Role of BoLA-DRB3 genetic diversity against resistance to mastitis in cattle: Review

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The major histocompatibility complex (MHC) is an organized cluster of tightly linked genes, present in all vertebrates, playing an important role in the immune system, except the jawless fish [1]. MHC was first identified during tissue transplantation studies in mice [2] and was first known for its role in histocompatibility. Consequently, the role of MHC was discovered in immune regulation [3] and several other functions [4,5]. The important function of the MHC is to code for specialized antigen-presenting receptor glycoproteins, also called as MHC molecules. The products of these genes are involved in the induction and regulation of immune response. These molecules bind processed peptide antigens and present them to T-lymphocytes, thereby triggering immune response.

The high levels of polymorphism observed in MHC region has been associated with the effectiveness of its immunological non-self or self-recognition function [6]. Cattle MHC molecules are, known as the bovine leukocyte antigen (BoLA) complex. It is located on chromosome 23 and spans approximately 2.5Mb of the cattle genome [7]. It has been estimated that the mammalian MHC contains over 200 genes. The genes are organized into three distinct classes (class I, II, and III). Each of these classes is divided into regions and sub-regions, containing pseudogenes. The BoLA complex is found in two separate regions of the chromosome rather than a single cluster of genes as seen in most mammals. The larger gene cluster is located at BTA23 band 22 and apparently contains all of the bovine class I and class II genes encoding both subunits of the classical class II proteins DQ and DR. The remaining BoLA class II loci (DIB, DNA, DOB, DYA, DYB, TCP1, LMP2, LMP7, and TAP2) are located in a cluster near the centromere at BTA23 band 12-13 [7]. A major rearrangement within the class II region has led to the division of the MHC into two distinct sub-regions, such as class IIa and class IIb, on chromosome 23. The class IIa sub-region contains the functionally expressed DR and DQ genes.

Among the three DRB genes, DRB3 is believed to be functionally important [8]. The DRB3 gene is the most polymorphic class II locus in cattle and influences both the magnitude and
epitope specificity of antigen-specific T-cell responses to infectious diseases. BoLA-DRB3.2, which is the second exon of the third DRB bovine gene, is responsible for the β1 domain of the only widely expressed DRB gene in cattle. There is an increasing efforts to characterize and document the bovine MHC allele frequencies by breed and location, because of the role of BoLA-DRB3 alleles in resistance and/or susceptibility to infectious diseases and immune response [9,10].

Structure and function of BoLA-DRB3 gene

Molecules of Class II are heterodimer glycoproteins consisting of two polypeptide chains (α and β), both anchored in the membrane. The sub designation α1, α2, etc. refers to separate domains within the MHC gene; each domain is usually encoded by a different exon within the gene, and some genes have further domains that encode leader sequences, transmembrane sequences, etc. These molecules have both extracellular regions as well as a transmembrane sequence and a cytoplasmic tail (Figure 1). The α1 and β1 regions of the chains come together to make a membrane-distal peptide-binding domain, while the α2 and β2 regions, the remaining extracellular parts of the chains, form a membrane-proximal immunoglobulin-like domain. The antigen binding groove, where the antigen or peptide binds, is made up of two α-helixes walls and β-sheet.

**Figure 1:** MHC class II molecule structure on antigen presenting cell membrane.

![MHC class II molecule structure on antigen presenting cell membrane](image1)

**Figure 2:** Schematic presentation of antigen presentation by MHC II molecules.

Under the DR region of the MHC complex DRA gene encodes α-chain, whereas highly polymorphic β chain encoded by DRB genes [11]. Three BoLA (bovine leukocyte antigen) DRB genes (DRB1, 2 and 3) are reported in bovines but only DRB3 is functional. DRB1 is a pseudogene, DRB2 is expressed at lower level and DRB3 gene is highly expressed and polymorphic as well [12]. DRB3 plays a central role in the immune system by presenting peptides derived from extracellular proteins.
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(Figure 2). DRB3-encoded molecules share the DR alpha chain and identical DRB3 amino acid sequences in regions important for T-cell receptor interaction. However, the few differences between DRB3*0101, DRB3*0202, and DRB3*0301 change the nature of the peptide-binding pockets and account for their differences in peptide selection and presentation [13]. Class II molecules are expressed in antigen presenting cells (APC: B lymphocytes, dendritic cells, macrophages). DRB3 being a member of MHC class II genes and being polymorphic, plays significant role in disease resistance and in immune responsiveness variability [14].

