Relationship between indirect genetic effects for growth, environmental enrichment, coping style and sex with the serum metabolome profile of pigs

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Including Indirect Genetic Effects (IGE) in breeding programs to reduce aggression in group housed animals has been proposed. However, the effect of selection for IGE for growth on animal metabolism and physiology is unknown. The purpose of this study was twofold: (1) To investigate the effects of this new breeding method along with two housing (barren and straw), coping style (high and low resisters) and sex (female and castrated males) options on the metabolome profile of pigs. (2) To identify and map biological processes associated with a regrouping test at 9 weeks of age. We used Nuclear Magnetic Resonance to quantify 49 serum metabolites at week 8, 9 and 22. Also, we quantified 3 catecholamines (tyramine, epinephrine, phenylethylamine) and serotonin and three water soluble vitamins (B2, B5 and B7). Overall, no significant differences were observed between negative and positive IGE animals. The magnitude of change (delta) of many metabolites as a response to the regrouping test was significantly affected by IGE, especially that of the amino acids (P < 0.05), being greater in positive IGE pigs. The regrouping test was associated with alteration in glycine, serine and threonine metabolism. In conclusion positive and negative IGE animals respond differently to the regrouping test.

Pork consumers are increasingly demanding good pork quality together with high animal welfare standards. Traits like behavior in relation to housing types, social interaction in group housed pigs, and different stress factors, are becoming economically important in breeding programs. Both environment and genetics have an impact on behavior and social interaction in group housed pigs. Modern pig production systems include regrouping of the animals at different stages e.g. weaning, and finishing stage. When regrouping occurs, pigs need to cope with abrupt changes in food supply, housing conditions and social environment. Animals adopt different coping strategies in response to these environmental stresses1. In relation to coping strategy, the backtest had been used to categorize pigs as high or low resisters. High-resisters (HR) pigs are characterized by greater number of struggles and vocalizations, when compared with low-resisters (LR)1–3. Moreover, pigs classified as HR and LR are characterized by differences in behavioral, physiological and endocrine responses to conflict situations2,3.

Current intensive indoor systems limit a pig's ability to perform rooting and chewing behaviors leading to boredom, frustration and to redirect exploratory behavior towards pen mates4 resulting in competition, aggression, tail biting and other negative social interactions. The positive impact of environmental enrichment on pig behavior has been extensively studied and reviewed5–7. Genetics also plays an important role in the behavior of animals. Indirect Genetic Effect (IGE) describes the impact of an individual on trait values of other individuals and was first introduced by Griffing8. Including IGE in breeding programs to reduce aggression in group housed animals has been proposed9–12. Pigs selected for a positive effect on the growth of their group members (positive IGE pigs) performed less non-reciprocal biting behavior and less aggression in a regrouping test, and less biting on enrichment material11,13. In addition, positive IGE pigs were less fearful than negative IGE pigs as determined

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by several novelty tests\textsuperscript{14,15}. The response of pigs in novelty tests seems to depend also on their sex, females being less fearful than castrated males\textsuperscript{14}. Female pigs have been found to have lower leukocytes and basal cortisol but higher haptoglobin concentrations when comparing to castrated males\textsuperscript{15}, suggesting physiological differences between females and castrated males.

Some evidence showed that at weaning the weight of positive and negative IGE pigs is not significantly different\textsuperscript{5}. However, opposite to expectation, positive IGE animals grew slower in the first 4 days after weaning than negative IGE pigs\textsuperscript{16}. Furthermore, the positive IGE animals had lower carcass weight and less muscle depth\textsuperscript{5}. At present, it is not clear whether selection for IGE on growth, combined with different housing conditions, sex and coping strategy, has any negative or positive impact on the animals’ metabolism, physiology or performance.

Metabolomics, although relatively new in animal sciences, has also shown great potential to identify indicators of economically important traits such as feed efficiency\textsuperscript{17}. Metabolomics approaches are expected to result in a better understanding of the physiological mechanisms underlying the traits of interests, as well as for pleiotropic effects and to predict which traits may be affected by the selection performed on another trait.

Based on the previous literature published using the same pigs and experimental design it has reported that pigs with positive and negative IGE differ in their behaviors, immunological status, and carcass traits suggesting differences in the metabolism. Therefore, in the present study we hypothesized that pigs selected for positive or negative IGE differ in their metabolome profile. The purpose of this study was two-fold: (1) To investigate the effects of IGE for growth, housing conditions, sex and coping style on the serum metabolome profile of pigs using a targeted metabolomics approach. (2) To identify and map biological processes associated with IGE for growth and a regrouping test.

### Results

The effect of time was significant for all metabolites except for ethanol and dimethyl sulfone concentration pointing to an effect of age on metabolite concentration. Batch was the second factor which affected a great number of metabolites and their ratios (forty-six) pointing to an effect of the season.

