Metabolic and endocrine profiles and reproductive parameters in dairy cows under grazing conditions: effect of polymorphisms in somatotropic axis genes

Gretel Ruprechter1*, Mariana Carriquiry1, Juan Manuel Ramos2, Isabel Pereira1 and Meikle Ana1

Abstract

Background: The present study hypothesized that GH-AluI and IGF-I-Snabl polymorphisms do change the metabolic/endocrine profiles in Holstein cows during the transition period, which in turn are associated with productive and reproductive parameters.

Methods: Holstein cows (Farm 1, primiparous cows, n = 110, and Farm 2, multiparous cows, n = 76) under grazing conditions were selected and GH and IGF-I genotypes were determined. Blood samples for metabolic/endocrine determinations were taken during the transition period and early lactation in both farms. Data was analyzed by farm using a repeated measures analyses including GH and IGF-I genotypes, days and interactions as fixed effects, sire and cow as random effects and calving date as covariate.

Results and Discussion: Frequencies of GH and IGF-I alleles were L:0.84, V:0.16 and A:0.60, B:0.40, respectively. The GH genotype was not associated with productive or reproductive variables, but interaction with days affected FCM yield in multiparous (farm 2) cows (LL yielded more than LV cows) in early lactation. The GH genotype affected NEFA and IGF-I concentrations in farm 1 (LV had higher NEFA and lower IGF-I than LL cows) suggesting a better energy status of LL cows.

There was no effect of IGF-I genotype on productive variables, but a trend was found for FCM in farm 2 (AB cows yielded more than AA cows). IGF-I genotype affected calving first service interval in farm 1, and the interaction with days tended to affect FCM yield (AB cows had a shorter interval and yielded more FCM than BB cows). IGF-I genotype affected BHB, NEFA, and insulin concentrations in farm 1: primiparous BB cows had lower NEFA and BHB and higher insulin concentrations. In farm 2, there was no effect of IGF-I genotype, but there was an interaction with days on IGF-I concentration, suggesting a greater uncoupling somatotropic axis in AB and BB than AA cows, being in accordance with greater FCM yield in AB cows.

Conclusion: The GH and IGF-I genotypes had no substantial effect on productive parameters, although IGF-I genotype affected calving-first service interval in primiparous cows. Besides, these genotypes may modify the endocrine/metabolic profiles of the transition dairy cow under grazing conditions.

* Correspondence: gruprechter@adinet.com.uy
1Faculty of Veterinary Medicine and Agronomy Sciences, University of Uruguay, Montevideo, Uruguay
Full list of author information is available at the end of the article

© 2011 Ruprechter et al; licensee BioMed Central Ltd. This is an Open Access article distributed under the terms of the Creative Commons Attribution License (http://creativecommons.org/licenses/by/2.0), which permits unrestricted use, distribution, and reproduction in any medium, provided the original work is properly cited.
Background

Energy balance of dairy cows becomes negative (NEB) during the transition period due to increased nutrient requirements that typically exceed dietary intakes. With the onset of lactation, plasma levels of non-esterified fatty acids (NEFA) and B-hydroxybutyrate (BHB) increase markedly, according to the magnitude of adipose tissue mobilization, to provide additional energy for maintenance and milk production [1-3]. Growth hormone (GH) is known to be responsible for galactopoiesis and persistency of lactation [1,4], and the uncoupled somatotropic axis (GH-insulin-like growth factor I axis, IGF-I) mediates nutrient partitioning for lactogenesis in high producing dairy cows [5]. Concentrations of GH are usually increased during early postpartum and its metabolic effects are antagonistic to insulin by enhancing lipolysis in the adipose tissue and gluconeogenesis in the liver [1,6,7]. Thus, insulin resistance develops to help direct nutrients from insulin-sensitive tissues to the lactating mammary gland [1]. Indeed, genetically-selected dairy cows had increased GH and reduced IGF-I and insulin concentrations during early lactation [8]. Since IGF-I and insulin affect ovarian function, low concentrations of these hormones during the postpartum period are associated with prolonged acyclicity [9-13]. As GH has proven to play a key role on the regulation of metabolism and milk production by modulating the expression of many genes, including IGF-I [14,15], these two genes - GH and IGF-I - could be considered candidate gene markers for productive and reproductive traits.

