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Predicting physiological imbalance in Holstein dairy cows by three different sets of milk biomarkers

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Abstract

Blood biomarkers may be used to detect physiological imbalance and potential disease. However, blood sampling is difficult and expensive, and not applicable in commercial settings. Instead, individual milk samples are readily available at low cost, can be sampled easily and analysed instantly. The present study sampled blood and milk from 234 Holstein dairy cows from six experimental herds in different European countries. The objective was to compare the use of three different sets of milk biomarkers for identification of cows in physiological imbalance and thus at risk of developing a metabolic or infectious disease. Random forests was used to predict body energy balance (EBAL), index for physiological imbalance (PI-index) and three clusters differentiating the metabolic status of cows created on basis of concentrations of plasma glucose, plasma β -hydroxybutyrate (BHB), plasma non-esterified fatty acids (NEFA) and serum IGF-1. These three metabolic clusters were interpreted as cows in balance, cows in physiological imbalance and "intermediate cows" with a physiological status in between. The three sets of milk biomarkers used for prediction were: milk Fourier transform mid-IR (FT-MIR) spectra, 19 immunoglobulin G (IgG) N-glycans and 8 milk metabolites and enzymes (MME). Blood biomarkers were sampled twice; around 14 days after calving (days in milk (DIM)) and around 35 DIM. MME and FT-MIR were sampled twice weekly 1-50 DIM whereas IgG N-glycan were measured only four times. Performances of random forests predictions for EBAL and PI-index were measured by the coefficient of determination (R2cv) and the root mean squared error (RMSEcv) from leave-one-cow-out (internal) cross-validation (CV). For metabolic clusters, performance was measured by sensitivity, specificity and global accuracy from this crossvalidation. Neither EBAL nor PI-index were sufficiently precise to be used as a management tool for identification of risk cows. The best prediction of PI-index was obtained by MME (R2CV = 0.40 at 14 DIM and 0.35 at 35 DIM) while FT-MIR showed a better performance than MME for prediction of EBAL (R2CV = 0.28 vs 0.21). Global accuracies of predicting metabolic clusters from MME and FT-MIR were at the same level and ranged from 0.54 to 0.65 for MME and 0.51 to 0.68 for FT-MIR. R2CV and accuracies were lower for IgG N-glycans. In conclusion, MME and FT-MIR can be used to predict the physiological status of the cows, while the use of IgG N-glycans for prediction still needs development.

Keywords	Metabolites; enzymes; FT-MIR; IgG N-glycans; metabolic clusters; random forests
Taxonomy	Animal Lactation, Dairy Cattle, Animal Energetics, Animal Metabolism
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To the Editor

Preventive Veterinary Medicine

Currently, the prediction at a large scale of physiological status of cows is of great interest in order to perform genetic studies and for the management of cows. The use of milk biomarkers seems a good strategy as it is easily accessible and already routinely collected. Enclosed please find our manuscript entitled "Predicting physiological imbalance in Holstein dairy cows by three different sets of milk biomarkers" authored by Foldager et al. This manuscript was developed in the frame of the GplusE project granted by the European Union, which sampled blood and milk from 234 Holstein dairy cows from six experimental herds in different European countries. The objective was to compare the use of three different sets of milk biomarkers for identification of cows in physiological imbalance and thus at risk of developing a metabolic or infectious disease. Milk biomarkers used are metabolites and enzymes, Fourier transform mid-infrared (FT-MIR) spectra and immunoglobulin G (IgG) N-glycans. Based on the same data, two other papers from the GplusE project (De Koster et al., 2019; Grelet et al., 2019) have considered the prediction of metabolic status (balanced/unbalanced) using metabolic clusters based on k-means clustering of four blood biomarkers; glucose, non-esterified fatty acids (NEFA) and β -hydroxybutyrate (BHB) in plasma and insulin-like growth factor-1 (IGF-1) in serum. A third paper from the GplusE project (Krogh et al., accepted 26 Sep 2019) focused on herd variation in the biomarkers. The present paper brings new knowledge by comparing random forest predictions of body energy balance (EBAL), index for physiological imbalance (PI-index) and the metabolic clusters just described. The paper goes deeper in the evaluation of the potential of milk metabolites and enzymes but also investigate the potential of IgG N-glycans as biomarker and contributes to the understanding of the clustering approach. The main objective was to compare the use of milk metabolites and enzymes, FT-MIR spectra and IgG N-glycans for identification of cows in physiological imbalance and thus at risk of developing a metabolic or infectious disease.

We hope you will consider this paper for publication in Preventive Veterinary Medicine.

Yours sincerely,

Leslie Foldager, PhD, MSc Senior Researcher Department of Animal Science Aarhus University, Tjele, Denmark

1 Highlights

- Identifying physiological imbalance/disease risk in dairy cows for herd
 management
- Blood biomarkers are relevant indicators but not generally applicable
- 5 commercially
- 6 Milk biomarkers can be taken automatically as in Herd Navigator™
- FT-MIR spectra and milk metabolites and enzymes appeared equally good as
 biomarkers
- IgG *N*-glycans suffered from fewer samples and completeness and needs

10 development

1	Predicting physiological imbalance in Holstein dairy cows by three different
2	sets of milk biomarkers
3	
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- 29
- 30

31 Abstract

Blood biomarkers may be used to detect physiological imbalance and potential 32 disease. However, blood sampling is difficult and expensive, and not applicable in 33 commercial settings. Instead, individual milk samples are readily available at low 34 cost, can be sampled easily and analysed instantly. The present study sampled 35 blood and milk from 234 Holstein dairy cows from six experimental herds in different 36 European countries. The objective was to compare the use of three different sets of 37 milk biomarkers for identification of cows in physiological imbalance and thus at risk 38 of developing a metabolic or infectious disease. Random forests was used to predict 39 body energy balance (EBAL), index for physiological imbalance (PI-index) and three 40 clusters differentiating the metabolic status of cows created on basis of 41 concentrations of plasma glucose, plasma β -hydroxybutyrate (BHB), plasma non-42 43 esterified fatty acids (NEFA) and serum IGF-1. These three metabolic clusters were interpreted as cows in balance, cows in physiological imbalance and "intermediate 44 cows" with a physiological status in between. The three sets of milk biomarkers used 45 for prediction were: milk Fourier transform mid-IR (FT-MIR) spectra, 19 46 immunoglobulin G (IgG) N-glycans and 8 milk metabolites and enzymes (MME). 47 Blood biomarkers were sampled twice; around 14 days after calving (days in milk 48 (DIM)) and around 35 DIM. MME and FT-MIR were sampled twice weekly 1-50 DIM 49 whereas IgG *N*-glycan were measured only four times. Performances of random 50

forests predictions for EBAL and PI-index were measured by the coefficient of 51 determination (R²_{cv}) and the root mean squared error (RMSE_{cv}) from leave-one-cow-52 out (internal) cross-validation (CV). For metabolic clusters, performance was 53 measured by sensitivity, specificity and global accuracy from this cross-validation. 54 Neither EBAL nor PI-index were sufficiently precise to be used as a management tool 55 for identification of risk cows. The best prediction of PI-index was obtained by MME 56 $(R^{2}_{CV} = 0.40 \text{ at } 14 \text{ DIM and } 0.35 \text{ at } 35 \text{ DIM})$ while FT-MIR showed a better 57 performance than MME for prediction of EBAL (R^{2}_{CV} = 0.28 vs 0.21). Global 58 accuracies of predicting metabolic clusters from MME and FT-MIR were at the same 59 level and ranged from 0.54 to 0.65 for MME and 0.51 to 0.68 for FT-MIR. R²_{CV} and 60 accuracies were lower for IgG N-glycans. In conclusion, MME and FT-MIR can be 61 used to predict the physiological status of the cows, while the use of IgG N-glycans 62 for prediction still needs development. 63

