

**In Press, Pre-Proof Version**

## **Sample storage and fasting times affect serum and plasma concentrations of metabolites in fasted and non-fasted broiler chickens**

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### **ABSTRACT**

Sample storage and fasting times leads to some changes of blood metabolite in broilers. Therefore, a study was conducted with the aim to assess the influence of storage and fasting times in serum and plasma fractions on glucose, total cholesterol, triacylglycerols, aspartate aminotransferase (AST), alanine aminotransferase (ALT), and gamma-glutamyl transferase (GGT) concentrations in broilers. A total of 70 male broiler chickens fasted at 7 times (0, 2, 4, 6, 8, 10, and 12 h) to collect blood fractions (serum and plasma) stored at -20 °C for 0, 30, and 60 days. Glucose and GGT were affected by fasting times×blood fraction. Serum glucose concentration decreased linearly ( $\cong 2.48 \text{ mg dL}^{-1}$ ), whereas total cholesterol and plasma GGT increased linearly ( $\cong 0.92 \text{ mg dL}^{-1}$  and  $\cong 0.19 \text{ IU L}^{-1}$ , respectively) with fasting time. There was a quadratic effect on plasma glucose and serum GGT (maximum at 3.95 h and minimum at 5.22 h of fasting, respectively), and triacylglycerol (minimum at 8.75 h of fasting) and ALT concentrations (maximum at 8.45 h of fasting). Glucose, total cholesterol, AST, ALT, GGT concentrations were higher in serum, while triacylglycerols were higher in plasma. Glucose concentration had the lowest values at 30 days, while ALT was higher on day 0. However, GGT concentrations were lower on days 0 and 30. **Samples of plasma for glucose**, ALT, and GGT stored at -20 °C for long periods should be avoided. In addition, serum samples and 6 h fasting are recommended for the assessment of blood biochemical metabolites in broilers.

**Keywords:** Blood fractions, Broiler, Fasting, Liver enzyme, Sample storage.

### **INTRODUCTION**

The assessment of biochemical parameters in poultry research can be better targeted, allowing the measurement of the concentration of several blood biochemical constituents that can be used as indicators of metabolic disorders, nutritional status, and in the diagnosis of

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33 diseases. Experiments involving broiler chickens commonly aim to assess bird performance,  
34 promoting satisfactory feed conversion without compromising the metabolism and health  
35 integrity of the birds. The changes that occur in the metabolism of birds can be caused by the  
36 inclusion of ingredients, additives, chemotherapeutics, or other components of the diet (Hagan  
37 *et al.*, 2022), in addition to environmental effects, which can alter the nutritional status and  
38 health of the animals. Therefore, blood biochemical assessments are an important tool for the  
39 diagnosis of diseases and metabolic disorders, providing efficient, fast, and safe diagnoses.

40 According to Gattani *et al.* (2016), the measurement of blood concentrations of glucose,  
41 total cholesterol, triacylglycerol, aspartate aminotransferase (AST), alanine aminotransferase  
42 (ALT), and gamma-glutamyl transferase (GGT) activities can be used to aid in the diagnosis of  
43 numerous metabolic disorders in broiler chickens. However, there is a wide divergence in the  
44 conditions under which blood samples are taken for analysis, as well as no standardization of  
45 the postprandial fasting period performed before blood collection. Córdova-Noboa *et al.* (2018)  
46 did not fast birds to collect blood samples, but Sadeghi *et al.* (2014), Behboudi *et al.* (2016),  
47 Zakaria *et al.* (2017), and Swarna *et al.* (2018) performed fasting between 2 to 12 h prior sample  
48 collection. Taking into account these variations in fasting times, it is not possible to state that  
49 the results obtained in biochemical blood analyses will not be influenced by a long postprandial  
50 fasting period, as recently demonstrated in a study conducted by Wachholz *et al.* (2023).

51 Biochemical parameters can be measured in serum samples, or plasma obtained with the  
52 use of anticoagulants. The difference between these two processes is that in plasma, a larger  
53 volume of supernatant is obtained compared to serum (Lumeij, 2008). Some divergence in the  
54 use of serum or plasma in research results in a lack of standardization of results. Some authors  
55 used samples of serum to measure blood biochemical parameters (Chand *et al.*, 2018, Gilani *et al.*  
56 *et al.*, 2018, Rehman *et al.*, 2018, Subhani *et al.*, 2018), while others assessed in plasma (Sharideh  
57 *et al.*, 2016, Yang *et al.*, 2017, Zhang *et al.*, 2017, Kim *et al.*, 2019).

