTS**
**Comparison of milk fat composition during the outdoor period (fresh forage-based diets)**

On average the total fat content was higher in milk from LI systems compared with the HI system, and was significantly higher for the NO-LI system compared with the HI system (Table 3). When the composition of milk fat was compared, on average, the percentage of SFAs in milk fat was lower, while percentages of both MUFA (of which >80% was oleic acid C18:1 cis9) and PUFA were higher in milk from LI systems, compared with the HI system, and was significantly higher for the NO-LI system compared with the HI system (Table 3).

Percentages of the nutritionally desirable FAs (α-LA and CLA9) were significantly higher, while levels of total n-6 PUFAs were significantly lower in milk from both LI systems, when compared with milk from HI farms (Table 3). As a result, the n3:n6 ratio was also higher in milk from LI systems (Table 3). CLA10 was found in low concentrations in milk from all production systems and was not affected by production system (Table 3). Differences between O-LI and NO-LI were generally smaller than those between HI and LI systems, but the percentage of CLA was significantly higher in milk from NO-LI systems and the percentage of total n-6 FA was significantly higher in milk from O-LI systems (Table 3).

The concentrations of most antioxidants (the RRR stereoisomer of α-tocopherol, β-carotene, lutein and zeaxanthin) were highest in milk from O-LI, at intermediate concentrations in milk from NO-LI, and lowest in milk from HI systems (Table 3) during the outdoor period. Concentrations of the 2R stereoisomer of α-tocopherol were not significantly different between systems, but were slightly lower in milk from NO-LI systems.

**Comparison of milk fat composition during the indoor period (conserved forage-based diets)**

Since the spring, block-calving NO-LI and O-LI systems did not produce milk during the indoor period only milk from all-year calving O-LI and HI systems was compared.
In contrast to results from the outdoor rearing period, there were few differences in milk composition during the housed period. The percentages of total SFA in milk fat were significantly higher (4%) and MUFA significantly lower (10%) in milk from HI systems compared with milk from HI systems (Table 4). There was also a significantly lower (24%) content of n-6 fatty acids and trends towards a higher content (38%) of α-linolenic acid (P = 0.052) and a higher (30%) lutein content (P = 0.081) in O-LI milk compared with HI milk (Table 4).

Comparison of milk fat composition during the grazing period between O-LI and NO-LI spring block calving dairy systems

Apart from CLA9 isomer (which was present in significantly higher percentages in milk from NO-LI farms on the August and May sampling dates), significant differences in FA composition between O-LI and NO-LI block calving systems were found only late in the outdoor grazing period (August and October sampling date, Fig. 1). The percentages of total SFA and αLA were higher in milk from O-LI systems, while percentages of MUFA, PUFA, VA and CLA9 were higher in milk from NO-LI systems. No significant differences in the percentages of CLA10 and n-6 FAs were detected (data not shown). There were also significant interactions between LI production system and date for PUFA (P = 0.020; Fig. 1(c)), VA (P = 0.029; Fig. 1(e)) and CLA (P = 0.030; Fig. 1(f)).

The concentration of most antioxidants changed significantly over time, and at specific dates significant differences in the concentrations of individual antioxidants between the two LI systems could be detected. Concentrations of 2R toc were significantly higher in milk from O-LI systems in May, while concentrations of 3R toc were significantly higher in NO-LI systems. No significant differences in the percentages of CLA10 and n-6 FAs were detected (data not shown). There were also significant interactions between LI production system and date for PUFA (P = 0.020; Fig. 1(c)), VA (P = 0.029; Fig. 1(e)) and CLA (P = 0.030; Fig. 1(f)).

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Fatty acid and fat-soluble antioxidant concentrations in milk

Figure 1. Effect of organic (black bars) and non-organic (white bars) low-input production systems on the fatty acid composition of milk fat. (a) SFA, saturated fatty acids; (b) MUFA, monounsaturated fatty acids; (c) PUFA, polyunsaturated fatty acids; (d) ALA, α-linolenic acid; (e) VA, vaccinic acid; (f) CLA, conjugated linoleic acid isomer C18:2 c9 t11; * means for organic and non-organic low input systems are significantly different according to Tukey's honest significant difference test. Error bars indicate standard error of mean values. Two-way ANOVA (with production system and date as factors) identified significant differences (a) between production systems for VA ($P = 0.041$) and CLA ($P = 0.012$) and (b) between dates for PUFA ($P = 0.028$), VA ($P = 0.005$) and CLA ($P < 0.0001$). Significant interactions between system and date were identified for PUFA ($P = 0.020$), VA ($P = 0.029$) and CLA ($P < 0.300$).

