Kv7/M-type potassium channels in rat skin keratinocytes

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Abstract Skin keratinocytes fulfil important signalling and protective functions. Immunocytochemical experiments revealed the unexpected presence of immunoreactivity for the M-type potassium channel subunit Kv7.2 in the keratinocyte layer of intact rat paw skin and in keratinocytes isolated from the skin of 1-day-old rats and cultured in vitro for 3–10 days. Application of the M-channel enhancer retigabine (3–10 μM) to isolated cultured rat keratinocytes: (a) increased outward membrane currents recorded under voltage clamp, (b) produced ~3 mV hyperpolarization at rest, (c) enhanced ~3-fold the release of ATP induced by the TRPV3 agonist carvacrol (1 mM) and (d) increased the amplitude of the carvacrol-induced intracellular Ca2+ transient measured with Fura-2. The effect of retigabine on ATP release was prevented by the M-channel blocking agent XE991. We conclude that rat skin keratinocytes possess M-channels that, when activated, can modify their physiological properties, with potential significance for their sensory and other biological functions.

Keywords Potassium channels · Keratinocytes · ATP release · Immunocytochemistry · Membrane currents · Calcium

Introduction

Skin keratinocytes express several types of ion channel, which are variably involved in the secretion of signalling molecules, keratinocyte differentiation and the regulation of skin permeability. These include voltage-gated Na+ channels [48]; epithelial ENaC (SCNN1) Na+ channels [7]; L-type (α1C) voltage-gated Ca2+ channels [11]; a large-conductance, Ca2+-independent K+ channel [19]; large, intermediate and small conductance Ca2+-activated K+ channels (BKCa [19, 40], hIK1 [22], hSK4 [21]); Kir6.1, Kir6.2 and SUR ATP-gated inward rectifier K+ channel subunits [8]; several twin-pore K+ channels, principally TREK-1, TREK-2 and TRAAK [20] and several transient receptor potential (TRP) cation channels [25], including TRPV1 [18], TRPV3 [35] and TRPV4 [29], TRPA1 [1, 4] and several TRPC channels [3, 16].

In the course of a recent study on the location and function of M-type Kv7 K+ channels in nociceptive sensory fibre terminals in the hairy skin of the rat paw [34], we noted that immunoreactivity for the Kv7.2 subunit was not restricted to the nerve fibres but could also be detected in the inner layer of the epidermis (J. Reilly, unpublished). In the present paper, we show that this results from its expression in skin keratinocytes and that these channels, which are normally associated with neurons, can play a functional role in modulating the stimulated release of ATP from the keratinocytes.

Methods

Keratinocyte preparation and culture

Primary epidermal keratinocytes were cultured according to the methods described by Dlugosz et al [15]. One-day-old rats were killed in accordance with schedule 1 of the UK Animals (Scientific Procedures) Act 1986. The skin from the back and abdomen was removed and sequentially
washed for 2 min in 0.01 M phosphate-buffered saline (PBS; Sigma UK)/1 % antibiotic–antimycotic (100×) liquid (Invitrogen, UK), 70 % ethanol and 0.01 M PBS (Sigma UK)/1 % antibiotic–antimycotic (100×) liquid (Invitrogen, UK). The subcutaneous fat was removed by gently scraping. The skin was floated dermis side down in trypsin 0.25 %–EDTA solution (Invitrogen, UK) overnight at 4 °C. The following day residual dermal tissue was removed by gently scraping. The epidermis was sliced into sections, placed into keratinocyte serum-free media (keratinocyte SFM) (Invitrogen, UK) supplemented with 10 % foetal bovine serum (Invitrogen, UK) and 1 % antibiotic–antimycotic (100×) liquid (Invitrogen, UK) and stirred for 30 min at room temperature. The keratinocyte suspension was passed through a nylon mesh cell strainer (70 μm; BD Falcon, USA) and the keratinocytes harvested by centrifugation at 168×g for 5 min. The resultant cell pellet was re-suspended in keratinocyte SFM supplemented with 1 % antibiotic–antimycotic (100×) liquid and plated equally onto Petri dishes or glass coverslips. The medium was replaced after 15 h with keratinocyte growth SFM and every 48 h subsequently. Keratinocytes were maintained for up to 3 weeks or until confluence was reached. Older rats (up to P7) gave much poorer yields of confluent keratinocytes.