64

65 **Abbreviations**

66 BHB, β-hydroxybutyrate; CV, cross-validation; DIM, days in milk; EBAL, body energy

⁶⁷ balance; FT-MIR, Fourier transform mid-IR; IgG, immunoglobulin G; LDH,

68 dehydrogenase; MME, metabolites and enzymes; NAGase, *N*-acetyl-β-D-

69 glucosaminidase; NEFA, non-esterified fatty acids; PI-index, index for physiological

⁷⁰ imbalance; R², coefficient of determination; RMSE, root mean squared error; VIM,

71 variable importance measures

72

73 Keywords

Metabolites; enzymes; FT-MIR; IgG *N*-glycans; metabolic clusters; random forests
 75

76 Introduction

Diseases at calving and during early lactation account for the majority of health and 77 welfare problems in dairy production (Ingvartsen et al., 2003). These include 78 production diseases such as fatty liver, ketosis, rumen acidosis and lameness. Most 79 of such diseases in periparturient cows are argued to be the result of physiological 80 imbalance (Ingvartsen, 2006). Correspondingly, infectious diseases such as mastitis 81 and metritis are included as the immune system is strongly interlinked with 82 physiological imbalance via the endocrine system and metabolites that must 83 accommodate to the demands for lactation facing the transition cow (Ingvartsen and 84 Moyes, 2015). The consequences of subclinical and clinical diseases are suboptimal 85 animal welfare and production and lower reproductive efficiency. Thus, physiological 86 imbalance leading to these subclinical and clinical diseases should have high priority 87 of being addressed with regard to development of management tools. 88

89

Cows in physiological imbalance have increased risk of developing diseases and 90 reduced production (Ingvartsen et al., 2003; Bjerre-Harpoth et al., 2012). Subclinical 91 92 stages of diseases can be detected by biomarkers while the cow may appear 93 completely healthy. A number of biomarkers in blood are well described but are currently less well characterized in milk. In the review of Ingvartsen (2006), it is 94 documented that plasma concentrations of glucose, non-esterified fatty acids (NEFA) 95 and β -hydroxybutyrate (BHB) are relevant indicators to determine subclinical ketosis. 96 LeBlanc et al. (2005) also identified blood NEFA and BHB as relevant indicators of 97 displaced abomasum in dairy cows. Piechotta et al. (2012) reported that 98 concentrations of serum NEFA and plasma IGF-1 prepartum are associated with 99 postpartum diseases, while IGF-1 postpartum was the best predictor of both left 100

displaced abomasum and risk of culling (Lyons et al., 2014). However, collecting and
analysing blood samples for measuring biomarkers is difficult and expensive, and not
applicable in commercial settings. Instead, individual milk samples are readily
available and milking systems even provide automatic sampling and measurement of
e.g. milk conductivity. Such automatic systems can be expanded to measure e.g.
milk BHB (e.g. Herd Navigator[™], <u>http://www.herdnavigator.com</u>).

107

Enjalbert et al. (2001) showed that subclinical ketosis can be identified by measuring 108 BHB in milk with enzymatic analysis or with Ketolac test strips. Other studies also 109 reported milk BHB to be a relevant indicator of subclinical and clinical ketosis (e.g. 110 Nielsen et al., 2005). Free glucose, glucose-6-phosphate (Larsen and Moyes, 2015), 111 and isocitrate (Larsen, 2014) reflect the nutrient availability and metabolic turnover in 112 the mammary gland that are linked to the blood levels and therefore potentially 113 indicators of physiological imbalance and risk of disease. Larsen et al. (2010) and 114 Kitchen et al. (1978), respectively, reported that the milk enzymes lactate 115 dehydrogenase (LDH) and N-acetyl-β-D-glucosaminidase (NAGase) performed 116 equally with somatic cell count and acute phase proteins as inflammatory indicators 117 118 of mastitis. In addition, Fourier transform mid-IR (FT-MIR) spectra of milk can be calibrated to estimate e.g. milk metabolites, and measures of milk immunoglobulin G 119 (IgG) *N*-glycans may be potential new biomarkers. 120

121

Based on the same data as here, two other papers (De Koster et al., 2019; Grelet et al., 2019) have considered the prediction of metabolic status (balanced/unbalanced) using metabolic clusters based on k-means clustering of four blood biomarkers; glucose, NEFA and BHB in plasma and IGF-1 in serum. The present paper

supplements these papers by comparing random forests predictions from three
different sets of milk biomarkers; metabolites and enzymes (MME), FT-MIR spectra
and IgG *N*-glycans. In addition to metabolic clusters, predictions of body energy
balance (EBAL) and index for physiological imbalance (PI-index) (Ingvartsen, 2006;
Moyes et al., 2013a, 2013b) were considered. Grelet et al. (2019) used a different
prediction method and only considered FT-MIR, De Koster et al. (2019) only used
multiparous cows and both studies only considered prediction of clusters.

133

The present paper focuses more on MME but also investigate the potential of IgG *N*glycans as a set of milk biomarkers and contributes to the understanding of the clustering approach. The main objective was to compare the use of MME, FT-MIR and IgG *N*-glycans for identification of cows in physiological imbalance and thus at risk of developing a metabolic or infectious disease.

139

140 Material and methods

Study design, sampling and analysis of milk as well as blood have been described in 141 De Koster et al. (2019), Grelet et al. (2019) and Krogh et al. (2019). In brief, six 142 143 experiments were conducted in Northern Ireland (UK), Denmark (DK), Belgium (BE), Italy (IT), Germany (DE) and Ireland (IE). These included a total of 234 Holstein dairy 144 cows (55 first parity, 66 second parity, and 113 in third or higher parity (3+), see 145 Supplementary Table S1). In four experiments, all cows were fed a standard diet 146 typical for the particular country. In the UK and DK experiments, a standard diet and 147 two different experimental diets were used. An overview of the diets is shown in table 148 1 of Krogh et al. (2019). 149

150

151 Derived measures

The calculation of EBAL was described in De Koster et al. (2019) and Krogh et al. (2019). EBAL was only calculated if both morning and evening yield was available for that day. Afterwards, three days (i.e. +/- 1 days in milk (DIM)) moving averages of EBAL were calculated and used for the analyses. The average live body weights within calendar week was used to smooth large day-to-day variation and measurement errors of scales. Summary statistics of EBAL are shown in supplementary tables of Krogh et al. (2019).

PI-index was calculated as [log₁₀(NEFA)] + [log₁₀(BHB)] - [glucose] (Moyes et al., 160 2013a), where plasma concentrations of the individual metabolites were standardised 161 to an overall mean of zero and variance of one (as indicated by square brackets). 162 Moyes et al. (2013a) used the natural logarithm (In) but since log₁₀ and In are 163 proportional, $ln(y) = ln(10)log_{10}(y)$, the standardised values will be exactly equal, i.e. 164 $[\ln(y)] = [\log_{10}(y)]$. Thus, since the manuscripts of Grelet et al. (2019) and De Koster 165 et al. (2019) applied log₁₀-transformations of NEFA and BHB we decided to continue 166 this approach. 167

168

169 Metabolic clusters

As an alternative phenotype to negative EBAL and PI-index, clusters were created by
use of the k-means method of Hartigan and Wong (1979) from standardised
measures of plasma glucose, plasma log₁₀(BHB), plasma log₁₀(NEFA), and serum
log₁₀(IGF-1). As mentioned in the Introduction, these four blood biomarkers mirror the
physiological status of the animal. Three clusters (k=3) were constructed for each
combination of three parities (1, 2 and 3+ lactations) and two periods in early

lactation (around 14 and 35 DIM) as visualised in Figure 1. Deciding on the number
of clusters can be intricate but in the present sample k=3 was found to be a fair
compromise between size and similarity (in terms of the within cluster sum of
squares, results not shown). Based on a graphical interpretation using boxplots of the
standardised concentrations of plasma glucose, NEFA and BHB and serum IGF-1
(see Figure 1) three metabolic clusters were defined as representing balanced,
intermediate and imbalanced cows.

