

# Assessing serum metabolite profiles as predictors for feed efficiency in broiler chickens reared at geographically distant locations

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- 1 Running head: Serum predictors for RFI in chickens
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3	Assessing serum metabolite profiles as predictors for feed efficiency in
4	broiler chickens reared at geographically distant locations
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#### 25 Abstract

26 1. Various physiological mechanisms contribute to feed efficiency (FE) in chickens. Blood

27 metabolite profiles may correlate to the animal's FE, but have rarely been evaluated in

- 28 chickens. The objective of this study was to investigate differences in growth performance,
- 29 serum intermediary metabolites, acute-phase-proteins and white blood cells in low, medium
- 30 and high residual feed intake (RFI) chickens. It was also assessed if the environment affects
- 31 the FE and FE-related performance and serum profiles of chickens.
- 32 2. Individual BW and feed intake (FI) were recorded from d 7 of life. At 5 weeks of age,
- female and male chickens (Cobb 500) were selected according to their RFI (L1: Austria; L2:
- 34 UK; n = 9/RFI group, sex and location) and blood was collected.
- 35 3. Chickens at L1 had similar FI but a 15%-higher (P < 0.001) BW gain compared to chickens
- at L2. The RFI values of female chickens were -231, 8 and 215 g and those of male chickens -
- 197, 0 and 267 g for low, medium and high RFI, respectively (P < 0.001).
- 4. Location affected serum glucose, urea, cholesterol, NEFA and ovotransferrin in females,
- and serum glucose and triglycerides in male chickens (P < 0.05). Serum uric acid and NEFA
- 40 linearly increased from low to high RFI in females, whereas in males cholesterol showed the
- 41 same linear response from low to high RFI (P < 0.05). Serum alpha-1-acid glycoprotein and
- 42 blood heterophil-to-lymphocyte ratio linearly increased by 35 and 68%, respectively, from
- 43 low to high RFI but only in male chickens at L1 (P < 0.05).
- 5. Regression analysis showed positive relationships between RFI and serum uric acid ( $R^2 = 0.49$ ; P < 0.001) and cholesterol ( $R^2 = 0.13$ ; P < 0.001).
- 46 6. We conclude that RFI-related variation in serum metabolites of chickens was largely
- similar for the two environments and that serum metabolite patterns could be used to predictRFI in chickens.
- 49

- 50 Key words: acute phase response, broilers, feed-efficiency predictor, residual feed intake,
- 51 serum metabolites

### 52 Introduction

Improving feed efficiency (FE) is a continuing goal since feed is the major cost in chicken 53 production. Improved FE is often associated with reduced feed intake (FI) (Bottje and 54 Carstens, 2009). As it is a heritable trait and is independent of production traits, the residual 55 feed intake (RFI) has become the metric of choice for studying physiological mechanisms 56 underlying variation in FE of chickens and other livestock species (Herd and Arthur, 2009; 57 Berry and Crowley, 2012). Generally, a chicken population from a commercial breed shows 58 considerable variation in RFI (van Eerden et al., 2004). As knowledge about RFI related 59 physiological mechanisms in poultry and other livestock species advances, the biological 60 61 basis of inter-animal variations associated with FE becomes clearer (Bottje and Carstens, 62 2009; Aggrey et al., 2014; Lee et al. 2015; Mignon-Grasteau et al., 2015; Zhuo et al., 2015). In beef cattle and pigs some plasma metabolites and hormones correlated with animal's RFI 63 and have been discussed as RFI predictors (Kelly et al., 2010; Le Naou et al., 2012; 64 Montagne et al., 2014). Some evidence for RFI-associated differences in serum intermediary 65 metabolites also exists for cockerels (Gabbarou et al., 1997; Swennen et al., 2007); however, 66 due to the short production cycle, these have not been satisfactorily studied in meat-type 67 chickens. In general, peripheral blood is more easily accessible than other body tissues and 68 69 can provide useful information to identify the main biological processes which are modulated by genetic selection or by feeding strategies (Jegou et al., 2016). 70

The question whether meat-type chickens of diverging RFI respond differently to stressors
which may affect growth performance (Zulkifli *et al.*, 2014) has also not been completely
answered. In pullets, for instance, differences in RFI-related stress responses are small (van
Eerden *et al.*, 2004). Overall, concentrations of blood corticosterone, acute-phase-proteins
(APP) and heterophil-to-lymphocyte ratio (H-to-L) correlate in poultry (Gross and Siegel,
1983). Hence, APPs and the H-to-L ratio are commonly used as indices of stress in chickens

(Zulkifli *et al.*, 2000, 2014) and may help understanding RFI-related stress responses in
broiler chickens.

In most studies, RFI was derived from one contemporary population of chickens (Bottje 79 and Carstens, 2009), whereas information regarding the impact of the rearing environment on 80 RFI-related variation is scarce. In considering that substantial batch-to-batch variation has 81 been reported for the chicken gut microbiota under controlled conditions at one experimental 82 83 setting (Stanley et al., 2013, 2016; Ludvigsen et al., 2016), it is feasible that the environment may modify RFI-related physiological responses. This is an important issue since any 84 predictors or biomarkers of FE must be applicable across multiple environments and the result 85 86 will influence the approaches used to measure and manipulate the underlying physiological 87 mechanisms to improve FE gain.

We therefore hypothesized that, despite being raised in different environments, chickens of equal RFI would be characterized by similar RFI-related profiles for performance and serum parameters. The first objective of this study were to investigate differences in growth performance, FE, serum intermediary metabolites, acute-phase-proteins and white blood cells in low, medium and high residual feed intake (RFI) chickens. The second objective was to assess if the environment in which chickens were raised affect chicken's FE and FE-related performance and serum profiles.

95

#### 96 Materials and Methods

## 97 Experimental design and chickens

Two chicken experiments using common protocols comprising the experimental setup, diet
formulation, data and sample collection were conducted at the Institute of Animal Nutrition
and Functional Plant Compounds [University of Veterinary Medicine Vienna, Austria;
location 1 (L1)] and at the Agriculture Branch of Agri-Food and Biosciences Institute
[Hillsborough, Northern Ireland, United Kingdom; location 2 (L2)] using a completely

randomized study design. At both locations three replicate batches were performed using day-103 104 old mixed-sex Cobb 500FF chicks, resulting in a total population of 78 females and 79 males at L1 and in a total population of 96 females and 96 males at L2. Within each replicate batch, 105 equal numbers of females and males, except for batch 2 with one more male at L1, were used. 106 Due to the geographic distance, chickens came from different commercial hatcheries. The 107 three chicken batches at each location were run in parallel. All animal experimentation 108 procedures were approved by the institutional ethics committee at the University of 109 Veterinary Medicine Vienna and the Austrian national authority according to paragraph 26 of 110 Law for Animal Experiments, Tierversuchsgesetz 2012 - TVG 2012 (GZ 68.205/0131-111 112 II/3b/2013). At Agri-Food and Biosciences Institute the animal procedures were conducted 113 under the project licence number PPL 2781 obtained from the Department of Health, Social Services and Public Safety (DHSSPS) which adhere to the Animals (Scientific Procedures) 114 Act 1986. 115

At hatch, chicks were sexed and transported to L1 and L2 within the first day of life, where 116 chicks were weighed and group-housed. From d 7 of life, chickens were separated and 117 individually housed in cages until the end of the experimental period. The cage floors were 118 made of wire mesh (10 mm  $\times$  10 mm) and padded with rubber tubing. The chickens received 119 a light-to-dark ratio of 23:1h on the day of arrival which was gradually decreased to 18:6h on 120 d 6 of life and was maintained throughout the experimental period. The temperature was 121 maintained at 33°C for the first 5 days after which it was gradually decreased to a temperature 122 of 21°C on d 21 of life. Each cage was equipped with one manual feeder and one drinker and 123 124 feed and demineralized water were freely available.

