The Relationship between Functional Inhibition and Binding for $K_{\text{Ca}2}$ Channel Blockers

David Charles Hammond Benton*, Monique Garbarg, Guy William John Moss
Department of Neuroscience, Physiology and Pharmacology, University College London, London, United Kingdom

Abstract
Small conductance calcium-activated potassium channels ($K_{\text{Ca}2.1,2.2,2.3}$) are blocked with high affinity by both peptide toxins (e.g. apamin) and small molecule blockers (e.g. UCL 1848). In electrophysiological experiments, apamin shows subtype selectivity with $IC_{50}$ of $\sim 100$ pM and $\sim 1$ nM for block $K_{\text{Ca}2.2}$ and $K_{\text{Ca}2.3}$ respectively. In binding studies, however, apamin appears not to discriminate between $K_{\text{Ca}2.2}$ and 2.3 and is reported to have a significantly higher ($\sim 20$–$200$-fold) affinity ($\sim 5$ pM). This discrepancy between binding and block has been suggested to reflect an unusual mode of action of apamin. However, these binding and electrophysiological block experiments have not been conducted in the same ionic conditions, so it is also possible that the discrepancy arises simply because of differences in experimental conditions. We have now examined this latter possibility. Thus, we measured $^{125}\text{I}$-apamin binding to intact HEK 293 cells expressing $K_{\text{Ca}2}$ channels under the same ionic conditions (i.e. normal physiological conditions) that we also used for current block measurements. We find that binding and block experiments agree well if the same ionic conditions are used. Further, the binding of apamin and other blockers showed subtype selectivity when measured in normal physiological solutions (e.g.$^{125}\text{I}$-apamin bound to $K_{\text{Ca}2.2}$ with $K_i$ 91±40 pM and to $K_{\text{Ca}2.3}$ with $K_i$ 711±126 pM, while inhibiting $K_{\text{Ca}2.2}$ current at $IC_{50}$ 103±2 pM). We also examined $K_{\text{Ca}2}$ channel block in Ca$^{2+}$ and Mg$^{2+}$ free solutions that mimic conditions reported in the literature for binding experiments. Under these (non-physiological) conditions the $IC_{50}$ for apamin block of $K_{\text{Ca}2.2}$ was reduced to 20±3 pM. Our results therefore suggest that the apparent discrepancy between blocking and binding reported in the literature can be largely accounted for by the use of non-physiological ionic conditions in binding experiments.

Introduction
Small conductance Ca$^{2+}$-activated potassium channels (SK or $K_{\text{Ca}2}$) are widely expressed in vertebrates and have a role in the gating by Ca$^{2+}$constitutively binds calmodulin, which is responsible for channel $K_{\text{Ca}2}$ subunits lie in the susceptibility of the channels they form to block by peptide toxins and small molecule inhibitors. For example, apamin has been reported to block the current carried by $K_{\text{Ca}2.2}$ channels with an $IC_{50}$ of $\sim 100$ pM while $K_{\text{Ca}2.3}$ channels were less sensitive ($IC_{50}$ $\sim 1$ nM) [11]. Human $K_{\text{Ca}2.1}$ channels, when expressed in mammalian cell lines, are even less sensitive to apamin with an $IC_{50}$ of $\sim 3$–$12$ nM [14,15]. However, a different picture emerges from some direct studies of the binding of $^{125}\text{I}$-apamin to heterologously expressed $K_{\text{Ca}2}$ channels. These suggest that apamin binds to all the $K_{\text{Ca}2.2}$ subtypes with very high affinity (in the low picomolar range) and shows a much smaller degree of selectivity. For example, Finlayson et al. [16] showed that in saturation binding experiments $^{125}\text{I}$-apamin bound with $K_i$ values of 6 pM, 8 pM and 270 pM for $K_{\text{Ca}2.2}$, $K_{\text{Ca}2.3}$ and $K_{\text{Ca}2.1}$ respectively. Similarly, Lamy et al. [17] reported a value of 6 pM for both $K_{\text{Ca}2.2}$ and $K_{\text{Ca}2.3}$. Thus, in these experiments, not only did apamin fail to show appreciable selectivity between $K_{\text{Ca}2.2}$ and $K_{\text{Ca}2.3}$, as seen in functional studies, but the absolute affinity of apamin for all subtypes was much higher ($\sim 20$–$200$-fold) than would have been expected from the concentrations observed to block $K_{\text{Ca}2}$ channels in intact cells. One suggestion is that these
differences reflect the complex mechanism of action of apamin, a view that has quickly gained favour (see Adelman et al. [1] for a review). However, it is possible that such discrepancies simply reflect differing experimental conditions. The aim of the present work was to examine this second possibility by comparing KCa2 channel binding and functional inhibition under near-identical experimental conditions.

