Distinct purinergic signaling pathways in prepubescent mouse spermatogonia

David Fleck,1 Nadine Mundt,1 Felicitas Bruentgens,1 Petra Geilenkirchen,2 Patricia A. Machado,1 Thomas Veitinger,1 Sophie Veitinger,1 Susanne M. Lipartowski,1 Corinna H. Engelhardt,1 Marco Oldiges,2 Jennifer Spehr,1 and Marc Spehr1

1Department of Chemosensation, Institute for Biology II, RWTH Aachen University, D-52074 Aachen, Germany
2Institute of Bio- and Geosciences (IBG), IBG-1: Biotechnology, Research Center Jülich, D-52425 Jülich, Germany

Spermatogenesis ranks among the most complex, yet least understood, developmental processes. The physiological principles that control male germ cell development in mammals are notoriously difficult to unravel, given the intricate anatomy and complex endo- and paracrinology of the testis. Accordingly, we lack a conceptual understanding of the basic signaling mechanisms within the testis, which control the seminiferous epithelial cycle and thus govern spermatogenesis. Here, we address paracrine signal transduction in undifferentiated male germ cells from an electrophysiological perspective. We identify distinct purinergic signaling pathways in prepubescent mouse spermatogonia, both in vitro and in situ. ATP—a dynamic, widespread, and evolutionary conserved mediator of cell to cell communication in various developmental contexts—activates at least two different spermatogonial purinoceptor isoforms. Both receptors operate within nonoverlapping stimulus concentration ranges, display distinct response kinetics and, in the juvenile seminiferous cord, are uniquely expressed in spermatogonia. We further find that spermatogonia express Ca2+-activated large-conductance K+ channels that appear to function as a safeguard against prolonged ATP-dependent depolarization. Quantitative purine measurements additionally suggest testicular ATP-induced ATP release, a mechanism that could increase the paracrine radius of initially localized signaling events. Moreover, we establish a novel seminiferous tubule slice preparation that allows targeted electrophysiological recordings from identified testicular cell types in an intact epithelial environment. This unique approach not only confirms our in vitro findings, but also supports the notion of purinergic signaling during the early stages of spermatogenesis.

INTRODUCTION

Spermatogenesis ranks among the most complex, yet least understood, developmental processes in postnatal life. Initiated 5–7 d postpartum in rodents (Kolas et al., 2012), this intricate course of mass cell proliferation and transformation events generates fertile haploid spermatozoa from diploid spermatogonial stem cells (SSCs). The seminiferous tubule represents the functional unit of the testis. Along its epithelium, spermatogenesis has been simplified morphologically by attribution of sequential cellular stages, which progress through coordinated and precisely timed cycles (Hess and de Franca, 2008). However, prepubescent immature seminiferous tubules/cords are exclusively built by three cell types, i.e., Sertoli cells, peritubular cells, and spermatogonia (Bellvé et al., 1977). Although difficult to discriminate both morphologically and molecularly (Jan et al., 2012), premeiotic germ cells of the spermatogonial lineage comprise a heterogeneous population (Chiarini-Garcia and Russell, 2002), including SSCs, proliferating Apaired and Aaligned cells, and differentiating A1-A4, intermediate and B spermatogonia (Kolas et al., 2012). Both Sertoli and germ cells have developed elaborate, yet ill-defined mechanisms of functional communication (Cheng and Mruk, 2002). Multidirectional interactions among germ cells as well as between germ and somatic cells balance SSC self-renewal and differentiation, synchronize stage transitions, regulate blood–testis barrier dynamics, and control epithelial cyclicity via autocrine, paracrine, and endocrine feedback (Heindel and Treinen, 1989). Precisely regulated cellular communication within the seminiferous epithelium is thus imperative for spermatogenesis and reproduction.

In numerous developmental processes, purinergic signaling is emerging as a critical component of paracrine communication networks (Abbracchio et al., 2009; Praetorius and Leipziger, 2009). As a widespread and evolutionary conserved route for cell to cell interactions, extracellular ATP targets members of the P2 purinoceptor family (Burnstock, 1990). P2 receptors divide into two distinct classes: metabotropic P2Y (Barnard et al., 1994) and ionotropic P2X receptors (Bean...
Ca²⁺-activated large conductance (BK) K⁺ channels of both P2X4 and P2X7 receptor isoforms as well as multidimensional ATP response pathway that consists pubescent mice both in vitro and in situ. We identify a target signaling mechanisms in spermatogonia from pre-meiotic germ cells is currently unclear. However, whether spermatogonia are also targets of paracrine purinergic signaling and, if so, which molecular machinery mediates ATP sensitivity in premeiotic germ cells is currently unclear.

Here, we used gene expression analysis, immuno- and bioanalytical chemistry, protein knockdown, and single cell electrophysiology to investigate purinoceptor signaling mechanisms in spermatogonia from prepubescent mice both in vitro and in situ. We identify a multidimensional ATP response pathway that consists of both P2X4 and P2X7 receptor isoforms as well as Ca²⁺-activated large conductance (BK) K⁺ channels. This channel profile endows spermatogonia with distinct and dynamic response features over a broad range of stimulus concentrations. Cooperatively activated by depolarization and increased cytoplasmic Ca²⁺ (Fakler and Adelman, 2008), BK channels, in turn, provide a negative feedback mechanism that counteracts the electrophysiological effects of P2X receptor activation. In addition, P2X signaling itself acts as a regulator of testicular ATP release, providing a parallel positive feedback pathway. Consistency of results from primary cell culture and an intact acute seminiferous tubule slice preparation substantiates the physiological relevance of our findings.

MATERIALS AND METHODS

Animals

All animal procedures were approved by local authorities and in compliance with European Union legislation (Directive 86/609/EEC) and recommendations by the Federation of European Laboratory Animal Science Associations (FELASA). C57BL/6 mice (Charles River) were housed in groups of both sexes (room temperature [RT]; 12:12 h light–dark cycle; food and water available ad libitum). If not stated otherwise, experiments used 7-d-old males.