125

### 126 Diets and Data Collection

127 Chickens were fed starter, grower and finisher diets based on corn and soybean meal (Table1) from d 1 to 10, d 11 to 21, and d 22 to 42 of life, respectively. Diets did not contain anti-

microbial growth promoters or coccidiostats. Starter, grower and finisher diets were mixed 129 according to the same diet formulation at each location. At each location, starter, grower and 130 finisher diets for the replicate batches came from the same batch of commercially prepared 131 crumbles (starter diet) and pellets (3 mm; grower and finisher diets) and were stored in cool 132 (<15°C) and dry conditions for a duration of no longer than 6 months. Feed intake (FI) was 133 determined weekly. Feed leftovers and spills were collected before recording feed intake on d 134 14, 21, 28, 35, 36 and 38 of life. Once a week (upon arrival, d 7, 14, 21, 28 and 35) and on the 135 selection day, BW of all chickens were recorded at both locations. 136

137

## 138 Selection procedure and calculation of FE

Due to the fact that chickens at L1 grew faster than chickens at L2, selection of chickens at L1 took place two days earlier on d 36 of life in order to achieve approximately similar BW at sacrifice and hence to minimize the effect of BW and body composition on parameters of interest. Chickens at L2 were weighed and ranked according to their RFI value on d 38 of life. The RFI was calculated for each chicken for the test interval between d 7 and d 36 of life at L1 and between d 7 and d 38 of life at L2, respectively. Data for net total FI (TFI), metabolic

145 mid-test metabolic weight and total BW gain (TBWG) were used to estimate RFI and residual

146 BW gain (RBG) values as the residuals over the test interval with a nonlinear mixed model in

147 SAS (SAS Stat Inc., version 9.2; Cary; NC) as described in Metzler-Zebeli *et al.* (2016):

148 The MMW was calculated as:

149 MMW =  $[(BW at d 7 of life (g) + BW at d 35 of life (g)) / 2]^{0.75}$ .

150 The RFI and RBG were calculated as:

151  $\operatorname{RFI}(g) = \operatorname{TFI} - (a_1 + b_1 \times \operatorname{MMW} + b_2 \times \operatorname{TBWG}),$ 

152 where a<sub>1</sub> is the intercept and b<sub>1</sub> and b<sub>2</sub> are partial regression coefficients of MMW and TBWG

153 on TFI, respectively. In addition, RBG, residual intake over gain (RIG) and feed conversion

154 ratio (FCR) for the test interval were calculated for the selected chickens:

- 155 RBG (g) = TBWG  $(a_2 + b_3 \times MMW + b_4 \times TFI)$ ,
- where a<sub>2</sub> is the intercept and b<sub>3</sub> and b<sub>4</sub> are partial regression coefficients of MMW and TFI on
  TBWG, respectively.
- 158 The RIG was calculated as:
- 159 RIG(g) = RBG(g) RFI(g).
- 160 The FCR was calculated as:
- 161 FCR (g/g) = TFI (g) / TBWG (g).

162 In each replicate, batch and location it was aimed to select the three chickens with the

lowest RFI (Low RFI), the three chickens with the highest RFI (High RFI), and the three

- 164 chickens with the medium RFI (Medium RFI; a RFI value close to zero), separately for
- 165 female and male chickens. Finally, at location 1, each RFI group was represented by 9
- 166 females and 9 males. At location 2, 6 low RFI, 11 medium RFI and 6 high RFI female
- 167 chickens and 10 low RFI, 9 medium RFI and 9 high RFI male chickens were selected. Only
- 168 the data of the selected chickens at both locations were used for the comparison of FI, growth
- 169 performance and FE. Moreover, blood samples were only collected from the selected
- 170 chickens. The remaining chickens were removed from the experiment. TFI and TBWG were

171 compared for the test interval from d 7 to 36 of life across locations.

172

## **Blood sampling**

Body weight of selected chickens was recorded before chickens were humanely killed for
blood sampling from d 37 to 42 of life. At L1, selected chickens were euthanized with an
overdose of sodium pentobarbital (450 mg/kg, Release, WTD-Wirtschaftsgenossenschaft
Deutscher Tierärzte, Garbsen, Germany) by i.v. injection into the caudal tibial vein from d 37
of life with three to six chickens per day, whereas at L2 selected chickens were sacrificed on d
41 and 42 of life. Immediately thereafter, blood from the vena jugularis at L1 and the heart at
L2 was collected into serum collection tubes (Sarstedt, Nürnbrecht, Germany) and placed on

ice until centrifugation (1 811 × g for 10 min and 1 500 × g at 4°C for 10 min at L1 and L2, respectively; Eppendorf Centrifuge 5810 R, Eppendorf, Hamburg, Germany), and stored at -20°C until analysis. At L1, 1 mL blood was additionally collected in tubes containing EDTA as anticoagulant (Sarstedt, Nürnbrecht, Germany) from which blood smears were prepared on glass slides (n = 4/chicken) to count white blood cells. The intestinal mucosa was checked for *Eimeria*-related lesions at the necropsy which could not be detected.

187

### 188 Chemical analysis and calculations

Proximate nutrient analysis of diet samples was performed according to standard protocols
(Naumann and Basler, 2012). Dry matter was determined after oven-drying for 4 h at 103°C
(method 3.1), crude ash by overnight incineration at 550°C (method 8.1), and CP (nitrogen ×
6.25) by the Kjeldahl method (method 4.1.1; Naumann and Basler, 2012). Diet samples were
further analyzed for EE (method 5.1.1B), CF (method 6.1.1), total starch (method 7.2.1),
sugar (method 7.1.1), calcium (method 10.3.2) and phosphorus (method 10.6.1; Naumann and

195 Basler, 2012).

196

## 197 Blood leukocyte counts, serum metabolites, and acute-phase proteins

198 Blood smears were stained using the May-Grünwald-Giemsa stain (Hemacolor Rapid staining of blood smear kit; Merck KGaA, Darmstadt, Germany). A total of 100 leukocytes, including 199 granular (heterophils, eosinophils, and basophils) and nongranular (lymphocytes and 200 201 monocytes), were counted per slide using light microscopy (Leitz Orthoplan, Leitz, Wetzlar, Germany) at 100-times magnification, and the H-to-L ratio was calculated (Gross and Siegel, 202 1983). Serum glucose, uric acid, triglycerides, cholesterol and NEFA were determined by 203 standard enzymatic colorimetric analysis using an autoanalyzer for clinical chemistry (Cobas 204 6000/c501; Roche Diagnostics GmbH, Vienna, Austria). Chicken specific commercial ELISA 205 kits were used to determine the APPs ovotransferrin (OVT; Cusabio, Wuhan, China) and 206

207 alpha-1-acid glycoprotein (AGP; Genway Biotech Inc., San Diego, CA, US) in serum

according to the manufacturers' instructions. Samples were diluted 2 to 5-fold for both assays

209 depending on the individual sample concentration. The intra- and interassay variability for the

210 OVT and AGP kits were less than 10%, respectively, and the detection limit was 0.039 ng/ml

- and 3.125 ng/ml. All serum parameters were analyzed together at L1.
- 212

#### 213 Statistical analysis

Feed efficiency, FI, growth performance and serum parameters from the selected low,

215 medium and high RFI chickens (location 1: n = 9 low, medium and high RFI female and male

chickens; location 2: n = 6 low RFI, n = 11 medium RFI and n = 8 high RFI females, and n = 11

217 10 low RFI, n = 9 medium RFI and n = 9 high RFI males) were first analysed for normality

using Shapiro-Wilk test with the PROC UNIVARIATE in SAS. The Cook's distance (Cook'sD) test was used to determine any influential observation on the model. Parameters of

individual RFI, performance, and serum metabolites, APPs and white blood cells were

analysed by ANOVA using the PROC MIXED in SAS. Two models were run. The first

accounted for the fixed effects of sex, batch, location and RFI. Because chickens were

sacrificed at different days of life and in order to consider that chickens were consecutively

sampled, the first model included the random effect of chicken nested within day of life and

chicken order at sacrifice. The effects of bird age and BW at 7 days of life were also

separately tested as covariates in the model. As both covariate effects showed no significant

influence on any response variable evaluated, these covariates were removed from the final