A polymorphic site of the GH gene that results in an amino acid change at position 127 - leucine, (L) to valine, (V) - detected by AluI, has been linked to milk production traits [16]. However, research results have been controversial as several authors [17-20], reported increased production traits associated with the L allele, while others [21-23] determined a favorable effect of the V allele on production. In contrast, Yao et al. [24] were not able to prove any association between this polymorphism and production traits. Very few studies have been performed regarding the relationship between GH-AluI genotype and reproduction [25-27]. Lechnniak et al. [25] reported that homozygous VV beef bulls tended increased production traits associated with the L allele, whereas no effect of this polymorphism in days to first postpartum ovulation in dairy cows. A polymorphic site in the first promoter region of the bovine IGF-I gene was found by Ge et al. [28]. This polymorphism was identified as a point mutation, T (allele A) to C (allele B) transition, also referred to as SnaBI by the same author. Unlike the abundant reports found in relation to GH-AluI genotype, scarce reports exist regarding the relationship between milk production and the IGF-I-SnaBI genotype. Siadkowska et al. [29] determined that Polish Holstein-Friesian cows carrying the AB genotype yielded more daily fat-corrected milk (FCM) than those of AA and BB genotypes, while Hines et al. [30] found no association between IGF-I-SnaBI genotype and production traits in Holstein cattle. In addition, the BB genotype has been associated with greater body weight at weaning in commercial beef lines of Bos taurus [31] and greater growth rates in Holstein-Friesian bulls [29]. We have not found reports of IGF-I polymorphism and bovine reproduction.

Few studies performed in different bovine breeds and physiological stages focused on the mechanism by which these GH or IGF-I genotypes affect metabolic and endocrine profiles [32,28,33,34]. Only one report on the mentioned GH and IGF-I polymorphisms in dairy cows during the transition period was found. Balogh et al. [33] could not demonstrate any effect of GH-AluI genotype on BHB, insulin, IGF-I, and leptin concentrations in one blood sample collected between 4 and 13 days postpartum in Holstein Friesian dairy cows. The present study hypothesized that GH-AluI and IGF-I-SnaBI genotypes do change the metabolic and endocrine profiles in Holstein cows during the transition period, which in turn may be associated with the productive and reproductive responses.

Materials and methods

Animals and experimental design

Holstein cows under grazing conditions from two commercial dairy herds in Uruguay were used. All procedures were carried out in accordance with regulations of the Animal Experimentation Committee (Veterinary Faculty, University of Uruguay, Uruguay). Blood samples collected by coccygeal venopunction into tubes Vacutainer® (Becton Dickinson, NJ, USA) containing K3EDTA were used to determine GH and IGF-I genotypes. Preliminary data of milk production and composition according to these GH and IGF-I genotype has been published before [35].

Farm 1

Primiparous Holstein cows that calved between March and May were randomly selected (n = 110) from a 700-cow herd. All cows grazed a mixture of ryegrass (Lolium multiflorum) in the morning and alfalfa (Medicago sativa) in the afternoon and were supplemented with 12 kg dry matter (DM) of corn silage, 5 kg DM of high-moisture corn grain, and 2 kg DM sunflower meal. The diet offered had 17% crude protein and 1.7 Mcal/kg DM of net energy of lactation (NRC, 2001). Cows were milked twice daily and milk yield and composition (fat
and protein) were measured once monthly until the end of lactation. Body condition score (BCS) was determined at -7 ± 4, and exactly at 30 and 60 days postpartum (dpp) using a 5- point scale [36]. At the same time, blood samples for metabolites and hormones analyses were collected by coccygeal venupuncture into heparinized tubes from 94 cows, centrifuged at 3000 Xg for 20 min and plasma was stored frozen (-20°C) until further analysis. The breeding period consisted of 4 months from June to September. Oestrus was detected twice a day and cows were artificially inseminated (AI), 12 hours after heat detection by the same inseminator. Pregnancy diagnosis was performed by rectal palpation 45 days after AI.

**Farm 2**

Multiparous Holstein cows (n = 60) that calved between September and November were randomly selected from 450-cow herd; cows had 1 (L2, n = 36) or 2 (L3, n = 24) previous lactations. During the last month before calving, cows grazed on a native pasture and received 11 kg DM/cow/day of a diet composed of 7 kg DM of sorghum silage, 3 kg DM of sorghum grain, 1 kg DM of sunflower meal (36% crude protein) and 100 g of urea. After calving, cows received a commercial mineral supplement and were managed under a rotational grazing system with supplementary feed added to maintain a pasture forage availability of 1,200 kg DM and an estimated total intake of 18 kg DM/cow/day. The diet offered had 17% crude protein and 1.5 Mcal/kg DM of net energy of lactation (NRC, 2001). Cows were milked twice daily and milk yield and composition (fat and protein) were measured once weekly for the first month of lactation and afterwards monthly until the end of lactation. Cow BCS was determined every 15 days from -30 dpp to calving, and then once a week up to 60 dpp. The breeding period consisted of 3 months from September to November: during the first two months AI was used, insemination 12 hours after oestrus detection twice a day, and natural mating was used during the last month. Pregnancy diagnosis was performed as described in farm 1.

**Laboratory analysis**

Genotyping of GH and IGF-1 and hormone (insulin and IGF-1) analyses were performed at the Nuclear Techniques Laboratory (Veterinary Faculty, University of Uruguay, Uruguay), while metabolites analyses (NEFA and BHB) were performed at DILAVE Laboratory (Pando, Uruguay).

