Fate of lactadherin P47 during post-testicular maturation and capacitation of boar spermatozoa

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Polyclonal avian antibody was used partially to characterize the pig sperm lactadherin P47. P47 is a mosaic protein, composed of two epidermal growth factor (EGF)-like domains and two C1/C2 domains. P47 is homologous to the bovine mammary gland protein MGP 53/57 and mouse milk fat globule protein. Expression of P47 along the male genital tract and its localization on spermatozoa during post-testicular maturation and capacitation were studied. P47 was detected in the testis and in all parts of the epididymis by immunohistochemistry and by western blots of tissue extracts. By indirect immunocytochemistry, P47 was localized at the apical ridge of the sperm head in testicular, epididymal and ejaculated spermatozoa. The fluorescence intensity progressed during sperm transit from caput to cauda epididymis, probably caused by the ongoing expression and subsequent accumulation of P47 on the sperm surface. During the time course of capacitation, P47 appears to be unmasked by the release of coating proteins and appears to migrate from the apical ridge onto the entire acrosomal region, showing an intensive fluorescence pattern after 3 h capacitation in vitro. The kinetics of signal changes during in vitro capacitation were different in epididymal and ejaculated spermatozoa, indicating accelerated capacitational plasma membrane destabilization in epididymal spermatozoa.

Introduction

Surface architecture evolves continuously during the lifetime of spermatozoa. Mammalian spermatozoa undergo a series of ultrastructural and biochemical changes when passing through the male and female genital tracts. Initial changes, collectively called ‘epididymal sperm maturation’, occur in spermatozoa during their passage through the epididymis and are characterized by extensive remodelling of the sperm surface. This process is accompanied by the association and integration of exogenous lipids and proteins, removal of associated proteins as well as alteration and migration of integral plasma membrane components (for review, see Cooper, 1992; Yanagimachi, 1994; Kirchhoff and Hale, 1996). At ejaculation caused by contact with secretions of the accessory sex glands, the sperm surface becomes complemented by proteins of seminal plasma (Yanagimachi, 1994; Töpfer-Petersen et al., 1998). This surface coat protects the sperm cell membrane from rapid destabilization, which initiates the series of functional changes leading to the acquisition of fertilizing ability, a process known as ‘capacitation’. In the female reproductive tract of a number of mammalian species, the removal of decapacitation factors and the priming of sperm fertilizing ability takes place in the caudal region of the isthmus where spermatozoa are held back due to their binding to the epithelium. At about the time of ovulation, by initiating the capacitation sequence, spermatozoa detach from the epithelium and swim to the site of fertilization, thereby continuing the capacitation process (Hunter, 1996, 2001; Suarez, 1999; Töpfer-Petersen, 1999; Töpfer-Petersen et al., 2002). Part of the capacitation process is an ongoing remodelling of the sperm surface. The dynamics of protein migration from one surface subdomain to another or within a distinct domain during capacitation have been demonstrated for several proteins (Töpfer-Petersen et al., 1990a,b; Blobel et al., 1992). For the integral protein P86/5, capacitating conditions lead to protein aggregation and re-distribution in the peri-acrosomal region (Töpfer-Petersen et al., 1990a,b). The re-distribution of proteins is a general phenomenon, leading to building of protein-free domains which appear to be involved in fusion between plasma membrane and outer acrosomal membrane (Aguas and Pinto de Silva, 1989; Töpfer-Petersen et al., 1990a,b; Benhoff, 1993). After binding to the zona pellucida via a multimeric ligand–receptor system (Shur, 1998) and penetration of the spermatozoa through the extracellular matrix as the result of the acrosome reaction, further ligand–receptor interactions...
between the spermatozoa and the egg cell membranes are required for fusion of both gametes. Along the route of spermatozoa to fertilization, the surface coat and externally oriented integral plasma membrane components confer on spermatozoa the ability to participate in this interaction sequence. Therefore, the identification and functional characterization of sperm-surface proteins are relevant for understanding the molecular basis of fertilization.

Several sperm-surface associated proteins that appear to participate in different steps of fertilization have been identified in a number of mammalian species (Töpfer-Petersen et al., 1995; Snell and White 1996; Töpfer-Petersen 1999). Ensslin et al. (1998) isolated a novel peripherally associated 47 kDa protein of pig spermatozoa, P47, by affinity chromatography from solubilized sperm plasma membrane proteins bound to immobilized zona pellucida glycoproteins. P47 displays a mosaic structure with two N-terminal epidermal growth factor (EGF)-like domains followed by two tandem repeats that are similar to the C1 and C2 domain of the blood clotting factors V and VIII. The second EGF-like domain contains the putative integrin-binding sequence.

