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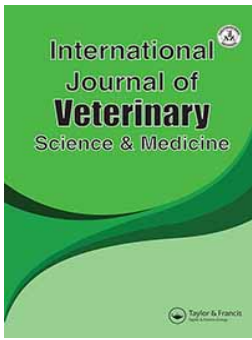
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





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Kick-starting ovarian cyclicity by using dietary glucogenic precursors in post-partum dairy cows: a review

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ABSTRACT

The objective of this review is to describe how dietary glucogenic precursors could stimulate ovarian activity in post-partum dairy cows and improve reproductive success. Although the nutrient requirements for the early resumption of ovarian cycles, and for follicle and embryo development are quantitatively small, reproductive success is deteriorated by post-partum negative energy balance. Since very little glucose is absorbed directly from the digestive tract of ruminants one of the targets for nutritional manipulation could be the glucogenic potential of the diet. This could be achieved by giving rumen-resistant starch or mono-propylene glycol. Both these adaptations increase glucose, insulin and insulin-like growth factor-1 plasma concentrations and stimulate ovarian follicle growth.

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1. Introduction

The objective of this review was to describe how dietary glucogenic precursors could stimulate ovarian activity in the post-partum (PP) dairy cow and improve reproductive success. As a result of increases in milk production obtained through advances in genetic selection and improved husbandry, reproductive efficiency in dairy cows has declined between 1975–1982 and 1995–1998 from 55.6% to 39.7% [1]. Although this negative trend has recently bottomed out and reproduction has begun to improve due to the inclusion of fertility traits in selection programmes [2], modern dairy cows still require an additional 30d to conceive when comparing results between 1999 and 2010 [3].

Feeding dairy cattle should always be optimized to cover requirements for milk production and maintain good health but it also may be possible through the choice of certain feedstuffs to target a particular physiological function, such as reproduction. The idea of targeting certain aspects of metabolism to stimulate reproduction in dairy cows was first proposed nearly twenty years ago with the use of glucogenic vs. lipogenic diets [4]. Further research has since been conducted.

The first section of this paper will outline the general metabolic context of the dairy cow PP. The second part will describe the rationale behind modifying the diet PP. The final section will describe how glucogenic supply can be increased to improve reproductive success.

2. Metabolism in the post-partum dairy cow

In the dairy cow, the negative energy balance (NEB) occurs PP [5] because the increase in feed intake after parturition, is not able to keep up with the rapid rise in energy requirements for milk production [6] even though the cows are fed ad libitum. The requirements for energy and protein of an average European dairy cow at peak milk production are multiplied by 3 to 5 compared to late gestation [7] and the peak in nutrient requirements occurs earlier (at 1 to 2 months PP) compared to the peak in feed intake (at 3 to 4 months PP) therefore inducing NEB [7]. The problem is physiological in relation to a lag in feed intake compared to nutrient requirements.

NEB can also exist in beef cattle PP but the situation is different compared to dairy cows because beef cows are often managed in low input systems. Nutrient requirements are not as high PP but farmers often use low quality forages or limit cow access to good quality forages therefore inducing NEB [7]. The problem is due to the farmer trying to reduce production costs by limiting nutrient intake compared to nutrient requirements.

As a result of NEB, insulin decreases and growth hormone (GH) increases during this period to promote lipolysis [8]. Despite high circulating GH, there is a decrease in insulin-like growth factor-1 (IGF1) because insulin is low and is no longer able to stimulate the expression of the GH receptor 1A. Without this receptor GH cannot stimulate IGF1 production. The somatotrophic axis is said to become “uncoupled”

[9]. Concomitantly, PP non-esterified fatty acids (NEFA) increase and this can lead to ketosis [10] and hepatic steatosis [11] if NEFA are not completely oxidized or exported. Hepatic steatosis caused by triglyceride accumulation reduces the ability of hepatocytes to synthesize glucose from propionate [12]. In conclusion, PP NEB results in low glucose, insulin and IGF1 and high GH, NEFA, β -hydroxybutyrate (BHB) and liver triglycerides [13].

Homeorhetic modifications occur PP to spare glucose and involve a decrease in insulin concentrations and tissue sensitivity to insulin [14]. Part of the mechanism is raised NEFA which reduce insulin sensitivity by provoking ceramide accumulation in plasma and liver [15,16]. When NEFA are mobilized palmitic acid increases [17] and it is a precursor of ceramides. Plasma ceramides were positively correlated with plasma NEFA and inversely correlated with insulin sensitivity in dairy cows in the peripartum period [18].