**Methods**

**Radio-ligand binding studies with $^{125}$I-apamin**

Radio-ligand binding studies were conducted using HEK 293 cell lines stably expressing KCa2.2 or KCa2.3. The cells were cultured in DMEM supplemented with 10% foetal calf serum, 2 mM L-glutamine, penicillin (200 units ml$^{-1}$) and streptomycin (100 µg ml$^{-1}$) in T500 flasks (Nunc). When confluent, these cells were harvested mechanically (to avoid the use of trypsin) into Ca$^{2+}$/Mg$^{2+}$ free HBSS. The cells were centrifuged at 50 g for 2 min, resuspended in DMEM at a density of approximately 2.5 x 10$^6$ cells ml$^{-1}$ and stored at 4°C until used (2 hr). Cell density was estimated using a haemocytometer.

Routinely, incubations were performed in a total volume of 250 µl comprising 100 µl cell suspension (~250000 cells), 100 µl $^{125}$I-apamin and 50 µl displacing agent or incubation medium. The incubation medium contained (in mM) NaCl 140, KCl 5, MgCl$_2$ 1, CaCl$_2$ 2, glucose 10 and HEPES 10. The pH was adjusted to 7.4 with 1 M NaOH. Non-specific binding was estimated in the presence of 100 nM UCL 1848, a potent KCa2 channel blocker which causes maximal inhibition at this concentration (see Hosseini et al. [18], Benton et al.[13]). Measurements for each test were performed in triplicate. Separation of cells from unbound ligand was achieved by rapid filtration through Whatman GF/B filters pre-treated with 0.3% v/v polyethyleneimine (100 µg ml$^{-1}$) in T500 flasks (Nunc). When confluent, these cells were harvested mechanically (to avoid the use of trypsin) into Ca$^{2+}$/Mg$^{2+}$ free HBSS. The cells were centrifuged at 50 g for 2 min, resuspended in DMEM at a density of approximately 2.5 x 10$^6$ cells ml$^{-1}$ and stored at 4°C until used (2 hr). Cell density was estimated using a haemocytometer.

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Saturation binding experiments

In order to establish a suitable incubation period for equilibrium binding studies we measured the time course of association of $^{125}$I-apamin with 50 µM KCa2 channel expressing cells. It was possible to obtain a straight line: $y = \frac{IC_{50}^{n}}{IC_{50}^{n} + [B]}$ where $y$ is the current in the presence of blocker at concentration [B] expressed as a percentage of control and $n$ is the Hill coefficient.

**Immunohistochemistry**

HEK 293 cells were plated on glass coverslips and transiently transfected with either KCa2.2 and GFP or KCa2.3 together with GFP, as described above. Cultures were then stained using rabbit polyclonal antibodies against KCa2.2 or KCa2.3 as previously described [20]. Briefly, cells were first washed in phosphate buffered saline (PBS, composition (mM): NaCl 136.9, KCl 2.7, Na$_2$HPO$_4$ 9.2, KH$_2$PO$_4$ 1.8, pH to 7.2 with HCl) and fixed in PBS containing 4% paraformaldehyde for 10 min. After rehydr-
**Results**