Materials and Methods

Chemicals and media

Unless otherwise stated, chemicals were obtained from Merck (Darmstadt), Roth (Karlsruhe) and Sigma Chemical Company (Steinheim) and were of suitably high purity. Enzymes were purchased from Roche Molecular Biochemicals, Mannheim (formerly Boehringer Mannheim) and the secondary antibodies Cy3-conjugated anti-chicken IgY from goat and anti-chicken IgY-alkaline phosphatase from goat were obtained from DianoVA (Hamburg).

A complete bicarbonate Tyrode’s balanced salt solution (pH 7.4, 300 mOsm kg⁻¹) consisting of 96 mmol NaClI⁻¹, 3.1 mmol KClI⁻¹, 5 mmol glucoseI⁻¹, 0.4 mmol MgSO₄I⁻¹, 15 mmol NaHCO₃I⁻¹, 2 mmol CaCl₂I⁻¹, 0.3 mmol NaH₂PO₄I⁻¹, 1 mmol sodium pyruvateI⁻¹, 21.6 mmol sodium lactateI⁻¹, 3 mg BSA ml⁻¹ and 20 mmol HepesI⁻¹, pH 7.6 (Tyrode’s medium; Harrison et al., 1993) was used after equilibration at 39°C, 5% CO₂ for 1 h for sperm capacitation. Swim-up buffer consisted of 95 mmol NaClI⁻¹, 5 mmol glucoseI⁻¹, 60 mmol saccharoseI⁻¹, 1 mmol MgCl₂I⁻¹, 1.5 mmol CaCl₂I⁻¹, 1 mmol KH₂PO₄I⁻¹, 5 mmol sodium pyruvateI⁻¹, 0.5% BSA and 20 mmol HepesI⁻¹ (pH 7.4, 290 mOsm kg⁻¹). All media were passed through a 0.2 μm single use filter unit (Minisart Sartorius, Göttingen) after preparation. TBS was composed of 50 mmol Tris–HClI⁻¹, pH 7.4, and 150 mmol NaClI⁻¹.

Isolation of bovine milk protein MGP and preparation of antibody

Bovine milk fat globule membrane protein (MGP) was isolated essentially as described by Aoki et al. (1994). Briefly, fresh bovine milk was skimmed by centrifugation at 4000 g for 15 min. The raw cream was washed three times with water to remove adhering skimmed milk components. The washed cream was frozen at −80°C for 1 h, thawed and stirred for 1 h at 40°C. After centrifugation (5000 g, 20 min at room temperature) the MGP-enriched fraction was recovered in the middle phase of the tube. After dialysis for 24 h at 4°C against distilled water, the protein fraction was subjected to gel filtration on a Superose 12 HR10/30 column (Amersham Pharmacia, Uppsala) in 20 mmol Tris–HClI⁻¹, pH 7.4, containing 150 mmol NaClI⁻¹ and 1% (w/v) CHAPS.
the morphologically abnormal forms were performed sperm concentration, sperm motility and assessment of estimation of sperm quality including evaluation of from the Institute's colony and used within 30 min.

Before immunization, bovine MGP 53/57 (300 μg ml⁻¹ in PBS or 1% (w/v) CHAPS) was treated with N-glycosidase-F (3.5 U), O-glycosidase (1 U) and neuraminidase (2 U) for 4 days at 37°C to remove most of the glycan chains. Deglycosylation was monitored by SDS-PAGE. After digestion, MGP 53/57 was separated from glycans on a NAP-10 column (Amersham Pharmacia) in 1% (w/v) CHAPS or PBS. The polyclonal antibody was generated by immunizing a pathogen-free cockerel with the purified and partially deglycosylated MGP 53/57. The antigen (104 μg: 1.5 ml) was emulsified with Freund's complete adjuvant (Difco, Detroit, MI) and injected into several subcutaneous sites. A second immunization (104 μg: 1.5 ml) was performed 4 weeks later. Blood was collected approximately 14 days after the second immunization. Pre-immune serum was collected about 14 days after the second immunization. Antibody was designated as anti-BT (Bos taurus) MGP 53/57. Specificity and crossreactivity were tested by SDS-PAGE and western blot analysis (see below).

