Epac Activates the Small G Proteins Rap1 and Rab3A to Achieve Exocytosis*

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Exocytosis of the acrosome (the acrosome reaction) relies on cAMP production, assembly of a proteinaceous fusion machinery, calcium influx from the extracellular medium, and mobilization from inositol 1,4,5-trisphosphate-sensitive intracellular stores. Addition of cAMP to human sperm suspensions bypasses some of these requirements and elicits exocytosis in a protein kinase A- and extracellular calcium-independent manner. The relevant cAMP target is Epac, a guanine nucleotide exchange factor for the small GTPase Rap. We show here that a soluble adenyl cyclase synthesizes the cAMP required for the acrosome reaction. Epac stimulates the exchange of GDP for GTP on Rap1, upstream of a phospholipase C. The Epac-selective cAMP analogue 8-pCPT-2′-O-Me-cAMP induces a phospholipase C-dependent calcium mobilization in human sperm suspensions. In addition, our studies identify a novel connection between cAMP and Rab3A, a secretary granule-associated protein, revealing that the latter functions downstream of soluble adenylyl cyclase/cAMP/Epac but not of Rap1. Challenging sperm with calcium or 8-pCPT-2′-O-Me-cAMP boosts the exchange of GDP for GTP on Rab3A. Recombinant Epac does not release GDP from Rab3A in vitro, suggesting that the Rab3A-GEF activation by cAMP/Epac in vivo is indirect. We propose that Epac sits at a critical point during the exocytic cascade after which the pathway splits into two limbs, one that assembles the fusion machinery into place and another that elicits intracellular calcium release.

During fertilization in eutherian mammals, the spermatozoon must penetrate the zona pellucida to reach the oolemma. Only sperm that have completed the acrosome reaction (AR)4 can successfully accomplish this task (1). The AR is a regulated exocytosis where the membrane of the acrosome, the single dense core secretary granule in sperm, fuses to the plasma membrane surrounding the anterior portion of the head. This process releases hydrolytic enzymes stored in the granule. These enzymes, together with the physical thrust derived from strong flagellar beating, enable sperm to penetrate the zona pellucida (1, 2). Physiological agonists accomplish the AR by inducing an influx of calcium from the extracellular medium and the assembly of a conserved proteinaceous fusion machinery that includes Rab3A, α-SNAP/NSF, synaptotagmin, complexin, and neurotoxin-sensitive SNAREs; the AR also requires an efflux of calcium from inside the acrosome through IP3-sensitive channels (reviewed in Refs. 3, 4).

In certain neurons, neuroendocrine and exocrine acinar cells, cAMP potentiates calcium-dependent exocytosis. Either cAMP-dependent protein kinase (PKA) or the exchange protein directly activated by cAMP (Epac) can be the targets of cAMP in the cAMP-regulated exocytosis. On the other hand, cAMP is the principal trigger of regulated secretion in various non-neuronal cells (5–7). Likewise, an elevation of cAMP alone is sufficient to trigger exocytosis in human sperm. Moreover, calcium relies on endogenous cAMP to accomplish acrosomal release, and it does so through a PKA-insensitive pathway involving Epac. The stimulation of endogenous Epac by the selective cAMP analogue 8-(p-chlorophenylthio)-2′-O-methyladenosine-3′,5′-cyclic monophosphate (8-pCPT-2′-O-Me-cAMP) is sufficient to trigger the AR even in the absence of extracellular calcium. Furthermore, when Epac is sequestered with specific antibodies, cAMP, calcium (8), and recombinant Rab3A (this study) are unable to elicit exocytosis.
Epac Activates Rap1 and Rab3A during Exocytosis

Epac1 and Epac2 are multidomain proteins that consist of an N-terminal regulatory region and a C-terminal catalytic region (9–11). The regulatory domain harbors the cAMP-binding site, which auto-inhibits the catalytic activity in the absence of cAMP (12–15). The catalytic portion bears a guanine-nucleotide exchange factor (GEF) activity specific for Rap1 and Rap2 (16, 17). Like all small G proteins, Raps cycle between an inactive GDP-bound and an active GTP-bound conformation. The GDP-GTP cycle is regulated by GEFs that induce the release of the bound GDP to be replaced by the more abundant GTP and by GTPase-activating proteins that coax the intrinsic GTPase activity to rapidly hydrolyze bound GTP, returning the G proteins to the inactive GDP-bound state (18, 19). Most small G proteins are linked to biological membranes via lipid modifications at their C terminus; for instance, Rap2A is farnesylated, and Rap1A/B, Rap2B, and Rabs are geranylgeranylated (20, 21). Guanine nucleotide dissociation inhibitors (GDIs) remove Rabs from membranes by sequesterization of their lipid tails (22).

Extracellular stimuli often result in the activation of cellular adenylate cyclases and an increase in CAMP levels. By serving as a cAMP-binding protein with intrinsic GEF activity, Epac couples cAMP production to a variety of Rap-mediated processes such as the control of cell adhesion and cell-cell junction formation, water resorption, cell differentiation, inflammatory processes, etc. (9–11). Many are the effectors of Epac and Epac-Rap signaling. Of particular interest to us is the observation that Epac stimulates phospholipase C (PLC) through the activation of Rap1 and -2, resulting in IP3-mediated release of calcium from internal stores (23, 24). PLCe is an unusual enzyme with two catalytic activities as follows: the typical phosphatidylinositol 4,5-bisphosphate hydrolyzing PLC activity plus a Rap-GEF activity. Thus, PLCe acts both downstream and upstream of Ras-like GTPases, perhaps to guarantee sustained Rap signaling (25).

During membrane fusion, Rab proteins direct the recognition and physical attachments of the compartments that are going to fuse (26, 27). This association, or tethering, represents one of the earliest known events in membrane fusion and is accomplished through the recruitment of tethering factors. Rab3A localizes to vesicles and secretory granules and is one of the isoforms directly implicated in regulated exocytosis of neurotransmitters and hormones (28). Rab3A interacts in a GTP-dependent manner with at least two effector proteins, rabphilin and Rim (29–31). Rab3A is present in the acrosomal region of human (32), rat (33), and mouse sperm (34). Rab3A (full-length recombinant protein or a synthetic peptide corresponding to the effector domain) stimulates human (32, 35) and ram (36) and inhibits rat sperm AR (33). Rab3A is required for the AR triggered by calcium (37, 38) and cAMP (8).

Epac is a multifunctional protein in which cAMP exerts its effects not only by promoting the exchange of GDP for GTP on Rap but also by allosterically regulating other molecules (10). In exocytosis for instance, a number of Rap-independent, Epac-linked signaling pathways have been described. They include the interaction of Epac2 with Rim2 (39) and the Rim2-related protein Piccolo (40). Epac2 also stimulates exocytosis by interacting with SUR1 (41). Finally, Epac2 controls ryanodine-sensitive calcium channels that are involved in calcium-induced calcium release (CICR) from internal stores in insulin-secreting cells (42).