The effect of indirect genetic effects on growth on metabolite concentration. Overall, no significant differences in the concentration of the metabolites were observed between negative and positive IGE animals. Negative IGE animals tended to have greater ratio of L-valine/isobutyric acid (69.30 ± 1.59 vs 65.50 ± 1.77; \( P = 0.09 \)), L-leucine/isobutyric acid (39.70 ± 0.96 vs 37.7 ± 1.06; \( P = 0.07 \)) and isoleucine/isobutyric acid (29.30 ± 0.70 vs 27.30 ± 0.78; \( P = 0.08 \)). These ratios decreased over time reaching the lowest values at week 22 (Supplementary Fig. S1).

The magnitude of change (delta) of many metabolites as a response to regrouping was significantly affected by IGE class, especially that of the amino acids (\( P < 0.05 \)). The significant metabolites are shown in Table 1. Negative IGE animals had a smaller delta of amino acids (AA) such as L-alanine, L-asparagine, L-aspartic acid, isoleucine, L-proline, L-serine and total glucogenic AA when compared to positive IGE animals (Table 1; \( P < 0.05 \)). After regrouping, L-lysine, L-glycine and the sum of branched-chain amino acids (BAA), increased in negative IGE but decreased in positive IGE pigs (Table 1; \( P < 0.05 \)). Moreover, negative IGE animals had a smaller delta of D-glucose, citric acid, choline, L-aspartic acid/L-glutamine ratio, creatine and creatine/creatinine ratio (Table 1; \( P < 0.05 \)).

| Metabolite \( \mu M \) | Negative IGE | Positive IGE | \( P \) value |
|-------------------------|-------------|--------------|---------------|
| L-alanine | −231.61 ± 44.31 | −386.61 ± 49.60 | 0.006 |
| L-asparagine | −17.73 ± 5.21 | −34.19 ± 5.84 | 0.005 |
| L-aspartic acid | −8.84 ± 1.83 | −16.35 ± 2.10 | 0.003 |
| L-isoleucine | −23.40 ± 7.41 | −43.90 ± 8.13 | 0.009 |
| L-proline | −65.90 ± 17.31 | −125.62 ± 20.10 | 0.01 |
| L-serine | −13.69 ± 7.21 | −51.18 ± 8.17 | 0.0001 |
| Glucogenic AA | −570.0 ± 151.0 | −928.0 ± 165.0 | 0.0002 |
| L-lysine | 62.92 ± 20.60 | −1.65 ± 22.80 | 0.004 |
| L-glycine | 20.80 ± 56.69 | −188.90 ± 59.50 | <0.0001 |
| BAA | 37.20 ± 28.89 | 48.0 ± 27.81 | 0.003 |
| D-glucose | −309.17 ± 244.26 | −974.93 ± 264.40 | 0.004 |
| Citric acid | −58.1 ± 9.39 | 80.0 ± 10.18 | 0.01 |
| Choline | −3.92 ± 1.17 | −6.92 ± 1.28 | 0.02 |
| L-aspartic acid/L-glutamine | −1.57 ± 0.57 | −2.81 ± 0.64 | 0.04 |
| Creatine | 91.5 ± 19.9 | 163.1 ± 22.7 | 0.009 |
| Creatine/creatinine | 1.03 ± 0.24 | 2.03 ± 0.27 | 0.002 |

Table 1. Least squared means and SE of the magnitude of change (delta) of metabolite concentrations of pigs that have an estimated relative positive genetic effect or negative genetic effect (IGE) on the growth of their pen mates after regrouping at 9 weeks of age.
Multivariate analysis, partial least-squares discriminant analysis (PLS-DA) showed that positive and negative IGE animals do not display differences in their metabolite profiles, but a clear effect of time was observed, with samples collected at week 8, 9, and 22 clustering separately (Supplementary Fig. S2).

Pathway analysis revealed that the top ranked pathways associated with the response to the regrouping test were: glycine, serine and threonine metabolism (compound members: L-serine, choline, L-glycine, L-aspartic acid and creatine), arginine and proline metabolism (compound members: L-arginine, L-aspartic acid, L-glutamine, L-proline, creatine), and L-lysine degradation (compound members: L-aspartic acid and L-lysine; Fig. 1).

Correlation analysis showed no significant correlation between the delta of metabolites and the delta of haptoglobin, IgG, IgM and leukocytes (P > 0.05).

Housing. Housing significantly affected seven metabolites. Overall, pigs housed in the enriched environment had a greater concentration of creatinine, D-glucose, dimethyl sulfone, ethanol and L-tyrosine when compared with barren housed pigs (P < 0.05). On the other hand, enriched housed pigs had lower mannose and 3-methyl-2-oxovaleric acid concentration when compared with those in the barren environment (P < 0.05; Table 2).

The delta of the metabolites 2-hydroxybutyrate, isobutyric acid, L-glutamine and vitamin B5 was positive in animals housed in the barren environment, while it was negative in animals housed in the enriched environment after the regrouping test (Table 3; P < 0.05).

