

## SLAUGHTER VALUE OF PIGS CARCASSES AS RELATED TO COLIPASE GENOTYPE (CLPS) CONSIDERING RYR1 GENE EFFECT

### WARTOŚĆ RZEŻNA TUSZ ŚWIŃ W ZALEŻNOŚCI OD GENOTYPU KOLIPAZY (CLPS) Z UWZGLĘDNIENIEM ODDZIAŁYWANIA GENU RYR1

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#### ABSTRACT

The experiment was carried out on 110 3-breed crossbreeds fatteners [ $\sigma^7$  Pietrain x ( $\text{♀}$  Polish Large White x Polish Landrace)]. Pigs were reared in standard conditions and slaughtered at an average body weight of 105 kg. Gene polymorphism was determined with the PCR-RFLP procedure (CLPS/DdeI and RYR1/HinPI). One day post-slaughter, the right carcass-sides were dissected and evaluated using the methods applied for the Polish Pig Testing Station (SKURTCh). The study evaluated the slaughter value of carcasses depending on colipase genotype (CLPS) with influence of major gene RYR1. Colipase gene differentiated carcass lean content only in RYR1<sup>CC</sup> pigs and in all pigs the neck weight as well as its tissue composition in RYR1<sup>CT</sup> pigs. Moreover, significant influence of major gene (RYR1<sup>T</sup>) on quantitative traits was confirmed.

KEY WORDS: pigs, colipase gene, RYR1 gene, carcass quality

#### STRESZCZENIE

Badaniem objęto 110 tuczników, mieszańców 3-rasowych [ $\sigma^7$  Pietrain x ( $\text{♀}$  Wielka Biała Polska x Polska Biała Zwisłoucha)]. Zwierzęta utrzymywane były w ujednoczonych warunkach i ubijane przy zbliżonej masie ciała, około 105 kg. Polimorfizm genów został określony przy użyciu metody PCR-RFLP (CLPS/DdeI i RYR1/HinPI). Dzień po uboju, prawe półtusze poddano dyssekcji i ocenie zgodnie z metodyką opracowaną dla polskich Stacji Kontroli Użytkowości Rzeźnej Trzody Chlewnej. Ocenie poddano wartość rzeźną tusz w zależności od genotypu kolipazy (CLPS), uwzględniając przy tym wpływ genu głównego RYR1. Gen kolipazy istotnie różnicował zawartość mięsa w tuszy tylko u tuczników o genotypie RYR1<sup>CC</sup> oraz u wszystkich świń masę karkówki i jej skład tkankowy u świń RYR1<sup>CT</sup>. Ponadto, potwierdzono wpływ genu głównego na cechy ilościowe tusz.

SŁOWA KLUCZOWE: świnie, gen kolipazy, gen RYR1, jakość tuszy

**STRESZCZENIE SZCZEGÓŁOWE**

Celem pracy było określenie wartości rzeźnej tusz świń w zależności od polimorficznych form genu kolipazy (CLPS) z uwzględnieniem wpływu genu głównego RYR1. Badaniem objęto 110 mieszańców 3-rasowych [♂ Pietrain x (Wielka Biała Polska x Polska Biała Zwisłoucha)]. W celu określenia genotypu kolipazy oraz genu podatności świń na stres w czasie trwania tuczu od zwierząt pobrano krew. Identyfikacji polimorfizmu genu CLPS dokonano metodą PCR-RFLP przy zastosowaniu endonukleazy DdeI według metody Baskina i Pompa [1]. Natomiast mutacji genu receptora ryanodiny (RYR1<sup>T</sup>) przy użyciu enzymu restrykcyjnego HinPI [4]. Uboju tuczników dokonywano przy masie ciała ok. 105 kg. Po 24 godzinnym wychłodzeniu tusz, prawe półtusze poddano szczegółowej ocenie zgodnie z metodyką opracowaną dla polskich Stacji Kontroli Użytkowości Rzeźnej Trzody Chlewnej [8]. Wykonano pomiary grubości słoniny grzbietowej w pięciu miejscach, pomiar powierzchni przekroju połówki oraz dokonano szczegółowego rozbioru wyrębów podstawowych na poszczególne elementy tkankowe (mięso, skórę ze słoniną oraz kości). Na podstawie dysekcji określono mięsność tusz. Uzyskane wyniki opracowano statystycznie wykorzystując program komputerowy STATISTICA 5.5 PL [9]. W badanej populacji mieszańców 3-rasowych wystąpiły wszystkie trzy genotypy kolipazy, a najliczniej reprezentowany był genotyp heterozygotyczny (frekwencja 0.56). Genotypy homozygotyczne wystąpiły w mniejszości: 0.34 (AA) i 0.10 (BB). Badana populacja mieszańców okazała się także zróżnicowana pod względem frekwencji genotypów RYR1, stwierdzono także wszystkie trzy jego formy (RYR1<sup>CC</sup>, RYR1<sup>CT</sup>, RYR1<sup>TT</sup>). Najwięcej badanych zwierząt wykazało genotyp RYR1<sup>CT</sup> (n=56), a najmniej genotyp RYR1<sup>TT</sup> (n=21) (tabela 1).