Sperm preparation

Ejaculates were collected from healthy fertile boars from the Institute's colony and used within 30 min. Estimation of sperm quality including evaluation of sperm concentration, sperm motility and assessment of the morphologically abnormal forms were performed as described by Petrunkina et al. (2001), according to the classification of Krause (1966). Sperm acrosomal status was evaluated by phase-contrast microscopy. Motile spermatozoa were selected by the swim-up procedure (for details see Petrunkina et al., 2001). For capacitation, sperm pellets (5 × 10⁷ cells ml⁻¹) were re-suspended with Tyrode's medium and incubated for 3 h at 39°C in 5% CO₂. The capacitation rates of spermatozoa were indirectly tested by inducing the acrosome reaction with lysophosphatidylcholin (LPC). Spermatozoa were incubated in Tyrode's medium for 3 h at 39°C before the development of immunoreaction. Identical MGP proteins were isolated from pig, horse and human milk.

Sperm motility was estimated (cauda and ejaculated spermatozoa) and only spermatozoa meeting normozoospermic criteria were used for the following experiments. Treated and untreated spermatozoa were then prepared for microscopic investigation or were used for extraction as described below.

Preparation of tissue and sperm extracts

Tissue samples from the testis, caput, corpus and cauda epididymides, bulbourethral gland, prostate gland, seminal vesicle, liver and salivary gland were used. The tissue samples were cut with scissors and homogenized in extraction buffer (TBS, pH 7.5; 2 mmol benzanidin l⁻¹; 1 mmol phenylmethylsulphonyl fluoride (PMSF) l⁻¹; and 1 mmol EDTA l⁻¹) with an Ultraturrax (Jahnke and Kunkel, Staufen). After centrifugation for 20 min (13,000 g at 4°C), the supernatants were collected as TBS extracts. The pellets were re-suspended in exchange buffer containing 1% (w/v) CHAPS. After shaking for 2 h at 4°C the samples were centrifuged at 13,000 g for 20 min. The supernatants were termed detergent extracts.

Boar sperm membranes were prepared from washed ejaculated spermatozoa using the nitrogen cavitation or centrifugation method and then solubilized as described by Ensslin et al. (1995).

Spermatozoa from caput, corpus and cauda epididymides and spermatozoa treated for capacitation were extracted in TBS, pH 7.5 (supplemented with 2 mmol benzanidin l⁻¹; 1 mmol PMSF l⁻¹; 10 mmol EDTA l⁻¹; and 1% (w/v) CHAPS), for 2 h at 4°C. Supernatants were collected after centrifugation (13,000 g for 20 min at 4°C) and stored at −20°C until use.

Sodium dodecyl sulphate-gel electrophoresis and western blot analysis

Samples (15 μg total protein per lane) of purified bovine and pig MGP 53/57, tissue and sperm extracts were subjected to sodium dodecyl sulphate (SDS) gel electrophoresis on 12.5% (w/v) acrylamide gels using a discontinuous buffer system according to Laemmli (1970). Proteins were transferred to PVDF membranes (Roche) using a semi-dry system (buffer composition: 39 mmol glycine l⁻¹; 48 mmol Tris base l⁻¹; 0.0375% (w/v) SDS; and 20% (v/v) methanol; 1 mA cm⁻² for 2 h at room temperature). The membranes were incubated overnight at 4°C with 10% (w/v) blocking reagent (Roche) dissolved in TBS. Strips were then sequentially incubated with anti-BT MGP 53/57 (1:4000, 1 h, 25°C) and alkaline phosphatase-conjugated chicken antibody (1:5000, 1 h at 25°C) in TBS containing 1% blocking reagent. Each incubation was followed by three washing steps in TBS containing 0.3% Tween 20 and two additional washing steps in buffer (0.1 mol Tris–HCl⁻¹, pH 9.5; and 0.1 mol NaCl⁻¹) before the development of immunoreaction.
Fig. 1. Specificity and cross-reactivity of antibody BT MGP 53/57 in the male genital tract in pigs. (a) Cow MGP; (b) pig MGP; (c) human MGP; (d) horse MGP; and (e) pig sperm membrane. (f) Inhibition of immunoreactivity of 47 kDa band of sperm membranes by pre-incubation of antibody BT MGP 53/57 with isolated MGP; (g–j) Cow MGP with antibody BT MGP 53/57 in different concentrations: (g) 1 : 1000; (h) 1 : 2000; (i) 1 : 4000; and (j) 1 : 8000.