Interaction between IGE on growth and housing conditions on the delta of metabolites. The interaction between IGE and housing conditions affected the delta of the amino acid L-threonine and body weight at week 10. In the barren environment negative IGE animals had a positive delta of L-threonine meanwhile positive IGE pigs showed a negative delta (Supplementary Fig. S3; P < 0.05). Overall, there was no significant difference in body weight between positive and negative IGE pigs, however at week 10, one week after

![Figure 1. The results of pathway analysis depicting the metabolome view, displaying all the matched pathways as circles. The color is based on its p value and size of each circle on its pathway impact value.](image)

| Metabolites µM | Enriched | Barren | P value |
|----------------|----------|--------|---------|
| Creatinine     | 94.20 ± 1.08 | 90.40 ± 1.18 | 0.01    |
| D-glucose      | 5533 ± 65.80  | 5340 ± 72.20  | 0.04    |
| Dimethyl sulfone| 16.40 ± 0.90  | 13.70 ± 0.99  | 0.04    |
| Ethanol        | 20.0 ± 0.69   | 17.70 ± 0.74  | 0.003   |
| L-tyrosine     | 124 ± 1.90    | 118 ± 2.12    | 0.04    |
| Mannose        | 52.60 ± 1.24  | 56.60 ± 1.42  | 0.02    |
| 3-Methyl-2-oxovaleric acid | 7.47 ± 0.16 | 8.07 ± 0.17 | 0.008   |

Table 2. Least squared means and SE of the concentration of significant serum metabolites in animals housed in enriched and barren environments at week 8, 9 and 22.
regrouping test, positive IGE pigs housed in the barren environment had a lower body weight when compared with positive IGE pigs housed in the enriched environment (Supplementary Fig. S3; P < 0.05).

**Coping style.** High resister pigs had an overall lower concentration of serotonin (P = 0.03), 1-methylhistidine (P = 0.02) and L-histidine (P = 0.04) when compared to LR (reactive) pigs (Fig. 2).

Acute stress due to regrouping had a larger impact on LR pigs. Pigs that were classified as HR had lower delta values of L-histidine, L-ornithine, ketoleucine, but greater delta values of vitamin B7, L-phenylalanine and L-tyrosine (Table 4; P < 0.05) when compared with pigs classified as LR.

**Sex.** Overall, female pigs had a greater concentration of serum 3-hydroxybutyric acid (BHBA; P = 0.01), and a lower serum serotonin concentration (P = 0.05) when compared to castrated male pigs (Fig. 3). In addition, females tended to have greater L-lactic acid when compared to castrated males (7973 ± 181 vs 7502 ± 182 µM; P = 0.06). After regrouping, aspartic acid decreased to a smaller extent in female animals when compared to castrated males (− 9.42 ± 1.92 vs − 15.47 ± 1.93 µM; P < 0.05).

Concentration of serum serotonin was negatively correlated with L-lactic acid (R = − 0.24, P < 0.001).

### Table 3. Least squared means and SE of the significant magnitude of change (delta) of metabolite concentration of pigs that were housed in enriched and barren environments after regrouping at 9 weeks of age.

| Metabolite µM | Enriched | Barren | P value |
|---------------|----------|--------|---------|
| 2-Hydroxybutyrate | − 3.69 ± 1.79 | 0.63 ± 1.76 | 0.01 |
| Isobutyric acid | − 0.32 ± 0.43 | 1.18 ± 0.42 | 0.001 |
| L-glutamine | − 0.40 ± 0.54 | 1.49 ± 0.53 | 0.001 |
| Vitamin B5 | − 0.14 ± 0.04 | 0.08 ± 0.04 | 0.0001 |

### Figure 2. Least squared means and SE of serum metabolites and serotonin concentration in pigs with a high resisting (HR) and low resisting (LR) coping style at week 8, 9 and 22.

### Table 4. Least squared means and SE of the magnitude of change (delta) of metabolite concentrations in pigs with high resisting (HR) and low resisting (LR) coping styles after regrouping at 9 weeks of age.

| Metabolite µM | HR | LR | P value |
|---------------|----|----|---------|
| L-histidine | 5.66 ± 2.32 | 15.81 ± 2.01 | 0.001 |
| Ketoleucine | − 0.25 ± 0.44 | 0.93 ± 0.41 | 0.005 |
| L-ornithine | − 3.34 ± 16.8 | 36.27 ± 15.1 | 0.04 |
| L-phenylalanine | − 14.77 ± 3.39 | 4.93 ± 2.05 | 0.02 |
| L-tyrosine | − 23.26 ± 5.32 | − 9.35 ± 4.71 | 0.04 |
| Vitamin B7 | − 0.0035 ± 0.0008 | 0.0010 ± 0.0007 | 0.01 |
In a more recent study on these same animals, Camerlink et al. 16 reported that positive IGE pigs had a greater latency to start eating and a lower weight gain over the first 4 days after weaning. Therefore, it is plausible that positive IGE pigs were mounting an immune response or they were temporarily in an energy deficiency state after the regrouping test. It was proposed that negative IGE pigs might have a more active immune system which, could lead to an increased need of amino acids e.g. for synthesis of haptoglobin 20. To test this hypothesis, we conducted correlation analysis between the delta of metabolites and delta of haptoglobin, IgG, IgM and leukocytes, reported by Reimert et al. 20. No significant correlation was found; therefore, it appears that the magnitude of change after the regrouping test might not be related with mounting an immune response. Another possible explanation of our results might be that the animals were temporarily in an energy deficiency status after a stressful event such as the regrouping test.