Extraction of DNA was performed according to Kawasaki [37] and DNA was stored frozen (-20°C) until further analysis. The GH-Alul genotype was determined by a polymerase chain reaction-restriction fragment length polymorphism (PCR-RFLP) according to Lucy et al. [18]. Primers designed to amplify a 428-bp sequence of the bovine GH gene, GH For.: 5'-CCGTGTCATGAGAACGC-3' and GH Rev.: 5'-TTCTTGAGCAGCGCGT-3' were used. A Digestion of PCR product was performed with 6U of AluI (Fermentas Inc., MD, USA) restriction endonuclease. Fragments of DNA were resolved in a 2% agarose gel stained with ethidium bromide (EtBr) and fragments of either leucine (L; 265, 96, 51 and 16 bp) or valine (V; 265, 147 and 16 bp) alleles were visualized under UV light (Cleaver Scientific, England).

The IGF-I-SnabI genotype was determined by PCR-RFLP according to Ge et al. [28]. Primers designed to amplify a 249-bp sequence of the bovine IGF-I gene, IGF-I For.: 5'-ATTCAAAAGCTGCCTGCCGCCC-3' and IGF-I Rev.: 5'-ACCTTACCCGTATGAAAGGAATATACGT-3' were used and PCR products digestion was performed with 5U of SnabI (Fermentas Inc., MD, USA) restriction endonuclease. The DNA fragments were resolved in a 3% agarose gel stained with EtBr and fragments of either T (A) (223 and 26 bp) or C (B) (undigested, 249 bp) alleles were visualized under UV light (Cleaver Scientific, England).

Plasma insulin concentrations were determined by a 125I-Insulin RIA kit (Diagnostic Products Co., Los Angeles, California, USA). The assay sensitivity was 1.3 μIU/mL and the intra-assay and inter-assay coefficients of variation were less than 8.2 and 10.1% for control 1 (4.2 μIU/mL) and 9.4 and 11.3% for control 2 (12.6 μIU/mL), respectively. Plasma IGF-I concentrations were determined by the IGFI RIACT (Cis Bio International, GIF SUR YVETTE CEDEX, France). The assay sensitivity was 16 ng/mL and the intra-assay coefficient of variation were 3.4 and 5.8% for control 1 (50.4 ng/mL) and 16 and 17% for control 2 (709 ng/mL), respectively.

Plasma NEFA and BHB concentrations were assayed by spectrophotometry using commercial kits: Kat. #FA 115 kit (Wako Chemicals, Richmond, VA, USA) and Kat. #RB 1007 (Randox Laboratories Ltd, Ardmore, UK), respectively. The intra-and inter-assay coefficient of variations for both metabolites were less than 7.3 and 9.7%, respectively.

**Statistical analyses**

Data were analyzed in a complete randomized design by farm using the SAS program (Statistical Analysis System; SAS Institute Inc., Cary, NC, USA). Univariate analyses were performed on all variables to identify outliers and inconsistencies and to verify normality of residuals. Production traits and hormone and metabolite concentrations were analyzed by repeated measures using the
The IGF-I genotype had no effect on the productive traits (calving-first service interval, number of services per conception and total pregnancy rate) as analyzed with a generalized linear model (GENMOD procedure) with a Poisson distribution and log transformation (calving-first service interval) or a binomial distribution and logit transformation (pregnancy rate). The model included the effect of GH and IGF-I genotypes as fixed effects and calving date as a covariate. Results are expressed as lsmeans (LSM) ± SE. For all results, means were considered to differ when P ≤ 0.05 and trends were identified when 0.05 < P < 0.10.

### Results

A chi² test showed that allele frequency and genotypes of GH and IGF-I were in Hardy-Weinberg equilibrium (P = 0.97) and did not differ between farms (P > 0.28). GH allele frequencies were L (0.84) and V (0.16), while IGF-I allele distribution were A (0.60) and B (0.40). The number of cows for each genotype was LL (n = 122), LV (n = 51) and VV (n = 4) for GH genotypes and AA (n = 63), AB (n = 98) and BB (n = 25) for IGF-I genotypes. Due to the unequal distribution of GH genotypes in our study (dominance of the L allele and low frequency of V allele) we exclude VV genotype from further analysis.

### Productive and reproductive responses

Correlations between predicted and observed values for all productive and reproductive variables were between 0.47 and 0.81. The GH genotype was not associated with productive variables in either of the farms (Tables 1 and 2, Figure 1A and 1B). While no effect of the interaction between GH genotype and dpp on productive variables was observed in farm 1 (primiparous cows), a trend was observed on 4%FCM yield (P = 0.09), as AB cows yielded more FCM than BB cows at 120 and 210 dpp (Figure 1C). In farm 2, IGF-I genotype tended (P = 0.09) to affect FCM yield (Table 1) as AB cows had greater FCM yield than AA cows (P = 0.03), while no differences were found between AB and BB cows. Fat-corrected-milk yield was numerically greater for BB than AA cows (21.9 vs. 19.7 ± 1.05 kg/d) but this difference did not reach significance (P = 0.16) (Table 3, Figure 1D).

