Increased muscle fatty acid oxidation in dairy cows with intensive body fat mobilization during early lactation


*Institute of Nutritional Physiology “Oskar Kellner,” Leibniz Institute for Farm Animal Biology (FBN), Wilhelm-Stahl-Allee 2, 18196 Dummerstorf, Germany
†Institute of Animal Science, Physiology & Hygiene Unit, University of Bonn, 53115 Bonn, Germany

ABSTRACT

The beginning of lactation requires huge metabolic adaptations to meet increased energy demands for milk production of dairy cows. One of the adaptations is the mobilization of body reserves mainly from adipose tissue as reflected by increased plasma nonesterified fatty acid (NEFA) concentrations. The capacity of the liver for complete oxidation of NEFA is limited, leading to an increased formation of ketone bodies, reesterification, and accumulation of triglycerides in the liver. As the skeletal muscle also may oxidize fatty acids, it may help to decrease the fatty acid load on the liver. To test this hypothesis, 19 German Holstein cows were weekly blood sampled from 7 wk before until 5 wk after parturition to analyze plasma NEFA concentrations. Liver biopsies were obtained at d 3, 18, and 30 after parturition and, based on the mean liver fat content, cows were grouped to the 10 highest (HI) and 9 lowest (LO). In addition, muscle biopsies were obtained at d −17, 3, and 30 relative to parturition and used to quantify mRNA abundance of genes involved in fatty acid degradation. Plasma NEFA concentrations peaked after parturition and were 1.5-fold higher in HI than LO cows. Muscle carnitine palmitoyltransferase 1α and β mRNA was upregulated in early lactation. The mRNA abundance of muscle peroxisome proliferator-activated receptor γ (PPARG) increased in early lactation and was higher in HI than in LO cows, whereas the abundance of PPARα continuously decreased after parturition. The mRNA abundance of muscle PPARδ, uncoupling protein 3, and the β-oxidative enzymes 3-hydroxyacyl-coenzyme A (CoA) dehydrogenase, very long-chain acyl-CoA dehydrogenase, and 3-ketoacyl-CoA was greatest at d 3 after parturition, whereas the abundance of PPARγ coactivator 1α decreased after parturition. Our results indicate that around parturition, oxidation of fatty acids in skeletal muscle is highly activated, which may contribute to diminish the fatty acid load on the liver. The decline in muscle fatty acid oxidation within the first 4 wk of lactation accompanied with increased feed intake refer to greater supply of ruminally derived acetate, which as the preferred fuel of the muscle, saves long-chain fatty acids for milk fat production.

Key words: dairy cow, lactation, transition period, muscle fatty acid oxidation

INTRODUCTION

The transition from gestation to lactation is associated with huge metabolic changes in dairy cows. Although energy demand increases up to 4-fold (Bobe et al., 2004), mainly due to enormous milk secretion, feed intake seems not to meet the resulting energy requirements. Energy intake may even decrease by parturition and increases only slowly in the first week of lactation, resulting in a negative energy and nitrogen balance (Bell et al., 2000; Drackley et al., 2001; Doepel et al., 2002; Kuhla et al., 2011). This deficiency is compensated for by the mobilization of body reserves, which contributes to supply lipids, proteins, and amino acids to the mammary gland. The key nutrient for milk production, however, is glucose and, thus, only a small amount of glucose is available as an energy source for peripheral tissues. During this state of hypoglycemia, lipids gain importance as the main energy source while lipogenesis is simultaneously reduced (Bauman and Currie, 1980; Grummer, 1993). The switch in systemic fuel selection is under endocrine control enabling large metabolic flexibility of many peripheral organs to support milk synthesis (Bauman and Currie, 1980).

The skeletal muscle is a major protein reserve and thus source of amino acids and its degradation may already start in late gestation (van der Drift et al., 2012). The extent of protein degradation, not only in muscle but also in other tissues, is supposed to be mirrored by plasma concentrations of branched-chain amino acids (BCAA) or 3-methyl histidine (3-MH; Doepel et al.,
Released amino acids are not only utilized for milk protein synthesis but are also required for hepatic gluconeogenesis (Bell et al., 2000). Triglycerides from adipose tissue are mobilized starting during the last week of gestation, resulting in an increased release of NEFA, which are oxidized by the liver and other tissues as an energy source (Grunmer, 1993). However, the hepatic capacity for NEFA utilization is limited, contributing to the development of fatty liver in early lactation. Thus, the oxidative capacity in other tissues such as skeletal muscle could diminish the metabolic load of FA on the liver, although muscle tissue itself is degraded.