The delta of amino acids and other metabolites was altered in both positive and negative IGE animals. However, the extent of increase or decrease was more pronounced in positive IGE animals. For example, the extent to which glucogenic amino acids, D-glucose and citric acid decreased in serum and the extent to which creatine increased was greater in positive IGE animals. During energy deficiency, glycolysis, lipolysis and proteolysis processes develop simultaneously. During proteolysis, muscle tissue is used as a source of amino acid. Glucogenic amino acids are converted into glucose which is used as energy source. In our study, glucogenic amino acids and D-glucose decreased to a greater extent in positive IGE pigs suggesting that positive IGE pigs were in a more pronounced state of energy deficiency when compared to negative IGE pigs.

Furthermore, pathway analysis showed that the top ranked pathway associated with the response to regrouping was glycine, serine and threonine metabolism. This pathway has been reported to be genetically correlated with the average daily gain in pigs 21. The amino acid L-glycine decreased in positive IGE animals after regrouping. L-Glycine acts as a precursor for creatine, therefore the decrease in L-glycine in positive IGE pigs could explain the increase in creatine in these animals after the regrouping test. Creatine is synthesized in the liver, kidney and pancreas 22. Approximately 95% of creatine is stored in skeletal muscle in the form of phosphocreatine. During energy demand, phosphocreatine transfers its phosphate group to ADP to form ATP and creatine. Creatine is then spontaneously transformed to creatinine, transported out of skeletal muscle via the blood and eliminated by the kidneys 23. The creatine/creatine-phosphate system plays an important role in the storage and transmission of phosphate-bound energy. The greater increase in creatine concentration and the ratio creatine/creatinine, might reflect a greater energy demand of positive IGE animals to cope with the regrouping test.

In commercial settings, pigs are often relocated and mixed with other unknown pigs at weaning and/or at grower and finisher stages. Behavioral studies conducted by Reimert et al. 15 demonstrated that pigs with a positive IGE were faster to interact with ropes provided in a novel rope test than pigs with a negative IGE. Moreover, in a more recent study on these same animals, Camerlink et al. 16 reported that positive IGE pigs had a greater latency to start eating and a lower weight gain over the first 4 days after weaning. Therefore, it is plausible that at regrouping, animals with positive IGE will explore the new environment first and spend more time establishing social relationships rather than approaching feed and maybe eating less, which in turn could have led to a temporary energy deficiency status as a response to regrouping.

**Figure 3.** Least squared means and SE of serum 3-hydroxybutyric acid and serotonin concentrations in female (F) and castrated male (M) pigs at week 8, 9 and 22.

**Discussion**

In this study we investigated the effect of divergent selection for IGE on growth, environmental enrichment, coping style and sex on the metabolome profile of pigs using a targeted metabolomics approach. As was expected, time of sampling affected the concentration of all, except for two, metabolites, pointing to an effect of age on metabolite concentration. Age-related metabolite changes have been previously reported in human plasma 8.

Overall, no significant differences in the concentration of the metabolites were observed between negative and positive IGE animals suggesting that this method of selection doesn't impact the metabolic profile of pigs in this study. Positive IGE animals tended to have lower ratios of L-valine/isobutyric acid, L-leucine/isobutyric acid and isoleucine/isobutyric acid when compared with negative IGE animals. Branched-chain volatile fatty acids including isobutyric acid are built up from the degradation products of branched amino acids and it has been suggested to be indicative of better utilization of dietary protein by the microbiota 19. However, our results showed only a tendency which deserve further future investigation.

Positive and negative IGE groups responded differently to acute stress induced by regrouping test. The extent to which the concentration of many metabolites changed after the regrouping test, especially that of the amino acids, D-glucose, citric acid, and creatine was greater in positive IGE pigs. These results suggest two possibilities: Either the positive IGE pigs were mounting an immune response or they were temporarily in an energy deficiency state after the regrouping test. It was proposed that negative IGE pigs might have a more active immune system which, could lead to an increased need of amino acids e.g. for synthesis of haptoglobin 20. To test this hypothesis, we conducted correlation analysis between the delta of metabolites and delta of haptoglobin, IgG, IgM and leukocytes, reported by Reimert et al. 20. No significant correlation was found; therefore, it appears that the magnitude of change after the regrouping test might not be related with mounting an immune response. Another possible explanation of our results might be that the animals were temporarily in an energy deficiency status after a stressful event such as the regrouping test.
A wealth of research has demonstrated that environmental enrichment exerts biological effects in many species, most of which are beneficial, and keeping the animals in a barren environment invokes chronic stress\textsuperscript{1,2}. Environmental enrichment has been shown to increase the concentration of brain-derived neurotrophic factor in pigs provided with foraging enrichment\textsuperscript{25} and decrease levels of lactate\textsuperscript{7}. In our study enriched pigs had greater concentrations of serum L-tyrosine when compared with animals in the barren environment. L-tyrosine is the precursor of the catecholamines and neurotransmitters\textsuperscript{28} and supplementation with tyrosine may reduce stress responses in swine\textsuperscript{27}. In a previous study that was carried out using the same pigs as the current one, enriched housed pigs were less stressed\textsuperscript{20} when compared with those animals housed in a barren environment. Therefore, a link between straw provision, stress response and tyrosine can not be excluded and deserves further investigation. This result also suggests that supplementing L-tyrosine might be a feeding strategy to reduce the stress response in pigs that are raised in barren environment.

