Association of BoLA-DRB3.2 alleles with fusobacteriosis in cows

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The Major Histocompatibility Complex (MHC) determines the immune response to pathogens, and its genes are promising candidates for the search of associations with diseases. A special role is played by BoLA-DRB3 gene, the product of which directly participates in the binding of alien antigens and conditions the specificity of the immune response. The second exon of this gene codes β-domain of class II antigens, which is necessary for binding a broad spectrum of alien antigens. Exon 2 of BoLA-DRB3 gene is extremely polymorphic, giving the possibility to search the associations of its alleles with various diseases. The article provides the results of the study on polymorphism of alleles of BoLA-DRB3.2 gene for detection of its associations with sensitivity to fusobacteriosis (necrobacteriosis) of cows. The survey was performed using PCR-RFLP method with DNA of blood from 176 cows of two herds of Ukrainian black-and-white dairy breed. As a result of the studies, in the first herd, 25 BoLA-DRB3.2 alleles were found. In the selections of necrobacteriosis susceptible and resistant cows, we found 22 and 21 variants respectively. In the second herd, in the general selection and group of healthy animals, 27 alleles were typed, and 22 in the group of susceptible cows. BoLA-DRB3.2*22 allele was the commonest in both herds in both general selections and groups of necrobacteriosis-resistant cows. In the selection of susceptible animals, the commonest was the variant BoLA-DRB3.2*16. We determined statistically significant associations of BoLA-DRB3.2 alleles with sensitivity to necrobacteriosis of cattle. BoLA-DRB3.2*03 and *22 alleles associate with necrobacteriosis-resistant, while *16 and *23 – with necrobacteriosis-susceptible cows of the both studied groups. Also, in the first herd, another allele was found – *24, indicating close relationship with the disease. The studies of polymorphism of BoLA-DRB3 gene expand the knowledge about genetic peculiarities of the Ukrainian black-and-white dairy breed. The identified molecular-genetic markers could be useful for breeders whose work is oriented towards the formation of herds which are resistant to diseases of the limbs in cattle.

Keywords: necrobacteriosis, Major Histocompatibility Complex; alleles; PCR-RFLP; lameness; cattle.

Introduction

Fusobacteriosis (necrobacteriosis) is a disease which tends to spread in the most countries of the world with developed animal husbandry. This infectious disease is characterized by purulent-neecrotic lesions mostly on the lower parts of the limbs, in some cases – in the oral cavity, in the udder, reproductive organs, liver, lungs, muscles and other tissues and organs (Vinnikov et al., 2019). The disease’s pathogen, Fusobacterium necrophorum, is a constant inhabitant of the gastrointestinal tract of both domestic and wild animals, common in the environment (animal housing, grazing areas, manure, soil, pastures, water from reservoirs without an outlet, etc.). The purpose of breeding dairy cattle is concentrated not only on the production characteristics, but also characteristics of health and life span, which necrobacteriosis directly affects. It causes death of up to 30% of adult cattle and 80% of young animals.

The main problem of necrobacteriosis in modern dairy herds is the need of lameness. It accounts for 40–60% of all diseases of the limbs of cattle (Skordoumou, 2007). Lesions of the limbs and hoofs are some of the main factors which drive dairy cows out of the working herd, comprising 10–15% of the unintended culling (Cha et al., 2010). Today, laminitis is the third cause, after mastitis and infertility (Hedges et al., 2001), of significant economic losses in dairy livestock farming (Ószvári, 2017). According to (Etterna & Østergaard, 2006), the losses from one case of lameness are estimated € 178–278. Losses from footrot are estimated in the range of $ 120–216, resulting in loss of 40% due to decrease in the milk production, 26% due to decrease of fertility and 34% for treatment (Cha et al., 2010). Recently, cattle more and more often have suffered from the hoof form of necrobacteriosis, because the mass transition to loose housing of animals caused specific problems, which farmers have not faced during stanchion-tied housing. Almost all domestic farms currently face the problem of necrobacteriosis. Spread of this disease often occurs in large farms housing highly productive animals. The causes of necrobacteriosis are multi-factorial: violations of the technology of animal husbandry and the norms of feeding of animals, uncontrolled import of cattle from abroad and mass holsteinization of the domestic breeds (Sokolovsky & Birukova, 2016; Suprovych et al., 2017).

