Introduction

Modern breeding of dairy cattle increasingly involves programmes based on the international trade of semen from elite bulls with high genetic merit. With the widespread use of advanced reproductive technologies, including artificial insemination and multiple ovulation embryo transfer, individual bulls are able to sire thousands of heifers in many countries (1). In that way, more than ten genetic diseases have spread, some of them resulting in fatal outcomes and causing significant economic losses. These include bovine leukocyte adhesion deficiency (BLAD), deficiency of uridine monophosphate synthase (DUMPS), and complex vertebral malformation (CVM) (2, 3, 4, 5, 6, 7).

With intensive livestock development, a single breed begins to dominate in separate animal species (e.g., the Holstein breed) among dairy cattle, and a small number of reproducers are used for fertilisation in such breeds. Consequently, genetic biodiversity is significantly reduced, many genes became homozygous, and abundant genetic diseases can occur. Animals carrying various genetic defects must be eliminated from the population so that the frequency of the undesired...
gene does not increase to the critical limit, at which point it becomes homozygous and manifests phenotypically. If the animal is only the carrier of unwanted genes, this gene is not phenotypically detectable, and the study can only be performed using genetic testing methods, so it is essential to analyse breeding livestock in order to prevent the spread of the undesired gene.

Known inherited disorders in cattle are mostly caused by autosomal recessive inherited genes. Heterozygous individuals can be identified using different methods, such as examination of progeny, clinical examination or necropsy, analysis of enzyme activity in blood, and genotyping of animals by genomic analysis. Recent developments within molecular genetics have made possible the efficient and rapid identification of heterozygous animals via genomic analysis. Knowing the molecular basis of a defect, the direct detection of carriers is possible at the genetic level, thus preventing the breeding of the ineligible animal (8). At present, there are identification records for several inherited bovine disorders, including bovine leukocyte adhesion deficiency (BLAD), deficiency of uridine monophosphate synthase (DUMPS), complex vertebral malformation (CVM), bovine citrullinaemia (BC), and factor XI deficiency (FXID).

The congenital calf immune deficiency BLAD is a lethal autosomal recessive hereditary disease manifested in Holstein breed calves. The disease was diagnosed for the first time by the American scientist Marco E. Kehrli in 1990 (9). Dale E. Shuster and other researchers have identified the CD18 gene nucleotide sequence of the healthy bovine and mutated CD18 gene (10). The mutation of the same gene was found in humans (11) and dogs (12). Heterozygous individuals are clinically healthy, but heterozygous bulls and cows have a 25% probability of producing homozygous calves with this disease. The BLAD disease gene is present in the BTA1 bovine chromosome. Due to the mutation, the gene responsible for the synthesis of glycoprotein-beta-integrin has been altered. This results in the synthesis of inadequate beta-integrin as in the position 128 of the protein molecule, the aspartic amine acid is replaced with glycine. Calves with two mutated alleles, which in homozygote status causes bovine BLAD, often have infections in the gastrointestinal tract and respiratory tract. Calves with BLAD are born with small body weight, grow and eat poorly, and their coat is not shiny. Calves are highly prone to infections, and are often suffering from enteritis, pneumonia, diarrhoea, ulcers, laryngitis, granulation stomatitis, gingivitis, peripheral lymphadenopathy, anaemia, and, if not treated, they usually die at 2-3 months of age. From the year 2000 to 2018 in Lithuania, the frequency of mutated allele, which in homozygote status causes bovine BLAD, decreased because of the intensive selection against this allele.