Enriched housed pigs had lower 3-methyl-2-oxovaleric acid when compared with barren housed pigs. This compound is a keto-acid that induces acidosis, that arises from the incomplete breakdown of branched-chain amino acids\textsuperscript{29}. To our knowledge, reports about 3-methyl-2-oxovaleric acid in pigs are absent. Enriched housed pigs could had a better utilization of branched amino acids and energy status, as also shown by the greater D-glucose concentration. Also, during the regrouping test (providing acute stress) housing pigs in an enriched environment could results in a better utilization of amino acids. In support of this idea, we found that delta of 2-hydroxybutyrate decreased in animals housed in the enriched environment. 2-hydroxybutyrate is produced by amino acid catabolism (L-threonine and L-methionine). In humans, 2-hydroxybutyrate appears at high concentrations in situations related to deficient energy metabolism and has been reported as an early marker for impaired glucose regulation\textsuperscript{29}. We think that providing straw has a positive effect on the ability of pigs to cope with acute stress during regrouping with other unknown pigs.

The molecular basis for how straw induces modulation of metabolites or other compounds are unknown, however there is some evidence that providing enrichment modulates the concentration of some hormones and metabolites\textsuperscript{22,25}. The purpose of providing enrichment (straw) is to serve as stimulus for pigs to perform exploratory, manipulative and rooting behavior. As pigs perform their natural behaviors, they also get more active which in turn leads to mobilization of different energy sources (carbohydrates, lipids and proteins). This could be one of the mechanisms that “kick start” the changes we found in the concentration of metabolites which are the intermediate or end product of the metabolism.

When in stressful situations animals can adopt coping strategies. Immunological differences between HR and LR pigs have been previously described\textsuperscript{30,39}. For example, HR pigs were reported to have greater complement activity via the alternative pathway (APW), therefore a more active innate immune system than LR pigs\textsuperscript{30}. In our study HR, had lower serum concentration of serotonin, histidine and 1-methylhistidine when compared to LR pigs. Interestingly, Hessing et al.\textsuperscript{31} reported that a difference in balance between T-helper 1 and T-helper 2 cells was underlying the difference between HR and LR pigs. Peripheral serotonin exerts pro-inflammatory as well as anti-inflammatory responses\textsuperscript{32,33}. Immune cells have the capacity to synthesize, transport, store, degrade and respond to serotonin which exerts an important role in the balance of T cell subsets\textsuperscript{32,34}. The differences in serum serotonin concentration between HR and LR suggest a difference in the serotonergic system between HR and LR pigs which is in agreement with Koolhaas et al.\textsuperscript{34}.

In our study, female pigs were found to have lower serum serotonin. Differences in the rates of brain serotonin synthesis between males and females have been reported in humans\textsuperscript{35} and rodents\textsuperscript{36}. In addition, female pigs had greater concentration of BHBA and tended to have greater concentration of L-lactic acid during the experiment. BHBA is a ketone body and together with the accumulation of lactic acid can lead to reduced serum pH. Interestingly, pigs and chickens fed acidified diets had depressed levels of serotonin in the brain\textsuperscript{37,38}. In addition, it was proposed that lactic acid increases serotonin uptake in human blood platelets in vitro by 40% thereby decreasing serotonin concentration in serum\textsuperscript{39}. In our study serum serotonin was negatively correlated with L-lactic acid and BHBA cannot be ruled out. We speculate that BHBA and lactic acid concentration could be one of the underlying physiological differences in serotonin concentration between females and castrated males.

In conclusion selection for positive or negative IGE for growth does not impact the metabolic profile and weight of the animals used in this study. However, animals selected for IGE responded differently to regrouping. The regrouping test (providing acute stress) had a greater impact on the metabolic profile of positive IGE animals. However, no differences in body weight were observed. The practical implication of this study is that while selection for IGE doesn’t have a negative impact on serum metabolites concentration, regrouping (mixing) is a critical process which can seriously impact the performance of the animals. More research is necessary to determine whether selection for more than one generation on IGE for growth, has any impact on metabolism, physiology, performance and response to acute stress. To our best knowledge this is one of the first studies to report that housing enrichment modulates metabolites concentration in pigs. Besides, differences in serum serotonin concentration, suggest a difference in the serotonergic system between HR and LR pigs and between females and castrated males.

