

# Towards a better understanding of the respective effects of milk yield and body condition dynamics on reproduction in Holstein dairy cows

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*The overall reproductive performance has decreased over the last decades, involving changes in cyclicity, oestrous behaviour and fertility. High milk yield (MY), low body condition score (BCS) and large body condition (BC) loss have been identified as risk factors. However, these effects are often confounded, as high MY and body lipid reserve mobilization are correlated. The aim of this study was to evaluate the respective effects of MY and BC on post-partum ovarian cyclicity, oestrus and fertility of Holstein cows. This study provides novel information, as MY and BC change were uncoupled in the overall dataset that included 98 lactations and milk progesterone profiles. Cows were assigned to two feeding-level groups: high feed, which achieved high MY and moderate BC loss throughout lactation (8410 kg, –1.17 unit from calving to nadir BCS), and low feed, which limited MY and triggered a large BC loss (5719 kg, –1.54 unit). MY and BC had different effects at different stages of the reproductive process. Cyclicity as well as non-fertilization and early embryo mortality were mainly driven by body lipid reserves, whereas oestrous behaviour and late embryo mortality were related to MY. The results point to possible uncoupling between cyclicity, oestrus and early and late embryo survival allowing compensation along the reproductive process and leading to similar final reproductive performance. In compact calving systems, which require high pregnancy rates within a short period, higher MY strategies appear unsuitable even where BCS is maintained, owing to depressed oestrous behaviour and probably increased late embryo mortality, which delays rebreeding. Similarly, strategies that compromise cyclicity and fertility by excessively low BCS are unsuitable.*

**Keywords:** dairy cows, reproduction, progesterone, milk yield, body condition

## Implications

The objective of this study was to improve our understanding of the respective effects of milk yield and body condition on *post-partum* ovarian activity, oestrus and fertility in Holstein dairy cows, whereas the originality of the dataset lies in an uncoupling of these two highly correlated factors. Cyclicity and early embryo survival appeared to be mainly driven by body lipid reserves, whereas oestrus and late embryo survival appeared to be mainly driven by milk production. These different relationships should be taken into account by physiologists to improve biological knowledge of dairy cows' reproduction and by zootechnicians to optimize breeding strategies.

## Introduction

Reproductive performance has decreased over the last 5 decades in dairy cows as milk production increased

(Rodriguez-Martinez *et al.*, 2008). All the stages of the reproductive process have been affected, with increased ovarian abnormalities (Royal *et al.*, 2000), depressed oestrus duration and intensity (Van Eerdenburg *et al.*, 1996; Kerbrat and Disenhaus, 2004), depressed AI success (Barbat *et al.*, 2005) and an increase in both early and late embryonic death (Diskin *et al.*, 2006; Humblot *et al.*, 2009).

Adverse effects of high milk yield (MY) were observed in many studies. High MY has been reported to delay first ovulation (Petersson *et al.*, 2006), to increase the risk of prolonged luteal phases (PLP; Opsomer *et al.*, 2000), to reduce standing oestrus duration (Lopez *et al.*, 2004) and to depress fertility (Mackey *et al.*, 2007) by increasing late embryo mortality (Grimard *et al.*, 2006). Some of these effects may be attributed to genetics, as genetic correlations between MY and several reproductive traits have been established (Boichard *et al.*, 2002; Royal *et al.*, 2002). More generally, body condition score (BCS) and body condition (BC) loss resulting from the negative energy balance after

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calving are thought to explain these adverse effects of high MY (Cutullic, 2010; Friggens *et al.*, 2010).

However, in most studies, MY and BC loss effects are confounded. Comparison of high-fed animals (high milk, moderate BC loss) with low-fed animals (low milk, large BC loss) can help to distinguish these effects, and Cutullic *et al.* (2011), using this design, observed that almost all the stages of the reproductive process appeared to be sensitive to feeding level in Holstein cows. Working on the same Holstein dataset, this study aims to quantify the respective effects of MY and BCS at each stage of the reproductive process.

## Material and methods

### Experimental design and dataset

From 2006 to 2008, a breed  $\times$  feeding system experiment was conducted on the INRA experimental farm in Le Pin-au-Haras (Normandy, 48.44°N, 0.09°E). The results per breed and feeding system are reported in another paper (Cutullic *et al.*, 2011). This trial is limited to Holstein dairy cows. Each year, 34 Holstein cows were equally distributed within two winter compact calving systems. At the beginning of the experiment, systems were balanced for parity, calving date, genetic indices for MY, fat and protein content, BCS, BW and previous rank of successful insemination. For each trait, the estimated genetic indices were evaluated using a Best Linear Unbiased Prediction animal model especially developed for this experiment by Larroque and Boichard (personal communication). The model included the sire and grandsire's genetic evaluation, the dam's performance over three lactations, the classical fixed environmental effects and the feeding treatment. Cows remained in their feeding group until culling (not pregnant, health problem or death). Replacement primiparous cows were balanced for genetic indices, BCS, BW and rank of successful insemination.

The yearly calendar experimental design is reported in Figure 1. The two feeding systems were grass-based: a high feed system (allowing high MY for a grass-based system and limiting BC loss) and a low feed system (without any concentrate, limiting MY and triggering a large BC loss).

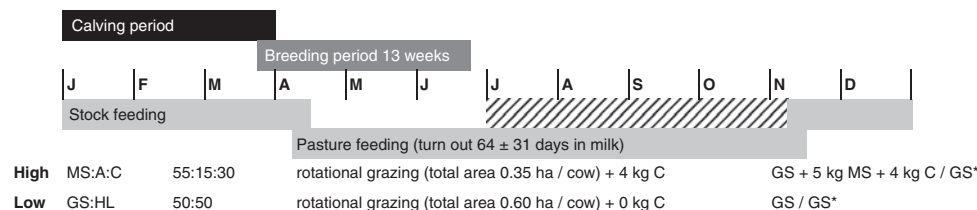
The cows were milked twice daily at 0630 and 1600 h. Individual MYs were recorded by flow meters (Metatron, Westfalia, Germany). Both morning and evening milk samples were collected and analysed for fat and protein content

three times a week using an infrared analyser (MilkoScan<sup>TM</sup>, Foss Electric, Hillerød, Denmark). From calving to 5 weeks after the end of the breeding period, morning milk samples were additionally collected on Monday, Wednesday and Friday, stored at  $-18^{\circ}\text{C}$  without preservative and assayed for progesterone ( $P_4$ ) using commercial ELISA kits (years 1 to 2: Ovucheck Milk, Biovet, Canada; year 3: Milk Progesterone ELISA, Ridgeway Science Ltd, England). From  $P_4$  profiles, commencement of luteal activity (C-LA), interovulatory intervals (ioi, estimate of cycle length), luteal phase lengths (lut) and interluteal intervals (ili) were determined (for details, see Cutullic *et al.*, 2011). The first ovulation was considered to be delayed if C-LA  $>50$  days. The luteal phase following the  $n$ -th ovulation was considered to be a PLP if  $\text{lut}_n > 25$  days. The  $n$ -th ovulation ( $n \geq 2$ ) was considered to be delayed if  $\text{ili}_{n-1} > 12$  days. The cycle following the  $n$ -th ovulation was considered to be a short cycle if  $\text{ioi}_n \leq 15$  days. *Post-partum* (*pp*)  $P_4$  profiles were classified as normal, PLP profile (if at least one PLP was observed), delayed (if C-LA  $>50$  days), interruption (if at least one ovulation of rank  $\geq 2$  was delayed) or disordered (if luteal activity appeared irregular but without any of the previously indicated abnormalities).