Figure 2. Effect of organic (black bars) and non-organic (white bars) low-input production systems on the levels of fat-soluble antioxidants in milk fat. (a) 2R α-toc, 2R stereoisomers of α-tocopherol; (b) 3R α-toc, 3R stereoisomers of α-tocopherol; (c) total carotenoids; (d) β-carotene; (e) lutein; (f) zeaxanthin; * means for organic and non-organic low-input systems are significantly different according to Tukey's honest significant difference test. Error bars indicate standard error of mean values. Two-way ANOVA (with production system and date as factors) identified significant differences (a) between production systems for β-carotene ($P = 0.003$), lutein ($P = 0.004$), zeaxanthin ($P = 0.027$) and total carotenoids (0.002), and (b) between dates for 2R α-toc ($P = 0.0005$), 3R α-toc ($P = 0.0005$), β-carotene ($P = 0.005$), lutein ($P = 0.0008$), zeaxanthin ($P = 0.002$) and total carotenoids (0.003). A significant interaction between system and date was only identified for 2R α-toc ($P = 0.003$).

DISCUSSION AND CONCLUSIONS
Effect of feeding regimes on milk fat composition: outdoor grazing period

The finding of lower percentages of SFA and contrasting higher percentages of MUFA in milk from the NO-LI system and higher PUFA (specifically α-LA and CLA9) and antioxidant content (α-tocopherol and carotenoids) of milk from both LI systems, compared with that from HI farms during the outdoor grazing period, is not surprising in view of the contrasting diets. The two LI systems used a high level of fresh forage (>80% of DMI), with only half that level (<40%) used in HI systems. Increasing the level of fresh forage by similar margins was previously shown to elevate nutritionally desirable PUFA, CLA, α-LA and antioxidant percentages in milk.}

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found in milk from LI and HI systems here. For example, CLA concentrations were previously shown to increase with the proportion of fresh grass intake, while high proportions of maize silage and/or cereal-based concentrates reduced CLA content.\textsuperscript{17,18,33} Also cutting and transport of grass to housed animals (a practice used to increase milk yield in zero-grazing systems) was also shown to decrease the CLA and VA content of milk by 50% and that of αLA content by 30%, compared to milk from cows grazing pasture.\textsuperscript{34} This response may have been due to rapid lipolysis of PUFA after harvest and/or a modification of rumen biohydrogenation.\textsuperscript{27}

The finding that concentrations of CLA9 were significantly higher in milk from LI than HI systems, while concentrations of CLA10 were similar in both systems, was likely to be caused by contrasting effects of LI and HI diets on the biosynthesis of CLA9 which is mainly (70–90%) generated from VA in the mammary gland, and that of CLA10 which is a minor intermediate of rumen biohydrogenation.\textsuperscript{19}

Previous studies have shown that VA in the rumen increases with increasing fresh forage and decreasing concentrate levels in dairy diets, while CLA10 generation in the rumen is relatively unaffected by changes in the diet except at very high levels of concentrate feeding.\textsuperscript{17,20}

The greater dietary contribution from fresh forage is also the most likely explanation of elevated levels of RRR tocopherol and carotenoids in milk from the LI herds during the grazing period, compared to the HI milk. Transfer of β-carotene and α-tocopherol into milk was reported to be directly proportional to dietary supply, being highest in spring grazing.\textsuperscript{21}