Depending on the tissue, glucose uptake requires insulin (insulin-dependent tissues, adipose tissue, muscle, ovary, hypothalamus) or does not require insulin (non-insulin dependent tissues as brain, heart and udder [19]). Glucose supply is important for ovarian metabolism because insulin-sensitive glucose transporters, GLUT1 and GLUT4, are present in sheep granulosa and theca cells [20] and GLUT4 in cumulus oophorus cells [21] and glucose is taken up by the ovary during the oestrous cycle [22]. Glucose uptake by the ovary may become limited for some cows because insulin is low and insulin-sensitive tissues are less responsive to insulin's action [23].

In conclusion, the NEB observed after calving activates homeorhetic adjustments to metabolism to divert nutrients towards milk production and this in turn reduces the availability of glucose for reproductive tissues.

3. Glucose precursors to improve reproductive efficiency

Changing the composition of the diet or adding mono-propylene glycol (MPG) can increase glucose precursors. The papers published on the effect of a glucogenic supplement (starch or MPG) on reproductive function are summarized in Tables 1–3.

3.1. Peri-partum period (ketosis and immunity)

A recent meta-analysis showed that the interval between calving-to-first-service was 8 d longer and calving-to-conception was 16 to 22d longer in cows with subclinical ketosis [24]. In periparturient dairy cows, MPG increased insulin and glucose while decreasing NEFA and BHB [25] and reduced the triacylglycerol content of the liver [26]. Therefore, MPG

Table 1. Summary results of the effect of a dietary glucogenic supplement in the form of starch or mono-propylene glycol (MPG) on reproductive function in grazing post-partum dairy cows.

Amount of glucogenic suppl. (ml/cow/d or % diet)	Source of starch	Duration of suppl. (days)	No. Cows /treatment	Metabolic effects			Energy balance	Effect of suppl. on reproductive parameters	References
				Glucose	Insulin	IGF1 NEFA			
17.8%	barley-corn	during 36d PP	17	↑	↑	↑	-	ICO ↓ ICAI → early CR trend ↑	[78, 79]
38.1%	-	during 6wk prior to planned mating	580-622	-	-	-	-	ICO ↓ CR at 12 and 16wk ↑ (200mL>2 x 200mL=0)	[80]
-	200mL 2 x 200mL	during 16wk PP	13-17	→	basal → ↑	→	-	ICO ↓ if low BCS at calving final ↑ if BCS low at calving LH pulse frequency ↑	[37]
-	2 x 250mL	during 21wk PP	13-18	↑	→	→	Good BCS	ICO →	[81]
NSC 22.5% NSC 34.7%	control = palm kernel meal-soy meal treatment = barley-corn	during 5-6wk PP stopping 4wk before breeding season	471-478	-	-	-	-	ICO → final CR tended ↓ in one of the herds used in expt → for other herds	[40]

NSC: non-structural carbohydrate, PP: post-partum, ICO: interval calving oestrous, ICAI: interval calving artificial insemination, LH: luteinizing hormone, P4: progesterone concentrations, CR: conception rate, BCS: body condition score, ↑: increase, →: no effect, ↓: decrease, -: not measured.

Table 2. Summary results of the effect of a dietary glucogenic supplement in the form of mono-propylene glycol (MPG) on reproductive function in non-grazing post-partum dairy cows.

