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Intra-flock variability in the body reserve dynamics of meat sheep by analyzing BW and body condition score variations over multiple production cycles

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Breeding for resilience requires a better understanding of intra-flock variability and the related mechanisms responsible for robustness traits. Among such traits, the animals' ability to cope with feed fluctuations by mobilizing or restoring body reserves (BR) is a key mechanism in ruminants. The objective of this work was to characterize individual variability in BR dynamics in productive Romane ewes reared in extensive conditions. The BR dynamics profiles were characterized by combining individual longitudinal measurements of BW and body condition scores (BCS) over several production cycles. Historical data, including up to 2628 records per trait distributed in 1146 ewes, underwent cluster analysis. Two to four trajectories were observed for BW depending on the cycle, while three trajectories were found for BCS, whatever the cycle. Most trajectories suggested that BR dynamics were similar but the level of BR may differ between ewes. Nevertheless, some trajectories suggested that both BR dynamics and levels were different for a proportion of ewes. Clustering on BW and BCS profiles adjusted for individual level trends, resulted in differences only in the level of BW or BCS, rather than differences in trajectories. Thus, the overall shape of trajectories was not changed considering or not the individual level trend across cycles. In addition to individual variability, the ewe's age at first lambing and litter size contributed to the distribution of the ewes between the trajectories. Regarding the entire productive life, three trajectories were observed for BW and BCS changes over three productive cycles. Increase in BW at each cycle suggested that ewes kept growing up until 3 to 4 years old in our conditions. Similar alternation of BCS gains and losses across cycles suggested BR dynamics might be repeatable. Many individual trajectories remained the same throughout a ewe's life, whatever the age at first lambing, parity or litter size. Our results demonstrate the relevance of using BW and BCS changes for characterizing the diversity of BR mobilization-accretion profiles in sheep in a long timespan perspective.

Keywords: body reserves, cluster analysis, longitudinal trajectories, ewe, Romane

Implications

We have demonstrated the existence of intra-flock variability in the dynamics of body reserves (BR) in extensively reared meat ewes over several production cycles. Our ultimate goal is to provide the basis for a future genetic improvement program for this character in sheep. These findings confirm our interest in such traits and represent a novel and significant step forward in improving sheep robustness.

Introduction

Current global livestock challenges require a shift toward more sustainable farming systems that meet economic, ecological and social demands while being able to overcome the unpredictable situations caused by climate change, financial crises or fluctuating market prices (Bocquier and González-García, 2010; Dumont *et al.*, 2014; Rojas-Downing *et al.*, 2017). Future production systems must thus change both their vision of farming and their implicit set of management practices, which includes working with resilient flocks (i.e. robust animals that combine productivity and adaptive capacity).

In this context, ruminant farming systems should rely in particular on the efficient use of natural resources by these animals, including in regions where weather conditions are

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less predictable. The dynamics of BR (i.e. lipid mobilization and accretion processes) could be considered a relevant trait. This trait characterizes the capacity of ruminants to efficiently overcome negative energy balance periods caused by feed shortages, forage seasonality (e.g. dry periods) or the requirements of specific physiological stages (e.g. late pregnancy, early lactation or suckling; Bauman and Currie, 1980; Nielsen et al., 2003: González-García et al., 2014). Such BR dynamics are linked in some extent to the BW and body condition score (BCS), which are typical and relatively easily monitored parameters (Molina et al., 1994; Álvarez-Rodríguez et al., 2012; Puillet and Martin, 2017). In sheep, numerous biotic and abiotic factors play significant roles in the BR dynamics of females, including feed availability, age, parity and litter size (Mendizabal et al., 2011; González-García et al., 2014 and 2015).

To our knowledge, no studies have investigated the different types of BW and BCS profiles in sheep longitudinally (i.e. covering several full production cycles throughout the ewes' lives). Furthermore, the accepted concept of 'breeding for robustness' justifies the necessity of gathering new insights into BR dynamics in order to characterize the degree of robustness at the individual level and the underlying determinant mechanisms (De La Torre *et al.*, 2015; Friggens *et al.*, 2017).

Considering the current lack of reports for grazing ruminants on this subject, and more specifically for sheep, our objective was to characterize the existing profiles of BR dynamics of ewes over successive production cycles and to evaluate the occurrence of intra-flock variability. For this, we used historical data (2002–2015) for BW and BCS that were recorded longitudinally in Romane meat ewes reared under extensive rangeland conditions. We hypothesized the existence of different BR dynamics subgroups within the same flock, whatever the production cycle and although managed in the same way. Such intra-flock variability would be affected both by the individual effect and to some extent by other biological factors such as the age at first lambing, parity, litter size or year of measurement.

Material and methods

Animals and experimental farming system

The BW (kg) and BCS measurements were collected from the Romane sheep flock at the INRA Experimental Farm La Fage (Causse du Larzac 43°54'54.52'' N; 3°05'38.11'' E; altitude approx. 800 m; Roquefort-sur-Soulzon, Aveyron), in the south of France. This flock consists of approximately 250 ewes with an annual renewal rate of 30%. The ewes are involved in a large number of genetic protocols and therefore rarely remain in the flock for more than three or four production cycles. The Romane breed is a hardy, prolific breed (Molénat *et al.*, 2005) obtained from an ancestral crossbred of the Romanov and the Berrichon-du-Cher (Ricordeau *et al.*, 1992). Before 2010, first mating of females occurred at 7 months of age for heavier ewe lambs (70%) and 19 months for slighter ewes (30%). After 2010, the reproduction system was changed and all females were aged 19 months at first mating (González-García and

Hazard, 2016). Mating took place in the autumn to obtain a peak of lambing at the beginning of the spring (usually mid-April). Weaning occurred at \sim 75 days.

Ewes were reared exclusively outdoors on about 280 ha of rangelands, a limestone plateau with thin soil covered by vegetation composed of about 25% of shrubs and 75% of grass. The rangelands included fertilized paddocks (15 ha) that were used during the suckling period (Molénat *et al.*, 2005). Further details on the climatic conditions and overall management practices used in this system were reported by Molénat *et al.* (2005), González-García *et al.* (2014) and González-García and Hazard (2016).

Historical database and variables

Individual BW and BCS measurements were recorded every year for the totality of the flock between 2002 and 2015. Data were entered in the INRA national database GEEDOC (https://germinal.toulouse.inra.fr/ ~ mcbatut/GEEDOC/). Individual BWs were measured using a conventional fixed scale balance. The BCS measurements were always performed by the same two trained observers and recorded giving a score on a scale from 1 to 5 with 0.1 increments, adapted from the original six-point scale described by Russel *et al.* (1969). Only ewes suckling at least one lamb over the entire suckling period were kept in the analyses. None of the data processed in the current study were deposited in an official repository.