Immunohistochemistry

**Intact skin** Sprague-Dawley rats (~150 g) were deeply anaesthetised with ketamine/xylazine hydrochloride solution (1 ml/kg, i.p., Sigma, UK) in accordance with schedule 1 of the Animals (Scientific Procedures) Act 1986 and transcardially perfused with ice-cold heparinised saline (9 % w/v NaCl (VWR, UK), 50,000 U/l heparin (Sigma, UK)) and then 4 % paraformaldehyde (EMS Diasum, USA) in PBS (Sigma, UK). The skin from either the hairy dorsal or plantar surface of the hind paw was removed, post-fixed for 3 h with 4 % paraformaldehyde in PBS, rinsed with PBS and then cryo-protected via immersion overnight in a 30 % sucrose (VWR, UK)/0.01 % Na azide (Sigma, UK)/PBS solution. Cryosections (40 μm) were obtained using a freezing microtome (Leica Microsystems, Germany) and free floating sections placed into PBS. Sections were incubated with blocking solution (PBS containing 3 % goat serum and 3 % Triton® X-100 (Sigma, UK)) followed by incubation overnight at 4 °C with primary antibody (see below). The sections were rinsed then incubated with a fluorophore-conjugated secondary antibody (Invitrogen, UK) for monoclonal antibodies, or a biotinylated secondary antibody (Sigma, UK) then with Cy3-conjugated streptavidin (Sigma, UK) for detecting polyclonal antibodies.

**Cultured keratinocytes** Cultured cells on cover slips were washed twice with 0.01 M PBS and then fixed in 4 % paraformaldehyde in PBS for 15 min. After fixation, the cells were washed (3×) for 10 min and then incubated for 1 h at room temperature with the above blocking solution or one containing 1 % (w/v) bovine serum albumin, plus 0.3 % (v/v) Triton X-100™ or 0.3 % TWEEN®20 (Sigma, UK). Cells were incubated overnight at 4 °C with the primary antibodies then treated with secondary antibodies or biotin-streptavidin as described above for tissue sections.

Staining of skin sections and cultured keratinocytes was visualised using a Leica CA TCS SP2 AOBS spectral confocal microscope (Leica, Germany). The gain, exposure time and wavelength were optimised for the excitation and emission spectra of each fluorophore. Control experiments performed without primary antibodies did not show labeling above background.

**Antibodies** The following primary antibodies, with final dilutions, were used: Alomone Kv7.2 rabbit polyclonal against rat Kv7.2 (epitope residues 578–593), 0.7 μg/ml (Alomone, Israel); Abcam Kv7.2 rabbit polyclonal against human Kv7.2 (epitope residues 850-end of C-terminus), 3 μg/ml (Abcam, Cambridge, UK) and Neuromab mouse Kv7.2 monoclonal against human Kv7.2 (1–50), 1:100 (Neuromab, Davis, CA, USA). The following secondary antibodies were used: biotinylated goat anti-rabbit (Vector Laboratories, UK), 1:333; biotinylated goat anti-mouse (Vector Laboratories, UK) goat anti-mouse rhodamine (Invitrogen, UK), 1:100 and goat anti-rabbit Rhodamine Red™-X (Invitrogen, UK), 1:100. In parallel experiments, the antibodies were tested in cultured rat sympathetic neurons [17]; in these, the specificity of the antibodies for the Kv7.2 protein was verified by adding a 10-fold excess of the appropriate epitope (where available) 1 h before adding the primary antibody (Supplementary Fig. S1). To assess secondary antibody specificity, in each keratinocyte experimental set a control was included in which the primary antibody was omitted.