183

Criteria to define the imbalanced metabolic cluster are the most important. We 184 defined the metabolic cluster as imbalanced if standardised plasma glucose and 185 186 serum IGF-1 concentrations were both lower than those of plasma BHB and plasma NEFA, and in addition both median BHB and NEFA were above 0.5 SD (Figure 1). 187 Intermediate and balanced metabolic clusters had less sharp definitions: The 188 intermediate metabolic cluster generally had lower standardised glucose and IGF-1 189 concentrations than BHB and NEFA, with NEFA and BHB boxes in the ±0.5 SD area 190 and glucose and IGF-1 around or below -0.5 SD. The balanced metabolic cluster had 191 standardised glucose and IGF-1 concentrations around 0.5 SD and standardised 192 193 NEFA and BHB concentrations below or equal to those of glucose and IGF-1, or all four approximately equal and around -0.5 SD. The metabolic cluster was also 194 considered balanced if all four boxes were inside the ±0.5 SD area. 195

196

197 Milk biomarkers

Three different sets of milk biomarkers (MME, FT-MIR spectra and IgG *N*-glycans)
were considered as predictors. Metabolites and enzymes consisted of six milk
metabolites (glycose-6-phosphate, free glucose, BHB, isocitrate, urea and uric acid)

and two enzymes (NAGase and LDH). Fourier transform mid-IR spectra from the 6 201 farms were standardised into a common format. FT-MIR data consisted of 202 absorbance values at 212 wavenumbers selected from a total of 1060 by removal of 203 areas known to be non-reproducible between instruments or non-informative due to 204 the water component in milk (Grelet et al., 2016). Finally, 19 peaks of IgG N-glycans 205 were manually identified and integrated. Each peak's percentage of the total area 206 under the 19 peaks was used as the measure for the statistical analyses. Further 207 details on the laboratory analysis are given in De Koster et al. (2019). 208

209

210 Random forests predictions

Each of the three sets of milk biomarkers were used to predict the responses (EBAL, 211 PI-index and metabolic clusters) separately for each parity and period by use of the 212 random forests algorithm (see below), i.e. in total 54 predictions. In addition, each of 213 the six plasma metabolites and serum IGF-1 were predicted. To make a more fair 214 comparison with IgG N-glycans, we also made a comparison using only data that 215 were complete across all three sets of milk biomarkers in relation to the two periods; 216 around DIM 14 and DIM 35. Random forests belongs to the field of machine learning 217 218 and is an ensemble of classification or regression trees (Breiman, 2001) with each tree being a set of decision rules. A short description of the algorithm is given below, 219 whereas we refer to Breiman (2001) for a technical presentation and introduction to 220 random forests. We generally used default settings of the implementation except that 221 we used 2500 trees (instead of the default 500) to stabilise estimates of accuracy. 222

223

224 Random forests algorithm

In summary, for each of a pre-specified number of trees (default: 500) a sample is 225 drawn from the original data by sampling with replacement (bootstrap sample). 226 These samples have the same size as the original data but contain on average 227 approximately two thirds of the individual records, since some are selected more than 228 once and some not at all. Each bootstrap sample is used for training an unpruned 229 tree. At each node of the tree, a set of predictors (default for binary classification: 230 square root number of predictors) are chosen at random as candidates for splitting 231 the data present at the current (parent) node into two chunks. The algorithm then 232 choose the candidate (categorical) or cut-point (continuous) that give the largest 233 reduction of the Gini index (Breiman et al., 1984), i.e. the most homogeneous child 234 nodes. Each tree is grown as large as possible. The random selection of candidate 235 predictors at each node protects from overfitting (Breiman, 2001) and pruning is not 236 necessary. When the random forest of trees have been developed, new records are 237 passed through each tree and majority voting or averaging predicts their classes or 238 values. 239

240

241 Statistical analysis

242 The statistical analyses were carried out using R version 3.6.1 (R Core Team, 2019). For k-means clustering the *kmeans* function of R was used. Random forests 243 modelling was carried out by use of the randomForest package (Liaw and Wiener, 244 2002). We evaluated performance of random forests predictions for metabolic 245 clusters by a leave-one-cow-out (internal) cross-validation strategy, i.e. in turn 246 preserving data from one cow as test set and using data from the other cows for 247 training of a random forests model. By use of the *confusionMatrix* function of the 248 caret package (Kuhn, 2008) we calculated global accuracy (proportion of correctly 249

classified samples, i.e. the diagonal of the 3 by 3 contingency table of predicted versus true cluster also known as the confusion matrix), sensitivity for each cluster (proportion correctly predicted to that cluster) and specificity (proportion correctly predicted not to be in that cluster). In addition, the precision of predictions for the individual blood biomarkers, EBAL and PI-index was measured by the coefficient of determination of cross-validation (R^2_{cv}) and the root mean squared error (RMSE_{cv}).

To explore the ranking of the individual MME biomarkers within parity and period, the variable importance measure (VIM) was calculated (Breiman, 2001) and plotted using *randomForests*. This measure is based on the internal out-of-bag samples, i.e. the third not picked to be included in each bootstrap sample, see Breiman (2001).

261

Characteristics and differences among metabolic clusters in milk metabolite concentrations, enzyme activities and daily milk yield were examined separately for parity 2 and 3+ at DIM 14 by ANOVA with F-tests. Since most health events and imbalances are expected to happen in the first and middle part of the early lactation period, we only focused on DIM 14 for this part. First parity cows were not given further attention since none of these were classified to the imbalanced cluster at DIM 14 and all were in clusters classified as balanced at DIM 35.

269

270 **Results**

Summary statistics for production, blood biomarkers and MME can be found in tables
and supplementary tables of Krogh et al. (2019).

273

274

275 Predictions of EBAL and PI-index by sets of milk biomarkers

The precisions (R^{2}_{CV} and $RMSE_{CV}$) of predicting measures of EBAL and PI-index by 276 the three sets of milk biomarkers as determined by leave-one-cow-out cross-277 validation are shown in Table 1. The best precision was obtained when predicting PI-278 index by MME with an R²_{CV} of 0.40 at 14 DIM and 0.34 at 35 DIM. For FT-MIR, the 279 corresponding R²_{CV} was 0.26 and 0.19. For EBAL, however, FT-MIR showed a better 280 performance than MME with an R²_{CV} of 0.28 vs 0.21. The RMSEs from MME and FT-281 MIR predictions were respectively 23.7 and 23.4 for EBAL and between 1.62 and 282 1.96 for PI-index. Predictions by IgG *N*-glycans had the lowest precisions, with R^2_{CV} 283 ranging between 0.01 and 0.06 and with RMSE_{CV} being 26.3 for EBAL and 2.04 for 284 PI-index. 285

286

287 *Predictions of individual blood biomarkers by sets of milk biomarkers*

Predictions of individual blood biomarkers are shown in Table 2. The best precisions 288 were obtained with MMEs for plasma urea (R^2_{CV} = 0.62 for 14 DIM and 0.59 for 35 289 DIM) and for plasma BHB (R^{2}_{CV} = 0.46 and 0.40). Interestingly, plasma cholesterol 290 was not predicted that well (R^2_{CV} = 0.09 and 0.12) whereas precisions of serum IGF-291 1 were at the same level as plasma BHB for DIM 35 (R^2_{CV} = 0.40) and a bit lower for 292 DIM 14 (R^2_{CV} = 0.32). The precisions by IgG *N*-glycans were always the lowest 293 whereas generally, FT-MIR were at the same level as MME but in some cases much 294 295 lower.