58 Regarding the storage times that each sample can be subjected to before the biochemical  
59 analyses, it is important to consider that the samples will usually be collected at different times  
60 from the day of analysis because factors such as distance between the laboratory and the  
61 experimental facility, time to obtain sufficient volume of samples, and transportation period,  
62 can delay the date of the measurements of the biochemical variables in blood (Livesey *et al.*,  
63 2008). This is supported by the study conducted by Wachholz *et al.* (2023), who assessed the  
64 effect of time and storage condition on biochemical metabolites in serum or plasma samples of  
65 broiler chickens and observed significant changes in the concentrations of glucose, total  
66 cholesterol, triglycerides, AST, ALT and GGT.

67 Here, a study was conducted based on the hypothesis that both independent variables tested  
68 can significantly alter blood metabolite concentrations and the results in broiler chicken  
69 experiments. Therefore, this study aimed to assess the influence of storage and fasting times in  
70 serum and plasma fractions on the concentrations of glucose, total cholesterol, triacylglycerols,  
71 AST, ALT and GGT in broiler chickens.

72

## 73 **MATERIALS AND METHODS**

### 74 **Place of study, birds, handling, housing, and diets**

75 The present study was conducted at Universidade Estadual do Oeste do Paraná (Unioeste),  
76 Marechal Cândido Rondon, PR, Brazil. The University Animal Ethics Committee approved the  
77 experiment under number 23/20. A total of 70 45-day-old Cobb 500<sup>®</sup> male broiler chickens  
78 were used, with an average body weight of 3,072±859 g. The birds were raised from 1 to 42  
79 days of age, receiving water and diet *ad libitum*, and the same care to management, lighting and  
80 ambient temperature recommended by the lineage manual. The diets for each phase (starter,  
81 grower, and finisher) were corn-soybean based supplemented with industrial amino acids,  
82 isonutritional and isoenergetic, and according to the nutritional requirements proposed by  
83 Rostagno *et al.* (2017).

84 At 42 days of age, the birds were assigned in a completely randomized design in 7 pens  
85 replicates (1.76 m<sup>2</sup>), with a masonry floor covered with 10 cm of pine wood shavings. Each pen  
86 was composed of 10 birds and a density of 5.7 birds per m<sup>2</sup>. The pens were equipped with a  
87 tubular feeder and nipple drinkers. The facility was equipped with electrical elements, hoods,  
88 evaporative pads to assist cooling and air exchange.

89

### 90 **Blood sampling, preparation and analysis**

91 After 3 days of adaptation at 45 days-old, the birds were fasted for 1 h, then fed for 30 min.  
92 This procedure was adopted so that all birds had the same postprandial feeding condition. After  
93 this period of feeding, the first blood collection was performed. At every 2-h interval, within a  
94 total period of 12 h (0, 2, 4, 6, 8, 10, and 12 h of fasting), blood was collected by puncture of  
95 the ulnar vein from 1 bird per pen ( $n = 7$  birds per fasting time randomly selected). Every 7  
96 birds were used only for a single sampling at a given fasting time. Blood collection was  
97 performed with the birds in the decubitus position lateral, using specific vacuum collection  
98 tubes (Vacutainer<sup>®</sup>, Curitiba, PR, Brazil) with a capacity of 10 mL, specific adapter and 25×0.8  
99 mm needles (21G 1”) (Labor Import brand, Maringá, PR, Brazil).

100 Two tubes of approximately 4 mL each were collected within the allowable for species and  
101 body weight (Kelly and Alworth, 2013). The first blood sample was collected to obtain the  
102 serum in a tube (BD Vacutainer<sup>®</sup>, Curitiba, PR, Brazil) with clot activator (silica powder)  
103 blasted on the tube wall to accelerate the process of coagulation. The second sample collected  
104 was to obtain plasma in a tube (BD Vacutainer<sup>®</sup>, Curitiba, PR, Brazil) with 5 mg of sodium  
105 fluoride as a glycolysis inhibitor and 4 mL EDTA-K<sub>3</sub> anticoagulant. These two draws collected  
106 per bird were consecutive. After collection, the samples remained for 15 min at room  
107 temperature in a horizontal position and then were centrifuged (Centrifuge Kasvi K14-4000,  
108 Kasvi, São Paulo, SP, Brazil) at 2,500 g for 10 min at room temperature.