**Electrophysiology**

Macroscopic currents and membrane potentials in isolated cultured keratinocytes were recorded using the voltage clamp perforated-patch technique as described in [38, 43]. The bath solution was N-2-hydroxyethylpiperazine-N′-2-ethanesulfonic acid (HEPES)-buffered Krebs’ solution containing (in millimolar): 144 NaCl, 2.5 KCl, 0.5 MgCl₂, 2 CaCl₂, 10 d-glucose and 5 HEPES; pH was adjusted to 7.4 using Trizma base. The pipette solution contained (in millimolars): 30 KCl, 80 K-acetate, 40 HEPES, 3 MgCl₂, 1 CaCl₂ and 3 ethylene glycol tetraacetic acid; free [Ca²⁺], was adjusted to 100 nM; pH was adjusted to 7.2 with KOH. Amphotericin B 300 μg/μl was used as a perforating agent. Pipette resistances were 5–10 MΩ when filled with the
pipette solution. Seal resistances were >1 GΩ. Series resistance was compensated 60–90 %. The mean capacitance of the cells studied was 15.6±1.9 pF \((n=9)\). All experiments were performed at a controlled room temperature \((22±0.5 ^\circ C)\). All recordings were made using an Axopatch 200A amplifier and Digidata 1440 A/D interface (Axon Instruments, Forster City, CA, USA) and with a pipette holder optimized for low noise recordings (G23 Instruments, UCL). All recordings were filtered with an eight-pole Bessel filter at 5 kHz and digitized at 10 kHz. For recording macroscopic currents in perforated-patch configuration, cells were held at −60 mV and commanded to voltages from −100 to +50 mV in ramps of 25 s duration.

ATP release

ATP release from the keratinocytes was measured using a modification of the standard luciferase reaction [30]. Keratinocytes were cultured until they reached confluence (1–3 weeks). The keratinocyte growth SFM was replaced with HEPES-buffered Krebs solution (composition as for electrophysiological experiments) and incubated for 1 h at room temperature \((-21–25 ^\circ C)\) so that any ATP released due to mechanical disruption or temperature change would be metabolised. Drugs were applied for 10 min. An equivalent volume of HEPES-buffered Krebs was added to control dishes to control for experimental errors resulting from possible mechanical disturbance to the keratinocytes.

ATP release was determined by adding a 50-μl sample of supernatant to an equal volume of CellTiter-Glo® Reagent (Promega, USA). This was vortexed and equilibrated at room temperature for 10 min. The sample was transferred to a disposable cuvette \((8 \times 50 \text{ mm})\), and light emission was measured in a luminometer (Model 20E, Promega, USA). A standard curve was constructed using known concentrations of ATP. The luminometer reading was linearly related to the ATP concentration over the range 0 to 3 μM, with a slope of 1.56 U output/nM (ATP). This was used to convert luminometer reading to concentration of ATP in 50 μl keratinocyte supernatant. We did not convert concentrations to amounts released since we do not know what fraction of released ATP might or might not have been metabolized and whether this was a constant fraction or not.

