cAMP/PKA-Dependent Increases in Ca Sparks, Oscillations and SR Ca Stores in Retinal Arteriolar Myocytes after Exposure to Vasopressin

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PURPOSE. To investigate the effects of arginine vasopressin (AVP) on Ca2+ sparks and oscillations and on sarcoplasmic reticulum (SR) Ca2+ content in retinal arteriolar myocytes.

METHODS. Fluo-4-loaded smooth muscle in intact segments of freshly isolated porcine retinal arteriole was imaged by confocal laser microscopy. SR Ca2+ store content was assessed by recording caffeine-induced Ca2+ transients with microfluorimetry and fura-2.

RESULTS. The frequencies of Ca2+ sparks and oscillations were increased both during exposure to, and 10 minutes after washout of AVP (10 nM). Caffeine transients were increased in amplitude 10 and 90 minutes after a 3-minute application of AVP. Both AVP-induced Ca2+ transients and the enhancement of caffeine responses after AVP washout were inhibited by SR 49050, a V1a receptor blocker. Forskolin, an activator of adenyl cyclase, also persistently enhanced caffeine transients. Rp-8-HA-cAMPS, a membrane-permeant PKA inhibitor, prevented enhancement of caffeine transients by both AVP and forskolin. Forskolin, but not AVP, produced a reversible, Rp-8-HA-cAMPS insensitive reduction in basal [Ca2+].

CONCLUSIONS. AVP activate a cAMP/PKA-dependent pathway via V1a receptors in retinal arteriolar smooth muscle. This effect persistently increases SR Ca2+ loading, upregulating Ca2+ sparks and oscillations, and may favor prolonged agonist activity despite receptor desensitization. (Invest Ophthalmol Vis Sci. 2010;51:1591–1598) DOI:10.1167/iovs.09-4401

Modulation of arteriolar resistance plays a central role in the control of regional blood flow and capillary pressure in the retina (see recent review in Ref. 1). To understand how flow is regulated to meet local needs, we must understand Ca2+ signaling within the smooth muscle of these vessels, since intracellular Ca2+ ([Ca2+]i) is a key regulator of contraction and thus of constriction in ocular vessels (reviewed in Ref. 2). Retinal arterioles are not autonomically innervated,3–4 and modulation of tone depends on local control mechanisms, such as those responsible for autoregulation of retinal blood flow in the face of changes in blood pressure or metabolic demand.5,6 Vasopressin, which constricts retinal arterioles in vitro,7 may play a role in such control. The endocrine actions of circulating vasopressin in the control of renal H2O reabsorption and vascular tone are well known (see reviews in Refs. 8, 9). However, locally released vasopressin acts as a paracrine signal within the brain itself.10 Vasopressin has been detected in rat retina,11 where it is localized to the ganglion cell layer close to the inner retinal vessels.12 It seems possible, therefore, that local release of vasopressin would contribute to retinal vascular tone. Retinal vasopressin levels are elevated during periods of exposure to light, when total retinal metabolism is reduced.11,13 The vasoconstrictor action of vasopressin may therefore help explain how this change in metabolic need is coupled to a reduction in retinal blood flow.14 Arginine vasopressin (AVP) stimulates a transient increase in smooth muscle intracellular [Ca2+]i, and vessel constriction when applied externally to retinal arteriole segments.7 In common with many other vascular preparations, this response desensitizes in pig (though not in rat) retinal vessels.1,15,16 We have reported that myocytes in freshly isolated segments of rat retinal arterioles generate asynchronous Ca2+ sparks and oscillations.17,18 Brief Ca2+ sparks summate to produce the more prolonged oscillations that are associated with muscle contraction, which leads us to propose that Ca2+ sparks are excitatory, proconstrictor events in retinal arterioles, rather than prodilatory, as in larger arteries.19 The purpose of the present study was to characterize the effects of AVP, a physiological vasoconstrictor, on Ca2+ sparks and oscillations in pig retinal arteriolar myocytes. These asynchronous, phasic, cellular and subcellular signaling events were increased in frequency both during application of AVP and 10 minutes after washout. SR Ca2+ loading was increased up to 90 minutes after AVP exposure, by a CAMP, protein kinase A (PKA)-dependent mechanism. These novel findings suggest that local or circulating AVP had prolonged effects on retinal vascular resistance, despite receptor desensitization.