Caudal vein blood samples were taken at 20 and 60 days in milk and at first insemination to assay non-esterified fatty acids and glucose plasma concentrations, using enzymatic colorimetry (Kone Instruments Corporation, Espoo, Finland).

BCS was evaluated monthly, always by the same two experienced assessors. It was scaled from 0 to 5, from extremely thin to extremely fat, in increments of 0.25 (Bazin *et al.*, 1984).

During the entire breeding season and the 4 previous weeks, experienced staff observed the cows at least 5 times a day and behaviours were noted down on a standardized card. An oestrus detection aid was used in 2007 and 2008 (Kamar Heat Mount Detector, KAMAR Inc., Steamboat Springs, CO, USA). For analyses, oestri detected by standing to be mounted (specific behaviour) were distinguished from oestri detected by proceptive and general signs (mounting, sniffing, licking, chin-resting, restless, clear mucous discharge, heat mount detector only). This information was linked with  $P_4$  profiles: (i) to classify oestri as 'true' if corresponding to an ovulation period or 'false' and (ii) to determine the ovulation detection rate. For each undetected ovulation, that is, not matching with a detected oestrus, an expected date of oestrus was



**Figure 1** Yearly calendar experimental design of the feeding system experiment conducted in the INRA experimental farm Le Pin-au-Haras, comparing a high feeding level system with a low feeding level system. MS, A, C, GS, HL: maize silage, dehydrated alfalfa pellets, concentrate, grass silage, haylage. Salt and mineral were included in the concentrate of high-fed cows and were additionally fed to low-fed cows. GS\*: during the dry period, cows were fed *ad libitum* with grass silage only. Hatched period: if the grass offered was insufficient, cows received additional MS (high) or GS (low) feeding (up to 8 kg/day), mostly in the high group, which had a 1.7 smaller maximal grazing area.

calculated. For each ovulation, we considered that there was another cow in oestrus at the same time if there was at least one other cow in true oestrus on the same or the previous day.

During the breeding period, artificial insemination (AI) was systematically used on spontaneous oestri after 42 days in milk. By combining progesterone assays and ultrasonography at 40 and 70 days after insemination, AI results were classified into four categories according to Humblot (2001): non-fertilization or early embryo mortality (low  $P_4$  before 25 days after AI); late embryo mortality (high  $P_4$  at 25 days, low before 50 days); foetal death or abortion (ultrasonography positive twice or  $P_4$  high more than 50 days and ultrasonography positive once); and recalving. In statistical analyses, the embryo survival category included the two categories recalving and foetal death or abortion following insemination.

All the major health problems were recorded throughout lactation with a focus on reproduction problems (calving difficulties, caesarean, retained placenta, metritis, etc.). In the case of a caesarean, retained placenta or abnormal involution, prostaglandin analogue injections (cloprostenol, Intervet-Schering Plough, Beaucouzé, France) were used. Severe metritis (purulent and foul-smelling vaginal discharge) was treated with an intrauterine infusion (Intervet-Schering Plough, Beaucouzé, France) and a cloprostenol injection. Anoestrous cows received no specific treatment.

#### Statistical analyses

**MY and BCS relationship.** For 98 lactations, the effects of feeding level on MY and BCS variables were investigated using variance–covariance analysis. The models included the

effects of lactation number (categorized as first, second, third or more), feeding level, year, first-order interactions with feeding level and MY index as a covariate for MY variables (Model MY-1). The relationship between BC change from calving to 120 days in milk ( $\Delta BC_{120}$ ) and peak MY was investigated using variance–covariance analysis. A first model included the factors lactation number (first, second, third or more), year, BCS at calving, feeding level and peak MY, and also the interaction between these two variables (Model MY-2). A second model did not include feeding level and its interaction with peak MY (Model MY-3).

**Variables influencing reproductive outcome.** The effects of independent variables on reproductive outcome were assessed using a forward stepwise regression approach. Unless stated otherwise, only variables that significantly improved the model were included ( $P < 0.05$ ). At each step, changes in the  $P$ -values of type III ANOVA tables were checked to detect confounding effects and we checked in the final models that significant variables remained significant regardless of which statistical unit of the population was removed. The investigated outcome variables relating to cyclicity, oestrus and fertility stages are listed in Table 1. They were analysed in a chronological manner in Model CYC-1-2-3, Model OES-1-2 and Model FER-1-2. Continuous output variables were analysed with linear models and categorical output variables with logistic regressions, either binomial (two output categories) or multinomial (three output categories). Overall, three kinds of independent variables were investigated in these models: general variables and lactation-level variables for all models, and event-level

**Table 1** Investigated reproductive outcome variables, their related population and raw characteristics (Models CYC, OES, FER)

Outcome variables	Model	<i>n</i>	Population	Mean $\pm$ s.d. or categories %
<b>Cyclicity</b>				
Log <sub>e</sub> (C-LA)	CYC-1	98	All lactations	3.53 $\pm$ 0.55 (i.e. 34.1 days)
PLP profile	CYC-2	98	All lactations	PLP (23%) v. not
Cycle type	CYC-3	189	Ovulations of ranks 1 to 4 not followed by an AI, $\leq$ 135 days in milk, not belonging to a disordered profile and not followed by a delayed ovulation <sup>1</sup>	Normal (72%) v. PLP (14%) v. short (14%)
<b>Oestrus</b>				
Ovulation detection	OES-1-1bis	255	Ovulations of ranks above 2 <sup>2</sup>	Detected (69%) v. not
Ovulation detection by standing oestrus	OES-2	241	Ovulations of ranks above 2 and with known behaviour <sup>2,3</sup>	Detected (44%) v. not
<b>Fertility</b>				
Embryo survival	FER-1	128	First and second AIs performed on true oestri <sup>4</sup>	Non-fert.–early embr. m. (39%) v. late embryo mortality (20%) v. embryo survival (41%)
Recalving	FER-2-2bis	128	First and second AIs performed on true oestri <sup>4</sup>	Recalving (37%) v. not

C-LA = commencement of luteal activity; PLP = prolonged luteal phase; AI = artificial insemination; non-fert.–early embr. m. = non-fertilization or early embryo mortality.

<sup>1</sup>Ovarian activity interruption and disordered profiles were too scarce ( $n = 3$  and  $1$ ) to be specifically analysed.

<sup>2</sup>First *post-partum* ovulations are known to be hardly detectable (17% detected v. 69% for higher ranks in this study,  $P < 0.001$ ) and were not included in ovulation detection analyses.

<sup>3</sup>14 detected ovulations with unknown behaviour were excluded.