Analiza wyników zestawionych w podgrupach (genotyp CLPS x genotyp RYR1) wskazuje na większą zawartość

mięsa w tuszach zwierząt o genotypie BB względem AA w grupie zwierząt odpornych na stres (RYR1<sup>CC</sup>);  $P \leq 0,05$  (tabela 2). Wykazano też wpływ polimorfizmu genu kolipazy na skład tkankowy karkówki oraz jej masę. Zwierzęta o genotypie AA oraz AB charakteryzowały się większą masą karkówki (odpowiednio: 5.13 kg i 5.04 kg) w stosunku do grupy BB (4.80 kg), ( $P \leq 0,01$ ) (tabela 3). Wyższą zawartością mięsa w tym wyrębie cechowały się zwierzęta z grupy AA (65.39%) niż zwierzęta z grupy BB (63.45%) (tabela 3). Analizując relacje między oddziaływaniem genu kolipazy i genu RYR1 wykazano wpływ genotypu kolipazy na umięśnienie i otłuszczenie karkówki tylko u świń heterozygot względem genu RYR1 (tabela 3). Ponadto potwierdzono statystycznie istotny wpływ genotypu RYR1<sup>TT</sup> na zawartość mięsa w tuszy oraz na wielkość powierzchni oka połówki (tabela 2) jak również na zawartość mięsa w karkówce (tabela 3).

**INTRODUCTION**

During the last two decades, several genes having a main effect on important production traits have been detected in farm livestock. Genes reported to have a major effect are called major genes. Examples for major genes in pig production are the pig stress susceptibility gene (RYR1) and the RN- gene influence on meat quality and the estrogen receptor locus (ESR) affecting litter size in pigs.

Moreover, exist also genes defined as being candidate. Polymorphism of candidate genes and their connections with commercial traits of breeding pigs is intensively studied in many laboratories on the world. One of the candidate genes is colipase gene (CLPS). The CLPS gene was located in the area of chromosome 7 [1] and considered a candidate gene for QTL affecting carcass fatness in pigs [10, 3].

During the study described in this paper we used the

Table 1. Frequency of particular genotypes of *CLPS* and *RYR1* gene in testing fatter groups  
Tabela 1 Rozkład genotypów *CLPS* oraz *RYR1* w badanej grupie tuczników

<i>RYR1</i> genotype		<i>CLPS</i> genotype			Total
		<i>AA</i>	<i>AB</i>	<i>BB</i>	
<i>CC</i>	number	10	20	3	33
	frequency	0.30	0.61	0.09	1.00
<i>CT</i>	number	22	28	6	56
	frequency	0.39	0.50	0.11	1.00
<i>TT</i>	number	6	14	1	21
	frequency	0.29	0.67	0.04	1.00
Total	number	38	62	10	110
	frequency	0.34	0.56	0.10	1.00

polymerase chain reaction (PCR) and RFLP to identify CLPS and RYR1 gene polymorphisms in crossbred pig populations with the aim of estimating the relationship between genotypes CLPS and slaughter value considering RYR1 gene effect.

### MATERIAL AND METHODS

The experiment was carried out on 110 crossbreds fatteners [♂ Pietrain x (♀ Polish Large White x Polish Landrace)]. Pigs were reared in standard conditions. In the course of fattening period blood samples were taken from the vena cava to test – tube with EDTA\*K<sub>3</sub> and freeze at -20°C. Determination of the colipase genotype was performed using the PCR-RFLP method with DdeI restriction enzyme [1] and RYR1/HinPI genotypes were identified with the method of Fujii et al. [4]. The fatteners were slaughtered at an average body weight of 105 kg.

One day post-slaughter, the right carcass-sides were dissected and evaluated using the methods applied for the Polish Pig Testing Station (SKURTCh) [8]. Thickness of the backfat was measured in five points: above the shoulder, between the last thoracic vertebra and the first lumbar vertebra and at the level of sacral vertebra – I, II, III. Major cuts were divided into tissue elements: meat, skin with backfat and bones. Lean content of carcasses was determined on the basis of dissection.