228 model and not accounted for in the further analyses. However, sex and batch as fixed effects

229 were found to be significant for many parameters. Therefore, data of female and male

chickens were analysed separately using a second model which was fitted to take into account

the fixed effects of RFI and location and their two-way-interaction. The random effect

232 considered the chicken nested within batch, day of life and chicken order at sacrifice. Since

white blood cell counts were only determined at L1, only the fixed effect of RFI was 233 234 considered. Moreover, in the second model, orthogonal polynomial contrast statement was used to evaluate linear relationships. Degrees of freedom were approximated by the method of 235 Kenward-Roger. Least squares means were computed and significance declared at  $P \le 0.05$ . 236 A trend was considered at  $0.05 < P \le 0.10$ . 237 In order to investigate whether sex-independent relationships between chicken's 238 individual RFI and serum metabolites existed linear discriminate analysis (LDA) and 239 regression analysis were applied. The LDA was performed using JMP10 software (SAS Stat 240 Inc.) with serum metabolites (glucose, urea, cholesterol, triglycerides and NEFA) as 241 242 covariates and RFI group as the categorical variable. The LDA results were visualised using 243 the first 2 principal components of the scores plot to identify characteristic trends or grouping among chickens of diverging RFI. Moreover, regression analysis (PROC REG of SAS) was 244 used to establish and quantify the relationships between individual serum metabolites, serum 245 APPs and blood H-to-L ratio and chickens' individual RFI values, irrespective of sex and 246 247 location. For this, mixed modelling (PROC MIXED of SAS) of each serum metabolite was performed including the fixed effects RFI, sex and location. The slope and intercept by RFI, 248 sex and location were included as random effects and the variance component structure was 249

- used as variance-covariance matrix. Significant relationships (P < 0.05) are shown in Fig. 1.
- 251

#### 252 **Results**

## 253 Chicken performance and feed efficiency

Sex did not affect BW on d 7 of life, whereas male chickens weighed approximately 300 g more on d 36 of life than females (P < 0.001; Table 1 and 2). Similarly, TFI and TBWG were higher (P < 0.001) in males compared to females. Location affected BW on d 7 and 36 of life. While female and male chickens weighed about 10 g more on d 7 of life at L2 compared to L1, they gained about 350 to 400 g less by d 36 at L2 compared to L1 (P < 0.001). In contrast, TFI between d 7 and 36 of life was not influenced by location. Likewise, location did not
affect the FE metrics RFI, RBG and RIG; providing similar values for female and male
chickens of the same RFI group, whereas FCR was about 12 % lower (P < 0.001) in chickens</li>
of L1 compared to chickens of L2.

The RFI ranged on average from -231 to 215 g in females and from -197 to 267 g in males 263 representing a difference of 330 and 500 g TFI between most and least efficient female and 264 male chickens (P < 0.001; Table 1 and 2). Body weight at d 36 and TBWG were similar 265 among chickens of diverging RFI. Likewise, the RBG of the selected chickens was similar 266 among the three RFI groups, whereas the RIG linearly decreased in the same range observed 267 268 for the increase in RFI from low to high RFI chickens, irrespective of sex. The FCR linearly increased from low to high RFI by on average 13% (P < 0.001). There was a sex effect and 269 location effect for FCR showing a 0.06 g/g-lower FCR in males compared to females as well 270 as a 0.19 g/g lower FCR in chickens at L1 compared to chickens at L2 (P < 0.001). 271

At sacrifice, male chickens at both locations had similar BW across locations (3.03 and 3.02 ± 0.062 kg at L1 and L2, respectively; P = 0.859) and RFI groups (3.04, 2.96 and 3.08 ± 0.076 kg for low, medium and high RFI, respectively; P = 0.535). By contrast, BW in female chickens at sacrifice differed across locations with females at L1 weighing about 270 g more than females at L2 (2.85 versus 2.58 ± 0.063 kg at L1 versus L2, respectively; P = 0.001), but their BW was not different among RFI groups (2.72, 2.71 and 2.73 ± 0.065 for low, medium and high RFI, respectively; P = 0.974).

279

## 280 Serum metabolite profiles and acute phase proteins

Results for serum metabolite profiles and acute-phase-proteins examined for female and male chickens are presented in Table 3 and 4, respectively. There was a location effect for serum OVT in females showing that chickens at L1 had a 2-fold higher serum OVT concentration than that of chickens at L2 (P < 0.05). Moreover, we observed a linear increase (P < 0.05) in

serum AGP from low to high RFI in male chickens at L1 but not at L2. Sex affected (P < 285 286 0.05) serum NEFA concentrations which were higher in males. Female chickens at L1 had a lower serum glucose and NEFA and higher serum urea and cholesterol than females at L2 (P 287 < 0.05). In males, serum glucose and triglycerides were lower at L1 compared to L2 (P <288 0.01). Despite differences in actual serum concentrations, FE-effects for glucose, uric acid 289 and cholesterol among RFI groups were similar at both locations in females. There was a 290 linear increase in serum uric acid (P < 0.05), and a tendency for a linear increase in serum 291 cholesterol and triglycerides (P < 0.1) from low to high RFI in female chickens. Serum NEFA 292 showed a FE  $\times$  location effect (P < 0.01) by increasing by 57% from low to high RFI at L2 293 294 but not in females at L1. Similar to the females, serum cholesterol linearly increased (P < 0.05) and triglycerides tended (P < 0.1) to increase by about 17 and 31% from low to high 295 RFI in male chickens, respectively. 296

297

### 298 White blood cell counts

White blood cell counts were only determined at L1 (Table 5). Females and males differed in their white blood cell counts with females having more lymphocytes but less monocytes and heterophils than males (P < 0.05). In females, FE tended to affect only monocyte counts with chickens of low RFI having less monocytes than chickens of medium and high RFI. In males, lymphocyte counts linearly decreased (P = 0.012) from low to high RFI, whereas heterophils linearly increased from low to high RFI (P = 0.031). Because of this, there was a linear (P =0.027) increase in the H-to-L ratio of 68% from low to high RFI in males.

306

## **307** Multivariate and regression analysis

308 The LDA plot of RFI groups and serum metabolites showed separate clustering for serum

309 metabolites for low and high RFI, whereas the 95% confidence intervals of medium RFI

overlapped with those of low and high RFI (Figure 1A). Serum glucose discriminated best for

low RFI, whereas serum triglycerides, uric acid and cholesterol correlated with high RFI. 311 312 When comparing locations (Figure 1B), the LDA showed clear clustering of serum metabolites between L1 and L2, whereby serum NEFA correlated to L2 and urea to L1. Due 313 to the separate clustering in the LDA together with trends for linear relationships between 314 some serum metabolites and RFI groups, relationships between serum parameters and the 315 individual RFI values of chickens from both sexes and locations were regressed. Regression 316 analysis showed positive relationships between serum cholesterol and RFI ( $R^2 = 0.13$ ; P < 0.13) 317 0.001; Figure 2A) and serum uric acid and RFI ( $R^2 = 0.49$ ; P < 0.001; Figure 2B). There was 318 also a weak positive relationship between the H-to-L ratio and RFI values for chickens at L1 319  $(R^2 = 0.15; P = 0.003;$  Figure 2C). 320

