Evaluation of ZP2 domains of functional importance with antisera against synthetic ZP2 peptides

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The mouse zona pellucida protein ZP2 plays an important role in the process of fertilization by mediating secondary sperm binding to mammalian oocytes. ZP2 primary structures are highly conserved as revealed by cDNA cloning. The aim of the study was to identify ZP2 domains of functional relevance. Antisera were raised against synthetic peptides that are either conserved in the structure of ZP2 from different mammalian species (AS ZP2–20) or present in the human ZP2 but not in the mouse ZP2 amino acid sequence (AS ZP2–26). Antibody binding to zona pellucida proteins was assessed by assaying the antisera with human hemizonae. Using human zonae pellucidae, we demonstrated that anti-ZP2 common antibodies and anti-ZP2 human peptide antibodies react with human zona pellucida antigens. For the first time, ZP2 domains of functional relevance for human sperm–oocyte interaction could be identified applying the competitive hemizona assay. Antiserum AS ZP2–20 significantly inhibited binding of spermatozoa to test hemizonae, whereas treatment of hemizonae with AS ZP2–26 did not influence sperm–oocyte interaction. These results show that antibodies against synthetic ZP2 peptides react with ZP2 protein and that AS ZP2–20 identifies a linear ZP2 epitope that is of possible functional importance for sperm–oocyte interaction.

Introduction

Gamete recognition and specific sperm binding to the oocyte occur through complementary molecules and are obligatory processes for successful fertilization (Yanagimachi, 1994). The mammalian oocyte is protected by the zona pellucida, an acellular coat surrounding the plasma membrane of the oocyte. The zona pellucida consists of different glycoproteins, depending on the species. One or more of the zona pellucida proteins are responsible for sperm binding and the subsequent induction of the acrosome reaction (Wassarman, 1990). Polypeptides similar to mouse ZP1 (ZPB), ZP2 (ZPA) and ZP3 (ZPC) proteins that are present as polypeptides with different apparent molecular masses on SDS-PAGE have been defined in various mammals (Liang and Dean, 1993; Harris et al., 1994). The human zona pellucida also consists of three glycoproteins, named ZP1 (90–110 kDa), ZP2 (64–78 kDa) and ZP3 (57–73 kDa) (Shabanowitz and O’Rand and, 1988).

As has been shown in the mouse, ZP1 is the homodimeric filament crosslinker held together by intermolecular disulphides (Greve and Wassarman, 1985). The murine ZP3 protein is responsible for species-specific binding of spermatozoa to the oocyte and for the induction of the acrosome reaction (Vazquez et al., 1989).

After ZP3-mediated acrosomal exocytosis, the spermatozoon is postulated to bind to ZP2 (Bleil et al., 1988). This interaction is required for the second step of sperm–zona pellucida interaction by maintaining the association of the spermatozoon with the zona as it progresses through the oocyte glyocalyx. Mouse ZP2 is a sulphated zona pellucida glycoprotein with an apparent molecular mass of 120 kDa. ZP2 cDNA clones and genes have been cloned from various mammals, including humans, mice, cats, dogs, pigs and rabbits (Harris et al., 1994). After fertilization, the ZP2 protein is modified in the ‘hardening reaction’ to prevent polyspermy. This modification is caused by cortical granule exocytosis.

The most probable complementary sperm protein(s) for ZP2 mediated secondary binding is proacrosin/acrosin (Töpfer-Petersen and Henschen, 1987; Lo-Leggio et al., 1994). These enzymes are believed to be located at the inner acrosomal membrane, which displays sites that recognize the zona pellucida (Yanagimachi, 1981) and bind to ZP2 (Bleil et al., 1988; Mortillo and Wassarman, 1991). Other studies demonstrated that a fucoidan-binding protein is located on the inner acrosomal membrane and either on or in the equatorial segment of acrosome-reacted guinea-pig spermatozoa (Huang and Yanagimachi, 1984). In boar spermatozoa the fucoidan-binding compound was identified as acrosin (Töpfer-Petersen and Henschen, 1987). Tesarik et al. (1990) reported that in acrosome-reacted human spermatozoa, acrosin is found

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primarily in the equatorial segment but also in the inner acrosomal membrane. However, another polypeptide, PH-20, a 64 kDa protein with hyaluronidase activity, may be a primary or secondary zona receptor for guinea-pig and cynomologous macaque spermatozoa (Overstreet et al., 1995).