(30–60 min) with nitro blue tetrazolium–5-bromo-4-chloro-3-indolyl phosphate (NBT/BCIP) used according to the manufacturer's recommendations (Roche). The reaction was stopped by washing with distilled water for at least 5 min and the strips were then air-dried and photographed. The first antibody (anti-BT MGP 53/57) was pre-incubated for 30 min with purified MGP 53/57 (15 μg ml⁻¹ antibody dilution) before use or replaced by the pre-immune serum at the same conditions to control the specificity. The 47 kDa band was identified by N-terminal analysis as described by Ensslin et al. (1995).

Immunohistochemistry

Tissues collected from testis, caput, corpus and cauda epididymides from slaughtered boars as well as liver and salivary gland as control tissues were used for immunohistochemical studies. Tissue samples (approximately 10 mm × 10 mm × 5 mm) were fixed for 24–48 h in 4% formaldehyde in PBS. After fixation, tissues were washed with running water for 6–8 h and 70% ethanol for 24 h. The preparations were dehydrated with increasing ethanol concentrations (70–100%) followed by acetic acid-N-butylester (ENBE; Roth), and embedded in paraffin wax. Sections (about 7 μm) were mounted on poly-lysin coated slides, de-paraffinized and rehydrated in decreasing ethanol concentrations (3 × 5 min in xylol, 5 min in 100% ethanol, 2 min in 96% ethanol and 30 min in 85% ethanol and 3 × 5 min in PBS). The non-specific binding was blocked by incubation with 10% goat serum in PBS (30 min, 37°C), followed by incubation with the primary antibody (anti-BT MGP 53/57, 1 : 400 v/v in PBS, 1 h, 37°C). After 3 × 5 min washing in PBS, the samples were incubated with alkaline phosphatase-conjugated anti-chicken IgY from goat (1 : 50 in PBS; 1 h, 37°C). Samples were then washed in PBS (3 × 5 min) and stained with NBT/BCIP (Roche) according to the manufacturer's recommendation. The reaction was stopped with distilled water, and the samples mounted with Kaiser's glyceringelatine (Merck). As a control, the successive sections were probed with the alkaline phosphatase-conjugated anti-chicken IgY from goat (1 : 50 in PBS; 1 h, 37°C) or with anti-BT MGP 53/57 pre-incubated with MGP 53/57 (15 μg ml⁻¹ antibody dilution).

The preparations were studied under a phase-contrast Axioskop microscope (Zeiss, Oberkochen) at ×200 and ×400 magnification.

Immunofluorescence microscopy

Sperm suspensions (10⁷ cells ml⁻¹) were smeared on slides, air-dried and fixed in 2% parafomaldehyde and 0.2% glutaraldehyde in PBS for 10 min. The reaction was stopped with 0.1% glycine in PBS for 3 min. After 2 × 5 min washing in PBS and 2 × 5 min washing in distilled water, the preparations were blocked with 20% goat serum in PBS in a wet chamber at 4°C overnight. The smears were then sequentially incubated with the primary antibody (anti-BT MGP 53/57, 1 : 500 in PBS containing 1% goat serum, 60 min, 37°C) and Cy3-conjugated anti-chicken IgY from goat diluted to 1 : 400 in PBS containing 1% goat serum. Each incubation was followed by washing (3 × 10 min in PBS) and two additional washing steps in PBS containing 0.1% Triton-X100 before mounting the smears with glycerol–PBS (9 : 1). The primary antibody was replaced by pre-immune serum or by 1% goat serum in PBS as a control.

The fluorescent slides were examined under an Axioskop microscope equipped with epifluorescence optics (Zeiss) using a 565 nm filter (magnification ×1000). At least 100 cells were evaluated.

Alternatively, P47 was immunolocalized in live spermatozoa. Ejaculated sperm suspensions (10⁷ cells ml⁻¹) were incubated with anti-BT MGP 53/57 (1 : 500 in PBS containing 1% goat serum) for 20 min at 37°C. Sperm suspensions were then gently washed and incubated with Cy3-conjugated anti-chicken IgY (1 : 400 in PBS and 1% goat serum) for 15 min at 37°C. As propidium iodide was not suitable for double staining, the samples were stained with Hoechst dye, the percentage of dead cells in digitonin-treated and untreated sperm suspensions from the same ejaculate was determined using propidium iodide stock solution (0.5 mg ml⁻¹) was added to 1 ml sperm suspension and incubated for 5 min. The percentage of stained cells (membrane defect) was then determined by counting.
Lactadherin localization in pig spermatozoa

Fig. 2. Immunohistochemical localization of pig lactadherin P47 with antibody BT MGP 53/57 in tissues of the male genital tract of the pig. (a) Testis; (b) caput; (c) corpus; and (d) cauda epididymides. The reaction in control experiments (antibody replaced by PBS or goat serum, or inhibited by pre-incubation with purified MGP 53/57) was abolished or markedly reduced.

