EXPRESSION OF TRANSIENT RECEPTOR POTENTIAL CHANNELS TRPC1 AND TRPV4 IN VENOATRIAL ENDOCARDIUM OF THE RAT HEART

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Abstract—The atrial volume receptor reflex arc serves to regulate plasma volume. Atrial volume receptors located in the endocardium of the atrial wall undergo mechanical deformation as blood is returned to the atria of the heart. The mechanosensitive channel(s) responsible for regulating plasma volume remain to be determined. Here we report that the TRP channel family members TRPC1 and TRPV4 were expressed in sensory nerve endings in the atrial endocardium. Furthermore, TRPC1 and TRPV4 were coincident with the nerve ending vesicle marker synaptophysin. Calcitonin gene-related peptide was exclusively confined to the myo-endocardium but appeared to be present in epithelial cells rather than sensory endings. In conclusion, we have provided the first evidence for TRPC1 and TRPV4 channels as potential contributors to mechanosensation in the atrial volume receptors. © 2014 The Authors. Published by Elsevier Ltd. on behalf of IBRO. This is an open access article under the CC BY license (http://creativecommons.org/licenses/by/3.0/).

Key words: TRPC1, TRPV4, synaptophysin, atrial volume receptor, mechanosensation, SK channels.

INTRODUCTION

Cardiovascular fluid homeostasis is essential for survival. The atrial volume reflex arc is an important contributor to the maintenance of bodily homeostasis, primarily responding to blood volume changes (Hainsworth, 1991). This reflex is initiated by cardiac mechanoreceptors known as atrial volume or low pressure receptors located at the venous atrial junction of the heart. The receptors are activated as blood is returned to the atria. Mechanical deformation of the atrial wall is transduced into a neural signal, which vagal afferent fibers (arising from the atrial volume receptors) conduct toward the nucleus tractus solitarii (Spyer, 1994). The information is integrated centrally, for example, in the paraventricular nucleus of the hypothalamus (Affleck et al., 2012) to bring about changes in the level of sympathetic activation that promotes water and electrolyte output from the kidney (Ledsome and Linden, 1964; Kappagoda et al., 1973). Paintal (1953) was probably the first to provide electrophysiological characteristics of the receptors. Information regarding their exact location and morphology is sparse and conflicting, although they are reported to be found mainly in the endocardial layer of the atrial wall (Woollard, 1926; Colendge et al., 1957; Holmes, 1957; Tranumjensen, 1975; Thoren et al., 1979; Hainsworth, 1991; Cheng et al., 1997). Details of the molecular machinery underpinning the transduction mechanism so far remain unknown.

Recent advances in understanding the molecular mechanisms of mammalian mechanotransduction systems have led to the identification of possible candidates for mechanosensitive channel proteins (Delmas et al., 2011). These include the epithelial Na channel/DEG/ASIC family in mechanotransduction in rat muscle spindles has recently been described (Simon et al., 2010). The TRP family proteins have been implicated in mechanosensation in the heart (Inoue et al., 2009). Baroreceptor terminals innervating the aortic arch and carotid sinuses in rats were found to express the γ subunit of ENaC (Drummond et al., 1998). In mice, ASIC1, 2 and 3 were found in aortic baroreceptor neurons in the nodose ganglia and their terminals in the aortic arch (Lu et al., 2009). Furthermore, ASIC2 null mice exhibit impaired baroreflex control and develop hypertension, thus providing evidence that compromised mechanosensing within the cardiovascular system could contribute to cardiovascular disease (Lu et al., 2009). The ASIC3 channel would appear to be important for blood volume control in mice (Lee et al., 2011) and be associated with calcitonin gene-related peptide (CGRP) in the nerve terminals in the venous atrial junction area.

Few studies have looked at TRP expression in mechanosensory organs and endings despite the fact they are considered to be strong candidates as mechanosensitive channels in mammals (Inoue et al., 2009).
A pressure-induced calcium influx that was gadolinium, but not lanthanide sensitive has been demonstrated in baroreceptor neurons from nodose ganglia of rats (Sullivan et al., 1997). These results suggest that stretch-activated channels, for which TRP channels are leading molecular candidates, could be the mechanotransducers in baroreceptors (Watanabe et al., 2008). TRPC1 and TRPC3–5 are present not only in the soma of nodose ganglion sensory neurons but also in the peripheral axons and mechanosensory endings that terminate as mechanosensitive receptors in the aortic arch of the rat (Glazebrook et al., 2005).