In this study, we piece together the analysis of two phenomena as follows: calcium mobilization and protein–protein interactions preceding exocytosis. To the best of our knowledge, this constitutes the first integrated molecular model that includes both the assembly of the fusion and intravesicular calcium release protein machineries during regulated exocytosis. By enquiring further into the signaling pathways operating during sperm exocytosis, we have found more players than previously suspected, and we discovered that the key components of these cascades are not arranged in a linear sequence. Epac sits at a central point of the signaling cascade after which the exocytotic pathway splits into two limbs as follows: one that assembles the fusion machinery into place, and another that elicits the release of calcium from the acrosome; both need to act in concert to achieve exocytosis. Our results identify Rab3A for the first time as a downstream target for Epac and place this small GTPase as an early component of the “fusion machinery” branch of the pathway. They also show that Epac stimulates the exchange of GDP for GTP on Rap1 and that this protein, as well as a PLC, drives intracellular calcium mobilization. Finally, our data reveal that a soluble adenyl cyclase (sAC) (43, 44) synthesizes the cAMP that activates Epac. Again, we believe that this is the first report linking sAC to an exocytic event.

EXPERIMENTAL PROCEDURES

Reagents—Recombinant streptolysin O (SLO) was obtained from Dr. Bhakdi (University of Mainz, Mainz, Germany). Spermatooza were cultured in human tubal fluid media (as formulated by Irvine Scientific, Santa Ana, CA) supplemented with 0.5% bovine serum albumin (HTF media). The rabbit polyclonal antibodies to Rap1A/B and Rab3A (purified IgG) were from Santa Cruz Biotechnology, Inc. (Santa Cruz, CA). The rabbit polyclonal anti-NSF (whole serum) were from Synaptic Systems (Gottingen, Germany), and rabbit polyclonal antibody to Rab11 was from Invitrogen. The rabbit polyclonal antibodies against Epac were generated by Genemed Synthesis, Inc. (San Francisco), using the synthetic peptide LREDNCHFLRVDK, and affinity-purified on immobilized Epac peptide (8). Anti-Rap1A/B rabbit polyclonal antibodies were raised against the peptide EDERVVGKEQGQNLC and affinity-purified on immobilized peptide (GenScript Corp., Piscataway, NJ). Horse-radish peroxidase-conjugated goat anti-rabbit IgG (anti-γ chain) was from Jackson ImmunoResearch (West Grove, PA). Glutathione S-transferase (GST)-tagged, human recombinant cAMP-specific phosphodiesterase 4D (PDE), and Rap1A were from Abnova Corp. (NeiHu, Taipei, Taiwan). KH7 was purchased from ChemDiv, Inc. (San Diego). 8-pCPT2′-O-Me-cAMP, N6-benzoyleadenosine-3′,5′-cyclic monophosphate (6-Bnz-cAMP), and 2′,3′-O-(N′-methylanthraniloyl)-GDP (mGDP) were from Biolog-Life Science Institute (Bremen, Germany). U73122 (1-(6-((17-yl)amino)hexyl)-1H-pyrole-2,5-dione) and the inactive analogue U73343 (1-(6-(17-yl)amino)hexyl)-2,5-pyrrolidine-dione) were from Biomol International L.P. (Plymouth Meeting, PA). Adenophostin A, hexasodium salt, and 2-aminoethoxydiphenyl borate (2-APB) from Calbiochem were purchased from Merck Quimica Argentina S.A.I.C. (Buenos Aires, Argentina). O-Nitrophenyl EGTA-
lysates were frozen until use. His6-NSF Y83E was expressed and except GST-Ral-GDS-RBD and GST-RIM, where bacterial were quantified on a 3550 Microplate Reader (Bio-Rad). Bovine serum albumin was used as a standard, and the results mined by the Bio-Rad protein assay in 96-well microplates. Rab3A proteins were prenylated for all but the were adjusted to 5–10 \times 10^6/ml before incubating for at least 2 h under capacitating conditions (HTF, 37 °C, 5% CO2, 95% air). For experiments involving intact sperm, we added increasing concentrations of KH7 and 10 \mu M A23187 or 5 \mu M proges- terone sequentially and incubated for 10–15 min at 37 °C after each addition. For experiments involving permeabilized cells, washed spermatozoa were resuspended in cold PBS containing 2.1 units/ml SLO for 15 min at 4 °C. Cells were washed once with PBS and resuspended in ice-cold sucrose buffer (250 mm sucrose, 0.5 mm EGTA, 20 mm Hepes-K, pH 7) containing 2 mm dithiothreitol. We added inhibitors and stimulants sequentially as indicated in the figure legends and incubated for 10–15 min at 37 °C after each addition. Where indicated, we preloaded SLO-permeabilized sperm with NP-EGTA-AM before incubating with inhibitors and/or calcium, carrying out all procedures in the dark. Photolysis was induced after the last incubation by exposing twice (1 min each time) to an UV transilluminator and mixing after each exposure. Intact and permeabilized sperm were spotted on Teflon-printed slides, air-dried, and fixed/permeabilized in ice-cold methanol for 1 min. Acrosomal status was evaluated by staining with FITC- coupled Pisum sativum (FITC-PSA) (52). At least 200 cells were scored using an upright Nikon microscope equipped with epifluorescence optics. Basal (no stimulation) and positive (0.5 mm CaCl2 corresponding to 10 \mu M free calcium estimated by MAX-CHELATOR, a series of program(s) for determining the free metal concentration in the presence of chelators) controls were included in all experiments. Acrosomal exocytosis indexes were calculated by subtracting the number of spontaneously reacted spermatozoa from all values and expressing the results as a percentage of the AR observed in the positive control. Data were evaluated using one way analysis of variance. The Tukey-Kramer post hoc test was used for pairwise comparisons. Differences were considered significant at the p < 0.05 level.

In Vitro Activation of Rab3A—Experiments were performed as described (50, 51), using Rab3A in addition to Rap1. Purified Rab3A or Rap1 (200 nM, GST removed with thrombin) loaded with the fluorescent GDP analogue mGDP was incubated in the presence of 20 \mu M unlabeled GDP plus 150 \mu M purified Epac1 or Epac2. Cyclic AMP (100–500 \mu M) was added as indicated. The nucleotide exchange was measured in real time as decay in fluorescence using a Cary Eclipse spectrofluorometer (Varian, Varian B.V., Middelburg, The Netherlands). The decay is caused by the release of protein-bound mGDP, which shows higher fluorescence intensity in the hydrophobic environment of the protein than in the buffer solution. The decay in the fluorescence signal is equal to the rate of nucleotide dissocia- tion, and nucleotide exchange should be accelerated in the presence of an active GEF. All data analysis, fitting, and plotting were done with the Graft 3.0 program (Erithacus software).

SLO Permeabilization and AR Assay—Human semen samples were obtained from normal healthy donors. Semen was allowed to liquefy for 30–60 min at 37 °C. We used a swim-up protocol to isolate highly motile sperm. Acrosomal status was evaluated by staining with FITC- coupled Pisum sativum (FITC-PSA) (52). At least 200 cells were scored using an upright Nikon microscope equipped with epifluorescence optics. Basal (no stimulation) and positive (0.5 mm CaCl2 corresponding to 10 \mu M free calcium estimated by MAX-CHELATOR, a series of program(s) for determining the free metal concentration in the presence of chelators) controls were included in all experiments. Acrosomal exocytosis indexes were calculated by subtracting the number of spontaneously reacted spermatozoa from all values and expressing the results as a percentage of the AR observed in the positive control. Data were evaluated using one way analysis of variance. The Tukey-Kramer post hoc test was used for pairwise comparisons. Differences were considered significant at the p < 0.05 level.