MATERIALS AND METHODS

Arteriole Preparation

Pigs were humanely killed at a registered slaughterhouse in compliance with the ethical and legal requirements specified by the Welfare of Animals (Slaughter or Killing) Regulations (Northern Ireland) 1996. The eyes were transported in low-Ca2+ solution at 1°C. The retinas were removed and the arterioles mechanically isolated according to a published method.7,17 The retinas were triturated in low-Ca2+ Hanks’ solution, centrifuged, washed with low-Ca2+ medium, and centrifuged.

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again. Vascular fragments were then incubated for 2 hours with the relevant Ca²⁺ indicator in low-Ca²⁺ solution. Arteriole segments (outer diameter, 30–100 μm) were anchored in position with tungsten wire slips and superfused with normal Hanks’ solution at 37°C.

### Ca²⁺ Imaging

High-speed Ca²⁺ imaging was performed, as described elsewhere. Retinal arterioles were incubated with 10 μM fluo-4 AM and imaged with a confocal scanning laser microscope (model MR-A1; Bio-Rad, Hercules, CA), mounted on an inverted microscope (Eclipse TE300; Nikon Instruments, Tokyo, Japan, with a PlanApo, 60, 1.4 NA, oil-immersion objective). The vessels were excited at 488 nm, and emitted light was filtered (530–560 nm) and detected with a photomultiplier tube. Data acquisition was controlled with commercial software (Time-course; Bio-Rad). Background-corrected fluorescence (F) was normalized to the resting fluorescence (F₀) in each cell and F/F₀ was calculated. Linescan images (500 lines second⁻¹) were limited to a maximum of 150 seconds because of photodamage.

To minimize observer bias, custom software has been developed for the automated detection and analysis of Ca²⁺ sparks. This method involves an ‘a trous’ wavelet transform for detecting small-amplitude events in noisy images. Events detected but not apparent on the image were manually deleted. Ca²⁺ spark amplitudes were recorded as the maximum increase in normalized fluorescence (ΔF/F₀) for a 5-pixel-wide region of interest centered on the point of maximum fluorescence. The software also extracted the full duration at half-maximum fluorescence (FDHM) of each event. Oscillations were analyzed with ImageJ (developed by Wayne Rasband, National Institutes of Health, Bethesda, MD; available at http://rsb.info.nih.gov/ij/index.html), to determine the amplitude and duration of manually defined events.

### Ca²⁺ Microfluorimetry

Changes in [Ca²⁺] within myocytes were recorded by using microfluorimetry, as previously described. Vessels were incubated in a 5-μM fura-2AM solution with DMSO (1% vol/vol), visualized on an inverted microscope (Diaphot 200; Nikon) and excited alternately at 340 and 380 nm (20 cyc/s). This method loads smooth muscle but not endothelium with the Ca²⁺ indicator in this preparation. Emitted fluorescence was recorded at 510 nm for each excitation wavelength (F₃₈₀ and F₅₁₀) with a spectrofluorometer (Ratiomaster RF-F3011; Photon Technology International, South Brunswick, NJ). The fluorescence ratio (R = F₃₈₀/F₅₁₀) was used to track changes in [Ca²⁺]). There were large changes in the absolute values of R during this project, although the proportional dynamic range, signal-to-noise ratio, and relative amplitudes of the caffeine responses were unaltered. To allow data to be summarized, changes in R (ΔR) have been normalized using the resting ratio in the same tissue at the beginning of the experiment (R₀). No attempt has been made to convert the data to [Ca²⁺], and analysis has been restricted to the changes in responses in internally controlled experiments.

### Solutions and Drugs

The bath perfusate had the composition (mM): NaCl, 140; KCl, 5; Na-glucose, 5; CaCl₂, 2; MgCl₂, 1.3; and HEPES, 10, with pH set to 7.4 with NaOH. Solution for vessel isolation contained 0.1 mM CaCl₂. Drugs were applied in a prewarmed bath solution to the outside of the arterioles. The drugs were obtained from Sigma-Aldrich (Poole, UK), with the exception of the V₁₃ antagonist, SR49059 (the kind gift of Claudine Serradeil-Le Gal; Sanofi, France).