\textbf{Effect of feeding regimes on milk fat composition: indoor period}

Few significant differences and trends in milk fat composition were found between HI and O-LI production systems during the indoor period when cows were fed conserved forage-based diets. This may have been due to feeding regimes used by O-LI and HI herds being more similar during the indoor compared with the outdoor feeding period. The higher SFAs and lower MUFA content of organic milk during this feeding period are difficult to explain, since previous studies have shown that fresh forage intake (24% in organic as opposed to none in conventional winter diets) increases dietary PUFA supply.\textsuperscript{20,27} However, some previous studies have reported lower biohydrogenation rates for high-concentrate indoor diets,\textsuperscript{17,20} suggesting that the higher proportion of concentrate in the HI diets results in lower biohydrogenation and thereby lower SFA and higher MUFA, and that this effect overrides the effect of higher fresh forage intake in the O-LI animals. In order to allow milk from organic or LI production systems to be marketed as having ‘added nutritional value’ throughout the year, efforts need to be made to achieve higher concentrations of at least some to the nutritionally desirable compounds during the indoor feeding period, if year-round grazing is not an option. This could be achieved by supplementation of conserved forage-based winter diet with oil seeds (e.g., rapeseed, linseed, sunflower seed), a practice shown to significantly improve α-LA, VA, CLA9 and/or fat-soluble antioxidant concentrations in milk.\textsuperscript{12,17,33–37}

Changes to the forage conservation methods may also increase the content of desirable FAs. For example, using hay rather than silage was also shown to increase the α-LA content in milk by up to 50%.\textsuperscript{33,36} It is interesting to note that in the UK it is very difficult to find farms feeding hay rather than silage, except among very traditional organic producers that work to biodynamic farming principles (which strongly recommend the use of hay for milking cows).

\textbf{Effect of vitamin feed supplements on antioxidant concentrations in milk}

Results of the study reported here suggest that the addition of synthetic vitamin/antioxidant supplements

\begin{table}[h]
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\begin{tabular}{llll}
\hline
Characteristic assessed & High-input & Low-input & ANOVA P-value \\
\hline
Number of samples & 21 & 10 & 0.0014 \\
Milk yield/cow (kg) & 26.5 (1.0) & 19.1 (1.3) & 0.041 \\
Protein content (g kg\textsuperscript{-1}) & 33.0 (0.3) & 33.1 (0.6) & 0.803 \\
Fat content (g kg\textsuperscript{-1}) & 40.8 (0.5) & 42.1 (0.7) & 0.235 \\
Fatty acid groups (g kg\textsuperscript{-1} milk fat) & & & \\
Total SFA & 712 (6) & 740 (11) & 0.028 \\
Total MUFA & 254 (5) & 228 (10) & 0.730 \\
Total PUFA & 53 (2) & 51 (4) & 0.114 \\
Omega 3 and 6 FA (g kg\textsuperscript{-1} milk fat) & & & \\
α-LA C18:3 c9 c12 c15 & 3.5 (0.3) & 7.3 (0.9) & 0.052 \\
γ-LA C18:3 c6 c9 c12 & 0.2 (0.02) & 0.2 (0.03) & 0.127 \\
Total n-6 & 21.7 (1.3) & 16.4 (0.7) & 0.018 \\
n-3:n-6 ratio & 0.30 (0.04) & 0.42 (0.06) & 0.139 \\
VA and CLA isomers (g kg\textsuperscript{-1} milk fat) & & & \\
VA C18:1 n11 & 16.4 (1.0) & 17.5 (2.3) & 0.636 \\
CLA C18:2 n9 n11 & 6.2 (0.04) & 7.8 (0.21) & 0.111 \\
CLA C18:2 n10 n12 & 0.31 (0.01) & 0.34 (0.02) & 0.139 \\
Fat-soluble antioxidants (mg kg\textsuperscript{-1} milk fat) & & & \\
α-Tocopherol & & & \\
2R α-toc & 3.5 (0.4) & 2.8 (0.4) & 0.360 \\
RRR α-toc & 20.4 (0.9) & 20.3 (1.5) & 0.776 \\
Total α-tocopherol & 23.9 (1.0) & 23.1 (1.6) & 0.513 \\
Carotenoids & & & \\
B-carotene & 5.49 (0.41) & 6.29 (0.64) & 0.359 \\
Lutein & 0.37 (0.03) & 0.48 (0.06) & 0.081 \\
Zeaxanthin & 0.12 (0.01) & 0.14 (0.01) & 0.265 \\
Total carotenoids & 5.98 (0.44) & 6.90 (0.68) & 0.314 \\
\hline
\end{tabular}
\caption{Fatty acid composition and fat-soluble antioxidant concentrations in milk from conventional high-input and organic and non-organic low-input dairy production systems, during the indoor conserved forage-based feeding period (mean values, with standard error of means in parentheses)}
\end{table}
to feed in HI systems has a relatively minor effect on antioxidant concentrations in milk. For example, milk from HI herds, which received high levels of vitamin E supplements (in our study between 450 and 750 IUs vitamin E per day) contained significantly lower concentrations of total α-tocopherol during grazing than milk from farms working to organic farming standards, which do not permit feed supplementation with synthetic vitamins. It is particularly interesting that the concentration of the 2R stereoisomer of α-tocopherol was not significantly higher in milk from the HI systems. The 2R stereoisomers account for most of the α-tocopherol in synthetic vitamin E supplements, but are virtually absent from natural sources of α-tocopherol such as forage. This indicates either poor uptake of the 2R stereoisomers in the gastrointestinal system and/or preferential/selective uptake/transfer of 3R stereoisomers from the blood into milk in the udder, as reported previously.22