Wild type HEK 293 cells do not express KCa2 channels

In order to rule out the possibility that our results might be complicated by the endogenous expression of KCa2.2 channels in HEK 293 cells, we performed a number of control experiments (Fig. 1). Firstly, we made patch-clamp recordings from wild type HEK cells in order to examine the endogenous currents. We saw no KCa2-like (voltage-independent) currents but instead saw a small, voltage-dependent current. This endogenous current has been studied by Zhu et al. [21] who concluded that it was predominantly carried by chloride channels. It is therefore, perhaps not surprising that we found it could not be inhibited by 10 nM apamin (Fig. 1B, C). To further confirm our finding we stained cells transiently transfected with GFP and either KCa2.2 or KCa2.3 expressing HEK 293 cells (Fig. 1D). Taken together, these observations show that there is no detectable expression of endogenous KCa2 channels in the HEK293 cell line used in the present study.

Binding of 125I-apamin to KCa2.2 channels in intact HEK 293 cells

We began by studying the binding of 125I-apamin to KCa2.2 expressed in HEK 293 cells suspended in a normal physiological buffer. In order to establish an appropriate incubation time for these experiments we first measured the kinetics of 125I-apamin binding. A typical example of the data obtained is shown in Fig. 2A. The value of $K_i$ for unlabelled apamin obtained from these experiments was 2 pM, similar to the value of $K_i$ for KCa2.2. As is clear from Fig. 1A, antibody staining is visible only in transfected cells (i.e. those expressing GFP). Finally, we were unable to demonstrate any inhibitable binding of 125I-apamin to wild type HEK cells (Fig. 1D). Taken together, these observations show that there is no detectable expression of endogenous KCa2 channels in the HEK293 cell line used in the present study.

Materials

Rat KCa2.2, subcloned into pTracer and a HEK 293 cell line stably expressing KCa2.2 were kindly provided by Professor L. Kaczmarek, Yale University and Professor William Joiner, UCSD. UCL 1684 and UCL 1848 were prepared in the laboratory of Professor C.R. Ganellin, UCL. Tissue culture reagents and Lipofectamine 2000 were purchased from Invitrogen. Apamin, gallamine, dequalinium, horse serum, bovine serum albumen and TRITC labeled goat anti-rabbit IgG were from Sigma. A stable HEK 293 KCa2.3 cell line was created using zeocin selection following transfection with the rat KCa2.3 subcloned into the pCDNA3.1 zeo plasmid (Invitrogen). [125I] mono-iodoapamin (125I-apamin) was supplied by New England Nuclear.

**Figure 1.** KCa2 channels are not expressed in wild type HEK 293 cells. A Immunostaining of HEK 293 cells transiently transfected with GFP and KCa2.2 (a,b,c) or KCa2.3 (d,e,f). GFP expressing cells are visible in a and d. Comparison with the brightfield images in c and f shows that only a proportion of the cells were transfected. Staining for KCa2 channels is shown in b and e where a signal is only visible in cells also expressing GFP. The scale bars indicate 10 μm. B Whole cell currents from a wild type HEK 293 cell recorded using a calcium containing internal solution before (■) and after exposure to 10 nM apamin (○). The endogenous current is not inhibited. C Current voltage relationship for recordings in B. D Binding of 125I-apamin to wild type, KCa2.2 and KCa2.3 expressing HEK 293 cells. Cells were incubated with 30 pM 125I apamin in the absence (T) or presence (N.S.) of 100 nM UCL 1848. There is no significant binding in wild type cells. In contrast, total binding is much higher in cells expressing KCa2 expressing cells and is almost completely inhibited by UCL 1848. doi:10.1371/journal.pone.0073328.g001
shows specific binding of 125I-apamin at times indicated on the graph shows total (constant from 3 separate experiments was 0.24 0.2 well to a single exponential (with apparent rate constant of results from one such experiment. Again the data fit reasonably appropriate incubation times were used. Figure 3A shows the from our saturation binding experiments with KCa2.3-expressing HEK cells are shown in Figure 3B. Fitting of these data provided for 125I-mono-iodoapamin binding to KCa2.2 by unlabelled apamin. Each point is the mean of triplicate observations from a single experiment. Data were fitted to the Hill equation (solid line). Estimates of K_i from saturation binding experiments were used to estimate K_i as described in the methods section and the derived estimates are given in Table 1.