Chemicals and solutions

The following solutions were used: (S1) HEPES-buffered extracellular solution containing (mM) 145 NaCl, 5 KCl, 1 CaCl₂, 0.5 MgCl₂, and 10 HEPES; pH = 7.3 (adjusted with NaOH); osmolarity = 300 mOsm (adjusted with glucose). (S₂) Oxygenated (95% O₂, 5% CO₂) extracellular solution containing (mM) 120 NaCl, 25 NaHCO₃, 5 KCl, 0.5 MgCl₂, 1.0 CaCl₂, 5 N,N-bis(2-hydroxyethyl)-2-aminoethanesulfonic acid (BES), and 10 glucose; pH = 7.3 (NaOH); 300 mOsm (glucose). (S₃) Extracellular low Ca²⁺ solution containing (mM) 145 NaCl, 5 KCl, 2.5 CaCl₂, 0.5 MgCl₂, 10 HEPES, and 5 EGTA; pH = 7.3 (NaOH); osmolarity = 300 mOsm (glucose), [Ca²⁺]free = ~110 nM. (S₄) Extracellular TEA solution containing (mM) 120 NaCl, 15 TEACl, 5 KCl, 1 CaCl₂, 0.5 MgCl₂, and 10 HEPES; pH = 7.3 (NaOH); osmolarity = 300 mOsm (glucose). (S₅) Standard pipette solution containing (mM) 143 KCl, 2 KOH, 1 EGTA, 0.3 CaCl₂, 10 HEPES, and 1 Na-GTP ([Ca²⁺]free = 110 nM); pH = 7.1 (adjusted with KOH); osmolarity = 290 mOsm (glucose). (S₆) Cs⁺-based pipette solution containing (mM) 143 CsCl, 2 CsOH, 1 EGTA, 0.3 CaCl₂, 10 HEPES, and 1 Na-GTP ([Ca²⁺]free = 110 nM); pH = 7.1 (adjusted with CsOH); osmolarity = 290 mOsm (glucose). (S₇) Standard block solution containing 5% (anti-P2X4, anti-P2X7, anti-Slo1) or 10% (anti-DAZL) normal goat serum (Thermo Fisher Scientific), 10 mg/ml BSA, 0.3% Triton X-100, and 0.02% NaN₃ in PBS−/- (100 mM). (S₈) Cell culture blocking solution containing 1% BSA and 0.1% Triton X-100 in PBS−/- (100 mM). (S₉) Standard washing solution containing 10 mg/ml BSA in PBS−/- (100 mM). (S₁₀) Standard staining solution containing 3% BSA (IgG free, protease free), 0.05% NaN₃, and Alexa Fluor 488 or 633 streptavidin conjugate (1:800; Thermo Fisher Scientific) in PBS−/- (100 mM). (S₁₁) Standard co-culture medium (adopted and modified from Iwanami et al. [2006]) consisting of DMEM (low glucose, pyruvate; Thermo Fisher Scientific) complemented with 10 ng/ml epidermal growth factor, 10% fetal calf serum (Thermo Fisher Scientific), 1 ng/ml FSH, 0.2 ng/ml growth hormone releasing factor, 5 µg/ml insulin, 10 ng/ml insulin-like growth factor, 1% MEM nonessential amino acid solution (Thermo Fisher Scientific).
nucleoside solution, 100 U/ml penicillin, 100 µg/ml streptomycin, 5 µM retinol acetate, 0.5 mM sodium pyruvate, 100 nM testosterone, and 5 µg/ml transferrin.

Free Ca$^{2+}$ concentrations were calculated using WEBMAXC Extended. If not stated otherwise, chemicals were purchased from Sigma-Aldrich. Alexa Fluor hydrazide was purchased from Thermo Fisher Scientific; 3-[[5-(2,3-Dichlorophenyl)-1H-tetrazol-1-yl]methyl]pyridine hydrochloride (AOPCP), and 1-Amino-4-(4-chlorophenyl)aminoantraquinone-2-sulfonic acid sodium salt (PSB 069) were purchased from Tocris Bioscience; and iberiotoxin was purchased from Abcam. PSB-POM144 (Lee et al., 2015) was provided by C.E. Müller (University of Bonn, Bonn, Germany). Final solvent concentrations were ≤0.1%. When high ATP concentrations (≥1 mM) were used, pH was readjusted. Solutions and pharmacological agents were applied either by the bath or from air pressure–driven reservoirs via an 8-in-1 multi-barrel "perfusion pencil" (Science Products). Changes in focal superfusion (Veitinger et al., 2011) were software controlled and, if required, synchronized with data acquisition by TTL input to 12V DC solenoid valves using a TIB 14S digital output trigger interface (HEKA).

Slice preparation
Male mice were sacrificed on postnatal day 7 (P7) by decapitation with sharp surgical scissors. Testes were rapidly dissected, each tunica albuginea was removed, and seminiferous tubules were embedded in 4% low-temperature agarose (Aagar) and placed in oxygenated S2. 200-µm slices were cut with a VT1000S vibratome (speed: 0.15 mm/s; frequency: 65 Hz; amplitude: 1 mm; Leica Biosystems) and transferred to a submerged, oxygenated storage container (S2; RT) until use.

Cell culture
After dissection of both testes on P7 and removal of the tunica albuginea, the seminiferous tubules from juvenile/prepubescent mice were placed in MEM (Thermo Fisher Scientific) containing 1 mg/ml collagenase (8 min; 37°C). Digestion was stopped by the addition of serum-containing DMEM (Thermo Fisher Scientific). The suspension was centrifuged (8 min; 400 g) and the supernatant discarded. The pellet was resuspended in 0.05% Trypsin-EDTA (Thermo Fisher Scientific), incubated (5 min, 37°C), and centrifuged (10 min; 400 g). After discarding the supernatant, the pellet was resuspended in culture medium (S11). Cells were plated on glass coverslips in 35-mm dishes at densities of ∼2.5 × 10^5 cells per dish and placed in a humidified incubator for 3 d (37°C; 5% CO2). On the fourth day in vitro (DIV4), incubation temperature was reduced (34°C; 5% CO2). After addition of 1 ml S11 on DIV1, one-third of the medium was replaced every day. Experiments were performed from DIV4 to DIV7.

Gene expression analysis
Total RNA from cultured spermatogonia, testis, brain, and spinal cord was isolated using the RNasy Mini kit (QIAGEN) according to the manufacturer’s instructions. cDNA was transcribed using RevertAid H Minus M-MuLV reverse transcription (Fermentas) according to the manufacturer’s instructions. Controls in which the reverse transcription step was omitted were routinely performed. PCR amplification was performed during 30 thermal cycles (94°C, 20 s; 52–63°C, 20 s; 72°C, 20 s). To confirm the anticipated size of the product, PCR products were visualized on an agarose gel via GelRed (VWR) staining. The specific primer pairs used for PCR amplification are listed in Table 1.

For qPCR, we used the QuantiFast SYBR Green PCR kit (QIAGEN) and an iQ5 thermal cycler (Bio-Rad Laboratories) according to the manufacturer’s instructions. α1A-tubulin was used to normalize the amount of target mRNA in the different cDNA populations tested. Relative mRNA levels were then calculated using the threshold cycle (Ct) value and the comparative Ct method. The primer pairs used for qPCR are listed in Table 2.

### Table 1. Specific primer pairs used for PCR amplification

<table>
<thead>
<tr>
<th>Target</th>
<th>Forward primer 5'-3'</th>
<th>Reverse primer 5'-3'</th>
<th>Size</th>
</tr>
</thead>
<tbody>
<tr>
<td>P2X1</td>
<td>GAGACTGCGGCACGAGCTTC</td>
<td>GCGAATCCAAAGCACCTGGA</td>
<td>bp</td>
</tr>
<tr>
<td>P2X2</td>
<td>TCCCTCCCCACACATGCAC</td>
<td>CACCACTGCTCAAGACGC</td>
<td>233</td>
</tr>
<tr>
<td>P2X3</td>
<td>GTCTCAACTCATCGCAAGAG</td>
<td>AATACCCAGACGCCACCC</td>
<td>149</td>
</tr>
<tr>
<td>P2X4</td>
<td>CCATTGTGGCCCATGATAT</td>
<td>CGTACCCTTTGATGAGTGT</td>
<td>150</td>
</tr>
<tr>
<td>P2X5</td>
<td>GGAAGATATCTGAGTGGTGA</td>
<td>TCTGACCAACTTCTCCAGT</td>
<td>145</td>
</tr>
<tr>
<td>P2X5b</td>
<td>GCTGGCTCCACTCGCAACC</td>
<td>ACCGCCCAAGACCATGAC</td>
<td>81</td>
</tr>
<tr>
<td>P2X6</td>
<td>TCCAGAACTCCCTCTGCTCC</td>
<td>GGGCAGCTTTCCAGTCTCA</td>
<td>253</td>
</tr>
<tr>
<td>P2X7</td>
<td>GACGAGGATATGGCCACCTGC</td>
<td>CCCACCCCTTGTACCTACTC</td>
<td>152</td>
</tr>
<tr>
<td>P2X7αex</td>
<td>GACCTCATCGGGGTCGGCT</td>
<td>ATGTCCCTGGGAGCCAGGCC</td>
<td>171</td>
</tr>
<tr>
<td>P2X7βex</td>
<td>GTCGCCCTAGCGGAGCAAGCG</td>
<td>TGGGTCCTGGTGATGAGTGC</td>
<td>116</td>
</tr>
<tr>
<td>β-actin</td>
<td>GTCTCCTCCCTCCATCTGCGG</td>
<td>TGGATGCACCGAGATC</td>
<td>237</td>
</tr>
</tbody>
</table>