Necrobacteriosis most often manifests as associated infection against the background of reduced resistance of the organism, immune deficiency of B-cellular link of the immunity, negative impact of exo- and endogenous stress factors on animals (Zhovnir et al., 2015). Therefore, there obviously is a necessity of the development of practical measures for the evaluation of genetic susceptibility (or by contrast resistibility) of the animals to necrobacteriosis.

Methods of molecular genetics combined with common tools of animal breeding are important for balancing the selection and optimization of the programs of breeding of animals. In scientific studies, the data on genetic aspects of necrobacteriosis are quite limited. Genetic selection for the reduction of hoof lesion is developing slowly, because the violations are not being recorded (Van der Spek et al., 2015). Systematic genetic evaluation of the health of hooves based on planned cleaning is being performed in some countries (Netherlands, Finland, Sweden, Canada) only since 2010 (Heringstad et al., 2018). Inheritance of the most often analyzed lesions of the hoof was usually low, ranging in the linear assessments 0.01–0.14 and threshold values 0.06–0.39. The assessed genetic correlations...
between the hoof lesions ranged 0.40–0.98 (Heringsstad et al., 2018). According to Boettcher et al. (1998), the assessment of inheritance of lameness equals around 0.1, which maximally reduces the attention of scientists to the genetic constituent of this problem. The assessment of laminitis inheritance, according to the results of the studies (Prakasa et al., 2019) was even lower (0.07 ± 0.01). One study (Okonomou et al., 2011) reports that for mastitis and laminitis, despite the low level of inheritance of these diseases, the selection based on genetic markers could be more important than the methods of typical selection. Svalve et al. (2014) revealed strong association of singular SNP on chromosome 21 with sensitivity to laminitis and assumes that genetic selection for improvement of resistance to laminitis is possible.

Currently, the molecular-genetic markers are essential in the sphere of animal genetics, because they allow determination of polymorphism at the level of DNA (DNA-markers). For cattle, there is a broad range of options of DNA-markers (Singh et al., 2014). During the selection of a marker needed for the identification of associations with different diseases of cattle, one should take into account its informativeness, the level of complexity of typing and cost of the work, therefore such studies should be conducted using a large number of animals.

From all the diversity of molecular-genetic markers, BoLA-DRB3 gene is unique. This is one of the genes of the Major Histocompatibility Complex (MHC). It is characterized by high level of polymorphism, associated with the formation of immune response of the organism to bacterial infections. Product of BoLA-DRB3 gene directly participates in binding of alien antigens and underlies the specificity of immune response. The second exon of BoLA-DRB3 gene which encodes Ii class β-domain is polymorphic due to the necessity of binding a broad range of alien antigens. The highest polymorphism of alleles of this gene indicates the possibility of high level of results from the search of possible markers associated with the diseases.

The significance of selection of BoLA-DRB3 gene as DNA-marker of sensitivity to various diseases of cattle is confirmed by the studies of its allele polymorphism in different breeds in many countries. The studies performed earlier allowed a large amount of data to be accumulated on the presence and the pattern of distribution of alleles and genotypes of BoLA-DRB3.2 gene for different populations, allowing performance of preliminary analysis regarding their use as DNA-markers for various diseases (Sulimova, 2006; Behl et al., 2012; Kumari et al., 2019). Variability of allele polymorphism in different breeds in many countries. The studies of sensitivity to various diseases of cattle, one should take into account its informativeness, the level of complexity of typing and cost of the work, therefore such studies should be conducted using a large number of animals.

The total amount of the mixture (20 µL) contained 60 nM Tris-HCL (pH 8.8), 2.5 mM MgCl₂, 20 mM KCl, 15 mM (NH₄)₂SO₄, 10 mM mercaptoethanol, 0.1% Triton X-100, 0.2 mM dNTP, 10 U of KlenTaq DNA polymerase, 10 µM of each primer, matrix DNA. For the first round, we used oligonucleotide primers:

HLO-30 (5'-3': TCCTCCTCTGACGACACTTCC),
HLO-31 (5'-3': ATTGCCGGTCCACTCCGGGCT); for the second round –
HLO-30 and HLO-32 (5'-3': TCCGGCTGGCAGTGAACACTTCC).