<sup>4</sup>17 inseminations performed on false oestri were excluded.

variables only for specific models (Table 2).  $\log_e(\text{C-LA})$  was also tested for the occurrence of PLP profiles (Model CYC-2), as early resumption of luteal activity is often associated with PLP (Opsomer *et al.*, 2000). For ovulation detection models (Model OES-1-2), observation period (before or during the breeding period and the 4 previous weeks) was included in the model to account for possible observation biases. For insemination result models (Model FER-1-2), parity and calving–AI interval ( $<$  or  $\geq 50$  days after calving) were also included in the model considering their known influence on fertility. Owing to the first results, we additionally tested the effects of the parameters of fitted Wood lactation curves over the first 14 weeks of lactation ( $\text{MY} = a \times \text{week}^b \times \exp^{-c \times \text{week}}$ ) on insemination results (Model FER-1). Feeding-level effect was not directly included in these models (CYC, OES, FER), as its effect is strongly correlated with the effects of

BCS and MY variables. In addition, for normal, PLP and delayed profiles, MY, BCS and BC change curves were compared using variance analysis; models included parity and profile category (Model CYCBCS).

*Linking cyclicity, oestrus and fertility.* In order to test the potential additional effects of cyclicity on oestrus and of cyclicity or oestrus on fertility, the following variables were investigated by including them in the two final models predicting ovulation detection (Model OES-1) and insemination results (Model FER-2):  $\log_e(\text{C-LA})$ , normal *v.* abnormal  $P_4$  profile, normal *v.* PLP *v.* other  $P_4$  profile, previous cycle length  $\leq 15$  *v.*  $> 15$  days, previous luteal phase  $\leq 25$  *v.*  $> 25$  days, current ili  $\leq 12$  *v.*  $> 12$  days, standing oestrus *v.* not. The new final models were labelled Model OES-1bis and Model FER-2bis, respectively (Table 1).

**Table 2** Independent variables included in statistical analyses to study their effectiveness to predict reproductive outcome variables (Models CYC, OES, FER)

General variables investigated in Models CYC, OES and FER	
Parity	Primiparous/multiparous
Year	2006/2007/2008
Genital health problem at/following calving (severe metritis, placenta retention, vaginal infection)	Yes/no
Lactation-level variables investigated in Models CYC, OES and FER	
<i>Genetic</i>	
MY index, MP content index, MF content index	MYindex ...
MY	MY <sub>3</sub> ... MY <sub>4 to 6</sub> ... MY <sub>max</sub>
MY (or FCMY) over weeks 1 to 3, 1 to 6, 1 to 14, 4 to 6, 7 to 14, at peak	
Milk content	
MP content over weeks 1 to 3, 1 to 6, at nadir	MP <sub>3</sub> ... MP <sub>min</sub>
MP/MF ratio over weeks 1 to 3, 1 to 6, at nadir	MP/MF <sub>3</sub> ... MP/MF <sub>min</sub>
BCS/BC change ( $\Delta\text{BC}$ )	
BCS 30 days before calving, at calving, 30, 60, 90, 120 days after calving, at nadir	BCS <sub>-30</sub> , BCS <sub>0</sub> ... BCS <sub>min</sub>
BC change from calving to -30, 30, 60, 90, 120 days, or to BCS <sub>min</sub> (can be BCS <sub>0</sub> )	$\Delta\text{BC}_{-30}$ , $\Delta\text{BC}_{30}$ ... $\Delta\text{BC}_{max}$
BC change from 30 to 90, 60 to 120 days after calving	$\Delta\text{BC}_{30 \text{ to } 90}$ , $\Delta\text{BC}_{90 \text{ to } 120}$
Metabolic status	
NEFA and glucose plasma concentrations at 20 and 60 d after calving	NEFA <sub>20</sub> , NEFA <sub>60</sub> ...
Event-level variables investigated in Models CYC-3, OES and FER only	
Days after calving	–
Pasture access	Yes/no (indoors only)
Ovulation rank	First/second or later
Observation during the breeding period or the 4 previous weeks	Yes (during)/no (before)
MY	
MY (or FCMY) over week(s) -2 to 0, -1 to +1, 0, 0 to +2 relative to event	MY <sub>week -2 to +2</sub> ...
MY dynamic	
MY (or FCMY) slope weeks -2 to 0, -2 to +2, -1 to +1, 0 to +2	MYslope <sub>week -2 to +2</sub> ...
Milk content	
MP content over weeks -2 to 0, -1 to +1, 0 to +2	MP <sub>week -2 to +2</sub> ...
MP/MF ratio over weeks -2 to 0, -1 to +1, 0 to +2	MP/MF <sub>week -2 to +2</sub> ...
BCS/BC change	
BCS at event (linear interpolation)	BCS <sub>event</sub>
BC change from calving to event	$\Delta\text{BC}_{event}$
BC dynamic	
BC change per 30 days from -30 <sup>1</sup> to +30, -30 <sup>1</sup> to 0, -15 <sup>1</sup> to +15, 0 to +30 relative event (day 0)	BCslope <sub>event -30 to +30</sub> ...
Metabolic status	
NEFA and glucose plasma concentrations at first and second inseminations	NEFA <sub>AIr</sub> , glucose <sub>AI</sub>

MY = milk yield (kg/day); MP = milk protein content (g/kg); MF = milk fat content (g/kg); FCMY = fat corrected milk yield (kg/day, milk fat content = 40 g/kg); BCS = body condition score (0 to 5 scale); BC = body condition; NEFA = non-esterified fatty acids.

<sup>1</sup>Left censored at calving to avoid a smoothing of the BC loss that occurs just after calving: for  $-a$  to  $+b$  case, the value for an event at time  $t$  is  $[\text{BCS at } t+b - \text{BCS at } \max(0, t-a)]/[t+b - \max(0, t-a)] \times 30$  to have a change expressed as BCS unit per 30 days.



**Statistical software and outputs.** Analyses were performed using the R statistical software (R Development Core Team, 2009; functions `lm`, `glm` and `multinom` for linear models, binomial logistic regressions and multinomial logistic regressions, respectively). When predicting a continuous output variable, linear model results are expressed as least-squared means for categorical independent variables or as slopes for continuous independent variables. When predicting categorical output variables with either two or three categories, the results are expressed as odds ratios for categorical independent variables (i.e.  $\exp(\text{coefficient of the linear predictor})$ ) or as transformed odds ratios for continuous independent variables. The transformed odds ratios reflect the effect of a 1 s.d. increase in the independent variable instead of the effect of a 1 unit increase (i.e.  $\exp(\text{coefficient of the linear predictor})^{\text{s.d.}}$ , with s.d. being calculated on the sample population, instead of  $\exp(\text{coefficient of the linear predictor})^1$ ). This transformation simplifies effect comparisons, as, for example, a 1 unit increase in peak MY is not comparable with a 1 unit increase in BCS at calving. Odds ratios interpretation remains the same: an odds ratio  $>1$  indicates that an increase in the independent variable increases the risk of the occurrence of a specific category rather than the occurrence of the reference category, whereas an odds ratio  $<1$  indicates that an increase in the independent variable decreases this risk. For example, in a binomial logistic regression predicting the occurrence of either category *B* or reference category *A*, an odds ratio  $>1$  indicates an increase in the risk of occurrence of category *B* rather than category *A*. In a multinomial logistic regression predicting the occurrence of category *C*, category *B* or reference category *A*, an odds ratio (*C* v. *A*)  $>1$  indicates an increase in the risk of occurrence of category *C* rather than category *A* and an odds ratio (*B* v. *A*)  $>1$  indicates an increase in the risk of occurrence of category *B* rather than category *A*. In these multinomial models, the reference category is thus important. To improve the understanding of the results of multinomial logistic regressions, the probabilities of each category were additionally calculated and plotted along the independent variable, so that  $\text{prob}(A) + \text{prob}(B) + \text{prob}(C) = 1$  at each given value of the independent variable.