We used antibodies directed towards synaptophysin (SYN) or CGRP to identify sensory nerve terminals within the endocardium in the right atrial cardiac tissue and combined these with anti-channel antibodies, focusing on proteins of the ENaC/DEG/ASIC and TRP families as potential contributors to the mechanotransduction process. Afferent output from rat muscle spindle endings is subject to autogenic modulation by glutamate that is Ca2+ dependant (Bewick et al., 2005) and Ca2+-activated K+ channels have been shown to regulate the discharge frequency from these endings (Banks et al., 2009). Furthermore in rat atria the Ca2+-activated K+ channel isoform SK4 is involved in the release of atrial natriuretic peptide (ANP) in response to stretch (Ogawa et al., 2009). Therefore we also tested for the presence of Ca2+-activated K+ channels SK1–4 as potential modulators of atrial volume receptor excitability.

**EXPERIMENTAL PROCEDURES**

**Ethics statement**

Three male Hooded Lister rats were humanely killed in strict accordance with the Code of Practice for the Humane Killing of Animals under Schedule 1 of the Animals (Scientific Procedures) Act 1986. The protocol was approved by Durham University Life Sciences Ethical Review Panel.

**Tissue preparation for immunohistochemistry**

The atria, including entrances of the major vessels, were removed and fixed overnight in 4% formaldehyde at 4 °C. Following cryoprotection in sucrose/phosphate-buffered saline (PBS), venoatrial regions known to contain atrial volume receptors were dissected out, frozen and 16-μm cryosections collected onto poly-l-lysine coated slides. Sections were allowed to air dry for 1 h at room temperature, the slides were then stored briefly at −20 °C before use. On the day of the assay, slides were removed from the freezer and air dried again for one hour.

**Immunohistochemistry**

Sections were blocked for 45 min at room temperature in PBS pH 7.4 containing 0.4% Triton X-100, 4% fetal calf serum (FCS) and 1% bovine serum albumin (BSA). The antibody diluent for both the primary and secondary antibodies was PBS pH 7.4 containing 0.1% Triton X-100, 1% FCS and 1% BSA. After the blocking stage sections were incubated for 48 h at 4 °C with a pair of primary antibodies (see Sections ‘Sensory neurons’, ‘Mechanosensitive channels’, and ‘Small conductance Ca2+-activated K+ channels (SK1–4)’ for details). Following this, the sections were rinsed and washed in PBS (3 × 10 min), and then Alexa Fluor (AF) secondary antibodies applied for 1 h (either goat anti-rabbit-AF 594 and goat anti-mouse-AF 488, or donkey anti-goat-AF 594 and donkey anti-mouse-AF 488). See Table 1 for details and concentrations of primary and secondary antibodies. The slides were rinsed and washed in PBS as before and mounted in PBS/glycerol.

**Sensory neurons.** Rabbit anti–CGRP and mouse anti-SYN antibodies were used to identify sensory neurons. Secondary antibodies were goat anti-rabbit-AF 594 and goat anti-mouse-AF 488. All secondary antibodies were used as described above (Section 'Immunohistochemistry').

**Mechanosensitive channels.** Anti-channel antibodies (ASIC2, ASIC3, αENaC, βENaC, γENaC, TRPC1, TRPC4/5, TRPC6, TRPV4) were used together with mouse anti-SYN. These were all rabbit antibodies, with the exception antibodies to ASIC2, ASIC3 and TRPC6, which were raised in goat. Secondary antibodies were either goat anti-rabbit-AF 594 and goat anti-mouse-AF 488, or donkey anti-goat-AF 594 and donkey anti-mouse-AF 488; as appropriate. Rabbit anti-TRPV4 and mouse anti-CGRP (goat anti-rabbit-AF 594 and goat anti-mouse-AF 488 as secondaries) was also used.