Table 1. Summary of IC_{50} for inhibition of KCa2 current and K_i for displacement of mono-iodoapamin in cells expressing KCa2.2 and KCa2.3.

| Compound      | IC_{50} | K_i | IC_{50}^\alpha | K_i  |
|---------------|---------|-----|----------------|------|
| Apamin        | 112±1 pM| 103±1.9 pM | 1.4±0.2 nM    | 350±83 pM    |
| UCL 1848      | 125±2.5 pM | 95±1.7 pM  | 2.1±0.3 nM    | 652±180 pM   |
| UCL1684       | 530±8 pM  | 1008±22 pM | 5.8±0.3 nM    | 7.7±1.6 nM   |
| Dequalinium   | 420±99 nM | 977±165 nM | 920±140 nM    | 865±160 nM   |
| Gallamine     | 7±0.8 μM  | 26±5 μM    | 98±0.7 μM     | 71±8 μM      |

^Data from Hosseini et al. 2000, using CHO cells.
doi:10.1371/journal.pone.0073328.t001

Inhibition of 125I-apamin binding by other KCa2 channel modulators

We next examined a range of small molecule blockers of KCa2 channels using the 125I-apamin assay, since these have also been reported to have a variety of potencies and even to have different rank orders of selectivity in binding versus block experiments. We thus examined KCa2.2 and then, in a separate experiment, KCa2.3, testing inhibition of 125I-apamin binding by UCL 1848, UCL 1684, dequalinium and gallamine. Inhibition curves for these compounds are shown in Fig. 4A (KCa2.2) and Fig. 4B (KCa2.3). Data for inhibition of 125I-apamin binding could be described by an inhibition curve with a Hill slope of one. Further, within the margins of error for this experiment, it was clear that specific binding of apamin could be completely inhibited. The values of K_i estimated above were used to calculate estimates of K_i for the compounds tested which are summarized in Table 1. The rank order of potency, apamin ~ UCL1848 > UCL1684 > dequalinium > gallamine was the same for both KCa2.2 and KCa2.3. However, all of the compounds showed selectivity for KCa2.2 over KCa2.3.

Inhibition of KCa2.2 current

The same compounds used in binding studies were next assessed for their ability to inhibit KCa2.2 current in a whole cell patch-clamp assay. We were careful to examine block of KCa2.2 current under conditions that were essentially identical to those used in binding. As illustrated in Fig. 5A, HEK cells expressing KCa2.2 exhibited robust, time-independent currents which reversed close to the predicted value of E_{K} (-85 mV) (Fig. 5A,C). In the presence of 10 nM apamin which, based on our binding experiments should be a near saturating concentration, this current was much reduced (Fig. 5A, right panel). Further, the residual current was time-dependent, showed weak outward rectification and reversed at ~40 mV. Thus, following the addition of 10 nM apamin, the only remaining current appeared to be the endogenous current (not itself inhibitable by apamin) (Fig. 5B, Fig. 1C). Thus it appears that apamin causes near-complete block of KCa2.2 current in our assay. Indeed, our results suggest that all of the compounds tested produced >90% inhibition of KCa2.2 current when applied at sufficiently high