Rab3-GTP and Rap1-GTP Precipitation Assays—Capaci- tated sperm (10–50 \times 10^6 cells) were washed twice and sus- pended in PBS. Sperm were treated with 100 \mu M 2-APB (an IP3-sensitive calcium channel blocker) to prevent cytosol/ membrane loss due to exocytosis before adding the AR inducers. Alternatively, sperm were permeabilized with SLO and treated with 2-APB and 20 \mu g/ml anti-Rap1 antibodies (GenScript) or 1 \mu g of His6- Rab3A before challenging with 50 \mu M 8-pCPT-2’-O-Me-cAMP or 0.5 mm CaCl2. After 10 min of incubation at 37 °C, cells were lysed in GST pulldown buffer (200 mm NaCl, 2.5 mm MgCl2, 1% (v/v) Triton X-100, 10% glycerol, 1 mm phenylmethylsulfonyl fluoride, 1X protease inhibitor mixture (P2714, Sigma), and 50 mm Tris-HCl, pH 7.4) by sonication on ice (two times for 15 s). We let proteins diffuse into the lysis buffer for 15 min at 4 °C. These whole cell deter-
gent extracts were clarified by centrifugation at 12,000 × g for 5 min and used immediately.

Glutathione-Sepharose beads were washed twice with GST pull-down buffer and incubated with bacterial lysates containing GST–Ral-GDS-RBD or GST–RIM for 1 h at 4 °C under constant rocking. Beads were washed twice with PBS and once with GST pull-down buffer and used immediately. Twenty μl of glutathione-Sepharose containing 10 μg of the appropriate fusion protein was added to sperm lysates in a total volume of 0.6 ml and incubated by rotation at 4 °C for 30 min. The resin was recovered by centrifugation at 4 °C (5 min at 10,000 rpm) and washed three times with ice-cold GST pull-down buffer. The resin-bound fractions were resolved by SDS-PAGE, and cellular GTP–Rap1 and GTP–Rab3A levels were analyzed by immunoblotting as described later.

**Triton X-114 Partition**—Partition experiments were conducted following standard procedures (53, 54). Briefly, capacitated sperm (100 × 10^6 cells) were washed twice with cold PBS, and proteins were extracted in 1 ml of lysis buffer containing 20 mM Tris-HCl, pH 7.5, 150 mM NaCl, 10% glycerol, 5 mM MgCl2, and 1% Triton X-114 by sonication on ice (three times for 15 s with 10-s intervals). The lysates were rocked for 45 min at 4 °C and spun at 14,000 × g for 20 min at 4 °C. These whole cell detergent extracts were incubated for 15 min at 30 °C and centrifuged at 3,000 × g for 2 min. Hydophobic proteins partitioned into the upper (aqueous) phase, whereas hydrophobic proteins were recovered from the lower (detergent) phase. Protein precipitation and removal of detergent were achieved via extraction with CC13H–CH3OH. Precipitated proteins were dissolved in sample buffer by heating once at 60 °C for 10 min and once at 95 °C for 3 min.

**SDS-PAGE and Western Blots**—Proteins were separated on SDS-gel electrophoresis (55) and transferred to 0.22-μm nitrocellulose membranes (Hybond, GE Healthcare). Nonspecific reactivity was blocked by incubation for 1 h at room temperature with 5% skim milk dissolved in washing buffer (PBS, pH 7.6, 0.1% Tween 20). Blots were incubated with the anti-Rap1 (0.5 μg/ml) or the custom-made anti-Rap1 (0.5 μg/ml in blocking solution) antibodies in blocking solution for 1 h at room temperature or overnight at 4 °C. Horseradish peroxidase-conjugated goat anti-rabbit IgG was used as secondary antibody (3 μg/ml) with 1-h incubations. Excess first and second antibodies were removed by washing five times for 7 min in washing buffer. Detection was accomplished with a chemiluminescence system (Western Lightning, PerkinElmer Life Sciences, Migliore Laclaustra, Buenos Aires, Argentina) and subsequent exposure to CL-XPosure film (Tecnolab, Pierce) for 1–10 min.

**Calcium Measurements**—Calcium levels were measured in sperm suspensions using the intracellular fluorescent probe Fluo-3. Motile sperm were adjusted to a concentration of 5–10 × 10^6 cells/ml and loaded with the permeable form of the dye (Fluo-3-AM, 2 μM) and 0.02% of pluronic acid for 30 min at 37 °C. Cells were washed once and resuspended in nominally calcium-free (~2 μM) medium (10 mM Hepes-Na, 120 mM NaCl, 4 mM KCl, 15 mM NaHCO3, 1 mM MgCl2, 5 mM d-glucose, 1 mM sodium pyruvate, 10 mM lactic acid, pH 7.4). Sperm suspensions were transferred to thermostated (37 °C) cuvettes for fluorescence measurements and stirred constantly. At the indicated times, 50 μM 8-pCPT-2′-O-Me-cAMP was added to the samples with or without previous incubation with 1 μM U73122 or U73343. Fluo-3 fluorescence (λ_ex = 505 nm, λ_em = 525 nm emission) was recorded in an Amino 8000 spectrofluorometer. Data were collected during 600 s at a frequency of 0.5 Hz. To calibrate the maximal response, [Ca2+]i was determined using Triton X-100 (0.1%). Measurements were performed at least five times with different batches of sperm. The criterion we used for inclusion of the results obtained with a given sample into the statistical analysis was the normal response of the said sample to 4 μM progesterone. The Tukey-Kramer post hoc test was used for pairwise comparisons. Differences were considered significant at the p < 0.05 level.

**RESULTS**

**sAC Synthesizes cAMP Required for Human Sperm Exocytosis**—Previous findings from our laboratory (8, 49) had led us to formulate the AR as summarized in Pathway 1, where single arrows indicate there is one step between the terms connected, and double arrows indicate that the number of steps taking place between the connected terms is unknown.

\[
\text{calcium} \rightarrow \text{cAMP} \rightarrow \text{Epac} \rightarrow \text{Rab3A} \rightarrow PTP1B \rightarrow \alpha\text{-SNAP/NSF} \rightarrow \text{SNAREs} \rightarrow \text{intracellular calcium mobilization} \rightarrow \text{exocytosis}
\]

**PATHWAY 1**
Epac Activates Rap1 and Rab3A during Exocytosis

This pathway begins with the sequence “calcium → cAMP → Epac.” In nonpermeabilized sperm, calcium enters the cells from the extracellular milieu when channels open in the plasma membrane in response to physiological agonists or when a calcium ionophore transports it to the cytosol; in permeabilized sperm, calcium enters through the SLO-generated pores. Because depletion of endogenous cAMP by an active PDE prevents the AR induced by calcium (8), we know that this cyclic nucleotide is required to accomplish exocytosis. Where does this cAMP come from? We found that the pool of cAMP relevant for the AR is synthesized by sAC, because the ARs of at least three independent experiments.