**Small conductance Ca2+-activated K+ channels (SK1–4).** SK1 and SK3 antibodies were raised in goat, SK2 and SK4 in rabbit. They were used with mouse anti-SYN as the sensory neuron marker. Secondary antibodies were either donkey anti-goat-AF 594 and donkey anti-mouse-AF 488 or goat anti-rabbit-AF 488 and goat anti-mouse-AF 594, as appropriate.

**Antibody specificity.** These are all commercial antibodies subject to routine quality assurance (see Table 1 for details). Where positive results were obtained the pattern of reactivity was distinctive to that particular antibody with specific structures consistently labeled by that antibody on repeat assays. Furthermore anti-TRPV4 immunoreactivity (IR) was absent in tissue from TRPV4+/− mice using this same commercial antibody (Girard et al., 2013). Similarly anti-TRPC1-specific staining in C2C12 myoblasts was significantly reduced following TRPC1 silencing with TRPC1–siRNA (Meacci et al., 2010). We have used the antibodies against ENaC/DEG/ASIC proteins (with the exception of anti-ASIC3) in a previous study investigating mechanotransduction in rat muscle spindles (Simon et al., 2010). We have used the SK 1–3 antibodies to examine the expression of SK channels in lanceolate endings of hair follicles in the rat (Shenton et al., 2010). Specificity of the secondary antibodies was confirmed by incubating control sections in PBS with the omission of primary antibodies. With this regime no staining was observed.
| Antibody | Host | Poly/monoclonal Purification method | Immunogen | Manufacturer | Catalog No. | Concentration or dilution | Manufacturer’s quality control |
|----------|------|-----------------------------------|------------|--------------|-------------|--------------------------|--------------------------------|
| **Sensory neurons** |      |                                   |            |              |             |                          |                                |
| Anti-SYN | Mouse | Mono clone SY38 protein A purified | Vesicular fraction of bovine brain | Millipore | MAB5258 | 1 µg/ml | WB mouse brain lysates, single band 38 kDa |
| Anti-CGRP | Mouse | Mono clone 4901 protein G purified | Rat alpha-CGRP | Abcam | Ab81887 | 1/80 | IHC DRG, pancreas and gut |
| Anti-CGRP | Rabbit | Polyclonal whole antiserum | Full length rat CGRP conjugated to bovine THY | Abcam | Ab43873 | 1/300 | ICC rat amygdala and spinal cord |
| **Mechanosensitive channels** |      |                                   |            |              |             |                          |                                |
| ASIC2 (E-20) | Goat | Polyclonal affinity purified | Epitope mapping at the N-terminus of human ASIC2 | Santa Cruz | sc-22333 | 5 µg/ml | WB rat brain, band at 65 kDa |
| ASIC3 | Goat | Polyclonal whole antiserum | Synthetic peptide from the extracellular domain of rat ASIC3 | Abcam | Ab101595 | 1/280 | WB recombinant peptide |
| αENaC (H-95) | Rabbit | Polyclonal proprietary method | aa 131-225 near the N-terminus of human αENaC | Santa Cruz | sc-21012 | 5 µg/ml | WB human recombinant αENaC fusion protein |
| βENaC (H-190) | Rabbit | Polyclonal proprietary method | aa 271-460 within an internal region of human βENaC | Santa Cruz | sc-21013 | 5 µg/ml | WB human βENaC transfected 293T cell lysates |
| γENaC (H-110) | Rabbit | Polyclonal proprietary method | aa 411-520 near the C-terminus of human γENaC | Santa Cruz | sc-21014 | 5 µg/ml | WB cell lysates: A549, Caki-1 and COLO 320DM cells |
| TRPC1 (H-105) | Rabbit | Polyclonal proprietary method | aa 689-793 mapping at the C-terminus of human TRPC1 origin | Santa Cruz | sc-20110 | 5 µg/ml | WB mouse and rat testis, band 83 kDa |
| TRPC4/5 (H-80) | Rabbit | Polyclonal proprietary method | aa 1-80 within an N-terminal cytoplasmic domain of human TRPC5 | Santa Cruz | sc-28760 | 5 µg/ml | IF mouse heart |
| TRPC6 (C-13) | Goat | Polyclonal affinity purified | Peptide mapping at the C-terminus of human TRPC6 | Santa Cruz | sc-19197 | 5 µg/ml | IHC human rectum tissue |
| TRPV4 | Rabbit | Polyclonal affinity purified | Synthetic peptide KLH conjugated, derived from within residues 850 to the C-terminus of mouse TRPV4 | Abcam | Ab39260 | 5 µg/ml | WB vs mouse brain lysates |
| **Small conductance Ca²⁺-activated K⁺ channels** |      |                                   |            |              |             |                          |                                |
| SK1 (A-13) | Goat | Polyclonal proprietary method | Epitope mapping near the C-terminus of human SK1 | Santa Cruz | sc-17991 | 5 µg/ml | WB rat and mouse brain |
| SK2 | Rabbit | Polyclonal affinity purified | aa 542-559 mapping within the intracellular C-terminus of rat SK2 | Alomone | APC-028 | 2 µg/ml | WB rat brain membranes |
| SK3 (H-17) | Goat | Polyclonal proprietary method | Epitope mapping within an internal region of human SK3 | Santa Cruz | sc-16027 | 5 µg/ml | WB PC-12 cell lysate |
| SK4 | Rabbit | Polyclonal affinity purified | aa 350-363 mapping within the intracellular C-terminus of rat SK4 | Alomone | APC-064 | 2 µg/ml | WB in rat and human tissue |
Image acquisition, visualization and analysis