One key regulator of the muscle’s oxidative metabolism is the phosphorylation status of the AMP-activated protein kinase (AMPK). Its activated (phosphorylated) form causes the repression of ATP-consuming processes and stimulates FA oxidation (FAO), whereas it lowers FA synthesis (Kahn et al., 2005). The FAO in skeletal muscle is also promoted by the adipokine leptin, which is predominantly produced by adipose tissue (Fuentes et al., 2010). Next to AMPK and leptin, other key regulators of FAO are well known. Most prominent is the peroxisome proliferator-activated receptor (PPAR) family (including the α, γ, and δ forms), with PPARδ being the most abundant member in skeletal muscle, but also PPARγ and its coactivator 1α (PPARGC1A) act as FA sensors and transcription factors regulating muscle FAO (Evans et al., 2004; Muoio and Koves, 2007; Moreno et al., 2010). Long-chain FA are transported into the mitochondria via the carnitine palmitoyltransferase 1 (CPT1) transporter system. In skeletal muscle, CPT1B is the predominant isoform (Price et al., 2003). Very long-chain acyl-CoA dehydrogenase (ACADVL; Zammitt, 1984) and, apart from its function in thermogenesis, uncoupling protein 3 (UCP3; Schrauwen and Hesselink, 2004) are also regulators for FA transport into and out of the mitochondria. Once in the mitochondria, FA are β-oxidized in a NADH-dependent process mediated by 4 major enzymes, including ACADVL. Elevated mRNA expression of these muscle β-oxidative enzymes has been described in early lactating sows (Ringseis et al., 2011), pointing to an increased muscle FAO, which should help to reduce the FA load on the liver. The extent to which the skeletal muscle of dairy cows participates in the degradation of FA during early lactation is not known. Our hypothesis was that skeletal muscle FA degradation adapts to the extent of fat mobilization during early lactation. To test this idea, we examined the expression of selected candidate genes at the mRNA level and their putative interaction with plasma metabolites in dairy cows with a different intensity of body tissue mobilization during early lactation.

**MATERIALS AND METHODS**

**Animals, Feeding, and Zootechnical Data**

All treatments were in accordance with the guidelines for the use of animals as experimental subjects of the State Government in Mecklenburg-Western Pomerania (Registration No. LALLF M-V/TSD/7221.3-2.1-021/09). The experimental design was described previously (Schäff et al., 2012). Briefly, 20 German Holstein cows in their second to fourth lactation were liver biopsied on d −34, −17, 3, 18, and 30 relative to parturition and grouped according to their mean liver fat content after parturition as a measure of body fat mobilization to the lowest [LO; liver fat content <24% (20.0 ± 2.0%) total fat/DM; n = 9] and highest [HI; liver fat content >24.4% (30.2 ± 1.8%) total fat/DM; n = 10] mobilizing cows. Due to its severe sickness during the trial, 1 LO cow was excluded from sampling.

Cows were housed in a tie-stall and fed twice daily (0600 and 1500 h) 1 of 3 different TMR according to the recommendations of the German Society of Nutrition Physiology (GfE, 2001). The TMR compositions are shown in Table 1. Samples of the individual silages used were taken weekly to determine the content of DM. Feed intake was recorded daily and BW was measured weekly to calculate DMI and energy balance (EB). Energy balance (expressed in MJ/cow × d) was calculated for the dry period as EB = NE_d intake − (0.293 × kg of BW0.75 + 0.044 × e0.0165 × days since conception ) (GfE, 2001). After parturition, cows were milked twice daily (0330 and 1500 h). Milk samples were taken weekly and analyzed for crude fat, CP, and lactose by an infrared spectrophotometric method (MilkoScan; Foss GmbH, Rellingen, Germany) to calculate ECM as follows:

ECM = [(0.038 × g of crude fat + 0.024 × g of CP + 0.171 × g of lactose) × kg of milk] ÷ 3.14 (Reist et al., 2001). Back fat thickness (BFT) and thickness of the musculus longissimus dorsi (MLDT) were measured weekly via ultrasound (SonoSite Inc., Bothell, WA).

**Blood Sampling and Analyses**

Weekly blood samples were taken from the jugular vein in EDTA-containing tubes. During the 2 wk before and 2 wk after parturition, 2 blood samples per week were obtained. Samples were immediately placed on ice and centrifuged within 30 min (at 1,565 × g for 20 min at 4°C). The obtained plasma was stored at −80°C until photometrical analyses for NEFA, BHBA, glucose, and lactate as described previously (Schäff et al., 2012).
Plasma leptin was determined by an enzyme immunoassay according to Sauerwein et al. (2004). Around parturition, weekly means were calculated from the 2 samples obtained. Plasma amino acid concentrations were measured in samples obtained in wk −4, −3, −1, 1, 2, 3, and 5 relative to parturition via HPLC (Schäff et al., 2012).

### Muscle Tissue Samples

Biopsies of the semitendinosus muscle were taken before morning feeding at d −17, 3, and 30 relative to parturition, alternating on the left and the right side (Kuhla et al., 2011). To this end, the skin was shaved, disinfected (Braunoderm; B. Braun Melsungen AG, Melsungen, Germany), and a sample was taken with a custom-made biopsy instrument as described earlier (Kuhla et al., 2011). Muscle tissue was liberated from skin and subcutaneous fat, immediately snap frozen, and stored at −80°C.