296

297 *Metabolic cluster changes*

The number of cows in each of the three metabolic clusters at DIM 14 and DIM 35 is reported in Table 3 with indication of changes between the two periods. All the 52

primiparous cows were interpreted balanced at DIM 35. Among the 28 parity 2 cows 300 in the intermediate cluster at DIM 14, 17 (61%) did not shift to a cluster deemed to be 301 more "balanced" at DIM35, staying in an intermediate cluster, while the rest changed 302 to a balanced cluster (N=11). Most of the 23 parity 2 cows in the balanced cluster at 303 DIM 14 stayed in a balanced cluster at DIM 35 (N=21) with only two cows shifting; 304 one to an imbalanced and one to an intermediate cluster at DIM 35. For 15 (4+11) 305 out of 18 (7+11) (83%) parity 2 and 3+ cows in the imbalanced cluster DIM 14, extra 306 attention may be relevant as they were also in an imbalanced cluster DIM 35. 307 Concerning parity 3+ cows in the balanced cluster DIM 14, 31 out of 38 (82%) were 308 still in a balanced cluster at DIM 35 while the rest changed to an imbalanced cluster. 309 310 Of the 54 parity 3+ cows in the intermediate cluster DIM 14, 39 (72%) changed to a balanced cluster at DIM 35, while the rest changed to an imbalanced cluster. 311

312

313 Prediction of metabolic clusters

Accuracies to predict the clusters from sets of milk biomarkers with random forests 314 models are presented in Table 4 for each combination of parity (1, 2 and 3+) and 315 period (DIM 14 and 35). As in Grelet et al. (2019) and De Koster et al. (2019), 316 317 including milk yield as a factor in the aim to help distinguishing between classes did not improve the accuracy (results not shown). Global accuracies from MME and FT-318 MIR were at the same level and ranged from 0.54 to 0.65 for MME and 0.51 to 0.68 319 for FT-MIR. Accuracies were lower for IgG *N*-glycans; ranging from 0.32 to 0.53. The 320 sensitivity for prediction of the imbalanced cluster was better with MME than with FT-321 MIR and IgG *N*-glycans. Unfortunately, examples of zero sensitivity (none predicted 322 correctly) were seen, likely due to a relatively low number of cows in the imbalanced 323 clusters, see Table 3. 324

325

Results from predictions using only data that were complete across all three sets of 326 milk biomarkers in each period are shown in Supplementary Table S2 and are less 327 stable with confidence intervals that are bit wider due to the smaller number of 328 observations. Nevertheless, predictions by IgG *N*-glycans tend to be less 329 unfavourable compared to MME and FT-MIR when judged on this reduced data set, 330 potentially giving a more fair comparison. Global accuracies tended to be lower with 331 the reduced data set and ranged from 0.39 to 0.59 for MME, 0.34 to 0.67 for FT-MIR 332 and 0.19 to 0.57 for IgG N-glycans. Using this reduced data set, we also examined 333 the pairwise agreement of predictions among the three sets of milk biomarkers, see 334 Supplementary Table S3. The best agreement with a global accuracy of 0.76 (95%) 335 CI: 0.62-0.87) was found between MME and FT-MIR for parity 3+ cows around DIM 336 14 but it should be noted that for these, none of the cows in the imbalanced cluster 337 were correctly determined by FT-MIR. The lowest agreement was seen between FT-338 MIR and IgG N-glycans for parity 3+ cows around DIM 35 with a global accuracy of 339 0.27 (0.16-0.41). Generally, the agreements were at the same level among all three 340 sets of milk biomarkers. 341

342

To ease comparison with table 6 in Grelet et al. (2019) and figure 5 in De Koster et al. (2019), we calculated the global accuracy for predicting the imbalanced cluster vs intermediate and balanced combined. For MME in parity 3+ this accuracy was 0.97 (0.92-0.99) and 0.82 (0.73-0.89) for DIM 14 and 35, respectively. For FT-MIR the corresponding accuracies were 0.89 (0.81-0.95) and 0.69 (0.59-0.78) and for IgG *N*glycans 0.92 (0.82-0.97) and 0.53 (0.40-0.66). These accuracies tend to be higher DIM 14 and at the same level or lower DIM 35 than those found in Grelet et al. (2019)

and De Koster et al. (2019). For parity 2, number of cows in the imbalance clusters
were quite low (see Table 3) and almost all sensitivity estimates were 0 and
specificities at or close to 1 (see Table 4). Thus, parity 2 accuracies are high (e.g.
0.93 (0.83-0.98) for MME at 14 DIM) but driven by specificity.

354

355 Differences in milk metabolite contents among metabolic clusters

Considering further the characteristics of parity 2 and 3+ cows at DIM 14, Table 5 356 presents quartiles for milk yield, metabolites and enzymes for each of the three 357 metabolic clusters. These results indicate that some of the milk metabolites and 358 enzymes were significantly different between the three metabolic clusters. The 359 360 concentration of free glucose was significantly lower in the imbalanced cluster while, generally, those of BHB and isocitrate were higher. For the parity 2 cows, glucose-6-361 phosphate, and free glucose concentrations were higher for the balanced cluster 362 than for the imbalanced, while for BHB, isocitrate and NAGase the concentrations or 363 activities were lower or tended (P = 0.07) to be lower for the balanced compared to 364 the imbalanced cluster. For parity 3+ cows, glucose-6-phosphate did not differ 365 between the metabolic clusters but otherwise the results were similar to those of 366 second parity cows. For parity 3+ cows, the urea concentration also tended (P=0.07) 367 to be higher for the imbalanced cluster compared with the balanced cluster. To 368 explore the ranking of importance within parity and period for the eight milk 369 metabolites and enzymes in the MME set of milk biomarkers, VIM plots are shown in 370 Supplementary Figures S1 to S4. BHB is among the most important for both the 14 371 and 35 DIM periods whereas isocitrate is important for both parity in the period 372 around DIM 14 but only for the oldest (3+) cows around DIM 35. For second lactation 373 cows around DIM 35, free glucose and LDH are marginally more important than BHB 374

which ranks third. For the oldest cows (3+) free glucose is more important than
isocitrate around DIM 14 whereas around DIM 35, uric acid and urea are also
important for the prediction of the metabolic clusters.

378

379 **Discussion**

The objective was to compare the use of three different sets of milk biomarkers for 380 identification of cows in physiological imbalance and thus at risk of developing a 381 metabolic or infectious disease. We defined a metabolic imbalanced cluster of cows 382 based on k-means clustering of four blood biomarkers; glucose, NEFA and BHB in 383 plasma and IGF-1 in serum. Random forests was used to predict individual blood 384 biomarkers, body energy balance (EBAL), index for physiological imbalance (PI-385 index) and the clusters differentiating the metabolic status of cows. Ideally, the 386 prediction algorithms should be validated using an external data set but this was not 387 possible in the present study. Therefore, internal cross-validation was used to 388 examine performance. 389

390

IgG *N*-glycans performed really poor compared to the other two sets of milk 391 392 biomarkers for predictions of individual blood biomarkers, EBAL, PI-index and metabolic clusters. This may partly be due to a less dense sampling of this milk 393 biomarker. Nevertheless, even when accounting for the difference in sampling 394 395 density IgG *N*-glycans had lower prediction accuracies than MME, FT-MIR or both. In addition, the analytical procedure is very complicated, expensive and with large 396 problems of getting reliable results. Thus, also in that respect more work is needed to 397 make this milk biomarker useful in herd health management. 398

399

The precision of predictions for the individual blood biomarkers, EBAL and PI-index was measured by the coefficient of determination of cross-validation (R^2_{cv}) and by the root mean squared error ($RMSE_{cv}$). These two measures of precision were interpreted with the recommendations from Alexander et al. (2015) in mind that as a rule of thumb the R^2 should higher than 0.6 and the RMSE within 10% of the outcome's range.