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role as an AR inducer. When added to SLO-permeabilized sperm, two different anti-Rap1 antibodies effectively abrogated calcium-triggered AR (Fig. 2B, bottom). The effect of the custom-made anti-Rap1 polyclonal was diminished by preincubation with the synthetic peptide against which the antibody was raised. Likewise, preincubation of the commercial anti-Rap1 antibody with recombinant GST-Rap1 decreased the ability of the antibody to prevent the AR. Neither a nonimmune rabbit IgG nor an antibody to an unrelated small G protein (anti-Rab11) had any effect (Fig. 2B). These data suggest that the inhibitory effect of the anti-Rap1 antibodies was specific and due to binding to endogenous Rap1. Taken together, these results reveal that Rap1 is necessary for human sperm acrosomal exocytosis.

Many hormones and agonists that elevate intracellular cAMP levels increase the amount of active, GTP-bound Rap1 in responsive cells. We therefore asked whether the onset of the AR induces Rap1 activation in human sperm. This was indeed the case because when we pulled down GTP-bound Rap1 with immobilized Ral-GDS-RBD we found that the amount of active Rap1 increased in response to 8-pCPT-2′-O-Me-cAMP (Fig. 2C). Next, we introduced Ral-GDS-RBD into SLO-permeabilized sperm before challenging with AR inducers, hypothesizing that this cassette would sequester Rap1-GTP if formed in response to a stimulus. 8-pCPT-2′-O-Me-cAMP was our positive control because we knew that activating Epac exchanged GDP for GTP on Rap1 (Fig. 2C). Rap1-GTP blocked the AR elicited by 8-pCPT-2′-O-Me-cAMP (Fig. 2D), indicating that Rap1-GTP was necessary for the AR and validating the use of the cassette to deplete the pool of active Rap1. Calcium was unable to achieve exocytosis in the presence of Ral-GDS-RBD (Fig. 2D), suggesting that this inducer also promoted the activation of Rap1 and required Rap1-GTP to elicit the AR. GST alone, used as negative control, had no effect on calcium-triggered exocytosis (Fig. 2D).

Rap1 Is Located Upstream of Calcium Mobilization from an Intracellular IP3-sensitive Store, Likely the Acrosome—The acrosome behaves as an internal store of releasable calcium (59, 60); efflux from this reservoir through IP3-sensitive channels is required for the AR initiated by calcium itself, Rab3A-GTP-γ-S, and cAMP (8, 35, 38, 59). To assess whether the requirement for Rap1 takes place prior to or following intra-acrosomal calcium efflux, we resorted to the photolabile calcium chelator NP-EGTA-AM. In our SLO-permeabilized human sperm model NP-EGTA-AM crosses the plasma and outer acrosomal membranes, accumulates inside the acrosome, and prevents the AR by sequestering intra-acrosomal calcium (59). UV photolysis of NP-EGTA-AM rapidly replenishes the acrosomal calcium pool, resuming exocytosis (Fig. 3A, gray bars). In combination with specific blockers, NP-EGTA-AM allowed us to place the requirement for the target of the blockers before or after the intra-acrosomal calcium-sensitive step. Briefly, we loaded SLO-permeabilized sperm with NP-EGTA-AM and challenged with an AR inducer (extracellular calcium in this case), allowing the fusion-related signaling pathways to advance up to the point where intra-acrosomal calcium

![FIGURE 3. Rap1 participates in a pathway that mobilizes calcium from an IP3-sensitive store. A, permeabilized spermatozoa were loaded with 10 μM NP-EGTA-AM (NP) for 10 min at 37 °C to chelate intra-acrosomal calcium. The AR was subsequently initiated by adding 0.5 mM CaCl2. After a further 15-min incubation at 37 °C to allow exocytosis to proceed up to the intra-acrosomal calcium-sensitive step, sperm were treated for 15 min at 37 °C with 20 μg/ml anti-Rap antibody (Santa Cruz Biotechnology) or 1 μg/ml Ral-GDS-RBD. All these procedures were carried out in the dark. UV flash photolysis of the chelator was induced at the end of the incubation period (hv), and the samples were incubated for 5 min (NP → calcium → inhibitor → hv, black bars). Several controls were run (gray bars) as follows: background AR in the absence of any stimulation (control); AR stimulated by 0.5 mM CaCl2 (calcium), inhibitory effect of NP-EGTA-AM in the dark (NP → calcium → dark), and the recovery upon illumination (NP → calcium → hv); and the effect of the inhibitors when present throughout the experiment (NP → inhibitor → calcium → hv). Sperrn were fixed; the AR was measured by FITC-PSA binding, and the data were normalized as described under “Experimental Procedures.” B, permeabilized spermatozoa were incubated with 15 μM U73343 or U73122 for 15 min at 37 °C. Acrosomal exocytosis was then initiated by adding 0.5 mM CaCl2 (calcium) or 50 μM 8-pCPT-2′-O-Me-cAMP (8pCPT) and a further 15-min incubation at 37 °C (black bars). Sperm were stained, and the AR scored as in A. Control permeabilized spermatozoa were incubated with 20 μg/ml anti-Rap antibodies (Santa Cruz Biotechnology), 1 μg/ml Ral-GDS-RBD, or 15 μM U73122 for 10 min at 37 °C. Exocytosis was initiated by adding 0.5 mM CaCl2. To demonstrate that Rap1-GTP and an active PLC were located upstream of the intracellular calcium efflux, 5 μM adenophostin was added to mobilize calcium from IP3-sensitive stores, and incubations proceeded for an additional 10 min at 37 °C (black bars). Controls (gray bars) included the following: background AR in the absence of any stimulation (control), AR stimulated by 0.5 mM CaCl2 (calcium), AR inhibition by 20 μg/ml anti-Rap antibodies, 1 μg/ml Ral-GDS-RBD, or 15 μM U73122, and AR unaffected by 5 μM adenophostin. Sperm were stained, and the AR scored as in A. Plotted results represent the mean ± S.E. of at least three independent experiments.]

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**Epac Activates Rap1 and Rab3A during Exocytosis**

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**A**

| Control | NP | Calcium |
|---------|-----|---------|
|         |     |         |

**B**

| Control | 8CPT | Calcium |
|---------|------|---------|
|         |      |         |