### Muscle Fat, Protein, Glycogen, and Glucose Content

Frozen muscle tissue was ground with a mortar and pestle under liquid N₂. Muscle tissue (50 mg) was dried in a muffle-type furnace at 105°C for 3 h to determine the DM content. Dried samples were analyzed for nitrogen and carbon content using a C/H/N/S TruSpec CHNS Microanalyzer (Leco Instrumente GmbH, Mönchengladbach, Germany). For the analysis of muscle glycogen, wet tissue (25 mg) was applied to an enzyme-based starch kit (no. 10207748035; Boehringer

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**Table 1.** Components and chemical composition of diets fed as TMR during the far-off dry period, close-up dry period, and early lactation

<table>
<thead>
<tr>
<th>Component</th>
<th>Far-off dry period¹</th>
<th>Close-up dry period¹</th>
<th>Early lactation</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ingredient (g/kg of DM)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Grass silage</td>
<td>744</td>
<td>302</td>
<td>202</td>
</tr>
<tr>
<td>Corn silage</td>
<td>37</td>
<td>399</td>
<td>350</td>
</tr>
<tr>
<td>Barley straw</td>
<td>115</td>
<td>54</td>
<td>16</td>
</tr>
<tr>
<td>Hay</td>
<td>92</td>
<td>39</td>
<td>26</td>
</tr>
<tr>
<td>Concentrate</td>
<td>1.4²</td>
<td>124²</td>
<td>296²</td>
</tr>
<tr>
<td>Molassed sugar beet pulp³</td>
<td>4.1</td>
<td>5.5</td>
<td>62</td>
</tr>
<tr>
<td>Extracted rapeseed meal</td>
<td>45</td>
<td></td>
<td>32</td>
</tr>
<tr>
<td>Mineral feed</td>
<td>7.7⁶</td>
<td>1.8⁵</td>
<td>6.7⁶</td>
</tr>
<tr>
<td>Nonlactating mineral feed with anionic salts⁷</td>
<td>12</td>
<td></td>
<td>0.7</td>
</tr>
<tr>
<td>Cattle salt⁶</td>
<td>1</td>
<td></td>
<td>8.8</td>
</tr>
<tr>
<td>Propylene glycol⁶</td>
<td>16</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Chemical analysis</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Utilizable CP¹⁰ (g/kg of DM)</td>
<td>126</td>
<td>137</td>
<td>163</td>
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<tr>
<td>Crude fat (g/kg of DM)</td>
<td>40</td>
<td>34</td>
<td>30</td>
</tr>
<tr>
<td>NEₑ¹⁰ (MJ/kg of DM)</td>
<td>5.8</td>
<td>6.5</td>
<td>7.1</td>
</tr>
<tr>
<td>NDF (g/kg of DM)</td>
<td>354</td>
<td>324</td>
<td>296</td>
</tr>
<tr>
<td>ADF (g/kg of DM)</td>
<td>202</td>
<td>140</td>
<td>135</td>
</tr>
</tbody>
</table>

¹Far-off dry period: wk 7 to 4 before parturition, close-up dry period: wk 3 before until parturition.  
²Universal 18/3 (Vollkraft): 20 to 40% cereals (triticale, rye, wheat, and barley), 25% rapeseed expeller, lower contents of malt germs, wheat gluten, wheat bran, peeled oat bran, beet pulp chips, molasses and glycerin, minerals, vitamins, 6.7 MJ of NEₑ/kg of DM, and 160 g of utilizable protein/kg of DM.  
³MF 2000 (Vollkraft, Güstrow, Germany): 33% extracted soy meal, 20% corn, 17% wheat gluten, 13% wheat, 8% extracted rapeseed meal, 5% sugar beet pulp, 2% sodium hydrogen carbonate, 1.3% calcium carbonate, 0.2% sodium chloride, 8.0 MJ of NEₑ/kg of DM, and 204 g of utilizable protein/kg of DM.  
⁴Trockenschnitzel (ATR Landhandel Arp, Thordsen, Rautenberg, Sollerupmühle, Germany): minerals, 7.3 MJ of NEₑ/kg of DM, and 153 g of utilizable protein/kg of DM.  
⁵Rinderstolz 9235 (Salvana Tiernahrung GmbH, Sparrieshoop, Germany): 75% crude ash, 4.5% calcium, 6% phosphorus, 10% sodium, 12% magnesium, and vitamins.  
⁶Rinderstolz 9522 (Salvana Tiernahrung GmbH): 92% crude ash, 20% calcium, 5% phosphorus, 6% magnesium, 8% sodium, and vitamins.  
⁷Anionen-Mix 1141(Salvana Tiernahrung GmbH): 14% calcium, 12.5% sulfur, 12% chlorine, 10% magnesium, 2% phosphorus, and vitamins.  
⁸esco SOLSEL Mineralleckstein mit Kupfer (esco GmbH & Co. KG, Hannover, Germany): 37% sodium, 1.6% calcium, 0.6% magnesium, and copper.  
⁹Propylenglycol USP (Dr. Pieper Technologie und Produktentwicklung GmbH, Wuthenow, Germany).  
mRNA Preparation and Real-Time Quantitative Reverse-Transcription PCR