Sections were examined using a Zeiss Axioskop 2 under epifluorescence. Digital images were captured with a Hamamatsu Orca 285 CCD camera controlled by Improvision Velocity (Acquisition, Restoration and Visualisation) software (v. 6.2.1). Images were obtained and processed in a similar manner to that described previously (Affleck et al., 2012). For selected images a series of z-stacks were acquired. The 3D volume was deconvolved using iterative restoration and the appropriate Point Spread Function which models the spread of light in the optical path that causes blur. This process reassigned out-of-focus haze without subtracting it from the data to improve resolution in the X, Y and Z planes. This produced a confocal-like quality image. The most appropriate single Z plane image was then converted to a 3-D Opacity image, which is a high-resolution render with options to display data in different views. The isosurface view is a form of indirect rendering, which identifies a surface around objects where all voxel intensity values are the same. Isosurface rendering generates a 3D non-transparent solid of the surface elements. The max intensity view applies direct maximum intensity projection rendering. The brightest intensity in the view path to the screen will form the image. These image manipulations can improve clarity and thereby help in the visualization of sites where differentially labeled proteins may coincide and interact.

The final images were imported into Adobe Photoshop (CS4 extended v. 11.02), which was used to adjust brightness and contrast. Processed images were grouped into plates and labeled in Adobe Photoshop.

RESULTS

Sensory neurons

**SYN labeling.** SYN-IR was widely expressed in all three layers of the atrial wall. In the endocardium, where atrial volume receptors are expected to be predominantly located SYN-IR was dense and complex in places (Fig. 1A) with an appearance characteristic of similar mechanosensory endings (Drummond et al., 1998; Maeda et al., 1999). In the myocardium SYN-IR was also present in the vicinity of structures with the appearance of small blood vessels (Fig. 1A). In the epicardium SYN-IR was found in nerves and around putative ganglion cells and blood vessels (not shown).

**CGRP labeling.** By contrast CGRP-IR was rarely seen in the endocardial layer (Fig. 1B). It was occasionally present in the myocardium, mostly in association with blood vessel-like structures (Fig. 1B) where there was overlap with SYN labeling (Fig. 1C); but also on some nerve endings within this layer (Fig. 1E). Immunoreactivity for CGRP was intense in the epicardium around putative ganglion cells and blood vessels, and within nerve fibers (not shown).

Mechanosensitive ion channels

**TRPC1 labeling.** Immunoreactivity for mechano-sensitive ion channels revealed the presence of TRPC1 (Fig. 2). Immunoreactivity for this channel was coincident with SYN-IR in the endocardium (Fig. 2C, C') and myocardium (not shown), although SYN-IR was always more prevalent. The isosurface view (Fig. 2C') suggests that while the TRPC1 and SYN labeling are closely associated, they may be localized to different compartments within the sensory nerve terminals. Immunoreactivity for TRPC1 was not found around ganglion cells in the epicardium (not shown), even where SYN labeling was evident.