Basal diet	Amount of glucogenic suppl. (g or mL/cow/d)		Duration of suppl. (days)	No. Cows/treatment	Metabolic effects				Energy balance	Effect of suppl. on reproductive parameters	References
	Basal diet starch and source	MPG			Glucose	Insulin	IGF1	NEFA			
TMR maize silage	?	0 mL 250 mL	Between 3–15d PP	9	-	-	-	-	-	ICO → ICAI ↓ CR at 1st AI →	[82]
TMR maize silage	Basal diet maize starch 21%	0 g 225g MPG + 225 g Ca propionate	During 6 wk PP	10	-	-	-	-	-	ICO ↓ CR at 1st AI ↑ ICF ↓ number AI/ fertilization ↓	[83]
TMR	?	0 mL 850 mL	-3 to 8d of induced oestrus at 60 DIM	13	-	-	-	-	-	P4 →	[84]
TMR prepartum hay/maize silage PP 33%/66% grass silage or lucerne hay/maize silage	Basal diet maize starch 20%	0 mL 300 mL	During 10d prepartum and on days 3, 6, 9 and 12 PP	19–20	-	-	↑	↑	-	At 90d PP number of acyclic cows ↓	[39]
50%/50% maize silage/lucerne hay + concentrates for milk production	Basal diet starch 15%	0 mL 500 mL	Between 7 and 42d PP	16–17	↑	↑	-	↓	No difference between groups	ICO ↓ length of 1st luteal phase ↑ (13.1 vs. 7.3d) P4 secretion → CR 1st AI → CR at 150d PP → Number of AI/conception →	[25]
50%/50% maize silage/ grass silage + concentrates for milk production	Basal diet maize starch 15%	0 mL 500 mL	Between 7 to 35–40d PP	17–18	↑	↑	-	↓	-	No effect on follicle dynamics no effect on LH secretion characteristics no effect on oocytes collected and their quality	[85]
TMR maize silage/grass hay/straw	?	0 mL 6 × 200 mL	Holstein cows at maintenance	17 from start of synchronization and superovulation until after 2 nd AI (embryos collection 6d after 2nd AI)	↑	↑	-	-	-	P4 at embryo recovery ↓ ovulation rate ↓ fertilization rate ↓ no effect on recovery rate and quality of recovered embryos	[86]
TMR 50%/50% maize silage/legume hay	Basal diet starch 23.5%	0 mL 500 mL	-10d to +25d PP	28	↑	↑	-	↓	Improved	No effect on follicle dynamics no effect on LH secretion characteristics	[36]
-	-	0 mL 267 mL	super ovulated heifers with AI and embryo collection cross-over	20 treatment over 10d during superovulation and AI period	-	↑	-	→	-	Number of transferable embryos → number of degenerated embryos → number of unfertilized oocytes recovered →	[62]
Hay plus concentrate	Basal diet maize starch 3%	0 mL 400 mL	Super ovulated heifers with OPU and embryo production cross-over	16 treatment over 5d during superovulation	↑	↑	↑	-	No difference in growth rate	Number of follicles, blastocysts and blastocyst quality ↑ in high AMH group	[61]

TMR: total mixed ration, PP: post-partum, ICO: interval calving oestrous, ICAI: interval calving artificial insemination, ICF: interval calving fertilization, DIM: days in milk, P4: progesterone concentrations, CR: conception rate, #: intensive blood sampling after MPG treatment, AMH: anti-Müllerian hormone, ↑: increase, →: no effect, ↓: decrease, -: not measured, ?: not indicated in publication.

Table 3. Summary results of the effect of a dietary glucogenic supplement in the form of starch on reproductive function in non-grazing post-partum dairy cows.

Basal diet	Amount of glucogenic suppl. (% diet)		Source of starch	Duration of suppl. (days)	No. Cows /treatment	Metabolic effects			Energy balance	Effect of suppl. on reproductive parameters	References	
	Basal diet starch	Starch				Glucose	Insulin	IGF1				NEFA
50%/50% grass /maize silage concentrates given individually 3.5-12kg	10.4%	26%	maize	between 3wk pre- to 9wk post-calving	42-44 for repro. 25-26 for metab.	→	↑ for MultiP cows → for Prim P cows	-	↓	no difference between groups	ICO ↓ no differences for parameters used to describe reproductive cycles (hormone levels, length ...)	[87]
TMR 77%/23% grass/maize silage	8.7%	13.5% 15.9% 18.3% 23.1%	wheat	between 40 and 70d PP	5	→	↑	→	↓	no consistent difference	P4 ↓ 3 to 5d post-ovulation small follicles ↑ pre- and post-ovulation size medium follicle ↓	[45]
TMR 66%/33% grass/maize silage	L = 9.8%	H = 18.2%	wheat	Between calving and 120d PP diets switched at first rise in P4 = HH, LH, LH and LL	15	→	↑	-	-	no difference between groups	total follicles at 60d PP ↑ Number of CL at 60d PP ↓ CR 1st AI ↑ Overall CR ↑	[60]
TMR 66%/33% grass/wheat silage	starch = 19% rumen by-pass starch: 7.1%	starch = 19% rumen by-pass starch: 8.2% 9.4% 10.5% 11.6%	wheat	between 40 and 70d post-calving	6	-	→	→	-	-	no differences for parameters used to describe reproductive cycles (P4 levels, follicle numbers ...) before and after synchronisation at 50d PP	[88]
TMR 75%/25% grass/maize silage or 25%/75% grass/maize silage	starch: 11.0% rumen by-pass starch: 4.6%	starch: 18.8% 19.1% 27.1% rumen by-pass starch: 8.0% 8.1% 14.4%	maize or wheat	between 40 and 70d post-calving	8	-	↑ by starch ↑ by maize vs. grass	→	-	-	no differences for follicle numbers before and after synchronisation at 50d PP P4 ↑ in grass vs. maize silage 3 to 5d post-ovulation FSH → follicle development → ICO ↓ CR →	[4]
TMR grass /wheat silage	10%	26%	wheat	between calving and 50d PP	10	-	↑	-	-	-	no difference in normal or abnormal resumption of cycling, ICCL, CL length, cycle length	[89]
TMR 60%/40% grass/maize silage concentrates for milk production	10.6%	21.5%	maize	between calving and 100d PP 2 diets x 3 dry periods	68-73	→	→	→	→	improved	no difference in normal or abnormal resumption of cycling, ICCL, CL length, cycle length	[90]
TMR 60%/40% grass/maize silage concentrates for milk production	10.6%	21.5%	maize	between calving and 100d PP 2 diets x 3 dry periods	total 130 cows over 6 treatments Repetition of Chen et al. [89] with same cows	↑	↑	↑	↓	no difference	no difference in normal or abnormal resumption of cycling and ICAI ICCL shorter Pregnancy rate ↓	[91]
TMR 66%/33% maize silage/ lucerne hay	19.20%	35.3%	maize	lactating cows cross-over after 14d	11	-	↑	-	-	improved	Cytochrome P 450 2C and CYP3A activity and mRNA expression ↓ (or tended to ↓) P4 half-life tended to be ↑	[52]