109 After centrifugation and separation of serum and plasma, the samples were identified and  
110 divided into three aliquots as technical triplicates, which were placed in 2 mL microtubes  
111 (Eppendorf<sup>®</sup> brand, Minispin<sup>®</sup>, Hamburg, Germany). An aliquot was immediately sent to the  
112 laboratory for analysis. The other aliquots (two microtubes as technical duplicates) were stored  
113 at -20 °C for times of 30 and 60 days for further analysis (Wachholz *et al.*, 2023).

114 Within each storage times, the samples were thawed under refrigeration (4 °C) and kept in  
115 a refrigerator for 24 h. Before performing the analysis, the samples were centrifuged in an  
116 Eppendorf microcentrifuge (Eppendorf<sup>®</sup> brand, Minispin<sup>®</sup>, Hamburg, Germany) to remove  
117 fibrin formation. Biochemical analyzes were performed using commercial kits and calibrators  
118 (Elical II multiparametric Calibrator, ref. CALI-0550), and measurement standards for birds  
119 (Elitrol I normal multiparametric control, ref. CONT-0060) (Elitech Clinical Systems, ELITech  
120 Group, Paris, France) in automatic calibration spectrophotometer (Elitech<sup>®</sup> brand, Flexor  
121 EL200 model, Puteaux, France).

122 The determination of glucose concentration (Glucose PAP) was performed by Trinder's  
123 method, enzymatic colorimetric kinetic (Trinder, 1969), total cholesterol was performed by  
124 Trinder's method, enzymatic colorimetric endpoint (Allain *et al.*, 1974), triacylglycerols  
125 (Triglycerides SL New), enzymatic colorimetric endpoint (Fossati and Prencipe, 1982), AST  
126 and ALT were performed according to International Federation of Clinical Chemistry method  
127 without pyridoxal phosphate, kinetic, UV (Schuman *et al.*, 2002a, Shuman *et al.*, 2002b), and  
128 GGT (GAMMA GT plus) was performed according the procedure Glupa-C substrate method,  
129 kinetic (Schuman *et al.*, 2002c).

130

### 131 **Statistical procedures**

132 Data were analyzed considering blood fraction (serum and plasma), fasting (0, 2, 4, 6, 8,  
133 10, and 12 h) and storage times of the sample (0, 30, and 60 days) as fixed effects. Residual

134 error was considered as a random factor. Each bird belonging to the same pen was considered  
135 an experimental unit. Data were subjected to normality analysis using the Shapiro-Wilk test.  
136 Afterward, the two-way analysis of variance (ANOVA) was performed, considering the  
137 isolated effects and interactions between the studied factors: fasting or storage times×blood  
138 fractions (serum and plasma) as fixed factors. All tests were performed at 5% significance level.  
139 In case of significance, F-test was performed for the blood fraction, regression analysis for  
140 fasting times (linear or quadratic models), and Tukey's post hoc test for storage times. The triple  
141 interaction was not tested. All statistical procedures were performed using the GLM procedure  
142 of SAS University Edition.

143

## 144 RESULTS AND DISCUSSION

145 Glucose concentrations were influenced by the interaction between fasting times×blood  
146 fractions (Table 1). Serum glucose concentration decreased linearly with fasting time, reducing  
147  $\cong 2.48 \text{ mg dL}^{-1}$  every 2 h of fasting. Total cholesterol concentrations increased linearly with  
148 prolonged fasting time, increasing  $\cong 0.92 \text{ mg dL}^{-1}$  (Figure 1). There was a quadratic effect on  
149 the concentrations of plasma glucose, with maximum concentrations at 3.95 h of fasting, and  
150 on triacylglycerol concentrations, with minimum concentration at 8.75 h of fasting. Glucose  
151 and total cholesterol concentrations were higher in serum than in plasma, while triacylglycerol  
152 concentrations were higher in plasma compared to serum. Glucose concentrations were  
153 influenced by storage times, with the lowest values at 30 days of storage compared to 0 and 60  
154 days.

