Function of complement regulatory proteins in immunity of reproduction: a review

J. Valentovičová, M. Simon, J. Antalíková

Institute of Animal Biochemistry and Genetics, Slovak Academy of Sciences, Ivanka pri Dunaji, Slovak Republic

ABSTRACT: Humoral immunity has an important role during the maturation and development of the functional properties of spermatozoa. Spermatozoa may be exposed to antisperm antibodies in semen and in cervical, ovarian follicular and fallopian fluid. Antisperm antibodies can be complement-fixing or non-fixing and may affect the reproductive functions in a number of ways. Although the antisperm antibody alone can cause sperm agglutination, complement fixation is required for their immobilization. Therefore, the complement activation might be a “keystone” for the better understanding of “sperm humoral immunity” and some types of infertility. Recently, three cell surface molecules (CD molecules – CD46, CD55, CD59) present on many tissues in male and female reproductive tracts and gametes have been identified. These proteins belong to the family of complement regulatory proteins which could regulate the function of a complement system by cleavage of complement cascade in discrete sites of both activation ways (classical and alternative). In this review, the particular mechanisms of activity of complement regulatory proteins are presented as well as their function in a fertilization process and expression in human and animal tissues and organs.

Keywords: complement regulation; cell surface antigens; CD46; CD55; CD59; spermatozoa

Immune surveillance of spermatogenesis

Reproduction is one of the most serious biological problems that markedly influence the efficiency of livestock production. Immunology is a powerful tool for studying both normal fertility and infertility of farm animals.

There are two basic functions of the immune surveillance of man and animals with respect to sperm immunity during the spermatogenesis and fertilization process: support of the elimination of the vast majority of “waste” spermatozoa that fail to fertilize and preservation of the tiny minority of surviving spermatozoa that are able to fertilize the oocytes. The primordial forms of spermatozoa arising in the seminiferous tubules of testes are preserved by the blood-testes barrier which is formed by Sertoli cells ringing the seminiferous tubules (Hunter, 1989). During maturation in the epididymis the spermatozoa acquire their antigenic properties. Simultaneously, the spermatozoa are coated with some proteins of seminal plasma having the immunosuppressive activity toward lymphocytes and macrophages involved in the immune reactions. The seminal plasma proteins accompany the migration of spermatozoa to the female genital tract (Matoušek, 1985).

The humoral immunity has an important role during the maturation and development of the functional properties of spermatozoa. Antibodies that recognize surface antigens of mature spermatozoa are present in sera of both sexes. The spermatozoa may be exposed to antisperm antibodies in semen and in cervical, ovarian follicular and fallopian fluid...
(Haas et al., 1982; D’Cruz et al., 1990). Antisperm antibodies may be complement-fixing or non-fixing and they affect reproductive functions in a number of ways. Although antibodies alone can cause sperm agglutination, complement fixation is required for their immobilization (Mathur et al., 1981). Therefore the complement activation might be a “keystone” for the better understanding of “sperm humoral immunity” and some types of infertility. The mechanism by which complement activation on spermatozoa and seminal plasma is controlled attracts much scientific interest. Recently, several specific complement-regulatory proteins, present also in male and female reproductive tracts and on gametes, have been identified. These proteins belong to the CD molecules family.

**Complement regulatory CD molecules (antigens)**

CD molecules are antigens appearing in the cell membrane in a specific stage of their development. They remain there in a specific development period or they remain there as a characteristic marker till destruction of the cell membrane. For these properties they are also marked as “differentiation antigens”.

To date, more than 200 surface antigens defined on human leucocytes are known and their number is still increasing. There is a large heterogeneity in the structure and functions of CD antigens. They comprise the receptors for antigens, MHC (Major Histocompatibility Complex) glycoproteins, adhesion molecules, receptors for immunoglobulins, receptors for complement, receptors for lymphokines and other growth and differentiation factors, membrane enzymes or transport molecules and other molecules, with well-characterized structure and expression but the function of which has not been defined yet (Hořejší, 1991; Barclay et al., 1997).

Majority of CD molecules are involved in the immune functions of organism. Some of them are also expressed on the sperm membrane (CD44, CD46, CD47, CD52, CD55, CD59 and CD117). Their role in sperm immunity is mostly unknown. It seems, however, that three CD molecules – CD46, CD55 and CD59 – expressed on spermatozoa (Rooney et al., 1992) are involved in the complement activation and consequently, they could participate in the regulation of complement (C) – mediated killing of spermatozoa (Figure 1). Moreover, the complement regulatory proteins (CD46, CD55 and CD59) are expressed on the fallopian tube endometrium, cervical mucosa, oocytes and embryos, and they may thus play an important role in protecting the transversing sperm and implanting blastocyst from complement-mediated damage (Fenichel et al., 1995; Jensen et al., 1995). There is some experimental evidence that CD molecules could protect human trophoblasts from a complement-mediated attack (Cunningham and Trichenor, 1995). Furthermore, CD46 may be involved in sperm binding to the zona pellucida of the female oocyte. Anti-CD46 monoclonal antibody inhibits the human spermatozoa-ovocyte interaction (Taylor et al., 1994).