**Potential effects of seasonal forage composition and availability on milk fat**

Differences in milk quality (both fatty acid profiles and antioxidant levels) were also detected between spring block calving O-LI and NO-LI systems which appeared to have very similar dietary regimes. These were more likely due to variation in the composition and/or total forage availability between the two systems over the season, since both systems grazed cows throughout the lactation and used very low levels of supplementary feeds such as conserved forage or concentrate. The finding that, in August, milk from O-LI systems had higher percentages of α-LA than milk from NO-LI systems is not surprising, and is likely to be due to a combination of two factors. Firstly, the use of mineral (especially N) fertilizers in the NO-LI system, a practice which has been shown to suppress the relative amounts of white clover in grass clover swards,39,40 and secondly, the impact of higher clover content causing elevation in concentrations of n-3 FAs in milk compared with ryegrass.27 However, it should be noted that most of the studies reviewed by Dewhurst et al.27 that compared the effect of clover and rye grass used ensiled forage, where reduced lipolysis in clover would have a greater influence over PUFA supply compared with fresh forage. The significantly higher CLA and antioxidants in milk from NO-LI systems are more difficult to explain, but may be related to differences in the nutritional composition of the herbage resulting from the grazing systems used (e.g., the length of time allowed for pasture regrowth between grazing periods), which has also been shown to affect the fatty acid composition of milk.27 Milk yields, protein and urea content in this study (data not shown) did not differ at times when differences in milk fat composition were detected between the two LI systems. This suggests that differences in milk fat composition were unlikely to be linked to contrasting energy or protein supply levels. However, since sward composition and total forage availability were not monitored in the study reported here this will have to be tested in future studies.

**Potential effects of dairy genotypes on milk fat composition**

The higher proportion of fresh forage in the dairy diet is likely to have been the main reason for the differences in milk composition. However, since contrasting dairy genotypes (breed index) were used in different production systems this may also have contributed to the differences in milk composition recorded between systems.

There is relatively little quantification of the effect of breed on fatty acid composition, although breed effects on CLA and antioxidant content have been reported to vary by up to 15–20% between breeds.21,35 This differential is considerably lower than the 60–99% for CLA9 and 30–140% for antioxidants measured between HI and LI systems recorded in this study.

The finding of substantial differences in milk fat composition between HI and LI systems during the outdoor grazing period, but similar milk composition during the indoor feeding period, also suggests that the differences in feeding regimes (rather than dairy genotypes) were the main factors responsible for the milk composition differences between systems. However, the exact influence of breed relative to dietary supply and possible interaction needs to be determined in future studies.