The different views and uncertainties about the chemical nature and site of the sperm counterpart(s) for the secondary ZP2 receptor make it difficult to understand the molecular mechanisms of mammalian sperm–oocyte interaction in detail. We have used an immunological approach to identify ZP2 epitopes that exhibit physiological relevance in sperm binding. One or more of those ZP2 domains might represent the counterparts for proteins (for example acrosin) localized in the sperm head. In the present study we demonstrate that antibodies against synthetic ZP2 peptides are useful tools for the assessment of defined linear ZP2 epitopes that reveal physiological function in the sperm–zona pellucida interaction process.

Materials and Methods

Animals

Female New Zealand white rabbits (approximately 4 kg) were obtained from Charles River Wiga (Sulzfeld).

Ethics of experimentation

Human oocytes were obtained from patients participating in the in vitro fertilization–embryo transfer programme at the University of Bonn (Bonn, Germany). The specimens represented cells that were not included in the programme and which would normally have been discarded. The investigations have been approved by the local ethical committee. Immunization of rabbits and generation of antisera (Hirsch et al. 1994) were approved by the German authorities and performed in accordance with the German legal requirements.

Materials

Keyhole limpet haemocyanin (KLH), peroxidase-conjugated anti-rabbit IgG from goat, o-phenylenediamine dihydrochloride (OPD), 3,3’-diaminobenzidine (DAB), H₂O₂, Tween 20, 0.1% (v/v) polyvinylpyrrolidone (PVP), human serum albumin (HSA) and bovine serum albumin (BSA), Hepes buffer, Ham’s F10 and haematoxylin were from Sigma (Deisenhofen). MgCl₂ was purchased from Mallinkrodt Chemical works (St Louis, MO). Anti-rabbit IgG antibodies and peroxidase-antiperoxidase complex were purchased from DAKO (Hamburg). Micro-well immunomodules (Polyisorp F 16) were from Nunc (Kamstrup). Dulbecco’s phosphate-buffered saline (PBS) was from Seromed (Berlin).

Methods

Preparation of anti-ZP2 peptide antisera. The sequences of the synthetic peptides (Fig. 1) were deduced from cDNA clones coding for mammalian ZP2 (EMBL ID: HSZP2GF; AC number: M90366). The antigens represented ZP2 domains (Liang and Dean, 1993) that were homologous in human and mouse ZP2 (amino acid residues 541–555, ZP2–20 peptide) or present in the human but not in the mouse ZP2 amino acid sequence (amino acid residues 505–517, ZP2–26 peptide). Selection and synthesis of ZP2 peptides as well as crosslinking of peptides to keyhole limpet haemocyanin (KLH) were performed as described by Hirsch et al. (1997).

Enzyme-linked immunosorbant assay (ELISA) with anti-ZP2 peptide antisera. ELISA was performed with horseradish peroxidase-labelled anti-rabbit antibodies, H₂O₂ as substrate and o-phenylenediamine dihydrochloride as colour reagent. Micro-well immunomodules were coated with synthetic peptide (1 mg in 100 ml of distilled water). Coated wells were air-dried at 37°C for 12 h. Subsequently, wells were washed with PBS supplemented with 0.05% (v/v) Tween 20. The coated wells were filled with antisera at appropriate dilutions in PBS (pH 7.4) supplemented with 1% (w/v) BSA. After 1 h of incubation with the antisera, wells were washed with PBS (pH 7.4) supplemented with 0.05% (v/v) Tween 20. Subsequently, the wells were filled with peroxidase-conjugated anti-rabbit IgG (diluted 1:6000). After incubation for 45 min, the wells were washed as described above. Finally, the wells were filled with 0.05 mol phosphate–citrate buffer 1 (pH 5.0) supplemented with 0.4 mg OPD ml⁻¹ and 0.001% (v/v) H₂O₂. Colour development was complete after 20 min. The absorbance was read at 492 nm using a Titertek Multiscan® ELISA reader (Titertek-Dynatech, Denkendorf).