Calcium imaging

Intracellular free Ca\(^{2+}\) was monitored using the cell permeable acetoxymethyl (AM) ester of Fura-2 (Fura-2 AM, Invitrogen, UK). The keratinocytes were incubated for 1 h at room temperature (22–25 °C) with 5 μM Fura-2 AM (final concentration) dissolved in DMSO with 10 % pluronic acid (Invitrogen, UK) and then perfused for 20 min prior to recording with HEPES-buffered Krebs containing 0.1 % (w/v) sulfipyrazone (to minimize loss of Fura-2; Sigma, UK). Cells were visualised on a Nikon Diaphot 300 inverted phase contrast microscope using an oil immersion \(x40\) fluorescent objective (Nikon Instruments, USA). Cells were excited at 350 and 380 nm from a xenon short arc lamp (XBO; Osram, UK) within a Polychrome II monochromator (TILL Photonics, Germany), controlled by Openlab 3.1 Image acquisition software (Improvision, UK) on an Apple Mac G4 computer. Excitation light intensity was regulated using quartz neutral density filters to reduce photo-bleaching of the fluorescent probe. Emitted light was passed via a dichroic mirror (400 nm) through a bandpass filter (520±20 nm) before being redirected to a 12-bit greyscale Hamamatsu C4880-80 camera (Hamamatsu, Japan). A Hamamatsu CCD camera controller enabled the control of exposure times and facilitated image capture. Suitable pseudo-colour palettes were overlaid onto the images to enhance visualisation. To minimise photo bleaching, experiments were performed in a dark room, and exposure times were kept to a minimum. Images acquired at 350 and 380 nm excitations were subjected to a ratiometric algorithm undertaken following background light subtraction at each excitation wavelength. For each cell, an area of cytoplasm was selected from which the changes in mean 350:380 ratio with time were plotted as a two-dimensional graph.

Results

Immunohistochemistry

**Intact skin** Intact hairy skin from the dorsal rat paw showed strong immunoreactive staining for Kv7.2 within the epidermal layer (Fig. 1 a) and around the hair follicles (Fig. 1 b). In plantar paw skin staining of Kv7.2 was apparent in the layers adjacent to the dermal–epidermal border, in the *stratum germinativum* and *stratum spinosum* (Fig. 2 a). Here, staining of individual keratinocytes could be clearly seen (Fig. 2 b). No staining was observed when the primary antibody was omitted (Fig. 2 c). Kv7.2 staining density in single keratinocytes was comparable to that in neighbouring peripheral nerve fibres (Supplementary Fig. S2).

**Cultured keratinocytes** Kv7.2 immunoreactivity was also observed in isolated keratinocytes cultured from the back skin of 1-day-old rats. Figure 3 shows immunoreactivity detected with two different antibodies, a polyclonal antibody (Abcam) and a monoclonal antibody (Neutromab). Immunoreactivity increased to a maximum after 5 days in culture (rows b and c). No immunoreactivity was observed in the absence of a primary antibody (row d). The transmitted light images of the keratinocytes (columns 2 and 4) showed similar morphology to those cultured to similar densities in other studies [24].
Membrane currents

Kv7.2 protein is a subunit of the M-channel [17, 44], a low-threshold, non-inactivating voltage-gated K⁺ channel [10]. To test for the presence of an M-current in keratinocytes, we recorded membrane currents under perforated-patch conditions from isolated cultured cells which had first been trypsinized then replated to minimize electrical coupling (see “Methods”). The experimental protocol we adopted is shown in Fig. 4a. We applied voltage ramps from −100 to +50 mV at two different speeds, fast (500 ms) and slow (25 s). With the fast ramp, an initial inward current appeared between −40 and −10 mV, followed by a large outward current (Fig. 4b). Using the slow ramp, the initial inward current was replaced by an outward current ‘hump’, followed by a more sustained outward current but of much reduced amplitude. This indicates a substantial inactivating component of the outward current, as shown by the response to the 3-s voltage step in Fig. 4a. The inward current and much of the outward current (including most of the ‘hump’ current) was inhibited by 100 μM La³⁺ (Fig. 4b2). This suggests that (a) the inward current was carried by a Ca²⁺-permeable cation channel such as a TRPV channel [32] (see “Introduction” and below) and (b) the ‘hump’ current and much of the current activated at more positive potentials was generated by Ca²⁺-activated K⁺ channels [21, 27, 28], with a possible contribution of Ca²⁺-activated Cl⁻ channels [21] (since ECl was around −35 mV).