## Results and discussion

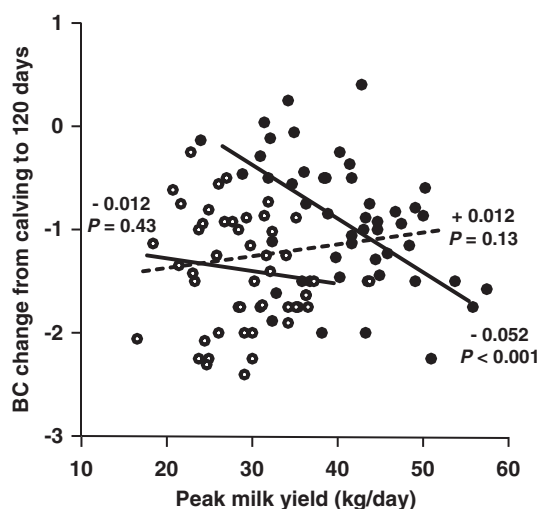
### *The link between MY and BC dynamic was disrupted in the entire dataset*

Cows in the low feed lost more BC, faster and for a longer period than those in high feed although they produced much less milk (Table 3; MY and BCS curves are reported by Cutullic *et al.*, 2011). The negative relationship between MY and BC change was observed within high feed (Figure 2). In low feed, BC loss was important irrespective of the MY. When the feeding level and the interaction between feeding level and MY are not included in the model, MY has no significant effect on BC loss: in the entire population, the link between MY and BC change was disrupted. In studies investigating the relationships between reproduction and MY or BC loss, these two factors are often confounded. Our

**Table 3** Adjusted MY and BCS (0 to 5 scale) for Holstein cows subject to a either high or low feeding level during lactation (Model MY-1)

	Feeding		Model		
	High	Low	r.s.e.	$R^2$	$P_{\text{feeding}}$
No. of lactations	50	48	–	–	–
44 weeks MY (kg)	8410	5719	851	79%	***
Peak MY (kg/day)	41.5	29.3	4.3	79%	***
BCS at calving	2.98	2.96	0.42	41%	0.85
BCS at nadir	1.81	1.42	0.50	33%	***

MY = milk yield; BCS = body condition score; r.s.e. = residual standard error. \*\*\*, \*\*, \*, †  $P < 0.001, 0.01, 0.05, 0.10$ .

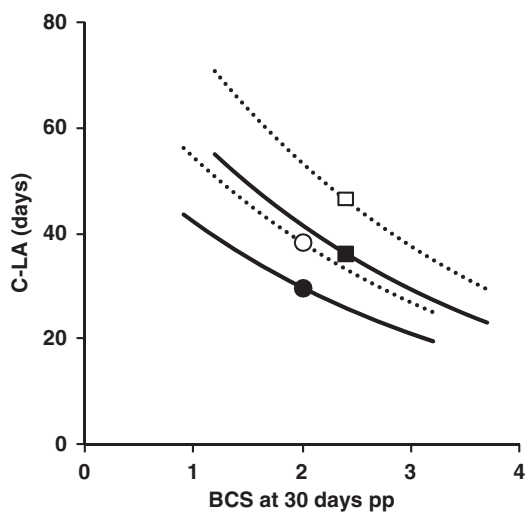


**Figure 2** Body condition (BC) change from calving to 120 days post partum (body condition score (BCS) on 0 to 5 scale) according to the peak milk yield (MY) for Holstein cows of the high (●,  $n = 50$ ) and low (○,  $n = 48$ ) feeding level groups. The solid lines represent the predicted regressions within feeding level when accounting for lactation number, year, BCS at calving, feeding level and peak MY, and the interaction between these two variables ( $R^2 = 51\%$ , r.s.e. = 0.46). The dashed line represents the predicted regression when not accounting for feeding level and its interaction with peak MY ( $R^2 = 29\%$ , r.s.e. = 0.55). Regression slopes and corresponding  $P$ -values are given on the plot.

design partially disrupted the relationship between these two factors and, therefore, provided an opportunity to disentangle the respective effects of MY and BC at each reproductive step in Holstein dairy cows. Owing to the well-documented relationships between reproduction and both MY and BCS (Roche *et al.*, 2009; Friggens *et al.*, 2010), we have assumed in our study that reproductive performance is mainly explained by the expressed MY and BCS level or change, although we cannot exclude any direct feeding treatment or genetic effect.

### *Pp cyclicity was linked to BCS and BC change variables*

*C-LA was mainly related to BCS in early lactation.* C-LA has been extensively studied in dairy cows, as a far too late first ovulation delays breeding time. The relationship between C-LA and parity, genital health problem and BCS pp is



**Figure 3** Predicted commencement of luteal activity (C-LA) according to the body condition score (BCS) at 30 days *post partum* for Holstein primiparous (■ □) and multiparous (● ○) cows with (---) or without (—) genital health problem at or following calving ( $n=98$ ). The corresponding equations are  $\log_e(\text{C-LA}) = 4.39 \pm 0.17$  if primiparous/multiparous  $\pm 0.13$  if genital health problem/no problem  $- 0.352 \times \text{BCS}_{30}$  ( $P = 0.002$ ,  $P = 0.027$ ,  $P < 0.001$ , r.s.e. = 0.49,  $R^2 = 25\%$ ; variable definitions are in Table 2).

consistently observed in the literature (Opsomer *et al.*, 2000; Roche *et al.*, 2009), whereas Figure 3 illustrates that a higher  $\text{BCS}_{30}$  is associated with a shorter interval to C-LA (Model CYC-1, slope  $-0.35$ , i.e.  $-0.22 \log_e$  days per 0.63 unit BCS increase (1 s.d.),  $P < 0.001$ ). Our analysis revealed that the protein content at nadir could also be included in the model as a higher protein content at nadir was associated with a shorter C-LA (slope  $-0.054$ , i.e.  $-0.10 \log_e$  days per 1.9 g/kg increase (1 s.d., mean 26.5 g/kg),  $P = 0.043$ ), whereas MY variables did not affect the model at all ( $P > 0.5$ ). All pp BCS variables had a greater effect than BC change variables. Among those variables,  $\Delta\text{BC}_{30}$  was the most predictive one when  $\text{BCS}_{30}$  was not included in the model. A shorter C-LA was associated with reduced BC loss (slope  $-0.44$ , i.e.  $-0.18 \log_e$  days per 0.40 unit BC change increase (1 s.d., mean  $-0.81$  unit),  $P < 0.001$ ,  $R^2 = 21\%$ ). In this model, the parity effect was no longer significant ( $P = 0.17$ ), as primiparous cows had both a longer C-LA and greater BC loss than multiparous cows. Among the metabolic variables, only the plasma glucose concentration at 20 days in milk had a significant effect when  $\text{BCS}_{30}$  was not included in the model (slope  $-0.019$ , i.e.  $-0.12 \log_e$  days per 6.2 mg/dl glucose concentration increase (1 s.d., mean 58.2 mg/dl),  $P = 0.044$ ,  $R^2 = 17\%$ ). Therefore, although pp BCS variables were more predictive than BC change variables, the influence of a negative energy balance (Butler *et al.*, 1981) was observed once more, as a large BC change within 30 days pp, low milk protein content at nadir and low plasma glucose concentration at 20 days pp also delay the resumption of luteal activity. The combined effects of both the level and the change of body lipid reserves may be expected, as reported in beef cows by Wright *et al.* (1992). A large BC loss may not prevent the cow from ovulating if the current BCS is still

high, namely if the *ante-partum* BCS was high enough, and is consistent with the concept of an 'endocrine memory' (Chagas *et al.*, 2007). However, in our dataset, including an interaction between BC change from calving to 30 days pp and categorized BCS at calving ( $> v. <$  mean within parity) did not improve the prediction of C-LA (results are not reported). Moreover, BCS at 30 days pp does not explain the delayed resumption of luteal activity for primiparous cows whereas BC change does. In comparison with multiparous cows, BCS may not represent the same amount of body lipid reserves for primiparous cows where prolonged uterine involution and the requirements to grow could also affect luteal activity independent of BCS status. In conclusion, the respective influences of the level and dynamic of body lipid reserves on C-LA remain to be quantified.