Competitive hemizona assay. The competitive hemizona assay (HZA) was carried out following the HZA described by Burkman et al. (1988) and used in previous studies (Oehninger et al., 1996). Five independent competitive hemizona assays were carried out using a total of 11 oocytes for antisperm AS ZP2–20 and 13 oocytes for AS ZP2–26. Human oocytes were collected from follicular aspirates. Adherent cumulus cells were removed mechanically using micropipettes, and oocytes were subsequently washed thoroughly. For storage, oocytes were placed in small plastic vials containing 0.5 ml of 1.5 mol MgCl₂, 1 (Mallinkrodt Chemical works, St Louis, MO), 0.1% (v/v) PVP, and 40 mmol Hepes buffer 1. Denuded salt stored human oocytes were equally microinjected using an Axiovert 100 microscope equipped with a micromanipulator (Zeiss, Frankfurt). Hemizonae were separated from oocyte particles by micropipetting. Matching hemizonae were then incubated with test sera (anti-ZP2 antisera) or pre-immune sera (diluted 1:30) for 2 h at 37°C in 5% CO₂ in air.

Human spermatozoa were obtained from fertile men; the motile sperm fractions were obtained using a swim-up technique. Spermatozoa were capacitated in the presence of 0.3% HSA (w/v) for 1 h in an incubator at 37°C in 5% CO₂ in air. Thoroughly washed hemizonae were placed in a sperm suspension containing 0.5 × 10⁶ motile spermatozoa ml⁻¹ for 4 h. After sperm–hemizona co-incubation, each hemizona was rinsed to remove loosely attached spermatozoa. Subsequently, the number of spermatozoa tightly bound to
Use of ZP2 antisera to identify functional ZP2 domains

MACRQRGGWSPSGWFNAQGVESTYSISLFFALVTSGNSIDVSQLVNP
AFPGEVTCDEEITEVEFSSSPGKKWHASVVDPLGDMPCNYLDEPE
KLTLRATYDNCRRVHGGHQMTIRVMNNSAALRHGAVMYQFFCA
MQVEETQGLASTICQKDFMSFLPVRFSLADSKGTQVMGWSIE
VDGARAKTLTLEAMKEGSLLLIDNHRMTFHPNATGTVTHYVQG
NSHLMVSLKLTISFPQQKVIFSSQACAIPDPVTCNATHMTLITEFPG
KLKSVSFENQIDVSLHGDIGLEATNGMLHFSKTLKTLSEKCL
LHQPFLALKLTLFPLRPETVSQITYEPCLESQPSVVTGELCQPQDFM
DVEVYSDQTQAPDLGTLRVRGNSQCPVPFEAOSQGLVRFHIPLNGCG
TRYKFEFFDKVYENEHIALWDTFPPSFISRSEFRMTVKSYSRNDM

AS ZP2-26
PLLINESLTPPSVASKLGPFTLILQSYDPNSYQQPYGEYFVLPVRFL
AS ZP2-20
QPIYMMEVRVLNRDDBNKLYLDDCVWATSNDMDPSFPQWNVVVDGC
AYGLDNQYTTFHPGVSSVTYDPHYQRDFNKAFVSEAHVLSSLVY
FHCALSICNRLSPDSLCPSCVTCPVSSHRATGATEAEMTVSLPGPL
LLSDDSSFRVGGSDLKASQSSGKEKSETGEVGRGMADTKGHKT
AGDVGSKAVAAAFAAGVATLFYIYLYEKRTVSNH

Fig. 1. Primary structure of the ZP2 amino acid sequence (Liang and Dean, 1995). The positions of amino acid sequences chosen for generation of anti-ZP2 synthetic peptide antisera AS ZP2-26 (bold and italics) and AS ZP2-20 (bold) are underlined. Amino acid sequences are given in the one-letter code.

the outer surface of each hemizona was counted. Finally, the hemizona index (HIZ) was calculated (HIZ = [number of sperm bound for test hemizona / number of sperm bound for control hemizona] × 100) (Burkman et al., 1988).