### Data Analysis

Summarized data are expressed as the mean ± SEM. Ca²⁺ spark and oscillation data for pig arterioles were not normally distributed (P < 0.001; Kolmogorov-Smirnov test), and so a nonparametric Mann-Whitney test was used. Differences between mean [Ca²⁺]transients in microfluorimetry experiments were assessed with a paired t-test for single comparisons, or repeated-measures ANOVA with a Tukey-Kramer post hoc test for multiple comparisons. The accepted significance level was P ≤ 0.05.

### RESULTS

#### Spontaneous Ca²⁺ Sparks and Oscillations under Control Conditions

Confocal linescan Ca²⁺ images from myocytes in porcine retinal arterioles revealed spontaneous Ca²⁺ sparks and oscillations.
Table 1. Summary Data for Ca\textsuperscript{2+} Sparks and Oscillations Recorded from Porcine Retinal Arteriolar Myocytes under Control Conditions and during the First 120 Seconds of AVP Exposure

<table>
<thead>
<tr>
<th>Ca\textsuperscript{2+} Events</th>
<th>Control</th>
<th>AVP</th>
<th>P</th>
</tr>
</thead>
<tbody>
<tr>
<td>Spark frequency, /10s</td>
<td>0.27 ± 0.03 (139 cells)</td>
<td>0.47 ± 0.19 (41 cells)</td>
<td>0.034</td>
</tr>
<tr>
<td>Spark amplitude, ΔF/Fo</td>
<td>0.47 ± 0.02 (48 events)</td>
<td>0.48 ± 0.01 (164 events)</td>
<td>NS</td>
</tr>
<tr>
<td>FDHM, ms</td>
<td>27 ± 3 (48 events)</td>
<td>24 ± 1 (164 events)</td>
<td>NS</td>
</tr>
<tr>
<td>Oscillation frequency, /10s</td>
<td>0.11 ± 0.04 (53 cells)</td>
<td>0.32 ± 0.08 (53 cells)</td>
<td>0.015</td>
</tr>
<tr>
<td>Oscillation amplitude, ΔF/Fo</td>
<td>0.45 ± 0.05 (14 events)</td>
<td>0.61 ± 0.03 (194 events)</td>
<td>NS</td>
</tr>
<tr>
<td>Duration, s</td>
<td>6.51 ± 1.15 (14 events)</td>
<td>4.94 ± 0.33 (194 events)</td>
<td>NS</td>
</tr>
</tbody>
</table>

The data were obtained from linescans of arterioles from 4 animals: 10 control vessels and 8 exposed to AVP. Average frequencies were calculated based on all the cells analyzed, including 0 values. P-values were calculated with the Mann-Whitney U-test. NS, P > 0.05.

Effect of AVP on Ca\textsuperscript{2+} Sparks and Oscillations

High-speed Ca\textsuperscript{2+} imaging revealed the complexity of cellular responses to AVP. AVP (10 nM) increased fluorescence, indicating a rise in baseline cytoplasmic [Ca\textsuperscript{2+}] elevations lasting many seconds (Fig. 1C). We use the term oscillation for these events, but recognize that they may include longitudinally propagating Ca\textsuperscript{2+} waves, which cannot be identified in our transverse linescans. Both spark and oscillation characteristics are summarized in Tables 1 and 2.