(bp) 233

Table 2. Primer pairs used for qPCR

<table>
<thead>
<tr>
<th>Target</th>
<th>Forward primer 5'-3'</th>
<th>Reverse primer 5'-3'</th>
<th>Size</th>
</tr>
</thead>
<tbody>
<tr>
<td>P2X4</td>
<td>TCCGCGTACGCATATTGGG</td>
<td>AGAAGTTGGTGGCACCCCA</td>
<td>116</td>
</tr>
<tr>
<td>P2X7</td>
<td>CCCAGATGGACTTCCGAC</td>
<td>GCACCTAGGGCCCACCTTCTT</td>
<td>116</td>
</tr>
<tr>
<td>αtubulin</td>
<td>TCCCAAGATGCTAATGCTG</td>
<td>CACAGGGAGGCTGTAAT</td>
<td>115</td>
</tr>
</tbody>
</table>

Protein knockdown
For small interfering RNA (siRNA) experiments, either Sertoli cells or spermatogonia were transfected 12–24 h before experiments using Lipofectamine 2000 (Thermo Fisher Scientific) according to the manufacturer’s instructions. Before transfection, the culture medium was replaced with antibiotic-free medium. siRNA (Applied Biosystems/Ambion) was cotransfected with BLOCK iT fluorescent oligo (FITC labeled; Thermo Fisher Scientific). Opti-MEM I Reduced Serum Medium (500 µl; Thermo Fisher Scientific) containing 6 µl Lipofectamine 2000, 502.5 ng siRNA (75 nM), and 1.875 µl BLOCK iT fluorescent oligo (75 nM) was added to each culture dish. Transfected cells were identified by FITC fluorescence. We used the following Silencer Select Pre-designed siRNA constructs to down-regulate receptor protein expression: P2X4 RNAi1, #s71184; P2X4 RNAi2, #s71185; P2X2 RNAi1, #s106892; P2X2 RNAi2, #s106893; and P2X7 RNAi, #s71189. As negative control, we used Silencer Select negative control #1 and #2 siRNA.

LC-MS/MS analysis
Seminiferous tissue samples were dissected as described above (Slice preparation). Individual sample weight was recorded for post hoc normalization. Samples and controls were exposed to either a defined concentration of ATP (10 µM), 2′-(3′)-O-(4-Benzoylbenzoyl) ATP (BzATP; 10 µM), or volume-matched physiological saline (S1) for 5 min. Next, samples were centrifuged (3 min; 4,000 g; 4°C) and filtered (10 min; 17,000 g; 4°C). All protocol steps were identical regardless of the sample/treatment type. Before analysis, samples were again filtered (10-kD cutoff; Centrifugal Devices NANOSEP).

Ion-pairing reverse-phase column chromatography (C18 Synergy hydro; Phenomenex) was used with eluent A (10 mM tributyramine aqueous solution and 5% methanol adjusted pH to 4.95 with 15 mM acetic acid) and eluent B (methanol) at 40°C. The following elution gradient was used: 2 min (100% A), 5 min (80% A), 8 min (80% A), 10 min (65% A), 14 min (0% A), 15 min (0% A), 15.5 min (100% A), and 18 min (100% A). The flow rate was set to 0.45 ml/min with 10-µl injection volume. Quantification was performed by isotope dilution mass spectrometry (Wu et al., 2005), adding 13C-labeled internal standard obtained from batch cultivation of Corynebacterium glutamicum with uniformly 13C-labeled glucose as the sole carbon source.

LC-MS/MS was carefully checked for cross-talk or false-positive detection of ATP originating from BzATP that was added in some experiments. For BzATP, stronger chromatographic retention was found showing clear baseline separation from ATP. Moreover, BzATP shows a unique MS/MS fragment pattern clearly separated from ATP with no interfering fragments; i.e., BzATP does not contaminate the mass trace of ATP or ATP fragments. However, the BzATP standard was found to contain ≤4% impurity of ATP, traces of ADP (below quantification limit), and no AMP. Measurement results were corrected accordingly, assuming maximum impurity.

Histology and immunocytochemistry
For immunocytochemistry of testicular cryosections, testes were fixed with 4% (wt/vol) paraformaldehyde (PFA) in Ca2+–Mg2+–free PBS−/− (10 mM, pH 7.4; ≥3 h for juvenile [P7] and ≥12 h for adult tissue; 4°C) and subsequently cryoprotected in PBS−/− containing 30% sucrose (≥24 h; 4°C). Samples were then embedded in Tissue Freezing Medium (Leica Biosystems), sectioned at 20 µm on a CM1950 cryostat (Leica Biosystems), and mounted on Superfrost Plus slides (Menzel). For blocking, sections were incubated in S3 (1 h; RT; gentle agitation). Cryosections were then washed (2 x 5 min; S3) and incubated with primary antibodies overnight (S3; 4°C) in a dark humidified chamber. Primary antibodies were anti-DAZL (1:500; Abcam), anti-Slo1 (extracellular; 1:500; Alomone Labs), anti-P2X4 (extracellular; 1:100, Alomone Labs), and anti-P2X7 (extracellular; 1:500; Alomone Labs). After washing in S3 (5×, 10 min), sections were incubated in S3 containing Alexa Fluor 488 goat anti–rabbit secondary antibody
scanning confocal microscope (TCS SP5 DM6000 CF5; Leica Microsystems) equipped with a 20× 1.0 NA water immersion objective (HCX APO L; Leica Microsystems). To control for nonspecific staining and to demonstrate antibody specificity, we performed (a) antigen preadsorption controls and (b) experiments in which the primary antibodies were omitted in parallel with each procedure. Digital images were uniformly adjusted for brightness and contrast using Photoshop CS6 (Adobe Systems).

For immunoblotting, both testes and control tissues (i.e., brain, olfactory bulb, muscle, and HEK293T cells) were homogenized in lysis buffer (100 µl; 0.1% Triton X-100; 4°C) in the presence of Complete Mini protease inhibitor cocktail tablets (Roche). The homogenate was sonicated (5 s) and centrifuged for 10 min; RT). After incubation, cells were washed/incubated in S8 (2 × 5 min, 1 × 60 min; RT). For nuclear counterstaining, cells were incubated in PBS− containing DAPI (1:100; 15 min; RT; Thermo Fisher Scientific) and washed again in PBS−.