Since M-currents are non-inactivating, we sought evidence for an M-current contribution to the outward currents by recording the effects of the M-channel enhancer, retigabine [42] and the M-channel blocker XE991 [44] on the current generated by the slow (25 s) voltage ramp (Fig. 5). Retigabine produced a small increase in mean outward current at depolarized potentials such that the average current amplitude measured at +50 mV was significantly greater than that of the control current or that recorded in the presence of XE991 (Fig. 5, inset). In contrast, mean outward currents recorded in the absence and presence of XE991 were not significantly different.

The variability in current amplitudes between cells might obscure otherwise consistent but small effects of the M-channel modulators. Therefore, since we used the same schedule of drug applications in each experiment (control–retigabine–XE991), we carried out a within-experiment comparison by measuring the current ratios at +50 mV retigabine/control, XE991/control and XE991/retigabine and

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Fig. 1  Kv7.2 antibody staining within keratinocytes in intact rat skin. Left side images fluorescence; right side transmitted light images. a Keratinocytes immunoreactive to Kv7.2 (Alomone) within the outer epidermal layer. b Keratinocytes immunoreactive to Kv7.2 (Abcam) within the epidermal layer surrounding the hair follicle. The arrows in the transmitted light images point to hair follicles. Scale bar 40 μm

Fig. 2  Immunoreactivity for Kv7.2 within the epidermis of plantar skin. a, b Kv7.2(Alomone) immunoreactivity within rat plantar skin with corresponding transmitted light images on the right. c Secondary only antibody. Scale=40 μm
assessing whether their means differed significantly from a null-hypothesis ratio of 1.0 (Table 1). The ratios retigabine/control and XE991/retigabine were significantly different from unity at $P<0.025$ and $P<0.0025$, respectively; that for the ratio XE991/control was marginally outside the 5 % significance level at $P=0.052$. Thus, it would seem likely that functional M-channels (presumably containing Kv7.2 subunits) are present in these keratinocyte membranes, contributing perhaps a quarter of the outward current at positive potentials, and that this current is enhanced ~20 % in the presence of retigabine.

Membrane potential

Several attempts were made to record the effects of M-channel modulators on the resting membrane potential of the cultured keratinocytes. In most (5+) cells successfully patched, the apparent resting potential was regarded as too positive ($\leq -25 \text{ mV}$) and/or too unstable, and recordings aborted or rejected. However, one cell showed a good and relatively stable resting potential (~75 mV; Fig. 6). In this cell, retigabine (10 μM) produced a clear hyperpolarization of ~3 mV, which was partly reversed on adding 3 mM XE991 and fully recovered on washing. This accords with the effect of retigabine on nerve cells, in which it produces a resting hyperpolarization by shifting the M-channel activation curve so that channels now open at negative potentials where they would normally be closed [42].

ATP release

To assess the maximum ATP-releasing capacity of cultured keratinocytes, we applied 50 μl NaOH to a series of keratinocyte cultures for 10 min to give a final bath concentration of 4 % [30]. This increased the ATP concentration in the surrounding medium by about 10-fold over that recorded on simply adding an aliquot of Krebs’ solution, from a mean of 34.6±5.2 nM ($n=5$) to 420±49.5 nM ($n=6$) (Fig. 7a).

To provide a more physiological stimulus for ATP release, we used the TRPV3 agonist carvacrol [46]. TRPV3 is

| Kv7.2 (Abcam) | Kv7.2 (Neuromab) |
|--------------|-----------------|
| Fluorescence | Transm. light   | Fluorescence | Transm. light |
| a            |                 | a            |                 |
| b            |                 | b            |                 |
| c            |                 | c            |                 |
| d            |                 | d            |                 |