PCR-products of the first round (2 µL) were used for the second round. PCR was conducted in the Tertyk amplifier of DNA-Technology firm. The first period: denaturation of DNA at 95 °C during 5 min., then 10 cycles of denaturation (94 °C during 1 min.), hybridization of the primers (62.5 °C during 2 min.), DNA synthesis (72 °C during 1 min.) and the final synthesis (72 °C during 7 min.). The second period: the initial denaturation (95 °C during 5 min.), 35 cycles of denaturation, hybridization of the primers and synthesis of DNA (94, 68 and 72 °C during 0.5 min.), and final synthesis (72 °C during 7 min.). To avoid the contamination, the reaction mixture for all types of analyses was collected in a separate room using an individual set of pipettes. The final product of PCR in the amount of 5 µL was subjected to electrophoresis on 1.8% agarose gel to test the quality and specificity of the DNA amplification fragments.

The products of PCR were treated with three restriction endonucleases: Rsal, HaeIII, XhoII (Promega, New England BioLabs and SibEnzim). Fragments of the restriction were divided using electrophoresis in 2% agarose gel and 9% polyacrylamide gel in TBE electrophoresis buffer (TBE: 0.089 M of boric acid, 0.002 M EDTA pH 8.0). Electrophoresis of
the agarose gel was performed in the horizontal chamber for 1–3 hours at the power of 2 V/cm of gel. The gels were stained using ethidium bromide (0.5 μL/mL) for 10 min. with their subsequent rinsing in distilled water. Fragments of DNA were visualized in UV light on the transilluminator at the wavelength of 280 nm. To measure the length of the fragments, we used the marker of the molecular weights GeneRuler™ Ultra Low Range DNA Ladder manufactured by Fermentas, Latvia. As a result, analyzable electrophoreograms were developed (Fig. 1).

Fig. 1. Fragments of electrophoreograms of the products of the amplification of exon 2 of BoLA-DRB3 gene from DNA of cows of Ukrainian black-and-white breed using endonucleases RsaI (a), HaellII (b) and HhalII (c); M – marker of the molecular weights GeneRuler™ Ultra Low Range DNA Ladder; numbers of blood samples are indicated at the top.

Amplification of exon 2 with the following analysis of the length of the restriction fragments and comparison of the DNA samples obtained using the three values of the restriction endonucleases allows 54 alleles to be identified.

The frequencies of alleles were measured taking into account the number of homozygotes (N_a) and heterozygotes (N_d) found for the corresponding allele according to the formula (N – amount of the selection):

\[ P(A) = \frac{0.5N_a + N_d}{N} \]  

(1)

The strengths of the associative connection are determined based on the relative risk (RR) which indicates by how many times the danger of disease increases or decreases, having or not the corresponding alleles:

\[ RR = \frac{\frac{n_{a+b}}{n_a} \times \frac{n_{c+d}}{n_c}}{\frac{n_{a+c}}{n_a} \times \frac{n_{b+d}}{n_b}} \]

If there was observed a slight deviation according to \( \chi^2 \) one of the conditions of the formula 2 was not fulfilled, then to determine the strength of the association, we performed a test using the Fisher exact test (p) and Pearson coefficient (C).

We hereby declare all ethical standards have been respected in preparation of the submitted article. Permission to use animals was approved by the State Agrarian and Engineering University in Podilya in accordance with the European Convention for the Protection of Vertebrate Animals used for Experimental and other Scientific Purposes.

**Results**

Our study revealed respectively 27 (herd 1) and 25 (herd 2) alleles (Table 1). Most often (average value for two selections) BoLA-DRB3.*22 allele (11.37%) was found. The first ten of the commonest alleles includes 6 variants in each one: *01, *03, *07, *08, *16, *22, *23, *24 and *28. Their total frequency accounted for 47.7% in herd 1, and 53.6% in herd 2. In the second, there were seen characteristic peculiarities of the distribution of alleles in the groups of infected and healthy cows. In the selection of susceptible animals in the both herds, 22 alleles in each were found, of which the commonest variant was the variant with BoLA-DRB3.*216 (16.1%). In the first ten of the commonest, the same 6 alleles in each were observed: *01, *08, *16, *22, *23 and *24, total frequency of which accounted respectively for 56.0% in herd 1 and 56.3% in herd 2.