Immunohistochemistry with human hemizonae

For evaluation of anti-ZP2 antibody binding to human zona pellucida proteins, immunohistochemical studies with human hemizonae were performed. The detection of hemizona-bound antibodies was essentially as described by Hinsch et al. (1994) and Oehninger et al. (1996). Briefly, human hemizonae were mounted on glass slides and air dried over night. Thereafter, test hemizonae were treated with antisera AS ZP2-20 (diluted 1:30) or AS ZP2-26 (diluted 1:30) for 1 h; the matching control hemizonae were treated with pre-immune serum (diluted 1:30). Specific binding was visualized with anti-rabbit IgG antibodies (diluted 1:100), peroxidase-antiperoxidase complex (diluted 1:100) and DAB as colour reagent. After fixation in graded alcohols, slides were mounted with a coverslip. Immunostaining was evaluated and photographed using a Zeiss Axioskop microscope (Zeiss, Frankfurt).

Statistical analysis

Results are presented as means ± SEM. Statistical analysis of the HZIs was carried out using the one sample t test.

Results

Antisera were generated either against synthetic peptides corresponding to a conserved ZP2 amino acid sequence or against an epitope present in human ZP2 (Fig. 1). Synthetic peptides were selected on the basis of their probability of being antigenic as predicted by analysis of the primary structure (Krchnak et al., 1987). Specificity of antibodies and antibody titres were determined by ELISA. The results show that the anti-ZP2 antibodies reacted with the respective peptide used as antigen. Antiserum AS ZP2-26 detected the ZP2 human synthetic peptide, displaying a typical sigmoidal titration curve (Fig. 2a). Antiserum AS ZP2-20 clearly reacted with the ZP2 common synthetic peptide (Fig. 2b). However, reaction of antiserum AS ZP2-20 with its antigen was considerably lower than the reaction of antiserum AS ZP2-26 with its corresponding synthetic peptide. Antibodies from the respective pre-immune sera exhibited no binding to synthetic ZP2 peptides.

The competitive hemizona assay revealed that anti-ZP2 common antibodies affect human sperm–zona pellucida binding. Test hemizonae treated with AS ZP2-20 revealed an average of about 15 spermatozoa bound to the outer surface, whereas the corresponding control hemizonae yielded a mean value of 28 tightly bound cells (Table 1). The calculated hemizona index (about 52 on average) and the statistical analysis revealed a significant inhibition of sperm binding to the zona pellucida caused by AS ZP2-20 antibodies compared with control hemizonae treated with pre-immune serum (P < 0.0001). In contrast to the results with AS ZP2-20, preincubation of hemizonae with antiserum AS ZP2-26 did
Table 1. Competitive hemizona assay with human gametes and anti-ZP2 peptide antisera

<table>
<thead>
<tr>
<th>Antiserum</th>
<th>Spermatozoa&lt;sup&gt;a&lt;/sup&gt; (test HZ)</th>
<th>Spermatozoa&lt;sup&gt;a&lt;/sup&gt; (control HZ)</th>
<th>HZI</th>
<th>P value</th>
<th>n</th>
</tr>
</thead>
<tbody>
<tr>
<td>AS ZP2-20</td>
<td>14.6 (± 3.0)</td>
<td>27.8 (± 5.5)</td>
<td>51.56</td>
<td>&lt; 0.0001</td>
<td>11</td>
</tr>
<tr>
<td>AS ZP2-26</td>
<td>87.7 (± 16.1)</td>
<td>92.3 (± 17.0)</td>
<td>100.82</td>
<td>&gt; 0.1</td>
<td>13</td>
</tr>
</tbody>
</table>

<sup>a</sup>Code numbers of antisera used in the present study are given.
<sup>b</sup>Values are mean numbers of human spermatozoa bound to outer human hemizona surface ± SEM.

n = Total number of human oocytes (number of hemizonae) used in five independent replicate assays; one test - hemizona was incubated with anti-ZP2 peptide antiserum, the matching control - hemizona was treated with pre-immune serum.