During each production cycle, seven BW and eight BCS records were collected at regular intervals depending on the physiological stage. The BW records were as follows: (1) at mating (15 days before, November; BW-M), (2) during early pregnancy $(40 \pm 15 \text{ days after mating (DAM)})$, December; BW-Pa), (3) at mid-pregnancy (80 ± 14 DAM, January; BW-Pb), (4) at lambing $(160 \pm 16 \text{ DAM}, \text{ April; BW-L})$, (5) at the beginning of suckling $(190 \pm 14 \text{ DAM}, \text{ April; BW-Sa})$, (6) at weaning $(250 \pm 16 \text{ DAM}, \text{ June}; \text{BW-W})$ and (7) at the end of the dry-off period $(310 \pm 12 \text{ DAM}, \text{August}; \text{BW-D})$. Similarly, eight BCS records were collected from mating to the dry-off stage as follows: (1) at mating (15 days before, November; BCS-M), (2) during early pregnancy (40 ± 15 DAM, December; BCS-Pa), (3) at mid-pregnancy (80 ± 14 DAM, January; BCS-Pb), (4) at lambing (160 \pm 16 DAM, April; BCS-L), (5) at the beginning of suckling $(190 \pm 14 \text{ DAM}, \text{April}; \text{BCS-Sa}), (6)$ at the end of suckling $(230 \pm 15 \text{ DAM}, \text{ June; BCS-Sb})$, (7) at weaning $(250 \pm 16 \text{ DAM}, \text{ June}; \text{ BCS-W})$ and (8) at the end of the dry-off period (310 \pm 12 DAM, August; BCS-D). Hence, up to 2628 records were recorded for each trait over this period, distributed in 1146 ewes. All these ewes were recorded during their first production cycle, among them 1068 were also recorded during their second production cycle and 414 were recorded during their third production cycle.

An adjusted trait was also calculated for BW and BCS in order to take into account the average individual level of the trait across cycles, as follow:

$$y'_{ijk} = y_{ijk} - \left[\frac{1}{mn} \sum_{(j=1)}^{m} \sum_{(k=1)}^{n} y_{ijk}\right]$$
 (1)

where y'_{ijk} was the adjusted trait of ewe *i* at physiological

stage *j* and cycle *k*, y_{ijk} was the raw value of the trait of ewe *i* at physiological stage *j* and cycle *k*, *j* is the physiological stage and *k* is the number of the cycle, *m* was the total number of measurements within cycle and *n* the total number of cycles for a given ewe.

Descriptive statistics

Deviations from normality were tested using the Kolmogorov-Smirnov test (Univariate procedure of SAS; SAS Institute Inc., Cary, NC, USA). None of the raw variables were transformed. All biologically and environmentally relevant effects and interactions were tested using analyses of variance taking into account the repeated measures (MIXED procedure of SAS software, version 9.4; SAS Institute Inc., Cary, NC, USA) to identify factors of variation on BW and BCS. Four fixed effects were tested in the study: the Age at first lambing and the Parity or production cycle of the ewe, the Litter size and the Year of measurement. The Age at first lambing effect took into account the age at which ewes lambed for the first time, which was related to the age at their first mating (i.e. 7 or 19 months old; González-García and Hazard, 2016): 1 or 2 years of age (classes 1 and 2, respectively). The Parity effect took into account first, second and third lambing (classes 1, 2 and 3, respectively). The Litter effect depended on the number of lambs born and kept with the dam during the suckling stage (i.e. class 1, singletons from lambing until weaning; class 2, ewes lambing twins and suckling one; class 3, ewes lambing and suckling twins; class 4, ewes lambing and suckling more than two lambs). The litter effect considered only the litter size of the corresponding cycle and not of the previous cycle, which means that, for a given ewe, Litter can vary between cycles. Finally, 14 Years were analyzed. The first-order interactions between Age \times Litter and Parity \times Litter were tested. An effect was considered significant if P < 0.05.

Clustering of individual profiles

Cluster analysis was performed in order to investigate the variability of individual BW and BCS profiles for each cycle without any assumptions as to factors of variation. Clustering could also create additional profiles for BW and BCS different from those described for the factors of variation. A first analysis was done within cycle with all the ewes present at the considered cycle. A second analysis was performed over the entire productive life (i.e. three cycles in our conditions) with all the ewes that completed three cycles. A third analysis was carried out with only ewes lambing and suckling twins in order to investigate clusters without the potential influence of the litter size. We choose ewes suckling twins because it was the group of ewes the best distributed across the three cycles.

A principal component (PC) analysis was performed on the BW and BCS data set, for each production cycle and for the entire lifetime, using the R package fdapace (Dai *et al.*, 2017). Individual profiles were smoothed on a defined number of nodes from the seven BW or eight BCS measurements recorded for each individual. This smoothed mean curve was calculated with fdapace, by using a local linear

Gaussian kernel regression between the seven or eight equidistant nodes that aggregate all the measurements together. The smoothed covariance matrix was used to perform eigen analyses in order to obtain the estimated PC scores (Yao et al., 2005). Then, using these PC scores, an unsupervised classification (i.e. a cluster analysis) was performed with the R package Rmixmod (Langrognet et al., 2016). This package allows to fit a mixture model of multivariate components to a data set, here the PC scores of the previous functional principal component analysis (FPCA). Each variance matrix of a cluster was decomposed in three types of parameters: one determining the orientation of the cluster, one determining its volume and the last one determining its shape (Lebret et al., 2014). The mixture parameter was obtained through the maximization of the log-likelihood by using the Expectation–Maximization (EM) algorithm. It provides iterative computation of the maximum likelihood estimation when the observed data are incomplete. Two to seven clusters were tested and the number of clusters selected was determined using Akaike information criteria (AIC) and Bayesian information criteria (BIC). The AIC (Akaike, 1974) is calculated as $AIC = 2k - 2 \times \ln(L)$ with k the number of parameters to estimate and *L* the maximum likelihood of the model. The BIC (Schwarz, 1978) is defined as $BIC = \ln(n) \times (k-2) \times \ln(L)$ with k the number of parameters and *n* the number of observations in the sample. The best model is the one with the lowest criteria value, but they do not determine the absolute adjustment quality of the model. The repeatability and stability of the clusters were also tested by repeating the EM algorithm three to four times. If the distribution of ewes between the clusters did not vary (i.e. within cycle for the first and second analyses), then the clusters were validated.

Once the optimal number of clusters was determined, each cluster was characterized depending on known biological and environmental effects. The best fitting GLM to explain the distribution of ewes between clusters within cycle was analyzed with:

Y = age at first lambing + litter + year + litter at previous cycle

where *Y* is the number of the ewes' cluster, *age at first lambing* the effect of the ewe's age at first lambing, *litter* the effect of the litter size and *year* the effect of the year of measurement as above described. The *litter at previous cycle* is the litter size of the ewe at cycle n - 1. This effect was not included in the model at cycle 1. The contribution of the effects to the model was estimated by comparing AIC and BIC criteria reach when the effect was removed from the model to those reach with the complete model.

The number of animals present in each cluster was expressed as the percentage of ewes for each cycle. The repeatability of cluster classes for production cycle n was given by indicating the proportion of animals from the clusters observed for the previous cycle n - 1.

Results

Preliminary results of the current work have been published in an abstract form (Macé et al., 2018a). The BW and BCS values measured in ewes at different physiological stages of the production cycle were significantly affected by the main fixed effects evaluated here (i.e. parity, age at first lambing, litter and year; Table 1). Globally, a significant increase in BW and BCS was observed as parity increased. On the contrary, an overall decrease in BCS was observed between mating and mid-pregnancy when comparing parity 1 and parity 2. The litter size effect on BW varied according to the physiological stage and type of litter (i.e. singletons or multiple lambs). The BCS, however, clearly decreased as the litter size increased (Table 1) except between mating (BCS-M) and early pregnancy (BCS-Pa). The age at first lambing had no significant effect on BW at BW-L, BW-Sa and BW-D and on BCS at BCS-M, BCS-Sa, BCS-Sb and BCS-W. Otherwise, BW and BCS increased as the age at first lambing increased. The year of measurement was always significant (P < 0.001) whatever the trait and the physiological stage. The interactions between fixed effects were not significant. When analyzing whole production cycles (i.e. from mating to dry-off), ewes seemed to gain weight from mating (BW-M, 51.76 ± 8.93) to mid-pregnancy (BW-Pb, 60.65 ± 9.40), then to decline and lose BW until the dry-off (BW-D, 54.75 ± 6.68), even if there is no significant effect of the overall physiological stage effect. Concerning BCS, ewes seemed to gain body condition from mating (BCS-M, 2.82 ± 0.24) to early pregnancy (BCS-Pa, 2.93 ± 0.25), and then to lose body condition until the end of suckling (BCS-Sb, 2.47 ± 0.22). Finally, ewes seemed to gain body condition after weaning (BCS-Sb, 2.50 ± 0.21 ; BCS-D, 2.59 ± 0.22).