Analysis of data

Data were analysed using Excel software and the statistics software package SAS (Version 7; SAS Institute Inc., Cary, NC). ANOVA was performed to analyse the influence of sperm origin (ejaculated or epididymal) and time of incubation in capacitation medium. Non-linear and linear modelling were performed and the corresponding model functions compared to describe the changes in kinetics of fluorescence signal shift during capacitation treatment. The rates of changes were compared using first derivatives of the model functions. Unless otherwise stated, data are presented as the mean ± SEM. Differences were considered to be significant if the calculated probability that they would occur by chance was less than 5% (P < 0.05).

Results

Expression of P47 in the male genital tract

Antibodies raised in chicken against bovine MGP 53/57 were shown to crossreact with a comparable crossreactivity with MGP proteins isolated from cow, pig, horse and human milk, and sperm membrane protein P47 in western blot analysis (Fig. 1). Positive reactions were observed only in late germ cells in the testis, indicating that P47 is already expressed in spermatids and differentiated spermatozoa (Fig. 2a). In contrast, the epithelial cells and the stereocilia in the caput and corpus epididymides showed strong immunoreaction (Fig. 2b,c). Signal intensity decreased in the cauda epididymides (Fig. 2d) and was absent in the ductus deferens, seminal vesicle and prostate gland (data not shown). Concomitantly, spermatozoa clustered in the lumen showed immunoreaction with a decreasing signal intensity from the caput to cauda epididymides. Liver and salivary gland were completely negative (not shown). Pre-absorbed primary antibody and secondary antibody alone showed essentially no reaction, indicating the specificity of the antibody. In agreement with these findings, immunoreactive bands at 47 kDa were detected in the detergent extracts of the epididymal tissues of each region after SDS-PAGE and western blotting (Fig. 3, lane b,c,d). The signal in the testicular tissue extract was missing or very faint (Fig. 3, lane a), corresponding to the immunohistochemical results that P47 is expressed only in late germ cells. An immunoreactive band of a similar molecular weight to that of P47 was observed in the seminal vesicle and liver. The results were identical for all boars tested (n = 3; Table 1).
Immunolocalization of P47 on epididymal and ejaculated spermatozoa

Indirect immunofluorescence microscopy using the anti-MGP 53/57 antibody was used to assess the topology of P47 on spermatozoa at different stages of maturation and capacitation.

Classification of fluorescence patterns

Immunolocalization on fixed preparations does not determine whether the epitopes are localized on the surface of live spermatozoa or exposed mainly in dead cells. Therefore, the distribution of P47 on live and permeabilized cell populations was tested by fluorescence microscopy. The percentage of cells in live cell populations that showed specific fluorescence signals (Fig. 4a) corresponded to the number of intact live cells determined separately by propidium iodide (77%). After digitonin permeabilization (100% of cells were propidium iodide-positive, as determined separately), no cells with P47-specific fluorescence were found, probably due to the loss of the plasma membrane as a consequence of digitonin treatment. In untreated sperm populations, spermatozoa stained with Hoechst 33258 (that is, spermatozoa with damaged plasma membrane) were P47-negative. In contrast, only unstained live spermatozoa showed the specific P47 fluorescence patterns (Fig. 4b,c), indicating that P47 is associated with the plasma membrane of spermatozoa.

As fluorescence patterns were shown to be identical in live and fixed sperm preparations, further studies were performed after fixation to determine the time course of the stage-dependent changes of the P47 localization.

In spermatozoa, two main fluorescence patterns were observed: (1) the fluorescence signal covering the entire acrosomal region (designated as A, Fig. 5a) and (2) the fluorescence signal at the apical ridge of the sperm head (designated as P, Fig. 5b,c). The fluorescence signals at the apical ridge were classified into three intensity levels: p1 (faint), p2 (moderate) and p3 (intense) to quantify the distribution of patterns and the changes in the distribution of patterns. The fluorescence was specific as shown by replacement of anti-MGP 53/57 antibody with pre-immune preparation and the use of pre-absorbed antibody.