The deficiency of uridine monophosphate synthase (DUMPS) is an autosomal recessive lethal Holstein breed bovine disease characterised by early embryonic mortality (13, 14, 15). UMPS (uridine monophosphate synthase) is an enzyme that is essential in the synthesis of pyrimidine nucleotides. It is indispensable for the normal growth and development of ruminants and other species of animals. The inactivation of this enzyme is caused by a point mutation in the UMPS gene (C → T), 405 codon of the 5 exon. This Holstein cattle disorder is characterised by a decrease of UMPS enzyme activity in the blood (16). DUMPS causes foetal death in the early stages of pregnancy and many reproductive problems in dairy herds. The gene responsible for lack of the enzyme uridine monophosphate is found on chromosome 1 of bovine animals. In mammalian cells, the final step of the pyrimidine nucleotide synthesis comprises the conversion of orotate to urinary monophosphate and is catalysed by the UMP synthase enzyme. UMP synthase is essential for the synthesis of novel pyrimidine nucleotides, which are components of DNA and RNA. Since pyrimidines are essential for nucleic acid synthesis during embryonic development, embryos that are homozygous for a recessive allele die up to the 40th days of gestation. Embryos are often absorbed during the first two months of pregnancy, and these cows return to oestrus. This leads to increased intervals between calving (13, 14, 15). Heterozygous animals are phenotypically normal, but only half of normal UMPS enzyme activity occurs, which causes an increase in the amount of orotic acid in their milk and urine (16). To date, no live animals have been found that are homozygous for the mutated allele (15).

Complex Vertebral Malformation (CVM) is an autosomal recessive Holstein cattle disease inherited as severe spinal degeneration. The syndrome was first identified in the year 2000 in the Danish Holstein population (3, 17, 18). The
disease was also found in the Wagyu breed. The first ancestor of the cattle that had this mutation was a bull named Carlin-M Ivanhoe Bell (19). Severe Vertebral Malformation (CVM) is determined with both aborted foetuses and prematurely born dead calves. Affected calves have abnormalities in the spinal column, such as not fully developed, broken, or unusually formed spinal vertebrae and ribs, scoliosis, and spinal synostosis. Low body weight and heart abnormalities are also observed (20). Such calves have shorter spine and chest areas, symmetrical contractions of meta-tarso-phalange joints on both sides and symmetrical arthrogryposis (17). Severe Vertebral Malformation (CVM) is caused by a mutation in the BTA3 chromosome, which replaces the amino acid sequence; instead of valine, phenylalanine is formed, in the 180 position of the protein uridine 5’-diphosphate-N-acetylglucosamine (21). This gene is responsible for the transportation of UDP-N-acetylglucosamine into the Golgi apparatus membrane; 80% of embryos that have inherited a gene mutation of CVM disease from both parents will be lost during the first three months of gestation. An aborted sick calve will have a shortened neck due to spinal cord injury, as well as altered ribs, limbs and interdigital joints. In addition, symptoms such as partial pulmonary hypoplasia, excessive liver segmentation, double gall bladder, as well as rectum and uterine atresia may occur (22). Heterozygous animals are carriers of the mutated gene.

In animal breeding, genetic disorders are one of the most imperative issues for breeders. Due to the negative influence of such disorders on animals, through abnormal anatomy or reduced production, breeders and breeding associations need to control the impact on the population.

Material and methods

The animals for this study were selected from dairy herds, in which breeding record and cattle productivity control were carried out. The farms were located in four different regions of Lithuania. In each dairy herd, 50 non-related (based on the documentation of origin) cattle (cows and heifers) were selected. In total, 200 dairy cattle were tested. They were healthy, and kept and fed according to hygiene norms; the conditions for keeping and feeding were in line with veterinary requirements.

Blood for genetic testing was collected in aseptic conditions from the cephalic vein in vacuum tubes (Vacutainer) with K2 EDTA (ethylenediamine tetraacetate) anticoagulant. The blood was stored at +4 °C until the test. DNA extraction from blood leukocytes was performed using the chloroform salt method, according to Miller et al. (23). Genomic DNA content and purity were determined using the spectrophotometric method (DNA/RNA Reader, Pharmacia).