321

## 322 Discussion

Our understanding of the physiological mechanisms underlying the FE of chicken's is steadily 323 advancing (e.g., Aggrey et al., 2014; Lee et al., 2015; Zhou et al., 2015). However, the 324 325 contribution of the rearing environment has not yet been sufficiently elucidated. In the current study, chickens from one hybrid line were raised using similar management protocols at two 326 distinct geographic locations to investigate if RFI-related performance traits and serum 327 profiles are affected by the rearing environment. Similar to Stanley et al. (2016), the present 328 chicken populations met or exceeded the expected average growth rate, and the range in TFI, 329 growth, and FE data recorded was consistent with previous studies in chickens selected for 330 RFI (e.g., Zhuo et al., 2015). Although the TFI from d 7 to 36 of life was similar across 331 locations, results indicated a marked location effect on TBWG of chickens between locations 332 333 which was apparent throughout all replicate batches and for both sexes. Furthermore, we could distinguish RFI-related profiles for certain serum intermediary metabolites, but not 334 acute-phase-proteins, in the current chicken populations, whereby RFI-effects were different 335 in males and females. The regression models implemented established linear relationships 336

between RFI and serum uric acid and cholesterol, suggesting them as predictors for RFI in the 337 338 current chicken populations irrespective of sex and location. Despite these relationships and clear clustering between low and high RFI in the LDA plots, the actual concentrations of 339 serum metabolites were location-specific which may render it difficult to predict universal 340 serum threshold values for low, medium and high RFI chickens. Moreover, as present 341 relationships between RFI and serum cholesterol and uric acid were weak to moderate, it may 342 be advisable to use serum metabolite patterns rather than individual metabolites to predict the 343 RFI in chickens. 344

Chicken RFI values were similar across locations, but it should be considered that 345 346 chicken's RFI values were determined two-days apart. The RFI is phenotypically independent of BW and level of production (e.g., ADG; Bottje and Carstens, 2009), and may have 347 therefore remained similar across locations in the current study despite differences in TBWG 348 and ADG. Similar observations were made for RBG and the combined metric RIG of the 349 selected chickens. Inconsistent findings exist in the literature for RFI-related differences in 350 351 BW and BW gain in low and high RFI chickens (van Eerden et al., 2004; Zhuo et al., 2015). Irrespective of location, chickens of diverging RFI could not be distinguished based on their 352 BW or TBWG. In contrast to some studies with short measurement periods of only one week 353 354 (e.g., Zhuo *et al.*, 2015), we determined the FE over a period of 29 and 31 days at L1 and L2, respectively. It is highly likely that this improved the accuracy of RFI prediction in the 355 present study as we observed slight differences in the FE and grouping of chickens according 356 to their RFI when assessed only on a weekly basis. Differences in TFI between low and high 357 RFI chickens were considerable and were already present at 21 days of life (Supplemental 358 Table 1). Notably, irrespective of the two-day difference in selection for RFI, location effects 359 were distinguishable when using the ratio metric FCR. This leads to the assumption that the 360 FCR may more accurately predict FE-related differences in growth performance among 361

362 chicken flocks, whereas the RFI may be the FE metric of choice to equally rank chickens363 independent from the environment.

The present environmental effects clearly suggest that physiological differences between 364 low and high RFI chickens may largely vary between farms due to environment-specific 365 factors. Parents' own FE essentially determines development and FE of the chicks post-hatch 366 (Bottje and Carstens, 2009; Romero et al., 2011). This may have been of less relevance in the 367 present study as chickens used in the present trials were not related within or between 368 locations (see Relationship analysis in Supplemental Material). The main environment-369 specific factors were likely the diet, even though it was of the same formulation, the housing 370 371 environment including environmental microbes at the hatcheries and rearing location as well as the personnel handling the chickens. The immediate colonization of chicken's intestine 372 post-hatch with microbes from the egg shell and environment is critical because it has a long 373 374 lasting effect on chicken's performance by influencing the further microbial colonization, intestinal development and priming of the immune system (Brisbin et al., 2008; Schokker et 375 al., 2015). The intestinal microbiota interacts with the host via several routes including 376 microbial metabolites and receptor-recognition pathways (Blaut, 2015). As a result, different 377 bacterial colonization patterns may have caused a more pronounced stimulation of the 378 379 immune system throughout the growing phase at one location which may have decreased the energy available for growth. Also, different bacterial colonization across locations may have 380 led to diverging profiles of intestinally produced short-chain fatty acids which, after being 381 382 absorbed, may have affected lipogenesis of the host and present serum profiles. Especially acetate serves as substrate for *de novo* lipogenesis in the liver, whereas propionate is used for 383 hepatic gluconeogenesis (Blaut, 2015). In general, due to the hygienic standards in modern 384 hatcheries, microbial colonization of the gastrointestinal tract of newly hatched chicks is more 385 influenced by microbes encountered in their wider environment (e.g., personnel, housing, 386 water and diet) than by the normal chicken gut microbiota (Stanley et al., 2013; Ludvigsen et 387

al., 2016). Because current chickens came from different hatcheries, the early microbial 388 389 colonization may have been one of the most influential factors for the variation between both locations. This would be supported by different RFI-associated bacterial microbiome profiles 390 in chickens between the two locations at 6 weeks of life (Siegerstetter et al., 2016). Moreover, 391 although the dietary formulations were the same and concentrations of most nutrients were 392 equal, natural differences in the raw materials, i.e. corn and soybean meal, between locations 393 (e.g. dietary fiber composition; Rodehutscord et al., 2016) may have altered digestive, 394 absorptive and fermentative processes. This probably affected the present results for growth 395 performance and serum metabolite profiles across locations. 396

397 The BW at sacrifice and thus body composition may have also contributed to the variation in serum parameters in female chickens across locations and were likely depicted in chickens' 398 serum metabolite and APP concentrations. Accordingly, serum profiles suggested that 399 400 chickens at L2 had either an increased intestinal glucose release or altered systemic glucose metabolism than those at L1, irrespective of sex. Moreover, differences in BW and thus 401 402 adipose tissue accretion likely led to the variation in serum lipids across locations. Moreover, the increased OVT response in females at L1 compared to L2 may indicate an increased 403 abundance of microbial stressors at L1. As an iron binding protein OVT provides 404 405 antimicrobial properties by sequestering iron and modulates heterophil and macrophage function in chickens (Murata et al., 2004). In spite of the observed location effects, the fact 406 that location × FE interactions were almost absent in our study allows assuming that RFI-407 408 related differences in performance traits and serum profiles were similar across locations. Although influenced by prandial activity, blood metabolites and hormones associated with 409 410 feed intake, growth, nutrient repartitioning and utilization may serve as potential physiological markers for FE in various livestock species (Richardson et al., 2004; Kelly et 411 al., 2010; Montagne et al., 2014; Jegou et al., 2016). Likewise, serum intermediary 412

chicken populations of the present study. Controversial results were previously reported for 414 serum triglycerides, NEFA and uric acid in cockerel lines selected for low and high RFI 415 (Gabbarou et al., 1997; Swennen et al., 2007), whereas, to our awareness, little information 416 exists for broiler chickens of diverging RFI. Although the selection strategy and age of the 417 chickens differed, Gabbarou et al. (1997) found a comparable increase in plasma triglycerides 418 and plasma glucose and uric acid concentrations in cockerels which corresponded to our 419 420 results in male chickens. According to the present linear FE-effects and regression analysis, serum concentrations of uric acid and serum cholesterol might be considered as predictors for 421 RFI in chickens. The higher FI in high RFI chickens should have increased the intestinal 422 423 glucose uptake and postprandial insulin level as well as peak duration. Accordingly, equal 424 serum glucose concentrations may indicate improved energy saving capacity or lower glucose uptake and metabolism of peripheral organs in low versus high RFI chickens (Bottje and 425 426 Carstens, 2009). Some authors (Richardson et al., 2004; Kelly et al., 2010) have proposed a decrease in insulin sensitivity in muscle tissue in energetically inefficient animals. 427 Concurrently, higher basal insulin concentrations in high-RFI animals may be linked to 428 greater fat deposition because insulin reduces lipolysis and stimulates lipogenesis in adipose 429 430 tissue (Kelly et al., 2010; Le Naou et al., 2012; Montagne et al., 2014; Zhuo et al., 2015). 431 Accordingly, Zhuo et al. (2015) showed that abdominal adipose tissue of high RFI chickens had a greater expression of lipid synthesis genes and decreased expression of triglyceride 432 hydrolysis and cholesterol transport genes. Moreover, in their study, low RFI chickens had a 433 potentially more active glucose-to-lipid conversion and different insulin signaling in adipose 434 tissue at transcriptome level compared to high RFI chickens (Zhuo et al., 2015). The latter 435 may explain the elevated postprandial serum triglycerides and cholesterol observed for high 436 RFI males and females compared to their low RFI counterparts in the present study. Varying 437 RFI-related serum profiles in males and females indicated that differences were more 438 pronounced in females than males. Despite not having measured serum insulin levels, 439

elevated serum uric acid and NEFA in high RFI females may confirm our assumption of
reduced insulin sensitivity since both metabolites are typically raised during insulin resistance
due to increased lipolysis and deamination of amino acids for energy provision (e.g., Yuan *et al.*, 2008; Ji *et al.*, 2012). In addition, raised serum uric acid in high RFI animals may also
suggest less efficient nitrogen recycling as recently shown for a different chicken line (Aggrey *et al.*, 2014).