Cluster analysis for BW and body condition score

Cluster analysis highlighted four clusters for production cycles 1 and 2 for BW whereas three clusters were found for adjusted BW. Two clusters were observed for BW and adjusted BW at cycle 3 (Figure 1) with 95% to 100% of variances explained by the two or three first PC of FPCA, for adjusted BW and BW, respectively (data not shown).

For BW at production cycles 1 and 2, the two major clusters for each cycle together included 85% and 91% of the ewes, respectively (clusters BW1, BW2 and BW5 and BW6). For adjusted BW at cycles 1 and 2, the two major clusters included 88% of the ewes (clusters BW11 and BW12) and 79% of the ewes (clusters BW14 and BW14), respectively. Cluster BW1 showed a BW gain from BW-M to BW-Sa and a stabilized BW from BW-Sa to the end of cycle 1. Clusters BW2 and BW3 showed similar profiles but ewes in cluster BW3 were heavier than ewes in cluster BW2. The BW2 and BW3 profiles were similar to BW1 from BW-M to BW-Pb, and were characterized by a decrease in BW from the end of pregnancy to BW-W and then an increase until the end of cycle 1. Ewes in BW2 were heavier than ewes in cluster BW1 until BW-Sa and then had the same BW until BW-W. Cluster BW11 showed a similar profile to BW2 and BW3. Cluster BW12

showed a combination of BW1 and BW4 profiles characterized by a progressive increase in BW from mating to beginning of suckling and a slight decrease thereafter. Cluster BW13 showed a similar profile to BW11 but with a greater increase of BW between BW-W and BW-D. For cycle 2, clusters BW5 and BW7 showed parallel profiles and were similar to cluster BW3 in cycle 1 except for a slight increase in BW from BW-L to BW-Sa and a slight decrease from BW-Sa to BW-D. Cluster BW6 showed a similar profile to cluster BW3 in cycle1 but a marked decrease in BW between BW-Pb and BW-W. Ewes in cluster BW5 were heavier than in cluster BW7, as were the ewes in BW6 compared to those in cluster BW7, except at BW-W when the ewes had a similar BW value. Cluster BW14 had a similar profile to BW6 with a slighter increase at the end of the cycle. Cluster BW15 showed a similar profile to BW5 but with a higher decrease at the end of the cycle. Cluster BW16 had a similar profile to BW7 but showed a more stable profile between BW-Sa and BW-D. Clusters BW9 and BW17 included 97% of the ewes measured for cycle 3 and showed a similar profile to clusters BW5, BW7 and BW14 in cycle 2 but with either lower BW values than BW5 (i.e. BW) or similar range of variation than BW14 (i.e. adjusted BW). Clusters BW4, BW8, BW10 and BW18 for cycles 1, 2 and 3, respectively, were different from the other clusters with a marked increase of BW at BW-Sa, with a gain of 10, 16, 18 and 23 kg, respectively, to finally reach higher levels than those measured in the other clusters. This increase was followed by a marked decrease between BW-Sa and BW-W with a final BW close to that of the other clusters.

Generally, the ewes in each cluster originated from different clusters in the previous cycle but some constancy was observed. For example, the ewes in the BW5 cluster of cycle 2 originated mostly from clusters BW1 (52%) and BW2 (28%) of cycle 1. The ewes belonging to BW2 and BW5 also belonged to BW9 and exhibited almost the same profiles at cycles 1, 2 and 3. The majority of the ewes in cluster BW6 originated from clusters BW2 (42%) and BW3 (30%) of cycle 1 (Figure 1). These ewes also belonged to BW9 and exhibited almost the same profiles at cycles 1, 2 and 3. The majority of the ewes in cluster BW14 originated mostly from cluster BW11 (60%), whereas ewes in cluster BW15 originated from cluster BW12 (63%). Most of the ewes that belonged to BW11 and BW14 also belonged to BW17 at cycle 3 and showed similar profile along the three production cycles.

Cluster analysis of BCS in the whole population highlighted three clusters for each production cycle (Figure 2) with 98% to 100% of variances explained by the two or three first PC of FPCA, for adjusted BCS and BCS, respectively (data not shown). Cluster analysis of adjusted BCS highlighted two clusters for each production cycle (Figure 2) with 86% to 90% of variances explained by the two first PC of FPCA. During production cycles 1 and 2 for BCS, the two major clusters for each cycle included 99% and 85% of the ewes, respectively (clusters BC1, BC2 and BC4, BC5, respectively). In production cycle 3, the major cluster (BC7) included 76% of ewes. For adjusted BCS at cycles 1 and 2, the major cluster

		Parity				Litter size					Age at first lambing			Year
Variables	п	1	2	3	Sign.	1	2	3	4	Sign.	1	2	Sign.	
n (%)		43.6	40.6	15.8		21.3	46.4	26.1	6.2		52.8	47.2		
BW-M	2627	45.8 (0.6)	54.9 (0.6)	60.7 (0.9)	***	53.5 (1.1)	52.6 (1.0)	54.1 (0.9)	55.0 (1.0)	***	51.8 (2.5)	55.8 (2.7)	***	* * *
BW-Pa	2385	48.5 (0.6)	57.0 (0.6)	62.9 (0.9)	***	55.9 (0.9)	55.1 (0.8)	56.4 (0.8)	57.1 (0.8)	***	54.6 (2.1)	57.6 (1.9)	* * *	* * *
BW-Pb	2378	53.7 (0.7)	63.2 (0.6)	69.6 (1.0)	***	60.9 (1.5)	61.4 (1.4)	62.3 (1.3)	64.2 (1.3)	***	60.7 (2.1)	63.7 (1.9)	* * *	* * *
BW-L	2573	52.1 (0.5)	59.6 (0.5)	64.2 (0.8)	***	59.9 (1.3)	57.4 (1.2)	59.2 (1.1)	58.0 (1.2)	***	58.4 (0.2)	58.8 (0.3)	NS	* * *
BW-Sa	2492	52.2 (0.5)	59.6 (0.5)	64.2 (0.8)	***	60.3 (1.4)	57.6 (1.2)	58.7 (1.2)	57.9 (1.2)	***	58.8 (0.2)	58.5 (0.2)	NS	* * *
BW-W	2594	50.1 (0.4)	57.1 (0.4)	61.1 (0.7)	***	57.8 (1.3)	56.0 (1.2)	55.2 (1.1)	55.2 (1.2)	***	56.3 (0.3)	55.8 (0.4)	*	* * *
BW-D	1883	51.9 (0.3)	58.4 (0.4)	60.7 (0.6)	***	58.1 (0.9)	56.7 (0.8)	56.4 (0.8)	56.8 (0.8)	***	57.0 (0.1)	57.0 (0.1)	NS	* * *
BCS-M	2448	2.92 (0.08)	2.75 (0.08)	2.79 (0.14)	***	2.83 (0.03)	2.85 (0.02)	2.80 (0.02)	2.80 (0.02)	***	2.81 (0.01)	2.83 (0.01)	NS	* * *
BCS-Pa	2627	2.97 (0.04)	2.89 (0.04)	2.93 (0.06)	***	2.94 (0.02)	2.95 (0.02)	2.92 (0.01)	2.92 (0.02)	*	2.90 (0.04)	2.96 (0.04)	***	***
BCS-Pb	2628	2.78 (0.04)	2.77 (0.04)	2.87 (0.06)	***	2.85 (0.04)	2.82 (0.03)	2.79 (0.03)	2.76 (0.03)	***	2.78 (0.04)	2.84 (0.04)	***	***
BCS-L	2628	2.56 (0.05)	2.62 (0.05)	2.69 (0.08)	***	2.74 (0.09)	2.61 (0.08)	2.63 (0.08)	2.52 (0.08)	***	2.61 (0.02)	2.64 (0.03)	* * *	* * *
BCS-Sa	2520	2.47 (0.03)	2.51 (0.03)	2.56 (0.05)	***	2.66 (0.12)	2.54 (0.10)	2.46 (0.10)	2.40 (0.10)	***	2.51 (0.01)	2.52 (0.01)	NS	* * *
BCS-Sb	857	2.56 (0.12)	2.64 (0.15)	2.66 (0.13)	***	2.74 (0.21)	2.66 (0.21)	2.56 (0.20)	2.52 (0.21)	***	2.61 (0.04)	2.63 (0.04)	NS	***
BCS-W	2434	2.44 (0.04)	2.50 (0.04)	2.52 (0.06)	***	2.59 (0.09)	2.54 (0.08)	2.43 (0.08)	2.40 (0.08)	***	2.48 (0.01)	2.49 (0.01)	NS	***
BCS-D	1717	2.53 (0.05)	2.63 (0.06)	2.63 (0.09)	***	2.67 (0.07)	2.64 (0.07)	2.55 (0.07)	2.52 (0.07)	***	2.55 (0.06)	2.64 (0.06)	* * *	***