Figure 2. Binding of 125I-apamin to KCa2.2. A Time course of 125I-apamin binding to HEK 293 cells stably expressing KCa2.2. The y-axis shows specific binding of 125I-apamin at times indicated on the x-axis. Each point represents the mean of triplicate observations from a single experiment. The data are fitted by a single exponential function (solid line) with an apparent rate constant of 0.3±0.03 min^{-1}. The mean rate constant from 3 separate experiments was 0.24±0.04 min^{-1} suggesting that binding reaches equilibrium well within 10 minutes. B Equilibrium binding of 125I-apamin to HEK 293 cells stably expressing KCa2.2. The graph shows total (■) and non-specific (●) binding in the presence of label. Each point is the mean of triplicate observations from a single experiment. Combined data yielded estimates of K_i and B_{max} of 91±40 pM and 6.4±1.3 fmol/10^6 cells (n = 3). Solid lines represent a linear fit (non-specific binding) or a fit using the modified Hill equation for B_{tot} (see methods). C Inhibition of 125I-mono-iodoapamin binding to KCa2.2 by unlabelled apamin. Each point is the mean of triplicate observations from a single experiment. Data were fitted to the Hill equation (solid line). Estimates of K_i from saturation binding experiments were used to estimate K_i as described in the methods section and the derived estimates are given in Table 1.

doi:10.1371/journal.pone.0073328.g002

Binding of 125I-apamin to KCa2.3 channels in intact HEK 293 cells

We next studied apamin binding to KCa2.3 channels. Again we began by examining the kinetics of binding to ensure that appropriate incubation times were used. Figure 3A shows the results from one such experiment. Again the data fit reasonably well to a single exponential (with apparent rate constant of 0.2±0.05 min^{-1}). Thus, again, a 10 minute incubation period was appropriate in order to ensure that equilibrium was reached. Data from our saturation binding experiments with KCa2.3-expressing HEK cells are shown in Figure 3B. Fitting of these data provided values of 711±226 pM and 175±18 fmol/10^6 cells for K_i and B_{max} respectively. This K_i value is significantly higher than the value we obtained for KCa2.2 and of the same order as the IC_{50} reported for inhibition of KCa2.3 current [11,18]. Finally, we assessed the ability of unlabelled apamin to inhibit mono-iodoapamin binding and found a K_i value of 350±83 pM (Fig. 3C). This is lower than the estimate of K_i for 125I-apamin, but within a factor of approximately two, which may reflect either experimental variability or some modest effect of iodination on apamin binding.
concentrations. IC50 values were obtained by fitting the Hill equation to the concentration-inhibition curves (Fig. 5C). The IC50 values are given in Table 1 for comparison with the 

The effect of changes in ionic composition on channel block

Given the close correspondence between the potency of apamin measured by electrophysiological block and ligand binding assays in identical (physiological) solutions, it seemed important to understand why such disparate values have been reported in the literature. One obvious possibility concerns the different composition of solutions used in previously published binding studies. In binding experiments on isolated membrane preparations it is common to use solutions which have low ionic strength and are free of Ca2+ and Mg2+. We therefore examined the effect of changing our extracellular solution to mimic these differences (Fig. 6A,B,C,D). In our standard bath solution 100 pM reduced KCa2.2 current to 51 ± 5% of control. In contrast, when the bathing solution was free of added Ca2+ and Mg2+ 100 pM apamin reduced the KCa2 current to 20 ± 0.9% of control (p < 0.05 compared with standard). When, in addition to removal of Mg2+ and Ca2+, NaCl was replaced by 280 mM sucrose to mimic even more closely conditions reported in the literature, 100 pM apamin reduced KCa2.2 current to 15 ± 7% of control (p < 0.05 compared with standard solution). The effect of removing Ca2+ and Mg2+ was further examined by constructing a concentration-inhibition curve for apamin in Mg2+ and Ca2+ free conditions (Fig. 6E). The IC50 and Hill slope were 20 ± 3 pM and 0.8 ± 0.06 respectively. Thus, simply removing divalent cations from the bath solution caused an approximately 5-fold increase in potency. These results suggest that the potency of apamin in binding to KCa2.2 channels is increased under ionic conditions similar to those frequently used in binding assays. This may account for a substantial part of the very high affinity 