$Scale \ bar=20 \mu M$
Fig. 4 Membrane currents recorded from an isolated cultured rat keratinocyte. a Voltage and current protocol (upper trace current, lower trace voltage). The cell was held at −60 mV, prepulsed to −100 mV and then commanded to +50 mV with a 25-s voltage ramp. After returning to −60 mV, two 3-s commands to −20 and +20 mV were applied. Finally, the voltage-ramp procedure was re-applied but at a faster speed (500 ms). Currents obtained before (black) and after (grey) adding 100 μM LaCl3 are shown. The zero current level in the upper (current) trace is indicated by the dashed line.

b Current–voltage relations for 500 ms (grey) and 25 s (black) ramps (1) before and (2) after adding LaCl3. Currents were leak-subtracted by subtracting extrapolated linear currents from −100 to −60 mV. Current scale pA/pF

Fig. 5 Superimposed averaged mean currents (pA/pF) generated by 25 s voltage ramps from −100 to +50 mV in 9 keratinocytes exposed sequentially to Krebs’ solution (control, black), 10 μM retigabine (amber) and 10 μM XE991 (violet). 100 μM LaCl2 was subsequently added to 7 of the same 9 cells (brown). Currents were leak-subtracted by subtracting extrapolated linear currents from −100 to −60 mV. Inset means±SEMs of current amplitudes at +50 mV. C control, R retigabine, X XE991. Means±SEMs (pA/pF) were: control, 1.06±0.23 (n=9); retigabine, 1.35±0.33 (n=9); XE991, 0.94±0.33 (n=9); La3+, 0.40±0.26 (n=7). For the bracketed columns, P values (two-tailed t test) were: * <0.05; ** <0.03
abundantly expressed in keratinocytes [25, 31], and its activation readily induces ATP release [26]. As shown in Fig. 7b, 1 mM carvacrol for 10 min produced a consistent and significant ($P < 0.01$) increase in ATP release of >2-fold over the 10-min exposure period, from 30.4±3.5 nM ($n=4$) with Krebs’ solution to 70.9±3.5 nM ($n=4$).

Addition of the M-channel enhancer retigabine (3 μM final bath concentration) to the carvacrol solution led to a further 3-fold increase in the amount of ATP released by carvacrol, from 191±31.8 nM ($n=6$) to 616±116 nM ($n=5$) (Fig 7c). Ten micromolars retigabine produced a similar effect (Fig 7d). This latter effect was annulled by prior addition of the M-channel blocking agent XE991 (10 μM; Fig. 7d) so presumably results from activation of M-channels. Retigabine (10 μM) did not affect spontaneous ATP release in the absence of carvacrol ([ATP]: controls 15.2±5.9 nM, $n=4$); retigabine 14.3±7.8 nM, $n=3$), nor did addition of 50 μl of the solvent (5 % DMSO) ([ATP]: controls 15.1±8.6 nM, $n=4$; DMSO 12.0±3.8 nM, $n=4$)). XE991 (10 μM) in the absence of retigabine had no significant effect on either the spontaneous release ([ATP]: controls 73.3±43.7 nM, $n=4$); XE991 98.4±19 nM, $n=3$) or the enhanced release produced by carvacrol ([ATP]: carvacrol 203.3±32.9 nM, $n=4$; carvacrol+XE991 359.8±84.5 nM, $n=4$).

Intracellular Ca2+ changes

Carvacrol has been reported to increase intracellular [Ca2+] in TRPV3-containing or expressing cells [46]. Since that increase required extracellular Ca2+, it presumably resulted from entry through the Ca2+-permeable TRPV3 channels [45]. In accordance with this, carvacrol increased intracellular [Ca2+] in cultured keratinocytes as measured from the 350:380 fluorescence ratio in Fura-2 loaded cells (Fig. 8). Fura-2 signals across different keratinocytes were highly variable, but signals with repeated carvacrol applications to the same keratinocyte were reasonably consistent, albeit with some desensitization as previously reported [46]. Hence, to test whether retigabine affected the Ca2+ signal, we applied carvacrol twice to the same cell then measured the ratio of the second to first responses (R2/R1) on adding either Krebs’ solution or 10 μM retigabine 5 min before the second response (Fig. 8a). As shown in Fig. 8b, the second response was smaller on average on adding Krebs solution (R2/R1=0.73±0.10, $n=33$) but was consistently increased after adding 10 μM retigabine (R2/R1=1.18±0.04, $n=24$); the difference between retigabine and Krebs’ solution was highly significant ($P<0.01$). Retigabine itself did not produce any consistent change in resting [Ca2+]..