**C**

| Control | Calcium |
|---------|---------|
|         |         |

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**FIGURE 3.** Rap1 participates in a pathway that mobilizes calcium from an IP3-sensitive store. A, permeabilized spermatozoa were loaded with 10 μM NP-EGTA-AM (NP) for 10 min at 37 °C to chelate intra-acrosomal calcium. The AR was subsequently initiated by adding 0.5 mM CaCl2. After a further 15-min incubation at 37 °C to allow exocytosis to proceed up to the intra-acrosomal calcium-sensitive step, sperm were treated for 15 min at 37 °C with 20 μg/ml anti-Rap antibody (Santa Cruz Biotechnology) or 1 μg/ml Ral-GDS-RBD. All these procedures were carried out in the dark. UV flash photolysis of the chelator was induced at the end of the incubation period (hv), and the samples were incubated for 5 min (NP → calcium → inhibitor → hv, black bars). Several controls were run (gray bars) as follows: background AR in the absence of any stimulation (control); AR stimulated by 0.5 mM CaCl2 (calcium), inhibitory effect of NP-EGTA-AM in the dark (NP → calcium → dark), and the recovery upon illumination (NP → calcium → hv); and the effect of the inhibitors when present throughout the experiment (NP → inhibitor → calcium → hv). Sperrn were fixed; the AR was measured by FITC-PSA binding, and the data were normalized as described under “Experimental Procedures.” B, permeabilized spermatozoa were incubated with 15 μM U73343 or U73122 for 15 min at 37 °C. Acrosomal exocytosis was then initiated by adding 0.5 mM CaCl2 (calcium) or 50 μM 8-pCPT-2′-O-Me-cAMP (8pCPT) and a further 15-min incubation at 37 °C (black bars). Sperm were stained, and the AR scored as in A. Control permeabilized spermatozoa were incubated with 20 μg/ml anti-Rap antibodies (Santa Cruz Biotechnology), 1 μg/ml Ral-GDS-RBD, or 15 μM U73122 for 10 min at 37 °C. Exocytosis was initiated by adding 0.5 mM CaCl2. To demonstrate that Rap1-GTP and an active PLC were located upstream of the intracellular calcium efflux, 5 μM adenophostin was added to mobilize calcium from IP3-sensitive stores, and incubations proceeded for an additional 10 min at 37 °C (black bars). Controls (gray bars) included the following: background AR in the absence of any stimulation (control), AR stimulated by 0.5 mM CaCl2 (calcium), AR inhibition by 20 μg/ml anti-Rap antibodies, 1 μg/ml Ral-GDS-RBD, or 15 μM U73122, and AR unaffected by 5 μM adenophostin. Sperm were stained, and the AR scored as in A. Plotted results represent the mean ± S.E. of at least three independent experiments.
release is required. Subsequently, we added AR blockers and illuminated the tubes. In this scenario, resistance to a blocker (revealed by unaffected exocytosis) implies that its target is required upstream of intra-acrosomal calcium efflux, whereas sensitivity to a blocker (revealed by inhibited exocytosis) suggests that its target is necessary after the intra-acrosomal calcium-sensitive step. The AR was unaffected by both anti-Rap1 antibodies and Ral-GDS-RBD (Fig. 3A, black bars), indicating that Rap1-GTP is necessary early in the fusion cascade, before calcium is released from the acrosome.

In HEK-293 cells, cAMP increases intracellular calcium concentrations and IP₃ synthesis by stimulating a PLC/H₂K₃ activity in an Epac1/Rap2B-dependent manner (23). Likewise, the pathway Epac-Rap1-PLC/H₂K₃ enhances calcium mobilization from the sarcoplasmic reticulum in ventricular cardiac myocytes (24). The AR triggered by 8-pCPT-2’-O-Me-cAMP relies on Epac, an active Rap1 (Fig. 2D), and on the mobilization of intravesicular calcium through IP₃-sensitive receptors (8). To ask whether sperm exocytosis requires the activity of a PLC, we introduced U73122, an inhibitor of multiple phosphatidylinositol-specific phospholipase C (PI-PLC) isoforms, into permeabilized sperm before challenging with calcium or 8-pCPT-2’-O-Me-cAMP. Treatment with U73122 (but not the inactive control U73343) decreased the exocytotic response by >90% (Fig. 3B), suggesting that the AR requires an active PLC downstream of Epac activation.

We hypothesized that signaling through Rap1 and PLC, with the ensuing synthesis of IP₃, drives the intracellular calcium mobilization required for the AR. If this was the case, then Rap (Fig. 2, B and D) and PLC (Fig. 3B) inhibitors blocked exocytosis because they prevented calcium mobilization. We tested this possibility by promoting intravesicular calcium release with the IP₃ receptor agonist adenophostin. The sole addition of adenophostin rescued exocytosis impaired by anti-Rap1 antibodies, Ral-GDS-RBD, and U73122 (Fig. 3C), supporting the notion that the end point of the Rap-PLC pathway is the mobilization of intracellular calcium.

8-pCPT-2’-O-Me-cAMP Augments the Concentration of Cytosolic Calcium in Human Sperm Suspensions—Human sperm bathed in a nominally calcium-free medium and loaded with the fluorescent calcium sensor Fluo-3-AM responded to 8-pCPT-2’-O-Me-cAMP with a transient increase in calcium concentration whose profile was similar to that elicited by progesterone under identical conditions (Fig. 4A). Preincubation with U73122, and to a much lesser degree with U73343, diminished the increase in calcium caused by the Epac activator (Fig. 4B). These data indicate that, in conditions identical to those in which it elicits exocytosis in sperm, 8-pCPT-2’-O-Me-cAMP increases the concentration of calcium in the cytosol through a mechanism involving an active PLC.

Based on the data collected here we reasoned that, in addition to the sequence depicted in pathway 1, the AR can also be

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**FIGURE 4. Epac signaling mobilizes calcium in a PLC-dependent manner.** Human sperm (nonpermeabilized) were loaded with 2 μM Fluo-3-AM and 0.02% pluronic acid and incubated for 30 min at 37 °C. At the indicated times (arrows) 50 μM 8-pCPT-2’-O-Me-cAMP (8pCPT) or 4 μM progesterone (Pg) were added with (B) or without (A) previous incubation with 1 μM U73122 or U73343. Maximal [Ca²⁺]ᵢ response was calibrated with 0.1% Triton X-100 (TX-100) at the end of the incubation period. Shown are traces representative of five experiments. The increase in fluorescence is expressed as (F/F₀) – 1 (maximum fluorescence intensity/initial fluorescence) – 1) versus time in seconds. Bars represent mean ± S.E. of five experiments; data were normalized against the calcium response to 4 μM progesterone (A) or 50 μM 8-pCPT-2’-O-Me-cAMP (B). ***, p < 0.001; *, p < 0.05 compared with 8 pCPT-2’-O-Me-cAMP.
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modeled to fit in a signaling pathway operating through the sequence shown in pathway 2,

\[
\text{calcium} \rightarrow -\rightarrow \text{cAMP} \rightarrow \text{Epac} \rightarrow \text{Rap-GTP} \rightarrow -\rightarrow \text{PLC} \rightarrow \text{IP}_3 \rightarrow \text{intracellular calcium mobilization} \rightarrow -\rightarrow \text{exocytosis}
\]

**PATHWAY 2**

In the following sections we will attempt to address the question of how do sperm signaling pathways summarized in pathways 1 and 2 integrate to accomplish exocytosis.