Fluorescent images were taken using an upright fixed stage scanning confocal microscope (TCS SP5 DM6000 DFS; Leica Microsystems) equipped with a 20× 1.0 NA water immersion objective and a three-position magnification changer (HCX IRA 25×/0.95W; Leica Microsystems; slices), respectively. Microscopes were equipped for either phase contrast or infrared-optimized differential interference contrast (IR-DIC) microscopy. Whereas cultured cells were observed using phase-contrast objectives (HCX PL FLUOTAR 20× 0.5 NA, 40× 0.6, and 63× 0.7; Leica Microsystems), acute sections were visualized using a 25× (HCX IRAPO L25×/0.95W) objective and a three-position magnification changer (0.35×, 1.25×, and 4.0×). Images were recorded using a cooled CCD camera (DFC365FX; Leica Microsystems). Samples were continuously superfused with S8 (∼3 ml/min; gravity flow; 23°C). Patch pipettes (5–7 MΩ) were pulled from borosilicate glass capillaries (1.50 mm OD/0.86 mm ID; Science Products) on a PC-10 micropipette puller (Narishige Instruments), fire polished (MF-830 Microforge; Narishige Instruments), and filled with pipette solution (Sf, depending on experimental design). In slice recordings, 20 µM Alexa Fluor 488 hydrazide (Thermo Fisher Scientific) was routinely added to the pipette solution to allow online evaluation of cell morphology and post hoc three-dimensional reconstruction, respectively. An agar bridge (150 mM KCl) connected reference electrode and bath solution. An EPC-10 USB amplifier controlled by Patchmaster 2.67 software (HEKA) was used for data acquisition. We monitored and compensated pipette and membrane capacitance (Cmem) as well as series resistance. Treated, to a first approximation, as a “biological constant” with a value of ∼1 µF/cm² (Gentet et al., 2000), Cmem values served as a proxy for the cell surface area and thus for normalization of current amplitudes (i.e., current density). Liquid junction potentials were calculated using
JCalcW software (Barry, 1994) and corrected online. Signals were low-pass filtered (analogue 3- and 4-pole Bessel filters [−3 dB]; adjusted to 1/4 to 1/5 of the sampling rate [5–20 kHz; depending on protocol]). If not stated otherwise, holding potential (V\text{hold}) was −40 mV. In some voltage-clamp experiments, leak currents were subtracted using a P/6 protocol at hyperpolarized V\text{hold}. For clarity, few exemplary current traces (Figs. 4 B and 9 B) were subsequently processed using a “box-7” smoothing algorithm implemented in IGOR Pro 6.5 software (WaveMetrics). Moreover, electric artifacts resulting from valve switching were removed. All data were recorded at RT. Individual voltage step and ramp protocols are described in the Results section.

Data analysis
All data were obtained from independent experiments performed on at least 2 d. Individual numbers of cells/ experiments (n) are denoted in the figure and/or captions. If not stated otherwise, results are presented as means ± SEM. Statistical analyses were performed using paired or unpaired t tests, one-way ANOVA with Tukey’s HSD post hoc test, or the Fisher exact test (as dictated by data distribution and experimental design). Tests and corresponding p-values that report statistical significance (≤ 0.05) are individually specified in the captions. Data were analyzed offline using FitMaster 2.67 (HEKA), IGOR Pro 6.5 (WaveMetrics), and Excel 2013 (15.0.4779.1001; Microsoft) software. Desensitization rates during successive ramp recordings (Fig. 6, B and D) were determined from linear regression fits to data plotted as current versus time (Fig. 6, C and E) in the presence of ATP and calculated as current reduction (%) over time (s; Fig. 6 G). Dose-response curves were fitted by the Hill equation.

Online supplemental material
Fig. S1 shows immunochemical analysis of channel/receptor protein expression in juvenile seminiferous cords. Fig. S2 shows dose-dependent inhibition of ATP responses by suramin in cultured Sertoli cells. Fig. S3 shows that posttranscriptional P2rx2 gene silencing diminishes Sertoli cell ATP responses. Fig. S4 shows reduced BzATP sensitivity of Sertoli cell P2X receptors. Fig. S5 shows quantitative LC-MS/MS analysis of [ATP]\text{ex} in seminiferous epithelia. Online supplemental material is available at http://www.jgp.org/cgi/content/full/jgp.201611636/DC1.

RESULTS
A Sertoli cell–germ cell co-culture from immature seminiferous tissue maintains spermatogonia for physiological analysis
To investigate purinergic signaling mechanisms in spermatogonia, we first established Sertoli–germ cell co-cultures from testicular tissue of prepubescent mice. Seminiferous tubules were isolated at P7 to prevent culture contamination with meiotic or postmeiotic cell types (Bellvé et al., 1977; Creemers et al., 2002). When testicular tissue at P7 is compared by freeze-fracture electron microscopy with samples from adult animals (Fig. 1, A and B), the relatively homogenous composition of the smaller immature seminiferous cords becomes apparent (Fig. 1 Bi). In corresponding cryosections (Fig. 1, C and D), immunohistochemical labeling of the spermatogonial marker protein DAZL (Deleted in AZoospermia-Like; VanGompel and Xu, 2011) reveals the expected circular staining in the basal compartment of adult seminiferous tubules (Fig. 1 C). In contrast, round DAZL-positive cells are diffusely distributed throughout the seminiferous cords in P7 mice (Fig. 1 D), confirming that most, if not all, germ cells in immature cords are spermatagonia. When Sertoli cell–germ cell co-cultures from immature seminiferous cords were performed according to established protocols (Veitinger et al., 2011; Hunter et al., 2012), apparent A\text{single} and A\text{paired} (Fig. 1 E) as well as larger clones of DAZL-positive A\text{aligned} spermatogonia (Fig. 1 F) resided on a confluent Sertoli cell feeder layer. Essentially all round cells in culture were labeled by the spermatogonial marker. Thus, the present co-culture allows maintenance of a relatively homogenous premeiotic germ cell population in vitro.

Spermatogonia are sensitive to purinergic stimulation over a broad concentration range
To address whether spermatogonia could be part of a paracrine purinergic signaling network (Praetorius and Leipziger, 2009), we performed whole-cell patch-clamp recordings from cultured germ cells (Fig. 2 A). For sample consistency, recordings were restricted to putative A\text{single} and A\text{paired} spermatogonia, which did not differ in any parameter investigated (i.e., both passive and active membrane properties). Spermatogonia exhibited a mean membrane capacitance (C\text{mem}) of 15.8 ± 7.4 pF and an input resistance (R\text{input}) of 1.24 ± 0.9 GΩ (means ± SD). When cells were exposed to 30 µM ATP at a negative holding potential (V\text{hold} = −80 mV), we consistently recorded a fast-activating inward current in 51–78% of all spermatogonia (Fig. 2 A) throughout the experimental timeframe (DIV4–DIV7). Upon prolonged exposure (>1 s), the current saturated and then monotonically declined in the presence of the stimulus (Fig. 2, A and B). This apparent desensitization was then examined in a multi-pulse paradigm (Fig. 2 C). When current amplitudes in response to a second stimulation were normalized to the initial signal, we observed essentially stable response magnitudes at interstimulus intervals (ISIs) >45 s. In contrast, a progressive decline in current amplitude emerged as a
function of reduced ISI. We therefore performed all subsequent recordings at an ISI of ≥1 min.