Discussion

The main points emerging from these experiments are that the M-channel Kv7.2 subunit protein is consistently expressed in rat skin epidermal keratinocytes and that these cells also possess functional (or potentially functional) membrane M-channels. Antibodies to Kv7.2 were selected as most helpful in seeking the presence of M-channel proteins because this subunit is not only a component of the heteromeric Kv7.2/7.3 M-channel observed in peripheral neurons [17, 44] (and indeed is essential for their functional expression [33]) but also because Kv7.2 subunits can form functional M-channels in the absence of other subunits [37–39]. Using Kv7.2 antibodies, we found a strong band of immunoreactivity in the keratinocytes in the epidermis and around hair follicles in intact rat skin and in keratinocytes from early postnatal rat skin when cultured for a few days in vitro. This is likely to be due to specific staining of Kv7.2 protein since (a) it was replicated by three antibodies directed at different epitopes and (b) where tested (supplementary Fig. S1), it was prevented by the subunit-specific epitope peptide.

Our primary evidence for functionality (or at least, potential functionality) of the Kv7.2 protein as a subunit of the M-channel rests on the effects of the selective M-channel enhancer retigabine. This drug (a) produced a small (~20 %) but consistent increase in outward membrane current, (b)

Table 1 Within-experiment measurements of current ratios at +50 mV from the experiments illustrated in Fig. 6

| Retigabine/ control | XE991/ control | XE991/ retigabine |
|---------------------|----------------|-----------------|
| Mean ratio          | 1.208          | 0.730           | 0.583           |
| SEM                 | 0.0896         | 0.147           | 0.107           |
| N                   | 9              | 9               | 9               |
| P (difference from 1.0, one tail) | 0.024          | 0.052           | 0.0022          |
hyperpolarized a keratinocyte with a high (−75 mV) resting potential, (c) augmented the release of ATP by the TRPV3 agonist carvacrol and (d) enhanced the increase in intracellular Ca²⁺ produced by carvacrol. The enhanced release of ATP by retigabine was suppressed by the M-channel blocking drug XE991, showing that this effect at least was unambiguously due to the action of retigabine on M-channels. On the other hand, XE991 alone did not produce a clearly significant (P=0.052) reduction in voltage-activated membrane current (which was dominated by Ca²⁺-activated outward currents) and did not significantly affect either the stimulated or resting release of ATP. This may suggest that, in these cultured cells at least, the density of functional M-channels is rather low compared with that in peripheral neurons and/or that very few of these voltage-gated channels are activated at the membrane potential of these cells and only carry appreciable current when their activation is enhanced by retigabine. On the other hand, the channels might well be present at a higher density in adult keratinocytes than in cultured immature keratinocytes and then might form a more substantial component of resting membrane current, since Kv7.2 immunoreactivity was appreciably stronger in adult rat skin keratinocytes than in the cultured cells and indeed comparable to that in neighbouring nerve fibres (Supplementary Fig. S2).

TRPV3 channels are Ca²⁺-permeable [45], and as reported previously for other TRPV3-expressing cells [46], their activation by carvacrol increased intracellular Ca²⁺ in our cultured keratinocytes (Fig. 8). Since raising intracellular Ca²⁺ in keratinocytes can release ATP [23], as in some
other cells (e.g. [6, 9, 23]), it seems reasonable to suppose that the rise in \([Ca^{2+}]_{in}\) is primarily responsible for the release of ATP by carvacrol, though other mechanisms for release are possible [2, 5]. Retigabine produced a significant increase in the carvacrol-induced rise in \([Ca^{2+}]_{in}\) though whether this is quantitatively sufficient to explain the increased ATP release is uncertain.