Powdered muscle tissue (50 mg) was homogenized with a FastPrep 120 centrifuge (Thermo Electron Corp., Saint-Herblain Cedex, France). Total RNA was extracted with TRIzol Reagent (Invitrogen Corp., Carlsbad, CA). The integrity and quality of total RNA was confirmed upon gel electrophoresis on agarose gel stained with ethidium bromide (Carl Roth GmbH, Karlsruhe, Germany) and by the measurement of the optical density 260:280 ratio (i.e., the ratio of optical density at 260 nm to the optical density at 280 nm). The absorption ratio was between 1.8 and 2.0 for all samples. Concentrations were detected spectrophotometrically on a NanoPhotometer (Implen GmbH, Munich, Germany). For cDNA synthesis, 1 μg of RNA was reverse-transcribed with 200 U of M-MLV Reverse Transcriptase, RNase H Minus, Point Mutant (Promega Corp., Madison, WI) and 100 pmol of random primers (Invitrogen Corp.). The cDNA was purified using a High Pure PCR Product Purification Kit (Roche Diagnostics, Laval, QC, Canada) and stored at −80°C. Primer pairs were designed with Primer3 software version 0.4.0 (http://bioinfo.ut.ee/primer3-0.4.0/primer3/), unless otherwise mentioned in Supplemental Table S1 (available online at http://www.journalofdairyscience.org/). Real-time quantitative reverse-transcription (qRT)-PCR was performed on a LightCycler 2.0 instrument with FastStart DNA MasterPLUS SYBR Green I Master Mix (Roche Diagnostics GmbH, Mannheim, Germany) with 2 μL of cDNA. Each cDNA sample was analyzed in duplicate. To verify specific PCR products, a melting curve analysis program was performed after the last amplification cycle. Furthermore, product purity and size were confirmed by agarose gel electrophoresis and by sequencing (MegaBACE 1000; GE Healthcare, Freiburg, Germany). Result analysis was performed with LightCycler analysis software 4.05. Data were quantified according to the 2^−ΔΔCT method (where CT is the cycle threshold; Livak and Schmittgen, 2001), using splicing factor 3 subunit 1 (SF3A1), which was recommended earlier as the reference gene for normalization (Pérez et al., 2008) and found to be nonregulated in our samples.

Phosphorylated AMPK ELISA

Powdered muscle tissue (20 mg) was extracted in 300 μL of lysis buffer containing 1 mM EDTA, 0.5% Triton-X-100, 5 mM sodium fluoride, 6 M urea, 10 μg/mL leupeptin, 10 μg/mL pepstatin, 100 μM phenylmethanesulfonyl fluoride, 3 μg/mL aprotinin, 2.5 mM sodium pyrophosphate, and 1 mM activated sodium orthovanadate dissolved in PBS. Extracts were diluted 6-fold in buffer containing 1 mM EDTA, 0.5% Triton-X-100, and 5 mM sodium fluoride dissolved in PBS. The protein content was determined according to the Bradford method, with BSA as standard. Diluted extracts (100 μL) were applied to the ELISA DuoSet IC (DYC3528-2; R&D Systems Inc., Minneapolis, MN) to determine the concentration of phosho-AMPKα1 (Thr174).

Statistical Analysis

All data were evaluated by repeated measures ANOVA using PROC MIXED of SAS/STAT software in the SAS System for Windows, release 9.2 (SAS Institute, 2009). Fixed effects for all analyses were group, time, and group × time; for qRT-PCR, technical replicate was additionally added as fixed effect. For blood metabolites and zootechnical data, analyses were performed separately for the prepartal period (wk −7 to −1 relative to parturition), the postpartal period (wk 1 to 5 relative to parturition), the transition period (wk −3 to 3 relative to parturition), and over the entire time period (wk −7 to 5 relative to parturition). Least squares means were calculated and presented with their standard errors for the entire time period, unless otherwise indicated. Differences were declared as significant at P < 0.05 and as trends at 0.05 ≥ P < 0.10. The mRNA abundance and plasma NEFA concentration differences between d 3 and d −17 as well as between d 30 and d 3 were calculated for each cow separately, yielding Δ-values. These Δ-values were tested for Pearson correlations with PROC CORR of SAS (SAS Institute, 2009) and correlations with P ≤ 0.05 were considered as significant.

RESULTS

Zootechnical and Milk Variables

The DMI increased with time (P < 0.001; Figure 1), but did not differ between HI and LO cows. Body weight, BFT, and MLDT were higher in HI cows (P = 0.003, P < 0.001, and P = 0.02, respectively; Figure 1) and in both groups higher before than after parturition (P < 0.001). Energy balance was not different between groups but reached a minimum, whereas liver fat content peaked after parturition (P < 0.001; Figure 1). Milk yield (P = 0.78) and ECM yield (P = 0.39) did not differ between groups. The HI cows had higher...
milk fat content ($P < 0.05$), whereas LO cows showed higher milk urea content ($P = 0.01$) and tended to have more milk lactose ($P = 0.09$). Milk protein content did not differ between HI and LO cows ($P = 0.8$; Table 2).

**Plasma Metabolites Around Parturition**

Before parturition, plasma NEFA concentrations did not differ between groups but thereafter were higher in HI compared with LO cows (Table 3). Plasma 3-MH concentrations peaked around parturition ($P < 0.001$; Table 3) without differences between groups. Plasma leptin concentrations tended ($P = 0.10$) to be greater for the HI cows, both prepartum and for the entire study. Plasma BHBA, glucose, and lactate concentrations were not different between groups. However, BHBA concentrations significantly increased during the transition period and thereafter ($P < 0.001$), whereas glucose ($P < 0.001$), lactate ($P = 0.01$), and leptin ($P < 0.001$) concentrations decreased after parturition, with the latter showing a group × time interaction in the transition period ($P = 0.02$; Table 3). Plasma BCAA tended to be higher in LO cows, before and after parturition ($P = 0.08$; Table 3).