**TRPV4 labeling.** TRPV4 labeling was particularly widespread in both the endocardium (Fig. 3B, E) and myocardium (Fig. 3E). In these layers TRPV4-IR and SYN-IR always coincided (Fig. 3C, C'). TRPV4-IR was far more abundant than CGRP-IR in both endocardium and myocardium, especially in endocardium where CGRP was rarely expressed (Fig. 3D–F'). Although far less prevalent, CGRP labeling was present on some TRPV4-positive endings (Fig. 3F, F'). The isosurface

![Fig. 1. Sensory neurons: synaptophysin and calcitonin gene-related peptide labeling. Double labeling with mouse anti-synaptophysin (green SYN A, C) and rabbit anti-calcitonin gene-related peptide (red CGRP B, C) antibodies. SYN-immunoreactivity (SYN-IR) (short arrows A, C) was present in all three layers of the cardiac wall. In the endocardium (ENDO) SYN-IR was dense and complex, compared to the myocardium (MYO) where it was sparse. SYN-IR was also evident in the location of blood vessel-like structures (BV). CGRP-IR was occasionally expressed in the myocardium, especially around blood vessels (long arrow B) where there was some overlap with SYN-IR (arrowhead C); but CGRP-IR did not appear to extend into endocardium. (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)

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views were again suggestive of compartmentalization of TRPV4 labeling compared with either SYN (Fig. 3C) or CGRP (Fig. 3F) expression on the same ending. In contrast to TRPC1, TRPV4-IR was also evident around ganglion cells in the epicardium where it coincided with SYN-IR labeling (not shown).

There was a complete absence of immunoreactivity for ENaC/DEG/ASIC family members: αENaC, βENaC, γENaC, ASIC2 or ASIC3. Also for the TRP family members, TRPV1 and TRPC4–6 again no immunoreactivity was evident.

Small conductance Ca\(^{2+}\)-activated K\(^{+}\) channels

**SK2 labeling.** For small conductance Ca\(^{2+}\)-activated K\(^{+}\) channel SK2 immunoreactivity (SK2-IR) was evident in nerve bundles within the epicardium, although it did not coincide with SYN-IR in this location (not shown). However, SK2-IR was only very occasionally present within the myocardium possibly extending into the endocardium (Fig. 4B) and here there was SK2-SYN co-labeling (Fig. 4C–C''), although the reactivity did not precisely overlap, suggestive of compartmentalization (Fig. 4C', C'').

**SK4 labeling.** Small conductance Ca\(^{2+}\)-activated K\(^{+}\) channel SK4 immunoreactivity (SK4-IR) was found in the endocardium giving small patches of diffuse labeling that appeared to be over epithelial cells (Fig. 4E). SK4-IR was clearly distinct from SYN-IR in this layer (Fig. 4F). There was an absence of SK1 or SK3 immunoreactivity throughout the heart tissue.

DISCUSSION

This study has provided the first evidence of the conductance channels present in putative atrial volume receptors within the endocardium of the right atria of the rat. The TRP family channels TRPC1 and TRPV4 were expressed, however there was no evidence to indicate the presence of ENaC/DEG/ASIC proteins. These findings suggest mechanotransduction in atrial volume receptors may rely on the TRP family of channels, which contrasts with the baroreceptor and muscle spindle where ENaC/DEG/ASIC channel families have been reported (Drummond et al., 1998; Simon et al., 2010) and are therefore candidates as the mechanotransducers in these endings.

Calcium-activated K\(^{+}\) channel, SK2 and SK4 were occasionally present in both the endo- or myocardial layers. Based on the appearance of the putative AVR’s identified by the sensory nerve terminal marker SYN (see Section ‘Synaptophysin labeling’), we suggest SK2 and SK4 may not form part of the molecular components of an AVR.