Methods

All experimental procedures were approved by the University of Alberta Animal Policy and Welfare Committee for Livestock as category “A” level of invasiveness (no animal manipulation as the animal work had already been completed and the present study involved only laboratory analysis). Animals had been cared for in accordance with the recommendations in the European Guidelines for accommodation and care of animals. The protocol was approved by the Institutional Animal Care and Use Committee of Wageningen University (Protocol Number:
Animals. Pigs were born at the experimental farm of Topigs Research Center IPG in Beilen, the Netherlands. Until weaning, they were housed in standard lactation pens of 3.8 m². At 3 days of age males were surgically castrated but tails were not docked. At four weeks of age, the piglets were weaned, and a total of 480 offspring Tempo × Topigs-20 pigs were introduced in five different batches (96 pigs/batch). Pigs were housed until slaughterc at 23 weeks of age in 16 pens of 6–7 m² located in one room per batch at the animal facility of Wageningen University, Wageningen, the Netherlands. Each pen consisted of six unrelated pigs, three females and three males. In addition, in each pen there were at least two pigs with a high-resisting (HR) coping style and two with a low-resisting (LR) coping style. Half of the pigs were housed in a barren environment, and the other half in a straw enriched environment (Supplementary Material and Method). All pigs were subjected to a regrouping test at week 9 of life to induce acute stress and to simulate a similar procedure which takes place in a farm setting. The details of behavioral tests (coping style classification and regrouping test) are presented in Supplementary Material and Method.

A total of 180 animals were included in the metabolomics analysis (animals from batch 1 and batch 2), balanced regarding sex and coping style, selected for positive IGE (n = 78) and negative IGE (n = 102) for growth of their pen mates and housed in barren or straw-enriched pens. The number of the animals in each group are summarized in Supplementary Table S1.

Blood collection and weight. Blood was collected three times; once in the week before regrouping (8 weeks of age), three days after regrouping (9 weeks of age), and at 22 weeks of age. Blood was taken by venipuncture from the jugular vein and collected in serum separating tubes and incubated for one hour at 37 °C after which they were centrifuged at 3200 rpm for 12 min at 20 °C. Sera were stored at −80 °C until further analysis.

For this study body weight records at 10, 17 and 23 weeks of age were considered in the statistical analysis.

Metabolomics analysis. Metabolomics analysis was performed at The Metabolomics Innovation Centre (TMIC) at University of Alberta following established protocols. Serum samples were used for Nuclear Magnetic Resonance (NMR) analysis to measure the concentration of 49 metabolites including amino acids, sugars, alcohols, organic acids, amines, TCA cycle intermediates, and short chain fatty acids.

Because catecholamines are an essential component of the stress responses of animals and serotonin is a neurotransmitter involved in regulation of emotional states such as fear and aggression we quantified 3 catecholamines (tyramine, epinephrine, phenylethylamine) and serotonin. In addition, three water soluble vitamins (B2, B5 and B7) which are involved in regulation of carbohydrate, fat and protein metabolism were quantified.

Three amino acid indexes were calculated: the sum of glucogenic, ketogenic, and branched AA. Glucogenic AA were calculated as the sum of all the amino acids, except L-leucine and L-lysine. Ketogenic AA were calculated as the sum of L-leucine and L-lysine. Branched AA were calculated as the sum of L-valine, L-leucine, and L-isoleucine. In addition, the sum of ketone bodies and the ratios L-valine/isobutyric acid, L-leucine/isobutyric acid, L-isoleucine/isobutyric acid, L-aspartic acid/glutamic acid, and creatine/creatinine were calculated because they are involved in the same metabolic pathway. The number of the variables that were included in the analysis are summarized in Supplementary Table S2.

Nuclear magnetic resonance compound identification and quantification. All serum samples were thawed on ice before use. A deproteinization step, involving ultra-filtration was introduced in the protocol to remove serum proteins. Prior to filtration, 3 KDa cut-off centrifugal filter units (Amicon Microcon YM-3) were rinsed five times each with 0.5 mL of 250 mM K2HPO4 + KH2PO4 pH 7.0 v/v containing 5 mM DSS (2,2-dimethyl-2-silcepentane-5-sulphonate), 5.8 mM 2-chloropyrimidine-5 carboxylate, and 0.1% NaN3 in H2O) was added to the sample. The filtrates were collected and the volumes were recorded and samples diluted to 200 µL with 50 mM potassium salt buffer (pH 7) as needed. The dilution factor was recorded and metabolite concentrations were corrected accordingly in the subsequent analysis. Subsequently, 50 µL of a standard buffer solution (54% D2O: 46% 250 mM K2HPO4 + KH2PO4, pH 7.0 v/v containing 5 mM DSS (2,2-dimethyl-2-silcepentane-5-sulphonate), 5.84 mM 2-chloropyrimidine-5 carboxylate, and 0.1% NaNO3 in H2O)) was added to the sample. The sample (250 µL) was then transferred to a 3 mm SampleJet NMR tube for subsequent spectral analysis. All 1H-NMR spectra were collected on a 700 MHz Avance III (Bruker) spectrometer equipped with a 5 mm HCN Z-gradient pulsed-field gradient cryoprobe. NMR pulse sequence parameters were as follows: acquisition time 4 s, mixing time 100 ms, recycle delay 10 ms, saturation delay 990 ms, transients (700 MHz) 128, steady state scans 8 and presaturation field width 80 Hz. 1H-NMR spectra were acquired at 25 °C using the first transient of the NOESY pre-saturation pulse sequence (noesypr1d), chosen for its high degree of quantitative accuracy. All Free Induction Decays were zero-filled to 250 K data points. The singlet produced by the DSS methyl groups was used as an internal standard for chemical shift referencing (set to 0 ppm) and for quantification all 1H-NMR spectra were corrected accordingly in the subsequent analysis.
processed and analyzed using the online Bayesil software package\textsuperscript{43}. The spectral fitting for each metabolite was done using the standard serum metabolite library. Most of the visible peaks were annotated with a compound name.