**BLAD genetic disease PCR-RFLP test method (10)**

For the identification of mutation c.383A>G (g.145114963A>G, rs445709131) in the CD18 gene, causing a change of adenine to guanine in the 383 position of the gene and manifesting as BLAD in cattle, the PCR-RFLP test method was used. We targeted a 357 bp long fragment of the fifth exon of the gene of the BLAD disease locus (Table 1).

10µl of DNA and 15µl of PCR mixture were poured into the tube. The PCR reaction was performed using a thermocycler (G-storm, United Kingdom). Reagents used: PCR Mix - 2.95µl ddH2O; 2.5µl 10xPCR buffer; 2.5µl of dNTP (2 mM); 1.5µl of MgCl2 (50 mM); 2.5µl of the forward primer (20 pmol); 2.5µl of the reverse primer (20 pmol); 0.25ml BSA; 0.3µl Taq polymerase (Thermo Scientific, Lithuania); in two places in the same column 10µl of the PCR product was digested with 10µl of the restriction mix (7.5µl ddH2O, 2µl 10 x buf., 0.5µl TaqI). Samples were left in a thermostat for 1 hour at 65 °C. The digested PCR products were fractionated using electrophoresis in 2.5% agarose gel, 100 V 40 min. The gel was stained with ethidium bromide for 15-20 minutes and analysed in UV light (wavelength 300 nm) with MiniBisPro Video Documentation (Herolab) (Table2).

**DUMPS genetic disease PCR-RFLP test method**

Testing method for nonsense (stop-gain) mutation c.1213 C>T in UMPS gene (g.69756880C>T) causing a change of cytosine to thymine in the 405 codon of the gene of the fifth exon and manifesting as DUMPS in cattle (Table 3).

10µl of DNA and 15µl of PCR mixture were poured into the tube. PCR reaction was performed
Table 1: Primers, PCR profile, PCR product size and restriction enzymes used for identification of genetic disease – BLAD

<table>
<thead>
<tr>
<th>Genetic defect</th>
<th>Primers</th>
<th>PCR profile</th>
<th>PCR product size</th>
<th>Restriction enzyme</th>
</tr>
</thead>
<tbody>
<tr>
<td>BLAD</td>
<td>F: 5’ GAAATGGCATCCTGCATCATCCACCA 3’ R: 5’ CTGAGGTTCCAGGGGAAGATGAGTAG 3’</td>
<td>94°C 3 min</td>
<td>357 bp</td>
<td>TaqI</td>
</tr>
<tr>
<td></td>
<td></td>
<td>94°C 30 s 33 cycles</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>65°C 30 s</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>72°C 30 s</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>72°C 5 min</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Table 2: CD18 gene c.383A>G DNA fragments sizes in bp after digestion with restriction endonuclease

<table>
<thead>
<tr>
<th>Bp</th>
<th>Homozygous for normal allele</th>
<th>Heterozygous</th>
<th>Homozygous for disease allele</th>
</tr>
</thead>
<tbody>
<tr>
<td>357</td>
<td>—</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>201</td>
<td>—</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>156</td>
<td>—</td>
<td>—</td>
<td>—</td>
</tr>
</tbody>
</table>

Table 3: Primers, PCR profile, PCR product size and restriction enzyme used for identification of genetic disease – DUMPS

<table>
<thead>
<tr>
<th>Genetic defect</th>
<th>Primers</th>
<th>PCR profile</th>
<th>PCR product size</th>
<th>Restriction enzyme</th>
</tr>
</thead>
<tbody>
<tr>
<td>DUMPS</td>
<td>F: 5’ GCAAATGGCTGAACATTCTG 3’ R: 5’ GCTTCTAACTGAACTCCTG 3’</td>
<td>94°C 5 min</td>
<td>108 bp</td>
<td>AvaI</td>
</tr>
<tr>
<td></td>
<td></td>
<td>94°C 60 s 40 cycles</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>58°C 60 s</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>72°C 90 s</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>72°C 5 min</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Table 4: UMPS gene c.1213 C>T DNA fragments sizes in bp of cattle DUMPS causing gene after digestion with restriction endonuclease

