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RNAseq reveals modulation of genes involved in fatty acid biosynthesis in chicken liver according to genetic background, sex and diet

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Manuscripts

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59 28 **Keywords:** fatty acid metabolism, transcriptome, poultry, local breeds, differentially expressed genes.
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29 ABSTRACT

30 Increases in chicken production are mainly due to specialised breeds. However, of increasing importance
31 are the local breeds, known for their ability to adapt to the environment and for their unique products.
32 Conventional poultry products contain lower levels of n-3 fatty acids compared to those obtained from
33 local breeds, therefore the aim of this study was to evaluate the modulation of expression of genes
34 involved in LC-PUFA biosynthesis pathways according to genetic background, diet conditions, and sex.
35 Animals from two local breeds and a commercial line were fed different diets: control and experimental
36 diet (10% linseed supplementation). For each breed and diet group, both males and females were reared.
37 The RNA was extracted from 36 liver samples and was sequenced by RNAseq method. Bioinformatic
38 analysis was carried out to find differentially expressed genes (DEGs) from different comparisons between
39 experimental groups. Results showed low impact of diet on DEGs related to fatty acid biosynthesis, but
40 linseed diet increased percentage of n-3 fatty acids of liver. Sex, particularly the female groups, and genetic
41 background determined the differential expression of genes related to LC-PUFA biosynthesis. Specifically,
42 females of local breeds shared 23 up-regulated genes when compared to their respective commercial line
43 groups. Some of the shared genes had a role in *de novo* triglyceride biosynthesis (*MTTPL* and *GPAM*), and in
44 genes involved in *de novo* FA biosynthesis (*ACACA* and *SCD*) was detected. In conclusion, sex and local
45 genetic background appear to have influence on the expression of genes related to LC-PUFA synthesis.

47 INTRODUCTION

48 To date, commercial chicken lines have dominated the poultry market so far, thanks to the fast growth and
49 high production performances. Besides the standard commercial chicken lines, of increasing importance
50 are the local breeds which are able to adapt to the environment where they live (Perini *et al.*, 2021), and
51 for their different quality products compared to conventional ones (Franzoni *et al.*, 2021). Unfortunately,
52 the productivity traits are critical for local breeds, which are instead appreciated for the diversity of their
53 products, which could include economically exploitable traits (Moula *et al.*, 2010). Currently, it is still

1
2
3 54 difficult to obtain economic feedback from breeders who decide to raise local breeds because of low
4
5 55 competitiveness on productive performances.
6
7 56 A potential valorisation strategy is the enrichment in terms of nutritional value of meat and eggs. Enhancing
8
9
10 57 the quantity of n-3 fatty acids in chicken meat and eggs through the diet can have important impacts: i)
11
12 58 supplementing the fatty acid intake in **the** human diet, as it is usually rich in n-6 and lacking in n-3 fatty
13
14 59 acids (Mariamenatu & Abdu, 2021); ii) chickens are able to convert precursors of n-3 in long chain (**> 20**
15
16 60 **carbon atoms**) polyunsaturated fatty acids (LC-PUFA); iii) **finally**, eggs and meat enriched with n-3 would
17
18
19 61 have increased market **potential** compared to standard animal products.
20
21 62 The LC-PUFAs, eicosapentaenoic acid (EPA, 20:5 n-3) and docosahexaenoic acid (DHA, 22:6 n-3) are the
22
23 63 main form of n-3. The fatty acid profiles of poultry meat and eggs can be readily enriched in EPA and DHA
24
25 64 through the diet (Cartoni Mancinelli *et al.*, 2022). Although conventional poultry meat contains low levels
26
27
28 65 of EPA and DHA, several factors such as sex, feed, and genetic background may influence their quantity. In
29
30 66 particular, genetic background is reported in the literature as a crucial factor: higher concentrations of n-3
31
32 67 LC-PUFA are synthesized by local breeds compared to commercial lines (Cartoni Mancinelli *et al.*, 2021).
33
34 68 The application of a diet rich in **precursors** of n-3 (α -linolenic acid: 18:3n-3, ALA) can affect the amount of
35
36
37 69 LC-PUFA in meat and eggs, and may change the expression level of genes involved in the process of lipid
38
39 70 biosynthesis and elongation. **In mammals**, these two metabolic pathways, mainly occur in the adipose
40
41 71 tissue, while in chicken the majority (90%) takes place in the liver (Nematbakhsh *et al.*, 2021). To elucidate
42
43 72 the complete expression profile of liver genes, RNAseq provides a useful tool. This transcriptomic approach
44
45 73 can accurately study tissue transcriptomes with high resolution and depth. Increasingly, the nutrigenomic
46
47
48 74 approach is used to investigate the effect of diets on metabolic processes also in poultry (Soglia *et al.*,
49
50 75 2022). A recent study of the liver transcriptome after folic acid supplementation in the diet of broiler
51
52 76 chickens found that peroxisome proliferator activated receptor (PPAR) signalling is the pathway most
53
54 77 activated by the enriched diet (Zhang *et al.*, 2021).
55
56
57 78 In recent years, many studies tried to elucidate which genes are principally involved in the biosynthesis
58
59 79 process of LC-PUFA. Studies focused on the evaluation of the presence/activity of elongase of very long
60

1
2
3 80 chain fatty acid (*ELOVL*) and fatty acid desaturase (*FADS*) enzymes (Cartoni Mancinelli *et al.*, 2022; Lee *et*
4
5 81 *al.*, 2016). Although **recently** many studies have highlighted some pathways involved in the biosynthesis of
6
7 82 LC-PUFAs (e.g. *PPAR* signaling), **there remains** a lack of scientific research in this area (Mihelic *et al.*, 2020).

8
9
10 83 The aim of this study was to determine **the modulation of expression of genes** involved in LC-PUFA
11
12 84 biosynthesis pathways, according to genetic background (local breed vs commercial line), diets, and sex.

13
14 85

16 86 **MATERIALS AND METHODS**

18 19 87 ***Animal ethics***

20
21 88 Birds were raised, handled and processed according to the European legislation for the protection of
22
23 89 chickens kept for meat production (European Commission, 2007), the protection of animals at the time of
24
25 90 killing (European Commission, 2009) and the protection of animals used for scientific purposes (European
26
27 91 Commission, 2010). The experimental protocol was positively evaluated and approved by the Ethical
28
29 92 Committee of the University of Perugia (ID: 62700_15/07/2020).

30
31 93

32 33 94 ***Experimental design***

34
35 95 The trial was carried out in the experimental section of the Department of Agricultural, Food and
36
37 96 Environmental Sciences (University of Perugia, Italy). Specifically, three different chicken genetic
38
39 97 **backgrounds** were used: Robusta Maculata (RM) and Bionda Piemontese (BP) as local **breeds** and Ross 308
40
41 98 (ROSS) as a commercial line. **Both RM and BP are dual-purpose breeds, in particular BP shows higher**
42
43 99 **aptitude for laying eggs (200 eggs/year vs 150 eggs/year of RM) (www.pollitaliani.it)**. One-day old chicks of
44
45 100 **each sex** were housed in an **environmentally**-controlled poultry facility and vaccinated against coccidiosis,
46
47 101 infectious bronchitis, Marek's Disease, Newcastle Disease and Gumboro. Until 21 days of age, all the
48
49 102 chickens received the same starter diet. **Then each genetic background was divided into two groups: one**
50
51 103 **fed a standard diet (control group, Ct), the second fed the standard diet supplemented with 10 % extruded**
52
53 104 **linseed (experimental group, L). The two diets were formulated to meet the nutritional recommendations**
54
55 105 **of Ross 308 birds (Table S1) (Aviagen, 2019). Three replicates/group were formed, each representing both**

1
2
3 106 **male and female chickens (5 males + 5 females/each replicate)**. Chickens were raised in different indoor
4
5 107 pens at the same stocking density (5 chickens/m²) and the temperature was set according to the age of the
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7 108 birds (20-32°C; Relative Humidity: 65-72%). Water and feed were provided *ad libitum*. At 81 days of age,
8
9
10 109 **live body weight was recorded (g)** for all the birds **which** were **later** slaughtered in a commercial
11
12 110 slaughterhouse. From each replicate, **two** birds (**one** for each gender) were selected. Table S2 schematically
13
14 111 explains the experimental design **used in** the present study.

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16 112
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18
19 113 ***Tissue collection and RNA extraction:***

20
21 114 Liver tissue was collected from all the animals. Around 1 g of tissue was collected in a 2 mL cryogenic vial
22
23 115 (Corning® Inc., Corning, NY), instantaneously frozen in liquid nitrogen, and stored at -80° C until RNA
24
25 116 extraction. **The remaining part of liver tissue was collected and stored at -20°C for further analysis of fatty**
26
27
28 117 **acids (FA) profile.**

29
30 118 At this stage, RNA was extracted only from liver tissue. Samples were prepared from 36 animals, with three
31
32 119 biological replicates for each experimental group (Table S2). RNA isolation was performed with **a**
33
34 120 NucleoSpin RNA Mini kit for RNA purification (Macherey-Nagel, Germany) as recommended by the
35
36
37 121 manufacturer, starting from 30 mg of tissue. A homogenization step was carried out using an Omni Tissue
38
39 122 Homogenizer (TH) - Omni, Inc) in ice. DNA contamination was removed with a DNase enzyme (included in
40
41 123 the kit) during **the** incubation step. RNA concentration was measured with a Qubit 3.0 fluorometer (Life
42
43 124 Technologies) and Qubit RNA HS Assay Kit (Life Technologies) according to manufacturer instruction. Total
44
45 125 RNA (~1.5 µg/sample) was sent to Genewiz (South Plainfield, NJ, USA) for quality check, library preparation
46
47
48 126 and Illumina sequencing. For testing RNA quality, the Agilent (Santa Clara, CA) 2100 Bioanalyzer Nano Kit
49
50 127 was used, and all the samples showed RNA integrity (RIN) numbers ≥6.5 (data not shown). Library
51
52 128 preparation was carried out through polyA + selection and paired-end (PE) sequencing was run on an
53
54 129 Illumina NovaSeq System that generated 150 bp PE reads.

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56 130
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58
59 131 ***Bioinformatic analysis***

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3 132 RNAseq data were quality checked by FastQC software (Brown *et al.*, 2017). Paired qualified reads were
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5 133 mapped to the chicken reference genome (GRCg6a, accession number: PRJNA13342) using STAR software
6
7 134 (Dobin *et al.*, 2015) with default parameters. The mapping rates for each sample were assessed using
8
9
10 135 Samtools 'flagstat' command (Heng Li *et al.*, 2009). The mapped BAM file was then used in featureCounts
11
12 136 software that, assigning raw alignments to annotated genes, generates the raw expression count file used
13
14 137 for gene expression quantification (Liao *et al.*, 2014). An overview of genomic distance in the dataset was
15
16 138 performed through Principal Component Analysis using the ggplot2 R package. Differential expression (DE)
17
18
19 139 analysis was performed using the DESeq2 R package (Love *et al.*, 2014). The P-value adjustment was made
20
21 140 using Benjamini and Hochberg's approach for controlling the False Discovery Rate (FDR). Genes with a P-
22
23 141 adjusted value < 0.05 and $-2 > \text{Log}_2$ fold change (LFC) > 2 were used as thresholds for significant DE by
24
25 142 DESeq2. Visualization of contrast between different experimental groups was performed by
26
27
28 143 EnhancedVolcano package in R (Blighe *et al.*, 2022). The overlap of significant genes among experimental
29
30 144 groups was assessed with Venn diagrams built with the Interactivenn web tool (Heberle *et al.*, 2015).
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32 145 Ingenuity Pathway Analysis (IPA, QIAGEN Redwood City, www.qiagen.com/ingenuity) was used to identify
33
34 146 gene ontology pathways, and regulatory networks to which DE genes belong, as well as upstream
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36
37 147 regulators. Ingenuity Pathway Analysis can transform a set of genes into a number of relevant networks
38
39 148 based on comprehensive records maintained in the Ingenuity Pathways Knowledge Base. **According to DEG**
40
41 149 **input, IPA software calculates a z-score value for each biological pathway. When a pathway has been**
42
43 150 **categorized with a z-score > 2, it resulted as an activated pathway.** Moreover, to confirm the IPA results,
44
45 151 gene ontologies (GO) were examined using the Panther (v17.0) database according to the Statistical over-
46
47
48 152 representation test. Fisher's test was used to correct for False Discovery Rate (Figure S1) (Mi *et al.*, 2019).
49
50 153 **Differential expression (DE) analysis was performed in individual contrasts according to diet, sex and**
51
52 154 **genetic background, separately. In particular we evaluated six diet contrasts, six contrasts for sex and four**
53
54 155 **contrasts for each local breed against the ROSS birds (Table S3).**
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59 157 **Fatty acids (FA) profile**
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1
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3 158 The fatty acids profile was determined from the same samples used for the RNAseq. Lipids were extracted
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5 159 from 5 g of liver/sample based on the methods described in Folch *et al.* (1957). The fatty acids were
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7 160 identified in the form of their methyl esters using a Varian Gas Chromatograph (CP-3800) and a DB wax
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9
10 161 capillary column (25 mm ϕ , 30 m long). The establishment of each fatty acid was done in relation to the
11
12 162 retention time with respect to fatty acid methyl ester standards (FAME, Sigma-Aldrich, Bellefonte, PA). The
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14 163 relative quantity of each fatty acid present in the liver was calculated using heneicosanoic acid (C21:0;
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16 164 Sigma-Aldrich) as the internal standard. Data were expressed as % of total FA. The average amount of each
17
18
19 165 FA was used to calculate the sum of total PUFA of the n-3 and n-6 series. After using the function of the
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21 166 Shapiro test (for normality test), data were analysed with ANOVA and with Tukey's *post-hoc* test in R
22
23 167 software. All results are expressed as mean \pm SEM, with the level of significance set at $p < 0.05$.

24 168

27 169 RESULTS

30 170 *Sequencing data analysis*

32
33 171 Average mapping rate was 85.11% across all samples, 98% of which were seen to be properly paired.
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35 172 Among all the samples, the average quality score for each base was $> Q30$ and the average GC content in
36
37 173 sequenced samples was around 52% (data not shown). After all quality checks, all 36 samples were taken
38
39 174 forward for further analysis. In order to shed light on genetic similarity, a principal component analysis
40
41
42 175 (PCA) plot was computed and visualized using R software. This showed genetic distance between the
43
44 176 samples according to the comparison of Principal Component 1 and 2 (PC1 and PC2) (Figure 1). Genetic
45
46 177 diversity was appreciable, especially between the two main clusters seen in Figure 1 representing male and
47
48 178 female birds. The male cluster showed less genetic variance compared to the female one, and the ROSS
49
50
51 179 birds grouped closest to each other within each cluster, because of their standard genetic background. On
52
53 180 the contrary, the two local breeds showed more genetic variance within the experimental groups and were
54
55 181 clearly separated from the ROSS cluster.

56 182

59 183 *Body weight and fatty acids profile of animals*

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2
3 184 The effect of diet was evaluated according to the results from live body weight (Figure S2) and the
4
5 185 percentage of n-3 and n-6 PUFA, and LC-PUFA (Figure 2). The diet had no statistical impact on the body
6
7 186 weight at slaughtering of the two local breeds, either in males or females. Instead, the L diet significantly
8
9
10 187 decreased the body weight in ROSS (male and female).

11
12 188 Figure 2 represent the n-6 and n-3 PUFA proportion in liver, and the sum of LC-PUFA. Figure 2a shows the
13
14 189 percentage of LC-PUFA was not statistically affected by the diet. On the contrary, the n-6 PUFA (ranging
15
16 190 from 18% to 34% approximately) showed higher level than n-3 PUFA (from 4% to 12). Moreover, the
17
18
19 191 control diet generally increased the n-6 percentage in both sexes of ROSS and in RM_M. On the contrary,
20
21 192 the L diet significantly enhanced the n-3 level in all groups, excepted for RM females.

22 23 193 24 25 194 ***Differentially expressed genes (DEGs)***

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27
28 195 Each experimental group was compared against one another to check differential gene expression. This was
29
30 196 done on the basis of: diet, sex, and genotype (Table S3). In particular, we were interested in comparing
31
32 197 local breeds to the broiler (ROSS) genotype. Appendix S1 lists all significantly differentially expressed genes
33
34 198 in each comparison. Moreover, all genes related to FA metabolism, and which were found to be
35
36
37 199 differentially expressed, are reported in Table S4.

38 39 200 40 41 201 ***Diet contrast***

42
43 202 For the evaluation of the possible effect of diet on gene expression between sexes and genetic background,
44
45 203 the linseed diet group was compared to the control diet group within each sex and within each genetic
46
47
48 204 background. Experimental groups formed by “genetic background” and “sex” were taking in consideration
49
50 205 individually. Linseed diet groups have been used as experimental groups, hence the up and down-regulated
51
52 206 genes and the GO analyses referring to them. Figure 3 shows the volcano plots resulting from each
53
54 207 comparison. Figures 2a-b showed the DEGs detected for the female (BP_F) and male (BP_M) Bionda
55
56
57 208 Piemontese experimental groups respectively. In BP_F, the diet played a limited role in differentiating gene
58
59 209 expression level, indeed Figure 3a had only eight differentially expressed genes with a P -value < 0.05 , of
60

1
2
3 210 which some involved in fatty acid metabolism (e.g. *LYG2*, *SCD*), being found down-regulated with the L diet.
4
5 211 Other important genes involved in FA synthesis (*ACACB*, *ACACA*, *FASN*) had a significant *P*-value but had
6
7 212 lower LFC values (between -2 and 2) (Appendix S1).
8
9
10 213 In the BP_M group, the diet contrast showed 280 significant genes, and the Gene Ontology study confirmed
11
12 214 that the most influenced pathways were cellular lipid metabolic process (GO:0044255) and lipid metabolic
13
14 215 process (GO:0006629) (Appendix S2). Regarding the Robusta Maculata breed, volcano plots for DEGs in
15
16 216 females (RM_F) and males (RM_M) are shown in Figures 2c-d respectively. Contrary to the BP birds, the
17
18 217 RM_F comparison showed more DEGs than the RM_M group, namely 83 and 13 genes. Gene ontology
19
20 218 analysis in each group showed that the RM_F group had biological processes which were significantly
21
22 219 enriched in the L diet, although not directly related with FA biosynthesis, and males did not show anything
23
24 220 significant (i.e. 'de novo' protein folding (GO:0006458)) (Appendix S2). Finally, the last genetic background
25
26 221 evaluated for the effect of diet was the commercial ROSS hybrid (Figure 3e-f). In both female and male
27
28 222 groups, a limited number of DEGs were identified (72 and 13, respectively). This is similar to the GO
29
30 223 biological annotations associated with FA metabolism. None of the six diet contrasts resulted in a
31
32 224 significantly activated biological pathway in IPA analysis.
33
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37 225
38

39 226 **Effect of sex**

40
41 227 The effect of sex on gene expression was assessed as follows: the experimental groups compared within
42
43 228 the same genetic background and diet groups but differing by sex. Female groups were taken as the
44
45 229 experimental group, hence the up and down regulated genes and GO analysis are referred to "F" group in
46
47 230 each contrast. Looking at Figure 4a, it is easy to appreciate the numerous DEGs up-regulated in females. For
48
49 231 each BP comparison, as shown in Figures 4a-b, the GO annotations on up-regulated genes clarify the sexual
50
51 232 dimorphism in the most typical way: cellular response to estrogen stimulus (GO:0071391) and response to
52
53 233 estrogens (GO:0043627) (Appendix S3). When comparing the ROSS broiler birds, and examining the linseed
54
55 234 diet group (L), significant biological enrichment was found in organo-nitrogen compound biosynthetic
56
57 235 process (GO:1901566), for example (Appendix S3). With regard to FA pathways, in BP birds there was
58
59
60

1
2
3 236 activation of lipid pathways in females, namely lipid transport (GO:0006869) and lipid localization
4
5 237 (GO:0010876). This result from Panther was also confirmed using IPA software. Fatty acid metabolism was
6
7 238 seen to be significantly enhanced in the two female groups of the BP breed. The significance of fatty acid
8
9
10 239 concentration pathway in RM chickens treated with linseed diet was also indicated (Appendix S3). In
11
12 240 particular, the data in Appendix S3 shed light on the genes that were differentially expressed in females
13
14 241 within the experimental group. The females expressed some genes strictly related to FA metabolism
15
16 242 differentially to males within respective groups. For examples: *ELOVL2* in BP_Ct, *ELOVL2*, *FADS1* and *FADS2*
17
18 243 in BP_L, and *SCD* in RM_L.