**Muscle Tissue and qRT-PCR**

Muscle fat content was higher in HI cows ($P = 0.04$), whereas muscle protein, glycogen, and glucose content were unaltered between groups ($P = 0.51, 0.46$, and 0.13, respectively; Table 4).

The *PPARA* mRNA abundance decreased over time ($P < 0.001$; Figure 2), with no differences between groups, whereas *PPARG* abundance increased from gestation to early lactation ($P = 0.02$) in HI but not in LO cows ($P = 0.04$; Figure 2). Abundance of *PPARD* was highest around parturition ($P < 0.001$) in both groups and tended to be higher in LO cows after parturition ($P < 0.07$), showing a group × time interaction ($P = 0.03$; Figure 2). Abundance of *PPARGC1A* markedly decreased after parturition ($P < 0.001$; Figure 2) in both groups. The isoforms *CPT1A* and *CPT1B* were upregulated in early lactation ($P < 0.001$; Figure 2) and a group × time interaction ($P = 0.05$) was observed for *CPT1B*. The mitochondria-specific FA-degrading enzymes 3-hydroxyacyl-CoA dehydrogenase (*HADH*), *ACADVL*, and 3-ketoacyl-CoA thiolase (*ACAA2*) showed no differences between groups but were highest in abundance around parturition ($P < 0.05$; Figure 2). Also, *UCP3* abundance increased from pregnancy to early lactation ($P < 0.001$) in both groups, but only in HI cows *UCP3* abundance remained elevated during the entire early lactation period (group × time interaction: $P = 0.02$; Figure 2). Phosphorylation of AMPK in muscle tissue did not differ between groups ($P = 0.6$) or with time ($P = 0.3$; Figure 2).

**Correlations Between mRNA Abundances and Plasma NEFA Concentrations**

From d −17 to 3, changes in *CPT1B* mRNA abundance correlated positively with *ACADVL* and *ACAA2*, the latter additionally correlated positively with *HADH* but negatively with changes in *PPARG* abundance. Changes in *UCP3* mRNA abundance correlated positively with *PPARGC1A* and NEFA (Figure 3). From d 3 to 30, significant correlations were found between changes in *CPT1B* and *ACAA2* and between changes in *ACAA2* and *PPARG* abundance. Furthermore, alterations in *CPT1B* abundance correlated positively with changes in *HADH* abundance. Expression changes for *CPT1A* were, in contrast to the prepartal period, positively correlated with changes in *PPARA*, *PPARD*, *PPARGC1A*, *UCP3*, and NEFA abundance. The latter additionally correlated positively with *PPARD*, *PPARGC1A*, and *UCP3* abundance.

**DISCUSSION**

Dairy cows investigated in the present study originated from one commercial farm before being brought to our institute where they were kept under identical environmental conditions. According to the LFC postpartum, we retrospectively grouped cows into HI and LO groups, whereas during the dry-off period, LFC did not differ between groups. However, HI cows had the greater BFT, muscle fat content, MLDT, and BW already before parturition, suggesting their greater potential to mobilize more body fat and body protein than LO cows. The extent of postpartum mobilization of subcutaneous fat, however, does not depend on the level of prepartum BFT, BCS, or BW (Janovick and Drackley, 2010; Weber et al., 2013). In the study by Kessel et al. (2008), it was shown that cows with an equal BFT and equal BW before calving may lose BW after calving to a different extent, likely because mobilization from other depots such as from visceral or inter- and intramuscular fat depots contributes to BW reduction. As the latter fat depots are hard to quantify, classifying cows with different extent of body fat mobilization based on LFC postpartum seems to be more appropriate, at least because LFC was positively related to BW changes but not to BFT changes after calving (Weber et al., 2013).

The mobilization of skeletal muscle protein reserves in dairy cows starts already in late gestation and reaches a maximum between wk 2 before and wk 4 after parturition (van der Drift et al., 2012). In accordance with
Figure 1. Dry matter intake (DMI), BW, back fat thickness (BFT), muscle (musculus longissimus dorsi) thickness (MLDT), energy balance (EB), and liver fat content (LFC) in cows with a low (○) or high (●) extent of body fat mobilization. Times of muscle biopsies taken are indicated with arrows. All parameters changed with time ($P < 0.001$) and differed between the highest-body-fat-mobilizing (HI) and lowest-body-fat-mobilizing (LO) cows for BFT, MLDT, EB, LFC ($P < 0.001$), and BW ($P = 0.003$) except DMI, which was equal in both groups ($P = 0.71$).
these earlier findings, we found a continuously decreasing MLDT, particularly for HI cows from wk 4 before until wk 4 after parturition. Despite the significantly greater MLDT in HI compared with LO cows, plasma 3-MH concentration, which has been proposed as a marker for muscle breakdown (Blum et al., 1985) did not differ between groups. Our finding is in contrast to earlier studies reporting higher 3-MH plasma concentrations in cows with a higher MLDT in the periparturient period (Doepel et al., 2002; van der Drift et al., 2012).