**AR Triggered by Recombinant Rab3A-GTP-γ-S Requires cAMP, Epac, Rap1-GTP, and a PLC Activity**—In addition to the early steps “calcium →→ cAMP → Epac,” pathways 1 and 2 share the late-acting steps “intracellular calcium mobilization →→→ exocytosis.” We examined these pathways further by investigating whether the AR triggered by recombinant Rab3A (Rab3A is a term in pathway 1) requires cAMP, Rap-GTP, and PLC. In human sperm, prenylated, GTP-γ-S-loaded, recombinant Rab3A induces a strong exocytotic response that relies on the same fusion machinery characterized for calcium-triggered exocytosis (32, 35). Here we show that the Rab3A-GTP-γ-S-sculpted AR also requires cAMP because pretreatment of permeabilized human sperm with KH7 or recombinant PDE impaired exocytosis (Fig. 5A). The PKA blocker H89 did not abolish the AR, in agreement with previous observations that this kinase is not the target of cAMP during sperm exocytosis (8). To investigate if Rab3A signals through cAMP/Epac, as is the case when calcium serves as the AR inducer, we introduced anti-Epac antibodies into permeabilized cells. The antibodies (6.7 nM) efficiently blocked the AR induced by calcium (Fig. 5B) (8). The AR triggered by recombinant Rab3A-GTP-γ-S was also abolished by anti-Epac antibodies, albeit at a higher concentration (134 nM, Fig. 5B). Next, we analyzed the contribution of the other players depicted in pathway 2, and we found that sequestering endogenous Rap1 with specific antibodies or Rap1-GTP with Ral-GDS-RBD prevented the AR elicited by Rab3A-GTP-γ-S. The same was true when we inhibited phosphoinositide-specific PLC activity with U73122 (Fig. 5B). Taken together, these data suggest that Rab3A-GTP-γ-S requires cAMP synthesized by sAC to achieve exocytosis and that Epac is the relevant cAMP target. Active Rap1 and PLC are also part of the pathway initiated by Rab3A-GTP-γ-S. In other words, recombinant Rab3A-GTP-γ-S relies on all the players summarized in pathway 2 to achieve exocytosis.

**Calcium Drives the AR through a Bifurcated Signaling Pathway**—The two simplest models we can think of to represent the AR are as follows: (i) a linear cascade consisting of an arrangement of all players depicted in pathways 1 and 2; (ii) a pathway that bifurcates after some common early steps into parallel cascades that converge later to achieve fusion. We favor the second alternative and hypothesize that following Epac activation, the pathway splits into a branch that begins with Rap1 and ends in intracellular calcium mobilization and another that starts with Rab3A and finishes with the assembly of SNARE proteins in loose trans complexes (38), both being necessary for exocytosis (see model in Fig. 9). We tested the validity of this proposal with adenophostin, predicting that it should not be able to rescue the effect of inhibitors whose targets are early common steps of the pathway (i.e. cAMP and Epac) or those required to assemble SNAREs (i.e. Rab3A, PTP1B, α-SNAP, and NSF). Data depicted in Fig. 6 show that this was indeed the case. Adenophostin failed to reverse the AR block caused by recombinant PDE (2 μg/ml), KH7 (10 μM), and anti-Epac antibodies (134 nM) (Fig. 6A). The activity of NSF is inhibited by tyrosine phosphorylation and de-repressed by tyrosine phosphatases (TPPs) in macrophages (61) and sperm (49). In the latter, PTP1B disassembles SNARE complexes through tyrosine dephosphorylation of NSF. As expected, adenophostin failed to rescue the AR block imposed by the substrate trapping mutant PTP1B D181A (300 nM) and the phosphomimetic NSF mutant NSF Y83E (186 nM). Similarly, adenophostin did not rescue the anti-Rab3A (20 μg/ml) antibody block (Fig. 6B). Interestingly, adenophostin did rescue exocytosis when the anti-Epac antibodies were added at 6.7 nM (Fig. 6A). We interpret the concentration-dependent behavior as indicative that at 6.7 nM anti-Epac antibodies prevent calcium-triggered exocytosis because they block only the “Rap1”
branch (the “Rab3A” branch proceeds normally), whereas at 134 nM they block both (Fig. 9).

The reversibility of the inhibition of the anti-Epac antibodies (6.7 nM) by adenophostin supplied us with an alternative strategy to test the bifurcation hypothesis. When added after the AR inducer (extracellular calcium in this case) in the presence of anti-Epac antibodies (rescued at the end of the incubation by adenophostin), neither anti-Rab3A nor anti-NSF antibodies blocked the AR (Fig. 6C, black bars). These results suggest that the AR inducer had driven the Rab3A branch of the pathway to completion even though the Rap1 arm had been halted by the anti-Epac antibodies. In other words, the steps catalyzed by Rab3A/NSF had already taken place when we added the antibodies, and therefore the system was refractory to their inhibitory effect. In the control condition, anti-Rab3A and anti-NSF antibodies were added prior to extracellular calcium in the presence of anti-Epac antibodies; both blocked the AR despite the fact that the end point of the Rap1 branch was achieved by adenophostin (Fig. 6C, gray bars).

8-pCPT-2′-O-Me-cAMP Activates a GEF for Rab3A in Human Sperm—A central aspect of our model is based on the observation that the AR triggered by cAMP (8) and calcium (38) requires Rab3A. We reasoned that if Rab3 serves as a key element in cAMP/Epac-mediated signaling, it must be activated in response to 8-pCPT-2′-O-Me-cAMP. To determine whether Epac exhibits GEF activity toward Rab3A, we incubated 150 nM recombinant Epac1 and Epac2 with fluorescent mGDP-loaded Rab3A (200 nM) in the presence of excess unlabeled GDP. We followed the exchange of guanine nucleotides in real time as a decrease in fluorescence. As shown in Fig. 7, there was no difference between the GDP release rates in the presence of Epacs (squares) and the intrinsic exchange rate for mGDP-Rab3A (black circles) measured...
Calcium was unable to achieve exocytosis in the presence of AR inducers. Calcium served as a positive control because it exchanges GDP for GTP on Rab3A (62). Before challenging with AR inducers, calcium (200 nM) was loaded with the fluorescent nucleotide analogue mGDP and incubated in the presence (squares) or absence (circles) of inactive or activated (500 nM CAMP treated) Rab3A (left) or Epac2 (center) after the addition of a 100-fold excess (20 μM) of unlabeled GDP, as indicated under "Experimental Procedures." For clarity of presentation and to avoid the overlay of symbols, in the left panel the data points of inactive Rab3A and Epac2 were shifted by 0.05 and 0.1 units, respectively. Likewise, in the center panel data points of inactive Rab3A and Epac2 were shifted by 0.05, 0.10, and 0.15 units, respectively. Please note that mGDP can be displaced very slowly from Rab3A (note the difference in time scales between the right panel and the rest) by an excess of GDP; thus the Rab protein is properly folded in a native, nucleotide-interacting state. The right panel demonstrates that the Epac1 (●) and Epac2 (▲) preparations were active and exhibited GEF activity toward Rap. Briefly, Rab1 (200 nM) was loaded with the fluorescent nucleotide analogue mGDP and incubated in the absence (○) or presence of activated (100 μM CAMP treated) Rab3A (●) or Epac2 (▲) after the addition of a 100-fold excess (20 μM) of unlabeled GDP, as indicated under "Experimental Procedures." The data represent the mean ± S.E. of at least two independent experiments.

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![Graph showing the dissociation of the Rab3A-mGDP complex monitored by a time-dependent decrease in fluorescence intensity.](graph)

* For clarity of presentation the data points of inactive Rab3A and Epac2 were shifted by 0.05 and 0.10 units, respectively.