20
21 244
22

23 245 **Genetic background comparison**

25 246 One of the aims of this study was to assess the impact of breed and genetic makeup on the profile of LC-
26
27
28 247 PUFA expression. In particular, the comparison between local breeds and the commercial line (same sex
29
30 248 and diet) was examined.
31
32 249 **Figure 5 represents** the DEGs from the BP vs ROSS comparison. **BP breed represented the experimental**
33
34 250 **group, which up and down regulated genes and GO analysis are referred to.** Figures 5a-b show females
35
36 251 treated with control and linseed diet respectively. According to results from Panther, many pathways
37
38
39 252 related to lipid metabolism were up-regulated in the BP breed, such as fatty acid biosynthetic process
40
41 253 (GO:0006633) and long-chain fatty acid metabolic process (GO:0001676) (Appendix S4). In the female
42
43 254 control diet group, the up-regulated genes found through IPA were related to FA metabolism: *SCD*, *CYP1A1*,
44
45 255 *FASN*, *ACACA*, *THRSP*, *FAR1*, *FADS1* (Appendix S4). **Figure 5b showed** the results for a genetic background
46
47
48 256 contrast with females fed the linseed diet. The results from GO analysis showed a large number of lipid
49
50 257 related pathways, the most important being lipid metabolic process (GO:0006629). Some of the genes
51
52 258 related to this, which were more highly expressed in BP relative to the commercial line include *FABP3*, *SCD*,
53
54 259 *GPAM*, *LPIN1*, *SREBF2* and *ACACA* (Appendix S4). Meanwhile, Figures 5c-d exhibit the contrast of BP with
55
56
57 260 ROSS birds, in males fed the two different diets. With the control diet, BP males showed higher expression
58
59 261 of genes involved in fatty acid metabolic process (GO:0006631) in comparison with ROSS. BP males fed the
60

1
2
3 262 linseed diet do not show DEGs significant in FA metabolism (e.g. *FASN*, *ACACA*, *ACACB*) when compared
4
5 263 with ROSS (Appendix S4), but amino acid biosynthetic processes are highlighted in the GO analysis.
6
7 264 We next wanted to compare Robusta Maculata (RM) vs Ross308 (ROSS) groups. The experimental groups
8
9
10 265 consisted of the RM breed, hence up and down regulated genes and GO analysis are referred to as “RM”
11
12 266 breed groups. The volcano plots are presented in Figure 6, of which the first two (Figure 6a-b) represented
13
14 267 the contrast in females, with control and experimental diets, respectively. Figure 6a showed 391 genes with
15
16 268 a P-value lower than 0.05, of which 234 genes had a value of $-2 > \text{LFC} > 2$, and 121 genes in RM with $\text{LFC} > 2$.
17
18
19 269 The Panther GO analysis clearly showed that within the up-regulated genes, fatty acid metabolic process
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21 270 (GO:0006631) was clearly enhanced, with similar pathways activated in female linseed diet groups. IPA
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23 271 analysis showed significance of LC-PUFA synthesis in RM birds fed the control diet and synthesis of FA in
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25 272 RM birds fed the linseed diet (Appendix S5). The RM female groups shared DEGs clearly involved in FA
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28 273 pathways, such as *SCD*, *ACACA*, and *THRSP*. On the contrary, in RM males (Figures 6c-d), no pathways
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30 274 related to FA appeared as significantly involved in birds fed either diet. Instead, immune pathways were
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32 275 active in Ct diet birds, while there were no significant pathways in birds fed the supplemented diet
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35 276 (Appendix S5).

36 37 277 38 39 278 **Comparison of local vs commercial females**

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41 279 According to the results from the sex comparison previously described, females, for the most part,
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43 280 appeared more specialized for FA production compared to males. Moreover, in the comparison of genetic
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45 281 backgrounds, all the female groups from local breeds showed higher activation of FA-related pathways
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48 282 compared to commercial hybrid birds. In order to understand which DEGs were common to the local
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50 283 breeds, a Venn diagram was generated (Figure 7). Here we took into consideration only the female birds,
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52 284 and only the genes differentially expressed in local breeds (BP and RM) compared to ROSS. Interestingly, 23
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54 285 genes were commonly up-regulated among local breeds (Appendix S6). A heat map was generated to
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57 286 graphically represent the 23 shared genes and their expression level among all female samples (Table 1)
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59 287 (Figure 8). GeneMania software was used to identify any connection between them, with *GPAM* being
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3 288 suggested as being directly related to *THRSP* and *SCD*. In fact, we saw *GPAM* as being up-regulated in both
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5 289 BP female groups and in RM females fed the control diet, but not the linseed diet (Figure S3).
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10 291 DISCUSSION

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13 292 Many studies have reported how the diet can modulate the expression of genes in different tissues (Sevane
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15 293 *et al.*, 2014; Szalai *et al.*, 2021). Here, we fed animals with a diet rich in the precursor of n-3 LC-PUFA. We
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17 294 designed the experiment assuming that diet plays a key role in changing lipid metabolism in the liver.
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19 295 Indeed, vitamin supplementation (Niu *et al.*, 2009) taurine (He *et al.*, 2019) and zinc oxide nanoparticles
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21 296 (Ramiah *et al.*, 2019) have been used to modulate gene expression of FA related genes, whereas the
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23 297 supplementation of dietary PUFA is widely used to increase LC-PUFA concentration in animal products of
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25 298 different animal species, such as dairy cows, rabbit and turkey (Castellini *et al.*, 2022; Kliem *et al.*, 2019;
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27 299 Szalai *et al.*, 2021). Chickens are usually an ideal target for diet supplementation, especially linseed based
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29 300 supplements (Head *et al.*, 2019; Jing *et al.*, 2013; Sevane *et al.*, 2014; Zhang *et al.*, 2021).
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33 301 In the present study, the different diets affected the body weight of commercial birds. This result is
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35 302 explained by the fact that during the experimental trial, the ROSS chickens showed problems of adaptation
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37 303 to the linseed diet, consuming around 7% less quantity of the L diet with respect to the Ct diet, both in
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39 304 males and in females. In other studies was already reported how extruded linseed diet could lead to lower
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41 305 body weight gain and lower feed intake in broiler (Avazkhanloo *et al.*, 2020; Anjum *et al.*, 2013). This
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43 306 outcome indicated once again how the local breeds can easily adapt themselves to different environmental
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45 307 factor (e.g. alternative diet sources), while commercial lines encounter difficulties. In the present study, n-
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47 308 3, n-6 and LC-PUFA of liver were examined. LC-PUFA, which are directly related to the desaturation and
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49 309 elongation ability, was not affected by diet, conversely to n-6 and n-3 levels. The percentage of n-6 was
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51 310 generally higher in the Ct diet because the Ct diet is higher in linoleic acid (LA), which is the precursors of n-
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53 311 6 PUFA. On the other hand, the L diet positively affected the n-3 PUFA, and indeed the n-3 is significantly
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55 312 higher in the L diet than in the corresponding Ct groups (Figure 2). Both these results are commonly found
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57 313 in the literature (Hang *et al.*, 2018; Head *et al.*, 2019; Meineri *et al.*, 2018). Head *et al.* (2019) described the
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3 314 modulation of FA profile in a given tissue through the diet, and also provided insights into low effect of the
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5 315 diet on the gene expression. The fact that the trend of LC-PUFA in the different experimental groups was
6
7 316 similar confirms this assertion. Thus, it seems that the entity of LC-PUFA produced by liver is mainly due to
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10 317 the level of dietary precursors. On the same time, the type of precursor furnished α -linolenic (ALA, n-3) or
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12 318 linolenic (LA, n-6) determined an alternative accumulation of the same PUFA series in the liver. Indeed, we
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14 319 found that the dietary supplementation of linseed affected expression of genes involved in FA metabolism
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16 320 in only one group (BP_M group – Appendix S2). The most highly expressed genes in the linseed diet group
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19 321 included *PPARGC1A*, and *LPIN1*. *PPARGC1A* is a co-activator of *PPAR γ* which is a transcription factor that
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21 322 participates in induction and stimulation of fat-specific genes and fatty acid bio-synthesis (Wang *et al.*,
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23 323 2017). *LPIN1* is involved in synthesis and transport of triacylglycerol, a major constituent of chicken lipids
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25 324 (Desert *et al.*, 2018), and also found to be over expressed in a chicken line selected for intramuscular fat
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28 325 content together with *PPAR γ* (Liu *et al.*, 2020). The remaining contrasts (n=5) for diet did not show enriched
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30 326 pathways, and for this reason, we can assert that diet is the factor with lowest impact on gene expression
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32 327 of fatty acid metabolism. Although no GO enrichment was discovered regarding FA metabolism in the other
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34 328 5 diet contrasts, there are some interesting genes down-regulated with the L diet belonging to the BP_F
35
36 329 and RM_F groups. Indeed, in BP_F the L diet showed down-regulation of *LYG2*, *ACACA*, *FASN*, *SCD* and in
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39 330 RM_F, down-regulation of *PLIN2* and *GPAM*, all important genes in FA metabolism and discussed later in
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41 331 this section. This is also in agreement with Head *et al.* (2019) who ascertained that the effect of linseed on
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43 332 the expression of 14 genes involved in FA metabolism in chicken liver was low. We therefore saw differing
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45 333 modulation of FA metabolism in liver between genetic backgrounds and sexes through diet.
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48 334 This study also evaluated the effect of sexual dimorphism and its impact on FA metabolism. A clear
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50 335 separation between male and female birds is appreciable in Figure 1, underlying a divergent level of
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52 336 expression in genes between the two groups. The background of sex effect on FA metabolism was
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54 337 investigated by Poureslami *et al.* (2010b) where they described the poor effect of sex on final concentration
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57 338 of MUFA and PUFA of the n-3 and n-6 series (Poureslami *et al.*, 2010a). Elsewhere, Lopez-Ferrer *et al.* used
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59 339 dietary linseed oil, finding that abdominal fat percentage in male chickens was significantly lower than in
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the female chickens (López-Ferrer *et al.*, 2001). This was corroborated in human where the differential ability to activate PPAR α together with the well-documented effect of sex hormones on lipid metabolism may highlight sex as a significant factor in plasma FA levels (Thifault *et al.*, 2013). Differences between male and female chickens have never been thoroughly described and hence our interest in including sex effect in our analysis. BP_F, in both diet groups, showed a greater enrichment for estrogen pathways because it was the most egg-layer type chicken in the dataset. Indeed, estrogen in laying hens stimulates the liver to synthesize fatty acids for egg enrichment (Hanlon *et al.*, 2022). As reported by Ayres *et al.* (2013), differential expression of genes related to the female W chromosome are essential for sex determination in chickens. Some of them (*NIPBL* and *UBAP2*) have also been found to be central to sex determination in Japanese quail (Caetano-Anolles *et al.*, 2015). Regarding lipid metabolism, some genes were found to be more highly expressed in females with respect to males. This is the case for *ELOVL2* (in both BP contrasts), *FADS1*, *FADS2* (in BP_L) and *SCD* (in RM_L, and in both ROSS groups) that code for enzymes directly involved in desaturation and elongation of FA in the diet, such as ALA (18:3n-3), resulting in LC-PUFA (Head *et al.*, 2019). Interestingly, the results can be appreciated in the comparison of females with males in the BP breed. Besides the finding of elongases and desaturase-related genes being more highly expressed in females, other genes directly involved in FA metabolism were also identified. For instance, *APOB* plays a vital role in the assembly and secretion of triacylglycerol-rich lipoprotein in the liver of egg-laying chickens (Ma *et al.*, 2017), in line with the BP breed having a major aptitude in laying, especially when compared to ROSS (meat type) and RM birds (dual-purpose). Moreover, in the BP_L group, the female highly expressed genes of the fatty acid-binding protein (*FABP*) family, particularly *FABP3* and *FABP1*, which are considered biomarkers for intramuscular fat content. Bongiorno *et al.* (2022) reported that the saturated fatty acid (SFA) of breast meat was mainly influenced by gender. On the other hand, in the RM breed the higher female expression of *THRSP* strongly suggests a control of lipogenic targets (Resnyk *et al.*, 2017). Activated estrogen pathways could explain differences in genes related to FA in females. In 2017 Zhang *et al.* found three miRNAs involved in down-regulation of genes related to FA metabolism. More specifically, estrogen eliminates the suppressive effect of miRNAs on the target gene *ELOVL5*. Interestingly, estrogen suppresses

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3 366 the host *SLIT2* gene, thus decreasing the expression of intronic miR-218-5p to promote hepatic synthesis of
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5 367 long-chain polyunsaturated fatty acids in the liver (Zhang *et al.*, 2017). Although *ELOVL5* was not found to
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8 368 be directly overexpressed in females in the present study, many genes functionally related to it (i.e.
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10 369 *ELOVL2*, *SCD*, *FADS1*, *FADS2*) were identified. It is possible that estrogen pathways could regulate FA
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12 370 metabolism in general through the down-regulation of miR-218-5p, which regulates the genes related to FA
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14 371 metabolism, thus enhancing FA biosynthesis. Interestingly, it is the same miRNA (miR-218-5p, regulated by
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16 372 estrogen) that has a role in *FADS1* regulation in liver (Hong Li *et al.*, 2016), and *ELOVL2* regulation in chicken
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18 373 muscle (Zhang *et al.*, 2018). A more comprehensive evaluation of estrogen modulation via miRNA of FA
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21 374 metabolism should be investigated in further studies.
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24 375 In the present study, we observed that the genetic background was the variable having the largest effect on
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26 376 differential gene expression. Initially, we expected differing expression of genes for FA metabolism
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28 377 between the local breeds and in particular when compared to ROSS broilers. In the BP breed, a huge effect
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31 378 was identified in all four experimental groups when compared against ROSS birds. According to gene
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33 379 expression rates, the two female groups for each diet were the most divergent. These results suggested not
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35 380 only that the females had higher expression of genes involved in FA metabolism, but also that the BP
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37 381 female exhibited higher expression of genes responsible for FA metabolism, when compared to the ROSS
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40 382 females. In particular, the female control diet group showed an enrichment for genes involved in the long-
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42 383 chain fatty acid metabolic process (GO:0001676) supported by both GO analysis and the related genes
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44 384 *ELOVL2*, *SCD*, *FADS1*, *FADS2*, *THRSP*, *FABP3* and *LPIN1*.
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46 385 With males belonging to the local breeds, an effect was only observed within the BP breed, suggesting the
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49 386 importance of FA processing in this breed. RM females also showed interesting results with regards the FA
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51 387 process. On the contrary, the comparison between RM_M and ROSS_M did not show significant differences
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53 388 in genes involved in the FA processes. This lower effect in males of local breeds could be due to the
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55 389 estrogen stimulus that was strongly activated in females from local breeds and results in the difference in
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58 390 expression of genes related to LC-PUFA. Comparing all the results from contrasts of BP and RM female
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60 391 groups against the ROSS birds highlights 23 genes that are shared between the female local breeds (Figure

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3 392 **7 and Figure 8, Appendix S6**). Some of these are still uncharacterized (*LOC101747680*, *LOC107053670*,
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5 393 *LOC426220*, *LOC101749589*, *LOC112532382*, *LOC107050519*, *LOC107053691*, *LOC112532439*), but three of
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7 394 them are already reported in the literature with regards FA metabolism related functions. The
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10 395 *LOC101747680* gene codes for a C-like protein and it is located on chromosome 11, and *LOC107053670* is a
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12 396 non-coding RNA situated on chromosome 6, each with roles which are still unclear. *LOC426220* is located
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14 397 on the W sex chromosome, and is an avidin-related protein 6-like, and with all other avidin-like molecules
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16 398 localized on chromosome Z, **suggesting** a sex-specific regulation. Moreover, two studies have found this
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18 399 gene strongly correlated with egg yolk, vitelline membrane, and white coloured eggs (Gloux *et al.*, 2019;
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20 400 Niskanen *et al.*, 2005). Regarding eggs and fertility, another three genes were found differentially expressed
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22 401 in local breed females: *WDFC8*, *CTSEAL*, and *ZP1* (Table 1). *WDFC8* is clearly related to “WAP four-
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24 402 disulphide core domain proteins”. Members of this family are involved in various aspects of mucosal
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26 403 immunity (Wilkinson *et al.*, 2011). *WDFC8* was also found highly overexpressed in the liver of laying hens
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28 404 (Gloux *et al.*, 2019) and has been shown to protect egg yolk precursors from proteolytic
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30 405 activities/inactivation in the plasma, on their way from the liver to the growing oocytes (Marie Bourin *et al.*,
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32 406 2012). Moreover, *WDFC8* has been found as a unique protein in egg yolk of chicken (Farinazzo *et al.*, 2009).
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34 407 The avian-specific *CTSEAL* gene was found to be overexpressed in liver of laying hens and the translated
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36 408 protein shares large sequence similarity with cathepsin D, suggesting *CTSEAL* as an accessory of cathepsin D
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38 409 in the processing of egg yolk precursors (Bourin *et al.*, 2012; Gloux *et al.*, 2019). *ZP1* gene was also up-
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40 410 regulated in the liver of hens, where the protein is synthesized and then transported to the ovary to be
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42 411 inserted in the perivitelline membrane surrounding the oocyte (Gloux *et al.*, 2019). Here, it plays a
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44 412 fundamental role in the first interactions between spermatozoa and the oocyte (Bausek *et al.*, 2004). These
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46 413 three genes were differentially expressed in liver of BP and RM birds, with an explanation being that these
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48 414 **are** multi-purpose breeds and are thus more suitable chickens for egg production than the ROSS broilers.
49
50 415 Table 1 **and Figure 8** also show other genes not known to be involved in a particular pathway or showing a
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52 416 functional relationship with others. An example is chitinase (*CHIA*), which is a gene coding for a major
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54 417 protease-resistant glycosidase with a physiological role as a digestive enzyme that breaks down chitin-
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3 418 containing organisms in the chicken gastrointestinal tract (Tabata *et al.*, 2018). Another example is
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5 419 Doublecortin (*DCX*) – a member of a family of microtubule-associated proteins that are required for
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8 420 neuronal migration during cortical development (Vermillion *et al.*, 2014). Tumor Necrosis Factor Receptor
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10 421 Superfamily, Member 14 (*TNFRSF14*) was also found differentially expressed. It mediates apoptosis
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12 422 signalling, and can allow cancer cells to escape the immune process (Guo *et al.*, 2020).
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14 423 The remainder of genes found differentially expressed in females of local breeds (compared to ROSS), are
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16 424 involved in FA metabolism. The *ABHD12B* and *ABHD5* genes are part of the α/β hydrolase domain-
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18
19 425 containing family appointed to mobilization of lipids. In particular, *ABHD12B* has been highlighted by Li et al
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21 426 (2020) as a hydrolyser of very long chain lysophosphatidylserine lipids in human cells (Li *et al.*, 2021).
22
23 427 *ABHD5* the most well-characterized gene of its family codes for a protein cofactor of the ATGL enzyme and
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25 428 stimulates triacylglycerol hydrolase activity. Ouyang et al (2016) reported that overexpression of *ABDH5*
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28 429 markedly decreased the triglyceride content of preadipocytes in chicken (Ouyang *et al.*, 2016). This
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30 430 mechanism is modulated via the two above-mentioned genes and might explain the question of why native
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32 431 breed hens raised in free-range systems have less fat but higher polyunsaturated fatty acids in their meat
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34 432 muscles (Sokołowicz *et al.*, 2016). We discovered other genes differentially expressed in local female breeds
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36 433 belonging to the cytochrome P450 family were also highlighted: *CYP2C45* and *RP11-400G3.5*. Cytochrome
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39 434 P-450 2C45 (*CYP2C45*) was the most highly expressed cytochrome P-450 isoform in chicken liver and is a
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41 435 unique isoform in avian species (Watanabe *et al.*, 2013). Moreover, Zhao *et al.* (2019) have shown that
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43 436 *CYP2C45* was overexpressed in liver of overfed geese, hence promoting hepatic steatosis. Eventually, this
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45 437 resulted in the up-regulation of some genes involved in FA metabolism (i.e. *PK* and *ALOX5*) acting via *PPAR*
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48 438 pathways (Zhao *et al.*, 2019). *RP11-400G3.5* is reported as a *CYP2C21*-like pseudogene and is the closest
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50 439 gene to *CYP2C45* on chromosome 6 (Watanabe *et al.*, 2013). In fact, both of these genes showed similar
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52 440 function, especially in arachidonic and linolenic acid metabolism in KEGG pathways (data not shown).
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54 441 Furthermore, in local breeds we found significantly more expression in two genes having a role in de novo
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56 442 triglyceride biosynthesis (*MTTPL* and *GPAM*), two genes involved in *de novo* FA biosynthesis (*ACACA* and
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443 *SCD*), confirmedly involved in desaturation of FA in comparison to ROSS birds. All those genes have been
444 seen to be involved in FA or lipid metabolism (Figure S3).