Another indicator suggested for muscle degradation or anabolism, respectively, may be BCAA (Nicastro et al., 2012). As high plasma BCAA concentrations go along with less protein breakdown (Zanchi et al., 2008) and LO cows tended to have higher BCAA concentrations, LO cows seemed to degrade less muscle protein during the periparturient period compared with HI cows. This difference is reflected by a different MLDT decrease in HI and LO cows in early lactation.

Subcutaneous BFT decreased after parturition. The extent was equal in cows possessing a different BFT before parturition, which agrees with an earlier study.

### Table 2. Milk yield and milk constituents of cows developing a low (LO) or high (HI) liver fat content (LFC) postpartum

<table>
<thead>
<tr>
<th>Variable, wk 2 to 5</th>
<th>LFC</th>
<th>ANOVA P-value</th>
<th>LFC</th>
<th>Time</th>
<th>LFC × time</th>
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</thead>
<tbody>
<tr>
<td>LO</td>
<td>SE</td>
<td>HI</td>
<td>SE</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Milk yield (kg)</td>
<td>41.72</td>
<td>1.70</td>
<td>41.15</td>
<td>1.59</td>
<td>0.78</td>
</tr>
<tr>
<td>Milk fat (%)</td>
<td>4.67</td>
<td>0.29</td>
<td>5.25</td>
<td>0.27</td>
<td>0.048</td>
</tr>
<tr>
<td>Milk protein (%)</td>
<td>3.13</td>
<td>0.08</td>
<td>3.11</td>
<td>0.08</td>
<td>0.84</td>
</tr>
<tr>
<td>Milk lactose (%)</td>
<td>4.81</td>
<td>0.05</td>
<td>4.72</td>
<td>0.05</td>
<td>0.087</td>
</tr>
<tr>
<td>Milk urea (mg/L)</td>
<td>219.09</td>
<td>16.35</td>
<td>174.11</td>
<td>15.27</td>
<td>0.012</td>
</tr>
<tr>
<td>ECM (kg)</td>
<td>44.60</td>
<td>1.85</td>
<td>46.31</td>
<td>1.72</td>
<td>0.38</td>
</tr>
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<td></td>
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</tbody>
</table>

1Data are presented as LSM with their SE.

### Table 3. Plasma metabolites of cows classified for either low (LO) or high (HI) liver fat content (LFC) postpartum calculated separately for the prepartal period (wk −7 to −1), the transition period (wk −3 to +3), the postpartal period (wk +1 to +5), and over the entire study (wk −7 to +5)

<table>
<thead>
<tr>
<th>Variable2</th>
<th>Period</th>
<th>LFC</th>
<th>ANOVA P-value</th>
<th>LFC</th>
<th>Time</th>
<th>LFC × time</th>
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<tbody>
<tr>
<td>BHBA (mmol/L)</td>
<td>Prepartal</td>
<td>0.39</td>
<td>0.04</td>
<td>0.37</td>
<td>0.04</td>
<td>0.56</td>
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<tr>
<td></td>
<td>Transition</td>
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<td>0.09</td>
<td>0.55</td>
<td>0.08</td>
<td>0.51</td>
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<tr>
<td></td>
<td>Postpartal</td>
<td>0.62</td>
<td>0.11</td>
<td>0.72</td>
<td>0.10</td>
<td>0.28</td>
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<td>0.23</td>
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<tr>
<td>Glucose (mmol/L)</td>
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<td>0.17</td>
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<tr>
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<tr>
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</tr>
<tr>
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<tr>
<td>BCAA (μmol/L)</td>
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<td>436.6</td>
<td>36.4</td>
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</tr>
<tr>
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</tr>
<tr>
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<td>5.94</td>
<td>0.53</td>
<td>7.43</td>
<td>0.78</td>
<td>0.10</td>
</tr>
</tbody>
</table>

1Data are presented as LSM with their SE.

23-MH = 3-methyl histidine; BCAA = branched-chain amino acids.
(Hammon et al., 2009). Accordingly, plasma NEFA concentrations peaked during early lactation and were higher in HI than in LO cows. During this time, released NEFA are partly used for the synthesis of milk fat, which was also higher in HI than in LO cows. The decrease in milk fat content parallels, therefore, the decrease in plasma NEFA concentrations (van Knegsel et al., 2007). The skeletal muscle may also contribute to the degradation of FA because intensive walking activity in postpartum dairy cows was negatively correlated with plasma NEFA concentrations (Adewuyi et al., 2006). Plasma leptin, a marker for triglyceride mobilization from adipose tissue, decreased after parturition in both groups and tended to be higher in HI compared with LO cows. These differences parallel the differences in BFT decrease and are in accordance with those of Hammon et al. (2009). Nevertheless, leptin’s role in stimulating FAO in skeletal muscle (Fuentes et al., 2010) around parturition seems to be marginal, because leptin concentrations decreased from late pregnancy to parturition and were not different between groups during transition (P > 0.15; Table 3), but this assumption needs to be confirmed in future studies (e.g., on leptin receptor signaling).