*PLP profiles were mainly related to C-LA and BC change.* PLP, which delay breeding by preventing the occurrence of ovulation and also by complicating the understanding of cows' cycles for the farmer, has become a major abnormality in Holstein cows (Royal *et al.*, 2000). Once accounted for  $\log_e(\text{C-LA})$  in logistic regression Model CYC-2, the best predictive variables for these abnormal profiles were  $\Delta\text{BC}_{60}$ ,  $\Delta\text{BC}_{90}$  and  $\Delta\text{BC}_{120}$  ( $P = 0.002$ , 0.007 and 0.002, respectively). The sooner luteal activity commenced and the deeper the  $\Delta\text{BC}_{60}$ , the greater the probability of having at least one PLP pp (Table 4). Whatever the model, neither MY variables, nor parity or genital health problem at/following calving had a significant effect ( $P > 0.2$ ). In other words, cows resuming luteal activity early and mobilizing reserves for a long time had PLP (lasting more than 25 days). According to the previously discussed predictors of C-LA, both high BCS at calving and intermediate BC change from calving to 30 days pp are probably required to maintain BCS at 30 days pp and to ovulate early. This result is supported by the different BCS curves observed for delayed-, normal- and PLP-classified cows (Figure 4). Delayed cows were thinner than normal cows after calving. PLP cows tended to have higher BCS before calving than delayed cows, but reached a similar low BC by 60 days in milk, owing to large and prolonged BC loss. Conversely, MY curves did not differ between the  $P_4$  profile categories. A higher BCS at calving for PLP cows is often observed (Pushpakumara *et al.*, 2003; Taylor *et al.*, 2003; Shrestha *et al.*, 2005) but is more rarely significant (Disenhaus *et al.*, 2002). This suggests a narrow optimum for BCS at calving: too thin and too fat are known to delay ovulation, whereas slightly over-conditioned animals are also susceptible to an increased risk of PLP.

Metritis is often considered to be a major risk factor of PLP (Opsomer *et al.*, 2000); however, the inclusion of genital health problems at calving did not improve models, even when distinguishing metritis from other events. Interestingly, all of the 15 cows that encountered severe metritis were well classified using a 0.25 cut-off probability ( $>0.25$  is PLP predicted) in the final model predicting the PLP profile with regard to  $\log_e(\text{C-LA})$  and  $\Delta\text{BC}_{60}$ : nine normal and four PLP profiles. Metritis and PLP may have similar risk factors,

**Table 4** ORs and 95% CIs for the binomial logistic regression model used to predict the occurrence of a PLP profile in 98 progesterone profiles of Holstein cows (decrease in the residual deviance,  $P = 5.8E-5$ ; Model CYC-2)

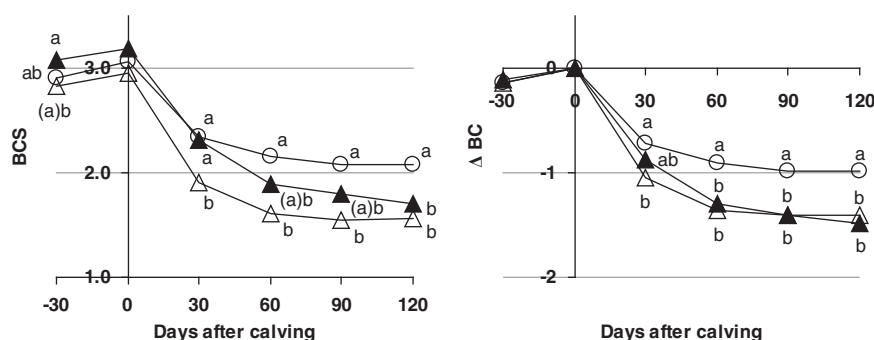
Risk factor	$P (>\chi^2)$	Mean $\pm$ s.d.	OR <sup>1</sup> (PLP v. no PLP)	95% CI <sup>1</sup>
$\text{Log}_e(\text{C-LA})$	<0.001	$3.53 \pm 0.55$	0.271**	0.118 to 0.620
BC change from calving to 60 days pp	0.002	$-1.08 \pm 0.54$	0.406**	0.213 to 0.776

OR = odds ratio; CI = confidence interval; PLP = prolonged luteal phase; C-LA = commencement of luteal activity; pp = *post-partum*.

The intercept of the model is 5.01.

\*\*\*, \*\*, \*, <sup>1</sup>OR is significantly or tends to be different from 1 ( $P < 0.001, 0.01, 0.05, 0.10$ ).

<sup>1</sup>For continuous variables, ORs are expressed as an increase in 1 s.d. instead of 1 unit (i.e.  $\exp(\text{coefficient of the linear predictor})^{\text{s.d.}}$ ).



**Figure 4** Curves of the fitted body condition score (BCS, 0 to 5 scale) and body condition change from calving ( $\Delta\text{BC}$ ) of Holstein cows according to the *post-partum* (pp) progesterone profile category ( $\circ$ , normal;  $\blacktriangle$ , prolonged luteal phase;  $\triangle$ , delayed;  $n = 53, 23$  and  $18$ , respectively). Models also accounted for parity. At the same pp interval, values with a common letter are not significantly different ( $P > 0.05$ ) and values with a common letter in brackets tend to be different ( $P < 0.10$ ).

but the elongation of the luteal phase can probably occur without any uterine disorder. In a  $2 \times 2$  nutritional experimental design, Villa-Godoy *et al.* (1990) induced overfattening or normal BC in dairy heifers and then restricted their feeding or not. The first and second luteal phases were lengthened by 3 and 5 days only for fat-restricted heifers. Thus, a combination of large body lipid reserves at calving and a strong energy deficit in early lactation may lengthen luteal phases.