<table>
<thead>
<tr>
<th>Bp</th>
<th>Homozygous for normal allele</th>
<th>Heterozygous</th>
<th>Homozygous for disease allele</th>
</tr>
</thead>
<tbody>
<tr>
<td>89</td>
<td>—</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>53</td>
<td>—</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>36</td>
<td>—</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>19</td>
<td>—</td>
<td>—</td>
<td>—</td>
</tr>
</tbody>
</table>

Table 5: Prevalence of BLAD, DUMPS and CVM diseases in Lithuanian Holstein population

<table>
<thead>
<tr>
<th>Genetic disease</th>
<th>Gene</th>
<th>Genotype frequency</th>
<th>Allele frequency</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>Homozygous for normal allele</td>
<td>Heterozygous</td>
</tr>
<tr>
<td>BLAD</td>
<td>CD18</td>
<td>0.995</td>
<td>0.005</td>
</tr>
<tr>
<td>CVM</td>
<td>SLC35A3</td>
<td>0.99</td>
<td>0.01</td>
</tr>
<tr>
<td>DUMPS</td>
<td>UMPS</td>
<td>1</td>
<td>0</td>
</tr>
</tbody>
</table>
Detection of bovine leukocyte adhesion deficiency, deficiency of uridine monophosphate synthase, and complex vertebral malformation (CVM) using a thermocycler (G-storm, United Kingdom). Reagents used: PCR Mix - 2.95ml ddH₂O; 2.5ml 10xPCR buffer; 2.5ml of dNTP (2 mM); 1.5ml of MgCl₂; (50 mM); 2.5ml of the forward primer (20 pmol); 2.5ml reverse primer (20 pmol); 0.25ml BSA; 0.3ml Taq polymerase (Thermo Scientific, Lithuania). 10μl of the PCR product was digested with 10μl of the restriction mix (7.5μl ddH₂O, 2μl 10 × buf., 0.5μl AvaI). Samples were left in a thermostat for 10 hours at 37 °C. The digested PCR products were fractionated using electrophoresis on 4% agarose gel, 100 V 40 min. The gel was stained with ethidium bromide for 15-20 minutes and analysed in UV light (wavelength 300 nm) using MiniBisPro Video Documentation (Herolab) (Table 4).

CVM genetic disease test method

Investigations of polymorphism analysis of the gene for CVM disease in cattle was performed at the Neogen Genomics Corporation Laboratory (USA). The study was performed using a sequencing method. CVM disease is determined by the missense mutation, present in the SLC35A3 gene. In the third cattle chromosome, a modified SLC35A3 gene with a single base conversion c.538G>T (g.43412427G>T, rs438228855) was detected.

Results

We tested cows for lethal autosomal recessive disorders widely distributed in the Lithuanian Holstein cattle population: bovine leukocyte adhesion deficiency (BLAD), deficiency of uridine monophosphate synthesis (DUMPS) and vertebral malformation (CVM). A recessive allele with point mutation c.383A>G, causing BLAD, was found in the Lithuanian cattle population with 0.0025 frequency; 0.5% of individuals were heterozygous while no homozygous animals with BLAD were found. Since 1999, all sires and selected cows used for breeding in Lithuania have the genotype for this disease recorded in the pedigree information. In Lithuania in the year 2000, 6.7% of key areas of the altered gene leading to BLAD disease were found among selected cows. Of the 146 bulls tested, 4 carriers were found. In 2002, 3% of selected cows were carriers, and in 2004, 2% of selected cows and 1 young bull were carriers. All breeding bulls and selected cows that are newly introduced among breeding animals are compulsorily tested for BLAD disorder. A similar mutated allele frequency as in our study was reported in Czech cattle: 0.82% (24), and in Chinese Holsteins (0.69%; 25). However, a much larger mutated allele frequency was reported for Indian Holstein Friesian cattle (2.99%) (26),

In the year 2000, the percentage of Lithuanian dairy cattle population who were heterozygote animals, BLAD disease gene carriers, was 5%; this percentage was 4% in 2002, and 2% in 2004. In 2018, we found 0.5% mutated allele carriers and distributors of disease genes in the population (Figure 1).