Table 1 Summary of least-squares means for BW and body condition score (BCS) variables (± standard error) at each physiological stage according to the parity, the litter size and the age of the ewe at first lambing

The significance probabilities for each fixed effect are provided. n = number of records; n (%) = proportion of ewes in each class of fixed effect; M = mating; Pa = early pregnancy; Pb = mid-pregnancy; L = lambing; Sa = beginning of suckling; Sb = end of suckling; W = weaning; D = dry-off. Sign. = significance probabilities. *P<0.05; ***P<0.001.

Intra-flock variability of sheep robustness



Figure 1 Cluster profiles for BW (on the left) and adjusted BW (on the right) in Romane ewes over their over the production cycles 1 (at the top), 2 (in the middle) and 3 (at the bottom). The scatter plots appearing on the left of each graph represents the individual coordinates on the two first principal components of the functional principal component analysis. The textbox on the right of each graph indicates the proportion of animals present in each cluster, and is given in percentage (%). The composition of each cluster at productive cycle *n* is given by indicating the proportion (%) of animals from clusters found at previous cycle n-1. M = mating; Pa = early pregnancy; Pb = mid-pregnancy; L = lambing; Sa = beginning of suckling; W = weaning; D = dry-off; Adjusted BW = BW taking into account average individual BW levels across cycles.

included 75% and 62% of the ewes (BC10 and BC12, respectively) and for cycle 3, ewes were distributed equally in the two clusters. Clusters BC1, BC2, BC4, BC5, BC6, BC7 and BC9 showed parallel within- and between-cycle profiles but different levels of BCS. The profiles for these clusters were characterized by an increase in BCS from BCS-M to BCS-Pa, a constant decrease from BCS-Pa to BCS-Sb and a progressive increase from BCS-Sb to the end of the production cycle. In cycle 3, cluster BC8 differed from clusters BC7 and BC9 by a very slight decrease of BCS between BCS-Pa and BC9-W and the highest BCS levels. Clusters for adjusted BCS showed very close ranges of variation within- and between-cycle and similar profiles to the main clusters found for BCS.

Ewes in cluster BC4 of cycle 2 originated mostly from clusters BC1 and BC2 (56% and 26%, respectively) of cycle 1. Ewes included in BC5 originated mainly from BC2 (53%) and

BC1 (36%) of cycle 1. Ewes in BC6 originated mostly from BC1 (64%) of cycle 1, and ewes in BC7 from BC4 (56%) and BC5 (28%) of the previous cycle. Ewes in clusters BC12 and BC13 originated mostly from BC10 (76% and 75%, respectively). Ewes in cluster BC8 originated mainly from BC4 (47%) and BC6 (33%) of cycle 2 and those in cluster BC9 mainly from BC5 (74%) of cycle 2. Ewes in clusters BC14 and BC15 originated mostly from BC12 (74% and 66%, respectively). In general, these ewes exhibited the same profiles at cycles 1, 2 and 3.

Figure 3 displays clusters observed considering ewes' BW and BCS over their first three productive cycles. Three clusters were found for each trait. Considering BW clusters, the three profiles showed similar and parallel profiles over the three cycles. The main cluster included 87% of ewes and indicated that majority of ewes were globally gaining weight until the



Figure 2 Cluster profiles for body condition score (BCS) (on the left) and adjusted BCS (on the right) in Romane ewes over their production cycles 1 (at the top), 2 (in the middle) and 3 (at the bottom). The scatter plot on the left of each graph represents the individual coordinates on the two first principal components of the functional principal component analysis. The textbox on the right of each graph indicates the proportion of animals in each cluster, which is given in percentage (%). The composition of each cluster at productive cycle *n* is given by indicating the proportion (%) of animals from clusters found at previous cycle n-1. BC = body condition; M = mating; Pa = early pregnancy; Pb = mid-pregnancy; L = lambing; Sa = beginning of suckling; Sb = end of suckling; W = weaning; D = dry-off; Adjusted BCS = BCS taking into account average individual BCS levels across cycles.

end of cycle 3 concomitantly with the observed alternation of weight gain and loss through each cycle. The two other clusters differed from the major one mainly at cycle 3 by higher or lower BW levels. Considering adjusted BW clusters, two of them (i.e. BW22 and BW24) showed similar and parallel profiles over the three cycles but with a crossing point between both trajectories during the second suckling period. The third cluster (i.e. BW23) differed from the two others in the alternation of weight gain and loss within- and between cycles. Considering BCS clusters, the main cluster (i.e. BC16, 76% of ewes) showed repeatable BCS gains and losses across the three first cycles while the two other clusters exhibited BCS gains and losses changing between cycles. Concerning adjusted BCS, the three clusters showed very close or overlapping trajectories and repeatable over the three cycles.

Figure 4 displays the BCS clusters observed considering only ewes lambing and suckling twins (no difference in the average age at first lambing between clusters within a cycle). Three clusters were identified for each production cycle. The two main profiles for each production cycle showed similar, parallel profiles, both within and between the cycles. The profiles were similar to those described for BCS clusters for the whole population. The two main clusters in cycles 1, 2 and 3 included 94.4%, 86.5% and 87.2% of ewes, respectively. There was 12% of the multiparous ewes that remained in clusters showing the highest BCS levels between cycles 2 and 3 (BC26 and BC29, respectively). Concerning the adjusted BCS for ewes lambing and suckling twins, only one trajectory was observed at cycles 1 and 3 and two trajectories at cycle 2 (data not shown), that presented the same profiles as the trajectories of the adjusted BCS (Figure 2).