To predict individual blood biomarkers, the best models were obtained by MME with 407 R_{CV}^2 of 0.62 and 0.59 for plasma urea at 14 and 35 DIM, respectively. These were 408 the only predictions reaching the 0.6 threshold mentioned above. Moreover, RMSE_{CV} 409 for MME predictions (0.72 and 0.78) were below 10% of the plasma urea range at 410 8.45 mM (supplementary tables of Krogh et al., 2019). The R²_{CV} for FT-MIR models 411 were generally lower than for MME and in some cases much lower, e.g. 0.06 (DIM 412 14) and 0.13 (DIM 35) for plasma urea. Correspondingly, the RMSE_{CV} were higher, 413 e.g. 1.08 and 1.13 for plasma urea at 14 and 35 DIM. Lower performances of the FT-414 MIR models, compared to Grelet et al. (2019), may possibly be explained by different 415 methodologies. In that study all DIM were combined into one global model, 416 417 distribution of data were artificially modified and partial least squares regression was used instead of random forests. These differences were one of the reasons for 418 redoing the FT-MIR predictions in the present paper. 419

420

For EBAL, FT-MIR showed a better performance than MME with an R^2_{CV} of 0.28 vs 0.21 whereas the opposite was the case when predicting PI-index with R^2_{CV} of 0.26 vs 0.40 at 14 DIM and 0.19 vs 0.34 at 35 DIM. Clearly these are below the 0.6 rule of thumb. The RMSEs from EBAL predictions (23.4 and 26.3) were lower than 10% of

⁴⁰⁶

the absolute range, whereas for PI-index only RMSEs from MME predictions (1.62
and 1.71) were around 10% of the absolute range.

427

Metabolic clusters were created as alternative phenotypes. The global accuracy of 428 predicting the metabolic clusters varied from 0.54 to 0.65 and 0.51 to 0.68 for MME 429 and FT-MIR predictions, respectively. Thus, the performance of MME and FT-MIR 430 was at an equal level. It should be noted that examples of sensitivity at zero and 431 specificity close to one were seen and may have biased the accuracy upwards. 432 There was no improvement of including daily milk yield in the prediction models, as 433 also concluded by Ingvartsen et al. (2003). It is not milk yield per se that increases 434 the risk of diseases but rather physiological imbalance reflecting difficulties for some 435 animals to adapt to the major physiological changes that occur particularly in the 436 transition cow. Moreover, this is in accordance with results in Grelet et al. (2019) and 437 De Koster et al. (2019) though comparison with these two studies is complicated by 438 differences in examined periods and parities. The present study did notice 439 differences in blood biomarker profiles among parities but more data would be 440 desirable for such differentiation. In this study, work has focused on the first 7 weeks 441 after calving and does not apply to cows at later stages. Since no clusters of 442 primiparous cows were considered imbalanced, it generally seems from the present 443 study that first parity cows do not require extra care and the attention should be on 444 the multiparous cows. Relatively few cows in the imbalance clusters were also 445 observed for parity 2 accompanied by sensitivity estimates at zero and specificities 446 close to one. Thus, neither first nor second parity cows were really informative for the 447 ability to predict the imbalanced cluster. 448

449

The purpose of the presented random forests algorithms were to identify cows in 450 physiological imbalance at risk of developing subclinical or more severe stages of 451 diseases. Such cows may need extra attention and potentially altered feeding or 452 other management actions to avoid that the physiological imbalance develop into 453 subclinical or more severe disease states. The required accuracy of detection is 454 obviously lower for this purpose since there is no risk of harm to the animal or of 455 needless use of medicine. The accuracies mentioned in this paper are likely too low 456 for diagnosing diseases that require medical treatment with e.g. antibiotics. 457 Generally, the required accuracy depends on the specific purpose and of e.g. 458 disease prevalence, costs associated with treatment and possible side-effects. The 459 required accuracy could be established by simulation methods. Possibly, a larger 460 461 data set for training prediction algorithms would improve the accuracies and the results presented here may be used to guide sample size decisions for future 462 studies. 463

464

Presently, no sensors are available to measure e.g. free glucose, isocitrate and 465 glucose-6-phosphate, but since FT-MIR algorithms tended to give as accurate 466 predictions as MME, FT-MIR may give the same opportunities to make relevant 467 classification of cows as balanced or in physiological imbalance (see also Grelet et 468 al., 2019 and De Koster et al., 2019). Moreover, it would also be interesting to 469 investigate direct prediction of udder inflammation from FT-MIR as opposed to the 470 use of e.g. LDH and NAGase enzymes that constitute an alternative for somatic cell 471 counts, helping in the detection of subclinical diseases (Kitchen et al., 1978; Larsen 472 et al., 2010; Hovinen et al., 2016). 473

474

475 Conclusion

Neither EBAL nor PI-index were sufficiently precise to be used as a management tool 476 for identification of risk cows. As an alternative, cows were divided into clusters 477 based on measures of glucose, BHB and NEFA in plasma and IGF-1 in serum. 478 These can be interpreted into metabolic clusters and the cluster of imbalanced cows 479 can be predicted equally well by MME and FT-MIR. Nevertheless, accuracies still 480 need to be improved and a larger data set for training the prediction algorithms would 481 probably be needed. Free glucose, isocitrate, glycose-6-phosphate, BHB and 482 NAGase measured in milk were significantly different among the three metabolic 483 clusters (balanced, intermediate and physiological imbalanced). Thus, if MME is the 484 preferred set of milk biomarkers to predict cows in physiological imbalance and at 485 risk of developing a production or infectious disease, the above mentioned 486 metabolites and enzyme should have high priority for inclusion. The use of IgG N-487 glycans for prediction still needs development. 488

489

490 Author's contribution

LF, CGa, MAK, MTS and KLI made the first draft of the paper. LF, CGr, MS, MH and 491 492 other partners from GC undertook data handling and data quality control. LF, CGr, MS and MH did the major parts of the statistical analyses including the conception of 493 the idea of using k-means clusters to combine selected blood biomarkers with 494 contribution to the latter from MTS and KLI. LF, CGa, MAK, MTS and KLI 495 collaboratively defined the metabolic interpretation of these clusters. MTS, MAC, KLI 496 and other partners from GC did the conception and designed the study. TL handled 497 storage of milk and blood samples and did lab analyses of milk metabolites, milk 498 enzymes and blood metabolites and assisted during the data guality control of these 499

500 biomarkers. CGr and other partners from GC undertook analyses and calibrations of

501 FT-MIR. EM, ROF, FC, MAC and other partners form GC did lab analyses and

502 interpretation of IgG *N*-glycans. All authors critically revised the first draft and

⁵⁰³ approved the final version of the manuscript.

504

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511

512 **Declaration of interest**

513 There is no direct financial interest of the authors and affiliations in the subject matter

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515

516 **Ethics statement**

517 The experiments were carried out in accordance with the standards recommended

518 by the EU Directive 2010/63/EU for animal experiments.

