Spermatogenesis I (635 - 640)

Conclusion: The intra-acrosomal alkalization took place during incubation conducive for sperm capacitation and induced to alter the nature of acrosomal contents to be ready for release at acrosome exocytosis.

635 A 32kDa Protein (IAM 32) is the Major Integral Protein of the Sperm Inner Acrosomal Membrane
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The inner acrosome membrane (IAM) participates in binding to the egg's zona pellucida and may also be involved in subsequent zona-penetration, yet its composition remains unknown. Our objective, therefore, was to obtain information on the integral protein constituents of the IAM. For this purpose we utilized sonicated and isolated bull sperm heads (SSPH) whose plasmalemma and acrosomal contents were missing but whose IAM remained bound to the SDS-insoluble permalecular theca. A Triton X-100 extract of the SSPH fraction revealed a major protein of 32kDa (IAM 32) and a less abundant 38kDa protein (SP 38) which could be eliminated by prior high salt extraction of the SSPH fraction. Anti-serum raised against and specific to bull IAM 32 cross-reacted with co-migrating proteins of rat sperm and immunogold labeled the IAM of spermatozoa and white blood and mature sperm, and of acrosome reacted rodent sperm. The amino terminal sequence of the isolated bull IAM 32 protein had no similarity to proteins in the NCBI protein data bases. In conclusion, we have uncovered the major integral membrane protein of the IAM that most contribute to the spermatozoa IAM matrix along with IAM 32, which is peripherally attached to the IAM after the acrosome reaction. (Supported by the National Science and Engineering Research Council of Canada (NSERC)).

636 Characterization of Rat Sperm Capacitation In Vitro and its Effect on Changes in Lysosome Membrane Markers
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Capacitation of rat sperm is not well-defined in the literature. Here we characterize rat sperm capacitation in vitro through analysis of tyrosine phosphorylation kinetics, cholesterol removal kinetics (with BSA and methyl-β-cyclodextrin (MBCD)) and motility. Using defined capacitation conditions, we also report effects on lipid raft (using GM, as a marker) and protein (CRISP-1) changes in the membrane. Quantification of cholesterol removal in cauda epididymal rat sperm capacitated in vitro shows that the amount of cholesterol extracted from the membrane increases directly with the concentration of MBCD; cholesterol extraction occurs rapidly, reaching peak values within 15 to 30 minutes. Following removal of cholesterol with MBCD or BSA, tyrosine phosphorylation of specific proteins increases, as determined by western blot analysis, with the rate dependent on the concentration and type of cholesterol binding agent. Concurrently, treatment with MBCD or BSA results in the initiation of hyperactive motility. Following capacitation, there is an increase in sperm membrane TaKaRa Stamenius stained with the actin-specific antibody that is a known marker of lipid rafts. Using chola toxin to label GM1, there is a shift from sharp, intense staining in the post-acrosomal sheath and equatorial band region of the head in sperm incubated in non-capacitating conditions to a diffuse, unrestricted staining across the entire head in sperm incubated in capacitating conditions. Western blot analysis and immunocytochemistry show that the amount of CRISP-1 on sperm capacitated with MBCD declines slightly during capacitation, but the majority remains on sperm that does not move within the plane of the microscope. (Supported by USPH grant HD-11962)