Biological factors

The main biological factors were tested to explain the distribution of the ewes between the clusters at each cycle (Tables 2 and 3). The Litter effect was the most significant effect for BCS clusters in the three production cycles. Ewes with the smallest average litter size were mostly included in clusters BC3, BC6 and BC8 that showed the highest BCS levels (Table 2). Ewes with larger litter sizes were included in clusters BC2, BC5 and BC9 that showed the lowest BCS in cycles 1, 2 and 3. When considering the adjusted BCS, all the variation factors (i.e. litter size, litter at previous cycle, age at first lambing and year) had similar effects on the trait. Ewes with larger litter sizes were included in clusters BC11, BC12

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Figure 3 Cluster profiles for BW (at the top left), adjusted BW (at the top right), body condition score (BCS) (at the bottom left) and adjusted BCS (at the bottom right) over the entire productive life of Romane ewes. The scatter plot on the left of each graph represents the individual coordinates on the two first principal components of the functional principal component analysis. The textbox on the right of each graph indicates the proportion of animals in each cluster, which is given in percentage. BC = body condition; M = mating; L = lambing; W = weaning; Adjusted BW = BW taking into account average individual BW levels across cycles; Adjusted BCS = BCS taking into account average individual BCS levels across cycles.

and BC14 that showed lowest BCS levels. Concerning BW and adjusted BW, ewes with the smallest average litter size were included in clusters BW1 and BW12 for cycle 1, BW5, BW7 and BW16 for cycle 2 and BW10 and BW18 for cycle 3 (Table 3).

Litter size at previous cycle had significant effect on ewes' distribution in adjusted BCS clusters at cycle 2. Ewes that had a bigger litter size at cycle 1 were included in cluster BC12 showing lower BCS level. Considering BCS, litter size at previous cycle had very little effect at cycles 2 and 3 (Table 2). For adjusted BW, the ewes with the biggest litter size at previous cycle were included in clusters BW14 and BW17 showing progressive decrease in BW between mid-pregnancy and weaning. Considering BW, litter size at previous cycle had a significant and important effect. Ewes with bigger litter size at the previous cycle were included in clusters BW6 and BW9 at cycles 2 and 3, respectively, showing progressive decrease in BW between mid-pregnancy and weaning (Table 3).

The age of the ewe was the most significant effect for BW clusters at cycle 1, ranked second for BCS clusters at cycles 2 and 3 and was observed to be the less significant effect for BCS clusters at cycle 1 and BW clusters at cycles 2 and 3. For BCS at cycle 1, ewes that lambed at 1 year of age were included in clusters BC1 and BC2 and there were no differences in the age at first lambing for adjusted BCS at cycle 1.

For the two other production cycles, the age at first lambing did not discriminate the clusters for BCS whereas, for adjusted BCS at cycle 2, ewes that lambed at 1 year of age were mostly included in BC12 (Table 2). For BW and adjusted BW at cycle 1, ewes that lambed at 1 year of age were included in clusters BW1, BW4, BW12 and BW13. For cycle 2, ewes that lambed at 1 year of age were included in BW8, BW15 and BW16 and the age at first lambing was not discriminating at cycle 3 (Table 3).

Discussion

The dynamics of body reserves as a trait for robustness in the current context

The accepted concept of 'breeding for robustness' (Kitano, 2004; Klopcic *et al.*, 2009; Molotsi *et al.*, 2017) justifies the necessity for gathering new insights into such traits in order to reveal the degree of individual robustness and the underlying determinant mechanisms (De La Torre *et al.*, 2015; Friggens *et al.*, 2017). The BR dynamics have been shown to be a relevant trait for studying the capacity to overcome negative energy balance periods (Bauman and Currie, 1980; Nielsen *et al.*, 2003; González-García *et al.*, 2014). This trait is being considered as an interesting biological component in the design of future livestock systems (Dumont *et al.*, 2014; Phocas *et al.*, 2016).

BCS 3.5 Cluster % BC22 (red) 54.8 BC23 (blue) 39.6 BC24 (green) 5.6 3.0 2.5 2.0 3.5 Cluster % BC22¹ BC23 BC24 BC25 (blue) 65.8 11 4 0 BC26 (red) 20.7 5 6 0 BC27 (green) 13.5 20 0 3.0 2.5 2.0 3.5 Cluster % BC25 BC26 BC27 BC28 (blue) 59.0 30 4 9 12 3 BC29 (red) 28.2 15 BC30 (green)12.8 13 27 0 3.0 2.5 2.0 M Pa Ph L Sa Sb W D

Figure 4 Cluster profiles for body condition score (BCS) of Romane ewes lambing and suckling twins (Litter 3) over their production cycles 1 (at the top), 2 (in the middle) and 3 (at the bottom). The scatter plot on the left of each graph represents the individual coordinates on the two first principal components of the functional principal component analysis. The textbox on the right of each graph indicates the proportion of animals in each cluster, which is given in percentage (%). The composition of each cluster at productive cycle *n* is given by indicating the proportion (%) of animals from clusters found at previous cycle n - 1. BC = body condition; M = mating; Pa = early pregnancy; Pb = mid-pregnancy; L = lambing;Sa = beginning of suckling; Sb = end of suckling; W = weaning; D = dryoff.

Relevance of BW and body condition score for interpreting body reserves dynamics

The BW and BCS are considered to be typical and relatively easily monitored parameters (Brown *et al.*, 2015), and are usually analyzed to describe BR dynamics (Thorup *et al.*, 2012; Morel *et al.*, 2016; Puillet and Martin, 2017). Nevertheless, there is a consensus about the subjectivity of BCS due to disagreements about the methods used and inconsistent intra- and inter-operator assessment skills (Russel *et al.*, 1969; Edmonson *et al.*, 1989; Kenyon *et al.*, 2014). One advantage of this study is the consistent assessment by the same two operators over the whole study period, with operators regularly attending training sessions to homogenize their scoring technique.

The main originality of our study was that we characterized the variability of existing BR dynamics profiles and described sequential and temporal BR changes, using longitudinal data covering up to three production cycles in a flock of ewes reared in extensive conditions. To our knowledge, this is the first report of longitudinal BR dynamics in several production cycles using real data, at least in grazing sheep.

Overall body reserves mobilization and accretion profiles

Both BW and BCS vary as an ewe progresses through the different physiological stages of the production cycle. The BRs are mobilized from mid-pregnancy until the end of the suckling period and this is reflected by a decrease in the BCS. After weaning, an increase in the BCS is observed (i.e. BR accretion) and lasts until the beginning of the next pregnancy. The BW and BCS profiles observed in our study population and conditions are consistent with the previously described overall dynamics (González-García et al., 2014). The simultaneous increase in BW and decrease in BCS observed between early to mid-pregnancy are likely to be due to the start of the fetus growth and development and the related increasing demand in energy (Bauman and Currie, 1980). The BW probably continued to increase during the second half of pregnancy but a BW measurement before lambing was missing in the present study to show the expected BW increase (González-García et al., 2014). This biased the tendency observed for BW between midpregnancy and lambing (Figure 1; missing point between mid-pregnancy – Pb- and lambing – L-).

During the 1st month of suckling, the decrease in BCS was probably due to an imbalance between energy intake and expenditure caused by the peak of milk production, a high priority given to the new-born lambs and the incapacity of the female to take in enough feed to meet energy requirements (Nielsen *et al.*, 2003; Smith *et al.*, 2017). The decreases in BCS and BW were observed despite the use of fertilized paddocks with enough quality biomass available to ensure the nutritional balance during the suckling period (Molénat *et al.*, 2005). An additional BCS measurement during the 2nd month of suckling could perhaps help to identify the exact endpoint of BR mobilization during this period.