445 In Table 1 we report the microsomal triglyceride transfer protein-like (*MTTPL*) gene located on
446 chromosome 6, which is an orthologue of *MTTP* on chromosome 4. *MTTP* is responsible for the assembly
447 and subsequent secretion of very low density lipoproteins from hepatocytes and controls the incorporation
448 of triglycerides into apolipoprotein B (Liu *et al.*, 2016). Moreover, *MTTPL* was found overexpressed in liver
449 of chickens fed with a diet rich in LC-PUFA precursor (Liu *et al.*, 2019; Tesseraud *et al.*, 2014). In the present
450 study, *MTTPL* was differentially expressed in the liver of local breeds, underlining their capacity for FA
451 metabolism. This was not the first time that genetic background has been shown to be a factor in liver
452 expression of *MTTPL*: in Hérault 2010 it is differentially expressed in Muscovy duck with respect to Pekin
453 duck fed *ad libitum* (Hérault *et al.*, 2010). Another gene involved in triglyceride metabolism is Glycerol-3-
454 phosphate acyltransferase (*GPAM*), not shown in Table 1, but found differentially expressed in 3 of the 4
455 comparisons depicted in Figure 7 (Appendix S1). In fact, *GPAM* was found significantly up-regulated in
456 RM_F_Ct, BP_F_Ct and BP_F_L groups, and plays a central role in *de novo* lipogenesis, particularly of
457 triglycerides (Figure 8) (Claire D'Andre *et al.*, 2013). It was observed that *THRSP* (alias *SPOT14*) was
458 discovered to have a reported role in *de novo* lipogenesis. Indeed, the *THRSP* expression level in liver is
459 correlated with its ability to synthesize lipids (Desert *et al.*, 2018). Furthermore, *SPOT14* has been shown to
460 be a direct target of the key lipogenic *SREBF1* transcription factor (Wu *et al.*, 2013) with its expression
461 levels being under control of estrogens in chicken (Ren *et al.*, 2017). However, the biochemical mechanism
462 linking *SPOT14* to *de novo* lipogenesis remains unclear. Another gene regulated by *SREBF1* is acetyl CoA
463 carboxylase (*ACACA*) also identified in this study, it is critically important for the synthesis of long chain
464 fatty acids (Resnyk *et al.*, 2017). The *ACACA* gene encodes for an enzyme which catalyses the conversion of
465 acetyl-CoA to malonyl- CoA, the substrate of the *de novo* lipogenesis (Nematbakhsh *et al.*, 2021). We also
466 identified stearoyl-CoA desaturase (*SCD*) that plays an important role in biosynthesis of LC-PUFA via the
467 PPAR signalling pathway. Along with *ACACA*, *SCD* could be regulated by *SREBF1*, with both genes being
468 involved in FA *de novo* biosynthesis (Resnyk *et al.*, 2017). *SCD*, thanks to its Delta-9 desaturase activity, can

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3 469 convert palmitic acid (C16:0) and stearic acid (C18:0) to palmitoleic (C16:1 n-7) and oleic acid (C18:1n-9),
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5 470 respectively. *SCD* is recognized as a gene responsible for FA metabolism and having a significant role in
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7 471 intramuscular fat deposition. The third gene regulated by *SREBF1* is not presented in Table 1, but was
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10 472 significant in the comparison between BP_F_Ct and ROSS birds. This is the *FASN* gene and it is involved in
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12 473 *de novo* FA biosynthesis along with *ACACA* and *SCD* (Nematbakhsh *et al.*, 2021).
13
14 474 In conclusion, we provide a comparison of differential gene expression in two native dual-purpose slow-
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16 475 growing chickens, namely RM and BP, compared to commercial ROSS, fed a diet high in n-3 PUFA and
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18 476 assert that high levels of PUFA precursors in the diet do not result in significant changes in expression of
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21 477 genes involved in FA metabolism, whichever the genetic background or sex studied. **The diet had an effect**
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23 478 **on phenotype as body weight, especially in ROSS birds which were significantly lighter in the L diet.** On the
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25 479 contrary, it is clear that sex is an important factor in FA processes. Indeed, females differentially express
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28 480 elongases and desaturase genes (*ELOVL2*, *FADS1*, *FADS2*, and *SCD*) with a central role in LC-PUFA
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30 481 production, particularly in the BP breed. **The BP breed is also suggested to be the genetic background with**
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32 482 **better capacity for modulation of expression of genes involved in FA metabolism.** The two local breeds
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34 483 compared with ROSS commercial birds, have shown a significantly higher modulation of gene expression in
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37 484 liver, which underlines a greater aptitude of local breeds in FA metabolism and final LC-PUFA production.
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39 485 **Finally, this study has shed light on the capability of local chicken breeds to modulate the expression of**
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41 486 **some of the genes involved in FA metabolism, as well as the gene expression in females compared to**
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43 487 **males. Furthermore, the local breeds had the ability to adapt easily to a different diet without change in**
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45 488 **final weight, enriching the tissue (liver in this case) with n-3, hence having the potential for the**
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48 489 **establishment of high-quality products.**
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6 496 **DATA AVAILABILITY STATEMENT**

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8 497 RNAseq raw data are available in NCBI (BioProject ID PRJNA865899).
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36 743 TABLES

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39 745 **Table 1: Differentially expressed genes** shared by BP and RM female groups when compared to ROSS birds.
40 746 **Genes are related to Figure 7 and 8 and Appendix S6.**
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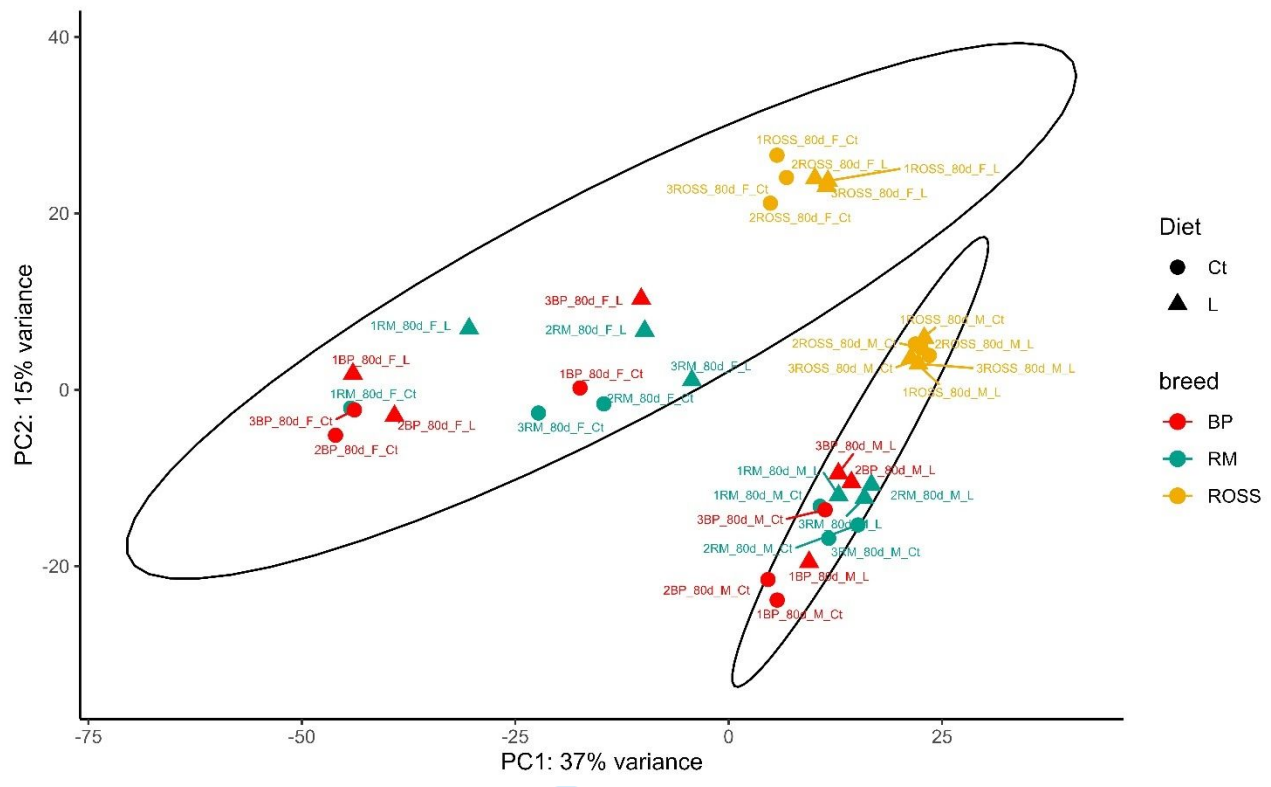
Gene ID	Gene name	Functions	Reference
ABHD12B	<i>Abhydrolase Domain-Containing Protein 12B</i>	Hydrolyzation of the very long chain lysophosphatidylserine lipids	(Z. Li <i>et al.</i> , 2021)
ABHD5	<i>Abhydrolase Domain Containing 5</i>	Crucial gene for fat mobilization and functions as a protein cofactor of ATGL	(Ouyang <i>et al.</i> , 2016)
ACACA	<i>Acetyl-CoA Carboxylase Alpha</i>	Catalysing carboxylation of acetyl-CoA to malonyl-CoA, which is the rate-limiting step in fatty-acid synthesis	(Pirany <i>et al.</i> , 2020)
CHIA	<i>Chitinase</i>	Chinase activity	(Tabata <i>et al.</i> , 2017)
CTSEAL	<i>Cathepsin E-A-like protein</i>	Process egg yolk proteins	(M. Bourin <i>et al.</i> , 2012)
CYP2C45	<i>Cytochrome P450 Family 2 Subfamily C Member 45</i>	Promotes hepatic steatosis by inducing glycolysis-related genes and PPAR pathway	(Zhao <i>et al.</i> , 2019)
RP11-400G3.5	<i>cytochrome P450 2C21-like (CYP2C21L)</i>	Unknown	(Watanabe <i>et al.</i> , 2013)
DCX	<i>Doublecortin</i>	Member of a family of microtubule-associated proteins that are required for neuronal migration during cortical development	(Vermillion <i>et al.</i> , 2014)
LOC101747680	<i>C-factor-like</i>	-	

LOC101749589	-	-	-
LOC107050519	-	-	-
LOC107053670	-	-	-
LOC107053691	-	-	-
LOC112532382	-	-	-
LOC112532439	-	-	-
LOC426220	<i>Avidin-related protein 6-like</i>	Correlated with egg yolk	(Gloux <i>et al.</i> , 2019)
MTTPL	<i>Microsomal triglyceride transfer protein-like</i>	Secretion and lipid metabolism	(Gloux <i>et al.</i> , 2019)
SCD	<i>Stearoyl-CoA desaturase</i>	Desaturation of palmitic and stearic acid to palmitoleic acid (C16:1) and oleic acid (C18:1), respectively	(Nematbakhsh <i>et al.</i> , 2021)
THRSP	<i>Thyroid hormone responsive</i>	Transcription factor involved in control of lipogenic enzymes	(Cui <i>et al.</i> , 2018)
TMEM30CP	<i>Transmembrane protein 30C</i>	Unknown	-
TNFRSF14	<i>Tumor necrosis factor receptor superfamily, member 14</i>	Mediated apoptosis, leading the cancer cells to escape during the immune process	(Guo <i>et al.</i> , 2020)
WFDC8	<i>WAP four-disulphide core domain protein 8</i>	Correlated with egg yolk	(Gloux <i>et al.</i> , 2019)
ZP1	<i>Zona pellucida sperm-binding protein 1</i>	Female fertility	(Bausek <i>et al.</i> , 2004)

FIGURES

Figure 1. Principal component analysis (PCA) of all samples. Colours define different genetic background (BP = Bionda Piemontese; RM = Robusta Maculata; ROSS = Ross 308); shapes represent different diet (Ct = Control diet; L = Linseed diet). The two clusters represent the sex: males (M) in the smallest cluster on bottom right, and females (F) in top left.

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Figure 2. Effect of diet on the percentage of total n-6 (a), total n-3 (b) and total PUFA (c) in relation to total FA in liver tissue. All the comparisons were assessed by experimental groups (same genetic background and sex, but different diet). *P < 0.05, ***P < 0.001 (post hoc Tukey's test). The data are presented as mean ± standard error.

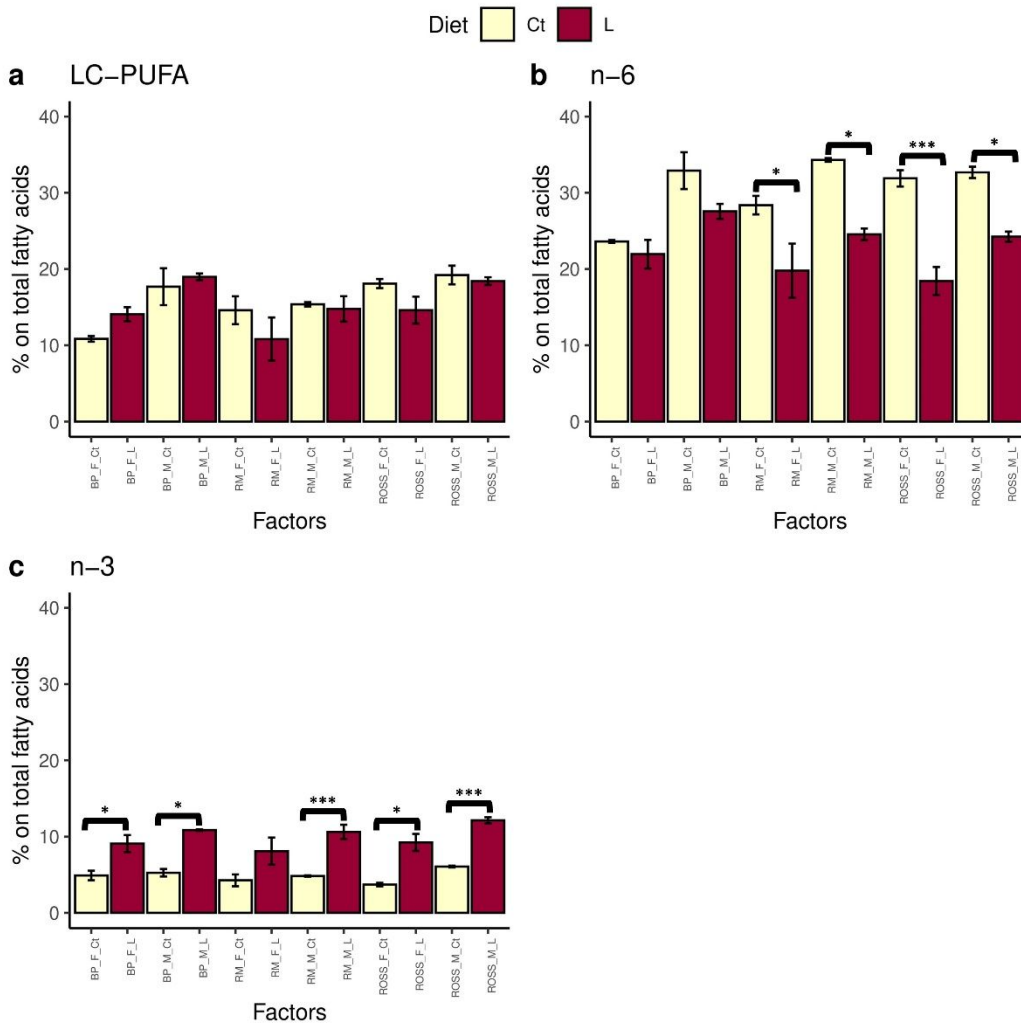
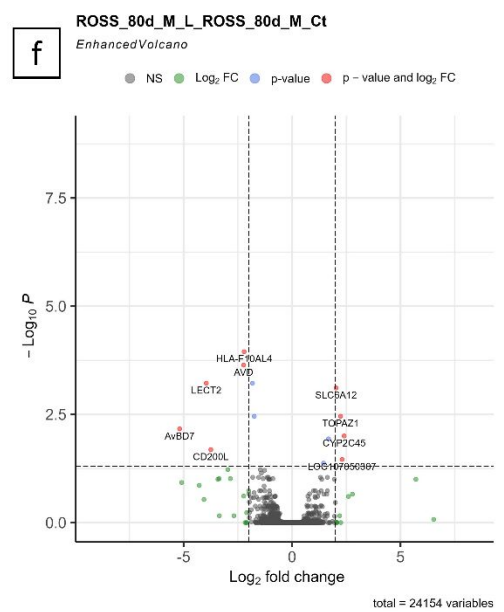
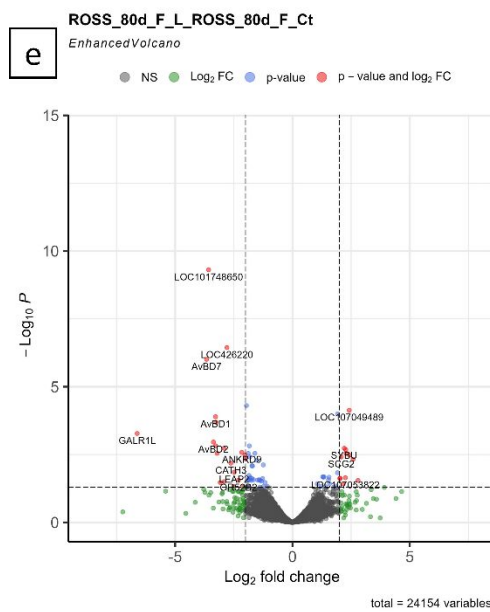
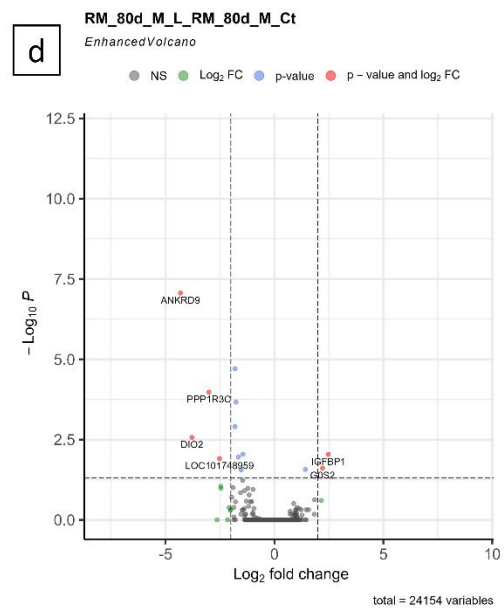
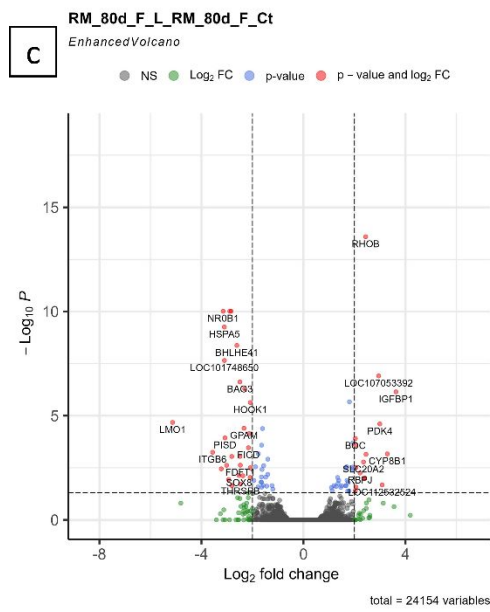
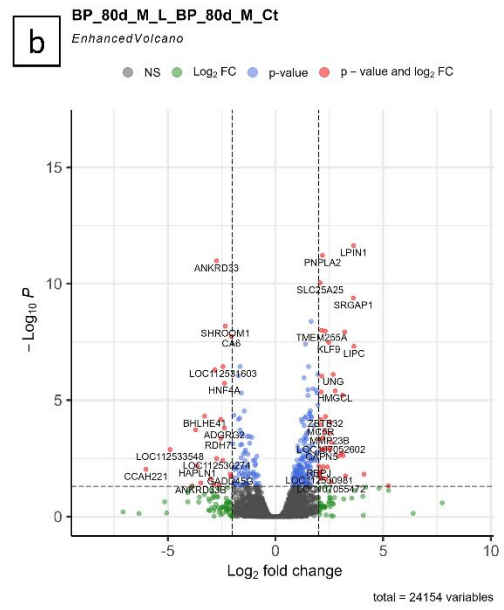
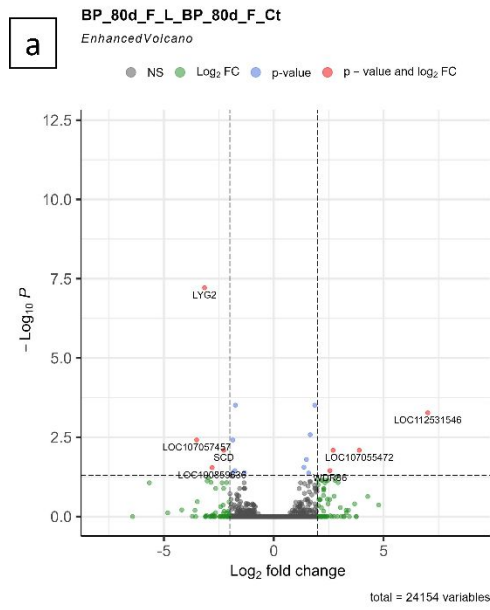