Inconclusive results exist on whether diverging RFI is accompanied by a change in the 446 stress response of meat-type chickens. As part of the physiological stress response via the 447 hypothalamic-pituitary-adrenal axis and sympathetic system, increased systemic levels of 448 449 corticosterone induces a general acute-phase response including OVT and AGP in chickens (O'Reilly and Eckersall, 2014; Zulkifli et al., 2014). Moreover, increased corticosterone 450 levels were associated with modified insulin sensitivity, reduced muscle protein accretion and 451 452 raised plasma lipids and uric acid in chickens (Dong et al., 2007; Yuan et al., 2008) which may have contributed to RFI-related metabolic alterations and serum metabolite profiles. 453 454 Present results for RFI-related differences in serum APPs were not, however, conclusive and only indicated a linear relationship between AGP and RFI in males at L1. Similar to AGP, the 455 H-to-L ratio showed the same RFI-related pattern in males at L1 only. AGP has an 456 457 immunoregulatory function by influencing T-cell function and thus white blood cell production (Murata et al., 2004). Since males and females were evenly distributed across the 458 experimental room for all three batches at L1, a greater immune response due to infectious 459 460 disease agents may be excluded as an explanation for the gender difference seen here. The question then arises as to whether the high RFI males at L1 showed a greater excitability or 461 aggressiveness compared to the female chickens. Despite the weak linear relationship 462 between RFI and serum H-to-L, its reliability to predict chicken's RFI should be evaluated in 463 further experiments since only data from L1 were available for regression analysis in the 464 present study. 465

In conclusion, the results of the present study demonstrate that chickens reared at two 466 geographically distinct locations showed similar RFI-related variation in serum intermediary 467 metabolites. Regression analysis confirmed the usefulness of serum metabolite patterns as 468 RFI predictors for the current chicken populations. Due to the environment-specific 469 differences observed here, further research is warranted to validate the reliability of serum 470 metabolites, such as uric acid and cholesterol, as RFI predictors in chickens. 471 472 Acknowledgements 473 This project (ECO-FCE) has received funding from the European Union's Seventh 474 475 Framework Programme for research, technological development and demonstration (grant no. 311794). The technical staff at the Institute of Animal Nutrition and Functional Plant 476 Compounds (University of Veterinary Medicine Vienna) and at the Agri-Food and 477 Biosciences Institute are gratefully thanked for their care of the animals and expertise when 478 conducting the experiment and for laboratory assistance. 479 480 **Disclosure statement** 481 The authors state no conflict of interest. 482 483 References 484 AGGREY, S.E., LEE, J., KARNUAH, A.B. & REKAYA, R. (2014) Transcriptomic analysis 485 of genes in the nitrogen recycling pathway of meat-type chickens divergently selected for feed 486 efficiency. Animal Genetics 45: 215-222. doi:10.1111/age.12098 487 BERRY, D.P. & CROWLEY, J.J. (2012) Residual intake and body weight gain: A new 488 measure of efficiency in growing cattle. Journal of Animal Science 90:109-115. 489

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		Residual feed intake (RFI) <sup>1,2</sup>				<i>P</i> -value <sup>3,4</sup>		
Item	Location	Low	Medium	High	SEM	FE	location	FE × location
BW, d 7 of life (g)	L1+2	145	145	147	2.6	0.805	0.001	0.802
	L1	141	138 <sup>y</sup>	141 <sup>y</sup>	3.6			
	L2	149	151 <sup>x</sup>	153 <sup>x</sup>	2.6			
BW, d 36 of life (g)	L1+2	2253	2187	2215	50.4	0.654	< 0.001	0.670
	L1	2392 <sup>x</sup>	2359 <sup>x</sup>	2420 <sup>x</sup>	68.9			
	L2	2115 <sup>y</sup>	2015 <sup>y</sup>	2009 <sup>y</sup>	73.2			
Total feed intake, d 7-36 of life (g)	L1+2	3447 <sup>b</sup>	3485 <sup>ab</sup>	3774 <sup>a</sup>	91.1	0.027*	0.479	0.566
	L1	3334 <sup>b</sup>	3510 <sup>ab</sup>	3751 <sup>a</sup>	123.2			
	L2	3559	3461 <sup>B</sup>	3797 <sup>A</sup>	131.0			
Total body weight gain, d 7-36 of life (g)	L1+2	2108	2042	2068	49.5	0.647	< 0.001	0.643
	L1	2251 <sup>x</sup>	2220 <sup>x</sup>	2279 <sup>x</sup>	67.7			
	L2	1966 <sup>y</sup>	1865 <sup>y</sup>	1856 <sup>y</sup>	72.0			
RFI (g)	L1+2	-231	8	215	20.1	< 0.001***	0.412	0.201
	L1	-195	18	197	27.5			
	L2	-267	-3	232	29.2			
RBG (g)	L1+2	-0.9	1.0	1.7	4.13	0.901	0.775	0.993
	L1	-2.1	0.5	1.3	5.65			
	L2	0.2	1.5	2.1	6.01			
RIG (g)	L1+2	230	-7	-213	20.2	<0.001***	0.380	0.195
	L1	193	-18	-196	27.6			
	L2	267	4	-231	29.3			

**Table 1.** Feed intake, growth performance and feed efficiency metrics in female broiler chickens raised at two different locations.

FCR (g/g)	L1+2	1.55	1.63	1.76	0.019	<0.001***	< 0.001	0.108
	L1	1.46 <sup>y</sup>	1.55 <sup>y</sup>	1.62 <sup>y</sup>	0.026			
	L2	1.65 <sup>x</sup>	1.71 <sup>x</sup>	1.89 <sup>x</sup>	0.028			

- 606 FE, feed efficiency; FCR, feed conversion ratio; RBG, residual BW gain; RIG, residual intake over gain; L1, University of Veterinary Medicine Vienna (Vienna, Austria); L2,
- 607 Agri-Food and Biosciences Institute (Hillsborough, Northern Ireland, UK).
- 608 <sup>1</sup>Values are least squares means  $\pm$  standard error of the mean (SEM).
- <sup>2</sup>Each RFI group represents n = 9 female chickens at location 1; n = 6 low RFI, n = 11 medium RFI and n = 8 high RFI females at location 2.
- 610  ${}^{3}P$ : probability level.
- 611 <sup>4</sup>Linear polynomial contrast:  $*P \le 0.05$ , and  $***P \le 0.001$ .
- 612 a-cLeast squares means within a row without a common lowercase superscript differ among RFI groups (P < 0.05).
- 613 <sup>A,B</sup>Least squares means within a row without a common uppercase superscript tend to differ among RFI groups (P < 0.1).
- 614 <sup>x,y</sup>Least squares means within a column without a common lowercase superscript differ between locations (P < 0.05).