The BW and BCS gains started again before the end of suckling. The stabilization or increase in BCS at this point could be related to a decrease in the ewes' energy requirements and dietary diversification in lambs. This is consistent with the findings of another study (Kharrat and Bocquier, 2010) that reported that BR replenishment was prioritized at the end of suckling in goats. Accretion continued during the autumn when ewes grazed the new herbage regrowth available in the native paddocks. It lasted during the dry-off period and until the beginning of the next pregnancy, including feed supplementation at mid-pregnancy (González-García *et al.*, 2014).

In addition to the physiological stages of the production cycle, various biological factors such as parity, age at first

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		п	Litter s	ize	Age at first lambing		Litter size at previous cycle	
Cycle	Cluster number		LSMeans	Contr.	LSMeans	Contr.	LSMeans	Contr.
1	BC1	703	2.03 (0.88)	* *	1.52 (0.50)	~	ND	ND
	BC2	434	2.31 (0.95)		1.47 (0.50)		ND	
	BC3	9	2.00 (1.00)		2.00 (0.00)		ND	
2	BC4	593	2.76 (1.06)	*	1.44 (0.50)	*	1.81 (1.06)	*
	BC5	329	3.03 (0.87)		1.36 (0.48)		1.98 (1.09)	
	BC6	146	2.02 (1.07)		1.55 (0.50)		1.76 (1.14)	
3	BC7	319	3.15 (1.00)	* * *	1.49 (0.50)	~	2.80 (1.11)	* *
	BC8	49	2.04 (1.04)		1.43 (0.50)		2.33 (1.26)	
	BC9	46	3.46 (0.75)		1.39 (0.49)		2.83 (1.14)	
1	BC10	1041	2.11 (0.91)	*	1.50 (0.50)	*	ND	ND
	BC11	105	2.31 (0.96)		1.57 (0.50)		ND	
2	BC12	824	2.78 (1.02)	*	1.33 (0.47)	* * *	1.77 (1.03)	**
	BC13	248	2.63 (1.14)		1.79 (0.41)		2.12 (1.20)	
3	BC14	235	3.26 (0.95)	**	1.43 (0.50)	*	2.77 (1.11)	* * *
	BC15	186	2.81 (1.13)		1.54 (0.50)		2.72 (1.18)	
1	BC22	113	2.00 (0.00)	ND	1.96 (0.21)	~	ND	ND
	BC23	83	2.00 (0.00)		1.98 (0.15)		ND	
	BC24	10	2.00 (0.00)		2.00 (0.00)		ND	
2	BC25	248	2.00 (0.00)	ND	1.42 (0.49)	~	1.86 (1.04)	* *
	BC26	78	2.00 (0.00)		1.45 (0.50)		1.75 (1.04)	
	BC27	51	2.00 (0.00)		1.39 (0.49)		2.00 (1.12)	
3	BC28	69	2.00 (0.00)	ND	1.36 (0.48)	*	2.68 (1.04)	* *
	BC29	33	2.00 (0.00)		1.48 (0.51)		2.76 (1.06)	
	BC30	15	2.00 (0.00)		1.80 (0.41)		2.43 (1.16)	

 Table 2 Least-square (LS) means (± standard error) for litter size, age at first lambing and litter size at previous cycle for ewes present in each cluster of body condition score (BCS) at each production cycle

n = number of ewes; ND = non-determined; Contr. = contribution of the effect to the model based on Akaike information criteria and Bayesian information criteria: ~, *, *** indicate no, low, medium and high significant contribution of the effect to the model; BC1 to BC9 = BCS clusters; BC10 to BC15 = adjusted BCS clusters (i.e. BCS taking into account average individual BCS levels across cycles); BC22 to BC30 = BCS clusters for the subgroup of ewes lambing and suckling twins.

parity, litter size or litter size at previous cycle had a significant impact on the BW and BCS levels of the females included in this study, as previously reported (María and Ascaso, 1999; González-García *et al.*, 2014 and 2015). A strong effect of litter size on BW loss was observed with the biggest litter sizes, since litter sizes were highly correlated between cycles, the BW loss may be related to two consecutive big litter sizes. The year/season also affected BR levels in our grazing system because the nutritional and energy balance are dependent on the weather and amount of biomass available.

Diversity of body reserves and body condition score profiles (cluster analysis)

In this study, we confirmed our hypothesis regarding the existence of different BW and BCS trajectories within a flock of Romane sheep reared in extensive conditions. Indeed, the characterization of variability in BCS profiles highlighted three separate trajectories for each production cycle. Most trajectories exhibited similar shapes (i.e. similar sequences of BCS gains and losses) but differed in terms of BW and BCS levels. This is confirmed by the two close trajectories observed with BCS taking into account average individual BCS levels across cycles (i.e. adjusted BCS). Indeed, trajectories found for adjusted BCS did

not differ in the levels within cycle but slightly in the shape. Consideration of the average individual level across cycles did not highlight strong differences of trajectory. Some trajectories, essentially those showing the highest BCS levels, displayed different dynamics in addition to different BCS levels. The latter trajectories represented generally clusters including a lower number of animals. In addition, all BCS and adjusted BCS trajectories, whatever the production cycle, also suggested that BR accretion occurred from weaning to the next early pregnancy as described for overall BR profiles. The BR accretion during the dry-off period was associated with a stabilization or slight increase in BW depending on the various trajectories and production cycle. Consistently with the increase of BCS observed during early pregnancy, all BW and adjusted BW trajectories showed an increase in BW at this phase and later during pregnancy as hypothesized for the overall profile. While some BW trajectories differed only in the levels and not in the shape, these trajectories were not found for BW taking into account average individual BW level across cycles (i.e. adjusted BW). Thus, differences in the shape of adjusted BW profiles suggested that various BW trajectories existed within flock. In addition, BCS trajectories showing similar shape exhibited higher amplitude of BCS variation in primiparous ewes than in multiparous ewes. This suggest that improved efficiency in the

Table 3 Least-square (LS) means (± standard error) for litter size, age at first lambing and litter size at previous cycle for ewes present in each cluster of BW at each production cycle

			Litter s	ize	Age at first lambing		Litter size at previous cycle	
Cycle	Cluster number	п	LSMeans	Contr.	LSMeans	Contr.	LSMeans	Contr.
1	BW1	569	1.72 (0.59)	*	1.10 (0.30)	***	ND	ND
	BW2	429	2.51 (0.99)		1.93 (0.26)		ND	
	BW3	116	2.75 (1.07)		2.00 (0.00)		ND	
	BW4	32	2.16 (0.81)		1.28 (0.46)		ND	
2	BW5	830	2.67 (1.04)	* *	1.37 (0.48)	*	1.84 (1.02)	* * *
	BW6	143	3.30 (0.88)		1.90 (0.31)		2.14 (1.30)	
	BW7	86	2.51 (1.18)		1.31 (0.47)		1.55 (1.26)	
	BW8	9	3.00 (1.00)		1.00 (0.00)		1.22 (0.67)	
3	BW9	404	3.07 (1.05)	* *	1.48 (0.50)	*	2.76 (1.13)	* *
	BW10	10	2.40 (1.17)		1.30 (0.48)		2.40 (1.51)	
1	BW11	546	2.55 (0.99)	*	1.92 (0.28)	***	ND	ND
	BW12	477	1.71 (0.63)		1.14 (0.35)		ND	
	BW13	123	1.86 (0.66)		1.07 (0.26)		ND	
2	BW14	430	3.07 (0.93)	*	1.79 (0.41)	* *	2.07 (1.27)	* * *
	BW15	408	2.61 (1.06)		1.15 (0.36)		1.60 (0.90)	
	BW16	234	2.39 (1.08)		1.26 (0.45)		1.90 (0.91)	
3	BW17	414	3.07 (1.05)	*	1.48 (0.50)	* * *	2.76 (1.13)	**
	BW18	7	2.29 (1.11)		1.43 (0.53)		2.14 (1.46)	

n = number of ewes; LSMeans = Least-square means; ND = non-determined; Contr. = contribution of the effect to the model based on Akaike information criteria and Bayesian information criteria: *, **, ***indicate no, low, medium and high significant contribution of the effect to the model; BW1 to BW10 = BW clusters; BW11 to BW18 = adjusted BW clusters (i.e. BW taking into account average individual BW levels across cycles).

use of BR could occur with time as the animal gains in 'biological experience'.