**Table 1** Frequency (%) of alleles of BoLA-DRB3.2 gene of Ukrainian black-and-white cows in the general selection, among those suffering from nectobacteriosis and healthy animals

| Alleles | General selection | infected | healthy |
|---------|------------------|----------|---------|
|         | herd 1 | herd 2 | herd 1 | herd 2 |
| *01     | (n = 85) | 4.71 | 9.34 | 2.63 | 5.00 | 6.38 | 12.80 |
| *03     | 7.06 | 7.14 | 2.63 | 2.50 | 10.60 | 10.80 |
| *06     | 1.76 | 2.20 | 1.32 | 3.75 | 2.13 | 0.98 |
| *07     | 4.71 | 3.30 | 1.32 | 2.50 | 7.45 | 3.92 |
| *08     | 11.20 | 4.95 | 9.21 | 7.50 | 12.80 | 2.94 |
| *10     | 0.00 | 3.85 | 0.00 | 6.25 | 0.00 | 1.96 |
| *11     | 6.47 | 0.00 | 5.26 | 0.00 | 7.45 | 0.00 |
| *12     | 2.35 | 1.65 | 2.63 | 0.00 | 2.13 | 2.94 |
| *13     | 2.35 | 2.20 | 3.95 | 2.50 | 1.06 | 1.96 |
| *14     | 0.59 | 2.75 | 0.00 | 5.00 | 1.06 | 0.98 |
| *16     | 9.41 | 8.24 | 15.80 | 16.30 | 4.26 | 1.96 |
| *18     | 1.76 | 2.75 | 2.63 | 5.00 | 1.06 | 0.98 |
| *21     | 1.18 | 1.65 | 1.32 | 2.50 | 1.06 | 0.98 |
| *22     | 12.40 | 10.40 | 5.26 | 5.00 | 18.10 | 14.70 |
| *23     | 5.29 | 7.69 | 9.21 | 15.00 | 2.13 | 1.96 |
| *24     | 8.24 | 9.34 | 14.50 | 7.50 | 3.19 | 10.80 |
| *26     | 1.18 | 2.20 | 0.00 | 2.50 | 2.13 | 1.96 |
| *28     | 8.24 | 6.59 | 9.21 | 3.75 | 7.45 | 8.82 |
| *32     | 1.18 | 1.10 | 0.00 | 0.00 | 2.13 | 1.96 |
| *33     | 1.76 | 0.00 | 2.63 | 0.00 | 1.06 | 0.96 |
| *36     | 1.18 | 2.75 | 1.32 | 1.25 | 1.06 | 3.02 |
| *37     | 0.59 | 2.75 | 1.32 | 1.25 | 0.00 | 3.02 |
| *39     | 1.18 | 0.00 | 2.63 | 0.00 | 0.00 | 0.00 |
| *45     | 3.53 | 0.00 | 1.32 | 0.00 | 5.32 | 0.00 |
| *51     | 1.18 | 0.00 | 2.63 | 0.00 | 0.00 | 0.00 |
| other   | 14.10 | 11.50 | 9.21 | 7.50 | 6.38 | 24.50 |

Note: other – alleles with the overall frequency of less than 2%.

In the groups of the nectobacteriosis-resistant cows, there were found respectively 21 and 27 alleles, of which the commonest was BoLA-DRB3.*22 allele (16.3%). The first ten of the commonest alleles included the same 6 variants in each: *03, *07, *08, *22, *23 and *28. Their total frequency was 59.6% in the herd 1, and 52.0% in the herd 2.

The main goal of the study was determining the molecular-genetic markers based on the alleles of gene BoLA-DRB3.2 of the nectobacteriosis-susceptible animals. Tables 2 and 3 demonstrate the results of the biometric calculations for alleles which have statistically significant associations with the diseases or resistance to it and may be potentially the DNA-markers of susceptibility or resistance to nectobacteriosis.

**Table 2** Alleles of gene of Ukrainian black-and-white dairy cows susceptible to nectobacteriosis (herd 1)

| Alleles | P(A) | RR | \( \chi^2 \) | Check for limited sample size |
|---------|------|----|-------------|-----------------------------|
|         |      |    |             | N | N | N | N |
| *03     | 0.071 | 4.87 | 4.44 | 5.36 | 55.1 | 32.6 | 40.4 |
| *16     | 0.094 | 4.96 | 7.32 | 7.15 | 10.40 | 30.9 | 38.2 |
| *22     | 0.124 | 4.82 | 7.43 | 9.39 | 8.40 | 28.6 | 35.4 |
| *24     | 0.082 | 5.98 | 7.78 | 6.26 | 9.06 | 31.7 | 39.3 |

In both selections, two alleles in each were found, indicating the statistical significance (\( \chi^2 > 3.84 \)) regarding susceptibility (RR ≥ 2) or resistance (RR ≤ 2) of the cows to nectobacteriosis.