Calculated were arithmetic means and their standard deviations. Two-way analysis of variance was performed using Duncan's test. STATISTICA 5.5 PL packet [9] was used.

### RESULTS AND DISCUSSION

Table 1 shows the frequency of genotypes CLPS and RYR1 genes. In whole crossbreds fatteners detected three genotypes of the CLPS/DdeI gene. Genotype AB was more frequent (0.56) than genotype AA (0.56) and BB (0.10). Similar frequency of genotypes colipase gene was obtained by Kurył et al. [7]. The least numerous was the homozygous BB (9.08% of research population) and the most numerous was heterozygous AB (57.27%). In investigations by Blicharski et al. [2] genotype AA was more frequent (69.80%) than BB genotype (2.01%) in pure breed Polish Large White populations.

Studied population of crossbred's fatteners was also differed as regards to RYR1 gene. Three genotypes – RYR1<sup>CC</sup>, RYR1<sup>CT</sup> and RYR1<sup>TT</sup> – were identified within tested populations. The most numerous were animals of RYR1<sup>CT</sup> (n=56) and the least those of genotype RYR1<sup>TT</sup> (n=21). Identification all genotypes of CLPS and RYR1 genes was profitable and make possible detailed analysis.

The main data traits connected with slaughter usability were presented on table 2. The percentage content of meat in carcasses at subgroups (CLPS genotype x RYR1 genotype) was higher in free of RYR1<sup>T</sup> (RYR1<sup>CC</sup>) animals with the BB genotype of CLPS than AA, P<0.05. There is a significant influence of major gene (RYR1) polymorphism on the quantitative traits. Higher meatiness was obtain at RYR1<sup>TT</sup> (55.16%) group than animals from least two groups (RYR1<sup>CT</sup> 51.87% and RYR1<sup>CC</sup> 50.74%), P<0.05. Similar results show Kortz

Table 2. Slaughter value of carcasses in relation to the CLPS and RYR1 genotypes  
Tabela 2 Wartość rzeźna tusz w zależności od genotypów CLPS oraz RYR1

Trait	RYR1 genotype	CLPS genotype			Average
		AA	AB	BB	
Carcass lean content, %	CC	49.36 <sup>a</sup> ± 4.73	50.98 ± 2.93	53.76 <sup>b</sup> ± 1.93	50.74 <sup>X</sup> ± 3.62
	CT	51.49 ± 2.80	52.60 ± 3.36	49.87 ± 2.67	51.87 <sup>X</sup> ± 3.30
	TT	55.74 ± 2.37	54.62 ± 3.24	59.13 ± 0.00	55.16 <sup>Y</sup> ± 3.05
Average		51.60 ± 3.85	52.53 ± 3.54	51.96 ± 3.81	-
Loin eye area, cm <sup>2</sup>	CC	41.71 ± 5.40	41.12 ± 3.63	42.70 ± 3.27	41.44 <sup>Xx</sup> ± 4.12
	CT	43.13 ± 5.32	44.59 ± 6.33	41.42 ± 4.28	43.67 <sup>y</sup> ± 5.77
	TT	48.77 ± 3.65	45.49 ± 4.35	54.40 ± 0.00	46.85 <sup>yy</sup> ± 4.57
Average		43.63 ± 5.51	43.67 ± 5.40	43.10 ± 5.36	-
Backfat thickness (average of 5 measurements), cm	CC	2.92 ± 0.38	3.07 ± 0.43	2.93 ± 0.48	3.01 <sup>x</sup> ± 0.41
	CT	3.00 ± 0.44	2.88 ± 0.54	3.26 ± 0.87	2.97 <sup>x</sup> ± 0.55
	TT	2.49 ± 0.26	2.75 ± 0.55	3.24 ± 0.00	2.70 <sup>y</sup> ± 0.50
Average		2.90 ± 0.43	2.91 ± 0.52	3.16 ± 0.70	-

Within rows means bearing different letter are related to CLPS genotypes; a, b at P<0.05; A, B at P<0.01;  
Within column means bearing different are related to RYR1 genotypes; x, y at P<0.05; X, Y at P<0.01

Table 3. Weight of neck and tissue content depending on *CLPS* and *RYR1* genotypes  
 Tabela 3 Masa i skład tkankowy karkówki w zależności od genotypów *CLPS* oraz *RYR1*