Discussion

Genomic selection is carried out not only to improve the genetic potential of cattle productivity, reproduction, and milk quality but also by assessing the signs of bovine health. In the dairy cattle population, due to the international breeding of males with high genetic value, despite also being the carriers of recessive genetic disease alleles, genetic diversity has been significantly reduced. As a result, many genes have become homozygous, producing abundant genetic diseases among the dairy cattle population, which causes significant economic losses, as many of them are lethal. After a retrospective assessment of the prevalence of the lethal genetic diseases of BLAD, DUMPS, and CVM in the dairy cattle population, the following was established.

CVM and BLAD have become some of the most common hereditary genetic defects of the Holstein breed in recent decades. Researchers found that in the year 2000 the mutant allele for BLAD disease frequency was 24%, and the rate of mutated alleles for CVM in the German Holstein population from 2001 to 2007 ranged from 9% up to 16% (6).

The mutated allele causing BLAD, c.383A>G, was found at a frequency of 0.0025% in the Lithuanian dairy cattle population; with 0.5 % of cattle being heterozygous while no homozygous animals with BLAD were found. Since 1999, all sires and selected cows used for breeding in Lithuania have the genotype for this disease recorded in the pedigree information. In Lithuania in the year 2000, 6.7% of key areas of the altered gene leading to BLAD disease were found among selected cows. Of the 146 bulls tested, 4 carriers were found. In 2002, 3% of selected cows were carriers, and in 2004, 2% of selected cows and 1 young bull were carriers. All breeding bulls and selected cows that are newly introduced among breeding animals are compulsorily tested for BLAD disorder. A similar mutated allele frequency as in our study was reported in Czech cattle: 0.82% (24), and in Chinese Holsteins (0.69%; 25). However, a much larger mutated allele frequency was reported for Indian Holstein Friesian cattle (2.99%) (26),
Iranian Holsteins (3.3%) (27), American Holsteins (8.2%), Polish Holsteins-Friesian (7.9%) (28) and Turkish Holstein cows (4.0%) (29).

The mutated allele causing the DUMPS genetic disorder in cattle (c.1213 C>T), manifested as early mortality of cattle embryos due to the shortage of uridine monophosphate synthase, was not found in the Lithuanian population. No carrier animals of the DUMPS genetic disorder were found in Turkey (2, 29, 30). Similar results were obtained in the research done by other scientists in Poland (13), the Czech Republic (24), Germany, India, Iran and Romania. However, the mutant DUMPS disease allele with a 1–2% frequency was found in the Holstein breed in the USA, 0.96% in Argentine Holstein bulls, and 0.06% in Chinese Holsteins.

The mutated allele causing CVM cattle disease (c.538G>T), manifested in malformations of the foetal spine, was found in the population of Lithuanian dairy animals at the frequency 0.005, while the number of heterozygous bovines was found to be 1%, and no homozygous bovine affected by CVM was found. In Turkey, the frequency of Holstein bovine CVM mutated allele carriers was 3.4% and 3.86% in the Chinese dairy population. A high frequency of mutated allele was found in Denmark with 31.0% of all cattle affected (21), in Poland at 24.8% (31), in Japan at 32.5% (32), in Sweden at 23.0% (33), in Germany, at 13.2% (34), and in China 15% (19). Since the year 2000, breeding programmes in most counties have been implemented to reduce the prevalence of CVM carriers. However, in some Holstein populations, the incidence of CVM disorder carriers remains high (Denmark, Poland, Japan). A study of Iranian Holstein cattle did not identify heterozygous bovine animals (18). This can be associated with the use of a small number of bulls carrying the mutated allele for breeding, during the formation of the breed, and further selection.