519

520 Software and data repository resources

521 None of the data were deposited in an official repository.

522

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622 Figure captions

623

- 624 **Figure 1** Box-and-whiskers plots for graphical interpretation (note that bars are
- 625 medians) of k-means clusters into metabolic clusters as indicated by colours:
- balanced cluster (magenta), intermediate cluster (orange) and physiological
- imbalanced cluster (yellow). Distribution of standardised blood metabolites and IGF-1
- in each cluster (1, 2 and 3), at 14 DIM (first row), at 35 DIM (second row), for
- 629 primiparous Holstein dairy cows (first column), second parity cows and for parity 3+
- 630 cows (last column). The horizontal lines indicate +/-0.5 SD.
- 631
- 632 Tables

Table 1 Precision of random forests predictions of EBAL and PI-index with three sets
 635 of milk biomarkers (milk metabolites and enzymes (MME). Fourier transform mid-IR 636 spectra (FT-MIR) and immunoglobulin G (IgG) N-glycans)¹ in Holstein dairy cows in 637 six herds. The performance was measured by the coefficient of determination of leave-638 one-cow-out cross-validation (R^{2}_{CV}) and by root mean squared error (RMSE_{cv}). 639 Individual milk biomarkers were standardised using all available data before matching. 640 In addition to sets of milk biomarkers, parity (1, 2 and 3+) as a factor and DIM (days in 641 milk) as continuous covariate were included as predictors for EBAL, whereas only 642 parity was added as predictor for PI-index. Number of cows (samples) are after 643 removal of records excluded due to missing values 644

Response	Period (DIM)	Sets of milk biomarkers	N_{cows} ($N_{samples}$)	R ² cv	RMSE _{cv}
EBAL (only using DK, IE and UK herds)	1-50	MME FT-MIR IgG	132 (1608) 132 (1230) 122 (328)	0.21 0.28 0.06	23.7 23.4 26.3
	14	MME FT-MIR IgG	216 201 133	0.40 0.26 0.01	1.62 1.86 2.04
PI-INDEX	35	MME FT-MIR IgG	218 195 134	0.34 0.19 0.05	1.71 1.93 2.04

¹ Milk biomarkers were matched with the EBAL closest in sampling date (+/- 3 days). For FT-MIR this matching 645 646 strategy was also applied to PI-index for the period noted in the column denoted "Period (DIM)". If no perfect match 647 (same day) was found, we proceeded as follows: Step 1 day backward first (day before milk biomarker sampling 648 date), then 2 days forward (i.e. 1 day after the sampling data), then 3 days back (corresponding to 2 days before 649 sampling), then 4 days forward, 5 days backward and 6 days forward. That is, closest match within 7 days (a week) 650 centred in the milk biomarker's sampling date. For IgG N-glycans, the measure from the period noted was used for 651 these two measurements. Averages of measures of milk metabolites and enzymes within the same week (Monday-652 Sunday) as blood sampling were used for PI-index.

Table 2 Precision (R² and RMSE by leave-one-cow-out cross-validation) of random forests predictions of plasma metabolites and serum IGF-1 with three sets of milk biomarkers (milk metabolites and enzymes (MME), Fourier transform mid-IR spectra (FT-MIR) and immunoglobulin G (IgG) N-glycans) in Holstein dairy cows. Individual milk biomarkers were standardised and the sample matching the blood sample date (+/- 3 days) was used. In addition, parity (1, 2 and 3+) was included as a predictor. Number of cows are after removal of those excluded due to missing values

Blood biomarker	Period (DIM)	Sets of milk biomarkers	N _{cows}	R ² _{cv}	RMSE _{cv}
		MME	213	0.12	16.9
	14	FT-MIR	198	0.11	17.2
Plasma fructosamine		lgG	131	0.03	17.6
		MME	214	0.18	16.4
	35	FT-MIR	191	0.02	18.5
		lgG	132	0.11	17.2
		MME	216	0.62	0.72
	14	FT-MIR	201	0.06	1.08
Plasma urea		lgG	133	0.01	1.07
		MME	218	0.59	0.78
	35	FT-MIR	195	0.13	1.13
		lgG	134	0.01	1.16
		MME	216	0.09	0.68
	14	FT-MIR	201	0.01	0.72
		lgG	133	0.01	0.72
Plasma cholesterol		MME	218	0.12	0.98
	35	FT-MIR	195	0.03	1.02
		lgG	134	0.04	1.02
		MME	216	0.13	0.25
	14	FT-MIR	201	0.10	0.26
		lgG	133	<0.01	0.26
Plasma log ₁₀ (NEFA)		MME	218	0.09	0.30
	35	FT-MIR	195	0.03	0.31
		lgG	134	0.01	0.32
		MME	216	0.29	0.41
	14	FT-MIR	201	0.23	0.43
		lgG	133	0.11	0.49
Plasma glucose		MME	218	0.32	0.43
	35	FT-MIR	195	0.19	0.48
		lgG	134	0.17	0.49
		MME	216	0.46	0.16
	14	FT-MIR	201	0.27	0.20
		lgG	133	0.04	0.24
Plasma log ₁₀ (BHB)		MME	218	0.40	0.17
	35	FT-MIR	195	0.25	0.19
		lgG	134	<0.01	0.22
		MME	216	0.32	0.27
	14	FT-MIR	204	0.36	0.26
		lgG	136	0.24	0.29
Serum log ₁₀ (IGF-1)		MME	216	0.40	0.21
	35	FT-MIR	197	0.35	0.22
		lgG	138	0.14	0.25

Table 3 Number of Holstein dairy cows per metabolic cluster (balanced,
intermediate, imbalanced) at DIM 14 and 35. Furthermore, the last column shows
which clusters the DIM 35 cows belonged to at DIM 14

Cluster and parity	Number	of cows	Cluster affiliation at DIM 14 for DIM 35 cours			
Cluster and parity	DIM 14	DIM 35	Cluster anniation at Dim 14 for Dim 55 cows			
Parity 1						
Balanced	38	52	38 Balanced + 14 Intermediate			
Intermediate	14	0				
Imbalanced	0	0				
Parity 2						
Balanced	23	32	21 Balanced + 11 Intermediate			
Intermediate	28	21	1 Balanced +17 Intermediate + 3 Imbalanced			
Imbalanced	7	5	1 Balanced + 4 Imbalanced			
Parity 3+						
Balanced	38	70	31 Balanced + 39 Intermediate			
Intermediate	54	0				
Imbalanced	11	33	7 Balanced +15 Intermediate + 11 Imbalanced			
Total	213	213				

Period	Cluster	Metabolic		Sensitivity			Specificity		Global accuracy ³ (98		5% CI)
and parity	number ¹	cluster ²	MME	FT-MIR	lgG	MME	FT-MIR	lgG	MME	FT-MIR	lgG
Parity 1											
DIM 14	1	Balanced	0.74	0.70	0.38	0.52	0.61	0.48	0.54	0.51	0.32
	2	Balanced	0.14	0.40	0.10	0.89	0.75	0.79	(0.39-0.68)	(0.37-0.65)	(0.17-0.51)
	3	Intermediate	0.60	0.31	0.45	0.84	0.87	0.70			
DIM 35	1	Balanced	0.63	0.25	0.00	0.98	0.90	1.00	0.62	0.68	0.43
	2	Balanced	0.68	0.83	0.69	0.63	0.71	0.21	(0.47-0.75)	(0.53-0.81)	(0.25-0.63)
	3	Balanced	0.53	0.69	0.18	0.73	0.87	0.68	. , ,		
Parity 2											
DIM 14	1	Imbalanced	0.50	0.00	0.00	0.98	0.98	1.00	0.55	0.59	0.46
	2	Balanced	0.50	0.70	0.42	0.68	0.65	0.70	(0.42-0.68)	(0.45-0.72)	(0.29-0.63)
	3	Intermediate	0.61	0.70	0.61	0.53	0.68	0.29			
DIM 35	1	Imbalanced	0.00	0.00	0.00	0.98	0.96	1.00	0.58	0.55	0.53
	2	Balanced	0.79	0.69	0.71	0.50	0.52	0.53	(0.44-0.70)	(0.40-0.69)	(0.35-0.70)
	3	Intermediate	0.36	0.50	0.44	0.70	0.71	0.60			. ,
Parity 3+											
DIM 14	1	Imbalanced	0.70	0.00	0.00	1.00	0.99	1.00	0.63	0.66	0.51
	2	Intermediate	0.74	0.76	0.74	0.51	0.63	0.17	(0.53-0.73)	(0.56-0.76)	(0.38-0.64)
	3	Balanced	0.46	0.70	0.17	0.78	0.76	0.74	, , , , , , , , , , , , , , , , , , ,	, , , , , , , , , , , , , , , , , , ,	, , , , , , , , , , , , , , , , , , ,
DIM 35	1	Imbalanced	0.71	0.59	0.10	0.87	0.74	0.73	0.65	0.59	0.44
	2	Balanced	0.71	0.63	0.71	0.68	0.82	0.70	(0.55-0.75)	(0.49-0.70)	(0.31-0.57)
	3	Balanced	0.50	0.56	0.45	0.90	0.83	0.73	-		. ,