**FIGURE 7. Epac is not a GEF for Rab3A.** The dissociation of the Rab3A-mGDP complex was monitored by a time-dependent decrease in fluorescence intensity. Rab3A (200 nM) was loaded with the fluorescent nucleotide analogue mGDP and incubated in the presence (squares) or absence (circles) of inactive or activated (500 nM CAMP treated; □) Rab3A (left) or Epac2 (center) after the addition of a 100-fold excess (20 μM) of unlabeled GDP, as indicated under "Experimental Procedures." For clarity of presentation and to avoid the overlay of symbols, in the left panel the data points of inactive Rab3A and Epac2 were shifted by 0.05 and 0.1 units, respectively. Likewise, in the center panel data points of inactive Rab3A and Epac2 were shifted by 0.05, 0.10, and 0.15 units, respectively. Please note that mGDP can be displaced very slowly from Rab3A (note the difference in time scales between the right panel and the rest) by an excess of GDP; thus the Rab protein is properly folded in a native, nucleotide-interacting state. The right panel demonstrates that the Epac1 (●) and Epac2 (▲) preparations were active and exhibited GEF activity toward Rap. Briefly, Rab1 (200 nM) was loaded with the fluorescent nucleotide analogue mGDP and incubated in the absence (○) or presence of activated (100 μM CAMP treated) Rab3A (●) or Epac2 (▲) after the addition of a 100-fold excess (20 μM) of unlabeled GDP, as indicated under "Experimental Procedures." The data represent the mean ± S.E. of at least two independent experiments.
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Epac proteins act as GEFs for Rap GTPases but not for Rab3, we next asked whether Epac-mediated Rab3 activation involved a signaling pathway downstream of Rap activation. To address this issue, we introduced anti-Rap antibodies into SLO-permeabilized human sperm before challenging with 8-pCPT-2‘-O-Me-cAMP. As shown in Fig. 8C (right), the antibodies had no effect on 8-pCPT-2’-O-Me-cAMP–induced Rab3 activation, although they potently inhibited Rap signaling during the AR (Fig. 2B). Taken together, these data suggest that Epac plays a central role in cAMP-dependent Rab3 activation but that Rap activity is not essential for this signaling cascade. In other words, upon the onset of exocytosis, Epac activates a GEF for Rab3 through a still undefined mechanism.

**DISCUSSION**

Epac proteins exert diverse effects on a variety of cellular functions and are thought to mediate many of the PKA-independent effects of cAMP-regulated signaling (5, 9–11). These functions include the modulation of secretory processes in pancreatic β-cells (39, 42, 64–66), kidney collecting duct (67), melanotrophs (68), neurons (see Ref. 69 and references therein), and granulosa cells (70). We owe most of our understanding of the molecular mechanisms that couple cAMP/Epac to exocytosis to recent work conducted in insulin-secreting cells (reviewed in Refs. 5, 10). In these cells, the activation of Epac by cAMP sensitizes intracellular (ryanodine and IP$_3$ receptors) calcium release channels, increasing cytosolic calcium concentrations and, subsequently, insulin secretion (42, 71). Direct sensitization of IP$_3$ receptors to IP$_3$ by cAMP has very recently been described in HEK-PR1 cells (72).

Epac is a GEF for Rap proteins. Rap1 belongs to the Ras subfamily of small GTP-binding proteins, which controls cell growth, differentiation, and survival. Although the best known Ras-independent Rap1 function is the control of adhesion-related events (73–75), there is evidence suggesting its involvement in insulin (76, 77) and amylase release (78). Until not too long ago, however, the reported protein-protein interactions linking Epac to insulin secretion did not include Rap (40, 66).

We now know that Rap1 is required for cAMP/Epac-dependent potentiation of insulin secretion (79), pancreatic amylase release (80), and a nonamyloidogenic soluble form of the amyloid precursor protein release (81). Epac proteins have been detected in mouse (82), hamster (83), and human sperm (8), as well as in mouse spermatids (84).

In human sperm, calcium triggers exocytosis in a PKA-independent manner through a pathway involving cAMP and Epac. By activating Epac, cAMP drives the whole cascade of events necessary to bring exocytosis to completion. These events include sequential protein-protein interactions as well as the mobilization of calcium from an intracellular store (4).

**FIGURE 8. Activation of Epac by 8-pCPT-2’-O-Me-cAMP stimulates a GEF for Rab3A during sperm exocytosis.** A, permeabilized spermatozoa were incubated for 10 min at 37 °C with 4 μM GDI (in all the experiments using GDI, the buffer was supplemented with 0.7 mM MgCl$_2$ and 0.2 mM GTP) followed or not by 300 nM Rab3A-GDP and further incubation for 10 min at 37 °C. Acrosomal exocytosis was initiated by adding 50 μM 8-pCPT-2’-O-Me-cAMP and incubating for an additional 10 min at 37 °C (black bars). Controls (gray bars) included the following background AR in the absence of any stimulation (control), AR stimulated by 0.5 mM CaCl$_2$ (calcium) and 50 μM 8-pCPT-2’-O-Me-cAMP (8pCPT), and lack of effect on the AR of 300 nM Rab3A-GDP and 5 μM GDI plus 300 nM Rab3A-GDP. Sperm were fixed; acrosomal exocytosis was evaluated by FITC-PSA binding, and data were normalized (mean ± S.E.) as described under “Experimental Procedures.” B, SLO-permeabilized spermatozoa were incubated for 10 min at 37 °C with 5 μg/ml RIM-(11–398) followed or not by 300 nM Rab3A-GDP and further incubation for 10 min at 37 °C. Acrosomal exocytosis was initiated by adding 0.5 mM CaCl$_2$ or 50 μM 8-pCPT-2’-O-Me-cAMP (8pCPT) and incubating for an additional 10 min at 37 °C (black bars). Controls (gray bars) included the following background AR in the absence of any stimulation (control), AR stimulated by 0.5 mM CaCl$_2$ (calcium) and 50 μM 8-pCPT-2’-O-Me-cAMP (8pCPT), and lack of effect on the AR of 300 nM Rab3A-GDP. Samples were processed, and the AR was scored as in A. C, capacitated human sperm were permeabilized with SLO, and the onset of the AR was prevented by 100 μM 2-APB. Sperm suspensions were subsequently incubated with (right) or without 20 μg/ml anti-Rap1 antibodies or with 1 μg of His$_6$-Rab3A (top left). AR inducers 50 μM 8-pCPT-2’-O-Me-cAMP (top left and right) and 0.5 mM CaCl$_2$ (bottom left) were added and incubations proceeded as described under “Experimental Procedures.” Whole sperm lysates were subjected to pulldown assay using GST-RIM-Sepharose, and the levels of GTP-bound Rab3A were determined by Western blot with an anti-Rab3A antibody. Shown are experiments representative of three repetitions; right, quantification (carried out with Image J, freeware from National Institutes of Health) is depicted above the immunoblot as mean ± S.E. from all replicates; different letters indicate statistical significance (p < 0.05).
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![Working model for the biochemical cascade leading to the AR.](image)

The diagram illustrates the biochemical cascade that leads to the acrosomal reaction (AR). Calcium enters the cell from the extracellular milieu when channels open in the plasma membrane (PM) in response to progesterone or other physiological inducers, when a calcium ionophore transports it to the cytosol, or through the SLO-generated pores. Cyclic AMP synthesized by sAC (or exogenously added 8-pCPT-2′-O-Me-cAMP) activates Epac, which directly activates Rap1. In turn, active Rap1 stimulates the synthesis of IP3 through the stimulation of a PLC activity. Direct activation of Rap1 by Epac and the intracellular calcium pathway proceeds as described. This IP3 (or exogenously added adenophostin) elicits the efflux of calcium from an IP3-sensitive store (likely the acrosome). A reaction taking place during or as a consequence of tethering recruit RIM to achieve tethering. The large quantities of RIM recruited by recombinant Rab3A-GTP supports the assembly of large macromolecular complexes. A reaction taking place during or as a consequence of tethering recruit RIM to achieve tethering. The large quantities of RIM recruited by recombinant Rab3A-GTP supports the assembly of large macromolecular complexes.