**Allelic diversity at BoLA-DRB3 gene locus**

Numerous studies have shown that polymorphism in the genes for BoLA class II molecules determines the specificity of the immune response and plays a significant role in conferring resistance or susceptibility to various diseases. Among BoLA class II genes, BoLA-DRB3 highly polymorphic functional genes, have been found to have a stronger association with resistance/susceptibility to various diseases. Behl et al. [15], have concluded that high polymorphism in DRB3.2 locus could assist in identification of superior haplotypes for disease resistance. Takeshima et al. [16], reported high levels of polymorphism in BoLA-DRB3 and identified 5 novel BoLA-DRB3 alleles in Philippine native cattle. While the study provided evidence for evolutionary relationships between the cattle breeds, based on BoLA-DRB3 alleles, it was concluded that allelic information discovered would be essential to understand the correlation between MHC and diseases in East Asian cattle. However, Das et al. [17], also documented a highly polymorphic BoLA-DRB3 exon2 with significant breed-specific genetic diversities in three *Bos indicus* cattle breeds of South Indian, hence recommended conservation to maintain native cattle genetic diversity. In Latin American Creole cattle breeds, Giovambattista et al. [18], reported high genetic diversity of MHC DRB3 which could be as a result of multiple sources of germplasm and could be maintained by balancing selection. Takeshima et al. [19], investigated bovine MHC class II DRB3 in South American Holstein cattle populations and reported high degree of genetic diversity with a low population genetic structure.

Sun et al. [10], investigated the effect of BoLA-DRB3.2 polymorphisms on lameness of Chinese Holstein cows and concluded that BoLA-DRB3 exon2 might be a candidate gene for lameness susceptibility in Chinese Holstein cows. It is important to note that genotype effect at the BoLA-DRB3 could not explain variation in somatic cell count and milk yield at an extent expected of a genetic marker as reported by Oprzadek et al. [20]. Based on their study the workers concluded that the locus is just an ordinary position of the polygene, when interpreting genetic variance of somatic cell count and milk. More than 100 alleles of DRB3 gene have been identified by DNA sequencing [15], data available in the IPD database (www.ebi.ac.uk/ipd/mhc). The allele frequencies of BoLA class II genes appear to vary between breeds of cattle, such as Jersey, Holstein, Argentine Creole, Japanese Shorthorn, Japanese Black, and Brazilian dairy Gir. Thus, estimating the frequency of BoLA class II alleles can help to identify the influence of selective forces such as symmetric balancing selection and positive selection, and differentiate between various populations of cattle [21,22].

**Association of BoLA-DRB3 with mastitis**

The proteins that are important for the functioning of immune system are encoded by MHC [23] and DRB locus is the most polymorphic among the MHC genes [24]. The DRB3 locus exists in the antigen presenting site and any variability in this region may lead to variability in the immune responsiveness of different individuals to particular pathogens. Due to this reason, the importance of the study of polymorphism of this locus has increased [25].
The MHC has been acknowledged to regulate the progression of many infectious diseases, so it is implied that development of markers for these loci may prove helpful in pinpointing superior haplotypes for disease resistance if the association between the trait and these markers can be established. The upstream regulatory region (URR) of the DRB3 gene lies approximately 200 bp upstream of the transcriptional start site and it has strong promoter/enhancer activity. This URR of the DRB3 gene consists of a series of sequence motifs like W, X, Y, CCAAT and TATA boxes. Since these motifs are highly conserved in all MHC Class II genes, their positions as well as spacings are crucial for precise transcription of BoLA genes [15]. DRB3 exon 2 is highly polymorphic with >100 identified alleles and encode the antigen recognition site of the DR [26]. The ability of DRB3 to exhibit high polymorphism makes it a strong candidate to be used as a marker in molecular genetics and phylogenetic studies [27].