In the Lithuanian dairy cattle population, lethal bovine diseases caused by recessive genes in the population were reduced because of an intense selection programme, eliminating the BLAD disease gene carriers and avoiding the use of BLAD, DUMPS, and CVM heterozygous bulls for breeding. The number of mutated allele carriers dropped from 6% in 2000 to 0.5% in 2018.

The incidence of hereditary diseases causes not only direct economic losses to livestock breeders but also leads to reductions in the genetic diversity of animal populations as a result of extensive culling of disease carriers. The identification of carriers with the use of molecular diagnostic tests is an important step in reducing the frequency of detrimental alleles and consequently lowering the incidence of hereditary diseases in the herd. This is particularly important when considering bulls inserted into progeny testing programmes because they can potentially sire thousands of progeny before the incidence of affected progeny can be associated with a particular animal.

**Conclusions**

An investigation of cows for the prevalence of the lethal genetic disorders BLAD, DUMPS, and CVM in the Lithuanian dairy cattle population showed 0.5% BLAD disease gene carriers and 1.0% CVM disease gene carriers. No DUMPS disease gene carriers have been found. In the Lithuanian Holstein cattle population, the number of carriers of heritable lethal diseases caused by recessive genes decreased, because of an intensive selection programme, eliminating mutated gene carriers, and avoiding the use of heterozygous bulls for the diseases BLAD, DUMPS, and CVM.

**References**

6. Schütz E, Scharfenstein M, Brenig B. Implication of complex vertebral malformation and


K. Morkūnienė, R. Bižienė, N. Pečiulaitienė, R. Ugenskienė

Povzetek: Namen raziskave je bil oceniti razširjenost najpogostejših smrtnih bolezni v populaciji litvanskega holštajnskega goveda. V študijo je bilo vključenih dvesto nesorodnih (na podlagi dokumentacije o poreklu) krav in telic. Izolacijo DNK iz krvnih levkocitov smo izvedli z metodo izolacije s soljo in kloroformom. V vzorcih DNK krav in telic smo preverili prisotnost mutacij v treh genih, ki povzročajo naslednje bolezni: motnjo prilepljanja govejih levkocitov (BLAD), pomanjkanje encima uridin monofosfat sintaze (DUMPS) ter kompleksna malformacije vretenc (CVM). Metoda PCR-RFLP je bila uporabljena za določanje polimorfizma gena CD18, ki je odgovoren za razvoj dedne bolezni BLAD. Recesivni alel s točkovno mutacijo A→G (383), ki povzroča bolezen BLAD, je bil ugotovljen v populaciji litvanskega goveda s frekvenco pojavljanja 0,0025. Bolezen CVM je povzročena z drugačno-pomensko mutacijo v genu SLC35A3. Prisotnost mutacije v tem genu smo izvedli z metodo sekvenciranja. Recesivni alel s točkovno mutacijo G→T (538), ki povzroča CVM, je bil ugotovljen v populaciji litvanskega goveda s frekvenco pojavljanja 0,005. Testna metoda PCR-RFLP je bila uporabljena za določanje polimorfizma gena UMPH, ki je odgovoren za razvoj dedne bolezni DUMPS. Recesivni alel s točkovno mutacijo C→T (1213), ki povzroča DUMPS, ni bil najden v litvanski populaciji holštajnskega goveda. Ker so se v preteklem desetletju izvajali intenzivni selekcijski programi, se je število prenašalcev dednih smrtnih bolezni znatno zmanjšalo.

Ključne besede: govedo; moteno prilepljanje govej levkocitov; pomanjkanje encima uridin monofosfat sintaze; kompleksna malformacije vretenc; BLAD; DUMPS; CVM