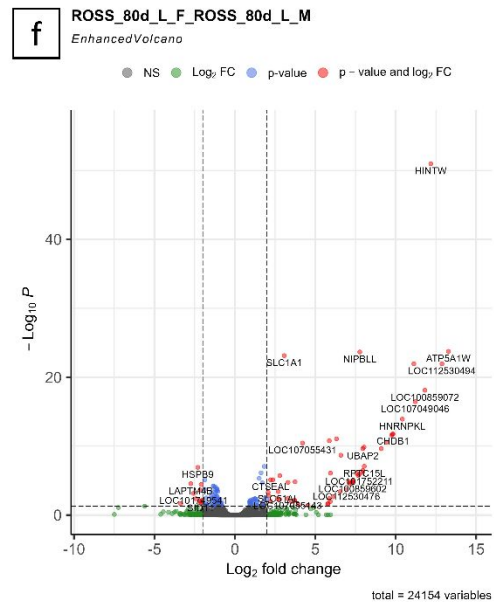
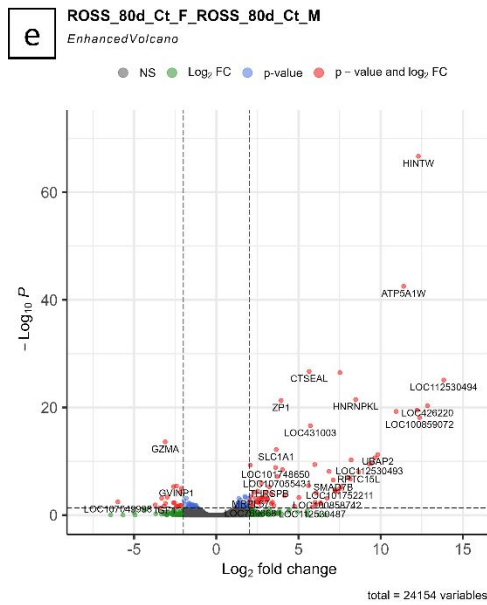
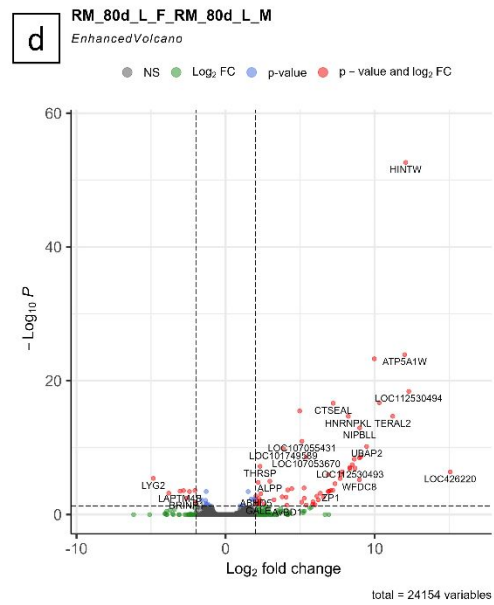
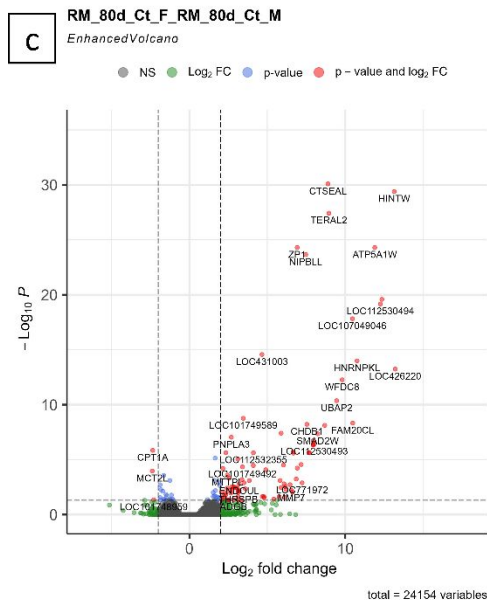
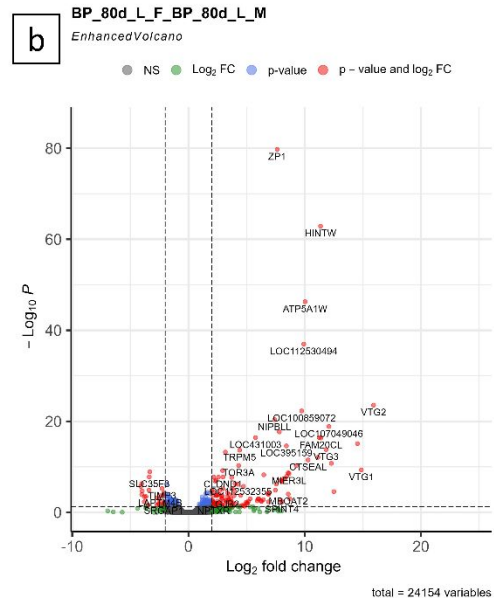
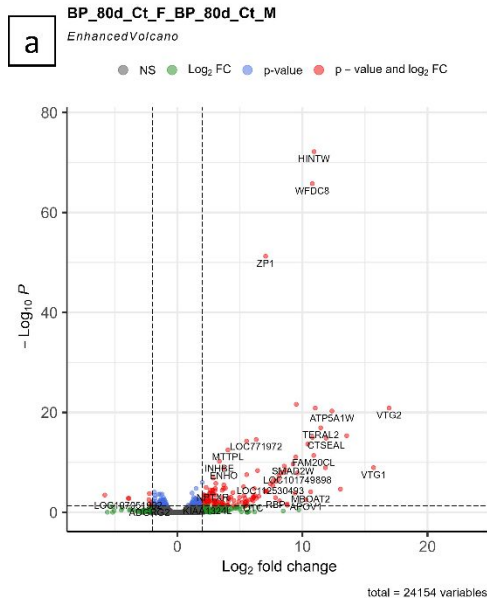
Figure 3. Volcano plot of differentially expressed genes (DEGs) between the Linseed diet (L) and Control diet (Ct) groups. Note: The red dots indicate genes with a P-adjusted value < 0.05 and $-2 > \text{Log}_2$ fold change > 2 . The red dots on the right quadrant of the figures are up-regulated in Linseed diet, the ones on the left are down-regulated. The blue dots represent the genes that reached the P-adjusted value < 0.05 but Log_2 fold change value is higher than -2 and lower than 2. The green dots represent the genes whose difference in expression level between the Ct and L groups did not reach significance ($\text{padj} > 0.05$).



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3 797 **Figure 4.** Volcano plot of differentially expressed genes (DEGs) between the Female (F) and Male (M)
4 798 groups. Note: The red dots indicate genes with a P-adjusted value < 0.05 and $-2 > \text{Log}_2$ fold change > 2 . The
5 799 red dots on the right quadrant of the figures are up-regulated for female, the ones on the left are down-
6 800 regulated. The blue dots represent the genes that reached the P-adjusted value < 0.05 but **Log₂ fold change**
8 801 **value is higher than -2 and lower than 2**. The green dots represent the genes whose difference in
9 802 expression level between the F and M groups did not reach significance ($\text{padj} > 0.05$).

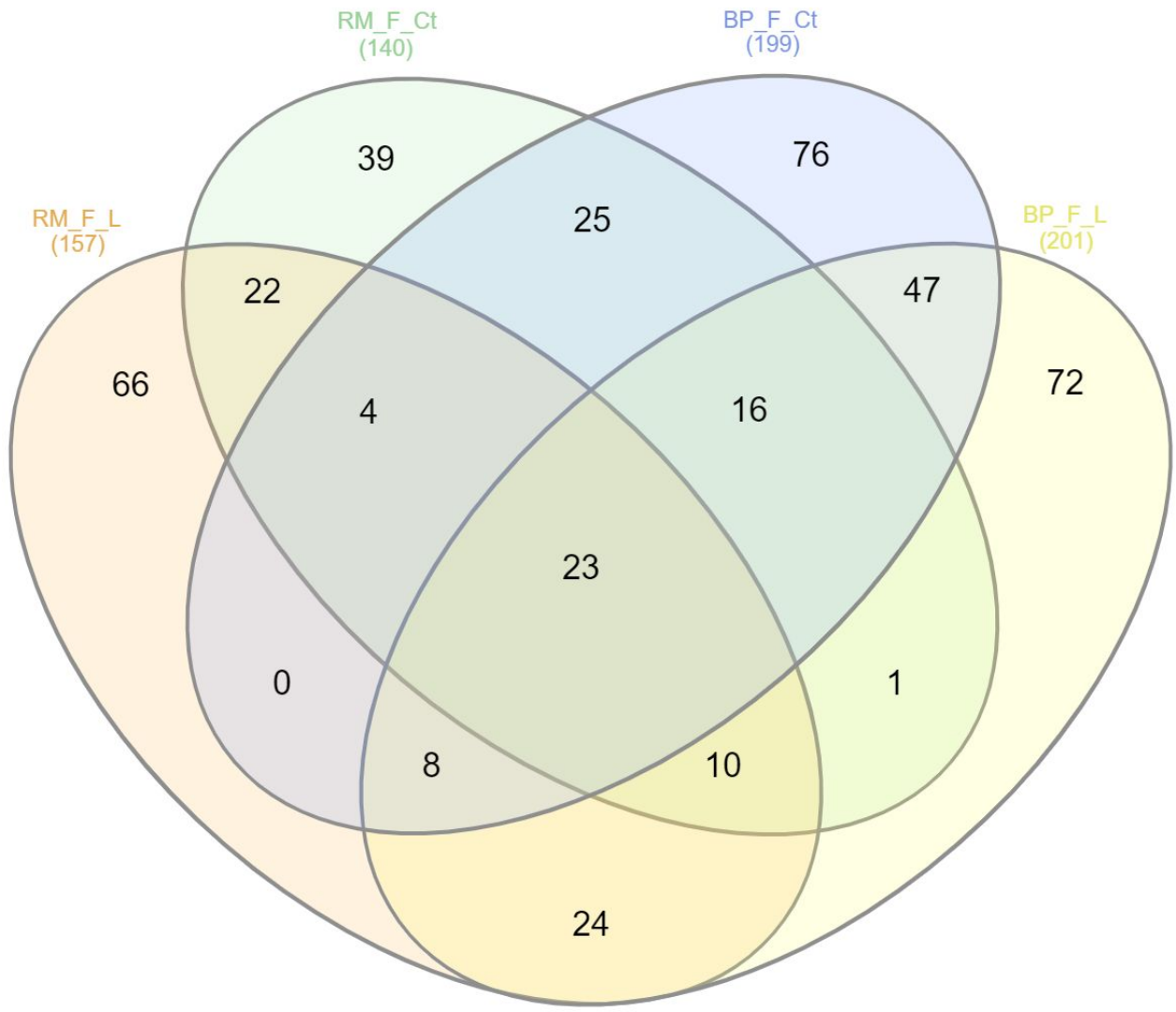
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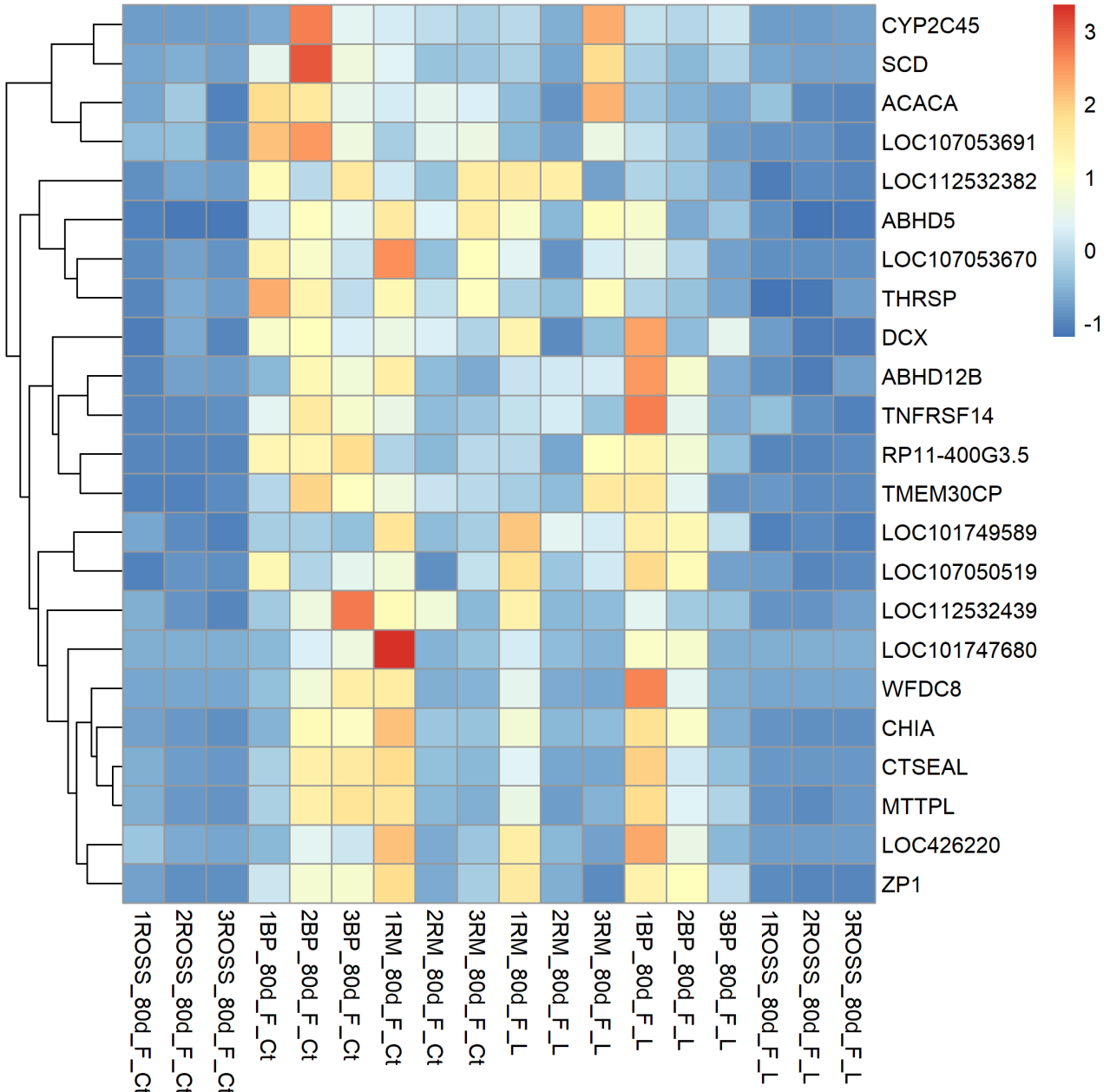
Figure 7. Venn diagram of up-regulated genes in Bionda Piemontese and Robusta Maculata birds compared to Ross308 (ROSS) chickens in female experimental groups.



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Figure 8. Heatmap showing the K-means clustering of transformed expression values for the 23 genes reported in common between female experimental groups in **Figure 7**. Red represents higher expression and blue represents lower expression.



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3 843 **SUPPLEMENTARY MATERIALS:**
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6 844 **Table S1.** Formulation and chemical analysis of chicken diet
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8 845 **Table S2.** **Outline** of experimental design. Birds reared for each experimental group. In brackets the liver
9 846 samples used for the RNA extraction.
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11 847 **Table S3.** Summary of contrasts made in differential expression analysis divided by Diet, Sex, Genetic
12 848 Background. In bold the groups used as experimental, with up and down regulated genes referred to them.
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14 849 **Table S4.** Representation of all genes involved in fatty acids metabolism found as differentially expressed in
15 850 all experimental groups and divided by the belonging contrast
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17 851 **Figure S1.** Step by step bioinformatic procedures adopted in the present study.
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19 852 **Figure S2.** Effect of diet (Ct = control, L = linseed) on Live Body Weight in g. ***P < 0.001 (post hoc Tukey's
20 853 test).
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22 854 **Figure S3.** GeneMANIA report of 23 genes (dashed circles) reported in common between female
23 855 experimental groups in Figure 7. Non-dashed circles represent genes commonly co-expressed with the 23
24 856 belonging to our dataset. Legends explain biological function in which the genes are involved.
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26 857 **Appendix S1.** DEGs of all the comparison. Six Diet comparisons (up and down regulated genes refer to the
27 858 Linseed group), six Sex comparisons (up and down regulated genes refer to the female group), four BP vs
28 859 ROSS comparisons (up and down regulated genes are referred to the BP group) and four RM vs ROSS
29 860 comparisons (up and down regulated genes refer to the RM group),
30 861 In each comparison is reported just genes with a P-value > 0.05. All sheets report Gene ID; baseMean as the
31 862 average of the normalized count values, dividing by size factors, taken over all samples; log2FoldChange
32 863 indicates how much the gene or transcript's expression seems to have changed between the experimental
33 864 and control groups. This value is reported on a logarithmic scale to base 2; lfcSE as the standard error
34 865 estimate for the log2 fold change estimate; stat as the value of the test statistic for the gene or transcript;
35 866 P-value of the test for the gene or transcript; and padj as Adjusted P-value for multiple testing for the gene
36 867 or transcript.
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40 868 **Appendix S2.** Gene Ontology for DEGs in Diet contrasts with significant values in Panther. Gene Ontologies
41 869 refer to up-regulated genes in the Linseed diet group. The considered contrast for each gene ontology is
42 870 reported in the title of each sheet. The GO tables are formed by *Gallus gallus* - REFLIST column that shows
43 871 how many genes are involved in a given pathway. Name of the contrast column shows number of genes
44 872 differentially expressed in the contrast belonging to a given pathway. Over/under represented column
45 873 represents whether the pathway is enriched or down-regulated. Fold enrichment column explain how many
46 874 times the pathway is enriched or down-regulated. Raw p-value and FDR (False discovery Rate) columns
47 874 represents the statistical output of the Statistical overrepresentation test in Panther.
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50 876 **Appendix S3.** Gene Ontology for DEGs in Sex contrasts with significant values in Panther and in IPA
51 877 software. Gene Ontologies refer to up-regulated genes in the female sex group. The considered contrast for
52 878 each gene ontology is reported in the title of each sheet. The GO tables are formed by *Gallus gallus* -
53 879 REFLIST column that shows how many genes are involved in a given pathway. Name of the contrast column
54 879 shows number of genes differentially expressed in the contrast belonging to a given pathway. Over/under
55 880 represented column represents whether the pathway is enriched or down-regulated. Fold enrichment
56 881 column explains how many times the pathway is enriched or down-regulated. Raw p-value and FDR (False
57 882 discovery Rate) columns represents the statistical output of the Statistical overrepresentation test in
58 882 Panther. IPA analysis used the Sex comparison for each group. Only pathways with a z-score > 2 (activated)
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885 are reported. Titles of each sheet explain the contrast under study and the pathways detected as activated.
886 Each IPA sheet reports ID and full name of genes involved in a given pathway, prediction of the behaviour
887 of a given gene when the pathway is activated, and log₂ Fold Change value for every gene in that contrast.

Appendix S4. Gene Ontology for DEGs in Genetic background contrasts (Bionda Piemontese vs Ross308) with significant values in Panther and IPA analyses. Gene Ontologies refer to up-regulated genes in the BP breed group. The considered contrast for each gene ontology is reported in the title of each sheet. The GO tables are formed by *Gallus gallus* - REFLIST column that shows how many genes are involved in a given pathway. *Name of the contrast column* showed number of genes differentially expressed in the contrast belonging to a given pathway. *Over/under represented* column represents whether the pathway is enriched or down-regulated. *Fold enrichment* column explain how many times the pathway is enriched or down-regulated. *Raw p-value* and *FDR* (False discovery Rate) columns represents the statistical output of the Statistical overrepresentation test in Panther. IPA analysis used the Breed comparison for each group. Only pathways with a z-score > 2 (activated) are reported.. Titles of each sheet explain the contrast under study and the pathways detected as activated. Each IPA sheet reports ID and full name of genes involved in a given pathway, prediction of the behaviour of a given gene when the pathway is activated, and log₂ Fold Change value for every gene in that contrast.

Appendix S5. Gene Ontology for DEGs in Genetic background contrasts (Robusta Maculata vs Ross308) with significant values in Panther and in IPA analyses. Gene Ontologies refer to up-regulated genes in the BP breed group. The considered contrast for each gene ontology is reported in the title of each sheet. The GO tables are formed by *Gallus gallus* - REFLIST column that shows how many genes are involved in a given pathway. *Name of the contrast column* shows number of genes differentially expressed in the contrast belonging to a given pathway. *Over/under represented* column represents whether the pathway is enriched or down-regulated. *Fold enrichment* column explain how many times the pathway is enriched or down-regulated. *Raw p-value* and *FDR* (False discovery Rate) columns represents the statistical output of the Statistical overrepresentation test in Panther. IPA analysis used the Breed comparison for each group. Only pathways with a z-score > 2 (activated) are reported. Titles of each sheet explain the contrast under study and the pathways detected as activated. Each IPA sheet reports ID and full name of genes involved in a given pathway, prediction of the behaviour of a given gene when the pathway is activated, and log₂ Fold Change value for every gene in that contrast.

Appendix S6. Gene lists of Venn diagram groups reported in Figure 7.

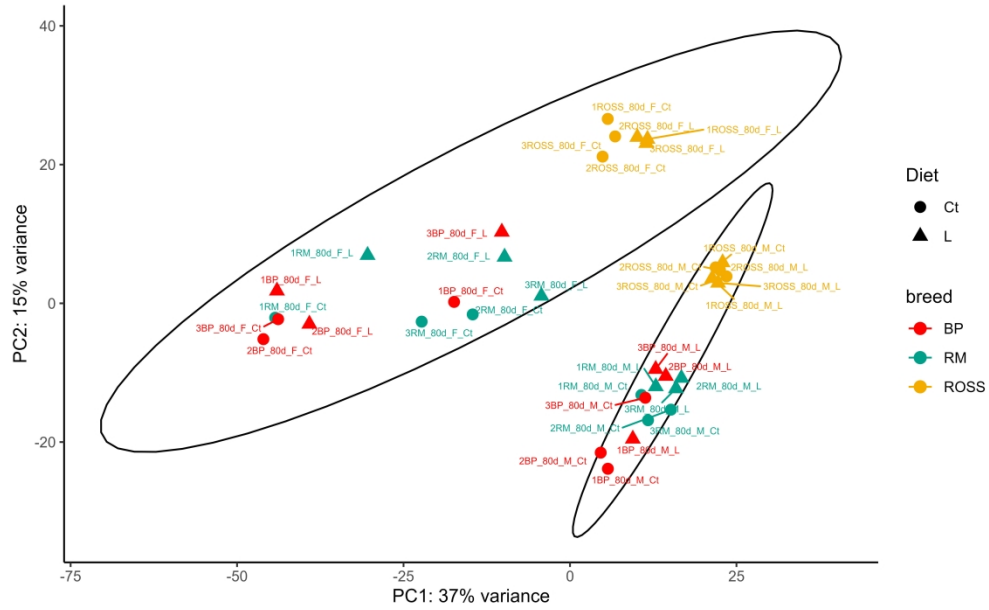


Figure 1. Principal component analysis (PCA) of all samples. Colours define different genetic background (BP = Bionda Piemontese; RM = Robusta Maculata; ROSS = Ross 308); shapes represent different diet (Ct = Control diet; L = Linseed diet). The two clusters represent the sex: males (M) in the smallest cluster on bottom right, and females (F) in top left.