		Residual feed intake (RFI) <sup>1,2</sup>			_	<i>P</i> -value <sup>3,4</sup>		
Item	Location	Low	Medium	High	SEM	FE	location	FE × location
BW, d 7 of life (g)	L1+2	145	145	148	2.2	0.704	< 0.001	0.919
	L1	139 <sup>x</sup>	140 <sup>x</sup>	141 <sup>x</sup>	3.0			
	L2	152 <sup>y</sup>	150 <sup>y</sup>	154 <sup>y</sup>	3.0			
BW, d 36 of life (g)	L1+2	2562	2483	2546	55.4	0.577	< 0.001	0.560
	L1	2712 <sup>x</sup>	2733 <sup>x</sup>	2756 <sup>x</sup>	79.0			
	L2	2380 <sup>y</sup>	2233 <sup>y</sup>	2367 <sup>у</sup>	79.4			
Total feed intake, d 7-36 of life (g)	L1+2	3753 <sup>b</sup>	3879 <sup>b</sup>	4253ª	70.1	<0.001***	0.340	0.573
	L1	3682 <sup>b</sup>	3901 <sup>b</sup>	4185 <sup>a</sup>	99.9			
	L2	3823 <sup>b</sup>	3857 <sup>b</sup>	4321ª	98.2			
Total body weight gain, d 7-36 of life (g)	L1+2	2401	2338	2414	54.5	0.582	< 0.001	0.560
	L1	2573	2593 <sup>x</sup>	2615 <sup>x</sup>	77.7			
	L2	2228	2083 <sup>y</sup>	2214 <sup>y</sup>	76.4			
RFI	L1+2	-197	0	267	21.8	<0.001***	0.149	0.610
	L1	-183	6	303	31.1			
	L2	-211	-6	231	30.6			
RBG	L1+2	5.5	-1.1	3.8	4.40	0.550	0.166	0.687
	L1	6.8	1.8	10.4	6.27			
	L2	4.2	-3.9	-2.7	6.16			
RIG	L1+2	202	-1.	-263	22.0	<0.001***	0.247	0.699
	L1	190	-4	-292	31.3			
	L2	215	2	-234	30.8			

**Table 2**. Feed intake, growth performance and feed efficiency metrics in male broiler chickens raised at two different locations.

	FCR	L1+2	1.50	1.58	1.70	0.019	<0.001***	< 0.001	0.774
		L1	1.41 <sup>y</sup>	1.48 <sup>y</sup>	1.61 <sup>y</sup>	0.028			
		L2	1.58 <sup>x</sup>	1.69 <sup>x</sup>	1.79 <sup>x</sup>	0.027			
617	FE, feed efficiency; RBG, residual BW gain; RIG, re	sidual intake ove	er gain; L1, Uni	versity of Veteri	nary Medicine	Vienna (Vier	nna, Austria); L2	, Agri-Food and	d Biosciences
618	Institute (Hillsborough, Northern Ireland, UK).								
619	<sup>1</sup> Values are least squares means $\pm$ standard error of the squares means $\pm$ standar	ne mean (SEM).							
620	<sup>2</sup> Each RFI group represents $n = 9$ male chickens at lo	cation 1; $n = 10$	low RFI, $n = 9$	medium RFI and	l n = 9 high RF	I males at loc	ation 2.		
621	<sup>3</sup> P: probability level.								
622	<sup>4</sup> Linear polynominal contrast: *** $P \le 0.001$ .								
623	<sup>a-c</sup> Least squares means within a row without a comm	on lowercase sup	erscript differ a	mong RFI group	P < 0.05).				
624	x,yLeast squares means within a column without a con	nmon lowercase	superscript diff	er between locat	tions ( $P < 0.05$ )	).			

		R	esidual feed inta	ke <sup>1,2</sup>	_	<i>P</i> -value <sup>3,4</sup>			
Parameter	Location	Low	Medium	High	SEM	FE	location	FE × location	
Glucose (mg/dl)	L1+2	304	283	310	16.8	0.450	0.002	0.920	
	L1	268 <sup>x</sup>	256 <sup>x</sup>	276 <sup>x</sup>	23.0				
	L2	340 <sup>Y</sup>	310 <sup>Y</sup>	344 <sup>y</sup>	24.4				
Urea (mg/dl)	L1+2	2.27 <sup>b</sup>	2.42 <sup>ab</sup>	2.83 <sup>a</sup>	0.182	0.101*	0.005	0.701	
	L1	2.46 <sup>b</sup>	2.76 <sup>abx</sup>	3.25 <sup>ax</sup>	0.248				
	L2	2.08	2.08 <sup>y</sup>	2.41 <sup>y</sup>	0.264				
Cholesterol (mg/dl)	L1+2	132	138	145	5.1	0.244†	0.002	0.628	
	L1	139	152 <sup>x</sup>	154	7.0				
	L2	125	125 <sup>Y</sup>	135	7.4				
Triglycerides (mg/dl)	L1+2	93 <sup>B</sup>	101	126 <sup>A</sup>	11.8	0.135†	0.802	0.882	
	L1	86 <sup>B</sup>	103	126 <sup>A</sup>	16.2				
	L2	99	99	127	17.2				
NEFA (µmol/l)	L1+2	204	241	269	11.6	0.002***	< 0.001	0.008	
	L1	199	214 <sup>y</sup>	208 <sup>y</sup>	15.8				
	L2	210°	269 <sup>bx</sup>	330 <sup>ax</sup>	16.8				
Ovotransferrin (µg/ml)	L1+2	13.2	10.8	14.1	0.34	0.761	0.031	0.226	
	L1	17.8	11.1 <sup>B</sup>	22.1 <sup>Ax</sup>	4.59				
	L2	8.5	10.6	6.0 <sup>y</sup>	4.83				
Alpha-1-acid glycoprotein (µg/ml)	L1+2	221.1	204.7	209.6	13.13	0.686	0.139	0.342	
	L1	240.7	223.8	205.5	18.04				
	L2	201.5	185.7	213.8	18.99				

# **Table 3.** Serum metabolites and acute-phase-proteins in female broiler chickens raised at two different locations.

- 627 FE, feed efficiency; L1 University of Veterinary Medicine Vienna (Vienna, Austria); L2, Agri-Food and Biosciences Institute (Hillsborough, Northern Ireland, UK).
- 628 <sup>1</sup>Values are least squares means  $\pm$  standard error of the mean (SEM).
- 629 <sup>2</sup>Each RFI group represents n = 9 female chickens at location 1; n = 6 low RFI, n = 11 medium RFI and n = 8 high RFI females at location 2.
- 630  ${}^{3}P$ : probability level.
- 631 <sup>4</sup>Linear polynomial contrast:  $*P \le 0.05$ ,  $***P \le 0.001$ , and  $\dagger P \le 0.10$ .
- 632 <sup>a-c</sup>Least squares means within a row without a common lowercase superscript differ among RFI groups (P < 0.05).
- 633 <sup>A,B</sup>Least squares means within a row without a common uppercase superscript tend to differ among RFI groups (P < 0.1).
- 634 <sup>x,y</sup>Least squares means within a column without a common lowercase superscript differ between locations (P < 0.05).
- 635 X,Y Least squares means within a column without a common uppercase superscript tend to differ between locations (P < 0.1).
- 636