Discussion

Binding and block for apamin

We find that when apamin binding and apamin block of KCa2.2 channels are measured using the same ionic conditions, the values of K_i and K_L for binding and the value of IC50 for inhibition (block) of current agree well. Whilst previous reports in the literature found differences of up to 20–200-fold, the largest discrepancy we found in comparing binding and block for both KCa2.2 and

Figure 3. Binding of 125I-apamin to KCa2.3. A Time course of 125I-apamin binding to HEK 293 cells stably expressing KCa2.3. The y axis shows specific binding of 125I-apamin at times indicated on the x-axis. Each point represents the mean of triplicate observations from a single experiment. The data are fitted by a single exponential function (solid line) with a rate constant of 0.2 ± 0.05 min⁻¹, suggesting binding reaches equilibrium within 10 minutes. B Equilibrium binding of 125I-apamin to HEK 293 cells stably expressing KCa2.3. The graph shows total (■) and non-specific (●) binding in the presence of label. Each point is the mean of triplicate observations from a single experiment. Combined data yielded estimates of K_L and B_max of 711 ± 226 pM and 175 ± 18 fmol/10⁶ cells. Solid lines represent a linear fit (non-specific binding) or a fit using the modified Hill equation for B_tot (see methods). C Inhibition of 125I-apamin binding to KCa2.3 by unlabelled apamin. Each point is the mean of triplicate observations from a single experiment. Data were fitted to the Hill equation (solid line). Estimates of K_i from saturation binding experiments were used to estimate K_i as described in the methods section and the derived values are given in Table 1. doi:10.1371/journal.pone.0073328.g003

Figure 4. Inhibition of 125I-apamin binding to KCa2.2 and 2.3 by known KCa2 inhibitors. Inhibition of 125I-apamin binding to KCa2.2(A) and KCa2.3(B) by the known KCa2 blockers UCL 1848 (▲), UCL 1684 (●), dequalinium (■) and gallamine (▼). Each point is the mean of triplicate observations from a single experiment. Data were fitted to the Hill equation (solid lines) and the values of IC50 used to estimate K_i as described in Methods. The pooled data are shown in Table 1.

doi:10.1371/journal.pone.0073328.g004

KCa2 Channel Binding and Block

PLOS ONE | www.plosone.org 5 September 2013 | Volume 8 | Issue 9 | e73328
Figure 5. Effects of apamin and known KCa2.2 inhibitors on KCa2.2 current. A Typical current record from a HEK 293 cell transiently transfected with KCa2.2. In control solution (left panel) robust, time-independent currents are seen. Application of 10 nM apamin causes a very substantial reduction (right panel). B Current-voltage relationships for the cell in A in the absence (●) and presence (○) of 10 nM apamin. In the absence of apamin the current reverses at −80 mV, close to the predicted value of $E_K$. In the presence of apamin the residual current is similar to that seen in an untransfected HEK 293 cell (□) suggesting that at this concentration apamin causes nearly complete inhibition of KCa2.2 current. C Concentration-inhibition curves for apamin (●), UCL 1848 (□), UCL 1684 (▲), dequaquium (●) and gallamine (▼). Each point is the mean of 3–5 observations, vertical bars show s.e.m. Fitted lines were drawn from the Hill equation with a value of 3 pM was reported for binding to hepatocyte membranes, causing 50% block in functional experiments. In marked contrast, a value of 390 pM [6], of the same order as the concentration needed to cause 50% block in functional experiments. In marked contrast, a value of 3 pM was reported for binding to hepatocyte membranes.