In the last decade a lot of data concerning the chemical structure of the cell and tissue distribution and the functional properties of complement regulatory proteins have been acquired. These data allow us to construct a possible mechanism of complement regulation in the immune system generally and especially in the sperm immunity.

**CD46 (MCP; membrane cofactor protein)**

CD46 is a transmembrane glycoprotein with the molecular weight of about 50–60 kDa (Van den Berg et al., 1997), consisting of four homologous short consensus repeats: serine/threonine/proline (STP) – enriched region, an area of undefined function, transmembrane hydrophobic domain, cytoplasmic anchor and cytoplasmic tail (Liszewski et al., 1991). It is important to note that due to the alternative “splicing” of STP- and cytoplasmic regions, various isoforms of MCP are expressed on somatic cells but no polymorphism was found on sperm.

In man, the MCP is expressed by acrosome-reacted, but not acrosome-intact spermatozoa (Anderson et al., 1989). Interestingly, the CD46 expressed on sperm is a 42 kDa molecule. It means that this form is by 20 kDa smaller than the isoforms expressed in other tissues and organs (Russel et al., 1992; Cervoni et al., 1993). MCP is not expressed on unfertilized oocytes but it appears at the 6–8 cell stage embryo. This glycoprotein is present on all human peripheral blood cells and platelets, but not on erythrocytes; on fibroblasts, endothelial and epithelial cells, and on tissues of reproductive system, including fallopian tube, uterine endometrium and placenta (Liszewski et al., 1991). It is also present in seminal fluid.

The main function of the glycoprotein membrane is to protect essentially the host cells from
Complement is a part of the mechanism of natural humoral immunity. This system is composed of more than 30 glycoproteins which are present in the blood serum and on the surface of some cells. The primary functions of these proteins are the production of inflammation, opsonisation of foreign materials for phagocytosis and mediation of direct cytotoxicity against various cells and microorganisms. Specific parts of the complement are activated gradually by a cascade mechanism – activation of the first component causes the arising of serine protease (proteolytic enzyme) which causes the cleavage of the subsequent component into two fragments (limited proteolysis). One of these fragments serves as an enzyme cleavage of subsequent component, etc. Some fragments do not serve as enzymes, they act as cofactors or they have bioregulatory functions. This system can be disrupted by the functional activity of CRP (CD46, CD 55, CD59) and further activity of the complement is interrupted. In Figure 1, the cleavage activity of complement regulatory proteins is presented and marked with linear arrows. Whereas CD46 cleaves C4b or C3b component of complement, CD55 affects the complement on the level of C3 or C5-convertases and causes their dissociation. CD59 prevents the assembly of the membrane attack complex and subsequently blocks the cell lysis.

Figure 1. Functional activity of complement-regulatory proteins
clonal antibodies caused a significant decrease in the number of oocytes showing sperm binding and formation of pronuclei (Taylor et al., 1994). It was reported that the blocking of human sperm by CD46 suppressed sperm binding to human and hamster oocytes (Okabe et al., 1990; Taylor and Johnson, 1996). Kitamura et al. (1997) proposed a new category of infertility – a sperm-specific CD46 aberration. The same authors reported that the sperm lost the ability to adhere to oocytes due to the observed defects in the sperm CD46, thereby causing a failure of fertilization.

**CD55 (decay accelerating factor, DAF)**

DAF is a 70 kDa glycolipid- an anchored membrane-bound complement regulation protein. Similarly like all complement regulation proteins DAF is mainly composed of tandemly repeated motif (short consensus repeats) of about 60 to 70 amino acids in length. In humans, multiple isoforms exist including glycosylphosphatidylinositol (GPI)-anchored, transmembrane and secreted forms with variable lengths of the serine/threonine-rich region.

Originally, DAF was described on human erythrocytes but later it was demonstrated on other circulating blood cells including neutrophils, lymphocytes, monocytes, granulocytes and platelets. While MCP is not present on the outer plasma membrane of freshly ejaculated spermatozoa, very low levels of DAF were detected in this site. Both proteins are strongly expressed on the inner acrosomal membrane, which is exposed on the surface of the cell following an acrosome reaction (approx. 7 h after entry to the female reproductive tract) (He et al., 2000).

In oogenesis, DAF is already expressed on the plasma membrane of oocytes at the primordial follicle stage and continues to be expressed throughout follicular development. The DAF isoform on oocytes seems to be very different from that on spermatozoa (He et al., 2000).