Long and very long-chain FA are transported into the mitochondria via CPT where they undergo β-oxidation. In muscle, the predominant form is CPT1B, which has a different enzyme kinetic [higher values of the Michaelis constant (Km) for palmitoyl-CoA and carnitine but lower Km values for malonyl-CoA] compared with CPT1A (Price et al., 2003). CPT1A and CPT1B were found upregulated in early lactation, whereas CPT1B was highest on d 3 and CPT1A was highest on d 30, suggesting an increased transport of FA from the cytosol into mitochondria in early lactation. The reduced CPT1B abundance from d 3 to 30 might be due to higher malonyl-CoA concentration, the latter the inhibitor for CPT1 and precursor for fat synthesis. Interestingly, muscle fat increased from d 3 to 30. The higher entrance of FA into the mitochondria was followed by an upregulation of the β-oxidative genes ACADVL and ACAAT2 but not HADH on d 3. Accordingly, strong positive correlations were observed between changes in ACADVL or ACAAT2, respectively, and CPT1B. However, it remains speculative if the increase in CPT1A and CPT1B mRNA abundances coincide with the postpartal NEFA peak because only 2 muscle samples were taken in early lactation.

Rising NEFA concentrations generally lead to increased mRNA abundance of the PPAR family, which are all key regulators of FAO or lipogenesis (Muoio and Koves, 2007; Brennan et al., 2009). We found increased abundance of the FA sensor PPARG after parturition which was higher in HI compared with LO cows, assuming a higher lipogenesis (Evans et al., 2004) in HI cows, although not reflected by the higher muscle fat content at d 30. In contrast, mRNA PPARA abundance decreased after parturition; however, PPARA is not the dominant form in muscle. The most abundant PPAR form in skeletal muscle is PPARD (Brennan et al., 2009), which may interact with PPARGC1A and thereby promotes complete FAO (Consitt et al., 2010). Although PPARD was highest in abundance on d 3, the abundance of PPARGC1A decreased after parturition, suggesting that either the PPARD promoting effect on FAO is not dependent on the interaction with PPARGC1A in early lactation, or that muscle FAO is incomplete during this time. Upregulation of PPARD on d 3 may promote the first steps of β-oxidation, whereas decreasing PPARD abundance by d 30 may rather impede complete FAO. Based on the group × time interaction for PPARD and CPT1B and the higher PPARD and CPT1B abundances in LO cows after parturition, we conclude that complete muscle

<table>
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<th>Variable</th>
<th>d</th>
<th>LO (%)</th>
<th>SE</th>
<th>HI (%)</th>
<th>SE</th>
<th>LFC</th>
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<tr>
<td>Muscle fat</td>
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<td>0.32</td>
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<td>1.42</td>
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<td>0.27</td>
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<td>0.11</td>
<td>0.07</td>
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<td>0</td>
<td>0</td>
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<tr>
<td>Muscle glucose</td>
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<td>0.51</td>
<td>0.59</td>
<td>0.03</td>
<td>0.13</td>
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<td>0</td>
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</tr>
</tbody>
</table>

Table 4. Muscle fat, protein, glycogen, and glucose content in percentage of wet weight of cows with a low (LO) or high (HI) liver fat content (LFC) postpartum.

1Data are presented as LSM with their SE.
Figure 2. Messenger RNA abundance of the peroxisome proliferator-activated receptors PPARA, PPARG, PPARD, the PPARG coactivator 1α (PPARGC1A), carnitine palmitoyltransferase 1α (CPT1A) and 1β (CPT1B), 3-hydroxyacyl-CoA dehydrogenase (HADH), very long-chain acyl-CoA dehydrogenase (ACADVL), 3-ketoacyl-CoA thiolase (ACAA2), and uncoupling protein 3 (UCP3), as well as the concentration of phosphorylated AMP-activated protein kinase (pAMPK) in skeletal muscle of cows with a different extent of body tissue mobilization [lowest-body-fat-mobilizing (LO) cows: white bars; highest-body-fat-mobilizing (HI) cows: black bars]. Messenger RNA abundance differed between HI and LO cows for PPARG (P < 0.05) and for all transcripts in time (for UCP3, PPARA, PPARD, PPARGC1A, CPT1A and CPT1B, and ACAA2: P < 0.001; for ACADVL: P < 0.01; for PPARG; P < 0.05) except for HADH (P = 0.7). A group × time interaction was found for UCP3 and PPARD (P < 0.05); PPARA and PPARG as well as CPT1B tended to interact between group and time (P = 0.08, P = 0.06, and P = 0.05, respectively). Muscle pAMPK was unaltered over time (P = 0.3) and between groups (P = 0.6). CT = cycle threshold.
FAO was more pronounced in LO than in HI cows during early lactation. Incomplete or limited FAO results in the formation of acylcarnitines or 3-hydroxy FA, thereby contributing to insulin resistance in peripheral tissues (Koves et al., 2008), which in turn facilitates net mobilization of amino acids from these tissues in the early lactation period (Bell et al., 2000).