Table 2. Summary Data for Ca\textsuperscript{2+} Sparks and Oscillations Recorded under Control Conditions and from Linescans of up to 130-Second Duration Recorded 10 Minutes after AVP Washout

<table>
<thead>
<tr>
<th>Ca\textsuperscript{2+} Events</th>
<th>Control</th>
<th>10 min WO after AVP</th>
<th>P</th>
</tr>
</thead>
<tbody>
<tr>
<td>Spark frequency, /10s</td>
<td>0.31 ± 0.04 (93 cells)</td>
<td>0.45 ± 0.05 (66 cells)</td>
<td>0.014</td>
</tr>
<tr>
<td>Spark amplitude, ΔF/Fo</td>
<td>0.54 ± 0.15 (251 events)</td>
<td>0.66 ± 0.02 (273 events)</td>
<td>NS</td>
</tr>
<tr>
<td>FDHM, ms</td>
<td>20 ± 1 (251 events)</td>
<td>21 ± 1 (273 events)</td>
<td>NS</td>
</tr>
<tr>
<td>Oscillation frequency, /10s</td>
<td>0.12 ± 0.02 (86 cells)</td>
<td>0.25 ± 0.03 (80 cells)</td>
<td>0.0008</td>
</tr>
<tr>
<td>Oscillation amplitude, ΔF/Fo</td>
<td>0.66 ± 0.04 (87 events)</td>
<td>0.75 ± 0.04 (180 events)</td>
<td>NS</td>
</tr>
<tr>
<td>Duration, s</td>
<td>7.09 ± 0.71 (87 events)</td>
<td>8.06 ± 0.54 (180 events)</td>
<td>NS</td>
</tr>
</tbody>
</table>

P-values were calculated with the Mann-Whitney U-test. NS, P > 0.05.

Effect of AVP on SR Ca\textsuperscript{2+} Loading in Retinal Arterioles

A possible cause of increased Ca\textsuperscript{2+} sparks and oscillations is increased SR Ca\textsuperscript{2+} loading. This possibility was tested by using caffeine to release intracellular Ca\textsuperscript{2+} stores by opening ryadnone receptors (RyRs). Caffeine (10 mM) produced a rapid Ca\textsuperscript{2+} transient followed by a variable undershoot under control conditions (Fig. 4). Subsequent superfusion with 10 nM AVP produced a smaller transient with slower kinetics. Cellular and subcellular signaling detail such as sparks and oscillations were not resolved. When caffeine was applied, 10 minutes after washout of AVP, a transient followed by a variable undershoot was observed (Fig. 3). The transient was not increased compared with the control.

Enhancement of Caffeine Responses by AVP Mediated by V\textsubscript{1A} receptors

Application of AVP failed to evoke an increase in [Ca\textsuperscript{2+}] when a V\textsubscript{1A} receptor blocker (SR 49059; 10 nM) was included in the bathing medium from 5 minutes before exposure to AVP.
application of AVP until 2 minutes after washout (Fig. 5A). The response to caffeine after AVP washout was similar to that during the control period. In 15 vessels from three animals, the mean caffeine response ($\frac{\Delta R}{R_0}$) was 0.77 ± 0.06 before AVP application and 0.78 ± 0.06 after a 10-minute washout (NS, paired t-test; Fig. 5B). This result suggests that both the direct effect of AVP on $[Ca^{2+}]_{i}$ and the persistent enhancement of the caffeine-evoked response were mediated via V1a receptors.

**Effect of Forskolin on Caffeine-Evoked Transients in Retinal Arterioles**

Experiments were performed to identify the signaling pathway involved in the increase in SR $Ca^{2+}$ loading. It has been demonstrated that forskolin, which activates adenylyl cyclase and elevates cAMP levels (see reviews in Refs. 26, 27), also increases caffeine responses in isolated vascular myocytes from cerebral arteries in both rats and mice.$^{27}$ Forskolin (10 $\mu$M)

![Figure 2](image2.png)

**Figure 2.** AVP increased $Ca^{2+}$ sparks and oscillations. (A) A 2-D confocal optical section through eight adjacent fluo-4-loaded myocytes in a rat retinal arteriole (left), indicating the relevant scanline. Linescan images are shown for the same cells (horizontal arrows: cell boundaries) under control conditions and during superfusion with AVP (10 nM). Normalized fluorescence ($F/F_0$) is plotted for the myocyte marked X; arrows: $Ca^{2+}$ sparks. AVP exposure commenced 60 seconds before acquisition of the rightmost linescan. Summary data (mean ± SEM) are for (B) spark and (C) oscillation frequencies under control conditions and during AVP exposure.