668 clustering (k=3) of standardised values of plasma glucose, $log_{10}(BHB)$ and $log_{10}(NEFA)$ and serum $log_{10}(IGF-1)$ in Holstein dairy cows

Table 4 Leave-one-cow-out cross-validation of performance for random forests predictions of metabolic clusters by milk metabolites

and enzymes (MME), Fourier transform mid-IR (FT-MIR) spectra and immunoglobulin G (IgG) N-glycans. Clusters based on k-means

⁶⁶⁹ ¹ The cluster numbers are arbitrary and cannot be compared among period/parity combinations.

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⁶⁷⁰ ² As interpreted from Figure 1. The metabolic clusters are comparable among period/parity combinations.

⁶⁷¹ ³ Proportion of correctly classified observations by the prediction, i.e. the diagonal of the confusion matrix.

Table 5 Characteristics¹ of milk yield, metabolites and enzymes and comparisons among the 672 three metabolic clusters (balanced, intermediate and physiological imbalanced) of Holstein dairy 673 cows at DIM 14 in parity 2 and 3+, respectively. Results of ANOVA F-tests for differences among 674

Milk measure and	Balanced (n=24) ⁴		Intern	nediate	(n=28)	Imba	Imbalanced (n=9)4			
parity	Q1	Q2	Q3	Q1	Q2	Q3	Q1	Q2	Q3	
Parity 2										
Glucose-6-P (mM)	0.17	0.22	0.28	0.14	0.18	0.20	0.16	0.18	0.23	*
Free glucose (mM)	0.18	0.25	0.28	0.17	0.22	0.26	0.07	0.12	0.15	**
log ₁₀ (BHB) ³	1.56	1.63	1.72	1.66	1.76	1.85	1.98	2.06	2.40	***
Isocitrate (mM)	0.15	0.17	0.19	0.17	0.19	0.20	0.19	0.28	0.29	**
Urea (mM)	2.47	3.15	3.83	2.16	3.18	3.79	2.66	2.82	4.90	ns
Uric acid (µM)	161	176	204	154	164	203	139	173	181	ns
log ₁₀ (NAGase) ³	0.24	0.35	0.46	0.18	0.26	0.41	0.41	0.42	0.46	ns
log ₁₀ (LDH) ³	0.37	0.46	0.63	0.42	0.56	0.68	0.46	0.57	0.72	ns
Milk yield (kg/day)	30.5	32.4	36.8	26.3	31.6	35.9	28.2	30.5	34.4	ns
	Balanced (n=39) ⁴			Intern	Intermediate (n=54)			Imbalanced (n=11)		
Parity 3+										
Glucose-6-P (mM)	0.15	0.19	0.24	0.15	0.17	0.22	0.16	0.18	0.20	ns
Free glucose (mM)	0.17	0.21	0.24	0.13	0.16	0.18	0.09	0.10	0.11	***
log ₁₀ BHB ³	1.55	1.66	1.74	1.66	1.74	1.92	2.05	2.12	2.23	***
Isocitrate (mM)	0.14	0.16	0.19	0.15	0.18	0.21	0.22	0.26	0.28	***
Urea (mM)	2.26	3.12	3.63	1.87	2.76	3.57	2.96	3.17	4.62	ns
Uric acid (µM)	126	166	200	114	155	187	144	174	203	ns
log ₁₀ (NAGase) ³	0.17	0.27	0.36	0.24	0.35	0.47	0.48	0.55	0.62	**
log ₁₀ (LDH) ³	0.28	0.41	0.61	0.38	0.48	0.67	0.55	0.64	0.73	ns
Milk yield (kg/day)	34.3	36.4	40.6	32.1	34.6	38.6	29.9	33.0	36.7	ns

metabolic clusters are indicated² 675

¹ Q1: first quartile, Q2: second quartile (median), Q3: third quartile, M: molar (mol/L). 676

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² ns P \geq 0.05; * P<0.05; ** P<0.01; *** P<0.001 ³ BHB (μ M), NAGase (units/L), LDH (units/L). ⁴ The difference in totals compared to Table 3 is due to cows only having measures DIM 14. 679



Predicting physiological imbalance in Holstein dairy cows by three different sets of milk biomarkers

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Supplementary Table S1. Number of Holstein dairy cows (row proportion) and summary statistics of parity (mean, SD, median and maximum) for each combination of parity, herd and diet, and pooled

			Parity			
Herd ¹	Diet ²	1	2	3+	Total	Mean (SD); median; max
UK	Low C	6 (0.30)	4 (0.20)	10 (0.50)	20	2.6 (1.5); 2.5; 7
	Standard C	6 (0.30)	2 (0.10)	12 (0.60)	20	2.9 (1.6); 3; 6
	High C	6 (0.29)	3 (0.14)	12 (0.57)	21	2.8 (1.6); 3; 7
	Pooled	18 (0.30)	9 (0.15)	34 (0.56)	61	2.7 (1.6); 3; 7
DK	High starch	5 (0.45)	2 (0.18)	4 (0.36)	11	2.5 (1.8); 2; 5
	High sugar	4 (0.40)	3 (0.30)	3 (0.30)	10	2.5 (1.8); 2; 6
	Standard	2 (0.14)	9 (0.64)	3 (0.21)	14	2.1 (0.6); 2; 3
	Pooled	11 (0.31)	14 (0.40)	10 (0.29)	35	2.3 (1.4); 2; 6
IE	Standard	2 (0.06)	11 (0.31)	23 (0.64)	36	3.3 (1.5); 3; 7
BE	Standard	13 (0.42)	9 (0.29)	9 (0.29)	31	2.3 (1.6); 2; 6
DE	Standard	3 (0.12)	8 (0.31)	15 (0.58)	26	2.5 (0.7); 3; 3
IT	Standard	8 (0.18)	15 (0.33)	22 (0.49)	45	2.6 (1.2); 2; 6
All	Pooled	55 (0.24)	66 (0.28)	113 (0.48)	234	2.6 (1.4); 2; 7

¹ UK (Agri-Food and Biosciences Institute, Northern Ireland, UK); DK (Aarhus University, Denmark); IE (UCD Lyons Research Farm, University College Dublin, Ireland); BE (Walloon Agricultural Research Centre, Belgium); DE (Leibniz Institute for Farm Animal Biology, Germany) and IT (Consiglio per la Ricerca in Agricoltura, Italy).

² C=concentrate.