637 Peripheral Bound Membrane Proteins Are Involved in the Maintenance of Boar Sperm Viability by Oviductal Apical Plasma Membrane Footprints and in Vitro
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We have previously shown that oviductal apical plasma membrane (APM) preparations enhance boar sperm longevity in vitro. In the current investigation we studied the nature and location of active factors in APM. Oviductal epithelium was scraped, homogenized and purified to prepare APM vesicles. A sample of APM was heat treated to destroy protein activity. A further sample of APM was incubated at 40°C in 1 M NaCl for 30 minutes, followed by separation of peripheral and integral membrane bound proteins by centrifugation at 100,000g for 1 hour. Washed swim in vitro studies in vitro (25 x 10⁶ sperm/ml) diluted with Tyrode's medium (25 µl) were added to 25 ml experimental APM with a final protein concentration of 200 µg/ml and to medium only (control). Treatments included heat-treated vs non-heat-treated APM and peripheral vs integral membrane bound proteins, alongside whole APM. These were incubated at 39°C and 5% CO₂ for 24 hours. Sperm viability was determined with Ethidium Homodimer-1 and SYBR 14. Results were expressed as percentage of the original viability (viability indexSEM). The heat treatment significantly (P<0.05) reduced the viability enhancing ability of APM (59±5, 78±5 and 65±5 for heat-treated, non-heat-treated and control respectively, n=8). There was a significant increase (P<0.05) in the percentage of viable sperm incubated with the peripheral fraction in comparison to all other treatments (79±2, 75±6, 85±2 and 58±5 for peripheral, integral, original APM fractions and control respectively, n=12). We suggest that active factor(s) in the peripheral fraction are involved in the maintenance of boar sperm viability in vitro of protein nature and peripherally bound to the APM of porcine oviductal epithelial cells. This study was funded by the Department for Environment, Food and Rural Affairs, UK.

638 SP 38: The Major Peripheral Protein of the Sperm Inner Acrosomal Membrane
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The consequence of the acrosome reaction is the exposure of the inner acrosomal membrane (IAM) as the leading surface of the sperm head, which then binds secondarily to the egg's zona pellucida, a prerequisite for zona-penetration. However, the IAM proteins involved have not been identified probably because the protein composition of the IAM was unknown. Our objective, therefore, was to obtain direct information on the peripheral protein constituents of the IAM. For this purpose, we devised a fractionation procedure to isolate apical tips of the rat sperm head, which consisted solely of the IAM bound to the SIDS-insoluble perforatorium. High salt incubation of this tip fraction extracted a major protein of 38 kDa coincident with partial removal of an electron dense layer attached to the IAM. A prominent 38kDa was also salt extractable from sonicated and isolated bull sperm heads that had reacted with rat IgG, a finding that is specific to this bull protein cross-reacted with the co-migrating 38kDa protein obtained from the rat tips and immunogold labeled the IAM of these tips, of sonicated and whole bull and mature sperm, and of acrosome reacted murine sperm. Cloning and sequencing of the 38kDa protein revealed its identity as SP 38 (Dev. Biol. 168:575-583, 1995), an acrosome protein with zona/ZP 2 binding capability whose precise location within the acrosome was unknown. In conclusion, the outermost membrane protein contributes to an electron dense layer or matrix that is peripherally attached to the IAM after the acrosome reaction. This suggests SP 38 is a strong candidate for secondary zona-binding. (Supported by NSERC).

639 Do D-3 Phosphonitrosides Signal Actin Polymerization during Ascidian Sperm Activation? 
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Sperm activation in the sea squirt Ascidia cerasoides is characterized by mitochondrial translocation (MUL), an actomyosin-dependent movement known to require elevation of both intracellular pH (pH) and free calcium ion concentration ([Ca²⁺]). Previously, we have shown that myosin activation requires G protein-mediated pathway involving inositol 1,4,5-triphosphate-mediated internal Ca release and a Ca-dependent internalisation of F-actin via the external Ca entry. Here, we explore signaling elements that are involved in triggering actin polymerization. In MTL assays, the actin polymerization inhibitor latrunculin (10µM) completely blocked high pH artificial sea water (ASW) induced sperm activation (positive control), and the actin polymerization inducer jasplakinolide (7.4µM) stimulated sperm activation equal to positive controls, an action blocked by latrunculin. Dual labeling with fluorescently tagged phalloidin and DNAseI revealed that filamentous actin was distributed mostly heavily on the mitochondrion whereas monomeric actin was also found along the length of the tail. Sperm activation appears to increase filamentous actin on the mitochondrion. In MTL assays, the phosphodiesterase 3-kinase (PI3K) inhibitor LY240402 (50µM) blocked sperm activation induced by pH 9.4 ASW but not that induced by the G protein activator mast (7.5µM) or the PKC activator OAG (50µM), agents shown to be part of the myosin activation pathway. Liposomes that incorporated phosphodiesterase 3,4,5-phosphate (PiP3) stimulated levels of sperm activation similar to positive controls. Indirect immunofluorescence using anti-profilin antibodies showed profilin to be present on the mitochondrion, providing a possible connection between PI3K-induced PiP3, production and actin polymerization. (Funded by NSF U1202-105 and the National Science Foundation Grant MH- NIH R25-GM65820 to RAK for LB, DB & AE; NIH R15HD36050 to RAK.)