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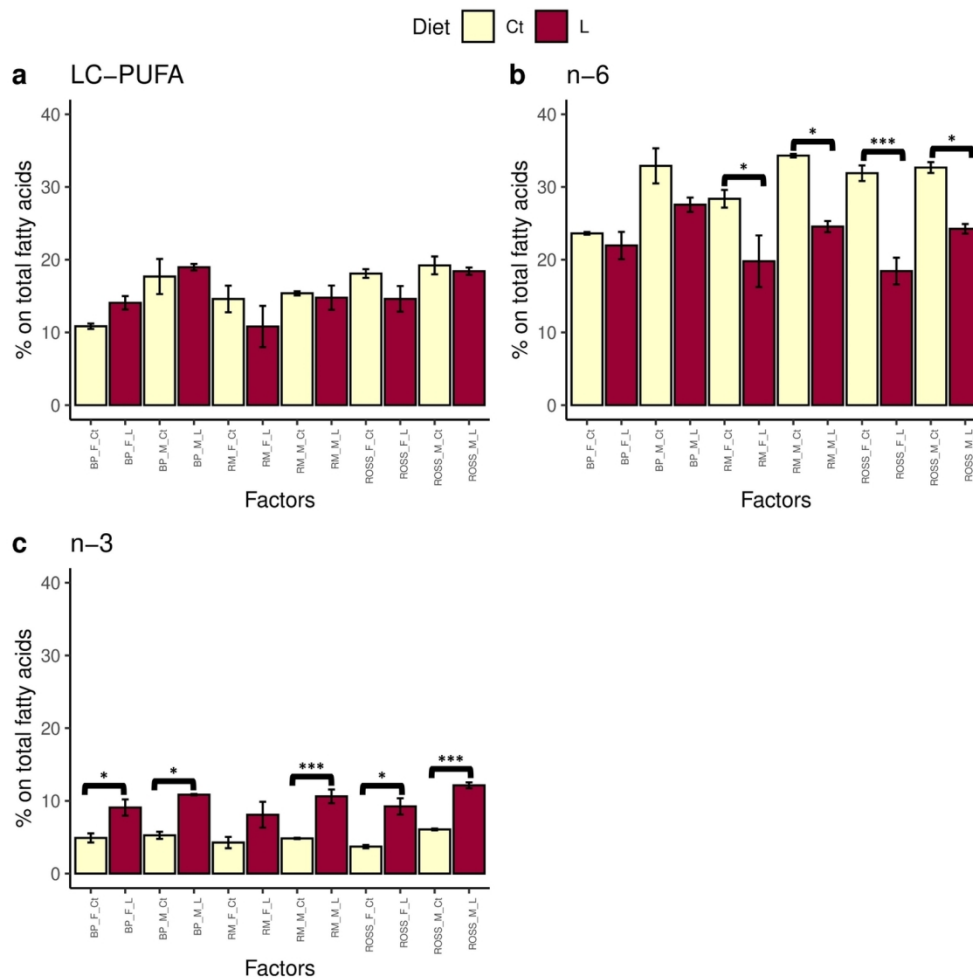


Figure 2. Effect of diet on the percentage of total n-6 (a), total n-3 (b) and total PUFA (c) in relation to total FA in liver tissue. All the comparisons were assessed by experimental groups (same genetic background and sex, but different diet). * $P < 0.05$, *** $P < 0.001$ (post hoc Tukey's test). The data are presented as mean \pm standard error.

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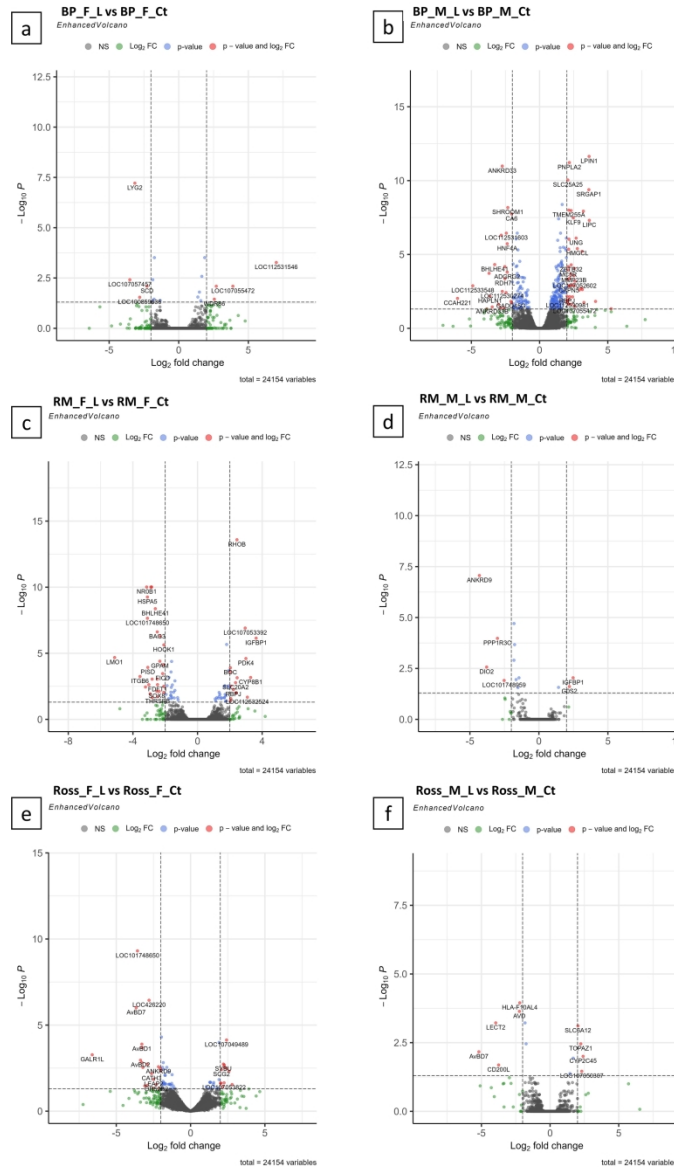


Figure 3. Volcano plot of differentially expressed genes (DEGs) between the Linseed diet (L) and Control diet (Ct) groups. Note: The red dots indicate genes with a P-adjusted value < 0.05 and >2 Log₂ fold change. The red dots on the right quadrant of the figures are up-regulated in Linseed diet, the ones on the left are down-regulated. The blue dots represent the genes that reached the P-adjusted value < 0.05 but Log₂ fold change value is higher than -2 and lower than 2. The green dots represent the genes whose difference in expression level between the Ct and L groups did not reach significance (padj > 0.05).

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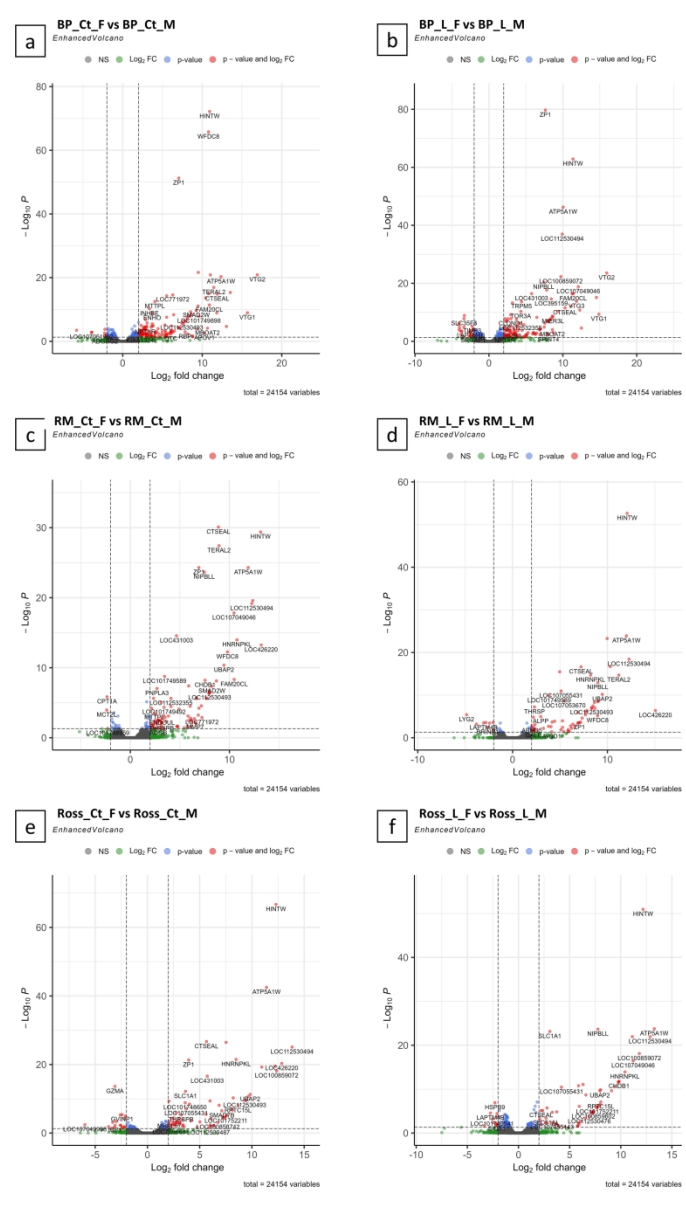


Figure 4. Volcano plot of differentially expressed genes (DEGs) between the Female (F) and Male (M) groups. Note: The red dots indicate genes with a P-adjusted value < 0.05 and $-2 > \text{Log}_2$ fold change > 2 . The red dots on the right quadrant of the figures are up-regulated for female, the ones on the left are down-regulated. The blue dots represent the genes that reached the P-adjusted value < 0.05 but Log_2 fold change value is higher than -2 and lower than 2 . The green dots represent the genes whose difference in expression level between the F and M groups did not reach significance ($\text{padj} > 0.05$).

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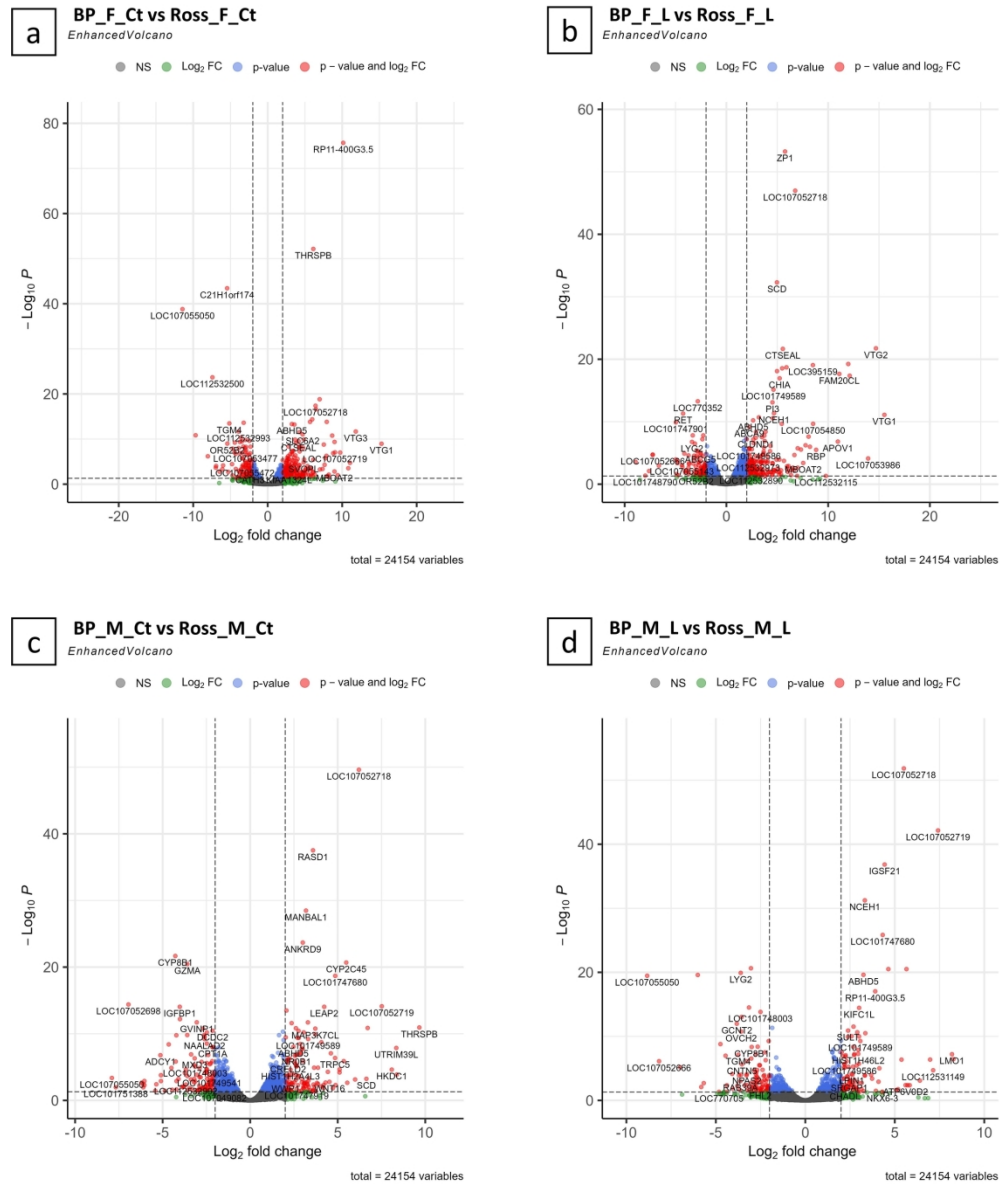


Figure 5. Volcano plot of differentially expressed genes (DEGs) between the Bionda Piemontese (BP) and Ross308 (ROSS) groups. Note: The red dots indicate genes with a P-adjusted value < 0.05 and $-2 > \text{Log}_2$ fold change > 2 . The red dots on the left quadrant of the figures are up-regulated in the Bionda Piemontese breed, the ones on the right are down-regulated. The blue dots represent the genes that reached the P-adjusted value < 0.05 but Log_2 fold change value is higher than -2 and lower than 2. The green dots represent the genes whose difference in expression level between the BP and ROSS groups did not reach significance ($\text{padj} > 0.05$).

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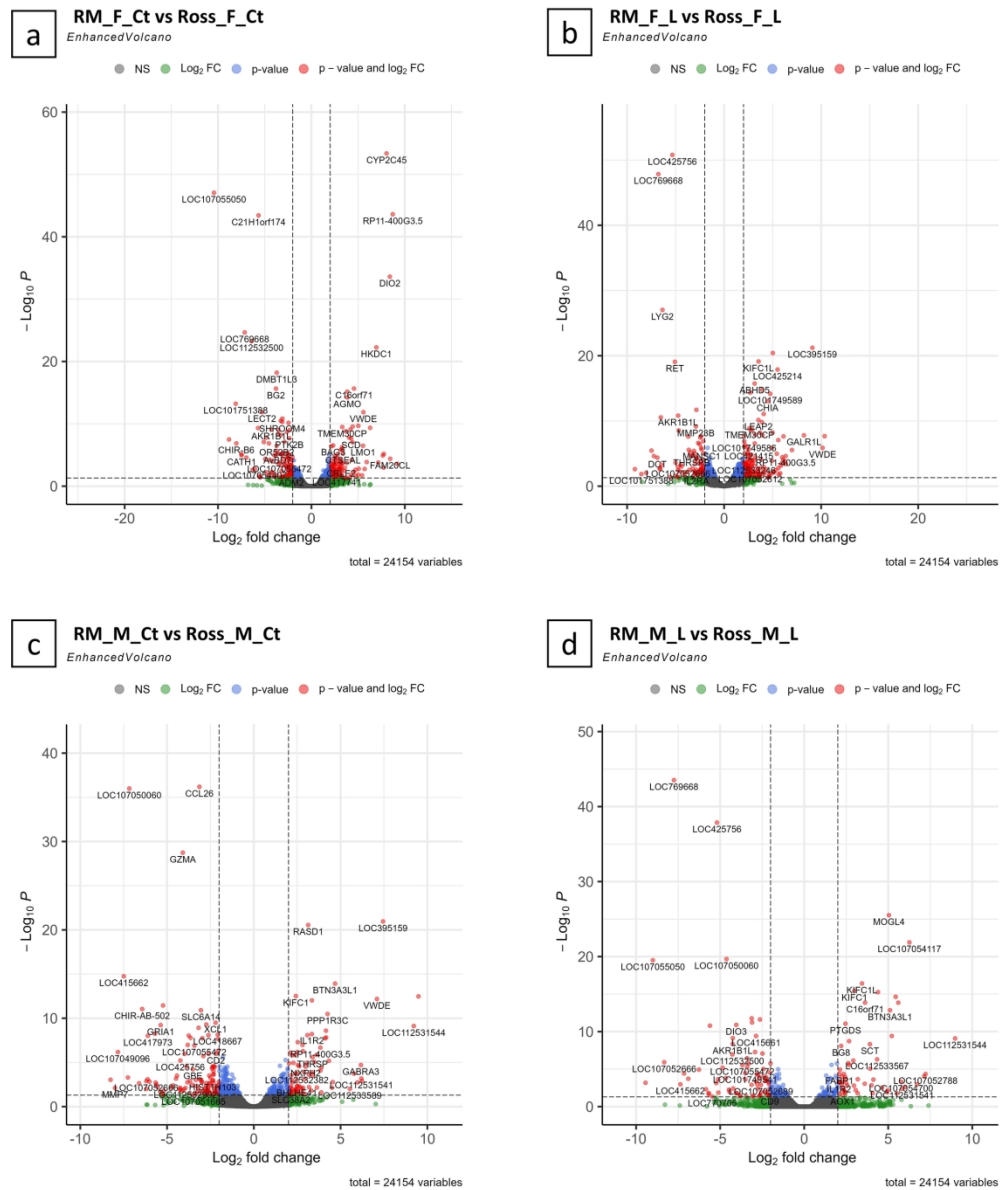


Figure 6. Volcano plot of differentially expressed genes (DEGs) between the Robusta Maculata (RM) and Ross308 (ROSS) groups. Note: The red dots indicate genes with a P-adjusted value < 0.05 and $-2 > \text{Log}_2$ fold change > 2 . The red dots on the left quadrant of the figures are up-regulated in the Robusta Maculata breed, the ones on the right are down-regulated. The blue dots represent the genes that reached the P-adjusted value < 0.05 but Log_2 fold change value is higher than -2 and lower than 2. The green dots represent the genes whose difference in expression level between the RM and ROSS groups did not reach significance ($\text{padj} > 0.05$).

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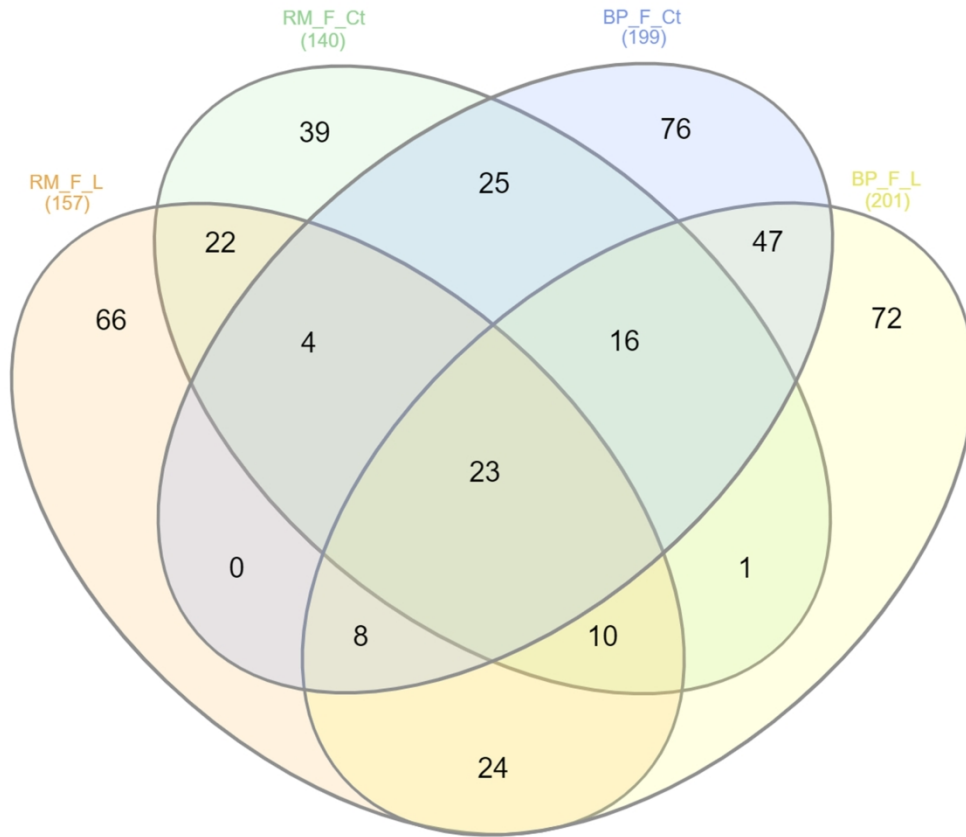


Figure 7. Venn diagram of up-regulated genes in Bionda Piemontese and Robusta Maculata birds compared to Ross308 (ROSS) chickens in female experimental groups.