Rising NEFA concentrations and the interaction between PPARD and PPARC1A may trigger the activation of UCP3 (Musa et al., 2012) from d −17 before to d 3 after parturition. Accordingly, we found changes of UCP3 mRNA abundance positively correlated with changes of NEFA concentrations throughout the periparturient period, as well as with changes of PPARC1A mRNA abundance before parturition. Considering that UCP3 is responsive to elevated fasting-induced NEFA concentrations in cattle (Brennan et al., 2009), the highest abundance of UCP3 together with PPARD and CPT1B on d 3 supports the idea that muscle metabolism switches from glycolytic to a more oxidative type by preferring FA as fuel (de Lange et al., 2007). Moreover, due to its primary role in transporting non-oxidizable FA out of the mitochondria, upregulation of UCP3 by d 3 may prevent myocytes from lipotoxicity during increased β-oxidation (Schrauwen and Hesselink, 2004). Taken together, upregulation of CPT1B, PPARD, UCP3, and β-oxidative enzymes from d −17 before to d 3 after parturition indicate induced muscle FAO. This conclusion is supported by the earlier described upregulation of muscle retinal dehydrogenase producing retinoic acid as a ligand for PPAR signaling in the second week of lactation (Kuhla et al. 2011). Therefore, increased muscle FAO around parturition could contribute in reducing the metabolic load of FA on the liver.

We further found a group × time interaction for UCP3. The higher abundance of UCP3 in HI than in LO cows in early lactation indicates a higher export of non-oxidizable FA from the mitochondria. Those non-oxidizable FA may be lipid peroxides that are produced in the reaction with reactive oxygen species generated during UCP3 mediated uncoupling (Hoeks et al., 2012). Considering the lower PPARD but higher UCP3 mRNA abundance in HI cows in early lactation, it seems that HI cows have the higher incomplete but lower complete muscle FAO. Similarly, myocytes from humans with a high fat load had the lower complete and the higher incomplete FAO compared with lean counterparts (Consitt et al., 2010).

From d 3 to 30 of early lactation, however, UCP3 mRNA abundance decreased again. Similarly, UCP3 mRNA abundance is reduced in the muscle of early lactating compared with nonpregnant rats (Xiao et al., 2004). This decrease is accompanied by a reduction in PPARA, PPARD, and CPT1B mRNA abundance by d 30, indicating decreasing muscle FAO, whereas plasma NEFA concentrations remain elevated. A limited increasing or decreasing muscle FAO by d 30 of lactation may save and direct FA for milk production (de Lange et al., 2007), but also allows the fat content of muscle tissue to increase again from d 3 to 30. This repletion of muscle fat is either due to reesterification of long-chain FA or de novo synthesis from acetate, which is sufficiently supplied by the increase in feed intake in

**Figure 3.** Correlations between differences (Δ) in mRNA abundance or plasma NEFA concentrations occurring from d −17 to 3 (A) or from d 3 to 30 (B). Solid lines indicate direct correlations, whereas dashed lines indicate inverse correlations, with $P ≤ 0.05$. ACAA2 = 3-ketoacyl-CoA thiolase; PPAR = peroxisome proliferator-activated receptor; PGC1α = PPARγ coactivator 1α; UCP3 = uncoupling protein 3; CPT1 = carnitine palmitoyltransferase 1; ACADVL = very long-chain acyl-CoA dehydrogenase; HADH = 3-hydroxyacyl-CoA dehydrogenase.
early lactation. Consequently, the reduced long-chain FA and also glucose utilization by the skeletal muscle in early lactation (Kuhla et al., 2011) could be compensated by increased acetate utilization as the preferred fuel source. Intragastrically administered acetate was shown to induce the expression of UCP and PPAR and CPT1 in the liver of mice (Kondo et al., 2009), but this effect has not been observed for the muscle in the present study. How UCP3, CPT, and PPAR are regulated in response to an increased acetate supply in the bovine muscle remains to be determined.

Another major player promoting fat oxidation is AMPK. Phosphorylated in times of energy deficiency, phosphorylated AMPK signals the inhibition of acetyl-CoA carboxylase, thereby suppressing malonyl-CoA synthesis and thus enabling FA transport via CPT1 in skeletal muscle (Kahn et al., 2005). However, we found no differences in AMPK phosphorylation between groups or during the periparturient period, suggesting that during this time, the muscle is sufficiently supplied with ATP, either via glycolysis, FAO, or by acetate oxidation. The switch in fuel selection of the skeletal muscle together with the degradation and repletion of muscle energy reserves during the periparturient period indicates the great metabolic flexibility of the muscle. This integrates the muscle into homeorhesis, which has been defined as the orchestrated and coordinated control of body tissue metabolism aligned to partition nutrients and energy to the dominant process of milk production (Bauman and Currie, 1980).

CONCLUSIONS

Taken together, we found the highest mRNA abundance of genes involved in skeletal muscle FA transport and oxidation 3 d after parturition. During this time, plasma NEFA concentrations still increased, reaching a maximum later in early lactation. Although mRNA abundance does not necessarily reflect protein abundance or enzyme activity, our results led us assume that transport of FA into muscle mitochondria and subsequent FAO respond to fat mobilization around parturition and thus help to protect the liver from excess fat load. The higher extent of fat mobilization, however, is associated with increased incomplete and reduced complete muscle FAO. From d 3 to 30 of lactation, mRNA abundance of genes involved in muscle FAO decreases, supporting repletion of muscle fat and sparing FA for milk fat synthesis.

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