155 The linear decreasing effect shown in serum glucose and quadratic for plasma glucose is  
156 explained by the decrease of glycogen in the liver not being enough to maintain blood glucose  
157 concentrations, causing its gradual reduction (Rodrigues *et al.*, 2017). The increase in total  
158 cholesterol concentration and the decrease in triacylglycerols is due to lipid mobilization, as  
159 lipids enter the bloodstream and are consequently transported by very-low-density lipoproteins,  
160 causing a greater circulation of total cholesterol in the blood and, hence, the use of  
161 triacylglycerols by the tissues (Lumeij, 2008).

162 In addition, blood concentrations of triacylglycerols are also influenced by fasting times  
163 because when a negative energy balance occurs, bird metabolism is stimulated to mobilize  
164 circulating triacylglycerols to provide fatty acids and glycerol via  $\beta$ -oxidation and glycolysis,  
165 respectively, which are used for energy production, lowering blood triacylglycerols  
166 concentrations (Coelho *et al.*, 2013). These results are in agreement with Vosmerova *et al.*

167 (2010), who observed that birds exposed to prolonged fasting times showed reduced glucose  
168 and triacylglycerol concentrations.

169 The storage times of serum and plasma can lead to a reduction in glucose concentrations.  
170 The present study showed variation in glucose concentrations caused by storage even at -20 °C.  
171 This suggested that the temperature used during storage was not sufficient to keep glucose  
172 concentrations stable for 60 days. The results of this study corroborate with the findings of  
173 Cuhadar *et al.* (2013) and Clark *et al.* (1990), who stored serum samples from humans at -20  
174 °C for 30 and 60 days, and observed a decrease in glucose concentrations.

175 The results observed for total cholesterol and triacylglycerols demonstrate that these  
176 analytes can remain stable for 60 days at a temperature of -20 °C without any change. According  
177 to Stokes *et al.* (1986), and Tiedink and Katan (1989), cholesterol and triacylglycerols present  
178 in plasma and serum do not change their concentrations stored at -20 °C for 19 to 27 weeks.  
179 The stability of triacylglycerols for long storage periods was also observed by Shimizu and  
180 Ichihara (2019) storing serum samples for 2 months at -20 °C.

181 Although the differences found between the concentrations of biochemical analytes in the  
182 blood fraction are statistically significant, Picheth *et al.* (2001) reported that these differences  
183 between fractions for glucose do not represent chemically significant differences, pointing out  
184 that both fractions can be used analytically for glucose.

185 Gamma-glutamyl transferase concentrations were affected by the interaction between  
186 fasting times×blood fractions (Table 2). Plasma GGT concentrations increased linearly with  
187 prolonged fasting time, increasing  $\cong 0.19$  IU L<sup>-1</sup> (Figure 1). There was a quadratic effect on the  
188 concentrations of serum GGT, with a minimum concentration at 5.22 h of fasting. In addition,  
189 there was a quadratic effect on ALT concentrations, with a maximum concentration at 8.45 h  
190 of fasting. Aspartate aminotransferase, ALT and GGT concentrations were higher in serum than  
191 in plasma. Alanine aminotransferase and GGT concentrations were influenced by storage times.  
192 It was observed that ALT concentration was higher on day 0 compared to the other storage  
193 times. However, GGT concentrations were lower on days 0 and 30 than on day 60.

194 The quadratic response for ALT enzyme activity corroborates the results described of Veiga  
195 *et al.* (1978), who reported that fasting time increases ALT concentrations in birds due to the  
196 greater hepatic influx of gluconeogenic substrates, but less correlated with hepatic  
197 gluconeogenesis than AST activity. The occurrence of a quadratic effect with the increase in  
198 serum GGT concentration, and the linear increase in plasma GGT is because the metabolism of  
199 the animals in prolonged fasting triggers the protein catabolism and, consequently, increases  
200 the transport of amino acids. This highlights the main function of GGT, which according to Yu

201 and Long (2016), has the function of acting in the extracellular environment by catalyzing the  
202 conversion of glutathione into glutamate dipeptides or cysteine.

203 Another metabolite stable to storage is AST, which did not change its concentration during  
204 the 2 months of storage at -20 °C. According to Kaneko *et al.* (2008), this enzyme is stable to  
205 storage during freezing. This response was also observed by Thoresen *et al.* (1995) and Oliveira  
206 *et al.* (2011), who studied the same enzyme, but in blood samples from dogs and lambs,  
207 respectively. However, ALT concentrations tend to decrease with storage times (Kaneko *et al.*,  
208 2008). This decrease in blood ALT concentration is the result of enzyme denaturation as  
209 reported by Ikeda *et al.* (2015) in a study conducted with human blood samples, who observed  
210 enzyme denaturation and a decrease in its concentrations at -20 °C due to its instability at this  
211 temperature.