![Figure 3](image3.png)

**Figure 3.** $Ca^{2+}$ sparks and oscillations were persistently increased after washout of AVP. (A) Confocal linescan records from separate regions of the same arteriole (Ai) under control conditions and (Aii) 10 minutes after washout of AVP (10 nM). XY frames to the left of each panel show six contiguous arteriolar myocytes and the relevant scanline (white). Horizontal arrows: cell boundaries. $F/F_0$ is plotted for a single myocyte in each linescan (a and b); arrows: sparks. (B) Summary data (mean ± SEM) are for spark frequency recorded before (control) and 10 minutes after washout of AVP from different regions in the same vessels. (C) Summary data (mean ± SEM) for oscillation frequency recorded under control conditions and 10 minutes after washout of AVP.
mimicked the effect of AVP on caffeine transients in pig retinal arterioles. Caffeine (10 mM) was applied under control conditions and again after a 10-minute superfusion with forskolin (Fig. 6A). Forskolin itself produced a small but consistent decline in the fluorescence ratio, which averaged 0.08 ± 0.02 relative to the control (P < 0.005, paired t-test). This decrease recovered during a 10-minute washout, after which the caffeine response was still enhanced relative to the control. In 12 vessels from six animals, the mean response to caffeine (ΔR/ R₀) was increased from 0.48 ± 0.06 under control conditions to 0.59 ± 0.05 after superfusion with forskolin for 10 minutes (P < 0.05 versus control, ANOVA, Fig. 6B). After a further 10-minute washout, caffeine transients were still enhanced, with a mean increase of 0.62 ± 0.06 (P < 0.01 versus control, ANOVA).

**Effect of Rp-8-HA-cAMPS on Enhancement of Caffeine-Evoked Transients by AVP and Forskolin**

Cyclic AMP exerts many of its effects through activation of protein kinase A (PKA; see review in Ref. 28). We tested this

![Figure 4](image1.png)  
**Figure 4.** Caffeine-evoked Ca²⁺ transients were enhanced after AVP treatment. (A) Sample record showing changes in the fura-2 fluorescence ratio, normalized to the resting ratio (ΔR/R₀) for an arteriole exposed to caffeine under control conditions, then approximately 10 minutes after washout of AVP, and again approximately 90 minutes after washout. Changes in the fura-2 fluorescence ratio (ΔR) have been normalized to the resting ratio in the same arteriole (R₀). (B) Summary data for six arterioles showing the mean responses (±SEM) to each of the caffeine stimuli during the protocol shown in (A). *P < 0.05 versus control. (C) Ca²⁺ transients recorded from an arteriole during three consecutive applications of caffeine (10 mM). The vessel was superfused with normal bath solution for approximately 10 minutes between caffeine applications. (D) Summary data for eight vessels show the mean response (±SEM) to each of three consecutive applications of caffeine.

![Figure 5](image2.png)  
**Figure 5.** AVP effects were blocked by SR49059. (A) A sample record of [Ca²⁺]ᵢ for an arteriole that was exposed to caffeine and then to the V₁₃ blocker SR49059. AVP was then applied, and caffeine was reapplied after washout of AVP and SR49059. (B) Summary data for 15 arterioles show the mean responses (±SEM) to the two caffeine stimuli during the protocol shown in (A).