640 The Dynamics of Inter-Sertoli (SI) Tight Junctions (TJ) Are Regulated by Transforming Growth Factor-β (TGF-β) via the p38 Mitogen-Activated Protein (MAP) Kinase Signaling Pathway
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Earlier studies have shown that the inter-Sertoli TJ dynamics are regulated, at least in part, by TGF-β3, possibly via its effects on oculoductalis-1/120-1 (Liu et al, Endocrinology 142:1865-1877, 2001). We now report that this TGF-β3 regulatory effect on TJ-functionality is mediated via the MEK/Erk/p38 MAP kinase signal
transduction pathway. Using SCs cultured in vitro to allow the assembly of TJs when the TJ-permeability barrier was monitored by the transepithelial electrical resistance (TER) across the SC epithelium, we have examined if the TGF-β3-induced TJ barrier formation is mediated via four upstream cascades, namely MEKKs, Smad2/Smad3, Cdc42/Rac, and Cellular distribution studies using RT-PCR have shown that both SC and germ cells (GC) express almost similar levels of mRNA encoding for MEKK2, Smad2, and small GTPase, respectively, N-Ras, Rac, and N-WASP. Cellular induced increase in MEKK2 expression, but not Smad2, Cdc42/Rac2, or N-Ras, was detected in SC during the assembly of the TJ barrier. The TGF-β3-mediated (3 ng/ml) inhibitory effect on the assembly of TJs could be reversed dose-dependently by 20B2190 at 0.1 nM-1M, a specific p38 MAPK kinase inhibitor. We next investigated the protein expression of p38-MAP kinase (activated phosphorylated form) versus total p38-MAP kinase (nonphosphorylated inactive form) using SC lysates by immunoblotting and specific antibodies against p38- and p38-MAP kinase with a chemiluminescence-based detection system. It was found that the presence of TGF-β3 indeed regulated the production of p38 MAP kinase protein during TJ assembly. In summary: the TGF-β3-mediated effects on the inter-Sertoli TJ dynamics and the blood-testis-barrier functionality are regulated via the p38-MAP kinase pathway. [Supported in part by grants from CONRAD (CICRC06-05-A to CYC), and HKRGC (HKU7245/S00 to WML/CYC)].