The IGF-I genotype had a significant effect on calving-first service interval only in farm 1 (Table 1), as AB cows had a shorter interval than BB cows (Table 3). No observed in 4%FCM yield (P = 0.09), as AB cows yielded more FCM than BB cows at 120 and 210 dpp (Figure 1C). In farm 2, IGF-I genotype tended (P = 0.09) to affect FCM yield (Table 1) as AB cows had greater FCM yield than AA cows (P = 0.03), while no differences were found between AB and BB cows. Fat-corrected-milk yield was numerically greater for BB than AA cows (21.9 vs. 19.7 ± 1.05 kg/d) but this difference did not reach significance (P = 0.16) (Table 3, Figure 1D).

The IGF-I genotype had a significant effect on calving-first service interval only in farm 1 (Table 1), as AB cows had a shorter interval than BB cows (Table 3). No

### Tables

#### Table 1 F-tests of fixed effects included in the model for productive/reproductive parameters and metabolic/endocrine variables and BCS of Holstein cows under grazing conditions in two commercial farms

| Parameter             | Farm 1                          | Farm 2                          |
|-----------------------|---------------------------------|---------------------------------|
|                       | GH IGF-I dpp                     | GH IGF-I dpp                     |
| Milk (L)              | 0.43 ± 0.02 < 0.01              | 0.41 ± 0.04 < 0.01              |
| FCM (L)               | 0.54 ± 0.02 < 0.01              | 0.56 ± 0.04 < 0.01              |
| Total solids (kg)     | 0.50 ± 0.23 < 0.01              | 0.44 ± 0.13 < 0.01              |
| Calving 1st service (days) | 0.26 ± 0.01 -                | 0.23 ± 0.17 -                  |
| Service/conception    | 0.74 ± 0.68 -                  | 0.45 ± 0.90 -                  |
| Pregnancy rates       | 0.97 ± 0.96 -                  | 0.80 ± 0.97 -                  |
| BCS                   | 0.42 ± 0.58 0.86              | 0.42 ± 0.95 0.01              |
| BHB (mmol/L)          | 0.31 ± 0.01 0.09              | 0.86 ± 0.77 0.01              |
| NEFA (mmol/L)         | 0.01 < 0.01 0.06              | 0.77 ± 0.44 0.01              |
| Insulin (µIU/mL)      | 0.99 0.02 < 0.01              | 0.53 ± 0.91 0.01              |
| IGF-I (ng/mL)         | 0.09 0.34 < 0.01              | 0.85 ± 0.95 0.01              |

Fixed effects are GH and IGF-I genotype and days post partum (dpp)

#### Table 2 Productive/reproductive parameters and metabolic/endocrine variables (LSM ± SE) for GH genotypes of Holstein cows in two commercial farms

| Parameter             | Farm 1 | Farm 2 |
|-----------------------|--------|--------|
|                       | GH genotype | GH genotype |
|                       | LL | LV | SE | LL | LV | SE |
| Milk (L)              | 17.9 | 17.5 | 0.50 | 23.1 | 22.1 | 0.80 |
| FCM (L)               | 15.9 | 15.5 | 0.45 | 21.6 | 21.0 | 0.90 |
| Total solids (kg)     | 1.14 | 1.11 | 0.03 | 1.60 | 1.54 | 0.06 |
| Calving 1st service (days) | 2.3 | 2.2 | 0.2 | 1.3 | 1.6 | 0.1 |
| Service/conception    | 80 | 78 | 7 | 72 | 64 | 10 |
| Pregnancy rates       | 0.93 | 0.41 | 0.02 | 0.34 | 0.33 | 0.03 |
| BHB (mmol/L)          | 0.25 | 0.26 | 0.01 | 0.63 | 0.60 | 0.06 |
| NEFA (mmol/L)         | 3.5 | 3.5 | 0.23 | 2.60 | 2.81 | 0.38 |
| Insulin (µIU/mL)      | 98.5 | 79.9 | 7.45 | 113.64 | 113.92 | 11.14 |
| BCS                   | 2.9 | 2.7 | 0.2 | 3.04 | 3.00 | 0.04 |

* Values marked with different letters are significantly different at P ≤ 0.05

Values marked with different letters are a trend at P < 0.10
other effect of IGF-I genotype was observed on reproductive variables (Table 3).

**Metabolic and endocrine profiles**

In farm 1, the GH genotype affected or tended ($P = 0.09$) to affect plasma NEFA and IGF-I concentrations, respectively, but did not affect BCS or any other metabolic parameters (Table 1, Figure 2 A-H). Cows carrying LV genotype had greater plasma NEFA and tended to present lower IGF-I concentrations than LL cows (Table 2). Although plasma IGF-I concentrations decreased ($P < 0.01$) after calving in both genotypes, LV cows presented lower IGF-I concentrations at 30 and 60 dpp ($P < 0.02$) than LL cows (Figure 2G). In farm 2, GH genotype did not affect BCS or any of the endocrine/metabolic profiles (Table 1).