TMR: total mixed ration, PP: post-partum, CL: corpus luteum, ICO: interval calving oestrous, ICAI: interval calving artificial insemination, ICF: interval calving fertilization, P4: progesterone concentrations, CR: conception rate, ICCL: interval calving corpus luteum appearance, ↑: increase, →: no effect, ↓: decrease, -: not measured.

reduces the risk of cows developing subclinical and clinical ketosis, and hepatic steatosis.

In addition, mastitis and endometritis can become a problem [27] because cows cannot fight oxidative stress and their immune system is depressed PP. Clinical mastitis delays ovarian activity [28], reduces conception rates [29] and increases embryonic losses [30]. In cases of infection, peripheral insulin sensitivity decreases, leading to decreased glucose uptake by insulin-dependent tissues such as skeletal muscle [31], adipose tissue [32] and probably the ovary in order to preserve glucose for the immune system. Indeed, it has been estimated that an activated immune system requires substantial quantities of glucose, 2 kg/day, in addition to lactation requirements [33]. Therefore, the immunologically challenged PP cow may benefit from a dietary supplement of glucose (starch or MPG).

3.2. Ovarian activity

3.2.1. Delayed resumption of ovarian cyclicity

Numerous growth factors (insulin and IGF1) and metabolites (glucose) influence gonadotropin-releasing hormone (GnRH) release from hypothalamic neurons [34] and both follicle-stimulating hormone (FSH) and luteinizing hormone (LH) are released from the anterior pituitary in response to GnRH [35]. FSH stimulates follicle recruitment and early follicle growth while pulsatile LH is required for continued growth and the development of the dominant ovulatory follicle. Butler [5] found that NEB is strongly associated with low levels of blood glucose, insulin and IGF1 and at the same time LH pulse frequency is reduced. Glucogenic precursors (starch supplement) did not influence FSH concentrations in non-grazing cows [4] and Butler et al. [36] showed that MPG in non-grazing cows had no effect on LH secretion characteristics. To our knowledge only one publication showed a positive effect of MPG on LH pulse frequency [37] and it was in grazing cows. Therefore, glucogenic precursors do not appear to modify FSH and LH secretion parameters. However, the pulsatile nature of their secretion may make studies difficult to undertake.

Britt [38] suggested that the negative effect of NEB on fertility could be explained by a carry-over effect of some metabolites on follicles during their development from inactive primordial follicles up to ovulation which takes between 60 to 80 days. Exogenous and endogenous lipogenic metabolites are acetate, butyrate and long-chain FA while glucogenic metabolites are propionate and starch. A glucogenic diet given between calving and 50 days PP increased plasma insulin and IGF1 compared with a lipogenic diet and this resulted in a greater proportion of cows ovulating by 50 days PP [4]. Rumen MPG fermentation produces propionate and MPG drenches modify

ovarian activity and hormones and metabolites [25]. Indeed, ovarian cycles started earlier in cows given MPG drenches PP compared with controls (38% acyclic vs. 58% acyclic at 90d PP) and IGF1 and cholesterol were higher while NEFA was lower although insulin was unaffected [39]. Butler et al. [36] were however unable to show an effect of MPG drenches on ovarian activity in the calving to 27d PP period. Other groups using grazing cows have also been unable to confirm the positive effect of MPG on reproduction [40].