The distribution of individual ewes between BCS and BW trajectories from one cycle to another was partially maintained. Between one-third and half of the ewes included in the biggest clusters remained in the same trajectory at the next cycle or even during the three cycles for BW, adjusted BW, BCS and adjusted BCS. Changes in the distribution of ewes between BCS trajectories in successive cycles are also probably related to changes in litter sizes. Indeed, the trajectory showing the highest BCS for each cycle included ewes with the smallest litter size, and conversely the trajectories with the low BCS levels included ewes with the largest average litter sizes. Nevertheless, ewes with similar litter size were also distributed in clusters showing different trajectories. This suggests that litter size, even if it was one of the most significant effect, was not the only factor explaining the distribution of ewes' trajectories in this study. In addition, individual variability in BCS management may contribute to how ewes are distributed between the trajectories. Indeed, considering cluster analysis for BCS specifically in ewes lambing and suckling twins, three well-defined trajectories were also described for primiparous or multiparous ewes. Those trajectories differed by the level of BCS and/or the shape while clusters did not differ in litter size and age at first lambing. This was in accordance with the trajectories of ewes lambing and suckling twins obtained with the adjusted BCS (data not shown), which differed mainly in the shape. These results support the hypothesis of individual variability that contributes to BR dynamics whatever the litter size, parity or

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age at first lambing. Recent genetic studies considering BW and BCS changes over time supported the hypothesis of genetic determinism for weight and fat changes across production cycle (Macé *et al.*, 2018b; Walkom *et al.*, 2014).

Depending on trajectories and cycle, ewes recovered or increased their initial BW (i.e. at the beginning of cycle) at the end of the cycle. The variations in BW observed during the cycle resulted in few overall changes in BW between the beginning and the end of the cycle. Nevertheless, two BW trajectories and adjusted BW trajectories for production cycle 1 showed overall higher BW gain and represented 53% of the primiparous ewes, either considering BW or adjusted BW. The females included in these trajectories were the youngest ewes, so we hypothesized that these females were still growing, as previously reported by González-García and Hazard (2016). Interestingly, when cluster analysis was performed for the ewes' entire productive life, the BW trajectory including most of the ewes (i.e. 87%) suggested that females kept growing during their three first cycles. Trajectories were even better defined when taking into account average BW level across cycles since the ewes in the three adjusted BW trajectories kept gaining weight across the first cycles, despite alternating periods of BW gain and loss. Moreover, the youngest ewes at first lambing were ewes that gained the higher amount of BW across the three cycles. This is in accordance with the study of Zygoyiannis *et al.* (1997) reporting growth in Greek ewes until they reached 3.5 years of age. It would be interesting to analyze BW trajectories in older ewes to determine at which age Romane ewes actually stop growing in this environment.

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Our 'ideal body reserves profile' definition

When analyzing the diverse BW and BCS profiles found in our study, it seemed important to define what we deemed the 'ideal profile' of a robust ewe to be. Regarding BR dynamics efficiency, a robust ewe would be an individual that avoids guick and drastic losses in BRs when facing a negative energy balance challenge and reacts by guickly replenishing BRs to recover at least its initial status once the shortage ended without affecting its performance (Bocquier and González-García, 2010; Martin and Sauvant, 2010). Interestingly, cluster analysis for BCS in ewes with similar litter sizes still showed three clusters, as described in the whole population (Figure 4; Supplementary Figure S1). We hypothesized that the 'robust trajectory' was the trajectory that showed intermediate BCS levels in primiparous ewes (red profile; BC22) and the highest BCS in multiparous ewes lambing and suckling twins (profiles BC26 and BC29; Figure 3). Ewes belonging to the robust trajectory for all the three production cycles would be the most interesting ewes. For now, 5% to 12% of the ewes followed the robust trajectory for two cycles but none of them followed this trajectory for the three cycles.

To better characterize robust individuals, new parameters should be considered in addition to those evaluated here (BW and BCS). On the one hand, the longitudinal analysis of a set of plasma metabolites and hormones could provide additional, specific data on each individual's energy metabolism. On the other hand, other zootechnical parameters related to the ewes' reproductive and productive performances will be needed to define individual robustness in a multicriteria assessment approach. Everything as a whole would add to the quality of the 'big-picture' for a relevant interpretation of such a complex and multifactorial trait.

Conclusions

In this study, the analysis of the measurements recorded in the INRA historical database over several production cycles confirmed that BW and BCS levels in Romane ewes were, as expected, significantly affected by the age at first lambing, parity, physiological stage, litter size and litter size at the previous cycle, as well as by the year of measurement. Our most relevant findings are the identification of different profiles or trajectories for BCS and BW suggesting different dynamics in BR mobilization-accretion. For a proportion of the ewes, the profiles remained consistent throughout the animal's life, whatever the age at first lambing, parity and litter size (i.e. repeatability of BR dynamics between successive cycles). Thus, the individual differences in profiles observed at a given stage illustrate the existence of proven intra-flock variability in the adaptive capacities of ewes to manage their BR. Such variability could probably be explained by a genetic component, which justifies further efforts to continue to explore the main genetic and physiological mechanisms behind this phenomenon.

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Declaration of interest

The authors fully declare, under their responsibility, that there are no conflicts of interest to declare.

Ethics statement

The care and management of all animals whose BW and BCS measurements were included in this study was approved by the INRA' Institutional and Regional Animal Care and Use Ethical Committee.

Software and data repository resources

None.

Supplementary materials

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References

Akaike H 1974. A new look at the statistical model identification. IEEE Transactions on Automatic Control 19, 716–723.

Álvarez-Rodríguez J, Estopañan G, Sanz A, Dervishi E, Govoni N, Tamanini C and Joy M 2012. Carry-over effects of body condition in the early pregnant ewe on peri-partum adipose tissue metabolism. Journal of Animal Physiology and Animal Nutrition 96, 985–992.

Bauman DE and Currie WB 1980. Partitioning of nutrients during pregnancy and lactation: a review of mechanisms involving homeostasis and homeorhesis. Journal of Dairy Science 63, 1514–1529.

Bocquier F and González-García E 2010. Sustainability of ruminant agriculture in the new context: feeding strategies and features of animal adaptability into the necessary holistic approach. Animal 4, 1258–1273.

Brown DJ, Savage DB, Hinch GN and Hatcher S 2015. Monitoring liveweight in sheep is a valuable management strategy: a review of available technologies. Animal Production Science 55, 427–436.

Dai X, Hadjipantelis PZ, Ji H, Mueller HG and Wang JL 2017. fdapace: functional data analysis and empirical dynamics. R package version 0.3. 0: CRAN. Retrieved on 9 January 2017 from https://github.com/functionaldata/tPACE/

De La Torre A, Recoules E, Blanc F, Ortigues-Marty I, D'Hour P and Agabriel J 2015. Changes in calculated residual energy in variable nutritional environments: an indirect approach to apprehend suckling beef cows' robustness. Livestock Science 176, 75–84.