240x248mm (300 x 300 DPI)

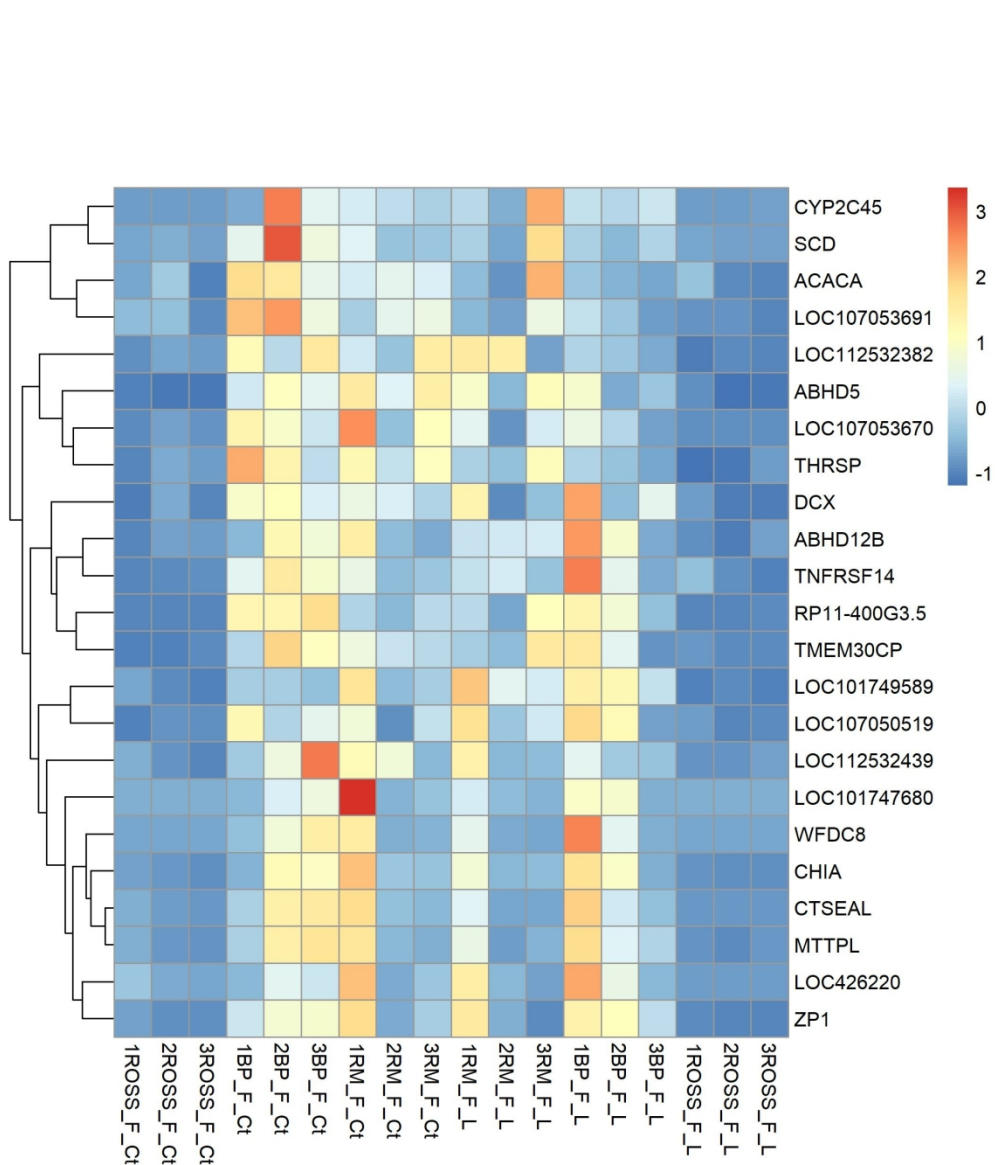


Figure 8. Heatmap showing the K-means clustering of transformed expression values for the 23 genes reported in common between female experimental groups in Figure 7. Red represents higher expression and blue represents lower expression.

177x177mm (300 x 300 DPI)

Table 1: Differentially expressed genes shared by BP and RM female groups when compared to ROSS birds. Genes are related to Figure 7, Figure 8 and Appendix S6.

Gene ID	Gene name	Functions	Reference
ABHD12B	<i>Abhydrolase Domain-Containing Protein 12B</i>	Hydrolyzation of the very long chain lysophosphatidylserine lipids	(Li <i>et al.</i> , 2021)
ABHD5	<i>Abhydrolase Domain Containing 5</i>	Crucial gene for fat mobilization and functions as a protein cofactor of ATGL	(Ouyang <i>et al.</i> , 2016)
ACACA	<i>Acetyl-CoA Carboxylase Alpha</i>	Catalysing carboxylation of acetyl-CoA to malonyl-CoA, which is the rate-limiting step in fatty-acid synthesis	(Pirany <i>et al.</i> , 2020)
CHIA	<i>Chitinase</i>	Chinase activity	(Tabata <i>et al.</i> , 2017)
CTSEAL	<i>Cathepsin E-A-like protein</i>	Process egg yolk proteins	(M. Bourin <i>et al.</i> , 2012)
CYP2C45	<i>Cytochrome P450 Family 2 Subfamily C Member 45</i>	Promotes hepatic steatosis by inducing glycolysis-related genes and PPAR pathway	(Zhao <i>et al.</i> , 2019)
RP11-400G3.5	<i>cytochrome P450 2C21-like (CYP2C21L)</i>	Unknown	(Watanabe <i>et al.</i> , 2013)
DCX	<i>Doublecortin</i>	Member of a family of microtubule-associated proteins that are required for neuronal migration during cortical development	(Vermillion <i>et al.</i> , 2014)
LOC101747680	<i>C-factor-like</i>	-	-
LOC101749589	-	-	-
LOC107050519	-	-	-
LOC107053670	-	-	-
LOC107053691	-	-	-
LOC112532382	-	-	-
LOC112532439	-	-	-
LOC426220	<i>Avidin-related protein 6-like</i>	Correlated with egg yolk	(Gloux <i>et al.</i> , 2019)
MTTPL	<i>Microsomal triglyceride transfer protein-like</i>	Secretion and lipid metabolism	(Gloux <i>et al.</i> , 2019)
SCD	<i>Stearoyl-CoA desaturase</i>	Desaturation of palmitic and stearic acid to palmitoleic acid (C16:1) and oleic acid (C18:1), respectively	(Nematbakhsh <i>et al.</i> , 2021)
THRSP	<i>Thyroid hormone responsive</i>	Transcription factor involved in control of lipogenic enzymes	(Cui <i>et al.</i> , 2018)
TMEM30CP	<i>Transmembrane protein 30C</i>	Unknown	-
TNFRSF14	<i>Tumor necrosis factor receptor superfamily, member 14</i>	Mediated apoptosis, leading the cancer cells to escape during the immune process	(Guo <i>et al.</i> , 2020)
WFDC8	<i>WAP four-disulphide core domain protein 8</i>	Correlated with egg yolk	(Gloux <i>et al.</i> , 2019)
ZP1	<i>Zona pellucida sperm-binding protein 1</i>	Female fertility	(Bausek <i>et al.</i> , 2004)

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2
3 1 **RNAseq reveals modulation of genes involved in fatty acid biosynthesis in chicken**
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6 2 **liver** according to genetic background, sex and diet
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11 4 Perini F.¹, Wu Z.², Cartoni Mancinelli A.¹, Soglia D.³, Schiavone A.³, **Mattioli S.¹**, Mugnai C.³, Castellini C.¹, Smith J.²,
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57 27
58
59 28 **Keywords:** fatty acid metabolism, transcriptome, poultry, local breeds, differentially expressed genes.
60

29 ABSTRACT

30 Increases in chicken production are mainly due to specialised breeds. However, of increasing importance
31 are the local breeds, known for their ability to adapt to the environment and for their unique products.
32 Conventional poultry products contain lower levels of n-3 fatty acids compared to those obtained from
33 local breeds, therefore the aim of this study was to evaluate the modulation of expression of genes
34 involved in LC-PUFA biosynthesis pathways according to genetic background, diet conditions, and sex.
35 Animals from two local breeds and a commercial line were fed different diets: control and experimental
36 diet (10% linseed supplementation). For each breed and diet group, both males and females were reared.
37 The RNA was extracted from 36 liver samples and was sequenced by RNAseq method. Bioinformatic
38 analysis was carried out to find differentially expressed genes (DEGs) from different comparisons between
39 experimental groups. Results showed low impact of diet on DEGs related to fatty acid biosynthesis, but
40 linseed diet increased percentage of n-3 fatty acids of liver. Sex, particularly the female groups, and genetic
41 background determined the differential expression of genes related to LC-PUFA biosynthesis. Specifically,
42 females of local breeds shared 23 up-regulated genes when compared to their respective commercial line
43 groups. Some of the shared genes had a role in *de novo* triglyceride biosynthesis (*MTTPL* and *GPAM*), and in
44 genes involved in *de novo* FA biosynthesis (*ACACA* and *SCD*) was detected. In conclusion, sex and local
45 genetic background appear to have influence on the expression of genes related to LC-PUFA synthesis.

47 INTRODUCTION

48 To date, commercial chicken lines have dominated the poultry market so far, thanks to the fast growth and
49 high production performances. Besides the standard commercial chicken lines, of increasing importance
50 are the local breeds which are able to adapt to the environment where they live (Perini *et al.*, 2021), and
51 for their different quality products compared to conventional ones (Franzoni *et al.*, 2021). Unfortunately,
52 the productivity traits are critical for local breeds, which are instead appreciated for the diversity of their
53 products, which could include economically exploitable traits (Moula *et al.*, 2010). Currently, it is still

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2
3 54 difficult to obtain economic feedback from breeders who decide to raise local breeds because of low
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5 55 competitiveness on productive performances.
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7 56 A potential valorisation strategy is the enrichment in terms of nutritional value of meat and eggs. Enhancing
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9
10 57 the quantity of n-3 fatty acids in chicken meat and eggs through the diet can have important impacts: i)
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12 58 supplementing the fatty acid intake in **the** human diet, as it is usually rich in n-6 and lacking in n-3 fatty
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14 59 acids (Mariamenatu & Abdu, 2021); ii) chickens are able to convert precursors of n-3 in long chain (**> 20**
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16 60 **carbon atoms**) polyunsaturated fatty acids (LC-PUFA); iii) **finally**, eggs and meat enriched with n-3 would
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18
19 61 have increased market **potential** compared to standard animal products.
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21 62 The LC-PUFAs, eicosapentaenoic acid (EPA, 20:5 n-3) and docosahexaenoic acid (DHA, 22:6 n-3) are the
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23 63 main form of n-3. The fatty acid profiles of poultry meat and eggs can be readily enriched in EPA and DHA
24
25 64 through the diet (Cartoni Mancinelli *et al.*, 2022). Although conventional poultry meat contains low levels
26
27
28 65 of EPA and DHA, several factors such as sex, feed, and genetic background may influence their quantity. In
29
30 66 particular, genetic background is reported in the literature as a crucial factor: higher concentrations of n-3
31
32 67 LC-PUFA are synthesized by local breeds compared to commercial lines (Cartoni Mancinelli *et al.*, 2021).
33
34 68 The application of a diet rich in **precursors** of n-3 (α -linolenic acid: 18:3n-3, ALA) can affect the amount of
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36 69 LC-PUFA in meat and eggs, and may change the expression level of genes involved in the process of lipid
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39 70 biosynthesis and elongation. **In mammals**, these two metabolic pathways, mainly occur in the adipose
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41 71 tissue, while in chicken the majority (90%) takes place in the liver (Nematbakhsh *et al.*, 2021). To elucidate
42
43 72 the complete expression profile of liver genes, RNAseq provides a useful tool. This transcriptomic approach
44
45 73 can accurately study tissue transcriptomes with high resolution and depth. Increasingly, the nutrigenomic
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47
48 74 approach is used to investigate the effect of diets on metabolic processes also in poultry (Soglia *et al.*,
49
50 75 2022). A recent study of the liver transcriptome after folic acid supplementation in the diet of broiler
51
52 76 chickens found that peroxisome proliferator activated receptor (PPAR) signalling is the pathway most
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54 77 activated by the enriched diet (Zhang *et al.*, 2021).
55
56
57 78 In recent years, many studies tried to elucidate which genes are principally involved in the biosynthesis
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59 79 process of LC-PUFA. Studies focused on the evaluation of the presence/activity of elongase of very long
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3 80 chain fatty acid (*ELOVL*) and fatty acid desaturase (*FADS*) enzymes (Cartoni Mancinelli *et al.*, 2022; Lee *et*
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5 81 *al.*, 2016). Although **recently** many studies have highlighted some pathways involved in the biosynthesis of
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7 82 LC-PUFAs (e.g. *PPAR* signaling), **there remains** a lack of scientific research in this area (Mihelic *et al.*, 2020).
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9
10 83 The aim of this study was to determine **the modulation of expression of genes** involved in LC-PUFA
11
12 84 biosynthesis pathways, according to genetic background (local breed vs commercial line), diets, and sex.
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14 85

16 86 **MATERIALS AND METHODS**

18 19 87 ***Animal ethics***

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21 88 Birds were raised, handled and processed according to the European legislation for the protection of
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23 89 chickens kept for meat production (European Commission, 2007), the protection of animals at the time of
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25 90 killing (European Commission, 2009) and the protection of animals used for scientific purposes (European
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27 91 Commission, 2010). The experimental protocol was positively evaluated and approved by the Ethical
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29 92 Committee of the University of Perugia (ID: 62700_15/07/2020).
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32 33 94 ***Experimental design***

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35 95 The trial was carried out in the experimental section of the Department of Agricultural, Food and
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37 96 Environmental Sciences (University of Perugia, Italy). Specifically, three different chicken genetic
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39 97 **backgrounds** were used: Robusta Maculata (RM) and Bionda Piemontese (BP) as local **breeds** and Ross 308
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41 98 (ROSS) as a commercial line. **Both RM and BP are dual-purpose breeds, in particular BP shows higher**
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43 99 **aptitude for laying eggs (200 eggs/year vs 150 eggs/year of RM) (www.pollitaliani.it).** One-day old chicks of
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45 100 **each sex** were housed in an **environmentally**-controlled poultry facility and vaccinated against coccidiosis,
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47 101 infectious bronchitis, Marek's Disease, Newcastle Disease and Gumboro. Until 21 days of age, all the
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49 102 chickens received the same starter diet. **Then each genetic background was divided into two groups: one**
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51 103 **fed a standard diet (control group, Ct), the second fed the standard diet supplemented with 10 % extruded**
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53 104 **linseed (experimental group, L). The two diets were formulated to meet the nutritional recommendations**
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55 105 **of Ross 308 birds (Table S1) (Aviagen, 2019). Three replicates/group were formed, each representing both**
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3 106 male and female chickens (5 males + 5 females/each replicate). Chickens were raised in different indoor
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5 107 pens at the same stocking density (5 chickens/m²) and the temperature was set according to the age of the
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7 108 birds (20-32°C; Relative Humidity: 65-72%). Water and feed were provided *ad libitum*. At 81 days of age,
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10 109 live body weight was recorded (g) for all the birds which were later slaughtered in a commercial
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12 110 slaughterhouse. From each replicate, two birds (one for each gender) were selected. Table S2 schematically
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14 111 explains the experimental design used in the present study.

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19 113 **Tissue collection and RNA extraction:**

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21 114 Liver tissue was collected from all the animals. Around 1 g of tissue was collected in a 2 mL cryogenic vial
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23 115 (Corning® Inc., Corning, NY), instantaneously frozen in liquid nitrogen, and stored at -80° C until RNA
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25 116 extraction. The remaining part of liver tissue was collected and stored at -20°C for further analysis of fatty
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27
28 117 acids (FA) profile.

29
30 118 At this stage, RNA was extracted only from liver tissue. Samples were prepared from 36 animals, with three
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32 119 biological replicates for each experimental group (Table S2). RNA isolation was performed with a
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34 120 NucleoSpin RNA Mini kit for RNA purification (Macherey-Nagel, Germany) as recommended by the
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36
37 121 manufacturer, starting from 30 mg of tissue. A homogenization step was carried out using an Omni Tissue
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39 122 Homogenizer (TH) - Omni, Inc) in ice. DNA contamination was removed with a DNase enzyme (included in
40
41 123 the kit) during the incubation step. RNA concentration was measured with a Qubit 3.0 fluorometer (Life
42
43 124 Technologies) and Qubit RNA HS Assay Kit (Life Technologies) according to manufacturer instruction. Total
44
45 125 RNA (~1.5 µg/sample) was sent to Genewiz (South Plainfield, NJ, USA) for quality check, library preparation
46
47
48 126 and Illumina sequencing. For testing RNA quality, the Agilent (Santa Clara, CA) 2100 Bioanalyzer Nano Kit
49
50 127 was used, and all the samples showed RNA integrity (RIN) numbers ≥6.5 (data not shown). Library
51
52 128 preparation was carried out through polyA + selection and paired-end (PE) sequencing was run on an
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54 129 Illumina NovaSeq System that generated 150 bp PE reads.

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59 131 **Bioinformatic analysis**

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3 132 RNAseq data were quality checked by FastQC software (Brown *et al.*, 2017). Paired qualified reads were
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5 133 mapped to the chicken reference genome (GRCg6a, accession number: PRJNA13342) using STAR software
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7 134 (Dobin *et al.*, 2015) with default parameters. The mapping rates for each sample were assessed using
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9
10 135 Samtools 'flagstat' command (Heng Li *et al.*, 2009). The mapped BAM file was then used in featureCounts
11
12 136 software that, assigning raw alignments to annotated genes, generates the raw expression count file used
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14 137 for gene expression quantification (Liao *et al.*, 2014). An overview of genomic distance in the dataset was
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16 138 performed through Principal Component Analysis using the ggplot2 R package. Differential expression (DE)
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19 139 analysis was performed using the DESeq2 R package (Love *et al.*, 2014). The P-value adjustment was made
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21 140 using Benjamini and Hochberg's approach for controlling the False Discovery Rate (FDR). Genes with a P-
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23 141 adjusted value < 0.05 and $-2 > \text{Log}_2$ fold change (LFC) > 2 were used as thresholds for significant DE by
24
25 142 DESeq2. Visualization of contrast between different experimental groups was performed by
26
27
28 143 EnhancedVolcano package in R (Blighe *et al.*, 2022). The overlap of significant genes among experimental
29
30 144 groups was assessed with Venn diagrams built with the Interactivenn web tool (Heberle *et al.*, 2015).
31
32 145 Ingenuity Pathway Analysis (IPA, QIAGEN Redwood City, www.qiagen.com/ingenuity) was used to identify
33
34 146 gene ontology pathways, and regulatory networks to which DE genes belong, as well as upstream
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36
37 147 regulators. Ingenuity Pathway Analysis can transform a set of genes into a number of relevant networks
38
39 148 based on comprehensive records maintained in the Ingenuity Pathways Knowledge Base. **According to DEG**
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41 149 **input, IPA software calculates a z-score value for each biological pathway. When a pathway has been**
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43 150 **categorized with a z-score > 2 , it resulted as an activated pathway.** Moreover, to confirm the IPA results,
44
45 151 gene ontologies (GO) were examined using the Panther (v17.0) database according to the Statistical over-
46
47
48 152 representation test. Fisher's test was used to correct for False Discovery Rate (Figure S1) (Mi *et al.*, 2019).
49
50 153 **Differential expression (DE) analysis was performed in individual contrasts according to diet, sex and**
51
52 154 **genetic background, separately. In particular we evaluated six diet contrasts, six contrasts for sex and four**
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54
55 155 **contrasts for each local breed against the ROSS birds (Table S3).**

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59 157 **Fatty acids (FA) profile**
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3 158 The fatty acids profile was determined from the same samples used for the RNAseq. Lipids were extracted
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5 159 from 5 g of liver/sample based on the methods described in Folch *et al.* (1957). The fatty acids were
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7 160 identified in the form of their methyl esters using a Varian Gas Chromatograph (CP-3800) and a DB wax
8
9
10 161 capillary column (25 mm ϕ , 30 m long). The establishment of each fatty acid was done in relation to the
11
12 162 retention time with respect to fatty acid methyl ester standards (FAME, Sigma-Aldrich, Bellefonte, PA). The
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14 163 relative quantity of each fatty acid present in the liver was calculated using heneicosanoic acid (C21:0;
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16 164 Sigma-Aldrich) as the internal standard. Data were expressed as % of total FA. The average amount of each
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19 165 FA was used to calculate the sum of total PUFA of the n-3 and n-6 series. After using the function of the
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21 166 Shapiro test (for normality test), data were analysed with ANOVA and with Tukey's *post-hoc* test in R
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23 167 software. All results are expressed as mean \pm SEM, with the level of significance set at $p < 0.05$.
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27 169 RESULTS

30 170 *Sequencing data analysis*

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33 171 Average mapping rate was 85.11% across all samples, 98% of which were seen to be properly paired.
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35 172 Among all the samples, the average quality score for each base was $> Q30$ and the average GC content in
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37 173 sequenced samples was around 52% (data not shown). After all quality checks, all 36 samples were taken
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40 174 forward for further analysis. In order to shed light on genetic similarity, a principal component analysis
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42 175 (PCA) plot was computed and visualized using R software. This showed genetic distance between the
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44 176 samples according to the comparison of Principal Component 1 and 2 (PC1 and PC2) (Figure 1). Genetic
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46 177 diversity was appreciable, especially between the two main clusters seen in Figure 1 representing male and
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48 178 female birds. The male cluster showed less genetic variance compared to the female one, and the ROSS
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51 179 birds grouped closest to each other within each cluster, because of their standard genetic background. On
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53 180 the contrary, the two local breeds showed more genetic variance within the experimental groups and were
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55 181 clearly separated from the ROSS cluster.
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60 183 *Body weight and fatty acids profile of animals*

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3 184 The effect of diet was evaluated according to the results from live body weight (Figure S2) and the
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5 185 percentage of n-3 and n-6 PUFA, and LC-PUFA (Figure 2). The diet had no statistical impact on the body
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7 186 weight at slaughtering of the two local breeds, either in males or females. Instead, the L diet significantly
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10 187 decreased the body weight in ROSS (male and female).