Changes in P47 localization during epididymal maturation

Most spermatozoa (about 80%) leaving the testis showed one of the fluorescence patterns of the apical ridge (p1, p2 or p3). The distribution of intensities shifted to the most strong fluorescence in the apical ridge (p3) as spermatozoa proceeded from caput to cauda epididymides (Fig. 6a). However, P47 remained strictly localized at the apical ridge in more than 80% of the cells. The observation of Ensslin et al. (1998) that P47 appears to cover the entire acrosomal membrane when passing the duct could not be confirmed statistically. The percentage of spermatozoa with acrosomal fluorescence (3.7 versus 2.3% of pattern A) remained unchanged.

Concomitantly, the P47 band at 47 kDa was detected in sperm extracts of each epididymal region by electrophoresis and western blot analysis (Fig. 6b, lanes 1–3).

Changes in P47 localization during in vitro capacitation

The same two patterns were observed in ejaculated spermatozoa, with almost identical quantitative pattern distribution. Under capacitation treatment over a period of 3 h, the percentage of spermatozoa of both functional state (ejaculated and epididymal spermatozoa) with fluorescence in the apical ridge (p1, p2 and p3) decreased over time. The percentage of spermatozoa expressing fluorescence over the entire acrosomal region (pattern A) or without any fluorescence signal (p0) increased. However, there were noticeable differences in the localization and intensity of P47 signals in the apical ridge during capacitation treatment between epididymal and ejaculated spermatozoa (P < 0.08), as well as a significant influence of incubation time on percentages of spermatozoa showing apical or acrosomal patterns (P < 0.05). There was a significant interaction of the origin of spermatozoa and treatment influence (P < 0.05). Therefore, the changes of distribution were analysed for epididymal and ejaculated spermatozoa...
Fig. 4. Localization of pig lactadherin P47 on live spermatozoa. (a) Phase-contrast photograph. (b) Hoechst-stained spermatozoa: top, spermatozoa stained in the post-equatorial segment, corresponding to live, membrane-intact spermatozoa; bottom, intensively stained spermatozoa, corresponding to spermatozoa with defective membrane. (c) Immunocytochemistry with anti-BT MGP 53/57, fluorescence in the entire acrosomal region (pattern A) observed in membrane-intact spermatozoa, no fluorescence (p0) in spermatozoa with defective membrane.

Fig. 5. Immunocytochemical localization of pig lactadherin P47 on spermatozoa. Classification of the fluorescence patterns: (a) pattern A, fluorescence in the entire acrosomal region; (b) pattern P, fluorescence in the apical ridge, intensity p3 (intense); and (c) pattern P, intensities moderate (p2, bottom) and faint (p1, top).

separately (Fig. 7). Within the group of epididymal spermatozoa, there was a significant decrease in spermatozoa with specific fluorescence in the apical ridge (from 92.3 to 51%; P < 0.0002), accompanied by changes in the distribution of intensities p1, p2 and p3: the percentage of spermatozoa showing the strongest intensity decreased significantly, and after 3 h of incubation spermatozoa showing the faint intensity (p1) were missing. Concomitantly, a significant increase in spermatozoa with fluorescence over the entire acrosomal region (pattern A, from 1.3 to 17.7%; P < 0.01) as well as a significant increase in non-stained spermatozoa (p0, from 5.0% after 3 min to 31.3% after 3 h incubation) was observed. These findings were paralleled by the observation that the intensity of the protein band at 47 kDa in western blots of the sperm extracts was markedly reduced after a 3 h treatment when compared with shorter incubation periods (Fig. 6b, lane 7, compared with lanes 5 and 6).

There was a similar decline in the total percentage of ejaculated spermatozoa showing specific fluorescence signal in the apical ridge (P < 0.03), although to a lower degree than in epididymal spermatozoa (to 75.7%). The tendency in re-distribution of fluorescence patterns p1, p2, p3 was similar, but not statistically significant. In addition, the tendency in the increase of spermatozoa showing fluorescence over the entire acrosomal region was not significant.