Dumont B, González-García E, Thomas M, Fortun-Lamothe L, Ducrot C, Dourmad JY and Tichit M 2014. Forty research issues for the redesign of animal production systems in the 21st century. Animal 8, 1382–1393.

Edmonson AJ, Lean IJ, Weaver LD, Farver T and Webster G 1989. A body condition scoring chart for Holstein dairy cows. Journal of Dairy Science 72, 68–78.

Friggens NC, Blanc F, Berry DP and Puillet L 2017. Review: deciphering animal robustness. A synthesis to facilitate its use in livestock breeding and management. Animal 11, 2237–2251.

González-García E, Gozzo de Figuereido V, Foulquie D, Jousserand E, Autran P, Camous S, Tesniere A, Bocquier F and Jouven M 2014. Circannual body reserve dynamics and metabolic profile changes in Romane ewes grazing on rangelands. Domestic Animal Endocrinology 46, 37–48.

González-García E and Hazard D 2016. Growth rates of Romane ewe lambs and correlated effects of being mated as hoggets or two-tooth ewes on first offspring performance. Livestock Science 189, 63–69.

González-García E, Tesniere A, Camous S, Bocquier F, Barillet F and Hassoun P 2015. The effects of parity, litter size, physiological state, and milking frequency

on the metabolic profile of Lacaune dairy ewes. Domestic Animal Endocrinology 50, 32–44.

Kenyon PR, Maloney SK and Blache D 2014. Review of sheep body condition score in relation to production characteristics. New Zealand Journal of Agricultural Research 57, 38–64.

Kharrat M and Bocquier F 2010. Impact of indoor feeding at late lactation stage on body reserves recovery and reproductive performances of Baladi dairy goats fed on pastoral system. Small Ruminant Research 90, 127–134.

Kitano H 2004. Biological robustness. Nature Reviews Genetics 5, 826-837.

Klopcic M, Reents R, Philipsson J and Kuipers A 2009. Breeding for robustness in cattle. Wageningen Academic Publishers, Wageningen, The Netherlands.

Langrognet F, Lebret R and Poli C 2016. Rmixmod: supervised, unsupervised, semi-supervised classification with MIXture MODelling (Interface of MIXMOD Software). R package version 2.1.1. Retrieved on 9 January 2017 from https:// cran.r-project.org/web/packages/Rmixmod/index.html

Lebret R, lovleff S, Langrognet F, Biernacki C, Celeux G and Govaert G 2014. Rmixmod: the R package of the model-based unsupervised, supervised and semi-supervised classification mixmod library. Journal of Statistical Software 67, 241–270.

Macé T, González-García E, Pradel J, Parisot S, Carrière F, Douls S, Foulquié D and Hazard D 2018b. Genetic analysis of robustness in meat sheep through body weight and body condition score changes over time. Journal of Animal Science 96, 450–4511.

Macé T, Hazard D, Carrière F, Douls S, Foulquié D, Robert-Granié C. and González-García E 2018a. Body weight and body condition score variations in Romane ewes: intraflock variability in their dynamics and magnitude over multiple production cycles. Journal of Dairy Science 101 (suppl. 2), 236.

María GA and Ascaso MS 1999. Litter size, lambing interval and lamb mortality of Salz, Rasa Aragonesa, Romanov and F1 ewes on accelerated lambing management. Small Ruminant Research 32, 167–172.

Martin O and Sauvant D 2010. A teleonomic model describing performance (body, milk and intake) during growth and over repeated reproductive cycles throughout the lifespan of dairy cattle. 1. Trajectories of life function priorities and genetic scaling. Animal 4, 2030–2047.

Mendizabal JA, Delfa R, Arana A and Purroy A 2011. Body condition score and fat mobilization as management tools for goats on native pastures. Small Ruminant Research 98, 121–127.

Molénat G, Foulquie D, Autran P, Bouix J, Hubert D, Jacquin M and Bibe B 2005. Pour un élevage ovin allaitant performant et durable sur parcours : un système expérimental sur le Causse du Larzac. INRA Productions Animales 18, 323–338.

Molina A, Gallego L, Torres A and Vergara H 1994. Effect of mating season and level of body reserves on fertility and prolificacy of Manchega ewes. Small Ruminant Research 14, 209–217.

Molotsi A, Dube B, Oosting S, Marandure T, Mapiye C, Cloete S and Dzama K 2017. Genetic traits of relevance to sustainability of smallholder sheep farming systems in South Africa. Sustainability (Switzerland) 9, 1–18.

Morel PCH, Schreurs NM, Corner-Thomas RA, Greer AW, Jenkinson CMC, Ridler AL and Kenyon PR 2016. Live weight and body composition associated with an increase in body condition score of mature ewes and the relationship to dietary energy requirements. Small Ruminant Research 143, 8–14.

Nielsen HM, Friggens NC, Løvendahl P, Jensen J and Ingvartsen KL 2003. Influence of breed, parity, and stage of lactation on lactational performance and relationship between body fatness and live weight. Livestock Production Science 79, 119–133.

Phocas F, Belloc C, Bidanel J, Delaby L, Dourmad JY, Dumont B, Ezanno P, Fortun-Lamothe L, Foucras G, Frappat B, González-García E, Hazard D, Larzul C, Lubac S, Mignon-Grasteau S, Moreno CR, Tixier-Boichard M and Brochard M 2016. Towards the agroecological management of ruminants, pigs and poultry through the development of sustainable breeding programmes: I-selection goals and criteria. Animal 10, 1749–1759.

Puillet L and Martin O 2017. A dynamic model as a tool to describe the variability of lifetime body weight trajectories in livestock females. Journal of Animal Science 95, 4846–4856.

Ricordeau G, Tchamitchian L, Brunel JC, Nguyen TC and François D 1992. La race ovine INRA 401: un exemple de souche synthétique. INRA Productions Animales hs (hs), 255–262.

Rojas-Downing MM, Nejadhashemi AP, Harrigan T and Woznicki SA 2017. Climate change and livestock: impacts, adaptation, and mitigation. Climate Risk Management 16, 145–163.

Russel AJF, Doney JM and Gunn RG 1969. Subjective assessment of body fat in live sheep. Journal of Agricultural Sciences 72, 451–454.

Schwarz G 1978. Estimating the dimension of a model. The Annals of Statistics 6, 461–464.

Smith GL, Friggens NC, Ashworth CJ and Chagunda MGG 2017. Association between body energy content in the dry period and post-calving production disease status in dairy cattle. Animal 11, 1590–1598.

Thorup VM, Edwards D and Friggens NC 2012. On-farm estimation of energy balance in dairy cows using only frequent body weight measurements and body condition score. Journal of Dairy Science 95, 1784–1793.

Walkom SF, Brien FD, Hebart ML, Fogarty NM, Hatcher S and Pitchford WS 2014. Season and reproductive status rather than genetics factors influence change in ewe weight and fat over time. 2. Spline analysis of crossbred ewes. Animal Production Science 54, 814–820.

Yao F, Mueller HG and Wang JL 2005. Functional linear regression analysis for longitudinal data. Annals of Statistics 33, 2873–2903. (Dense data).

Zygoyiannis D, Stamataris C, Friggens NC, Doney JM and Emmans GC 1997. Estimation of the mature weight of three breeds of Greek sheep using condition scoring corrected for the effect of age. Animal Science 64, 147–153.