11
12 188 Figure 2 represent the n-6 and n-3 PUFA proportion in liver, and the sum of LC-PUFA. Figure 2a shows the
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14 189 percentage of LC-PUFA was not statistically affected by the diet. On the contrary, the n-6 PUFA (ranging
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16 190 from 18% to 34% approximately) showed higher level than n-3 PUFA (from 4% to 12). Moreover, the
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19 191 control diet generally increased the n-6 percentage in both sexes of ROSS and in RM_M. On the contrary,
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21 192 the L diet significantly enhanced the n-3 level in all groups, excepted for RM females.

193 194 ***Differentially expressed genes (DEGs)***

195 Each experimental group was compared against one another to check differential gene expression. This was
196 done on the basis of: diet, sex, and genotype (Table S3). In particular, we were interested in comparing
197 local breeds to the broiler (ROSS) genotype. Appendix S1 lists all significantly differentially expressed genes
198 in each comparison. Moreover, all genes related to FA metabolism, and which were found to be
199 differentially expressed, are reported in Table S4.

200 201 ***Diet contrast***

202 For the evaluation of the possible effect of diet on gene expression between sexes and genetic background,
203 the linseed diet group was compared to the control diet group within each sex and within each genetic
204 background. Experimental groups formed by “genetic background” and “sex” were taking in consideration
205 individually. Linseed diet groups have been used as experimental groups, hence the up and down-regulated
206 genes and the GO analyses referring to them. Figure 3 shows the volcano plots resulting from each
207 comparison. Figures 2a-b showed the DEGs detected for the female (BP_F) and male (BP_M) Bionda
208 Piemontese experimental groups respectively. In BP_F, the diet played a limited role in differentiating gene
209 expression level, indeed Figure 3a had only eight differentially expressed genes with a P -value < 0.05, of
210

1
2
3 210 which some involved in fatty acid metabolism (e.g. *LYG2*, *SCD*), being found down-regulated with the L diet.
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5 211 Other important genes involved in FA synthesis (*ACACB*, *ACACA*, *FASN*) had a significant *P*-value but had
6
7 212 lower LFC values (between -2 and 2) (Appendix S1).
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10 213 In the BP_M group, the diet contrast showed 280 significant genes, and the Gene Ontology study confirmed
11
12 214 that the most influenced pathways were cellular lipid metabolic process (GO:0044255) and lipid metabolic
13
14 215 process (GO:0006629) (Appendix S2). Regarding the Robusta Maculata breed, volcano plots for DEGs in
15
16 216 females (RM_F) and males (RM_M) are shown in Figures 2c-d respectively. Contrary to the BP birds, the
17
18 217 RM_F comparison showed more DEGs than the RM_M group, namely 83 and 13 genes. Gene ontology
19
20 218 analysis in each group showed that the RM_F group had biological processes which were significantly
21
22 219 enriched in the L diet, although not directly related with FA biosynthesis, and males did not show anything
23
24 220 significant (i.e. 'de novo' protein folding (GO:0006458)) (Appendix S2). Finally, the last genetic background
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26 221 evaluated for the effect of diet was the commercial ROSS hybrid (Figure 3e-f). In both female and male
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28 222 groups, a limited number of DEGs were identified (72 and 13, respectively). This is similar to the GO
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30 223 biological annotations associated with FA metabolism. None of the six diet contrasts resulted in a
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32 224 significantly activated biological pathway in IPA analysis.
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39 226 **Effect of sex**

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41 227 The effect of sex on gene expression was assessed as follows: the experimental groups compared within
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43 228 the same genetic background and diet groups but differing by sex. Female groups were taken as the
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45 229 experimental group, hence the up and down regulated genes and GO analysis are referred to "F" group in
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47 230 each contrast. Looking at Figure 4a, it is easy to appreciate the numerous DEGs up-regulated in females. For
48
49 231 each BP comparison, as shown in Figures 4a-b, the GO annotations on up-regulated genes clarify the sexual
50
51 232 dimorphism in the most typical way: cellular response to estrogen stimulus (GO:0071391) and response to
52
53 233 estrogens (GO:0043627) (Appendix S3). When comparing the ROSS broiler birds, and examining the linseed
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55 234 diet group (L), significant biological enrichment was found in organo-nitrogen compound biosynthetic
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57 235 process (GO:1901566), for example (Appendix S3). With regard to FA pathways, in BP birds there was
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3 236 activation of lipid pathways in females, namely lipid transport (GO:0006869) and lipid localization
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5 237 (GO:0010876). This result from Panther was also confirmed using IPA software. Fatty acid metabolism was
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7 238 seen to be significantly enhanced in the two female groups of the BP breed. The significance of fatty acid
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10 239 concentration pathway in RM chickens treated with linseed diet was also indicated (Appendix S3). In
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12 240 particular, the data in Appendix S3 shed light on the genes that were differentially expressed in females
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14 241 within the experimental group. The females expressed some genes strictly related to FA metabolism
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16 242 differentially to males within respective groups. For examples: *ELOVL2* in BP_Ct, *ELOVL2*, *FADS1* and *FADS2*
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18 243 in BP_L, and *SCD* in RM_L.

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Genetic background comparison

One of the aims of this study was to assess the impact of breed and genetic makeup on the profile of LC-PUFA expression. In particular, the comparison between local breeds and the commercial line (same sex and diet) was examined.

Figure 5 represents the DEGs from the BP vs ROSS comparison. BP breed represented the experimental group, which up and down regulated genes and GO analysis are referred to. Figures 5a-b show females treated with control and linseed diet respectively. According to results from Panther, many pathways related to lipid metabolism were up-regulated in the BP breed, such as fatty acid biosynthetic process (GO:0006633) and long-chain fatty acid metabolic process (GO:0001676) (Appendix S4). In the female control diet group, the up-regulated genes found through IPA were related to FA metabolism: *SCD*, *CYP1A1*, *FASN*, *ACACA*, *THRSP*, *FAR1*, *FADS1* (Appendix S4). Figure 5b showed the results for a genetic background contrast with females fed the linseed diet. The results from GO analysis showed a large number of lipid related pathways, the most important being lipid metabolic process (GO:0006629). Some of the genes related to this, which were more highly expressed in BP relative to the commercial line include *FABP3*, *SCD*, *GPAM*, *LPIN1*, *SREBF2* and *ACACA* (Appendix S4). Meanwhile, Figures 5c-d exhibit the contrast of BP with ROSS birds, in males fed the two different diets. With the control diet, BP males showed higher expression of genes involved in fatty acid metabolic process (GO:0006631) in comparison with ROSS. BP males fed the

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3 262 linseed diet do not show DEGs significant in FA metabolism (e.g. *FASN*, *ACACA*, *ACACB*) when compared
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5 263 with ROSS (Appendix S4), but amino acid biosynthetic processes are highlighted in the GO analysis.
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7 264 We next wanted to compare Robusta Maculata (RM) vs Ross308 (ROSS) groups. The experimental groups
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10 265 consisted of the RM breed, hence up and down regulated genes and GO analysis are referred to as “RM”
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12 266 breed groups. The volcano plots are presented in Figure 6, of which the first two (Figure 6a-b) represented
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14 267 the contrast in females, with control and experimental diets, respectively. Figure 6a showed 391 genes with
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16 268 a P-value lower than 0.05, of which 234 genes had a value of $-2 > \text{LFC} > 2$, and 121 genes in RM with $\text{LFC} > 2$.
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19 269 The Panther GO analysis clearly showed that within the up-regulated genes, fatty acid metabolic process
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21 270 (GO:0006631) was clearly enhanced, with similar pathways activated in female linseed diet groups. IPA
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23 271 analysis showed significance of LC-PUFA synthesis in RM birds fed the control diet and synthesis of FA in
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25 272 RM birds fed the linseed diet (Appendix S5). The RM female groups shared DEGs clearly involved in FA
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28 273 pathways, such as *SCD*, *ACACA*, and *THRSP*. On the contrary, in RM males (Figures 6c-d), no pathways
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30 274 related to FA appeared as significantly involved in birds fed either diet. Instead, immune pathways were
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32 275 active in Ct diet birds, while there were no significant pathways in birds fed the supplemented diet
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35 276 (Appendix S5).

36 37 277 38 39 278 **Comparison of local vs commercial females**

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41 279 According to the results from the sex comparison previously described, females, for the most part,
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43 280 appeared more specialized for FA production compared to males. Moreover, in the comparison of genetic
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46 281 backgrounds, all the female groups from local breeds showed higher activation of FA-related pathways
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48 282 compared to commercial hybrid birds. In order to understand which DEGs were common to the local
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50 283 breeds, a Venn diagram was generated (Figure 7). Here we took into consideration only the female birds,
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52 284 and only the genes differentially expressed in local breeds (BP and RM) compared to ROSS. Interestingly, 23
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55 285 genes were commonly up-regulated among local breeds (Appendix S6). A heat map was generated to
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57 286 graphically represent the 23 shared genes and their expression level among all female samples (Table 1)
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59 287 (Figure 8). GeneMania software was used to identify any connection between them, with *GPAM* being
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3 288 suggested as being directly related to *THRSP* and *SCD*. In fact, we saw *GPAM* as being up-regulated in both
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5 289 BP female groups and in RM females fed the control diet, but not the linseed diet (Figure S3).
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10 291 DISCUSSION

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13 292 Many studies have reported how the diet can modulate the expression of genes in different tissues (Sevane
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15 293 *et al.*, 2014; Szalai *et al.*, 2021). Here, we fed animals with a diet rich in the precursor of n-3 LC-PUFA. We
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17 294 designed the experiment assuming that diet plays a key role in changing lipid metabolism in the liver.
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19 295 Indeed, vitamin supplementation (Niu *et al.*, 2009) taurine (He *et al.*, 2019) and zinc oxide nanoparticles
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21 296 (Ramiah *et al.*, 2019) have been used to modulate gene expression of FA related genes, whereas the
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23 297 supplementation of dietary PUFA is widely used to increase LC-PUFA concentration in animal products of
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25 298 different animal species, such as dairy cows, rabbit and turkey (Castellini *et al.*, 2022; Kliem *et al.*, 2019;
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27 299 Szalai *et al.*, 2021). Chickens are usually an ideal target for diet supplementation, especially linseed based
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29 300 supplements (Head *et al.*, 2019; Jing *et al.*, 2013; Sevane *et al.*, 2014; Zhang *et al.*, 2021).
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33 301 In the present study, the different diets affected the body weight of commercial birds. This result is
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35 302 explained by the fact that during the experimental trial, the ROSS chickens showed problems of adaptation
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37 303 to the linseed diet, consuming around 7% less quantity of the L diet with respect to the Ct diet, both in
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39 304 males and in females. In other studies was already reported how extruded linseed diet could lead to lower
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41 305 body weight gain and lower feed intake in broiler (Avazkhanloo *et al.*, 2020; Anjum *et al.*, 2013). This
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43 306 outcome indicated once again how the local breeds can easily adapt themselves to different environmental
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45 307 factor (e.g. alternative diet sources), while commercial lines encounter difficulties. In the present study, n-
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47 308 3, n-6 and LC-PUFA of liver were examined. LC-PUFA, which are directly related to the desaturation and
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49 309 elongation ability, was not affected by diet, conversely to n-6 and n-3 levels. The percentage of n-6 was
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51 310 generally higher in the Ct diet because the Ct diet is higher in linoleic acid (LA), which is the precursors of n-
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53 311 6 PUFA. On the other hand, the L diet positively affected the n-3 PUFA, and indeed the n-3 is significantly
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55 312 higher in the L diet than in the corresponding Ct groups (Figure 2). Both these results are commonly found
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57 313 in the literature (Hang *et al.*, 2018; Head *et al.*, 2019; Meineri *et al.*, 2018). Head *et al.* (2019) described the
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3 314 modulation of FA profile in a given tissue through the diet, and also provided insights into low effect of the
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5 315 diet on the gene expression. The fact that the trend of LC-PUFA in the different experimental groups was
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7 316 similar confirms this assertion. Thus, it seems that the entity of LC-PUFA produced by liver is mainly due to
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10 317 the level of dietary precursors. On the same time, the type of precursor furnished α -linolenic (ALA, n-3) or
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12 318 linolenic (LA, n-6) determined an alternative accumulation of the same PUFA series in the liver. Indeed, we
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14 319 found that the dietary supplementation of linseed affected expression of genes involved in FA metabolism
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16 320 in only one group (BP_M group – Appendix S2). The most highly expressed genes in the linseed diet group
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19 321 included *PPARGC1A*, and *LPIN1*. *PPARGC1A* is a co-activator of *PPAR γ* which is a transcription factor that
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21 322 participates in induction and stimulation of fat-specific genes and fatty acid bio-synthesis (Wang *et al.*,
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23 323 2017). *LPIN1* is involved in synthesis and transport of triacylglycerol, a major constituent of chicken lipids
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25 324 (Desert *et al.*, 2018), and also found to be over expressed in a chicken line selected for intramuscular fat
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28 325 content together with *PPAR γ* (Liu *et al.*, 2020). The remaining contrasts (n=5) for diet did not show enriched
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30 326 pathways, and for this reason, we can assert that diet is the factor with lowest impact on gene expression
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32 327 of fatty acid metabolism. Although no GO enrichment was discovered regarding FA metabolism in the other
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34 328 5 diet contrasts, there are some interesting genes down-regulated with the L diet belonging to the BP_F
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37 329 and RM_F groups. Indeed, in BP_F the L diet showed down-regulation of *LYG2*, *ACACA*, *FASN*, *SCD* and in
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39 330 RM_F, down-regulation of *PLIN2* and *GPAM*, all important genes in FA metabolism and discussed later in
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41 331 this section. This is also in agreement with Head *et al.* (2019) who ascertained that the effect of linseed on
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43 332 the expression of 14 genes involved in FA metabolism in chicken liver was low. We therefore saw differing
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46 333 modulation of FA metabolism in liver between genetic backgrounds and sexes through diet.
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48 334 This study also evaluated the effect of sexual dimorphism and its impact on FA metabolism. A clear
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50 335 separation between male and female birds is appreciable in Figure 1, underlying a divergent level of
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52 336 expression in genes between the two groups. The background of sex effect on FA metabolism was
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55 337 investigated by Poureslami *et al.* (2010b) where they described the poor effect of sex on final concentration
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57 338 of MUFA and PUFA of the n-3 and n-6 series (Poureslami *et al.*, 2010a). Elsewhere, Lopez-Ferrer *et al.* used
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59 339 dietary linseed oil, finding that abdominal fat percentage in male chickens was significantly lower than in
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3 340 the female chickens (López-Ferrer *et al.*, 2001). This was corroborated in human where the differential
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5 341 ability to activate PPAR α together with the well-documented effect of sex hormones on lipid metabolism
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7 342 may highlight sex as a significant factor in plasma FA levels (Thifault *et al.*, 2013). Differences between male
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10 343 and female chickens have never been thoroughly described and hence our interest in including sex effect in
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12 344 our analysis. BP_F, in both diet groups, showed a greater enrichment for estrogen pathways because it was
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14 345 the most egg-layer type chicken in the dataset. Indeed, estrogen in laying hens stimulates the liver to
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16 346 synthesize fatty acids for egg enrichment (Hanlon *et al.*, 2022). As reported by Ayres *et al.* (2013),
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19 347 differential expression of genes related to the female W chromosome are essential for sex determination in
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21 348 chickens. Some of them (*NIPBL* and *UBAP2*) have also been found to be central to sex determination in
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23 349 Japanese quail (Caetano-Anolles *et al.*, 2015). Regarding lipid metabolism, some genes were found to be
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25 350 more highly expressed in females with respect to males. This is the case for *ELOVL2* (in both BP contrasts),
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28 351 *FADS1*, *FADS2* (in BP_L) and *SCD* (in RM_L, and in both ROSS groups) that code for enzymes directly
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30 352 involved in desaturation and elongation of FA in the diet, such as ALA (18:3n-3), resulting in LC-PUFA (Head
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32 353 *et al.*, 2019). Interestingly, the results can be appreciated in the comparison of females with males in the BP
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34 354 breed. Besides the finding of elongases and desaturase-related genes being more highly expressed in
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37 355 females, other genes directly involved in FA metabolism were also identified. For instance, *APOB* plays a
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39 356 vital role in the assembly and secretion of triacylglycerol-rich lipoprotein in the liver of egg-laying chickens
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41 357 (Ma *et al.*, 2017), in line with the BP breed having a major aptitude in laying, especially when compared to
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43 358 ROSS (meat type) and RM birds (dual-purpose). Moreover, in the BP_L group, the female highly expressed
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46 359 genes of the fatty acid-binding protein (*FABP*) family, particularly *FABP3* and *FABP1*, which are considered
47
48 360 biomarkers for intramuscular fat content. Bongiorno *et al.* (2022) reported that the saturated fatty acid
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50 361 (SFA) of breast meat was mainly influenced by gender. On the other hand, in the RM breed the higher
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52 362 female expression of *THRSP* strongly suggests a control of lipogenic targets (Resnyk *et al.*, 2017). Activated
53
54 363 estrogen pathways could explain differences in genes related to FA in females. In 2017 Zhang *et al.* found
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56
57 364 three miRNAs involved in down-regulation of genes related to FA metabolism. More specifically, estrogen
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59 365 eliminates the suppressive effect of miRNAs on the target gene *ELOVL5*. Interestingly, estrogen suppresses
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3 366 the host *SLIT2* gene, thus decreasing the expression of intronic miR-218-5p to promote hepatic synthesis of
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5 367 long-chain polyunsaturated fatty acids in the liver (Zhang *et al.*, 2017). Although *ELOVL5* was not found to
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7 368 be directly overexpressed in females in the present study, many genes functionally related to it (i.e.
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10 369 *ELOVL2*, *SCD*, *FADS1*, *FADS2*) were identified. It is possible that estrogen pathways could regulate FA
11
12 370 metabolism in general through the down-regulation of miR-218-5p, which regulates the genes related to FA
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14 371 metabolism, thus enhancing FA biosynthesis. Interestingly, it is the same miRNA (miR-218-5p, regulated by
15
16 372 estrogen) that has a role in *FADS1* regulation in liver (Hong Li *et al.*, 2016), and *ELOVL2* regulation in chicken
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18 373 muscle (Zhang *et al.*, 2018). A more comprehensive evaluation of estrogen modulation via miRNA of FA
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21 374 metabolism should be investigated in further studies.
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23
24 375 In the present study, we observed that the genetic background was the variable having the largest effect on
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26 376 differential gene expression. Initially, we expected differing expression of genes for FA metabolism
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28 377 between the local breeds and in particular when compared to ROSS broilers. In the BP breed, a huge effect
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30 378 was identified in all four experimental groups when compared against ROSS birds. According to gene
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32 379 expression rates, the two female groups for each diet were the most divergent. These results suggested not
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34
35 380 only that the females had higher expression of genes involved in FA metabolism, but also that the BP
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37 381 female exhibited higher expression of genes responsible for FA metabolism, when compared to the ROSS
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39 382 females. In particular, the female control diet group showed an enrichment for genes involved in the long-
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42 383 chain fatty acid metabolic process (GO:0001676) supported by both GO analysis and the related genes
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44 384 *ELOVL2*, *SCD*, *FADS1*, *FADS2*, *THRSP*, *FABP3* and *LPIN1*.
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46 385 With males belonging to the local breeds, an effect was only observed within the BP breed, suggesting the
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48 386 importance of FA processing in this breed. RM females also showed interesting results with regards the FA
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51 387 process. On the contrary, the comparison between RM_M and ROSS_M did not show significant differences
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53 388 in genes involved in the FA processes. This lower effect in males of local breeds could be due to the
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55 389 estrogen stimulus that was strongly activated in females from local breeds and results in the difference in
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57 390 expression of genes related to LC-PUFA. Comparing all the results from contrasts of BP and RM female
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59 391 groups against the ROSS birds highlights 23 genes that are shared between the female local breeds (Figure