**SYN and CGRP as markers of sensory nerve terminals: putative atrial volume receptors**

The vesicle marker SYN was abundantly expressed in all three layers of the cardiac wall. SYN is used to identify the synaptic vesicles in presynaptic endings. Furthermore synaptic-like vesicles have commonly been described in mechanosensory endings of vertebrate and invertebrate animals (Katz, 1966). The pattern of SYN-IR observed in the endocardium was characteristic of sensory endings (Drummond et al., 1998; Maeda et al., 1999) and therefore likely to be labeling atrial volume receptors. Previous histological studies describe atrial
Fig. 3. Mechanosensitive ion channels: Transient Receptor Potential Vanilloid 4 (TRPV4) labeling in endocardium and myocardium. Double labeling with rabbit anti-TRPV4 (red TRPV4 B, C, E, F, F') and either mouse anti-SYN (green SYN A, C, C') or mouse anti-CGRP (green CGRP D, F, F') antibodies. TRPV4 immunoreactivity (TRPV4-IR) (long arrows B, E) was widespread in the endocardium (ENDO) and also extended into the myocardium (MYO). Nerve endings identified by SYN-IR (short arrows A) were colabeled with anti-TRPV4 antibodies (arrowheads C). Panel C' is the isosurface presentation of a Volocity 3D slice to illustrate the close relationship between TRPV4 and SYN labeling (black arrows C', indicate concurrent TRPV4–SYN labeling). CGRP-IR was only rarely found in either endocardium or myocardium (short arrows D). However, on the occasions when it was present the endings were also TRPV4 positive (arrowhead F). Panel F' is the isosurface presentation of a Volocity 3D slice to illustrate the presence of CGRP labeling on TRPV4-positive endings (black arrows F', indicate dual labeling). The isosurface views (C', F') are again indicative of anti-channel and sensory nerve labeling occurring in distinct compartments within the same ending. (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)
volume receptors as predominantly within the endocardial layer ideally placed to sense blood volume changes (Coleridge et al., 1957; Holmes, 1957; Tranumjensen, 1975; Thoren et al., 1979; Cheng et al., 1997). Based on the detailed description of vagal afferents carried out by Cheng et al. (1997), the SYN-IR we observed around ganglion cells in the epicardium was probably efferent rather than afferent innervation. SYN-IR within the myocardium was slightly less dense than in the other two layers. Presumptive atrial volume receptors have also been shown to extend into myocardium (Tranumjensen, 1975) and it has been suggested that atrial volume receptors in these various locations could account for the different types based on electrophysiological classification (Kappagoda et al., 1976).

In our hands CGRP-IR was not as widespread as SYN-IR and it seldom extended into the endocardium. CGRP–IR was strongest in the epicardium surrounding ganglion cells, blood vessels and within nerve fibers. These findings are in agreement with a study in which only 9% of vagal afferent neurons, identified by injecting the retrograde tracer cholera toxin B-subunit into the pericardium, were found to be CGRP positive (Corbett et al., 2005). There was seldom co-localization of CGRP and SYN-IR in any of the tissue layers and this, together with the distribution pattern, suggests that CGRP is not a marker for mechanosensory afferents here. CGRP– labeling of volume receptor nerve terminals in the venoatrial junction has been found co-localized with ASIC3-IR (Lee et al., 2011). However, it is not certain that the endings described here were in the endocardium.

CGRP- has been associated with the regulation of sympathetic activity (Oh-hashi et al., 2001). However, our observations suggest that in respect to blood volume at least, this is more likely to be mediated via CGRP-positive neurons located in the venoatrial junction and has been found co-localized with ASIC3-IR (Lee et al., 2011). However, it is not certain that the endings described here were in the endocardium.

Mechanosensitive ion channels

Immunoreactivity for TRPC1 and TRPV4 was found within the atrial tissue and in particular the endocardial layer. Where TRPV4-IR was observed around ganglion cells in the epicardium, the innervation is most probably efferent (Cheng et al., 1997). Around ganglion cells in the epicardium TRPV4 reactivity was observed, but not
TRPC1-IR. Again this innervation around ganglion cells is most probably efferent (Cheng et al., 1997). The TRPC1 and TRPV4-IR always coincided with SYN-IR, suggestive of labeling of putative atrial volume receptors where it was present within the endocardium. Since our CGRP-labeling experiments suggested this peptide was not expressed in atrial volume receptor afferents (see Section ‘Discussion’), only limited assays were undertaken using anti-mechanosensitive ion channel antibodies in combination with anti-CGRP antibodies.