**Potential nutritional impacts of differences in milk fat composition**

Differences in nutritionally desirable FA and antioxidants between HI and LI systems during the grazing period were generally quite large (65% and 45% for α-LA, 60% and 99% for CLA9, 33% and 50% for α-tocopherol, 30% and 74% for β-carotene, 67% and 148% for lutein and 46% and 82% for zeaxanthin, for O-LI and NO-LI systems, respectively). This confirms previously published comparisons of conventional and organic, low-input production systems carried out in Germany, Italy and the UK.14–16

Consumption of milk and milk products from LI systems produced during this period may therefore contribute significantly to increasing the intake of these compounds in line with nutritional recommendations. Importantly, the higher percentages of nutritionally desirable PUFA (CLA9 and α-LA) found in milk from LI systems did not coincide with a significant increase in nutritionally less desirable PUFA (e.g., CLA10, total n-6 FA). Also, the higher n-3 FA and lower n-6 FA percentages found in milk from LI systems resulted in a higher n-3:n-6 FA ratio, which is also considered nutritionally desirable.4,10,12,27,41

Even if trends of elevated α-LA and lutein in organic milk produced during housing were confirmed, it is clear that consumption of organic milk produced during the indoor winter period will not increase the intake of nutritionally desirable compounds to the same extent as low-input milks produced during the outdoor grazing period.
While CLA9 and n-3 FA have been linked to a range of beneficial impacts on health,\textsuperscript{10–13} it should be pointed out that it is currently uncertain whether the main n-3 FA found in milk, $\alpha$-linolenic acid ($\alpha$LA; C18:3 $\text{c9 c12 c15}$), has similar effects on human health as the long-chain n-3 FAs found mainly in fish oil (C20 or longer), which have been shown to protect against coronary heart disease, associated with improved neurological function and linked to reduced risk of type 2 diabetes, hypertension and certain cancers.\textsuperscript{10,12,41,42} These long-chain n-3 fatty acids are known to be present at low levels in milk fat\textsuperscript{42} and were not determined in this study. However, there is now both direct and indirect evidence that significant levels of longer-chain n-3 FAs, especially eicosapentaenoic acid (EPA; C20:5 n-3) and to a lesser extent docosahexaenoic acid (DHA; C22:6 n-3), are generated from $\alpha$LA in humans.\textsuperscript{42}

The impact of fat-soluble antioxidants/vitamins on human health has been reviewed extensively.\textsuperscript{24,43–45} Beneficial effects of increased dietary $\alpha$-tocopherol (a compound belonging to the vitamin E group) intake to reduce oxidative stress, which was shown to be a risk factor for a number of chronic health conditions including cardiovascular disease, cancer, impaired immunity and premature ageing.\textsuperscript{45} Carotenoids can act as precursors for vitamin A, although a range of health benefits were linked to their antioxidant properties, and thought to be independent from their contribution to vitamin A generation.\textsuperscript{46}

With respect to the current availability of milk from LI systems for consumers, it should be emphasized that milk from organic producers is identifiable and widely available, while milk from the non-organically certified LI farms is currently mixed with milk from HI conventional systems in the supply chain and is not available to consumers. Given the apparently high nutritional quality of milk produced in NO-LI-systems it is important that this practice is reviewed in order to take advantage of the price premiums that can currently be achieved by ‘nutritionally enhanced’ food products.\textsuperscript{47}

When data for all sampling dates were pooled, the concentration of $\alpha$-LA was elevated by 60% and that of CLA9 by 64% in the organic compared to HI milk ($\alpha$-LA; mean = 9.4, SE = 0.3 \textit{versus} mean = 5.7, SE = 0.3 g kg\textsuperscript{-1} fat, \(P < 0.001\) and CLA9; mean = 12.2, SE = 0.7 \textit{versus} mean = 7.5, SE 0.4 g kg\textsuperscript{-1} fat, \(P < 0.001\) for O-LI and HI milk, respectively). These data may help explain why consumption of organic dairy produce has been shown to have a significant impact on the CLA content of breast milk in lactating women\textsuperscript{48} and on the eczema risk during the first 2 years of life.\textsuperscript{49} It is now important to (a) identify exactly those production system components in organic, LI and conventional farming systems that are responsible for differences in milk composition and (b) to allow agronomic strategies in dairy production to be optimized further with respect to compounds that can be linked to positive health impacts.

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Fatty acid and fat-soluble antioxidant concentrations in milk