Likewise, the mechanism whereby increasing M-channel activity with retigabine increases the carvacrol-induced Ca\(^{2+}\) transient and ATP release is, as yet, equally unclear. One possibility is that an M-current-induced hyperpolarization increases the driving force for Ca\(^{2+}\) entry through the TRPV3 channels. The hyperpolarization at rest potential was rather small (~3 mV; Fig. 6) but since the channels are voltage-sensitive, the retigabine-enhanced channels may have a much more significant effect in reducing the TRPV3-induced depolarization (essentially clamping the membrane potential near to the resting potential) and hence increasing the Ca\(^{2+}\) driving force at the peak of TRPV3 activation. Additionally, the activated M-channel might have some more direct effect on the TRPV3 channel to enhance its activity or sensitivity to carvacrol. Thus, Kv7.2 and TRPV1 have been reported to co-precipitate when co-expressed in HEK 293 cells, though in this case without effect of retigabine on the action of the TRPV1 ligand, capsaicin [47]. Further tests for Kv7.2-TRPV3 interaction might be worthwhile.

Physiologically, TRPV3 (and TRPV4) channels in keratinocytes are suggested to mediate keratinocyte thermosensation, probably through the release of ATP and activation of subjacent sensory fibres [25]. Thus, in cocultures of dorsal root ganglion neurons with keratinocytes, the response of the neurons to an applied heat ramp has been shown to result from activation of the keratinocyte TRPV3 channels and the release of ATP onto the adjacent neurons, rather than to a direct effect on the neurons [26]. Assuming that retigabine has a comparable effect on the response of keratinocyte TRPV3 channels to heat as that to carvacrol observed in the present study, retigabine would be expected to enhance skin thermosensation. Thus, the fact that retigabine exerted the opposite (suppressant) effect on thermally-induced skin peripheral afferent discharges in situ [34] would seem to rule out the possibility that the latter effects were mediated indirectly via the keratinocytes. On the other hand, the effects of retigabine on keratinocyte responses might imply some limitation to the prospective utility of retigabine in suppressing pathological nociceptive activity [36].

Apart from thermal sensation, keratinocyte TRPV3 channels and local ATP release may be involved in other aspects of skin function including vasodilator responses and the regulation of skin structure and permeability [12, 25]. For example, openers of some other K\(^+\) channels have been reported to accelerate epidermal barrier recovery after barrier disruption [13]. Keratinocytes also possess Gq-coupled receptors such as purinergic P2Y receptors [14] and bradykinin B2 receptors [41], which would be expected to modulate M-channel activity [10]. Thus, the presence of M-channels and their effects noted in the present paper may have rather broad implications for skin biology and possibly some therapeutic implications for wound healing.

**Fig. 8** Retigabine enhances the increase in intracellular \([Ca^{2+}]_{in}\) in cultured keratinocytes produced by carvacrol. a Representative Ca\(^{2+}\) signals, measured as increases in the 350:380-nm fluorescence signal in Fura-2 preloaded cells and recorded from two keratinocytes following two applications to each of 1 mM carvacrol for 3 min. The top trace shows representative responses to two successive applications of carvacrol. In the lower trace, the second carvacrol application was preceded by the addition of retigabine (Ret: bath concentration 10 μM). b The mean ratio of the second-to-first responses (R2/R1) following addition of Krebs’ solution (Carv+Krebs) or retigabine (Carv+Ret) before the second response. Bars are SEM, number of cells in brackets. The asterisk (**) indicated that the R2/R1 ratio is significantly greater when R2 was preceded by retigabine than when it was not (P<0.01).
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