Surprisingly, no immunoreactivity for ENaC (αENaC, βENaC, γENaC) or ASIC (ASIC2, ASIC3) was evident in any area of the atrial tissue examined. Amiloride sensitive channels play a role in mechanotransduction in rat muscle spindles (Simon et al., 2010) and these are likely to be ENaC/DEG/ASIC-related proteins. The antibodies used in this present study were the same as those used by ourselves in rat muscle spindles where we detected immunoreactivity to ENaC (αENaC, βENaC, γENaC), and ASIC2 in primary mechanosensory afferents (Simon et al., 2010). Therefore, our evidence would suggest that mechanotransduction in the atrial volume receptors is dependent upon a different set of channel proteins.

In muscle spindle terminals primarily a Na⁺ current underpins mechanotransduction (Hunt et al., 1978). TRP channels are non-selective Ca²⁺ channels (Ramsay et al., 2006), therefore the ionic basis for the current in atrial volume receptors is likely to be different from that observed in spindle terminals. Interestingly, gadolinium, a non-selective blocker of stretch-activated ion channels, reduced blood volume expansion effects on urine flow, neural activation, and ANP release both in ASIC3−/− and ASIC3+/+ control mice (Lee et al., 2011). Furthermore, gadolinium also blocks TRP channels including TRPV4 (Jung et al., 2003; Becker et al., 2005): suggesting that other molecular components, for example TRP channels, in addition to ASIC3 can contribute to blood volume control.

The ASIC2 channel located in the nodose ganglia and aortic arch has been shown to be an important component of the arterial baroreceptor reflex in mice (Lu et al., 2009). A pressure-induced calcium influx has been demonstrated in baroreceptor neurons from the nodose ganglia of rats with characteristics compatible with TRP channels (Sullivan et al., 1997). The TRPC1 channel appears to be important for cardiovascular regulation, affecting vascular tone and smooth muscle depolarization in response to pressure load (Inoue et al., 2009). Our finding of TRPC1 immunoreactivity in putative mechanosensory endings in the right atrium, suggests that TRPC1 activation in this location could in addition initiate or regulate the atrial volume receptor reflex arc.

It is probable that TRPV channels function as part of multiprotein complexes (Suzuki et al., 2005; Fu et al., 2006; Huai et al., 2012), where different components of the complex may regulate TRPV expression and underlie tissue-specific sensing modalities (D’Hoedt et al., 2008). For example, in mice, within the parvocellular neurons of the paraventricular nucleus, TRPV4 and SK channels are involved in osmosensing (Feetham and Barrett-Jolley, 2012). Interestingly, it has recently been demonstrated that TRPV4 forms heteromeric channels with TRPC1 in vascular endothelial cells (Ma et al., 2010) and this TRPV4-TRPC1 complex mediates flow-induced endothelial Ca²⁺ influx. Functional evidence also supports a role for TRPV4 in blood pressure regulation in rats. Administration of the TRPV4 selective activator 4α-phorbol 12,13-didecanoate (4α-PDD) leads to dose-dependent decreases in blood pressure (Gao et al., 2009). Here application of specific blockers, demonstrates the hypotensive effect of TRPV4 is mediated, at least in part, by activation of Ca²⁺-activated K⁺ channels (blocked by apamin and charybdotoxin) and CGRP receptors (blocked with CGRP8-37) responding to CGRP release from sensory nerves (Gao and Wang, 2010). We found scant evidence for CGRP-IR in the endocardial and myocardial layers of the atrial wall. Where CGRP-IR was occasionally evident on nerve endings in these layers there was coincident TRPV4 labeling and these endings could contribute to volume sensing. Activation of TRPV4 in this location could potentially result in an inhibition in sympathetic nerve activity and a reduction in blood pressure via the well-characterized CNS circuits known to regulate the volume reflex arc (Ledsome and Linden, 1964; Yang and Coote, 2003).