Insulin stimulates follicle recruitment [41] as well as follicular growth and differentiation [42]. Moreover, insulin stimulates *in vitro* proliferation and function of granulosa [43] and thecal cells [44]. Starch addition to the diet of lactating dairy cows increased insulin concentrations [45] and increased insulin in follicular fluid of preovulatory follicles in high producing dairy cattle [46].

3.2.2. Steroid production

A short luteal phase is often observed during the first oestrus cycle PP. This short luteal phase was prevented by giving MPG drenches which increased insulin and restored normal progesterone (P4) concentrations [25]. In addition, P4 is necessary for the uterine secretion of nutrients and growth factors that are essential for early embryonic development.

Circulating steroid hormone concentrations are affected by their rate of production and clearance (hepatic blood flow and catabolic enzyme activities). In goats, weekly administration of insulin prior to and during gestation increased circulating P4 [47]. Insulin may increase P4 production by stimulating cholesterol uptake across the ovary since there was a strong correlation between glucose and cholesterol uptake by the ovary in ruminants [48].

Moriel et al. [49] showed in ovariectomized cows given a P4 intra-vaginal implant that when dietary treatment increased insulin, P4 concentrations were also higher. P4 is inactivated in the liver by cytochrome P450 2 C (CYP2 C) or cytochrome P450 3A (CYP3A) [50,51]. Elevated insulin concentrations produced by dietary manipulation (high starch vs. high fibre) decreased P4 clearance and prolonged P4 half-life in lactating dairy cows [52] without any changes in liver blood flow. CYP2 C activity was decreased and CYP2 C mRNA expression tended to be decreased, and CYP3A activity tended to be reduced and CYP3A mRNA expression was unaffected (starch vs. fibre [53]). Finally, Lemley et al. [54] demonstrated that MPG or insulin infusion decreased the abundance in liver biopsies of mRNA for enzymes responsible for hepatic P4 catabolism. In conclusion, insulin appears to increase circulating P4 concentrations by increasing cholesterol uptake by the ovary and by reducing hepatic steroid clearance.

3.3. Oestrus expression

High producing cows have shorter oestrus periods and lower plasma oestradiol (E2) concentrations than those producing less milk [55]. These observations are partly explained by an increase in hepatic clearance since high milk production is associated with high feed intake [56]. *In vivo* Butler et al. [57] using a hyperinsulinaemic-euglycemic clamp in PP dairy cows showed that NEFA decreased and, IGF1 and E2 increased. Further experiments have also confirmed the positive effect of insulin on E2 production in superovulated goats [58]. Therefore, insulin appears to have positive effects on E2 production and may improve the expression of oestrous.

3.4. Pregnancy rate

Although insulin has positive effects on follicle growth, it is important to reduce insulin levels during the insemination period. Indeed, high insulin induced by high starch diets during the insemination period had negative effects on oocyte quality and blastocyst development rate [59] and pregnancy rate tended to be reduced [60]. Gamarra et al. [61] showed that MPG drenches during superovulation in heifers improved the production of grade 1 oocytes, expanded blastocysts and embryos after ovum pick-up (OPU), *in vitro* maturation (IVM), fertilization and culture. The collected oocytes were no longer under the influence of high insulin during fertilization and culture since they had been collected and placed in culture medium. Recently using a similar model, Dupras et al. [62] showed that MPG during superovulation and up to the first 4d after artificial insemination (AI) did not influence the number of transferable embryos collected 6d after AI. This finding supports the conclusions of Fouladi-Nashta et al. [59] and Garnsworthy et al. [60].

In conclusion, increasing glucogenic nutrients in the early PP cow could stimulate follicle growth (via increased glucose, insulin and IGF1), limit lipolysis and ceramide production (via insulin) and support P4 concentrations.