**Catecholamines and serotonin quantification.** Catecholamine quantification was carried out following the protocol reported by Zheng et al.\textsuperscript{44} All serum samples were thawed on ice in the dark before use and transferred to a 96-well filter plate. Ten μL of the internal standard mixture (ISTD) solution and 25 μL of samples (PBS for “zero” samples, calibration curve standards, and serum samples) were directly onto the center of each well. The whole plate was evaporated under nitrogen flow for 45 min to dry. Fifty μL of phenylisothiocyanate (PITC) derivatization solution was then added to each well, and the reaction was kept at room temperature for 20 min. The samples were then again dried in a nitrogen evaporator for 90 min to remove the liquid, especially the excess PITC, followed by the addition of 75 μL of methanol containing 5 mM ammonium acetate. The plate was covered and shaken at 450 rpm for 30 min at room temperature, and then spun by centrifuge for 15 min at 1200 rpm. To each well of the collection plate, 75 μL of water was added and mixed thoroughly, and 50 μL was injected for LC–MS/MS analysis. An Agilent 1100 series HPLC system (Agilent, Palo Alto, CA) and an Agilent reversed-phase Zorbax Eclipse XDB C18 column (3.0 mm × 100 mm, 3.5 μm particle size, 80 Å pore size) with a Phenomenex (Torrance, CA, U.S.A.) SecurityGuard C18 pre-column (4.0 mm × 3.0 mm) were used for online LC–MS/MS analysis with an AB SCIEX QTRAP® 4000 mass spectrometer (Sciex Canada, Concord, ON.). The controlling software for the LC–MS system was Analyst® 1.6.2. For the HPLC work, solvent A was 0.2% (v/v) formic acid in water, and solvent B was 0.2% (v/v) formic acid in ACN. The gradient profile for the HPLC solvent run was as follows: t = 0 min, 0% B; t = 0.5 min, 0% B; t = 5.5 min, 95% B; t = 6.5 min, 95% B; t = 7.0 min, 0% B; and t = 9.5 min, 0% B. The column oven was set to 50 °C. The flow rate was 500 μL/min, and the sample injection volume was 40 μL. The mass spectrometer was set to a positive electrospray ionization mode with multiple reaction monitoring (MRM). The IonSpray voltage was set at 5500 V and the temperature at 500 °C. The curtain gas (CUR), ion source gas 1 (GAS1), ion source gas 2 (GAS2) and collision gas (CAD) were set at 20, 40, 50 and medium, respectively. The entrance potential (EP) was set at 10 V. Likewise, the declustering potential (DP), collision energy (CE), collision cell exit potential (CXP), MRM Q1 and Q3 were optimized and set individually for each analyte and each corresponding deuterated ISTD\textsuperscript{44}.

**Vitamin quantification.** Vitamins B2, B3 and B7 were quantified. Samples were thawed on ice, transferred to a 96-well plate and centrifuged at 10,000 rpm for 3 min. Ten μL of the ISTD solution was added to all wells of the plate except the blank well. A seven-point calibration curves was used for quantification. Three QCs (synthetic mixtures, low, medium and high) were used. Fifty μL of calibration standards was pipetted onto the center of wells starting from the lowest concentration to the highest concentration. Fifty μL QCs was added onto the center of wells starting from the lowest concentration to the highest concentration and 50 μL of sample was added onto the center of the remaining wells. In addition, 100 μL of extraction solvent (19 mg ammonium acetate in 50 mL HPLC grade methanol) was added to each well. The plate was covered with the plastic lid and shaken for 30 min at 150 rpm and centrifuged for 5 min at 500 g. All the wells in the capture plate were diluted with 50 μL of HPLC grade water and sealed. The plate was shaken for 2 min at 150 rpm and run using LC–MS. The running time was 7 min. An Agilent 1100 series HPLC system (Agilent, Palo Alto, CA) and a Phenomenex (Torrance, CA, USA) Kinetex C18 (2.6 μm, 3.0 × 100 mm, 100A, S/N: 669679-8) with a Phenomenex SecurityGuard C18 pre-column (4.0 mm × 3.0 mm) were used for online LC–MS/MS with an AB SCIEX QTRAP® 4000 mass spectrometer (Sciex Canada, Concord, ON.). The controlling software for the LC–MS system was Analyst® 1.6.2. For the HPLC work, solvent A was 10 mM AmFA in water, and solvent B was 10 mM AmFA in 90:10 ACN:water. The gradient profile for the HPLC solvent run was as follows: t = 0 min, 0% B; t = 2.0 min, 30% B; t = 3.5 min, 60% B; t = 5.0 min, 75% B; t = 5.5 min, 75% B; t = 5.6 min, 0% B; and t = 7.0 min, 0% B. The column oven was set to 40 °C. The flow rate was 500 μL/min, and the sample injection volume was 5 μL. The mass spectrometer was set to a positive electrospray ionization mode with MRM. The IonSpray voltage was set at 5500 V and the temperature at 450 °C. The curtain gas, ion source GAS1, ion source GAS2 and CAD were set at 20, 40, 60 and medium, respectively. The EP was set at 10 V. Likewise, DP, CE, CXP, MRM Q1 and Q3 were optimized and set individually for each analyte and each corresponding deuterated ISTD.