![Figure 6](image3.png)  
**Figure 6.** Caffeine-evoked Ca²⁺ transients were enhanced after exposure to forskolin. (A) A sample record of [Ca²⁺]ᵢ for an arteriole which was exposed to caffeine (10 mM) under control conditions, after a 10-minute exposure to forskolin (10 μM), and finally, approximately 10 minutes after washout of forskolin. Changes in the fura-2 fluorescence ratio (ΔR) were normalized to the resting ratio in the same arteriole (R₀). (B) Summary data for 12 arterioles showing the mean responses (±SEM) to each of the caffeine stimuli during the protocol shown in (A). *P < 0.05; **P < 0.01; versus control in both cases.
effect by using the Rp-isomer of the compound 8-hexylamino-
adenosine-3', 5'-monophosphorothioate (Rp-8-HA-cAMPS), a lipophilic PKA inhibitor.\textsuperscript{29} Tissues were pre-exposed to the drug (10 \(\mu\)M) for a minimum of 2 hours. Brisk, caffeine-evoked responses were still seen but were no longer enhanced after treatment with AVP or forskolin (Fig. 7). In a series of 19 vessels (from eight animals) preincubated with Rp-8-HA-cAMPS, the mean caffeine-induced increase in fluorescence ratio (\(\Delta R/R_0\)) was 0.93 \pm 0.09 during the control period and 0.82 \pm 0.08 after a 10-minute washout after AVP (NS, paired \(t\)-test; Fig. 7B). [Ca\(^{2+}\)]\(_i\) transients were still evoked by AVP itself in these experiments, with a mean increase in normalized ratio of 0.31 \pm 0.04 (Fig. 7A). Rp-8-HA-cAMPS also blocked the increase in caffeine responses after exposure to forskolin (Fig. 7C). In 12 vessels from six animals, the mean caffeine response was 0.76 \pm 0.12 during the control period, 0.76 \pm 0.08 after 10 minutes of forskolin superfusion, and 0.75 \pm 0.07 after a further 10-minute washout (Fig. 7D; NS, ANOVA versus control). The forskolin-induced reduction in baseline was not affected in these experiments (Fig. 7C), suggesting that the effect was not PKA dependent.

**DISCUSSION**

In the present study, we demonstrated for the first time in any native vascular myocytes that AVP persistently increases the frequency of Ca\(^{2+}\) sparks and oscillations in retinal arterioles, as well as raising baseline [Ca\(^{2+}\)]\(_i\), (Table 1, Fig. 2). The fact that AVP, which constricts retinal arterioles,\textsuperscript{50} also increases sparks and oscillations, is consistent with our proposal that sparks can act as building blocks for global Ca\(^{2+}\) signals during excitation–contraction coupling in these microvessels.\textsuperscript{18} In pig retinal vessels, as in many other preparations, AVP responses desensitize relatively rapidly.\textsuperscript{1,15,16} Persistent upregulation of Ca\(^{2+}\) sparks and oscillations after washout of AVP (Fig. 3) may prolong agonist effects despite receptor desensitization. In rat retinal arteriolar myocytes, sparks and oscillations reflect release of SR Ca\(^{2+}\) via RyR2-gated channels,\textsuperscript{31} and brief localized sparks can summate to generate more prolonged and global Ca\(^{2+}\) oscillations.\textsuperscript{18} Spark and oscillation characteristics in porcine vessels (Tables 1, 2) were similar to those previously observed in the rat.\textsuperscript{17,18} It should be noted that the apparent increase in oscillation duration in this study reflects a change in the measure used (from FDHM to full duration) rather than any biological difference. Ca\(^{2+}\) oscillations were also observed to result from spark summation in pig arterioles (Figs. 1C, 2A, 3A). Oscillations are associated with myocyte contraction in rat retinal vessels\textsuperscript{17,18} and, although transverse linescans cannot detect cell shortening, the narrowing of the cell profile during the second oscillation in Figure 1C almost certainly resulted from cell movement due to contraction. Taken together, this evidence again suggests that Ca\(^{2+}\) sparks promote constriction in these microvessels, rather than relaxation, as in larger arteries.\textsuperscript{1,19}

Our results support a model in which increased SR Ca\(^{2+}\) content after AVP exposure accounts for the increase in Ca\(^{2+}\) sparks and oscillations. RyR opening, the Ca\(^{2+}\) release event responsible for sparks,\textsuperscript{18} can be stimulated by increases in cytoplasmic or SR [Ca\(^{2+}\)].\textsuperscript{12,53} Rapid application of a high concentration of caffeine activates RyR release channels maximally and has been widely used as an indirect measure of SR Ca\(^{2+}\) content.\textsuperscript{27,54,55} The amplitude of caffeine-induced Ca\(^{2+}\) transients was increased both 10 and 90 minutes after AVP removal, suggesting a persistent increase in SR loading (Fig. 4). Given that spontaneous Ca\(^{2+}\) release in the form of sparks is increased under these conditions (Fig. 5), it seems likely that SR Ca\(^{2+}\) uptake must also be increased to account for the increased SR load (Fig. 4). A possible target molecule is phospholamban, which inhibits uptake by the SR Ca\(^{2+}\) ATPase when phosphorylated by PKA.\textsuperscript{27,56} Cyclic nucleotide-dependent increases in Ca\(^{2+}\) store loading, with consequent increases in spontaneous Ca\(^{2+}\) release and activation of large-conductance, Ca\(^{2+}\)-sensitive K\(^+\) channels, are believed to contribute to the action of some vasodilators.\textsuperscript{27,57} However, we believe this is the first demonstration that the cAMP/PKA pathway can persistently increase Ca\(^{2+}\) store content, spark, and oscillation frequency in response to a vasoconstrictor such as AVP.