More precisely, the ongoing cycle type was related to the BC dynamic at the time of ovulation. The final model used to predict the occurrence of a PLP cycle, a short cycle ( $\leq 15$  days) or a normal cycle following ovulation (Model CYC-3) included ovulation rank (first or later), BC slope at ovulation and BC change from calving to ovulation. Irrespective of the ovulation rank, severely declining BC around ovulation increased the probability of a PLP (Figure 5, Table 5). Conversely, a slightly declining or an increasing BC around ovulation increased the probability of a short cycle, but at first ovulation only, as short cycles are scarce anyway at subsequent ovulations (only 2% of ovulations of ranks 2-3-4 were followed by a short cycle). The probability of a short cycle was all the more high as BC change from calving to first ovulation was negative. Of the PLP events, half occur at first ovulation, but can occur up to the fourth ovulation. According to our results, ovulations that occur while mobilization is still ongoing are more likely to be followed by a PLP. Although requiring further investigation, our results

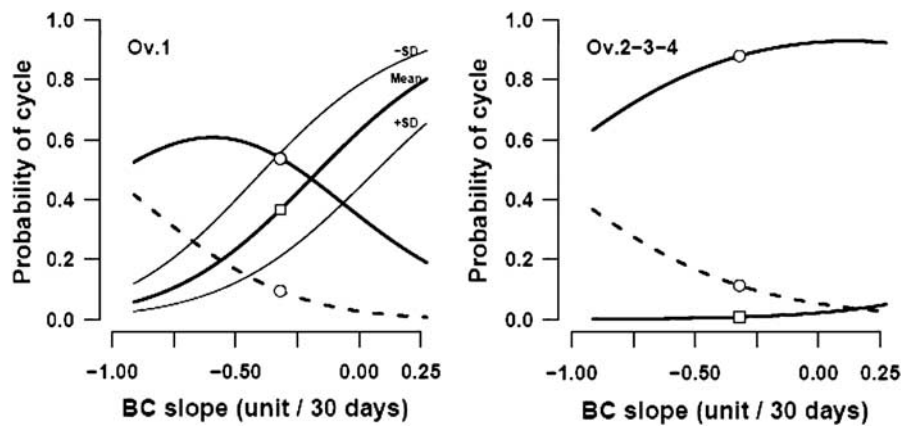
indicate that this BC change effect remains even when accounting for days after calving.

The fact that short cycles conversely occur while the cow is gaining BC close to ovulation is of interest. To our knowledge, there is no literature investigating the relationship between BC change and the realization of short and long luteal phases at first ovulation. Butler *et al.* (1981) considered that transitory  $P_4$  elevation may occur when the cow returns to a positive energy balance, but concluded that it was not related to luteal activity. The probability of short cycle increases as BC change from calving to ovulation widens, suggesting a possible carry-over effect of the negative energy balance period on follicular activity, but only at first ovulation. The risk of short cycles also increases as days in milk increase; however, part of this effect may relate to genetics, as C-LA and the length of the first luteal phase are negatively correlated genetically (Royal *et al.*, 2002).

Overall, these BC effects on ovarian activity are consistent with the much better ovarian activity in breeds with higher BCS throughout lactation and smoother BC change, such as the Normande breed (Cutullic *et al.*, 2011).

#### Oestrus detection relied on current MY

Oestrus detection is a key element of the reproductive performance, especially in compact calving systems. Theoretically, during a 12-week breeding period, cows only have one to four ovulations according to their calving date. In the final Model OES-1 used to predict ovulation detection, the odds of being detected in oestrus were multiplied by 3.0 when at



**Figure 5** Probabilities of the occurrence of either a normal cycle (○-○-), a short cycle (□-□-) or a prolonged luteal phase cycle (-○-○-), predicted at first ovulation (Ov.1) and second–third–fourth ovulations (Ov.2-3-4) not followed by an insemination in 189 cycles of Holstein cows, according to the body condition (BC) slope from –30 to +30 days relative to the day of ovulation (left censored at calving), predicted for a BC change from calving to ovulation of –0.83 (Mean), –1.35 (Mean–1 s.d.) or –0.31 (Mean+1 s.d.) for the short cycles category. Model characteristics are given in Table 5.

**Table 5** ORs and 95% CIs for the multinomial logistic regression model used to predict the occurrence of a normal cycle, a short cycle or a PLP cycle following ovulation in Holstein cows. Prediction relied on 189 ovulations of ranks 1 to 4, occurring before 135 days in milk, not preceding a delayed ovulation of rank above 2, not belonging to a disorder classified progesterone profile and not followed by an insemination (decrease in the residual deviance,  $P = 7.6E-12$ ; Model CYC-3)

Risk factor	$P(>\chi^2)$	Category % Mean $\pm$ s.d.	OR <sup>1</sup> PLP v. normal cycle	95% CI <sup>1</sup>	OR <sup>1</sup> short v. normal cycle	95% CI <sup>1</sup>
Ovulation rank	<0.001					
2-3-4		52%	1		1	
First		48%	1.37	0.42 to 4.54	78.3***	13.1 to 469
BC slope <sup>2</sup>	<0.001	–0.32 $\pm$ 0.30	0.47*	0.26 to 0.84	2.52**	1.38 to 4.62
$\Delta$ BC calving to ovulation	0.012	–0.83 $\pm$ 0.52	1.11	0.63 to 1.97	0.47**	0.27 to 0.82

PLP = prolonged luteal phase; OR = odds ratio; CI = confidence interval; BCS = body condition score; BC = body condition. Intercepts of the model are –2.54 for PLP and –2.78 for the short cycle; the normal cycle is the reference category.

\*\*\*, \*\*, \*, †OR is significantly or tends to be different from 1 ( $P < 0.001, 0.01, 0.05, 0.10$ ).

<sup>1</sup>For continuous variables, ORs are expressed as an increase in 1 s.d. instead of 1 unit (i.e.  $\exp(\text{coefficient of the linear predictor})^{\text{s.d.}}$ ).

<sup>2</sup>BC slope from –30 to +30 days relative to the day of ovulation (left censored at calving), expressed in BCS unit per 30 days.

least one other cow was in oestrus, were divided by 1.7 for an increase in MY by 7.8 kg/day (1 s.d.) and were multiplied by 1.6 for an increase in BCS at ovulation by 0.66 unit (1 s.d.; Table 6). Whereas all MY variables had a significant effect ( $P < 0.001$ ), MY dynamic and general variables had no effect. It was similar for the probability of detecting ovulation by standing oestrus (Model OES-2), except that BCS at ovulation had no significant effect ( $P > 0.4$ ; Table 6). Thus, as observed in our previous work (Cutullic *et al.*, 2009b), an increase in MY at ovulation depresses oestrous behaviour independent of BC change. This deleterious effect of MY is consistent with descriptive studies that possibly confound MY and BC loss effects (Lopez *et al.*, 2004; López-Gatius *et al.*, 2005). However, too low a BCS may also suppress oestrous behaviour, possibly by altering ovarian activity. Indeed, when we investigated the possible links between cyclicity variables and ovulation detection (Model OES-1bis), we observed that delayed ovulations were scarcely detected. Among the variables related to the previous cycle, only having a current delayed ovulation could affect the model

previously reported in Table 6, as none of the six delayed ovulations were detected. When accounting for this variable in the model, the effect of BCS at ovulation narrowed (odds ratio 1.8, 95% CI 1.1 to 3.0). The results suggest that a threshold may exist, under which oestrous behaviour is compromised and above which oestrous behaviour is related to MY. Cutullic *et al.* (2009b) observed a positive effect of losing more BC on oestrous signs; however, the winter total mixed ration low-feeding level was higher and cows did not reach as low a BCS as in this work. In that study, the  $2 \times 2$  factorial design (high v. low feeding in winter  $\times$  high v. low feeding at grazing, allowing cows to switch from high-to-low or low-to-high MY) suggested an instantaneous MY effect. This may help in achieving strategies to improve oestrus detection around the breeding period.