The IGF-I genotype affected BHB, NEFA, and insulin concentrations in farm 1 (Table 1) as BB cows presented lower plasma NEFA and BHB and greater insulin concentrations than AA and AB cows (Table 3, Figure 3 A, C, E). While insulin concentrations declined ($P < 0.01$) from -7 to 30 and 60 dpp for AA and AB cows, plasma insulin was maintained during the study in BB cows; being insulin concentrations at 30 dpp greater in BB than AA and AB cows ($P < 0.01$) (Figure 3 E). In farm 2, the interaction between IGF-I genotype and dpp tended ($P = 0.06$) to affect IGF-I concentrations as AA cows tended ($P < 0.07$) to present lower prepartum IGF-
suggested that the L allele appeared to have an additive effect on milk production only at the beginning of the lactation. Besides, Lucy et al. [18] reported that cows carrying LL genotype yielded more milk, fat, and protein than LV cows. No effect of the interaction between GH genotype and dpp was found in primiparous cows (farm 1), and this could be associated with the level of production and/or a differential role of GH genotype in growing animals. In contrast, Dybus et al. [20] determined an effect of GH genotype on milk, fat and protein yield in primiparous but not in multiparous cows, and they suggested that the observed differences could have resulted from another source of variation (e.g. effects of herd, sires) not considered in the study.

The IGF-I genotype tended to affect FCM yield in farm 2 (multiparous cows), and the interaction between IGF-I genotype and dpp tended to affect FCM yield in farm 1 (primiparous cows), as AB cows yielded more FCM than BB (farm 1) or AA (farm 2) cows. Similarly, Siadkowska et al. [29] reported that AB cows yielded more FCM than AA and BB cows. In contrast, Hines et al. [30] did not find any effect of IGF-I genotype on productive parameters. In our study, cows were under grazing conditions and were average producing cows (17 L/day in farm 1 and 22 L/day in farm 2). Previous studies that could not find any effect of the genotype on milk production stated that genotype differences might not be expressed at this level of production [38,27]. In addition to this, Chilibroste et al. [39] and Kolver and Muller [40] reported that DM intake is not enough to achieve the genetic potential on grazing milk production systems.

In our study there was no effect of GH genotype on reproductive parameters in none of the farms. Balogh et al. [27] found no effect of the GH genotype on the time of the first postpartum ovulation. Lechniak et al. [25] reported a tendency for greater non-return rates of VV beef bulls at 60 dpp and Lechniak et al. [26] found no effect of the GH genotype on oocyte number. No data as such has been found for the relationship between IGF-I genotype and reproduction in dairy cows. For IGF-I genotype there was a significant effect on calving-first service interval only in farm 1, as BB cows had a longer interval than AB cows.

We found only one report regarding the effects of GH genotype [33] and none of IGF-I genotype on metabolic and/or endocrine profiles in the transition dairy cow. Balogh et al. [33] did not found either an effect of GH genotype on plasma BHB, insulin, and IGF-I concentrations, but they performed only one postpartum determination (4 to 13 dpp). In the present study we have included pre and postpartum determinations which in our understanding, allowed a better comprehension of the metabolic endocrinology during the peripartum period.
In farm 1 (primiparous cows), NEFA and IGF-I concentrations were affected by GH genotype, as LL cows had lower NEFA and greater IGF-I concentrations than LV cows. Since NEFA and IGF-I are both indicators of the metabolic status [5], these data suggest that LL cows presented a better energy status than LV cows. It is supposed that bovine GH with Leu127 stimulate the release of IGF-I more than other variants of bGH [41] which is consistent with the results found in the present study. In contrast, Schlee et al. [32] observed that Simmental
LV bulls presented greater IGF-I concentrations. This differential metabolic/endocrine environment was not reflected on productive/reproductive traits, which could be due to the level of production of primiparous cows as discussed before and/or to the extra energy demands for growth in these cows. In farm 2 (multiparous cows) there was no effect of GH genotype on any of the metabolites and hormones studied. Although a reduced number of animals were included in this farm, there are more metabolic/endocrine time measurements which allowed a better metabolic description of the NEB. Indeed, NEFA and \( \beta \)-hydroxybutirate concentrations increased around calving reflecting fat mobilization as reported before [1-3]. As expected, insulin
concentrations decreased around calving as has previously been observed [13]. This decrease in plasma insulin is a metabolic adaptation to cope with the energy demands of lactation as reported earlier [42,43], since low insulin concentrations favours gluconeogenesis and lipolysis [44] (e.g. homeorhetic effect). The decrease in IGF-I concentrations at calving confirmed the uncoupled somatotropic axis (GH-IGF-I), which mediates nutrient partitioning for lactogenesis [5]. We have no obvious explanation for the differential effect of GH genotype on metabolic/endocrine profiles found in primiparous cows (farm 1) vs. multiparous cows (farm 2), but as previously suggested it could be due to the differential role of GH during growth and development. Indeed, primiparous cows present greater insulin and IGF-I concentrations than multiparous cows [42].