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3 392 **7 and Figure 8, Appendix S6**). Some of these are still uncharacterized (*LOC101747680*, *LOC107053670*,
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5 393 *LOC426220*, *LOC101749589*, *LOC112532382*, *LOC107050519*, *LOC107053691*, *LOC112532439*), but three of
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7 394 them are already reported in the literature with regards FA metabolism related functions. The
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10 395 *LOC101747680* gene codes for a C-like protein and it is located on chromosome 11, and *LOC107053670* is a
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12 396 non-coding RNA situated on chromosome 6, each with roles which are still unclear. *LOC426220* is located
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14 397 on the W sex chromosome, and is an avidin-related protein 6-like, and with all other avidin-like molecules
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16 398 localized on chromosome Z, **suggesting** a sex-specific regulation. Moreover, two studies have found this
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18 399 gene strongly correlated with egg yolk, vitelline membrane, and white coloured eggs (Gloux *et al.*, 2019;
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20 400 Niskanen *et al.*, 2005). Regarding eggs and fertility, another three genes were found differentially expressed
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22 401 in local breed females: *WDFC8*, *CTSEAL*, and *ZP1* (Table 1). *WDFC8* is clearly related to “WAP four-
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24 402 disulphide core domain proteins”. Members of this family are involved in various aspects of mucosal
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26 403 immunity (Wilkinson *et al.*, 2011). *WDFC8* was also found highly overexpressed in the liver of laying hens
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28 404 (Gloux *et al.*, 2019) and has been shown to protect egg yolk precursors from proteolytic
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30 405 activities/inactivation in the plasma, on their way from the liver to the growing oocytes (Marie Bourin *et al.*,
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32 406 2012). Moreover, *WDFC8* has been found as a unique protein in egg yolk of chicken (Farinazzo *et al.*, 2009).
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34 407 The avian-specific *CTSEAL* gene was found to be overexpressed in liver of laying hens and the translated
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36 408 protein shares large sequence similarity with cathepsin D, suggesting *CTSEAL* as an accessory of cathepsin D
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38 409 in the processing of egg yolk precursors (Bourin *et al.*, 2012; Gloux *et al.*, 2019). *ZP1* gene was also up-
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40 410 regulated in the liver of hens, where the protein is synthesized and then transported to the ovary to be
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42 411 inserted in the perivitelline membrane surrounding the oocyte (Gloux *et al.*, 2019). Here, it plays a
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44 412 fundamental role in the first interactions between spermatozoa and the oocyte (Bausek *et al.*, 2004). These
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46 413 three genes were differentially expressed in liver of BP and RM birds, with an explanation being that these
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48 414 **are** multi-purpose breeds and are thus more suitable chickens for egg production than the ROSS broilers.
49
50 415 Table 1 **and Figure 8** also show other genes not known to be involved in a particular pathway or showing a
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52 416 functional relationship with others. An example is chitinase (*CHIA*), which is a gene coding for a major
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54 417 protease-resistant glycosidase with a physiological role as a digestive enzyme that breaks down chitin-
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3 418 containing organisms in the chicken gastrointestinal tract (Tabata *et al.*, 2018). Another example is
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5 419 Doublecortin (*DCX*) – a member of a family of microtubule-associated proteins that are required for
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7 420 neuronal migration during cortical development (Vermillion *et al.*, 2014). Tumor Necrosis Factor Receptor
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10 421 Superfamily, Member 14 (*TNFRSF14*) was also found differentially expressed. It mediates apoptosis
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12 422 signalling, and can allow cancer cells to escape the immune process (Guo *et al.*, 2020).
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14 423 The remainder of genes found differentially expressed in females of local breeds (compared to ROSS), are
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16 424 involved in FA metabolism. The *ABHD12B* and *ABHD5* genes are part of the α/β hydrolase domain-
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18
19 425 containing family appointed to mobilization of lipids. In particular, *ABHD12B* has been highlighted by Li et al
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21 426 (2020) as a hydrolyser of very long chain lysophosphatidylserine lipids in human cells (Li *et al.*, 2021).
22
23 427 *ABHD5* the most well-characterized gene of its family codes for a protein cofactor of the ATGL enzyme and
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25 428 stimulates triacylglycerol hydrolase activity. Ouyang et al (2016) reported that overexpression of *ABDH5*
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28 429 markedly decreased the triglyceride content of preadipocytes in chicken (Ouyang *et al.*, 2016). This
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30 430 mechanism is modulated via the two above-mentioned genes and might explain the question of why native
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32 431 breed hens raised in free-range systems have less fat but higher polyunsaturated fatty acids in their meat
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34 432 muscles (Sokołowicz *et al.*, 2016). We discovered other genes differentially expressed in local female breeds
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36 433 belonging to the cytochrome P450 family were also highlighted: *CYP2C45* and *RP11-400G3.5*. Cytochrome
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39 434 P-450 2C45 (*CYP2C45*) was the most highly expressed cytochrome P-450 isoform in chicken liver and is a
40
41 435 unique isoform in avian species (Watanabe *et al.*, 2013). Moreover, Zhao *et al.* (2019) have shown that
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43 436 *CYP2C45* was overexpressed in liver of overfed geese, hence promoting hepatic steatosis. Eventually, this
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45 437 resulted in the up-regulation of some genes involved in FA metabolism (i.e. *PK* and *ALOX5*) acting via *PPAR*
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48 438 pathways (Zhao *et al.*, 2019). *RP11-400G3.5* is reported as a *CYP2C21*-like pseudogene and is the closest
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50 439 gene to *CYP2C45* on chromosome 6 (Watanabe *et al.*, 2013). In fact, both of these genes showed similar
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52 440 function, especially in arachidonic and linolenic acid metabolism in KEGG pathways (data not shown).
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54 441 Furthermore, in local breeds we found significantly more expression in two genes having a role in de novo
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56 442 triglyceride biosynthesis (*MTTPL* and *GPAM*), two genes involved in *de novo* FA biosynthesis (*ACACA* and
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443 *SCD*), confirmedly involved in desaturation of FA in comparison to ROSS birds. All those genes have been
444 seen to be involved in FA or lipid metabolism (Figure S3).

445 In Table 1 we report the microsomal triglyceride transfer protein-like (*MTTPL*) gene located on
446 chromosome 6, which is an orthologue of *MTTP* on chromosome 4. *MTTP* is responsible for the assembly
447 and subsequent secretion of very low density lipoproteins from hepatocytes and controls the incorporation
448 of triglycerides into apolipoprotein B (Liu *et al.*, 2016). Moreover, *MTTPL* was found overexpressed in liver
449 of chickens fed with a diet rich in LC-PUFA precursor (Liu *et al.*, 2019; Tesseraud *et al.*, 2014). In the present
450 study, *MTTPL* was differentially expressed in the liver of local breeds, underlining their capacity for FA
451 metabolism. This was not the first time that genetic background has been shown to be a factor in liver
452 expression of *MTTPL*: in Hérault 2010 it is differentially expressed in Muscovy duck with respect to Pekin
453 duck fed *ad libitum* (Hérault *et al.*, 2010). Another gene involved in triglyceride metabolism is Glycerol-3-
454 phosphate acyltransferase (*GPAM*), not shown in Table 1, but found differentially expressed in 3 of the 4
455 comparisons depicted in Figure 7 (Appendix S1). In fact, *GPAM* was found significantly up-regulated in
456 RM_F_Ct, BP_F_Ct and BP_F_L groups, and plays a central role in *de novo* lipogenesis, particularly of
457 triglycerides (Figure 8) (Claire D'Andre *et al.*, 2013). It was observed that *THRSP* (alias *SPOT14*) was
458 discovered to have a reported role in *de novo* lipogenesis. Indeed, the *THRSP* expression level in liver is
459 correlated with its ability to synthesize lipids (Desert *et al.*, 2018). Furthermore, *SPOT14* has been shown to
460 be a direct target of the key lipogenic *SREBF1* transcription factor (Wu *et al.*, 2013) with its expression
461 levels being under control of estrogens in chicken (Ren *et al.*, 2017). However, the biochemical mechanism
462 linking *SPOT14* to *de novo* lipogenesis remains unclear. Another gene regulated by *SREBF1* is acetyl CoA
463 carboxylase (*ACACA*) also identified in this study, it is critically important for the synthesis of long chain
464 fatty acids (Resnyk *et al.*, 2017). The *ACACA* gene encodes for an enzyme which catalyses the conversion of
465 acetyl-CoA to malonyl- CoA, the substrate of the *de novo* lipogenesis (Nematbakhsh *et al.*, 2021). We also
466 identified stearoyl-CoA desaturase (*SCD*) that plays an important role in biosynthesis of LC-PUFA via the
467 PPAR signalling pathway. Along with *ACACA*, *SCD* could be regulated by *SREBF1*, with both genes being
468 involved in FA *de novo* biosynthesis (Resnyk *et al.*, 2017). *SCD*, thanks to its Delta-9 desaturase activity, can

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3 469 convert palmitic acid (C16:0) and stearic acid (C18:0) to palmitoleic (C16:1 n-7) and oleic acid (C18:1n-9),
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5 470 respectively. *SCD* is recognized as a gene responsible for FA metabolism and having a significant role in
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7 471 intramuscular fat deposition. The third gene regulated by *SREBF1* is not presented in Table 1, but was
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10 472 significant in the comparison between BP_F_Ct and ROSS birds. This is the *FASN* gene and it is involved in
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12 473 *de novo* FA biosynthesis along with *ACACA* and *SCD* (Nematbakhsh *et al.*, 2021).
13
14 474 In conclusion, we provide a comparison of differential gene expression in two native dual-purpose slow-
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16 475 growing chickens, namely RM and BP, compared to commercial ROSS, fed a diet high in n-3 PUFA and
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19 476 assert that high levels of PUFA precursors in the diet do not result in significant changes in expression of
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21 477 genes involved in FA metabolism, whichever the genetic background or sex studied. **The diet had an effect**
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23 478 **on phenotype as body weight, especially in ROSS birds which were significantly lighter in the L diet.** On the
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26 479 contrary, it is clear that sex is an important factor in FA processes. Indeed, females differentially express
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28 480 elongases and desaturase genes (*ELOVL2*, *FADS1*, *FADS2*, and *SCD*) with a central role in LC-PUFA
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30 481 production, particularly in the BP breed. **The BP breed is also suggested to be the genetic background with**
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32 482 **better capacity for modulation of expression of genes involved in FA metabolism.** The two local breeds
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34 483 compared with ROSS commercial birds, have shown a significantly higher modulation of gene expression in
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37 484 liver, which underlines a greater aptitude of local breeds in FA metabolism and final LC-PUFA production.
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39 485 **Finally, this study has shed light on the capability of local chicken breeds to modulate the expression of**
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41 486 **some of the genes involved in FA metabolism, as well as the gene expression in females compared to**
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43 487 **males. Furthermore, the local breeds had the ability to adapt easily to a different diet without change in**
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46 488 **final weight, enriching the tissue (liver in this case) with n-3, hence having the potential for the**
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48 489 **establishment of high-quality products.**

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54

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56

57 493 Bando 2017 Prot. 2017S229WC. Title: USE OF LOCAL CHICKEN BREEDS IN ALTERNATIVE PRODUCTION

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60 494 CHAIN: WELFARE, QUALITY AND SUSTAINABILITY

DATA AVAILABILITY STATEMENT

RNAseq raw data are available in NCBI (BioProject ID PRJNA865899).

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36 743 TABLES

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40 745 **Table 1:** Differentially expressed genes shared by BP and RM female groups when compared to ROSS birds.
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42 746 Genes are related to Figure 7 and 8 and Appendix S6.
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47 749 FIGURES

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53 751 **Figure 1.** Principal component analysis (PCA) of all samples. Colours define different genetic background
54 752 (BP = Bionda Piemontese; RM = Robusta Maculata; ROSS = Ross 308); shapes represent different diet (Ct =
55 753 Control diet; L = Linseed diet). The two clusters represent the sex: males (M) in the smallest cluster on
56 754 bottom right, and females (F) in top left.
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Figure 2. Effect of diet on the percentage of total n-6 (a), total n-3 (b) and total PUFA (c) in relation to total FA in liver tissue. All the comparisons were assessed by experimental groups (same genetic background and sex, but different diet). *P < 0.05, ***P < 0.001 (post hoc Tukey's test). The data are presented as mean \pm standard error.

Figure 3. Volcano plot of differentially expressed genes (DEGs) between the Linseed diet (L) and Control diet (Ct) groups. Note: The red dots indicate genes with a P-adjusted value < 0.05 and $-2 > \text{Log}_2$ fold change > 2 . The red dots on the right quadrant of the figures are up-regulated in Linseed diet, the ones on the left are down-regulated. The blue dots represent the genes that reached the P-adjusted value < 0.05 but Log_2 fold change value is higher than -2 and lower than 2. The green dots represent the genes whose difference in expression level between the Ct and L groups did not reach significance (padj > 0.05).

Figure 4. Volcano plot of differentially expressed genes (DEGs) between the Female (F) and Male (M) groups. Note: The red dots indicate genes with a P-adjusted value < 0.05 and $-2 > \text{Log}_2$ fold change > 2 . The red dots on the right quadrant of the figures are up-regulated for female, the ones on the left are down-regulated. The blue dots represent the genes that reached the P-adjusted value < 0.05 but Log_2 fold change value is higher than -2 and lower than 2. The green dots represent the genes whose difference in expression level between the F and M groups did not reach significance (padj > 0.05).

Figure 5. Volcano plot of differentially expressed genes (DEGs) between the Bionda Piemontese (BP) and Ross308 (ROSS) groups. Note: The red dots indicate genes with a P-adjusted value < 0.05 and $-2 > \text{Log}_2$ fold change > 2 . The red dots on the right quadrant of the figures are up-regulated in the Bionda Piemontese breed, the ones on the left are down-regulated. The blue dots represent the genes that reached the P-adjusted value < 0.05 but Log_2 fold change value is higher than -2 and lower than 2. The green dots represent the genes whose difference in expression level between the BP and ROSS groups did not reach significance (padj > 0.05).

Figure 6. Volcano plot of differentially expressed genes (DEGs) between the Robusta Maculata (RM) and Ross308 (ROSS) groups. Note: The red dots indicate genes with a P-adjusted value < 0.05 and $-2 > \text{Log}_2$ fold change > 2 . The red dots on the right quadrant of the figures are up-regulated in the Robusta Maculata breed, the ones on the left are down-regulated. The blue dots represent the genes that reached the P-adjusted value < 0.05 but Log_2 fold change value is higher than -2 and lower than 2. The green dots represent the genes whose difference in expression level between the RM and ROSS groups did not reach significance (padj > 0.05).

Figure 7. Venn diagram of up-regulated genes in Bionda Piemontese and Robusta Maculata birds compared to Ross308 (ROSS) chickens in female experimental groups.

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Figure 8. Heatmap showing the K-means clustering of transformed expression values for the 23 genes reported in common between female experimental groups in **Figure 7**. Red represents higher expression and blue represents lower expression.

SUPPLEMENTARY MATERIALS:

Table S1. Formulation and chemical analysis of chicken diet

Table S2. **Outline** of experimental design. Birds reared for each experimental group. In brackets the liver samples used for the RNA extraction.

Table S3. Summary of contrasts made in differential expression analysis divided by Diet, Sex, Genetic Background. In bold the groups used as experimental, with up and down regulated genes referred to them.

Table S4. Representation of all genes involved in fatty acids metabolism found as differentially expressed in all experimental groups and divided by the belonging contrast

Figure S1. Step by step bioinformatic procedures adopted in the present study.

Figure S2. Effect of diet (Ct = control, L = linseed) on Live Body Weight in g. ***P < 0.001 (post hoc Tukey's test).

Figure S3. GeneMANIA report of 23 genes (dashed circles) reported in common between female experimental groups in **Figure 7**. Non-dashed circles represent genes commonly co-expressed with the 23 belonging to our dataset. Legends explain biological function in which the genes are involved.

Appendix S1. DEGs of all the comparison. Six Diet comparisons (up and down regulated genes refer to the Linseed group), six Sex comparisons (up and down regulated genes refer to the female group), four BP vs ROSS comparisons (up and down regulated genes are referred to the BP group) and four RM vs ROSS comparisons (up and down regulated genes refer to the RM group), In each comparison is reported just genes with a P-value > 0.05. All sheets report Gene ID; baseMean as the average of the normalized count values, dividing by size factors, taken over all samples; log2FoldChange indicates how much the gene or transcript's expression seems to have changed between the experimental and control groups. This value is reported on a logarithmic scale to base 2; lfcSE as the standard error estimate for the log2 fold change estimate; stat as the value of the test statistic for the gene or transcript; P-value of the test for the gene or transcript; and padj as Adjusted P-value for multiple testing for the gene or transcript.

Appendix S2. Gene Ontology for DEGs in Diet contrasts with significant values in Panther. Gene Ontologies refer to up-regulated genes in the Linseed diet group. The considered contrast for each gene ontology is reported in the title of each sheet. The GO tables are formed by *Gallus gallus* - REFLIST column that shows how many genes are involved in a given pathway. Name of the contrast column shows number of genes differentially expressed in the contrast belonging to a given pathway. Over/under represented column represents whether the pathway is enriched or down-regulated. Fold enrichment column explain how many times the pathway is enriched or down-regulated. Raw p-value and FDR (False discovery Rate) columns represents the statistical output of the Statistical overrepresentation test in Panther.

Appendix S3. Gene Ontology for DEGs in Sex contrasts with significant values in Panther and in IPA software. Gene Ontologies refer to up-regulated genes in the female sex group. The considered contrast for each gene ontology is reported in the title of each sheet. The GO tables are formed by *Gallus gallus* -

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3 839 *REFLIST* column that shows how many genes are involved in a given pathway. *Name of the contrast column*
4 840 shows number of genes differentially expressed in the contrast belonging to a given pathway. *Over/under*
5 841 *represented* column represents whether the pathway is enriched or down-regulated. *Fold enrichment*
6 842 column explains how many times the pathway is enriched or down-regulated. *Raw p-value* and *FDR* (False
7 843 discovery Rate) columns represents the statistical output of the Statistical overrepresentation test in
8 844 Panther. IPA analysis used the Sex comparison for each group. Only pathways with a z-score > 2 (activated)
9 845 are reported. Titles of each sheet explain the contrast under study and the pathways detected as activated.
10 846 Each IPA sheet reports ID and full name of genes involved in a given pathway, prediction of the behaviour
11 847 of a given gene when the pathway is activated, and log₂ Fold Change value for every gene in that contrast.

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15 848 **Appendix S4.** Gene Ontology for DEGs in Genetic background contrasts (Bionda Piemontese vs Ross308)
16 849 with significant values in Panther and IPA analyses. *Gene Ontologies* refer to up-regulated genes in the BP
17 850 breed group. The considered contrast for each gene ontology is reported in the title of each sheet. The GO
18 851 tables are formed by *Gallus gallus* - *REFLIST* column that shows how many genes are involved in a given
19 852 pathway. *Name of the contrast column* showed number of genes differentially expressed in the contrast
20 853 belonging to a given pathway. *Over/under represented* column represents whether the pathway is enriched
21 854 or down-regulated. *Fold enrichment* column explain how many times the pathway is enriched or down-
22 855 regulated. *Raw p-value* and *FDR* (False discovery Rate) columns represents the statistical output of the
23 856 Statistical overrepresentation test in Panther. IPA analysis used the Breed comparison for each group. Only
24 857 pathways with a z-score > 2 (activated) are reported.. Titles of each sheet explain the contrast under study
25 858 and the pathways detected as activated. Each IPA sheet reports ID and full name of genes involved in a
26 859 given pathway, prediction of the behaviour of a given gene when the pathway is activated, and log₂ Fold
27 860 Change value for every gene in that contrast.

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31 861 **Appendix S5.** Gene Ontology for DEGs in Genetic background contrasts (Robusta Maculata vs Ross308) with
32 862 significant values in Panther and in IPA analyses. *Gene Ontologies* refer to up-regulated genes in the BP
33 863 breed group. The considered contrast for each gene ontology is reported in the title of each sheet. The GO
34 864 tables are formed by *Gallus gallus* - *REFLIST* column that shows how many genes are involved in a given
35 865 pathway. *Name of the contrast column* shows number of genes differentially expressed in the contrast
36 866 belonging to a given pathway. *Over/under represented* column represents whether the pathway is enriched
37 867 or down-regulated. *Fold enrichment* column explain how many times the pathway is enriched or down-
38 868 regulated. *Raw p-value* and *FDR* (False discovery Rate) columns represents the statistical output of the
39 869 Statistical overrepresentation test in Panther. IPA analysis used the Breed comparison for each group. Only
40 870 pathways with a z-score > 2 (activated) are reported. Titles of each sheet explain the contrast under study
41 871 and the pathways detected as activated. Each IPA sheet reports ID and full name of genes involved in a
42 872 given pathway, prediction of the behaviour of a given gene when the pathway is activated, and log₂ Fold
43 873 Change value for every gene in that contrast.

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47 874 **Appendix S6.** Gene lists of Venn diagram groups reported in Figure 7.
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