Physiologically, based on standing oestrus duration quantifications (i.e. ovulations without standing oestrus were excluded), Wiltbank *et al.* (2006) suggested that this deleterious effect of MY could be because of increased  $17\beta$ -oestradiol and  $P_4$  hepatic clearance rates, owing to



**Table 6** ORs and 95% CIs for the two binomial logistic regression models used to predict ovulation detection ( $n = 255$ ; Model OES-1) and ovulation detection by standing oestrus ( $n = 241$ ; Model OES-2) in ovulations of ranks  $\geq 2$  of Holstein cows (decrease in residual deviance,  $P = 1.6E-7$  and  $P = 2.4E-6$ )

Risk factors	Detection				Detection by standing oestrus			
	$P (>\chi^2)$	Category % Mean $\pm$ s.d.	OR <sup>1</sup>	95% CI <sup>1</sup>	$P (>\chi^2)$	Category % Mean $\pm$ s.d.	OR <sup>1</sup>	95% CI <sup>1</sup>
Herdmate in oestrus	<0.001				0.027			
Absence		35%	1			37%	1	
Presence		65%	3.01***	1.66 to 5.48		63%	1.91*	1.07 to 3.40
MY ovulation week	<0.001	29.8 $\pm$ 7.8	0.57***	0.43 to 0.77	<0.001	29.6 $\pm$ 7.8	0.59***	0.44 to 0.79
BCS at ovulation	0.001	1.99 $\pm$ 0.66	1.64**	1.20 to 2.25	–	–	–	–
Observation period	0.427				0.005			
Before		9%	1			9%	1	
During		91%	1.52	0.54 to 4.27		91%	4.92*	1.36 to 17.7

OR = odds ratio; CI = confidence interval; MY = milk yield; BCS = body condition score.

Intercepts are 0.46 for the detection model and  $-0.15$  for the detection by standing oestrus model.

\*\*\*, \*\*, \*, †OR is significantly or tends to be different from 1 ( $P < 0.001, 0.01, 0.05, 0.10$ ).

–, Not included in the final model.

<sup>1</sup>For continuous variables, ORs are expressed as an increase in 1 s.d. instead of 1 unit (i.e.  $\exp(\text{coefficient of the linear predictor})^{\text{s.d.}}$ ).

higher dry matter intake. However, if assumed to be the only effect, the dry matter hypothesis is inconsistent with both the similar ovulation detection rate and the similar proportion of oestri detected by standing behaviour, which we observed for Normande and Holstein cows (Cutullic *et al.*, 2011). Indeed, Holstein cows have higher dry matter intakes than Normande cows in grass-based feeding systems (Dillon *et al.*, 2003) and should therefore have poorer oestrus expression. Such a dry matter effect should thus be considered within breed. We may also wonder whether low BC can enhance oestradiol production (Cutullic *et al.*, 2009a), possibly by reducing the inhibitory effect of leptin on steroidogenesis (Spicer, 2001). Direct links between the mammary gland, ovaries and brain may also exist, but they are still poorly understood.

While investigating possible links between cyclicity and oestrus (Model OES-1bis), we also observed that the odds of ovulation detection were divided by 2.6 (95% CI 1.2 to 5.5) for cows with a PLP profile compared with cows with a normal profile. Thus, PLP cows were more at risk of having undetected ovulations, regardless of whether or not these ovulations followed the PLP itself. This result is consistent with the concomitant increase in the PLP frequency (Royal *et al.*, 2000) and the decrease in oestrus duration and intensity (Van Eerdenburg *et al.*, 1996) over the last decades in Holstein cows. PLP occurrence appears to be related to body lipid reserves, whereas oestrus behaviour is associated with MY. This suggests that some effects on oestrus are not accounted for by the MY level and requires further investigations.

#### *Embryo survival was linked to both BCS and MY*

Recalving requires fertilization, early embryo survival, but also late embryo survival and foetal survival. Both early embryo mortality and late embryo mortality are thought to be the major risk factors for pregnancy failure in a temperate climate, in varying proportions according to regions, cow types and

management strategies (Diskin *et al.*, 2006; Humblot *et al.*, 2009). An increase in BCS<sub>min</sub> significantly decreased the risk of non-fertilization or early embryo mortality rather than embryo survival at first and second inseminations (Table 7, Model FER-1). However, an increase in BCS<sub>min</sub> only tended to increase the recalving rate (Table 7, Model FER-2). In fact, in our study, late embryo mortality did not appear to be BCS related, but rather MY related. Indeed, an increase in the slope of MY around ovulation (i.e. a flatter MY slope around ovulation, as MY slopes are mostly negative) decreased the risk of late embryo mortality rather than embryo survival. Significant effects were also found for most of MY variables at the beginning of lactation ( $P < 0.05$ ), suggesting a link to the shape of the lactation curve. Indeed, the Wood curve persistency parameter  $c$  had a significant effect on insemination results ( $P = 0.003$ ). In order to illustrate this effect, four low-persistency curves and four high-persistency curves (within the range of our dataset) were drawn and the corresponding predicted probabilities of insemination results were calculated (Figure 6). Regardless of the MY level, cows with a low persistency after peak were more likely to encounter late embryo mortality, whereas cows with flat curves were more likely to encounter non-fertilization or early embryo mortality. In the final Model FER-1 (non-fertilization or early embryo mortality *v.* late embryo mortality *v.* embryo survival), glucose also tended to have an effect while replacing BCS<sub>min</sub> ( $P = 0.065$ ,  $n = 122$ , six missing values). An increase in glucose concentration at insemination by 5.7 mg/dl (1 s.d., mean 62.8 mg/dl) divided by 2.0 the odds of non-fertilization or early embryo mortality rather than late embryo mortality (95% CI 1.1 to 3.7).

Our results suggest that low BCS increases the risk of early embryo mortality, even though MY is low, whereas the effects of peak lactation increase the risks of late embryo mortality (Figure 6) even when a correct BCS is maintained. MY and BCS may thus affect oocyte and embryo development at different stages.

**Table 7** ORs and 95% CIs for the two logistic regression models used to predict either the occurrence of nf-eem v. lem v. embryo survival (Model FER-1, multinomial) or the occurrence of recalving (Model FER-2, binomial) following 128 artificial inseminations of ranks 1 and 2 of Holstein cows (decrease in the residual deviance,  $P = 0.005$  and  $0.085$ )

Risk factors	Category % Mean $\pm$ s.d.	Model FER-1				Model FER-2			
		$P (> \chi^2)$	OR <sup>1</sup> nf-eem v. survival	95% CI <sup>1</sup>	OR <sup>1</sup> lem v. survival	95% CI <sup>1</sup>	$P (> \chi^2)$	OR <sup>1</sup> recalve v. not	95% CI <sup>1</sup>
Parity		0.845	1		1	0.704	1		
Primiparous	41%		1.24	0.52 to 2.98	1.28		0.86	0.39 to 1.90	
Multiparous	59%	0.100	1		1	0.091	1		
Calving-AI	17%		0.38	0.12 to 1.27	2.13		2.87	0.75 to 10.9	
<50 days	83%		0.94	0.62 to 1.42	0.43**		–	–	
$\geq 50$ days		0.006	0.63*	0.40 to 0.99	0.95	–	–	–	
MY slope <sup>1,2</sup>	$-0.071 \pm 0.141$	0.098				0.089		0.94 to 2.08	
BCS <sub>min</sub> <sup>1</sup>	$1.70 \pm 0.55$								

OR = odds ratio; CI = confidence interval; nf-eem = non-fertilization-early embryo mortality; lem = late embryo mortality; MY = milk yield. \*\*\*, \*\*, \* OR is significantly or tends to be different from 1 ( $P < 0.001$ ,  $0.01$ ,  $0.05$ ,  $0.10$ ).