641

T3-Regulated Expression of a Novel Attachment Factor, PB-cadherin, may be Critical for Development of Neonatal Testicular Stem Cells
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In the rodent testis, contact-mediated interactions between gonocytes, or neonatal stem cells, and Sertoli cells are critical for development. Previously, we showed that Thyroid Hormone (T3) regulates expression in neonates of at least one Sertoli cell-specific factor, N-WASP. Sequencing and cDNA microarray and detected expression of another factor, short-type PB-cadherin (STPB-C) in neonatal Sertoli cell- gonocyte co-cultures. PB-cadherin is a novel cadherin mainly expressed in primary germinal and blood of adult mice. PB-cadherin is involved in development by regulating Ca2+-dependent cell-cell adhesion. Therefore, our present aims were (1) to explore expression of STPB-C in vivo, and (2) to localize STPB-C mRNA in co-cultures with in situ hybridization, and (3) to determine if expression of STPB-C is regulated by T3. RT-PCR was used to generate cDNA for STPB-C from total RNA isolated from co-cultures, cDNA was cloned into pcDNA3(+) cloning vector, and plasmid DNA was isolated and sequenced to confirm the fidelity of the STPB-C cDNA portion of the plasmid. In subsequent Northern analysis of testicular RNA, expression of STPB-C was strong on day 1, then diminished appreciably by day 3, because barely detectable by day 15, and disappeared in testes of adults. When neonatal co-cultures were treated with T3 (10 mM, 24 hr) or vehicle, STPB-C mRNA was strongly expressed by neonatal stem cells and weakly by Sertoli cells in situ, while Northern analysis indicated that expression of STPB-C was down-regulated by T3 in vitro. Thus, regulation of PB-cadherin by T3 during the early neonatal period may be critical in development of the stem cell population from which alluring germ cells subsequently arise. (Supported by NIH HD-15563).

642

Regulation of Catenins in The Rat Epidermis
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Adhering junctions are essential for the formation and regulation of tight junctions. In the epidermis, tight junctions between adjacent principal cells form the blood-epidermal barrier which creates a specific environment for the lumen of the epidermis necessary for sperm maturation. Adhering junctions in the epidermis are composed of a transmembrane protein, cadherin, and catenins (alpha, beta and pl120). The objective of this study was to determine the effects of testis and testicular androgens on the immunolocalization of cadenins (alpha, beta and pl20) in the epidermis. In intact control adult rat epididymis, each of the three catenins were localized along the lateral plasma membranes of adjacent principal cells as well as between principal and basal and clear and basal cells. Twenty one days following orchidectomy there was a marked increase in the cytosolic staining of both alpha- and beta-catenins, particularly in the corpus and cauda epididymides, suggesting a loss in the integrity of the adhering junctions. Interestingly, immunostaining for pl20 appeared to be unaltered by orchidectomy. In orchidectomized rats that had been given testosterone implants at the time of orchidectomy, the immunolocalization of alpha- and beta-catenins was maintained along the lateral plasma membrane of epidydimal principal cells. These data suggest that androgens can maintain the integrity of adhering junctions in the epidermis and may represent a mechanism by which androgens can regulate tight junctions and the blood-epididymal barrier in adult rats.

643

Regulation of Sertoli(SC)-Germ(GC) Cell Anchoring Junction (AJ) Dynamics by RhoB GTPase
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During spermatogenesis, there are extensive AJ restructuring, however, the mechanistic pathway that regulates AJ dynamics is not known. Rho GTPases have been implicated in the actin organization and cytoskeletal control. For instance, Rho GTPases are known to regulate AJ functionality by redistributing cadherins during AJ assembly. Using RT-PCR, both SC and GC were found to express RhoB. Moreover, the assembly of SC-GC AJs, but not SC TJs, was associated with a transient induction of RhoB expression. These changes were confirmed with a monospecific RhoB antibody for immunoblots using cell lysates and a chemiluminescence-based detection system. Disruption of AJs in SC-GC cocultures by hypertonic treatment also induced a surge in RhoB expression, which became visible within 5 min. Moreover, when SC-GC AJs in vitro was disrupted by treatment of rats with a single dose of 1-2,4-dichlorobenzyl)-indazole-3-carboxyhydrate (DICC, 300 mg/kg b.w. by gavage), a surge in RhoB expression by blot was also detected within 1 hr. This is long before the depletion of germ cells from epithelium become visible, which required ~10 days. When GCs were added to the SC epithelium (GCSC, 1:1), which had been cultured for 5 days at 0.5x10^6 cells/cm2 on Matrigel-coated dishes, and cultured for 2 days to allow the assembly of AJs; inclusion of DICC at 250 mg/ml to the cocultures also induced a surge in RhoB expression ~5 min-1 hr. These results thus illustrate that RhoB is activated during the assembly of AJ, and prior to the actual disruption of AJs. These changes are possibly needed because the RhoB regulates redistribution of AJ-proteins, such as cadherin. In summary, RhoB may be an important intracellular molecule that regulates AJ dynamics in the testis; and ii) the DCIC-induced GC loss from the epithelium is mediated via the Rho GTPase signaling pathway.