Insulin-like growth factor-I genotype affected NEFA and BHB concentrations in farm 1 (primiparous cows); as BB cows had lower concentrations than AA and AB cows. In accordance to the better energy balance in BB cows reflected by these metabolites, these cows presented greater insulin concentrations at 30 dpp. Unexpectedly, IGF-I concentrations were not affected by IGF-I genotype. Ge et al. [28] and Maj et al. [34] reported lower or greater IGF-I blood concentrations in BB young Angus cattle or BB Holstein -Friesian young bulls and heifers, respectively. Since this polymorphism is located in the promoter region of the IGF-I gene, a variety of responses in gene expression may result depending on the physiological and/or nutritional status of the animal. This differential energy balance is not consistent with the calving first service interval, since it is known that cows in a better energy balance have also better reproductive performance [45], and in this study BB cows presented longer calving-first service interval than AB cows (103 vs 73 days, respectively). Unfortunately, we do not have the endocrine/metabolic profiles at the time of the initiation of the services which could clarify these contradictory results; indeed the endocrine system changes dynamically according to the nutritional and productive status. Moreover, BB cows had not only a reduced reproductive performance but they tended to yield less FCM at 120 and 210 dpp than AB cows. Staples and Thatcher [46] reported that cows with more DM intake, present not only greater milk production, but also better reproductive performance. In farm 2 (multiparous cows), there was no effect of IGF-I genotype on plasma NEFA, BHB, and insulin concentrations. On the other hand, IGF-I profiles suggest a greater uncoupling of the somatotropic axis in AB and BB cows than AA cows which is in accordance with the greater FCM yield of AB than AA cows.

**Conclusions**

In summary, the GH - Alul and IGF-I - Snal1 genotypes did not have a relevant effect on productive parameters, although the latter genotype affected calving-first service in primiparous cows. On the other hand, this study demonstrated that these genotypes do alter the endocrine and metabolic profiles of the transition dairy cow under grazing conditions.

**Acknowledgements and funding**

The present study received financial support from the National Institute of Agricultural Research to A.M. (INIA FFTA 214). We would like to thank Dr. G. Uriarte and P. Nicolini for their technical advice.

**Author details**

1. Faculty of Veterinary Medicine and Agronomy Sciences, University of Uruguay, Montevideo, Uruguay. 2. University of the Enterprise, Montevideo, Uruguay.

**Authors’ contributions**

GR lead the experimental designs, carried out the hormone and the metabolites determinations and genotyping, and drafted the manuscript. IP and JM contributed with the experimental designs. MC contributed with the statistical analysis and helped together with AM on data interpretation and manuscript corrections. All authors read and approved the final manuscript.

**Competing interests**

The authors declare that they have no competing interests.

Received: 12 April 2011 Accepted: 2 June 2011 Published: 2 June 2011

**References**

1. Bell AW: Regulation of organic nutrient metabolism during transition from late pregnancy to early lactation. *J Anim Sci* 1995, 73:2804-2819.
2. Chilliard Y: Metabolic adaptations and nutrient partitioning in the lactating animal. In Biology of Lactation. Collection Mieux Comprendre Edited by: INRA, Paris (FRA), Martinet J, Houdéleme LM, Head HH 1999, 503-552.
3. Fenwick MA, Fitzpatrick R, Kenny DA, Diskin MG, Patton J, Murphy JJ, Watkins DC: Interrelationships between negative energy balance (NEB) and IGF regulation in liver of lactating dairy cows. *Domest Anim Endocrinol* 2008, 34:31-44.
4. Svennersten-Sjauinja K, Olsson E: Endocrinology of milk production. *Domest Anim Endocrinol* 2005, 29:241-258.
5. Lucy MC, Veerkerk GA, Whyte BE, Maconald KA, Burton L, Cursons RT, Roche JR, Holmes W: Somatotropic axis components and nutrient partitioning in genetically diverse dairy cows managed under different feed allowances in pasture system. *J Dairy Sci* 2009, 92:1526-39.
6. Block SS, Butler WR, Ethhardt RA, Bell AW, Van Amburgh ME, Bosclar YR: Decreased concentration of plasma leptin in periparturient dairy cows is caused by negative energy balance. *J Endocrinol* 2001, 171:339-348.
7. Ingvarsen KL: Feeding- and management-related diseases in the transition cow. Physiological adaptations around partum and strategies to reduce feeding-related diseases. *Anim Feed Sci and Technology* 2006, 126:175-213.
8. Gong JG, Lee WJ, Garnsworthy PC, Webb R: Effect of dietary-induced increases in circulating insulin concentrations during the early postpartum period on reproductive function in dairy cows. *Reproduction* 2002, 123:419-427.
9. Huseiniczia Gy, Haraszi J, Molnár’ L, Solti L, Fekete S, Eleki K: Some metabolic characteristics of dairy cows with different post partum ovarian function. *J Vet Med* 1988, 35:506-15.
10. Butler WR, Smith RD: Interrelationships between energy balance and postpartum reproductive function in dairy cattle. *J Dairy Sci* 1989, 72:767-83.
11. Staples CR, Thatcher WW, Clark JH: Relationship between ovarian activity and energy balance during the early postpartum period of high producing dairy cows. *J Dairy Sci* 1990, 73:938-47.
12. Rhodes FM, Clark BA, Nation DP, Taufa VK, MacMillan KL, Day ML: Factors influencing the prevalence of postpartum anoestus in New Zealand dairy cows. *Proc NZ Soc Anim Prod* 1998, 58:79-81.
13. Mekle A, Kúlsar M, Chilliard Y, Febel H, Delavaud C, Cavestany D, Chilibroste P: Effects of parity and Body Condition Score at Calving on Endocrine and Reproductive Parameters of the Dairy Cow under Grazing Conditions. Reproduction 2004, 127:721-737.