**Statistical analysis.** Univariate statistical analysis was performed using R (version 3.5.2)\textsuperscript{45}. The response variables were the metabolite concentrations and weight. Statistical analysis for weight was carried out considering the animal weight at 10, 17 and 23 weeks of age. Prior to the analysis, when appropriate, metabolites were log transformed to obtain normality of residuals. Fixed effects were tested for their significance by fitting a Mixed Model for Repeated Measurements assuming a first-order autocorrelation variance–covariance structure (CovAR1) which allows for unequally spaced time observations with the final model being:

\[
\gamma = \mu + \text{IGE} + \text{time} + \text{sex} + \text{batch} + \text{housing} + \text{coping style} + \text{IGE} \times \text{housing} + \text{pen} + e,
\]

where \(\gamma\) is the response variable (metabolites and weight), \(\text{IGE}\) (positive or negative), \(\text{time}\) (week 8, week 9, week 22), \(\text{sex}\) (females and castrated males), \(\text{batch}\) (1 and 2), \(\text{housing}\) (barren and enriched), \(\text{coping style}\) (HR, LR), and \(e\) refers to the common error term. To test differences between IGE classes, housing condition, coping style and sex the Least Square Means (LSMs) for each pair-wise comparison were estimated and adjusted using false discovery rate (FDR). In addition, statistical analysis for weight were done separately for week 10, 17 and 23 with the Linear Mixed Model being:
$y = \mu + \text{IGE} + \text{sex} + \text{batch} + \text{housing} + \text{coping style} + \text{IGE} \times \text{housing} + \text{pen} + e,$

where IGE class, sex, batch, housing condition, coping style and housing condition and its interaction with IGE, were included as fixed effects, pen as a random effect and $e$ refers to the common error term.

To investigate physiological differences in response to the regrouping test, the delta of metabolites between week 9 and week 8 was calculated and analyzed with a Linear Mixed Model:

$y = \mu + \text{IGE} + \text{sex} + \text{batch} + \text{housing} + \text{coping style} + \text{IGE} \times \text{housing} + \text{pen} + e,$

where IGE class, sex, batch, housing condition, coping style and housing condition and its interaction with IGE, were included as fixed effects, pen as a random effect and $e$ refers to the common error term.

In addition, we performed correlations analysis between: 1-serotonin and lactic acid and 1- between the delta of metabolite concentrations and the delta of haptoglobin, IgG, IgM and leukocytes, reported by Reimert et al. Significant results were considered if $P \leq 0.05$ and $P > 0.05$ and $\leq 0.10$ it was considered as a tendency.

Multivariate analysis was performed considering the metabolites’ concentration at week 8, 9 and 22 separately. Before multivariate analysis each metabolite concentration was adjusted for the significant fixed effect (sex, batch, housing and coping style) other than time and IGE. For each metabolite, the adjusted values were extracted and used for Partial Least Squares- Discriminant Analysis (PLS-DA). Multivariate analysis was performed using a web-based tool MetaboAnalyst (https://www.metaboanalyst.ca/) according to previously published protocols.

To understand biological processes involved in the response to regrouping in the two IGE classes, pathway analysis was performed using the list of significant “delta” of metabolites and pathway-associated metabolite sets were selected as metabolite library. Metabolic pathway analysis calculates the impact on the pathways based on network topology analysis. For this analysis we used Homo sapiens (SMPDB) as the library because no library for pigs exists. In order to estimate the importance of a compound within a given metabolic network, we selected global test algorithm and relative-betweeness centrality which measures the number of shortest paths going through the node of interest.

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Author contributions

E.D performed data curation, formal analysis, wrote, revised and reviewed the manuscript. I.R, L.Z and P.M provided the samples, and contributed in Statistical analysis and reviewed the manuscript. E.K supervised the analysis and reviewed the manuscript. G.S.P conceptualized, supervised the project, and reviewed data analysis and the manuscript. All authors reviewed and approved the final manuscript.

Competing interests

The authors declare no competing interests.

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