644

The dynamics of Sertoli (SC) -germ (GC) anchoring junctions (AJs) are regulated by E-cadherin, beta-catenin, N-cadherin, and SrC catenin. Nikki Pai Yue Lee, Will M Lee, C Yan Cheng. Population Council, 1230 York Avenue, New York, NY 10021, 1Department of Zoology, The University of Hong Kong, Hong Kong

Earlier studies have shown that the assembly of AJs between SC is associated with a transient induction in N-cadherin/beta-catenin expression. Moreover, the junctional dynamics in the tests are regulated by the interplay of phosphatases and kinases that regulates the intracellular phosphoprotein content. To expand these earlier studies, we have assessed the role of E-cadherin (SC), N-cadherin (SC), and SrC (SC) associated signaling molecule) in AJ dynamics. When SC were cultured at 0.5x10^6 cells/cm2 or SC-GC were cocultured (SC-GC ratio at 1:1, SC at 0.5x10^6 cells/cm2) in vitro on Matrigel-coated dishes to allow the assembly of AJs, it was associated with a transient induction in the expression of E-cadherin, N-cadherin and SrC. Similar changes were detected when cell lysates were prepared from these samples for immunoblots using the corresponding antibodies and a chemiluminescence-based detection system. Cellular distribution studies by semi-quantitative RT-PCR revealed that while SC expressed almost twice as much N-cadherin when compared to GC, GC expressed almost 3 times as much E-cadherin as SC, suggesting GC play an important role contributing to the AJ-associated protein pool in the testis. Also, the expression of SC E-cadherin and N-cadherin were stimulated by testosterone and DHT by ~3-10 fold at 10^{-9}M, suggesting androgens may also regulate AJ functionality via their effects on AJ-associated proteins. Work is now in progress to assess whether these changes are changes in the phosphorylation state of these proteins during AJ assembly. In summary, these results demonstrate that the dynamics of AJs are regulated, at least in part, by N-cadherin, E-cadherin, SrC and androgens. [Supported in part by grants from the CONRAD Program (CICRC06-05-A to CYC) and Hong Kong Research Grant Council (HKU7245/S00 to WML/CYC)].

645

The role of inducible nitric oxide synthase (iNOS) and endothelial nitric oxide synthase (eNOS) in the dynamics of tight (TJ) and anchoring junctions (AJ)
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NOS catalyzes the enzymatic oxidation of L-arginine to nitric oxide (NO), which plays a critical role in a variety of bioregulatory functions including junction assembly. Also iNOS and eNOS are present in the tests and are implicated in the regulation of spermatogenesis. However, their roles in junction dynamics in the tests has not been explored. When SC were cultured at 0.5x10^6 cells/cm2 on Matrigel-coated dishes for up to 7 days, there was a transient induction in eNOS expression, but not iNOS, coinciding with the assembly of inter-Sertoli TJ permeability barrier. When SCs were cultured at 0.5x10^6 cells/cm2 on Matrigel-coated dishes for 5 days to allow the assembly of both TJs and AJs, freshly isolated GCs were then added onto the SC epithelium at a SC-GC ratio of 1:1 to induce SC-GC AJ assembly. There was an increase in iNOS expression, but not eNOS, coinciding with the assembly of SC-GC AJs. To further explore the involvement of iNOS in SC-GC AJ dynamics, an in vivo model was used in which 1,2,4-dichlorobenzyl)-indazole-3-carboxyhydrate (DICC) was used to induce GC depletion from the epithelium. A significant increase in iNOS, but not eNOS,