14. Summantran VH, Tsai ML, Schwanitz J: Growth hormone induces c-fos and c-jun expression in cells with varying requirements for differentiation. Endocrinol 1992, 130:2016-2024.

15. Lincoln DT, Snawatz F, el-Hifnawi E, Hughes RL, Waters M: Evidence of a direct role for growth hormone (GH) in mammary gland proliferation and lactation. Anat Histol Embryol 1995, 24:107-115.

16. Lucy MC, Hauser SD, Eppard PJ, Kriev GG, Clark JH, Bauman DE, Collier RJ: Variants of somatotropin in cattle: gene frequencies in major dairy breeds and associated milk production. Domest Anim Endocrinol 1993, 10:325-333.

17. Furu LM, Katsmter GW, Zinn SA, Rynowich S: Somatotropin Mspl and Alul polymorphism relative to indicators of the genetic merit of Holstein Aisire. J Anim Sci 1988, 76:75.

18. Lucy MC, Hauser SD, Eppard PJ, Kriev GG, Collier RJ: Genetic polymorphism within the bovine somatotropin (bST) gene detected by polymerase chain reaction and endonuclease digestion. J Dairy Sci 1991, 74(Suppl 1):284.

19. Shariflou MR, Moran C, Nicholas FW: Association of the Leu (127) variant of the bovine growth hormone (bGH) gene with increased yield of milk, fat, and protein in Australian Holstein-Friesians. Australian J Agric Res 2000, 51:515-522.

20. Dybus A: Associations between Leu/Val polymorphism of growth hormone gene and milk production traits in Black-and-White cattle. Arch Tierz 2002, 45:421-428.

21. Grochowska R, Sarsens P, Zwierzchowski L, Snuchowski M, Levendahl P: Genetic variation in stimulated GH release and in IGFI of young dairy cattle and their associations with the leucine/valine polymorphism in the GH gene. J Anim Sci 2001, 79:450-476.

22. Zwierzchowski L, Krzyzewski J, Straszewska N, Sidakowska E, Ryniewicz Z: Effects of polymorphism of growth hormone (GH), Pit-1, and leptin (LEP) genes, cow’s age, lactation stage and somatic cell count on milk yield and composition of Polish Black-and-White cows. Anim Sci Pap Rep 2002, 20:213-227.

23. Kovács K, Völgyi-Cik J, Zsolnai I, Győrök I, Fésüs L: Associations between the Alu polymorphism of growth hormone gene and production and reproduction traits in a Hungarian Holstein-Friesian bull dam population. Arch Tierz 2006, 49:265-249.

24. Yao J, Aggrey SE, Zadworny D, Hayes JF, Kuhenlein U: Sequence variations in the bovine growth hormone gene characterized by singlestrand conformation polymorphism (SSCP) analysis and their association with milk protein traits in Holsteins. Genetcs 1996, 144(4):1809-1816.

25. Lechniak D, Machnik G, Szadkowski M, Switkowski M: Growth hormone gene polymorphism and reproductive performance of AI bulls. Theriogenology 1999, 52:145-152.

26. Lechniak D, Adamowicz T, Stanislawska D, Kaczmarek D: In vitro maturation and fertilization of bovine oocytes in relation to GH gene polymorphism (Leu/Val) Reprod Nutr Dev 2002, 42:275-280.

27. Balogh O, Kovács K, Kúlsar M, Gáspárady A, Zsolnai A, Kárai L, Pécsi A, Ferüsz L, Butler WR, Huszenicza Gy: Alul polymorphism of the bovine growth hormone (GH) gene, resumption of ovarian cyclicity, milk production and loss of body condition at the onset of lactation in dairy cows. Clomining 2000, 71:553-559.

28. Ge W, Davis ME, Hines HC, Irvin KM, Simmon RCM: Association of genetic markers with blood serum insulin like growth factor-I concentration and growth traits in Angus cattle. J Anim Sci 2001, 79:1757-1762.