Liszewski et al. (1991) reported that MCP and DAF protect autologous cells from a complement-mediated attack by regulating the activation of C3/C5 convertases. DAF increases the dissociation of C3/C5 convertases (Figure 3) whereas MCP facilitates their degradation by proteolysis of C3b and C4b (Lublin and Atkinson, 1989).

**Figure 2. Mechanisms of CD 46 function**

CD 46 binds to C4b (classical complement pathway) or C3b component of C (alternative pathway) and causes cleavage of these proteins and hence incapability of forming C3-convertases and inhibition of further complement activation.
CD59 (protectin)

Similarly like DAF, CD59 is an 18–20 kDa membrane bound glycoprotein linked to the membrane by a GPI-anchor. Unlike CD46 and CD55, CD59 can be transferred between cells via fluid phase vesicles (prostasomes) and non-membranous complexes, which can explain its presence in many body fluids including blood plasma, saliva, amniotic, seminal fluid and urine (Rooney et al., 1993b).

CD59 is expressed on leucocytes, erythrocytes, platelets, a variety of endothelial and epithelial cells, placenta and spermatozoa. The high levels of CD59 observed on all trophoblast subpopulations suggest that its MAC-inhibitory activity is of extreme importance in the maintenance of placental function (Rooney et al., 1993a).

On spermatozoa, CD59 is expressed on the caput on the surface of acrosome-intact spermatozoa (Rooney et al., 1992). Thus the CD59 molecule seems to be capable of protecting intact spermatozoa in absence regulators of C3-convertases (CD46 and CD55) (Figure 4). CD59 is strongly expressed on the outer plasma membrane of spermatozoa and may prevent membrane attack complex (MAC)-mediated damage to uncapacitated spermatozoa in the female reproductive tract (Rooney et al., 1993a).

To get together the cleavage activity of the three complement regulatory proteins in the inactivation of the complement, the whole procedure of complement regulation in both classic and alternative way could be constructed. In Figure 1 the cleavage sites of complement regulation proteins are demonstrated.

Complement regulatory proteins of farm animals

A majority of studies concerning the complement regulatory proteins was performed in man. Due to the intensive study, man is the single species where CD46, CD55 and CD59 were unequivocally identified. The question remains open whether the complement regulation proteins and a similar mechanism of complement activation exist in other species such as farm animals.

There are some evidences for the existence of CD46 in primates (Nickells and Atkinson, 1990). In the rabbit, tentative identification of CD46 on platelets was made based on its binding to affinity columns of rabbit C3 (Manthei et al., 1988). Analogous molecules to human MCP with preferential expression in testes and testicular germ cells were identified in guinea-pig (Hosokawa et al., 1997) and mice (Tsujimura et al., 1998). Moreover, the guinea-pig epididymal spermatozoa carry CD55 on their surface (He et al., 2000). More detailed studies were performed in pig with a panel of monoclonal antibodies raised against pig lymphocytes. N-terminal amino acid analysis through the first 28 residues showed a 43% homology with the human complement regulatory molecule membrane cofactor protein. The purified protein showed cofactor activity for factor I-mediated cleavage of human and pig C3b, confirming its identity as the pig analogue of human MCP (Van den Berg et al., 1997). Moreover, the cofactor activity of pig MCP toward the human C3b is particularly important in the context of recent strategies for xenotransplantation. Transgenic pig expressing human complement inhibitors on their tissues are bred as organ donors in the expectation that their organs will be less susceptible to hyperacute rejection in the human recipient. MCP of pig is abundantly expressed throughout all tissues examined with particularly strong staining on the vascular endothelium. Unlike human analogues, CD 46 in the pig and orangutan MCP is abundantly expressed on erythrocytes (Van den Berg et al., 1997). Connective tissue elements within liver and testes are also strongly stained by anti-pig MCP antibodies (Peré de la Lastra et al., 1999). Recently, the cattle CD46 molecule has also

![Figure 4. Mechanism of CD 59 function](image)

CD 59 blocks the assembly of MAC. It eliminates the binding of C 9 to C5b-8 complex and prevents the formation of the membrane attack complex that would mediate the cell lysis.
been identified. It was shown that MCP could serve as a cellular receptor for bovine viral diarrhoea virus (Maurer et al., 2004). However, no information is available about the expression of CD46 on mature and capacitated sperm or its presence in the liquids of male and female reproductive tracts. More complex study of the structure and function of the complement regulation proteins of farm animals could help us to understand better the immune protection and/or selection of spermatozoa as well as the interpretation of some types of infertility.

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Corresponding Author

Ing. Jana Valentovičová, Institute of Animal Biochemistry and Genetics, Slovak Academy of Sciences, Moyzesova 61, 900 28 Ivanka pri Dunaji, Slovak Republic
Tel. +421 245 943 151, e-mail: jana.valentovicova@savba.sk