A number of studies have reported the association of one or more of the BoLA-DRB3.2 alleles with susceptibility/resistance to the infectious diseases in cattle. Kulberg et al., [28], studied the association of BoLA-DRB3.2 alleles with clinical mastitis in Norwegian Red cows. Genotyping of bovine leucocyte antigen DRB3.2 (BoLA-DRB3.2) in a total of 523 Norwegian Red (NR) cows from two groups selected for high protein yield and low clinical mastitis, respectively, identified 27 previously reported BoLA-DRB3.2 alleles across the groups. Significant differences in BoLA-DRB3.2 allele frequencies were found between the selection groups. Contradictory results from different studies investigating associations between BoLADR3.2 alleles and mastitis indicate that future studies should focus on associations of mastitis with BoLA haplotypes rather than with single BoLA genes.

Characterization of different allelic variants of the MHC class-II gene of DRB3.2 has been investigated in mastitis and healthy cattle using PCR- RFLP by Nandedkar et al. [29]. They identified specific genotype for mastitis and healthy animals. Their study gives evidences of usefulness of DRB3 locus allelic diversity in marker assisted selection of animals on the basis of mastitis susceptible and resistant genotypes. Association analysis of BoLA-DRB3 alleles with mastitis resistance and susceptibility in Japanese Holstein cows have been carried out by Yoshida et al. [30]. On the basis of somatic cell count they classify the animal into healthy and mastitis affected cow and they reported specific alleles associated with resistance to mastitis. Oprzadek et al. [31], evaluated the suitability of the BoLA-DRB3 gene polymorphism to define the phenotypic value of somatic cell count of 808 Polish Holstein cows. The cows were fathered by 190 sires. Significant relationship between the occurrence of the BoLA-DRB3 gene alleles and somatic cell count were reported. Recently, Suprovych et al., [32], explored the BoLA-DRB3 gene polymorphism in the two cattle breeds having commercial importance: Ukrainian black-pied dairy (UBPD) and the Ukrainian red-pied dairy (URPD) and its association with mastitis. For genotyping, PCR-RFLP technique was applied using Rsal, HaeIII and XhoII restriction enzymes by the workers. They observed that in 276 UBPD cows total 32 alleles were present, out of which six alleles (*03, *08, *10, *22, *24 and *28) were identified having a frequency of more than 5% (total amount of 50.4%). The most frequently occurring allele was BoLA-DRB3.2*24 with a frequency of 19.2% and they reported that four BoLA-DRB3.2 were truly associated with mastitis namely *24 and *26 with susceptibility and *13 and *22 with resistance. Similarly, in 117 URPD cattle, they identified 22 alleles from which the most frequent alleles were *07, *22, *11, *24, *01, *03 and *16 (total frequency 64.5%). Allele BoLA-DRB3.2*07 (present in 25.6% of cows) was the most commonly found. In this population, four alleles truly associated with mastitis were identified. Animals susceptible to mastitis were
having alleles *07 and *08, and resistant animals had alleles *22 and *24.

Karthikeyan et al., [33,34], have investigated the genetic variability of mastitis resistance and concluded that it may be attributed by the polygenic control of the many involved genes. Polygenic nature of this trait initiated the need to explore the candidate genes which are responsible for mastitis resistance which may aid in producing the herd with the animals having genetically mastitis resistance alleles.

Though lot of work has been done to understand the genetic diversity at MHC class II DRB locus in cattle, but complexity of the structure in terms of duplications and high levels of polymorphism poses a challenge and makes it difficult to work on these molecules. At least most clear about DRB3 locus is that the higher allelic diversity in a herd is definitely linked to the fitness of the animals. Intense breeding programs are probably leading to loss of genetic diversity at this locus and thus appearance of mastitis and other infectious diseases is more common now. Therefore, periodic screening of animals for allelic diversity and introduction of animals, bulls in particular having desirable alleles should implemented. For that reason, native cattle breeds of tropical regions which are not under intensive selection program have an upper hand.

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