Trait	<i>RYR1</i> genotype	<i>CLPS</i> genotype			Average
		<i>AA</i>	<i>AB</i>	<i>BB</i>	
Neck weight, kg	<i>CC</i>	5.08 <sup>A</sup> ± 0.38	5.06 <sup>A</sup> ± 0.29	4.55 <sup>B</sup> ± 0.16	5.02 ± 0.34
	<i>CT</i>	5.12 ± 0.34	5.05 ± 0.31	4.91 ± 0.29	5.06 ± 0.32
	<i>TT</i>	5.27 ± 0.48	4.98 ± 0.49	4.94 ± 0.00	5.06 ± 0.48
Average		5.13 <sup>A</sup> ± 0.37	5.04 <sup>A</sup> ± 0.35	4.80 <sup>B</sup> ± 0.29	-
Meat in neck, %	<i>CC</i>	63.91 ± 2.73	63.94 ± 2.09	64.91 ± 1.55	64.11 <sup>xx</sup> ± 2.27
	<i>CT</i>	65.11 <sup>a</sup> ± 2.55	65.61 <sup>Aa</sup> ± 2.95	62.00 <sup>Bb</sup> ± 4.86	65.03 <sup>x</sup> ± 3.18
	<i>TT</i>	68.85 ± 1.35	65.75 ± 3.02	64.78 ± 0.00	66.59 <sup>yy</sup> ± 2.92
Average		65.39 <sup>a</sup> ± 2.88	65.10 ± 2.79	63.45 <sup>b</sup> ± 4.16	-
Backfat of neck, %	<i>CC</i>	22.58 ± 3.58	22.33 ± 2.21	20.71 ± 1.71	22.26 <sup>X</sup> ± 2.64
	<i>CT</i>	21.58 <sup>a</sup> ± 3.35	20.67 <sup>a</sup> ± 3.59	24.70 <sup>b</sup> ± 4.76	21.46 ± 3.76
	<i>TT</i>	17.55 ± 2.74	20.71 ± 3.38	21.95 ± 0.00	19.87 <sup>Y</sup> ± 3.41
Average		21.21 ± 3.64	21.21 ± 3.21	23.23 ± 4.12	-

Within rows means bearing different letter are related to *CLPS* genotypes; a, b at  $P \leq 0.05$ ; A, B at  $P \leq 0.01$ ;  
 Within column means bearing different are related to *RYR1* genotypes; x, y at  $P \leq 0.05$ ; X, Y at  $P \leq 0.01$

et al. [6] and Kapelański et al. [5]. Moreover, carcasses fatteners with *RYR1*<sup>TT</sup> (46.85 cm<sup>2</sup>) genotype had higher loin eye area and also thinner backfat thickness (2.70 cm) in comparison with other groups ( $P \leq 0.01$ ;  $P \leq 0.05$ ).

Table 3 presents weight of neck and tissue content depending on colipase and *RYR1* genotype. A significant effect of the colipase gene polymorphism was demonstrated with regard to these traits. Animals of *AA* and *AB* genotype showed higher weight neck (5.13 kg and 5.04 kg, respectively) in comparison with *BB* group (4.80 kg);  $P \leq 0.01$ . Higher content of meat in this cut had animals from *AA* group (65.39%) than animals from *BB* group (63.45%). According to research carried out by Kurył et al. [7], *CLPS/DdeI* genotype influenced significantly the weight of neck, neck with shoulder and also meat of neck.

Received results were analyzed in subgroups. Higher weight of neck had animals of *AA* and *AB* genotype in comparison with animals *CLPS/BB* in group of fattening pigs of *RYR1*<sup>CC</sup>,  $P \leq 0.01$ . Meat content in neck was higher in *AB* genotype animals (65.61%) and *AA* (65.11%) than *BB* (62.00%) only in animals *RYR1*<sup>CT</sup> group. Interesting it seems to be that the animals with arrangement of genotypes *TT AA* shows about 7% more meat in neck and less fat in this cut than animals with *CT BB* genotype ( $P \leq 0.01$ ).

The stress susceptibility gene polymorphism differentiated significantly the proportion of meat and fat in neck. Animals from *RYR1*<sup>TT</sup> group (66.59%) were characterized by larger content of meat in this cutting than animals from remaining two groups (*RYR1*<sup>CC</sup> 64.11% and *RYR1*<sup>CT</sup> 65.03%), and also lower fat content in neck (respectively: 19.87%, 21.46% and 22.64%).

Summarizing, the results of presented work obtained on a population of three-breed [*P* x (*PLW* x *PL*)] crossbred porkers, demonstrated a significant relation between the polymorphism in *CLPS* gene and the some important production traits. Interesting it seems to be that the polymorphism of *CLPS* gene influence on carcass lean content only in animals of genotype *CC/RYR1* but it should be treated as introductory information because of different number of genotypes in subgroup.

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