212 Storage times can negatively interfere with the results of blood analyses in experiments  
213 because researchers can often make conclusions based on results influenced by storage times.  
214 The concentrations of glucose, total cholesterol, AST, ALT, and GGT evaluated in the current  
215 study were higher in the serum sample compared to those observed in the plasma. This higher  
216 concentration in serum can be attributed to the action that low molecular weight anticoagulants,  
217 such as sodium fluoride, can exert on the samples. The osmotic effect of these anticoagulants  
218 tends to remove an amount of water from inside the erythrocytes and transfer it to the plasma,  
219 which would cause a dilution of the plasma concentrations of some metabolites (Alper *et al.*,  
220 1974). According to Grande *et al.* (1964), this can cause dilution of the plasma constituents,  
221 resulting in lower values in this blood fraction and, therefore, an anticoagulant that does not  
222 interfere with the analysis results should be chosen. In addition, another important point to note  
223 is that an osmotic redistribution between cells and plasma can occur, and this can interfere with  
224 the analyte results.

225 The use of sodium fluoride as an anticoagulant can interfere analytically in  
226 spectrophotometric assessments because this anticoagulant can cause fibrin formation in  
227 samples during their collection (Fernandez *et al.*, 2013, Al-Kharusi *et al.*, 2014, Bonetti *et al.*,  
228 2016). Thus, after plasma separation, the formation of hemolysis must be observed and, if  
229 possible, to perform a new collection or make a higher centrifugal force.

230 Gamma-glutamyl transferase concentrations showed different effects on blood fraction  
231 during the storage times. The difference in GGT concentrations between serum and plasma, as  
232 well as a decrease in metabolite values in plasma analysis, are due to the use of sodium fluoride  
233 anticoagulant, which reduces GGT activity (Burtis *et al.*, 2012).

234 **Du et al. (2023)** examined serum blood metabolites in broilers at 49-day-old on restricted  
235 feeding and 16-h fasting and found lower average values than our findings for total cholesterol  
236 (67.14 vs. 136 mg dL<sup>-1</sup>), triacylglycerols (7.92 vs. 42 mg dL<sup>-1</sup>), and glucose (151 vs. 233 mg  
237 dL<sup>-1</sup>) and ALT (2.06 vs. 9.51 IU L<sup>-1</sup>) compared to the 12-h fasting in our study. However, the  
238 values for AST (587 vs. 412 IU L<sup>-1</sup>) and GGT (24.78 vs. 22.59 IU L<sup>-1</sup>) were higher than those  
239 of the present study. Corroborating the results of the present study, Hagan *et al.* (2022) reported  
240 serum reference values of 197 to 299 mg dL<sup>-1</sup> for glucose, 129 to 297 mg dL<sup>-1</sup> for total  
241 cholesterol, and average values of 96.15 mg dL<sup>-1</sup> for triacylglycerol concentrations in 56-day-  
242 old Cobb and Ross broiler chickens fasted for 12 h.

243 Based on the criteria assessed in the current study, the results indicated that the  
244 determination of AST is not influenced by the sample storage and fasting times. Total  
245 cholesterol and triacylglycerol measurements can be performed on samples stored for up to 60  
246 days at -20 °C. Plasma glucose, ALT, and GGT analyses stored at -20 °C for long periods should  
247 be avoided. In addition, serum samples and 6 h fasting are recommended for the assessment of  
248 blood biochemical metabolites in broilers.

249

## 250 **ACKNOWLEDGEMENTS**

251 The authors gratefully acknowledge the Coordenação de Aperfeiçoamento de Pessoal de Nível  
252 Superior (CAPES), Fundação Araucária and the Conselho Nacional de Desenvolvimento  
253 Científico e Tecnológico (CNPq).

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407 **Table 1.** Concentrations (mg dL<sup>-1</sup>) of glucose, total cholesterol and triacylglycerols in serum and  
 408 plasma stored at different times in 45-day-old broiler chickens subjected to different fasting times.