In the first herd, BoLA-DRB3.*216 (P < 0.01) and *24 alleles (P < 0.01) indicate close relationship with the pathology. With the resistance to the diseases, BoLA-DRB3.2 *03 (P < 0.05) and *22 alleles (P < 0.01)
were related. In the second herd, the BoLA-DRB3.2*03 (P < 0.05) and *22 alleles (P < 0.05) manifested associative relationship with the resistance to nectobacteriosis, and allele BoLA-DRB3.2*16 and BoLA-DRB3.2*23 showed quite strong association (P < 0.001) with the disease.

Table 3
Alleles of BoLA-DRB3.2 gene of cows of Ukrainian black-and-white dairy breed susceptible to nectobacteriosis (herd 2)

| Alleles          | P(A)  | RR  | \( \chi^2 \) | (a+b+c+d) | (a+b+c+d) | (a+b+c+d) | (a+b+c+d) |
|------------------|-------|-----|-------------|-----------|-----------|-----------|-----------|
| *08              | 0.071 | 5.23| 5.03        | 0.03      | 5.71      | 6.0       | 34.3      | 43.7      |
| *16              | 0.082 | 11.8| 13.30       | 0.09      | 6.59      | 10.2      | 33.4      | 42.6      |
| *22              | 0.014 | 3.75| 5.11        | 0.22      | 8.35      | 8.35      | 31.7      | 40.4      |
| *23              | 0.077 | 10.5| 11.70       | 0.06      | 6.15      | 9.38      | 33.9      | 43.2      |

Note: *22 – parameters of which the value exceeds 0.05.

The analysis of the biometric parameters of others alleles showed at least two variants worth attention regarding the possibility of using them as molecular-genetic markers among the alleles found in this study (Table 4).

Table 4
Alleles of BoLA-DRB3.2 gene of Ukrainian black-and-white dairy cows susceptible to nectobacteriosis which have limitations during detection of the associations

| Alleles | P(A)  | RR  | \( \chi^2 \) | (a+b+c+d) | (a+b+c+d) | (a+b+c+d) | (a+b+c+d) |
|---------|-------|-----|-------------|-----------|-----------|-----------|-----------|
| *23 (hand 1) | 0.053 | 5.06| 4.45        | 4.06**    | 5.51      | 34.0      | 42.0      |
| *01 (hand 2) | 0.093 | 3.08| 3.54**      | 4.74      | 7.85      | 32.5      | 41.5      |

Note: ** – parameters of which the value exceeds the acceptable thresholds.

BoLA-DRB3.2*01 allele is limited from the perspective of the biological studies (\( \chi^2 < 3.83; P = 0.06 \)), and BoLA-DRB3.2*23 allele did not stand up to the test for the size of selection due to low number of the animals with the variants a = 7 and b = 2.

For the evaluation of the statistical significance of the strength of the association of the sensitivity to nectobacteriosis, Fisher’s exact test was performed with testing the strength of the association according to the Pearson coefficient. For allele *23, we determined that \( P = 0.031 \) and \( C = 0.223 \), and for allele *01 – \( P = 0.038 \) and \( C = 0.194 \). Both variants have the test with the exact criterion (P < 0.05), allowing us to propose them for the practical use as DNA-markers. However, mean and weak strength of the association according to the Pearson coefficient suggests that the approach in interpreting such opportunity must be cautious. For the final decision on the possibility of using these alleles as DNA-markers associated with nectobacteriosis, additional studies should be undertaken.