The IC50 values calculated for each compound are given in Table 1. doi:10.1371/journal.pone.0073328.g005

KCa2.3 values is a factor of only approximately four, and in that case the comparison is to IC50 values using a different cell line (CHO cells) in a study that was not part of the current work. Further, the IC50 values we observed are, overall, very similar to the values obtained by others using electrophysiological approaches to measure inhibition [11,13,15,17,22]. In contrast, previously published values of $K_I$ for the binding of labelled apamin to membrane preparations of KCa2.2-expressing HEK cells have been much lower, of the order of 6 pM [16,17]. Moreover, whilst it is often reported that apamin is significantly more potent as a blocker of KCa2.2 than KCa2.3 (see e.g. Kohler et al.[11]) binding studies on isolated membranes suggest that apamin does not discriminate between the two sub-types [16,17]. In the present study we found that labelled apamin bound to KCa2.3 with a $K_I$ of 71 ± 226 pM and a $K_I$ value of 350 ± 83 pM. Even taking the lowest of these estimates the affinity of apamin is still significantly less for KCa2.3 than KCa2.2 ($K_I$ 90 ± 40 pM, $K_I$ 103 ± 2 pM). This is in keeping with IC50 values from functional experiments (see Table 1).

Other evidence supports our conclusion that the ionic conditions used in binding measurements markedly affect the outcome. For example, in intact guinea pig hepatocytes the $K_I$ for apamin binding in normal physiological solutions was reported as 390 pM [6], of the same order as the concentration needed to cause 50% block in functional experiments. In marked contrast, a value of 3 pM was reported for binding to hepatocyte membranes, an experiment performed in low ionic strength [23]. Similarly, in primary cultures of intact cortical neurones a value of 60 to 120 pM was reported by Seagar et al. [24] in normal physiological solutions, compared with a value of 6 pM for isolated cortical membranes, again at low ionic strength [16]. Finally, Hugues et al. [5] reported a similar discrepancy between binding of labelled apamin to, and inhibition of, the apamin-sensitive AHP in NIE115 neuroblastoma cells. Further work by this group showed that the binding of labelled apamin was inhibited by divalent cations including Mg2+ and Ca2+ and by Na+. In keeping with this observation we have now demonstrated that removal of Ca2+ and Mg2+ caused an increase in the blocking action of apamin. In experiments on isolated membranes it is standard practice to use a low ionic strength solution (e.g., 10 mM Tris, 5.4 mM KCl, pH 7.5) as employed by Lamy et al. [17]. Taken together, the evidence now available suggests that ionic environment is highly important in determining the affinity of apamin for its binding site. Whether

Figure 6. The effect of the ionic composition of bathing solution on the potency of apamin. Representative recordings of KCa2 current in HEK 293 cells transiently transfected with KCa2.2. The recordings were made in standard solution (A), nominally Ca2+- and Mg2+ free solution (B) and Na+, Ca2+ and Mg2+ free solution in which NaCl was replaced by 280 mM sucrose (C). In each case the left panel shows currents recorded before and the right panel after the application of 100 pM apamin. D Summary of data for the inhibition caused by 100 pM apamin in the solutions used in A, B and C. Vertical bars show s.e.m. * denotes that the values were significantly different from the effect of apamin in standard solution (p<0.05, unpaired t-test).

E Concentration-inhibition curve for apamin applied in Ca2+- and Mg2+ free solution. Each point is the mean of 3–5 observations with vertical bars indicating s.e.m. The solid line was drawn from a fit of the Hill equation with IC50 = 20 ± 3 pM and n = 0.8 ± 0.06. The dashed line represents the concentration-inhibition curve for apamin in standard solution shown in Figure 4. doi:10.1371/journal.pone.0073328.g006
this is due to charge screening or other changes to the channel-toxin interaction remains to be determined. Nor is it possible to exclude the possibility that the conformation and charge distribution in apamin itself changes in a low ionic strength solution devoid of Ca\(^{2+}\), Mg\(^{2+}\) and Na\(^{+}\). It is interesting in relation to this to note that binding/activity discrepancy seen under such experimental conditions is not observed with all blockers e.g. UCL 1684 [25], even though this compound carries two positive charges to mimic the charges that are thought to be important in apamin binding.