Supplementary Table S2 Leave-one-cow-out cross-validation of prediction performance for milk metabolites and enzymes (MME), Fourier transform mid-IR spectra (FT-MIR) and immunoglobulin G (IgG) N-glycans predictions of metabolic clusters based on *k*-means clustering (k=3) of standardised values of plasma glucose, plasma log₁₀(BHB), plasma log₁₀(NEFA), and serum log₁₀(IGF-1) in Holstein dairy cows. Data with the restriction that all three milk biomarkers were successfully measured in the period

Period	Cluster	Metabolic		Sensitivity			Specificity		Global accuracy ³ (95% CI)		
and parity	number1	cluster ²	MME	FT-MIR	lgG	MME	FT-MIR	lgG	MME	FT-MIR	lgG
Parity 1											
DIM 14	1	Balanced	0.62	0.69	0.38	0.39	0.53	0.16			
	2	Balanced	0.00	0.10	0.00	0.82	0.73	0.64	0.39	0.34	0.19
	3	Intermediate	0.44	0.11	0.11	0.82	0.74	0.91	(0.22-0.58)	(0.19-0.53)	(0.07-0.36)
DIM 35	1	Balanced	0.00	0.00	0.00	1.00	0.92	1.00	0.50	0.07	0.44
	2	Balanced	0.79	0.86	0.57	0.46	0.77	0.31		(0.35-0.75) (0.46-0.83)	0.41 (0.22-0.61)
	3	Balanced	0.40	0.60	0.30	0.71	0.76	0.59	(0.35-0.75)		
Parity 2											
DIM 14	1	Imbalanced	0.00	0.00	0.00	0.96	0.96	1.00			
	2	Balanced	0.33	0.75	0.42	0.63	0.71	0.76	0.39	0.67	0.48
	3	Intermediate	0.50	0.81	0.69	0.27	0.76	0.29	(0.22-0.58)	(0.48-0.82)	(0.31-0.66)
DIM 35	1	Imbalanced	0.00	0.33	0.00	1.00	0.96	1.00	0.50	0.50 0.04	0.57
	2	Balanced	0.64	0.64	0.45	0.59	0.71	0.82	0.50	0.01	
	3	Intermediate	0.50	0.64	0.79	0.50	0.64	0.36	(0.31-0.69)	(0.41-0.79)	(0.37-0.76)
Parity 3+											
DIM 14	1	Imbalanced	0.00	0.00	0.00	1.00	1.00	1.00			
	2	Intermediate	0.85	0.76	0.70	0.19	0.50	0.32	0.59	0.65	0.53
	3	Balanced	0.24	0.61	0.33	0.86	0.78	0.70	(0.45-0.72	(0.51-0.78)	(0.39-0.66)
DIM 35	1	Imbalanced	0.50	0.00	0.12	0.79	0.71	0.87	0.50	0.05	0.45
	2	Balanced	0.47	0.58	0.63	0.66	0.64	0.78	0.50	0.35	0.45
	3	Balanced	0.68	0.42	0.58	0.89	0.67	0.53	(0.41-0.69)	(0.22-0.49)	(0.32-0.59)

¹ The cluster numbers are arbitrary and cannot be compared among period/parity combinations.

² As interpreted from Figure 1. The metabolic clusters are comparable among period/parity combinations.

³ Proportion of correctly classified observations by the prediction, i.e. diagonal of the confusion matrix.

Supplementary Table S3 Pairwise comparisons of agreement by leave-one-cow-out cross-validation among milk metabolites and enzymes (MME), Fourier transform mid-IR spectra (FT-MIR) and immunoglobulin G (IgG) N-glycans for prediction of metabolic clusters based on k-means clustering (k=3) of standardised values of plasma glucose, plasma log₁₀(BHB), plasma log₁₀(NEFA), and serum log₁₀(IGF-1) in Holstein dairy cows. Data with the restriction that all three milk biomarkers were successfully measured in the period

Period	Cluster	Metabolic cluster ²	Sensitivity			Specificity			Global accuracy ³ (95% CI)		
and parity	number1		MME / FT-MIR	MME / IgG	FT-MIR / lgG	MME / FT-MIR	MME / IgG	FT-MIR / IgG	MME / FT-MIR	MME / IgG	FT-MIR / IgG
Parity 1											
DIM 14	1	Balanced	0.76	0.65	0.52	0.57	0.45	0.36	0 55	0.50	0.00
	2	Balanced	0.14	0.25	0.13	0.88	0.91	0.75	0.55	0.52	0.38
	3	Intermediate	0.43	0.33	0.00	0.79	0.75	0.76	(0.36-0.73)	(0.33-0.70)	(0.21-0.56)
DIM 35	1	Balanced	0.00	_4	_4	1.00	1.00	0.93	0.56	0.37	0.48
	2	Balanced	0.73	0.53	0.53	0.42	0.10	0.40	(0.25, 0.75)		
	3	Balanced	0.40	0.10	0.40	0.71	0.53	0.65	(0.35 - 0.75)	(0.19-0.56)	(0.29-0.00)
Parity 2											
DIM 14	1	Imbalanced	0.00	_4	_4	0.97	0.97	0.97			
	2	Balanced	0.31	0.30	0.50	0.61	0.62	0.57	0.42	0.48	0.52
	3	Intermediate	0.53	0.57	0.52	0.29	0.30	0.50	(0.25-0.61)	(0.30-0.67)	(0.34-0.69)
DIM 35	1	Imbalanced	0.00	_4	_4	1.00	1.00	0.93	0.54	0.57	0.64
	2	Balanced	0.58	0.63	0.75	0.56	0.55	0.70	(0.34, 0.72)	(0.27.0.76)	(0.44, 0.91)
	3	Intermediate	0.57	0.55	0.60	0.57	0.63	0.75	(0.34-0.72)	(0.37-0.76)	(0.44-0.81)
Parity 3+											
DIM 14	1	Imbalanced	_4	_4	_4	1.00	1.00	1.00			
	2	Intermediate	0.94	0.81	0.63	0.39	0.12	0.29	0.76	0.59	0.53
	3	Balanced	0.39	0.12	0.29	0.94	0.81	0.63	(0.62-0.87	(0.45-0.72)	(0.39-0.66)
DIM 35	1	Imbalanced	0.27	0.29	0.14	0.70	0.70	0.79	0.20	0.20	0.07
	2	Balanced	0.29	0.47	0.25	0.53	0.66	0.46		0.39	U.Z/ (0.16.0.44)
	3	Balanced	0.32	0.36	0.32	0.69	0.73	0.59	(0.10-0.44)	(0.20-0.53)	(0.16-0.41)

¹ The cluster numbers are arbitrary and cannot be compared among period/parity combinations.

² As interpreted from Figure 1. The metabolic clusters are comparable among period/parity combinations.

³ Proportion of predictions that are the same between methods, i.e. diagonal of the confusion matrix.

⁴ None predicted in the cluster by the "reference" milk biomarker (last mentioned, e.g. IgG).



Supplementary Figure S1 *Plot of the variable importance measure (VIM) from the random forests algorithm predicting metabolic clusters by a milk biomarker set of eight milk metabolites and enzymes measured around 14 days after calving (DIM14) in second parity cows.*



Supplementary Figure S2 *Plot of the variable importance measure (VIM) from the random forests algorithm predicting metabolic clusters by a milk biomarker set of eight milk metabolites and enzymes measured around 14 days after calving (DIM14) in cows with three or more lactations (parity 3+).*



Supplementary Figure S3 *Plot of the variable importance measure (VIM) from the random forests algorithm predicting metabolic clusters by a milk biomarker set of eight milk metabolites and enzymes measured around 35 days after calving (DIM35) in second parity cows.*



Supplementary Figure S4 *Plot of the variable importance measure (VIM) from the random forests algorithm predicting metabolic clusters by a milk biomarker set of eight milk metabolites and enzymes measured around 35 days after calving (DIM35) in cows with three or more lactations (parity 3+).*