Next, we asked whether ATP sensitivity is dose dependent. At extracellular ATP concentrations ([ATP]_{ex}) of up to 300 µM, spermatogonia exhibited a sigmoidal dose–response relationship with a threshold [ATP]_{ex} of ∼3 µM and half-maximal activation at 13.5 µM (Fig. 2, D and E). Although we observed apparent saturation at [ATP]_{ex} ≥100 µM, massively increased stimulus concentrations (≥1 mM) seemed to recruit a second low-affinity response mechanism. Notably, currents activated by these large concentrations lacked signs of desensitization (Fig. 2 D), supporting the notion of two independent signaling mechanisms that operate at opposite ends of the dose–response spectrum.

To examine the current–voltage relationship of “high-affinity” ATP-induced currents, we next challenged spermatogonia with 30 µM ATP at various membrane potentials (Fig. 2 F). Immediately activated currents reversed at ∼0 mV and exhibited pronounced inward rectification (Fig. 2 G). Surprisingly, we observed an additional outward current that developed with considerable delay (Fig. 2 F) and reversed at approximately −80 mV (Fig. 2 G).

Together, our data reveal that cultured mouse spermatogonia express a distinct set of receptors and channels that confer both sensitivity and functional specificity in response to a universal auto/paracrine signal (i.e., ATP).

BK channels balance electrophysiological effects of ATP stimulation

Given the negative reversal potential (E_{rev}) of the ATP-dependent delayed conductance (and because measurements were conducted in symmetric Cl⁻ solutions), we hypothesized that this delayed current (I_{del}) is largely carried by K⁺. To test this, we again recorded current–voltage curves in response to ATP (10 µM), replacing intracellular K⁺ with Cs⁺ (S6; Fig. 3 A). Under these conditions, the immature seminiferous cords exhibit a more homogenous cellular composition devoid of a lumen. (C and D) Confocal fluorescence images of immunostainings against DAZL, a marker of premeiotic germ cells. Cryosections from adult (C) and prepubescent (D) mice reveal DAZL-positive cells in the periphery of the adult seminiferous epithelium (C), whereas spermatogonia are scattered throughout the immature seminiferous cords (D) and appear to make up the only spherical seminiferous cell type at P7, as visualized in merged fluorescence and DIC micrographs (Ci and Di). (E and F) Scanning electron microscopy (E) and confocal dual-channel fluorescence (F) images of Sertoli cell–germ cell co-cultures from seminiferous cords of prepubescent mice (P7). Relatively small spherical cells reside on a confluent flat layer of large Sertoli cells (E). Immunostaining against DAZL (Fi) and nuclear counterstaining (DAPI; Fii) reveals spermatogonial marker expression in essentially all spherical cells. Note that cultured putative spermatogonia are found as single and paired cells (E) or as aligned groups of up to 16 cells (F).
conditions, instantaneous ATP-mediated currents ($I_{\text{inst}}$) were still present in 40% of spermatogonia and appeared similar to control recordings (Fig. 2 G). In contrast, $I_{\text{del}}$ was absent (Fig. 3, A–C), suggesting that this current is largely carried by K⁺.

To identify the molecular correlate of $I_{\text{del}}$, we next determined the current's pharmacological profile (Fig. 3, D and E). First, 15 mM TEA strongly inhibited $I_{\text{del}}$. Second, $I_{\text{del}}$ was essentially blocked when extracellular Ca²⁺ was reduced (110 nM), whereas $I_{\text{inst}}$ was significantly increased under these conditions. Both Ca²⁺ dependence and TEA sensitivity are hallmarks of Ca²⁺-activated BK channels (Marty, 1981; Pallotta et al., 1981; Yellen, 1984). If BK (Slo1) channels mediate $I_{\text{del}}$, the current should be blocked by the selective BK channel inhibitor iberiotoxin (Galvez et al., 1990). Indeed, iberiotoxin strongly reduced $I_{\text{del}}$ without affecting $I_{\text{inst}}$. Moreover, an antibody against an extracellular epitope of the BK channel α₁ subunit (anti-Slo1) specifically labeled spermatogonia-like cells in cryosections from P7 testes (Fig. S1, A–D). Our data thus indicate that, upon ATP exposure, spermatogonial BK channels are cooperatively activated by depolarization and increased cytoplasmic Ca²⁺.

We next investigated how ATP stimulation translates into changes in membrane potential ($V_{\text{mem}}$). In current-clamp measurements, transient ATP exposure induced a biphasic $V_{\text{mem}}$ response (Fig. 3, F and G). Fast depolarization was followed by gradual repolarization and transient hyperpolarization. Thus, ATP-dependent sequential activation of $I_{\text{inst}}$ and $I_{\text{del}}$ is mirrored by coun-
teracting Vmem fluctuations between each current’s characteristic $E_{\text{rev}}$. BK channels thus appear to narrow the temporal window of ATP-induced depolarization, and therefore, these channels constitute a potential mechanism of negative feedback control in spermatogonial purinergic signaling.

Spermatogonial high-affinity ATP responses are mediated by P2X4 receptors

We next aimed to identify the ion channel or channels underlying $I_{\text{inst}}$. Consistent with previous results (Veitinger et al., 2011), reverse transcription PCR–based expression profiling of all seven P2X receptor isoforms revealed transcripts for P2X2, P2X4, and P2X7 receptors in both Sertoli–germ cell co-cultures and juvenile testes at P7 (Fig. 4 A). Compared with other family members, P2X7 receptors exhibit substantially reduced ATP sensitivity ($EC_{50} \geq 300$ µM [Chessell et al., 1998; Young et al., 2007; Casas-Pruneda et al., 2009]). Subsequent pharmacological/biophysical fingerprinting of $I_{\text{inst}}$ was thus performed using low [ATP]ex (10 µM), Cs+-based pipette solution (S1, S6), or iberiotoxin (100 nM; $E_{\text{hold}} = -40$ mV; S1), respectively (2 min preincubation; right). (E) Bar chart quantifying the effects of pharmacological treatment and reduction of extracellular Ca$^{2+}$ on both $I_{\text{inst}}$ and $I_{\text{del}}$ (peak responses normalized to control conditions). Data are means ± SEM. Numbers of cells analyzed are indicated above bars. Asterisks (*) denote statistical significance, $P < 0.005$ (paired t tests comparing peak currents before [control] and during treatment). (F and G) ATP stimulation (100 µM; 3 s) affects spermatogonial Vmem. (F) Representative whole-cell current-clamp recording (S1, S5) illustrating a biphasic change in Vmem. From a resting potential of approximately $-40$ mV, ATP triggers rapid depolarization toward 0 mV (reversal potential of $I_{\text{inst}}$; B), which is followed by gradual repolarization and transient hyperpolarization. (G) Quantification of current-clamp recordings. Mean Vmem(±SEM; n = 23) is plotted at rest (without ATP; $-39.4 \pm 0.4$ mV) as well as upon maximum depolarization ($-18.1 \pm 1.4$ mV) and hyperpolarization ($-53.1 \pm 1.7$ mV), respectively. For comparison, resting Vmem was adjusted in a few cells to match $-40$ mV by small current injections.
spermatogonia (Fig. 4, B and F). Next, stimulations were performed in the presence of Cu²⁺, which potentiates P2X2 (Xiong et al., 1999) but blocks P2X4 (Coddou et al., 2003). Spermatogonial ATP responses were essentially abolished by Cu²⁺ exposure (Fig. 4, C and F). Similar data resulted from extracellular acidification (pH 6.3; Fig. 4, D and F), which potentiates P2X2 (King et al., 1996; Veitinger et al., 2011) but inhibits P2X4 (Stoop et al., 1997). Finally, we stimulated spermatogonia in the absence and presence of ivermectin (Fig. 4, E and F), which selectively potentiates P2X4 (Khakh et al., 1999; Silberberg et al., 2007; Casas-Pruneda et al., 2009). The macrocyclic lactone mediated a dramatic increase in spermatogonial Iₙₙₙ. Together, these data are most consistent with functional expression of homomeric P2X4 receptors in cultured mouse spermatogonia.