Discussion

The high level of polymorphism and heterozygosity of BoLA-DRB3 gene helps the immune system to react to a broad range of pathogens. The uniqueness of this area of DNA has concentrated the attention of scientists. Over the recent 20 years, using PCR-RFLP and PCR-SBT, allele spectrum of over 30 breeds of cattle around the globe has been identified (Ramnubjør et al., 2017). Currently, 330 DRB3-alleles have been recorded in the Immuno Polymorphism Database (IPD)-MHC database (www.ebi.ac.uk/ipd/mhc/group/BoLA). Most studies focused on the identification of associations of BoLA-DRB3.2 alleles and the following diseases: mastitis (Kulberg et al., 2007), leucosis (Gutiérrez et al., 2017), hoof-and-mouth disease (Lei et al., 2012), diseases caused by Acari (Dua-...

The Ukrainian black-and-white breed is diverse in genealogical structure, which is confirmed by the results of typing of blood samples of both herds. Polymorphism of alleles of BoLA-DRB3.2 gene ranges within a broad spectrum, both according to the quantity and the nomenclature. In our study, in the herd 1, 27 variants were found, while in the herd 2 – 25. This is an average indicator. A maximum of 35 alleles of a possible 54 PCR-RFLP variants were identified for Mongolian cattle (Ruzina et al., 2010). Greater diversity was characteristic of only the Filipino cattle, in which 71 alleles were detected using PCR-SBT method (Takeshima et al., 2014). According to the review (Behl et al., 2012), the lowest polymorphism was determined for Jersey and Rath breed, 13 alleles of BoLA-DRB3.2 gene for each. It should be noted that the general frequency of the commonest alleles in cows of the Ukrainian black-and-white breed was lower than in other industrial breeds. The high level of consolidation of alleles indicates impoverishment of the genetic diversity of the breed. The pressure from breeders leads to allele funds of different breeds becoming more similar and genetically homogeneous with time. In our study, the total frequency of such alleles was within 50–60%, whereas the accumulation of alleles most often detected in American Holstein cattle accounted for 79.6% (Dietz et al., 1997), and 91.3% in Canadian cattle (Sharif et al., 1998).

The data of the study on the associations of MHC genes with the limb disorders of ruminants are quite limited. Genetic advantage is determined by the heredity of a feature, intensity of the selection, reliability of the selective values and the interval of the generation. Even if the heredity is low, genetic differences exist (Van der Linde et al., 2010). This is indicated by the results of the studies we published earlier (Suprovych et al., 2016) regarding the associations of alleles of BoLA-DRB3.2 gene with the nectobacteriosis of Ukrainian black-and-white breed on the basis of the breeding herd Kozatska Dolya 2006, performed using the methods and on the basis of the Laboratory of Genetics of the Vavilov Institute of General Genetics (Moscow). The allele polymorphism was studied on the blood samples from 114 cows (43 animals diagnosed nectobacteriosis).

Total of 4 alleles were determined to have susceptibility to the disease: *16 (RR = 24.10; P < 0.001), *23 (RR = 4.41; P < 0.05), *03 (RR = 7.70; P < 0.05) and *22 (RR = 3.57; P < 0.05). It has to be noted (Suprovych, 2013) that BoLA-DRB3.2*22 allele which manifested itself as a marker of resistance to nectobacteriosis has also a close relationship with the resistance to mastitis in cows of Ukrainian black-and-white (RR = 2.52, P < 0.05) and red-and-white dairy breeds (RR = 4.66, P < 0.001).

The relationship between mastitis and lameness of cows is reported in the study undertaken in 10 dairy farms in South-West England (Peeler et al., 1994). The assessment of the relationships between the periphery diseases of 3,603 cows over three lactations using logistic regression revealed that the clinical lameness is related to 1.4 times increase in the chances of clinical mastitis.

Currently, among other studies, only one work is known which was oriented towards the identification of associations of BoLA-DRB3 gene with laminitis in a dairy breed, performed by Chinese scientists (Sun et al., 2013). They studied the relationship between polymorphism of gene and lameness of Chinese Holstein cows (n = 435) and came to the conclusion that RFLP alleles of exon 2 may be candidates of susceptibility to limb diseases in that population. The study was based on identifying genotypes of cows using the restrictases BstUI, BstYI and HaeIII (MBI fermentas, China). Analysis of seven RFLP genotypes with HaeIII restriction showed presence of statistically significant associations with laminitis in the Chinese cattle.