Data were fitted to two kinetic models to confirm that the changes in P47 fluorescence patterns obey the different temporal dependence in epididymal and ejaculated sperm populations (Fig. 8). The changes in the ejaculated spermatozoa could be described by a linear model (P < 0.05), and the changes in epididymal
The sperm samples from three boars were treated with LPC after incubation in capacitation medium for 3 h to examine whether the increase in fluorescence signal over the entire acrosomal region was characteristic for capacitating spermatozoa. The percentage of spermatozoa with different patterns of P47 fluorescence was determined by simultaneous evaluation of the acrosomal state. After stimulation of the acrosome reaction, the percentage of spermatozoa showing pattern A decreased rapidly (from 12 to 4%). The percentage of spermatozoa with fluorescence in the apical ridge decreased in patterns p3 and p2 (from 28% in each class to 11 and 12%, respectively). The percentage of spermatozoa with pattern p1 increased from 27 to 38%, whereas the intensity of this pattern became merely distinguishable after LPC treatment. The percentage of spermatozoa without fluorescence signal progressed from 6 to 34%. The spermatozoa with fluorescence signal were identified as acrosome-intact by phase-contrast microscopy after LPC treatment.

Discussion

Lactadherin-like proteins have been identified in the male genital tracts of pigs, cattle, mice, humans (Ensslin et al., 1998) and horses (E. Töpfer-Petersen and M. Gentzel, unpublished). The wide distribution of messenger RNA coding for lactadherin P47 (testis, epididymis, mammary gland, uterus, adrenal gland, tonsil, muscle, heart, lymphatic gland, thymus and kidney) indicates that their action is not restricted to the particular tissue (Ensslin et al., 1998). These results are in agreement with those of Andersen et al. (1997) who demonstrated that homologous bovine milk fat globule lactadherins (formerly PAS 6/7 and MGP 53/57) are expressed in a number of tissues essentially similar to those detected by Ensslin et al. (1998).

As the sequence identity is about 80% between cow and pig lactadherins, a polyclonal antibody directed against purified bovine MGP 53/57 showed strong cross-reactivity with the members of the lactadherin family in pigs. Lactadherin P47 was identified by immunological approaches in testis and in all parts of the epididymis in pigs. The epithelial cells stained positive for the protein in the epididymis, whereas in the testis the protein was found to be associated solely with late germ cells, indicating that spermatozoa released into the lumen and transported into the epididymis already carry P47 molecules on the surface. P47 was localized in the plasma membrane over the apical ridge in most spermatozoa of testicular origin. A small portion showed P47 at the entire acrosomal region. These results confirm the findings of Ensslin et al. (1998). In the present study, however, P47 localization during epididymal maturation and in vitro capacitation was studied by quantifying the intensity and distribution patterns.

Interpreting the data (low amount of protein on the testicular sperm surface and in the testicular sperm extracts), P47 appears to be already associated with the peri-acrosomal plasma membrane when leaving the testis, possibly being masked by testicular proteins. It can be hypothesized that the strong interaction of P47 with the sperm membrane is mediated by the C-terminal peptide of the C2-like domain that is homologous to the phosphatidylserine-binding region of factor VIII (Gilbert and Baleja, 1995) and is highly conserved within the lactadherin family (Andersen et al., 2000).
The increase in signal intensity at the apical region of the sperm head during epididymal transit may be caused by the unmasking of the protein already present on testicular spermatozoa, or by interaction of the protein secreted by the epididymis with the spermatozoa (subsequent association of P47 with the sperm surface), possibly as the result of a changing lipid architecture in the apical membrane domain (Töpfer-Petersen et al., 1990a,b; Christova et al., 2002). An opposite dynamic of changes on the sperm surface was observed in both epididymal and ejaculated sperm populations during capacitation. A shift of the P47 signal to the entire acrosomal region occurred and this was accompanied by an increase in a cell portion showing no fluorescence. Whereas P47 loss is related to acrosome reaction occurring at the end of in vitro capacitation, the acrosomal appearance of the protein can be due to the combined effects of unmasking by the release of coating proteins and the migration of P47 molecules from the apical subdomain into the acrosomal domain. The apical head plasma membrane in pigs seems to form a specialized subdomain. Gadella and Harrison (2002) demonstrated in boar spermatozoa the exposure of aminophospholipids and the subsequent migration of cholesterol into the apical plasma membrane as the result of bicarbonate-induced lipid re-organization under conditions that promote capacitation. The migration of the transmembrane proteins P86/5 into the apical subdomain may be closely related to these processes (Aguas and Pinto DaSilva, 1989; Töpfer-Petersen et al., 1990a,b). The abolition of the barrier may also allow the migration of lipid-bound proteins from the apical subdomain into the acrosomal region.