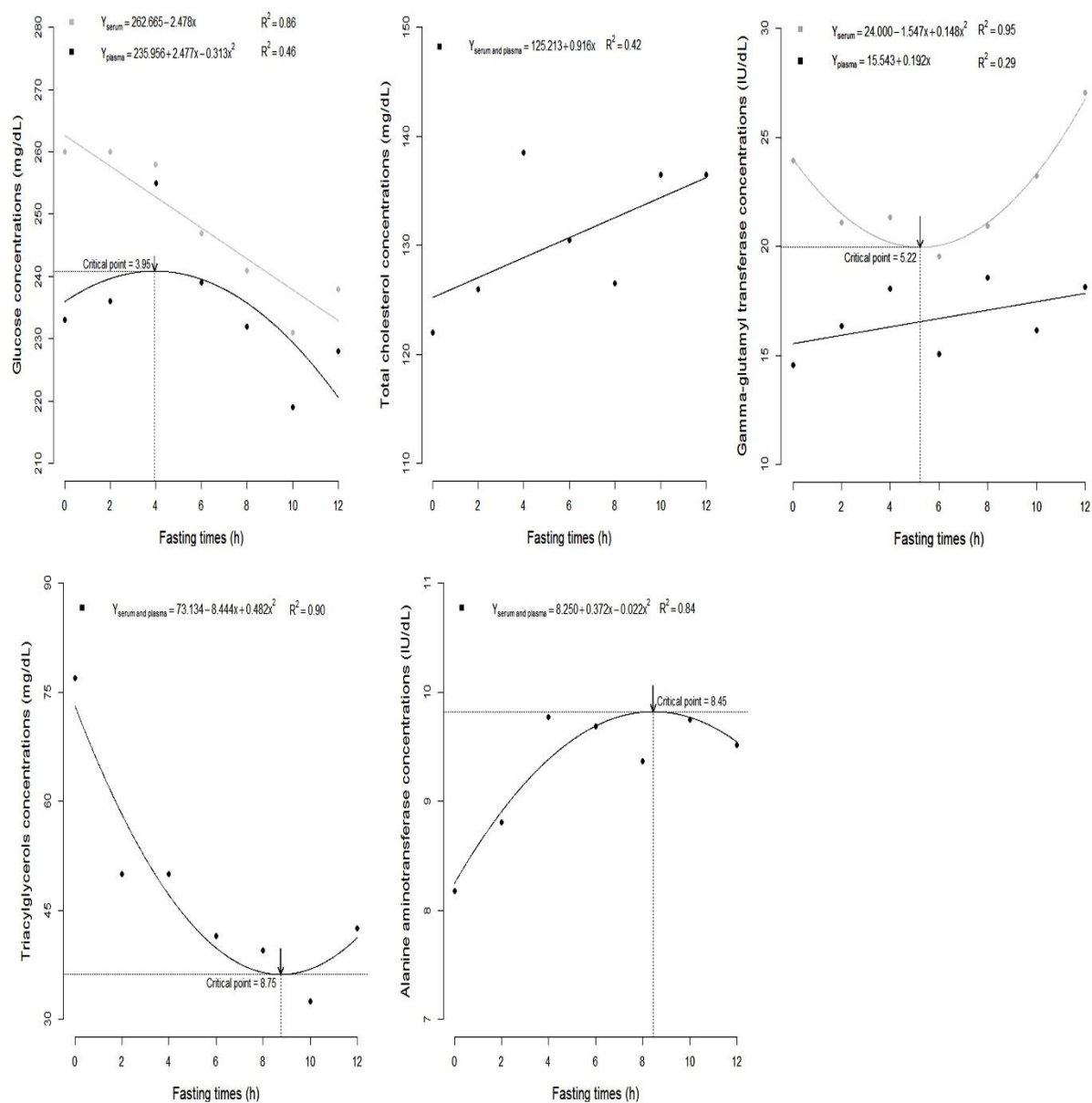
Fasting times (h)	Glucose		Total cholesterol		Triacylglycerols	
	Serum	Plasma	Serum	Plasma	Serum	Plasma
0	260±12	233±15	130±13	114±10	76±19	78±19
2	260±18	236±13	134±13	118±13	48±10	52±12
4	258±21	255±17	141±14	136±13	47±12	53±13
6	247±21	239±12	138±12	123±12	41±14	42±11
8	241±11	232±10	132±11	121±12	38±6	41±08
10	231±03	219±10	146±14	127±11	32±7	33±06
12	238±21	228±17	145±14	128±13	42±5	43±06
Storage times (days)						
0	251±20	238±16 <sup>A</sup>	137±14	124±14	44±19	49±20
30	243±17	230±15 <sup>B</sup>	138±14	125±12	47±17	50±71
60	249±23	235±19 <sup>A</sup>	138±14	122±15	47±16	48±18
Average	248±20 <sup>a</sup>	234±17 <sup>b</sup>	138±14 <sup>a</sup>	124±14 <sup>b</sup>	46±17 <sup>b</sup>	49±18 <sup>a</sup>
SEM	0.961		0.765		0.847	
Fasting times	<0.001		<0.001 (L)		<0.001 (Q)	
Blood fraction	<0.001		<0.001		0.017	
Storage times	<0.001		0.501		0.364	
Fasting times×blood fraction	<0.001		0.061		0.899	
Linear	<0.001	<0.001				
Quadratic	<0.001	<0.001				
Storage times×blood fraction	0.848		0.594		0.355	