Among the reports on search of genetic markers in relation to lameness of cattle, we should draw attention to the studies (Buitenhuis et al., 2007) where the genome was scanned for QTL on the basis of 29 autosomes using microsatellite markers and (Okononmoyou et al., 2011) who studied the influence of polymorphism of loci of STAT5A and FGF2 genes on the diseases of limbs of Holstein cows. The study (Ferchichi et al., 2018a) presents the results of research of the genetic polymorphism of FGF-2 gene and its influence on the lameness of Holstein cows in Tunisia. In another work (Ferchichi et al., 2018b), the influence of polymorphism of leptin gene on the lameness of cows (n = 412) was studied. The researches revealed that the animals with AB genotype have less lameness than cows with homozygote genotypes BB and AA.

Genome-wide association study showed significant amount of SNP associated with different disorders of hooves. In general, 11 significant and 46 suggestive SNP-associations related to the pathology of hooves on 20 chromosomes were found (Van der Spek et al., 2015). Also, there is a report about the search of similar associations in sheep. The animals were
selected into the groups of correspondence to the inherited paternal haplo-
type (Escayg et al., 1997) for studying the relationships between MHC locus of class II and the status of footrot. The study revealed the associa-
tions of MHC haplotype and the status of the disease (P = 0.005). Another research work (Vailionis et al., 2016) provides the data of studying the polymorphisms of Ovar-DRB1 and BMC5221 genes for the identifica-
tion of animals resistant to lameness (footrot) among Chezrel sheep. Statis-
tical analysis of Ovar-DRB1 alleles demonstrated high frequencies of FF and AA genotypes in animals resistant to lameness (P < 0.01), whereas more common frequencies of DD genotype were seen in susceptible animals (P < 0.01). In BMC5221 locus, higher frequencies of A1A3 genotype were observed in the animals resistant to lameness (P < 0.01), while higher frequencies of A3A3 and A2A2 genotypes were observed in susceptible animals (P < 0.01).

The future of selective breeding of cattle cannot be based only on the short-term purposes oriented towards the profitability of the production of animal origin. Traditionally, the programs of livestock farming had a special focus on the genetic improvement of economically important characteristics: milk, fat and protein. Genetic progress has been quite successful regarding these indicators. After introduction of complex models of genetic assessment and improvement of genetic selection, milk production increased more than twice over the last 40 years. But as the genetic ability of cows to produce milk heightened, their health and wel-
fare were significantly damaged. Today, dairy cows suffer from serious problems such as mastitis, lameness and infertility. Also, the productive period of use of these animals decreased. This means that currently, main-
taining the level of usefulness requires higher standards of management and costs.

The fact that the actual (phenotypical) relationships between health and production, usually, are much lower than the genetic ones indicates that this problem is being successfully solved. However, if genetics behind health- and life span-related problems is ignored, the welfare of herds will decrease below the acceptable level. Therefore, the genetic data on the peculiarities of health and welfare of animals is becoming more and more valuable. Moreover, the genetic improvements are very economical, constant and accumulate over generations. Finally, the accurately imple-
mented genome technologies using DNA-markers, including the ones based on alleles of BoLA-DRB3.2 gene, could give new solutions of this increasingly complex problem of animal selective breeding, where we often face controversial genetic wishes of maximum output of animal pro-
duction compared with resistance to diseases.

Conclusions

We found associations of alleles of exon 2 of BoLA-DRB3 gene with nectobacteriosis in cattle. BoLA-DRB3.2*03 and *22 alleles associated with resistance, while *16 and *23 are associated with susceptibility to nectobacteriosis of cows in both studied herds. Furthermore, the first herd was found to have one more allele *24, which indicates strong relationship with the disease. Similar results were obtained in earlier studies on the Ukrainian black-and-white dairy breed, confirming the genetic relation between BoLA-DRB3 gene and nectobacteriosis.

It is determined that molecular-genetic markers are potentially useful, because they allow controlling possible manifestations of nectobacteriosis, which is the main cause of lameness of dairy cows, at the early stages of ontogenesis. Genetic selection for resistance to nectobacteriosis and health improvement of dairy herds against lameness could be based on the found BoLA-DRB3 alleles.

The results of the study once again confirm the importance of MHC in the animal husbandry and significantly extend our knowledge of diver-
sity and polymorphism in this locus. To obtain knowledge about the role of gene in the mechanism of immu-
ne protection against limb diseases caused by nectobacteriosis, other breeds of cattle need to be additionally studied.

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