		R	esidual feed inta	ke <sup>1,2</sup>	_	<i>P</i> -value <sup>3,4</sup>			
Parameter	Location	Low	Medium	High	SEM	FE	location	FE × location	
Glucose (mg/dl)	L1+2	295	312	317	15.9	0.585	< 0.001	0.377	
	L1	270	257 <sup>x</sup>	272 <sup>x</sup>	23.1				
	L2	320	368 <sup>y</sup>	362 <sup>y</sup>	21.8				
Urea (mg/dl)	L1+2	2.30	2.38	2.66	0.194	0.406	0.126	0.665	
	L1	2.34	2.61	2.93	0.283				
	L2	2.27	2.16	2.40	0.267				
Cholesterol (mg/dl)	L1+2	134 <sup>A</sup>	142 <sup>A</sup>	157 <sup>B</sup>	5.2	0.010**	0.133	0.453	
	L1	143	142	162	7.5				
	L2	125ª	142 <sup>ab</sup>	153 <sup>b</sup>	7.2				
Triglycerides (mg/dl)	L1+2	91 <sup>B</sup>	102	119 <sup>A</sup>	11.9	0.248†	0.001	0.226	
	L1	84	71 <sup>x</sup>	86 <sup>x</sup>	17.3				
	L2	98	133 <sup>y</sup>	153 <sup>y</sup>	16.3				
NEFA (µmol/l)	L1+2	253	295	293	25.9	0.429	0.354	0.126	
	L1	244	318	238 <sup>x</sup>	37.7				
	L2	262 <sup>B</sup>	273	348 <sup>yA</sup>	35.6				
Ovotransferrin (µg/ml)	L1+2	7.61	11.86	13.24	3.06	0.394	0.743	0.904	
	L1	7.79	11.71	14.96	4.22				
	L2	7.43	12.01	11.52	4.33				
Alpha-1-acid glycoprotein (µg/ml)	L1+2	202.1	227.1	235.0	16.59	0.338	0.164	0.246	
	L1	195.3 <sup>b</sup>	241.7 <sup>ab</sup>	$267.9^{aX}$	24.46				
	L2	208.8	212.5	202.1 <sup>Y</sup>	23.50				

# **Table 4.** Serum metabolites and acute-phase-proteins in male broiler chickens raised at two different locations.

- 638 FE, feed efficiency; L1, University of Veterinary Medicine Vienna (Vienna, Austria); L2, Agri-Food and Biosciences Institute (Hillsborough, Northern Ireland, UK).
- 639 <sup>1</sup>Values are least squares means  $\pm$  standard error of the mean (SEM).
- 640 <sup>2</sup>Each RFI group represents n = 9 male chickens at location 1; n = 10 low RFI, n = 9 medium RFI, and n = 9 high RFI males at location 2.
- 641  ${}^{3}P$ : probability level.
- 642 <sup>4</sup>Linear polynomial contrast contrast: \*\* $P \le 0.01$ , and  $\dagger P \le 0.10$ .
- 643 <sup>a-c</sup>Least squares means within a row without a common lowercase superscript differ among RFI groups (P < 0.05).
- 644  $^{A,B}$ Least squares means within a row without a common uppercase superscript tend to differ among RFI groups (P < 0.1).
- 645 <sup>x,y</sup>Least squares means within a column without a common lowercase superscript differ between locations (P < 0.05).
- 646 X,YLeast squares means within a column without a common uppercase superscript tend to differ between locations (P < 0.1).

647	Table 5. White blood cells in female and male broiler chickens raised at location 1.
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	I	Residual feed intake <sup>1,2</sup>			
Parameter	Low	Medium	High	SEM	FE, <i>P</i> -value <sup>3</sup>
Females					
Lymphocytes (%)	86.3	83.8	84.9	1.36	0.465
Heterophils (%)	12.1	13.4	13.2	1.25	0.730
Basophils (%)	0.07	0.17	0.03	0.05	0.160
Monocytes (%)	1.57	2.52	1.92	0.27	0.064
H-to-L proportion (%)	14.2	16.3	15.8	1.76	0.680
Males					
Lymphocytes (%)	83.6	82.6	75.3	2.14	0.023*
Heterophils (%)	13.9	15.0	20.6	2.06	0.067*
Basophils (%)	0.14	0.00	0.23	0.08	0.121
Monocytes (%)	2.37	2.32	2.81	0.39	0.629
H-to-L proportion (%)	17.0	18.7	28.5	3.44	0.057*

648 FE, feed efficiency; location 1, University of Veterinary Medicine Vienna (Vienna, Austria).

649 <sup>1</sup>Values are least squares means  $\pm$  standard error of the mean (SEM).

650 <sup>2</sup>Each RFI group represents n = 9 chickens females and males.

651  ${}^{3}P$ : probability level.

- 652 <sup>4</sup>Linear polynominal contrast:  $*P \le 0.05$ ;  $**P \le 0.01$ ,  $***P \le 0.001$ , and  $\dagger P \le 0.10$ .
- 653 <sup>5</sup>Nitrogen × 6.25.

## 654 **Figure captions**

**Figure 1.** a) Linear discriminant analysis of RFI groups and serum metabolites: low RFI

- group ( $\bigcirc$ ), medium RFI group ( $\diamondsuit$ ), and high RFI group ( $\bigcirc$ ). b) Linear discriminant analysis
- of location and serum metabolites: location 1 (Austria (●)), and location 2 (UK (○)). Circles
- 658 indicate 95% confidence intervals.
- 659
- 660 Figure 2. Quantification of relationships between RFI values and serum metabolites in male
- and female chickens from both locations (A-C). Relation between chicken's RFI value (x) and
- serum concentration (y) of cholesterol (A) and serum uric acid (B): linear regression, A) y =
- 663  $140.72 + 0.039 \times x$ , RMSE = 20.652,  $R^2 = 0.13$ , P < 0.001 and B)  $y = 2.34 + 0.00070 \times x$ , root
- 664 mean square error (RMSE) = 0.143,  $R^2 = 0.49$ , P < 0.001. Relation between RFI value (x) and
- blood heterophil-to-lymphocyte proportion in chickens at location 1 (C): linear regression, y =
- 666  $17.98 + 0.018 \times x$ , RMSE = 8.358,  $R^2 = 0.15$ , P = 0.003.
- 667

### 1 Metzler-Zebeli et al. – Supplemental Material

Item	Starter <sup>1</sup>	Grower <sup>2</sup>	Finisher <sup>3</sup>
Ingredient (g/kg as-fed)			
Corn	612	660	679
Soybean meal	331	282	260
Soybean oil	17.5	20.6	27.7
Limestone flour	11.0	9.8	7.0
Salt	2.0	2.0	2.3
Dicalcium phosphate	16.1	15.0	13.4
Vitamin/mineral-premix	11.0	11.0	10.0
Analyzed chemical composition (g/kg DM)	at L1		
Dry matter	926	923	914
Crude protein	243	223	216
Ether extracts	50	52	59
Crude fiber	31	27	28
Crude ash	69	62	55
Starch	462	506	514
Sugar	40	46	49
Calcium	11.9	10.7	8.9
Phosphorus	8.2	7.8	6.9
Analyzed chemical composition (g/kg DM)	at L2		
Dry matter	908	902	902
Crude protein	221	219	209
Crude ash	94	81	72
Metabolizable energy <sup>4</sup> (MJ/kg)	13.7	14.3	14.6

2 **Supplemental Table 1.** Ingredients and chemical composition of diets.