To verify the aforementioned conclusion, we immunohistochemically confirmed endogenous P2X4 expression in spermatogonia-like cells of juvenile seminiferous cords (Fig. S1, E–H), and we assessed the effect of selective P2X4 posttranscriptional gene silencing (knockdown) on spermatogonial ATP signaling by administration of siRNAs in vitro. Cells were transiently transfected with a fluorescent marker and either of two targeting siRNA constructs, a nontargeting negative control siRNA containing a nonspecific sequence or a construct selectively targeting P2X2, respectively (Fig. 5). Effective P2X4 knockdown by either targeting siRNA was confirmed by quantitative real-time PCR (Fig. 5 A). Relative to nontargeting siRNA controls, P2X4 transcript levels were reduced to 47 ± 4% (RNAi1) and 35 ± 5% (RNAi2), respectively. Readily identified by fluorescent labeling (Fig. 5 B), we next recorded Iₙₙₙ from spermatogonia transfected with either siRNA construct (Fig. 5 C). Nontargeting negative control siRNA did not alter peak ATP responses. The same holds true for spermatogonia expressing P2X2-specific targeting constructs, for which we confirmed efficacy in Sertoli cell recordings (Fig. S3). In contrast, compared with both nontransfected cells and spermatogonia expressing nontargeting siRNA controls, selective P2X4 knockdown significantly reduced maximum Iₙₙₙ (Fig. 5 D). Together, gene silencing experiments substantiate the notion that, at relatively low stimulus concentrations, ATP-dependent Iₙₙₙ is predominantly, if not exclusively, mediated by P2X4 receptors.

**P2X4 receptors mediate low-affinity ATP responses in spermatogonia**

The two-step dose–response curve we observed when challenging spermatogonia with a broad range of [ATP]ₑₓ (1 µM to 10 mM; Fig. 2 F) strongly suggests a second low-affinity ATP response mechanism that operates at [ATP]ₑₓ ≥300 µM. Therefore, we next aimed to determine the molecular correlate of this low-affinity cur-
rent. Again, to avoid confounding BK channel currents, experiments were performed using Cs+-based pipette solution (S₆). We first recorded currents induced by prolonged and repeated exposure to high [ATP]ᵅ (1 mM; Fig. 6 A). In sharp contrast to the relatively small, desensitizing inward currents initiated by lower [ATP]ᵅ (10 µM), high stimulus concentrations induced lasting currents that gradually increased during prolonged agonist exposure. By measuring I–V curves at a rate of 2 Hz before, during and after stimulation with either low [ATP]ᵅ (100 µM; Fig. 6, B and C) or high [ATP]ᵅ (1 mM; Fig. 6, D and E), several distinctive features of the low-affinity current emerged, i.e., (a) little rectification, (b) large amplitude, (c) slow development, and (d) no desensitization (Fig. 6, F and G). In concert with our expression profiling results (Fig. 4 A) and the considerably increased [ATP]ᵅ required to activate the current, the aforementioned properties all indicate that P₂X₇ serves as the spermatogonial low-affinity ATP receptor (Surprenant et al., 1996; Chessell et al., 1998; Donnelly-Roberts et al., 2009). If so, the low-affinity current should (a) exhibit enhanced sensitivity when stimulated with the ~10-fold more potent agonist BzATP (Surprenant et al., 1996), (b) gradually increase upon repeated short stimulations (Yan et al., 2008), and (c) be insensitive to selective P₂X₄ knockdown. This is indeed the case (Fig. 7, A and B). When repeatedly exposed to brief pulses of intermediate [ATP]ᵅ (300 µM), currents progressively decreased and mean amplitudes were strongly diminished in spermatogonia transfected with siRNA against P₂X₄. In contrast, successive stimulation with the same concentration of BzATP activated a larger current that increased with multiple stimulations and remained essentially unchanged by P₂X₄ knockdown. This effect, however, was restricted to spermatogonia because BzATP, as compared with ATP, proved much less potent when Sertoli cells were stimulated (Fig. S4), matching a previous report of reduced BzATP sensitivity of the P₂X₂ receptor (Evans et al., 1995).

Further evidence for functional expression of P₂X₇ in spermatogonia emerged from immunochemical analysis, pharmacological inhibition, and posttranscriptional gene silencing. Antibody staining against an extracellular P₂X₇ epitope revealed receptor expression in round cells of the juvenile seminiferous cord (Fig. S1, I–L). In whole-cell recordings, the selective P₂X₇ receptor antagonist A-438079 (McGaraughty et al., 2007) strongly reduced BzATP-mediated currents (Fig. 7, C and D), whereas responses to lower ATP concentrations were unaffected. Moreover, receptor knockdown significantly reduced the relative amount of P₂X₇ transcripts (Fig. 7 E) and had profound effects on current development upon prolonged ATP (1 mM) stimulation (Fig. 7 F). Although the initial desensitizing and inwardly rectifying current was largely unaffected by P₂X₇ gene silencing (Fig. 7, F, G, and I; and arrowhead in F), the slowly increasing essentially nonrectifying current that only saturated after tens of seconds (Fig. 7, F, H, and J; and
Asterisk in F) was substantially diminished in spermatogonia transfected with siRNA constructs targeting P2X7. Together, our findings suggest that P2X7 receptors are crucial mediators of the low-affinity ATP-dependent current in cultured mouse spermatogonia.

Multiple mechanisms of cellular ATP release in response to mechanical or biochemical stimulation have been discussed, including vesicular secretion (Bodin and Burnstock, 2001), active exocytosis of lysosomes (Zhang et al., 2007), and passive transport via connexin or pannexin hemichannels (Cotrina et al., 1998; Bao et al., 2004), large-conductance anion channels (Bell et al., 2003), or voltage-gated ATP-release channels (Taruno et al., 2013). Intriguingly, the P2X7 receptor has also been shown to function as an ATP-induced ATP release route (Pellegatti et al., 2005; Suadicani et al., 2006). Although FSH-dependent ATP release from Sertoli cells has been shown (Gelain et al., 2003, 2005) and ATP secretion from germ cells has been postulated (La-levée et al., 1999), a potential self-enhancement mechanism via P2X7 receptors within the seminiferous epithelium has never been addressed. Therefore, we performed quantitative LC-MS/MS analysis of [ATP]_{ex} in seminiferous tissue samples (Fig. S5). First, we controlled the method's accuracy by adding 10 µM ATP to either controls (water) or tissue samples (± a combination of different ectonucleotidase inhibitors). LC-MS/MS detected 9.93 ± 0.97 µM ATP in "no-tissue" controls (n = 6), suggesting a high degree of analytical accuracy. Seminiferous tissue that retained ectonucleotidase activity showed slightly, though not significantly reduced [ATP]_{ex} (8.4 ± 1.88 µM), whereas broad range inhibi-
tion of ectonucleotidases resulted in slightly increased ATP levels (11.07 ± 0.76 µM; Fig. S5 A). Next, we compared the extracellular ATP content of seminiferous tissue samples (± ectonucleotidase inhibition) in the absence or presence of BzATP (Fig. S5 B). Control experiments showed (a) that BzATP does not contaminate the mass traces of ATP and its metabolites and (b) that LC-MS/MS results had to be corrected for up to 4% chemical impurity (not depicted). When stimulated with BzATP, tissue weight–corrected ATP levels massively increased by factors of 38 (− blocking agents) and 21 (+ inhibitor cocktail), respectively. These results thus indicate that purinoceptor activation in seminiferous