29. Sidorakowa E, Zwierzchowski L, Oprzaddek J, Straszewska N, Baginska E, Krzyzewski J: Effect of polymorphism in IGF-I gene on production traits in Polish Holstein-Friesian cattle. Anim Sci Pap Rep 2006, 3:225-236.

30. Hines HC, Ge W, Zhao Q, Davis ME: Association of genetic markers in growth hormone and insulin-like growth factor-I loci with lactation traits in Holsteins. Anim Genetics 1998, 29(Suppl 1):161.

31. Li C, Basarab J, Smelting WM, Benkel B, Murdnoch B, Hansen C, Moore SS: Assessment of positional candidate genes myf5 and IGF1 for growth on bovine chromosome 5 in commercial lines of Bos taurus. J Anim Sci 2004, 82:1897-1904.

32. Schlée P, Gräml R, Schallenberger E, Schams D, Rottmann Q, Olbrich-Biuda A, Pirchler F: Growth hormone and insulin-like growth factor-I concentrations in bulls of various growth hormone genotypes. Theor Appl Genet 1994, 88:497-500.

33. Balogh O, Kovács K, Kúlsar M, Gáspárady A, Febel H, Zsolnai A, Fésüs L, Delavaud C, Chilliard Y, Gilbert RO, Huszenicza Gy: Interrelationships of growth hormone Alu polymorphism and hyperketonemia with plasma hormones and metabolites in the beginning of lactation in dairy cows. Livestock Science 2009, 123:180-186.

34. Maj A, Snochowski M, Sidakowska E, Rowinska B, Lisowski P, Robakowska-Hyzoerek D, Oprzaddek J, Grochowska R, Kochman K, Zwierzchowski L: Polymorphism in genes of growth hormone receptor (GHR) and insulin-like growth factor-I (IGF1) and its association with both the IGF1 expression in liver and its level in blood in Polish Holstein-Friesian cattle. Neuro Endocrinol Lett 2008, 29(6):981-989.

35. Ruprechter G, Nicolini P, Mekle A, Carriquiy M: Marcadores moleculares de hormona de crecimiento y factor de crecimiento similar a la insulina-1 como predictores del desempeño productivo en vacas Holando bajo condición pastoril. Veterinaria. Montevideo 2009, 45:173-176.

36. Ferurgeon JD, Galligan DT, Thomsen N: Principal descriptors of Body Condition Score in Holstein Cows. J Dairy Sci 1994, 77:2695-2703.

37. Kawasaki ES: Simple preparation from blood, cells and others fluids. In PCR Protocols: A guide to Methods and application. Edited by: Innis MA, Gelfand DH, Sninsky JJ, White TJ. New York: Academic Press; 1990:3-12.

38. Lee BK, Lim GJ, Cooker BA, Murgauma NP, Hansen LB, Chester-Jones H: Association of somatotropin (bST) gene polymorphism at the 5th exon with selection for milk yield in Holstein cows. Domest Anim Endocrinol 1996, 13:373-381.

39. Chilibroste P, Mekle A, Mattaouda DA, Bentancor C, Soca P: The American Holstein Dairy Cow During Early Lactation: Grazer or Browser? In: An overview of research and pastoral-based system in the Southern part of South America. 1 edition. Edited by: Machado C, Wade M Carneiro Da Silva S, Aguensni M, De Faccio Carvalho M, Morris S, Beskow W. Tandil: Universidad Nacional del Centro de la Provincia de Buenos Aires; 2010:154-167, ISBN 978-950-658-239-5.

40. Kolver ES, Muller LD: Performance and Nutrient Intake of High Producing Holstein Cows Consuming Pasture or a Total Mixed Ration. J Dairy Sci 1998, 81:1403-1411.

41. Vanderkoor W, Vandeheij MA, Sharma BK, Binelli M, Tucker HA, Aikins RM, Moseley WM: Comparison of growth hormone-releasing factor and somatotropin: the somatotropic axis in lactating primiparous cows. J Dairy Sci 1995, 78:2140-2149.

42. Taylor VJ, Cheng Z, Pushpakumara PGA, Beever DE, Whetts DC: Relationships between the plasma concentration of insulin-like growth factor-I in dairy cows and their fertility and milk yield. The Veterinary Record 2004, 155:583-588.

43. Watkins DC, Bourne N, Cheng Z, Mann GE, Taylor VJ, Coffey MP: Multiple correlation analysis of metabolic and endocrine profiles with fertility in primiparous and multiparous cows. J Dairy Sci 2007, 90:1310-1325.

44. Herst T: Ruminat adaptation to negative energy balance: influence on the etiology of ketosis and fatty liver. Metabolic disorders of ruminants. Veterinary Clinical North American. Food Animal Practice 2000, 16(2):215-230.

45. Lucy MC: Reproductive loss in high-producing dairy cattle: where will it end? J Dairy Sci 2001, 84:1277-1293.

46. Staples CR, Thatcher WW, Clark JH: Relationship between ovarian activity and energy status during the early postpartum period of high producing dairy cows. J Dairy Sci 1990, 73:938-947.