#### 3 <sup>1</sup>Premix provided per kilogram of starter diet: vitamin A as retinyl acetate, 13,000 IU; vitamin D<sub>3</sub> as 4 cholecalciferol, 5,000 IU; vitamin E as alpha-tocopherol-acetate, 80 IU; vitamin K, 3 mg; thiamin, 3 mg; 5 riboflavin, 9 mg; pyridoxine, 4 mg; vitamin B<sub>12</sub> 20 µg; biotin, 0.15 mg; calcium pantothenate, 15 mg; nicotinic 6 acid, 60 mg; folic acid, 2 mg; 500 mg choline chloride; methionine, 3,405 mg; threonine, 745 mg; lysine, 2,812 7 mg; I, 1 mg as calcium iodate; Se, 0.35 mg as sodium selenite; Fe, 40 mg as ferrous sulphate; Mo, 0.5 mg as 8 sodium molybdate; Mn, 100 mg as manganous oxide; Cu, 15 mg as copper sulfate; Zn, 100 mg as zinc oxide. 9 <sup>2</sup>Premix provided per kilogram of grower diet: vitamin A as retinyl acetate, 10,000 IU; vitamin $D_3$ as 10 cholecalciferol, 5,000 IU; vitamin E as alpha-tocopherol-acetate, 50 IU; vitamin K, 3 mg; thiamin, 2 mg; 11 riboflavin, 8 mg; pyridoxine, 3 mg; vitamin B<sub>12</sub>, 15 µg; biotin, 0.12 mg; calcium pantothenate, 12 mg; nicotinic 12 acid, 50 mg; folic acid, 2 mg; 400 mg choline chloride; methionine, 3,018 mg; threonine, 726 mg; lysine, 2,831 13 mg; I, 1 mg as calcium iodate; Se, 0.35 mg as sodium selenite; Fe, 40 mg as ferrous sulphate; Mo, 0.5 mg as 14 sodium molybdate; Mn, 100 mg as manganous oxide; Cu, 15 mg as copper sulfate; Zn, 100 mg as zinc oxide. 15 <sup>3</sup>Premix provided per kilogram of finisher diet: vitamin A as retinyl acetate, 10,000 IU; vitamin $D_3$ as 16 cholecalciferol, 5,000 IU; vitamin E as alpha-tocopherol-acetate, 50 IU; vitamin K, 3 mg; thiamin, 2 mg; 17 riboflavin, 6 mg; pyridoxine, 3 mg; vitamin B<sub>12</sub>, 15 µg; biotin, 0.12 mg; calcium pantothenate, 10 mg; nicotinic 18 acid, 50 mg; folic acid, 1 mg; 350 mg choline chloride; methionine, 2,514 mg; threonine, 361 mg; lysine, 1,779

- 19 mg; I, 1 mg as calcium iodate; Se, 0.35 mg as sodium selenite; Fe, 40 mg as ferrous sulphate; Mo, 0.5 mg as
- 20 sodium molybdate; Mn, 100 mg as manganous oxide; Cu, 15 mg as copper sulfate; Zn, 100 mg as zinc oxide.
- 21 <sup>4</sup>Calculated according to NRC (1994).

## 23 Supplemental Table 2. Body weight, feed intake and growth performance between d 7 and 21 of life of female and male broiler chickens raised at

Item	Residual feed intake <sup>1,2</sup>					$P^{3,4}$		
	Location	Low	Medium	High	SEM	FE	location	FE × location
Females								
Body weight, d 7 of life (g)	L1+2	145	145	147	2.6	0.805	0.001	0.802
	L1	141	138 <sup>y</sup>	141	3.6			
	L2	149	151 <sup>x</sup>	153 <sup>x</sup>	2.6			
Body weight, d 21 of life (g)	L1+2	906	848	893	21.56	0.133	< 0.001	0.817
	L1	972 <sup>x</sup>	895 <sup>x</sup>	852 <sup>x</sup>	29.48			
	L2	840 <sup>y</sup>	801 <sup>y</sup>	834 <sup>y</sup>	31.35			
Total feed intake, d 7-21 of life (g)	L1+2	1009 <sup>b</sup>	$1023^{abB}$	1083 <sup>aA</sup>	23.07	0.067*	0.339	0.391
	L1	1001 <sup>b</sup>	1059 <sup>ab</sup>	1094ª	31.55			
	L2	1017	987 <sup>B</sup>	1073 <sup>A</sup>	33.55			
Total body weight gain, d 7-21 of life (g)	L1+2	761	703	746	21.02	0.131	< 0.001	0.845
	L1	831 <sup>x</sup>	757 <sup>x</sup>	811 <sup>x</sup>	28.75			
	L2	691 <sup>y</sup>	650 <sup>y</sup>	681 <sup>y</sup>	30.57			
Males								
Body weight, d 7 of life (g)	L1+2	145	145	148	2.173	0.704	< 0.001	0.919
	L1	139 <sup>x</sup>	140 <sup>x</sup>	141 <sup>x</sup>	3.010			
	L2	152 <sup>y</sup>	150 <sup>y</sup>	154 <sup>y</sup>	3.047			
Body weight, d 21 of life (g)	L1+2	920	928	933	19.09	0.895	< 0.001	0.046

24 two different locations.

	L1 L2	933 908	999 856	1005 861	27.23 26.76			
Total feed intake, d 7-21 of life (g)	L1+2	1049 <sup>b</sup>	1088 <sup>ab</sup>	1155ª	20.15	0.002***	0.985	0.042
	L1	1008 <sup>b</sup>	1096 <sup>ab</sup>	1187 <sup>a</sup>	28.74			
	L2	1089 <sup>ab</sup>	1080 <sup>b</sup>	1123ª	28.25			
Total body weight gain, d 7-21 of life (g)	L1+2	775	782	785	17.77	0.914	< 0.001	0.032
	L1	794 <sup>B</sup>	859 <sup>x</sup>	863 <sup>xA</sup>	25.34			
	L2	756	705 <sup>y</sup>	707 <sup>y</sup>	24.91			

25 FE, feed efficiency; RFI, residual feed intake; L1, University of Veterinary Medicine Vienna (Vienna, Austria); L2, Agri-Food and Biosciences Institute (Hillsborough, Northern

26 Ireland, UK).

27 <sup>1</sup>Values are least squares means  $\pm$  standard error of the mean (SEM).

28 <sup>2</sup>Each RFI group represents n = 9 female and male chickens at location 1; n = 6 low RFI, n = 11 medium RFI, and n = 8 high RFI females as well as n = 10 low RFI, n = 9

29 medium RFI, and n = 9 high RFI males at location 2.

30  ${}^{3}P$ : probability level.

- 31 <sup>4</sup>Linear polynominal contrast:  $*P \le 0.05$ , and  $***P \le 0.001$ .
- 32 a-cLeast squares means within a row without a common lowercase superscript differ among RFI groups (P < 0.05).
- 33 <sup>A,B</sup>Least squares means within a row without a common uppercase superscript tend to differ among RFI groups (P < 0.1).
- 34 x,yLeast squares means within a column without a common lowercase superscript differ between locations (P < 0.05).

## 35 Metzler-Zebeli et al. - Supplemental Material

## 36 Relationship analysis

Single nucleotide polymorphism genotypes were used to examine the genetic relationship 37 of all birds within and between each population received. In order to achieve the genetic 38 relationship of each pair of samples supplied, a G-matrix was established using the PreGS 39 program by Prof I. Misztal (Animal Breeding and Genetics group, University of Georgia, 40 Athens, GA, USA). Supplemental Table 2 lists the relationship statistic per population. 41 These data indicate that there is very little genetic relationship between any two birds 42 within replicate batch 1 and replicate batch 2 from the location 1. In replicate batch 3 at 43 44 location 1, two birds appeared to be half-sibs (relationship of 0.25). Similarly, the replicate batch 1 from location 2 appeared to contain two birds that are half-sibs (relationship of 0.20). 45 The overall relationships within and between populations has been plotted and is illustrated in 46 Supplemental Figure 1. 47

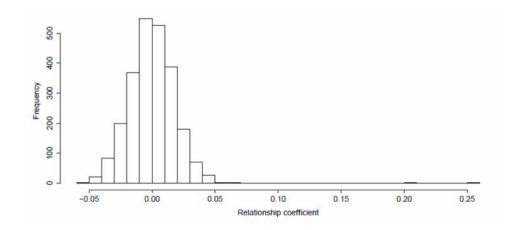
48

### 49 Supplemental Table 3. Genomic relationships among chickens.

	genomic relationships comparisons among birds						
		mean	sd	min	max		
Location 1 + 2	2415	0	0.02	-0.05	0.25		

50

51 Supplemental Figure 1. G-relationships among chickens from both locations.





# Assessing serum metabolite profiles as predictors for feed efficiency in broiler chickens reared at geographically distant locations

Metzler-Zebeli, B., Magowan, E., Hollman, M., Ball, E., Molnar, A., Lawlor, P. G., ... Zebeli, Q. (2017). Assessing serum metabolite profiles as predictors for feed efficiency in broiler chickens reared at geographically distant locations. BRITISH POULTRY SCIENCE, 1-10. DOI: 10.1080/00071668.2017.1362688

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