The AVP-evoked Ca\(^{2+}\) transient was inhibited by a V\(_{1a}\) antagonist (Fig. 5). V\(_{1a}\) receptors are normally coupled via a G protein of the \(G_{q/11}\) type to phospholipase C, stimulating the generation of inositol 1,4,5-trisphosphate (IP\(_3\)).\textsuperscript{38,39} This, in
turn, releases SR Ca\textsuperscript{2+} via IP\textsubscript{3} receptors (IP\textsubscript{3}R\textsubscript{s}). Since IP\textsubscript{3}-mediated Ca\textsuperscript{2+} release can stimulate RyR opening in smooth muscle,\textsuperscript{25,41} these events may underpin the increase in Ca\textsuperscript{2+} sparks and oscillations during acute application of AVP (Fig. 2). The persistent increase in SR store load was also V\textsubscript{1a} dependent (Fig. 5). This part of the response appeared to involve cAMP and PKA, as the response was mimicked by forskolin (Fig. 6) and blocked by RpAMPs (Fig. 7A). RpAMPs did not inhibit the AVP-induced Ca\textsuperscript{2+} transient, indicating that different signaling pathways are responsible for AVP-induced Ca\textsuperscript{2+} release and the persistent increase in store loading. Activation of V\textsubscript{1a} receptors has been shown to stimulate diverse signaling pathways in vascular cell lines.\textsuperscript{42,43} AVP treatment can amplify cAMP production when vascular smooth muscle cells are treated with a \beta-adrenoceptor agonist, although AVP alone does not elevate cAMP.\textsuperscript{44} This effect is blocked by V\textsubscript{1a}, but not V\textsubscript{2} antagonists, and appears to be dependent on the AVP-induced rise in [Ca\textsuperscript{2+}].\textsuperscript{14,45} Some such signal cross-talk may explain the observations in our experiments. Caffeine itself can act as a phosphodiesterase inhibitor and so increase cAMP levels,\textsuperscript{46} but we found no evidence of increased store loading after exposure to caffeine alone (Fig. 4). Smooth muscle responses secondary to stimulation of V\textsubscript{1a} receptors on endothelial cells cannot be completely ruled out.\textsuperscript{37}

It is interesting to speculate on the possible physiological significance of AVP in the regulation of tone in retinal arterioles. How much access circulating AVP would have to retinal myocytes across the endothelial tight junctions of the inner blood–retinal barrier is unclear,\textsuperscript{48} but vasopressin has also been identified within the retinal ganglion cell layer, adjacent to retinal vessels.\textsuperscript{12} Local release of vasopressin may contribute to retinal vascular tone. AVP may act as a cellular switch, increasing vascular resistance and reducing retinal blood flow on going from dark to light. Retinal vasopressin levels vary cyclically with a light/dark cycle and are almost 100% higher in the light.\textsuperscript{11} It is possible, therefore, that increased SR loading and Ca\textsuperscript{2+} release due to retinal AVP decreases retinal blood flow in daylight, when total retinal metabolism and O\textsubscript{2} consumption are reduced.\textsuperscript{13} This remains to be determined experimentally.

References


33. Gyurke I, Gyurke S. Regulation of the cardiac ryanodine receptor channel by luminal Ca2+ involves luminal Ca2+ sensing sites. *Biophys J.* 1998;75:2801–2810.