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shown that the acrosome behaves as a calcium-storing organelle (59, 60). Release of intra-vesicular calcium takes place in response to AR inducers and is necessary for human sperm exocytosis (59). By using a photosensitive calcium chelator in combination with inhibitory antibodies and a Rap1-GTP sequestering cassette, we have determined that Rap1 is required before the release of calcium from the acrosome (Fig. 3A). Furthermore, this release is the end point of the pathway driven by Rap1, because adenophostin, by eliciting an efflux of intracellular calcium through IP3-sensitive channels, can bypass Rap1 blockers and rescue the AR (Fig. 3C).

A link between cAMP/Epac and calcium signaling implicating Rap2B and a PLCγ activity in HEK-293 and neuroblastoma cells was reported a few years ago (23). More recently we learned that the pathway Epac-Rap1-PLCγ enhances CICR from the sarcoplasmic reticulum in ventricular cardiac myocytes (24). PLCβ (87), PLCγ (88), PLCδ (89), and PLCζ (90) have been described in mammalian sperm (but not PLCε so far). An active PI-PLC operates downstream of Epac during the human sperm AR, because 8-pCPT-2′-O-Me-cAMP could not elicit exocytosis (Fig. 3B) or mobilize calcium (Fig. 4B) in the presence of U73122. A PI-PLC activity is also required by calcium (Fig. 3B) and Rab3A-GTPγ-S (Fig. 5B) to achieve the AR. We believe this PI-PLC, whatever its molecular identity might be, is responsible for the synthesis of the IP3 that promotes efflux of calcium from an intracellular store in sperm, because adenophostin can rescue the block imposed by U73122 on the AR (Fig. 3C).

On reviewing the literature, we became aware of the existence of systems where sAC synthesizes the cAMP that culminates in a Rap1-activating cascade (91, 92). This calcium- and bicarbonate-stimulated cyclase is the predominant form in mammalian sperm (44, 56, 93, 94) and is critical for fertility (95, 96). It was therefore not surprising to discover that sAC was not needed for human sperm exocytosis (44). The calcium- and bicarbonate-stimulated cyclase is the predominant form in mammalian sperm (44, 56, 93, 94) and is critical for fertility (95, 96).

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Interplay between calcium and cAMP in human sperm appears to be different because cAMP achieves exocytosis in the presence of high concentrations of EGTA, which chelates extracellular calcium, preventing an influx from the outside, but not of low concentrations of 1,2-bis(2-amino-phenoxy)ethane-N,N,N′,N′-tetraacetic acid-acetoxymethyl ester, which chelates intracellular calcium, preventing mobilization from stores (8). Interestingly, cAMP-elevating agents such as GLP-1 stimulate the release of calcium stored in insulin-containing secretory granules (85). In this study we show that cAMP/Epac activate intracellular mobilization in human sperm via a pathway involving Rap1 and a PLC. Briefly, Rap1 was present and required for regulated exocytosis in human sperm (Fig. 2, A and B). Three AR inducers, an Epac-selective cAMP analogue, recombinant Rab3A-GTPγ-S, and calcium, promoted the activation of Rap1 in human sperm (Fig. 2, C and D, and Fig. 5B). Likewise, 8-pCPT-2′-O-Me-cAMP activates Rap1 in mouse germ cells (82, 84). The importance of Rap1 in male fertility has been highlighted by the discovery that transgenic mice expressing an inactive mutant in differentiating spermatids are severely subfertile (86). We and others have determined that Rap1 is required before the release of calcium from the acrosome (Fig. 3A). Furthermore, this release is the end point of the pathway driven by Rap1, because adenophostin, by eliciting an efflux of intracellular calcium through IP3-sensitive channels, can bypass Rap1 blockers and rescue the AR (Fig. 3C).

A link between cAMP/Epac and calcium signaling implicating Rap2B and a PLCγ activity in HEK-293 and neuroblastoma cells was reported a few years ago (23). More recently we learned that the pathway Epac-Rap1-PLCγ enhances CICR from the sarcoplasmic reticulum in ventricular cardiac myocytes (24). PLCβ (87), PLCγ (88), PLCδ (89), and PLCζ (90) have been described in mammalian sperm (but not PLCε so far). An active PI-PLC operates downstream of Epac during the human sperm AR, because 8-pCPT-2′-O-Me-cAMP could not elicit exocytosis (Fig. 3B) or mobilize calcium (Fig. 4B) in the presence of U73122. A PI-PLC activity is also required by calcium (Fig. 3B) and Rab3A-GTPγ-S (Fig. 5B) to achieve the AR. We believe this PI-PLC, whatever its molecular identity might be, is responsible for the synthesis of the IP3 that promotes efflux of calcium from an intracellular store in sperm, because adenophostin can rescue the block imposed by U73122 on the AR (Fig. 3C).

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E. Teves and L. Giojalas, personal communication.
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The AR was independent of PKA activity (Fig. 5). Rab3A-GTP-activation, the AR inducers calcium, cAMP, and recombinant AR by themselves. In addition to intracellular calcium mobilization, the identity of this protein(s) is presently unknown. Protein(s) is necessary to couple Epac activation to Rab3A signaling. The same was true for PDE, KH7, anti-Rab3A antibodies, PTP1B D181A, and NSF Y83E (Fig. 6A and B). We could conceivably fit some of these data into a linear model whereby cAMP/Epac promoted a release of calcium from an intracellular store prior to (and not in parallel with) the tethering, priming, and docking steps. The results depicted in Fig. 6C go against this model because they suggest that, even while maintaining the intracellular calcium mobilization pathway artificially off with anti-Epac antibodies, calcium assembles the fusion machinery into place before anti-Rab3A and anti-NSF antibodies have the opportunity to prevent it.

Therefore, we suggest that the AR is organized as a bifurcated pathway, with two separate limbs that diverge downstream of cAMP/Epac (Fig. 9). The end point of one limb (Rap-PLC) is the mobilization of intracellular calcium, whereas the other (Rab3Aα-SNAP/NSF-SNAREs) assembles the fusion protein machinery so that the outer acrosomal and plasma membranes become physically connected. Both pathways operate in a concerted fashion and converge at or downstream of intracellular calcium mobilization to accomplish exocytosis; once calcium is released into the cytosol, the protein machinery, likely through synaptotagmin (48), senses the rise in concentration, and the whole system evolves to open the fusion pores.

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