It is also interesting to compare the values of $K_i$ for the binding of apamin obtained in this study with values for the $IC_{50}$ obtained in preparations natively expressing KCa2.2 channels of known subtype. KCa2.2 has been shown to underlie the Ca\(^{2+}\)-activated K\(^{+}\) current seen in Jurkat T cells [26] and these native channels are blocked by apamin with an $IC_{50}$ of 300 pM [27], a value that is much higher than would be expected from membrane binding experiments performed in non-physiological ionic conditions. Similarly, rat sympathetic ganglion neurones and dorsal vagal neurones display a post-spike after-hyperpolarisation which is known to be mediated by KCa2.3 and is inhibited by apamin with an $IC_{50}$ of ~2 nM [18,28], again a value that is much higher than expected from binding studies of isolated membranes. However, in both of these cases the values are similar to those obtained from functional studies of heterologously expressed channels and from the values of $K_i$ obtained in binding experiments (using physiological solutions) in the present study. Further, looking at the different values for native KCa2.2 and KCa2.3 channels it appears that subtype selectivity is also observed in this setting and may thus be useful for elucidating the function of KCa2 channels in the future.

Investigation of KCa2 pharmacology by ligand binding and electrophysiology

We also compared the potency of some known KCa2 channel blockers in electrophysiological and binding inhibition experiments. With cells expressing KCa2.2 there was good agreement between $K_i$ and $IC_{50}$ values for all the compounds tested (UCL1848, UCL 1684, dequalinium and gallamine; Table 1). The largest difference between $K_i$ and $IC_{50}$ was a factor 3.7 for gallamine, the least active of the set and therefore the most likely to show non-specific actions. Similar agreement was seen with KCa2.3 (Table 1). The implication is that provided the binding studies are done using bathing fluids with ionic composition not far removed from physiological, they can give a good prediction of biological activity.

Implications for interpreting the mechanism of apamin block of KCa2 channels

Early experiments suggested that apamin might simply act as a pore blocker for KCa2 channels because TEA (a known small molecule pore blocker), prevented the binding of apamin [16,29]. Nonetheless, mutation studies have shown that interactions of apamin with the channel involve a number of amino acid residues, some of which are outside the central pore region [22,30,31]. This and other evidence has led to the suggestion that apamin does not act as a ‘classical’ pore blocker but instead inhibits channel opening by an allosteric mechanism [17,32]. The discrepancy between the $K_i$ for binding of apamin (measured in low ionic strength) and $IC_{50}$ for inhibition of KCa2.2 current (measured in normal physiological solutions) has been cited as evidence in support of this idea. Our findings suggest that caution must be exercised when making this kind of comparison. In particular, it is clear that ligand binding experiments agree much more closely with functional experiments when conducted under identical physiological conditions. Of course, agreement between $IC_{50}$ and $K_i$ values does not exclude an allosteric mode of action and indeed it is quite difficult to imagine an allosteric mechanism that does not predict close accord in the concentration dependence of both binding and block (otherwise block would still be able to increase when binding had already saturated). On the other hand, some allosteric mechanisms would predict incomplete block of current even at high concentrations of apamin. We see no clear evidence of this, in keeping with a number of other reports [11,13,22]. Nonetheless, two groups do report partial block [17,33] so that more work is needed to clarify this aspect of the action of apamin.

In summary, we have found that there is a close relationship between the binding and block of KCa2 channels for both apamin and other small molecule blockers when these are measured under identical conditions. While this observation is consistent with both allosteric and non-allosteric models it suggests that caution must be exercised when comparing electrophysiological block data obtained in normal (physiological) conditions with binding data obtained in non-physiological conditions. Our findings also suggest that more work may be needed to establish precisely the mechanism of action of apamin and other compounds known to block KCa2 channels.

Acknowledgments

We are grateful to Prof. D.H. Jenkinson and Prof. C.R. Ganellin for their help in the preparation of this manuscript.

Author Contributions

Conceived and designed the experiments: DCHB MG GWJM. Performed the experiments: DCHB MG. Analyzed the data: DCHB MG. Wrote the paper: DCHB GWJM.

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