It appears important that the kinetics of signal loss or shift differed for epididymal and ejaculated spermatozoa. The rate of signal changes was lower in ejaculated spermatozoa in the second half of incubation:

\[
\frac{d(Ep)}{dt} = -0.0025t^2 + 0.25t + 89.5; R^2 = 0.93 \\
\frac{d(Ej)}{dt} = -0.12t + 96.7; R^2 = 0.91
\]

Changes in epididymal spermatozoa progress more rapidly than in ejaculated spermatozoa in the second half of incubation:

\[
\frac{d|Ep|}{dt} > \frac{d|Ej|}{dt} \quad \text{when } t > 74 \text{ min.}
\]
spermatozoa (threefold at the beginning of the third hour of incubation: \(\frac{\Delta s}{\Delta t}_{[0.3]} = -0.12\) in ejaculated spermatozoa against \(-0.36\) in epididymal spermatozoa), providing evidence for accelerated capacitation-induced membrane remodelling in epididymal spermatozoa. It is unlikely that these kinetics are related solely to the incompletely matured morphology of epididymal spermatozoa. Indeed, the subpopulation differences in response to bicarbonate, caused by variable efficiencies in epididymal maturation within a sperm population as judged by cell morphology, have been reported by Flesch et al. (2001). Only spermatozoa that have accomplished the process of morphological maturation respond to bicarbonate by positive merocyanine staining, indicating the occurrence of cholesterol efflux from the sperm membrane initiating the capacitation process. Therefore, a lower response to capacitating conditions in epididymal spermatozoa would be expected than in ejaculated spermatozoa if the degree of epididymal maturation plays a crucial role. It appears more likely that in the present study such differences were caused by completion of the surface coat at the time of ejaculation. Epididymal spermatozoa that did not have contact with fluids of accessory glands seemed to be able to undergo capacitation-related loss of signal in the apical ridge at a higher rate, as shown by the observed kinetics. Similarly, more rapid functional destabilization of the plasma membrane of epididymal spermatozoa during long-term incubation under fertilizing conditions was detected by changes in osmotic behaviour of cell volume in an earlier study (Petrunkina and Töpfer-Petersen, 2000). Epididymal spermatozoa demonstrated rapid aberration in response to osmotic conditions and marked deviation from the model of perfect osmometer at approximately the same time as in the present study, that is, during 3 h of incubation.

The function of lactadherin in association with spermatozoa is largely unknown. The possible function of boar membrane P47 as an integrin RGD-dependent ligand as suggested by Ensslin et al. (1998) was previously indirectly supported by studies of Andersen et al. (1997), who showed that interaction between bovine lactadherin and integrin \(\alpha_v\beta_3\) was RGD-dependent. Moreover, these investigators have recently determined that lactadherin can act as a link between two surfaces by binding to integrin receptors through its N-terminal RGD-binding sites in the second EGF-like domain and to phospholipids through its C-terminal C1/C2-like domains (Andersen et al., 2000). Taylor et al. (2000) demonstrated that human lactadherin (formerly BA46) expressed in human milk and breast carcinomas promotes RGD-dependent cell adhesion via integrins. The proposed hypothesis of active participation in gamete interaction which takes place only after completing the acrosome reaction via the integrin binding site can be ruled out as lactadherin is not present on acrosome-reacted spermatozoa. However, the presentation of lactadherin on spermatozoa entering the oviduct indicates the interaction with integrin receptors localized on the oviductal epithelium (Chegini et al., 2001), thereby regulating cross-talk between spermatozoa and the oviductal epithelium. In addition, the results of the present study give an indication that lactadherin is involved in other aspects of sperm physiology such as capacitation and acrosome reaction, although it cannot be ruled out that the re-localization of the protein occurs as a consequence of these processes. The still unknown molecular mechanisms behind the observed functional changes need to be determined. One possible mechanism is an interaction with the EGF receptor (EGFR, found on the human spermatozoa; Damjanov et al., 1993) to which EGF-like domains such as those at the N-terminus of P47 bind specifically. Members of the EGFR family are transmembrane tyrosine kinases, and are involved in intracellular signalling. The signal transduction pathways that induce changes in the migration and unmasking of lactadherin epitopes require further investigation. Further study is also required to determine whether lactadherin epitopes are triggered or integrated in a further signal cascade priming acrosome reaction and preparation of sperm–egg fusion.

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