**Figure 7.** P2X7 constitutes the low-affinity spermatogonial ATP sensor. (A) Exemplary whole-cell voltage-clamp recordings of currents triggered by repetitive stimulation (1 s; ISI = 3 s; Si, Sj with ATP (300 µM; left) or BzATP (300 µM; right), respectively. Traces from untransfected spermatogonia (black) and cells transfected with either nontarget negative control

siRNA (gray) or a construct targeting P2X4 (green) are overlaid. (B) Quantification of results from repetitive stimulation experiments as shown in A. Peak current densities (means ± SEM) are color coded and plotted as a function of stimulus repetition. Cells were analyzed under control conditions (n = 33/13 [ATP/BzATP]; black), after transfection with nontargeting siRNA (n = 16/23 [ATP/BzATP]; gray), and after P2X4 knockdown (n = 16/23 [ATP/BzATP]; green). Inset (left) shows the data delimited by the dashed rectangle at an increased scale. Asterisks (*) indicate statistical significance, P < 0.05 (one-way ANOVA with Tukey’s HSD post hoc test). (C and D) Selective pharmacological inhibition of BzATP responses by 10 µM A-438079. (C) Representative whole-cell voltage-clamp recordings show individual responses to either 300 µM ATP (top) or 300 µM BzATP (bottom) in the absence or presence of the P2X7 receptor antagonist A-438079 (preincubation >60 s; Si, Sj). (D) Bar chart quantifying the efficacy of A-438079 as a function of agonist type (ATP vs. BzATP). Data are means ± SEM; normalized to control conditions. Numbers of cells are indicated above bars. Asterisk (*) denotes statistical significance, P < 0.005 (paired t test). (E) When transfected with transcript-specific siRNA (green), posttranscriptional gene silencing of P2X7 expression is confirmed by quantitative PCR. Relative transcript levels (means ± SEM) are normalized to mRNA quantities in untransfected spermatogonia and compared with cells treated with nontargeting siRNA controls (gray). Asterisks (*) denote statistical significance, P < 0.005 (one-way ANOVA with Tukey’s HSD post hoc test). (F) Representative current density plot over time. Each dot represents current measured at −80 mV obtained from sequential voltage ramps as in G and H. Data correspond to spermatogonia transfected with either targeting P2X7 siRNA (green) or a nontargeting control construct (black). When challenged with 1 mM ATP, a fast but relatively small inward current develops and shows a transient peak (first Imax; red arrowhead). After apparent desensitization, current amplitudes gradually increase over tens of seconds until a secondary maximum is reached (second Imax; red asterisk). (G and H) Exemplary current–voltage curves (−100 to 100 mV; 250 ms; 1 Hz repetition; S1, S2) in response to 1 mM ATP, corresponding to time points (first G vs. second H Imax) as indicated in F (red arrowhead and asterisk). Color code as for E and F. Data were corrected by digital offline subtraction of averaged leak controls recorded before stimulation. (I and J) Quantification of mean current densities measured at −80 mV at first (I) versus second (J) Imax; see F–H. Bar graphs display means ± SEM. Numbers of cells as shown above bars. Asterisks (*) indicate statistical significance, P < 0.05 (one-way ANOVA with Tukey’s HSD post hoc test).
tissue can trigger a self-enhancing ATP release pathway, possibly via spermatogonial P2X7 receptors.

An acute seminiferous cord preparation links in vitro findings to endogenous (electro)physiological profiles of male germ cells

So far, our data show that, in a Sertoli cell–germ cell co-culture, spermatogonia from juvenile mice detect elevations in [ATP]_ex over a broad concentration range using both P2X4 and P2X7 receptors as well as BK channels that somewhat balance the ATP-dependent depolarization. To determine how these findings compare with data obtained in situ, we established an acute seminiferous cord slice preparation (Fig. 8 A) from juvenile mice. These sections preserve much of the tissue integrity, leaving the structure of, for example, Sertoli–germ cell junctions (Cheng and Mruk, 2002) or spermatogonial cytosolic bridges (Chiarini-Garcia and Russell, 2002) largely intact. Transverse section planes, in particular, allow for patch pipette access to both putative Sertoli cells (Fig. 8 B) and spermatogonia (Fig. 8 D) under visual control. Diffusion loading of target cells with a fluorophore during whole-cell recordings enables online visualization of cell morphology as well as post hoc three-dimensional reconstruction. ATP-induced currents from large irregularly shaped, columnar cells—putative Sertoli cells—displayed the nondesensitizing waveform of currents mediated by P2X2 receptors, irrespective of low or high [ATP]_ex (Fig. 8 C). In contrast, recordings from smaller round cells—putative spermatogonia—revealed relatively rapid, desensitizing responses upon exposure to low [ATP]_ex (10 µM) and slow, gradually increasing responses when challenged with high [ATP]_ex (1 mM; Fig. 8 E). Comparative quantification of spermatogonial currents demonstrated dose-dependent peak responses as well as pronounced desensitization in response to low [ATP]_ex versus no desensitization during stimulation with high [ATP]_ex (Fig. 8 F). These data indicate that, in an intact environment, spermatogonia express at least two different ATP-dependent conductances that share kinetic hallmarks of P2X4 and P2X7 receptors.

If the observed in situ response profiles emerge from P2X2 receptor expression in Sertoli cells and/or expression of P2X4 and P2X7 receptors in spermatogonia,
respectively, ATP responses in acute sections should exhibit pharmacological characteristics similar to recordings from cells in Sertoli cell–germ cell co-culture. To address this, we recorded whole-cell currents from putative Sertoli cells and spermatogonia in response to both low and high \([\text{ATP}]_{\text{ex}}\) (10 \(\mu\)M and 1 mM, respectively) in the absence and presence of different P2X receptor antagonists (Fig. 9). Although suramin strongly inhibited low \([\text{ATP}]_{\text{ex}}\) responses in putative Sertoli cells, spermatogonial currents were essentially unaffected (Fig. 9, B and D). In contrast, the selective P2X7 receptor blocker A-438079 reduced spermatogonial responses (Fig. 9, A and D). Together, these findings support the endogenous functional expression of cell type–specific purinoceptor profiles within the premeiotic seminiferous epithelium. Our data also indicate that electrophysiological in vitro phenotypes are largely mirrored by both somatic and germ cells in an intact seminiferous cord preparation. Moreover, acute seminiferous cord/tubule slices provide an attractive experimental platform for direct analysis of diverse physiological processes in male reproductive biology.