409 <sup>a-b</sup> Lowercase letters in the same row differ by F test (P< 0.05); <sup>A-B</sup> Capital letters in the same column differ by  
 410 Tukey's post hoc test (P< 0.05); SEM: Pooled standard error of the mean; L: Linear effect; Q: Quadratic effect.

411  
 412 **Table 2.** Concentrations (IU L<sup>-1</sup>) of aspartate aminotransferase (AST), alanine aminotransferase  
 413 (ALT) and gamma-glutamyl transferase (GGT) in serum and plasma stored at different times  
 414 in 45-day-old broiler chickens subjected to different fasting times.

Fasting times (h)	AST		ALT		GGT	
	Serum	Plasma	Serum	Plasma	Serum	Plasma
0	353±60	322±56	9.17±2.43	7.18±2.13	23.95±6.97	14.57±2.62
2	408±87	407±81	9.34±2.29	8.28±2.83	21.10±6.78	16.36±4.86
4	450±76	440±70	9.95±2.84	9.59±2.85	21.32±5.49	18.05±4.01
6	436±63	403±71	10.25±2.45	9.12±2.15	19.55±6.01	15.08±2.67
8	413±66	380±73	9.93±2.00	8.80±2.30	20.96±5.10	18.57±4.02
10	460±67	417±70	10.65±2.33	8.85±2.91	23.25±6.07	16.14±4.22
12	436±71	387±65	10.65±3.01	8.38±2.75	27.04±5.63	18.15±2.99
Storage times (days)						
0	409±71	388±65	11.14±2.40	9.76±2.26 <sup>A</sup>	16.80±3.85	16.66±4.08 <sup>B</sup>
30	431±74	397±71	9.40±2.42	8.02±2.64 <sup>B</sup>	16.48±3.69	16.03±4.43 <sup>B</sup>
60	426±85	396±94	9.44±2.37	8.04±2.64 <sup>B</sup>	24.22±6.45	16.82±4.32 <sup>A</sup>
Average	422±77 <sup>a</sup>	394±77 <sup>b</sup>	9.99±2.52 <sup>a</sup>	8.60±2.64 <sup>b</sup>	22.46±6.41 <sup>a</sup>	16.70±3.94 <sup>b</sup>
SEM	3.826		0.130		0.296	
Fasting times	0.140		<0.001(Q)		<0.001	
Blood fraction	<0.001		<0.001		<0.001	
Storage times	0.135		<0.001		0.009	
Fasting times×blood fraction	0.483		0.352		0.002	
Linear					0.032	0.004
Quadratic					<0.001	0.010
Storage times×blood fraction	0.703		0.998		0.005	
Unfolding					<0.001	0.851

415 <sup>a-b</sup> Lowercase letters in the same row differ by F test (P< 0.05); <sup>A-B</sup> Capital letters in the same column differ by  
 416 Tukey's post hoc test (P< 0.05); SEM: Pooled standard error of the mean; L: Linear effect; Q: Quadratic effect.



417

418 **Figure 1.** Serum and plasma concentrations of glucose, total cholesterol, triacylglycerols,  
 419 alanine aminotransferase and gamma-glutamyl transferase in broiler chickens at 45-day-old  
 420 subjected to different fasting times ( $R^2$ : coefficient of determination).  
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421