Ca²⁺-activated K⁺ channels

For SK2 most expression was confined within nerves in the epicardium; while wispy SK4-IR was present over limited regions of endocardium, there was no relationship between SK4 and SYN labeling. Immunoreactivity for SK2 was only occasionally found on nerve endings within the myocardium possibly continuing into the endocardium, here it was associated SYN reactivity but the labeling did not strictly coincide. The absence of expression within the putative atrial volume receptor afferents, suggests these channels are unlikely to directly influence afferent sensitivity in this instance. However, SK channels are known to regulate neuronal excitability and levels of intracellular Ca²⁺ (Stocker, 2004). Murine atrial myocytes have been reported to contain SK2 channels where they contribute to cardiac depolarization (Li et al., 2009). By contrast we did not find SK2-IR in the myocardium localized to nerves or myocytes, but there was strong SK2-IR in nerves of the epicardium. We have found SK2-IR both in the terminals of muscle spindle primary endings and in lanceolate endings (F.C. Shenton, unpublished observations) where the SK2 reactivity co-localized with SYN-IR. The SK2 channel was also present in the satellite glial cells with which sensory terminals of the lanceolate endings are very closely associated. However in this present investigation there was no evidence of SK2 channels expressed in putative atrial volume receptors.

A specific SK4 inhibitor has been shown to significantly decrease both basal and stretch-stimulated
ANP secretion from an isolated rat atrial preparation (Ogawa et al., 2009). We have demonstrated the presence of SK4-IR in endocardium of the venoatrial junction, where it appeared to be expressed in endothelial cells; thus we postulate this channel could influence the release of ANP from atrial myocytes. We failed to detect immunoreactivity with either anti-SK1 or anti-SK3 antibodies. The SK1 channel is reported to be mostly restricted to the brain (Stockert, 2004) although Tutje et al. (2005) have also found it expressed in isolated rat cardiomyocytes. In our hands the anti-SK3 antibody stains lanceolate endings surrounding hair follicles (Shenton et al., 2010); however this immunoreactivity appeared to be predominantly confined to satellite glial cells surrounding the sensory endings and close to the interface between the endings and the glial cell processes, rather than in the terminals themselves.

Compartmentalization

The image acquisition and visualization process described in Section 'Image acquisition, visualization and analysis' produced images of a confocal-like quality. The isosurface rendering in particular aids in illustrating the correlation between differently labeled proteins. These views revealed that anti-channel labeling did not precisely coincide with SYN-IR on sensory neurons, suggesting compartmentalization of TRP and SYN proteins within the ending. This was also evident where SK2-SYN co-labeling was seen. SYN is expressed in synaptic-like vesicles in sensory endings (de Camilli et al., 1988; Bewick et al., 2005), while one might expect channel proteins to be preferentially concentrated in the terminal membranes.

CONCLUSION

This study provides the first evidence as to the possible identity of the mechanosensitive channels that detect blood volume changes and transduce that stimulus into a nervous impulse to initiate the atrial volume reflex arc. Therefore, TRPC1 and TRPV4 proteins should be a focus of future functional investigations to confirm their role in mechanotransduction in atrial volume receptors. Elucidation of the process whereby changes in returning blood volume are detected and signaled to the CNS is necessary to further our understanding of how normal cardiovascular homeostasis malfunctions in heart failure and hypertension. This will require a comprehensive knowledge of the channels and proteins involved. Our study has identified important channel differences between those sensory endings we putatively identify as atrial volume receptors, and the baroreceptor and muscle spindle. These differences may reflect the nature of the stimulus being detected and requires further investigation.

Mechanotransduction mechanisms in vertebrates are certain to be diverse and complex, reflecting the broad range of functions they underpin. Other channels and regulatory proteins, in addition to the primary mechanosensitive channels, contribute to the process (Delmas et al., 2011); nevertheless they may broadly function in a similar manner, while differing in the detail. Bewick et al. (2005) demonstrated autogenic modulation of mechanoreceptor excitability by glutamate release from SYN positive, synaptic-like vesicles in rat muscle spindle primary sensory endings, providing an elegant regulatory model that may pertain to similar mechanosensitive endings. Our observation of TRPC1 and TRPV4 coincident with abundant SYN-IR in the venous atrial junction would be compatible with such a model.

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