**DISCUSSION**

Spermatogenesis, the complex, lifelong process of mass germ cell proliferation and transformation, is fundamental to reproduction and fertility, yet physiologically ill defined. Although extensive communication among and between Sertoli and germ cells coordinates spermatogenesis (Cheng and Mruk, 2002), the function of numerous paracrine factors that have been implicated in the seminiferous cycle remains elusive (Schlatt and Ehmcke, 2014). Here, we postulate a role for ATP in seminiferous epithelial communication, a key signaling component of several paracrine networks involved in other developmental processes (Abbracchio et al., 2009; Praetorius and Leipziger, 2009). In vitro and in situ experiments identify both P2X4 and P2X7 receptors as well as BK channels as critical mediators of purinergic responses in juvenile mouse spermatogonia. Cell- and stage-specific testicular expression of several purinoceptor subunits has been reported in Sertoli cells (Veitinger et al., 2011), Leydig cells (Foresta et al., 1996; Antonio et al., 2009), postmeiotic germ cells (Glass et al., 2001), and mature spermatozoa (Navarro et al., 2011), albeit with somewhat contradictory results. An emerging concept implicates purinergic signaling mechanisms in paracrine control of gonadotropin effects on Sertoli and Leydig cells (Filippini et al., 1994; Meroni et al., 1998; Lalveé et al., 1999; Gelain et al., 2003, 2005), including, but not limited to, steroidogenesis and testosterone/17\(\beta\)-estradiol secretion (Foresta et al., 1996; Rossato et al., 2001). So far, the lack of an intact in situ preparation impeded functional analysis of the testicular purinergic signaling network in a physiological setting. The cellular complexity of the adult seminiferous epithelium as well as the lack of cell type–specific live cell markers have made it particularly difficult to analyze the distinct physiological properties of individual germ cell populations. The combination of a Sertoli–germ cell co-culture with acute seminiferous tubule sections from prepubescent mice overcomes this limitation as it allows a detailed physiological characterization of a defined germ cell population both in vitro and in situ. Our findings add spermatogonia to the seminiferous purinergic signaling circuit, suggesting that local \([\text{ATP}]_{\text{ex}}\) elevations will affect premeiotic male germ cells both electrophysiologically, by transient de- and hyperpolarization, and biochemically, by triggering substantial \(\text{Ca}^{2+}\) influx (unpublished data). Our data also indicate that one result of seminiferous

![Figure 9. Pharmacological profile of purinergic signals in the intact seminiferous epithelium.](image-url)
[ATP]_{ex} elevation is paracrine signal amplification by increased ATP release, although the cellular identity of the ATP source or sources is currently unclear. The impact of this positive feedback on each testicular cell population will depend on the cell type– and stage-specific purinoceptor expression profile, as different isoforms vary dramatically with respect to agonist affinity, response kinetics, and desensitization (Coddou et al., 2011; Khakh and North, 2012).

The mechanism or mechanisms of ATP-induced ATP release remain elusive. ATP-induced cytosolic Ca^{2+} transients—potentially mediated by Ca^{2+} influx through P2X receptors, P2Y receptor-dependent Ca^{2+} release from storage organelles (Gelain et al., 2005; Veitinger et al., 2011), and/or activation of voltage-gated Ca^{2+} channels (Liévano et al., 1996)—could trigger exocytotic release of vesicles (Bodin and Burnstock, 2001) and other secretory granules (Zhang et al., 2007). Although nanomolar pericellular ATP concentrations are generally found in quiescent cells (Falzoni et al., 2013), the ATP concentration in secretory vesicles is claimed to be in the high millimolar range (Fields and Burnstock, 2006). However, alternative ATP release pathways, such as connexin or pannexin hemichannels (Cotrina et al., 1998; Bao et al., 2004), ATP-binding cassette transporters (Lohman et al., 2012), voltage-gated release channels (e.g., CALHM1 (Taruno et al., 2013), large-conductance anion channels (Bell et al., 2003), or even P2X7 receptors themselves (Pellegrini et al., 2005; Suadicani et al., 2006), cannot be excluded. Rapid enzymatic degradation by ectonucleotidases renders the half-life of extracellular ATP relatively short and thus narrows its paracrine radius to a few hundred micrometers (Fitz, 2007). Signal spread, however, is fast as ATP should diffuse ~1 µm in <10 ms (Khakh, 2001). Given the spatial dimensions of the seminiferous tubule (Cheng and Mruk, 2002), ATP is thus ideally suited for a role as a fast local signal in paracrine testicular communication (Praetorius and Leipziger, 2009).

Which physiological roles might be included in the functional portfolio of testicular ATP signals? Evidently, the complexity and heterogeneity of the purinoceptor family allows for highly dynamic signaling over broad spatiotemporal scales. Short-term effects might include remodeling of the blood–testis barrier (Cheng and Mruk, 2002), synchronization of epithelial stage transitions (Heindel and Treinen, 1989; Syed and Hecht, 2002), control of bidirectional exchange between Sertoli and germ cells (McLachlan et al., 1995), and spontaneous epithelial Ca^{2+} oscillations (Sánchez-Cárdenas et al., 2012). Importantly, purinoceptor-dependent cytosolic Ca^{2+} elevations could also work on prolonged time scales, outlasting the short-term effects of the extracellular ATP surge (Khakh, 2001). For instance, Ca^{2+}-dependent recruitment of transcription factors and resulting long-term changes in gene expression could affect proliferation, differentiation, migration, and epithelial cell turnover. Because ATP release from apoptotic cells is well documented (Elliott et al., 2009) and because the P2X7 receptor, in particular, has been attributed a key role in cell death regulation (Surprenant et al., 1996), it is tempting to speculate that balancing the density-dependent germ cell death/survival ratio could also be a major function of the testicular purinergic signaling toolkit. Apoptosis is a vital process during spermatogenesis (Print and Loveland, 2000) as up to 75% of all germ cells undergo apoptosis under physiological conditions (Huckins, 1978).

With the exception of P2X1, which is critical for vas deferens contraction (Mulryan et al., 2000), no severe fertility deficits have been observed in P2X receptor–null mutants. In fact, phenotypic alterations in P2X knockout mice are rather subtle. Mice deficient for either P2X4 or P2X7 mostly show some immunological, cardiovascular, and/or neurological deficits as well as increased susceptibility to neuropathic pain (Coddou et al., 2011). However, given the versatile and often critical roles of purinoceptors in numerous physiological processes, deficiency compensation by alternative P2X isoform is likely to obscure the interpretation of gross phenotypic observations from knockout models.

In summary, we describe functional expression of P2X4 and P2X7 receptors as well as BK channels in immature male germ cells. Spermatogonia are thus equipped to respond to local testicular [ATP]_{ex} elevations over a broad dynamic range with distinct and variable electrophysiological features, including negative feedback regulation of V_mer fluctuations by BK channel recruitment. In this scenario, ATP may function as a self-amplifying signal. Future studies will aim to determine the potential role or roles of purinoceptor signaling during meiosis and postmeiotic germ cell maturation.

ACKNOWLEDGMENTS

We thank Christa E. Müller for kindly providing reagents, Ingo Scholz (RWTH Aachen University) for helping with scanning electron microscopy, Annika Triller (University of Bonn) and Hanns Hatt (Ruhr-University Bochum) for help during the early stage of the project, Ralf Hausmann and Günther Schmalzing (RWTH Aachen University) for helpful comments and suggestions, and all members of the Spehr laboratory for stimulating discussions.

This work was funded by the Volkswagen Foundation (1/83533 to M. Spehr). M. Spehr is a Lichtenberg Professor of the Volkswagen Foundation.

The authors declare no competing financial interests.

Sharona E. Gordon served as editor.

Submitted: 7 June 2016
Accepted: 22 July 2016
Purification and characterization of a unique, potent, peptidyl probe for the high conductance calcium-activated potassium channel from venom of the scorpion Bothus tamarus. J. Biol. Chem. 265: 11083–11090.