3.5. Inconsistencies in results

Not all experiments have shown a positive effect of glucogenic precursors on reproductive success. Several factors are identified to explain these discrepancies (Tables 1–3): sampling frequency, type of feeding system, the genetic background of the cows and a lot of the studies were under-powered. Infrequent sampling (weekly) often resulted in no visible effect of the glucogenic supplement on circulating hormones and metabolites while frequent sampling did. The feeding system modifies the glucogenic profile of the basal diets. Grazing would provide a more lipogenic profile

(high sugar and fibre levels) compared with conserved forages (maize silage). Lastly, the genetic background of the cows was different in the studies: New Zealand Holstein and Jersey-Holstein crosses compared with North American Holstein cows. The latter have been shown to produce more milk and mobilize more body reserves. Part of the effect of an increase in milk production was attributed to a reduction in insulin sensitivity in North American cows compared with New Zealand cows [63].

4. Practical suggestions to manipulate insulin concentrations

4.1. Limit ketosis and steatosis

It has recently been estimated that the average cost of a case of clinical ketosis and a case of sub-clinical ketosis were respectively, €709 and €150 [64]. Mono-propylene glycol was first reported to be useful in the treatment of ketosis in the 1950's [65]. McArt et al. [66] showed that oral drenching with MPG decreased hyperketonaemia in early lactation dairy cows. While Rukkamsuk et al. [67] showed that drenching with 400 mL MPG once daily from 7 days prior to expected calving until 7 days after calving reduced steatosis. Therefore, cows with a higher body condition score (BCS) than recommended prior to calving (≥ 3.5 on a 5 point scale) could be given (–1 to +2 weeks) MPG daily (300 mL/cow/d [66],) either mixed with the concentrates of the diet or given as a drench. MPG will limit adipose tissue lipolysis and steatosis by stimulating insulin secretion and promoting NEFA catabolism.

4.2. Encourage ovarian cyclic activity

The idea is to “kick-start” normal ovarian activity in the period +2 to +8 weeks prior to insemination to improve conception rate [68].

Firstly, increase dietary starch level. High rumen fermentable dietary starch is one of the risk factors for acidosis. Maize and sorghum are high in “rumen protected” starch compared with wheat (195–215 g/kg DM vs. 65 g/kg DM [69]). Sauvant et al. [70] calculated that there was no risk of acidosis if dietary rumen digestible starch was below 25% of dry matter. Climate change, currently characterized by increased atmospheric CO₂, rising temperatures and above all an alteration in the pattern of precipitation [71], may mean that growing sorghum is easier than maize to provide starch since sorghum is much more resilient to low rainfall than maize [72]. In addition to the choice of grain type, the preparation method [73] as well as maturity of grain at harvest are important [74]. Mature ground or rolled grain is recommended [75]. Rumen resistant starch may not be completely hydrolysed in the small intestine [76] due to starch increasing small intestine viscosity

[75]. Therefore, the total starch level of the diet should be lower than 35–40% to avoid acidosis and allow complete starch hydrolysis [77]. To ensure complete hydrolysis the quantity of starch reaching the small intestine should not exceed ≈ 2.5 kg/d [76].

Secondly MPG could be added to the diet. MPG does not cause acidosis. However, MPG is relatively un-palatable and may reduce feed intake if top-dressed on forage. The time course of MPG action is probably different from starch because its effects on glucose and insulin are relatively short-lived, 2–3 h, and of large amplitude. MPG can either be given as a drench or mixed with a concentrate in an automatic concentrate feeder such as in robotic milking systems. In conclusion, a supplement of ≈ 300 mL MPG/cow/d [66] can be given to stimulate reproductive function. Nielsen and Ingvarsten [65] concluded in their review that at levels of supplementation below 500 g/d unwanted side-effects should not be observed although individual cow responses were variable.

5. Conclusion

Glucogenic treatments have a dual role in the improvement of reproductive success. Firstly, through effects on metabolism and secondly, through direct effects on reproductive function.

Glucogenic treatments affect metabolism by reducing the risk of ketosis and steatosis by decreasing lipomobilisation and stimulating ketone oxidation. Limiting lipomobilisation reduces circulating palmitic acid and ceramide production. The latter can cause insulin-resistance and reduce the availability of glucose for the ovary therefore limiting ovarian function.

Glucogenic precursors appear to affect reproductive function by a local (ovary) rather than central mechanisms since they do not influence FSH and LH secretory characteristics. At the local level glucogenic precursors increase follicle recruitment, growth and differentiation, increase E2 concentrations (through improved granulosa and theca cell proliferation and function) and P4 concentrations (increased secretion by the corpus luteum and reduced clearance by the liver) and generally improve oocyte quality. However, maintaining high insulin around insemination may decrease oocyte quality and embryo survival.

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