–, Not included in the final model.

Intercepts of Model FER-1 are 2.00 for the nf-eem category and  $-2.06$  for the lem category; the intercept of Model FER-2 is  $-2.45$ .

<sup>1</sup>For continuous variables, ORs are expressed as an increase in 1 s.d. instead of 1 unit (i.e.  $\exp(\text{coefficient of the linear predictor}^{\text{s.d.}})$ ).

<sup>2</sup>Milk yield slope over weeks  $-2$  to  $+2$  relative to the week of ovulation, expressed in kg/day.

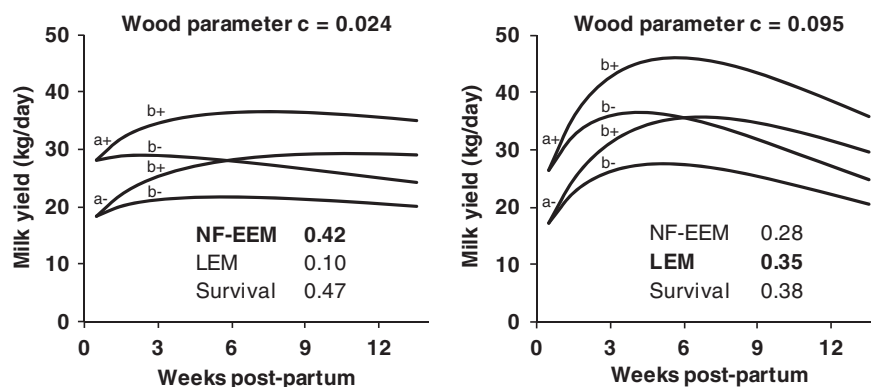
A negative energy balance pp is widely considered to affect oocyte and embryo developmental competence in high-yielding dairy cows (Leroy *et al.*, 2008) and low BCS or large BC change are often reported as major risk factors of infertility (Roche *et al.*, 2007). Little is known about the potential direct deleterious effects of current MY or lactation curve shape on late embryo survival, which may be related to over-exposure to metabolic or galactopoietic hormones. In a field study, Grimard *et al.* (2006) previously reported a deleterious effect of high peak MY on embryo-foetal survival. This shape effect may explain the reduced late embryo mortality rates in breeds with flatter lactation curves, such as the Normande breed (Cutullic *et al.*, 2011). However, further investigation is needed to confirm this result. Similarly, the conflicting impact of plasma glucose concentrations on early and late embryo development remains to be confirmed.

While investigating the possible links between cyclicity, oestrus and fertility (Model FER-2bis), it appeared that fertility at first and second inseminations was not considerably influenced by previous steps. The Model FER-2 already included parity, calving-AI interval  $< 50$  days,  $\text{MYSlope}_{\text{oestrus week } -2 \text{ to } +2}$  and  $\text{BCS}_{\text{min}}$  (Table 7).  $\text{Log}_e(\text{C-LA})$  tended to have an effect when  $\text{BCS}_{\text{min}}$  was removed from the model ( $P = 0.082$ ). An increase in  $\text{Log}_e(\text{C-LA})$  by 0.47 (1 s.d., mean 3.49) multiplied the odds of non-fertilization or early embryo mortality rather than embryo survival by 1.6 (95% CI 1.0 to 2.5). The occurrence of a previous short cycle also tended to improve the model ( $P = 0.098$ ), as the 13 inseminations following short cycles resulted in only two embryo survivals. Other previous cycle characteristics including  $P_4$  profile category and standing behaviour detection did not affect the model ( $P > 0.4$ ) and therefore good oestrus expression was not associated with improved fertility. This result seems logical with regard to the previously observed effects of milk production on oestrus and of BC on fertility and suggests that oestrus can be compromised without compromising fertility, whereas the converse is also true. In our previous study (Cutullic *et al.*, 2009b), the clearer the behavioural signs were, the higher was the recalving rate. However, low-fed cows were not challenged to the same intensity as in this study (Delaby *et al.*, 2009), and the oestrus-fertility relationship probably did not reach uncoupling as we have observed here.

Overall, our results point to a possible uncoupling between cyclicity, oestrus and fertility and allow for compensations leading to similar final reproductive performance in many studies that compared feeding levels (Horan *et al.*, 2004; Pedernera *et al.*, 2008; Pollott and Coffey, 2008; Delaby *et al.*, 2009).

## Conclusion

This study provides new information on the respective effects of MY and of body lipid reserve level and change on the overall reproductive process. The results indicate that these effects may be different at different stages, with some reproductive steps related to body lipid, whereas others may relate to MY.



**Figure 6** Examples of eight Wood lactation curves ( $MY = a \times week_b \times \exp^{-c \times week}$ ), with the persistency parameter  $c$  equal to either 0.024 (i.e. population mean  $-1$  s.d.) or 0.095 (i.e. mean  $+1$  s.d.) and the corresponding probabilities of non-fertilization – early embryo mortality, late embryo mortality and embryo survival at first and second inseminations in Holstein cows ( $n = 128$ ). These insemination result probabilities are predicted for a calving–artificial insemination (AI) interval  $\geq 50$  days and a Wood curve parameter  $c$  equal to either 0.024 or 0.095, in a multinomial logistic model accounting for calving–AI interval ( $P = 0.015$ ) and Wood curve parameter  $c$  ( $P = 0.003$ ; model decrease in residual deviance,  $P = 2.7E-3$ ). For Wood lactation curves, parameters  $a$ ,  $b$  and  $c$  control the milk yield scale, rise and persistency, respectively. Within each graph, that is, for each fixed persistency parameter  $c$ , the four lactation curves are generated by fixing the scale parameter  $a$  to its mean either  $+1$  s.d. ( $a+$ ) or  $-1$  s.d. ( $a-$ ) and by fixing the rise parameter  $b$  to its predicted value either  $+1$  r.s.e. ( $b+$ ) or  $-1$  r.s.e. ( $b-$ ) in a multiple regression model including parameters  $a$  and  $c$ .

Cyclicality and early embryo survival were associated with BCS level and change with deleterious consequences arising from excessively low BCS or high BC change. Conversely, oestrous behaviour and late embryo mortality were mainly associated with MY, level and change, respectively. These results also point to a possible uncoupling between cyclicality, oestrus and fertility and suggest that there is potential for compensation along the reproductive process. In compact calving systems requiring high and compact pregnancy rates, high MY strategies appear to be unsuitable even if BCS is maintained, owing to depressed oestrous behaviour and probably increased late embryo mortality that delays rebreeding. Similarly, strategies that compromise cyclicality and fertility by excessively low BCS are unsuitable. The results of this study also indicate that the thresholds wherein reproductive processes become compromised remain to be determined in order to optimize reproductive performance.

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