We also investigated whether anti-ZP2 antibodies bind to human hemizonae. A hemizona incubated with AS ZP2-20 exhibited staining throughout the zona pellucida (Fig. 3a). A brown ring was observed at the edge of the bisected zona pellucida; the amount of substrate deposition in the central part of the hemizona was slightly lower. The periphery of the hemizona revealed a ragged staining pattern towards the outer border, whereas the central portion of the hemizona exhibited a more homogeneous deposition of coloured substrate. Staining of the hemizona incubated with AS ZP2-26 appeared as an intense and clustered pattern of substrate deposition close to the edge of the hemizona (Fig. 3c). Hardly any staining was observed when matching hemizonae were treated with corresponding pre-immune sera (Fig. 3b,c).

**Discussion**

In this study, we have investigated whether antibodies against defined ZP2 epitopes bind to ZP protein in human hemizonae and whether the respective ZP domains detected by anti-ZP2 antibodies are of functional importance for sperm–zona pellucida interaction.

Antibodies against defined ZP2 epitopes were generated and characterized based on ZP2 cDNA sequences (Liang and Dean, 1993). The use of synthetic ZP2 peptides as immunogen provides several advantages over other ZP2 antigens. First, synthetic peptides can be synthesized in large quantities and with sufficient homogeneity. Second, no contaminating proteins will interfere with the outcome of the immunization and generate non-specific antibodies. Third, antisera against synthetic peptides can be used for the identification of ZP proteins that are only defined by molecular cloning. Finally, antisera against homologous ZP2 peptide sequences may allow the identification of unknown ZP2 variants, for example alternative splicing products. However, despite the advantages of synthetic peptide antigens, it must be remembered that antisera raised against synthetic ZP2 peptides can react only with the protein backbone of the ZP2 protein. Therefore, post-translational modifications (such as glycosylation) of zona pellucida proteins during zona pellucida assembly might prevent binding of antibodies to their antigen and lead to inconsistent results.

Fig. 2. Specificity of antisera raised against ZP2 peptides as determined by ELISA. The dilution factor of antisera is shown on the abscissa, the absorbance (492 nm) on the ordinate. One microgram of peptide antigen was used per well. (a) ZP2–26 peptide: ▲ AS ZP2–26, ◇ pre-immune serum 26; (b) ZP2–20 peptide: ■ AS ZP2–20, □ pre-immune serum 20.

not influence sperm–zona interaction. An average of 88 bound spermatozoa was counted when hemizonae incubated with AS ZP2-26 were used; the number of spermatozoa attached to control hemizonae was almost identical (about 92 spermatozoa on average). The hemizona index (about 101 on average) shows that anti-ZP2 human peptide antibodies do not influence sperm–zona binding. No significant change in sperm binding was calculated ($P = 0.89$).
An ELISA developed for the detection of antibodies against synthetic ZP2 peptides was used and anti-ZP2 antisera were shown to react strongly with the synthetic ZP2 peptides that were used as antigens. The antisera did not crossreact with other synthetic peptides; thus, the specificity of antibody reaction could be demonstrated. However, it is essential to demonstrate that the antisera also detected ZP2 protein. It has been shown that antisera AS ZP2-20 and AS ZP2-26 both reacted with ZP2 protein in immunoblots and that the antisera also specifically detected zona pellucida protein in sections of the human ovary (Hinsch et al., in press).

If antibodies are postulated to interfere with sperm–zona pellucida binding in vitro, evidence must be given that the antisera also detect ZP2 protein that is not chemically modified through western blot procedures or immunohistochemical treatment. In the study reported here, immunohistochemical studies with intact human zonae pellucidae were performed which showed that hemizonae that were incubated with AS ZP2-20 and AS ZP2-26 antibodies displayed intense immune peroxidase reactions, indicating that the antisera recognize intact zona pellucida protein. Although antiserum AS ZP2-26 bound strongly to
rehydrated salt stored hemizones (most probably interacting with a defined linear ZP2 domain), this antiserum did not block sperm–zona pellucida interaction.

The human competitive hemizona assay was used to demonstrate that highly specific anti-ZP2 peptide antibodies and human gametes can be used for the study of sperm–zona pellucida interaction. Anti-ZP2 common antibodies significantly inhibited sperm–zona binding, whereas the antiserum against the ZP2-human synthetic peptide did not interfere. The hemizona index revealed that AS ZP2-20 only decreased sperm binding to the hemizones by about 50%. This result might indicate that the concentration or the affinity of antibodies was not sufficient for a total block of sperm binding. However, higher concentrations of antiserum and pre-immune serum resulted in non-specific interaction of pre-immune antibodies with the zona pellucida and thus did not result in a lower hemizona index (not shown). A possible explanation for this finding is that more than one ZP2 domain is responsible for secondary binding.

It is possible that antisera against synthetic ZP2 peptides allow the characterization of defined linear epitopes of discrete ZP2 protein antigens. They can be used as tools for the determination of possible epitopes of possible biological relevance, such as domains for sperm–zona binding. However, it has to be noted that the epitope detected by AS ZP2-20 might not be identical to the domain of physiological importance. Steric hindrance of antibodies could influence a different ZP2 domain which is closely located to the AS ZP2-20 epitope. Two reasons make this assumption less probable. First, AS ZP2-26 detects an epitope close to the epitope detected by AS ZP2-20 but does not interfere in sperm–oocyte binding. Second, in the bovine competitive hemizona assay, F(ab), fragments derived from affinity purified AS ZP2-20 IgG block sperm binding to the same extent as whole IgG (E. Hinsch, unpublished).

The AS ZP2-20 epitope is also present in the mouse and pig amino acid sequence and differs in one amino acid from cat, dog, and rabbit ZP2. Inhibition of sperm–zona pellucida binding indicates that antibodies against this homologous ZP2 sequence block a species-independent sperm binding site. Koyama et al. (1991) generated monoclonal antibodies (MAB-5H4) against porcine zona pellucida protein. In vitro assays revealed that MAB-5H4 antibodies inhibited binding of human spermatozoa to the homologous zona pellucida. Epitope mapping of MAB-5H4 showed that the antibodies detect an amino acid sequence close to the N-terminal of the porcine zona pellucida glycoprotein pZP4; in contrast, antisera AS ZP2-20 and AS ZP2-26 detect epitopes near the C-terminal region. The pZP4 domain recognized by MAB-5H4 displays sequence identity to human ZP2 (8 out of 10 amino acids) but differs from mouse ZP2 in 4 out of 10 amino acids. Antibodies directed against a synthetic pZP4 peptide that corresponds to the MAB-5H4 epitope recognized zona pellucida from pigs, humans and rabbits but not from mice (Hasegawa et al., 1995). The antisera inhibited porcine fertilization in vitro but had no effect on sperm–zona binding immediately after insemination. These results and our data suggest that antibodies against two different ZP2 epitopes inhibit secondary binding of spermatozoa.

In conclusion, the conserved peptide sequence that was evaluated in this study may reflect a ZP2 domain that is important for secondary sperm binding. The results contribute to the understanding of the important events that occur during sperm–zona pellucida interaction. Anti-ZP2 peptide antibodies may be useful for the investigation of the physiology of sperm–oocyte interaction. The evaluation of a physiological role of ZP2 domains detected by antisera used in this study in species other than humans as well as the properties of the respective synthetic peptides are under investigation.

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