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Benzamil sensitive ion channels contribute to volume regulation in canine chondrocytes

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BACKGROUND AND PURPOSE
Chondrocytes exist within cartilage and serve to maintain the extracellular matrix. It has been postulated that osteoarthritic (OA) chondrocytes lose the ability to regulate their volume, affecting extracellular matrix production. In previous studies, we identified expression of epithelial sodium channels (ENaC) in human chondrocytes, but their function remained unknown. Although ENaC typically has Na⁺ transport roles, it is also involved in the cell volume regulation of rat hepatocytes. ENaC is a member of the degenerin (Deg) family, and ENaC/Deg-like channels have a low conductance and high sensitivity to benzamil. In this study, we investigated whether canine chondrocytes express functional ENaC/Deg-like ion channels and, if so, what their function may be.

EXPERIMENTAL APPROACH
Canine chondrocytes were harvested from dogs killed for unassociated welfare reasons. We used immunohistochemistry and patch-clamp electrophysiology to investigate ENaC expression and video microscopy to analyse the effects of pharmacological inhibition of ENaC/Deg on cell volume regulation.

KEY RESULTS
Immunofluorescence showed that canine chondrocytes expressed ENaC protein. Single-channel recordings demonstrated expression of a benzamil-sensitive Na⁺ conductance (9 pS), and whole-cell experiments show this to be approximately 1.5 nS per cell with high selectivity for Na⁺. Benzamil hyperpolarized chondrocytes by approximately 8 mV with a pD₂ 8.4. Chondrocyte regulatory volume decrease (RVI) was inhibited by benzamil (pD₂ 7.5) but persisted when extracellular Na⁺ ions were replaced by Li⁺.

CONCLUSION AND IMPLICATIONS
Our data suggest that benzamil inhibits RVI by reducing the influx of Na⁺ ions through ENaC/Deg-like ion channels and present ENaC/Deg as a possible target for pharmacological modulation of chondrocyte volume.

Abbreviations
AQP, aquaporin; Deg, degenerin; ENaC, epithelial sodium channel; OA, osteoarthritis; RMP, resting membrane potential; RVD, regulatory volume decrease; RVI, regulatory volume increase

Introduction
Chondrocytes are the cells of cartilage that synthesize and maintain the substance of cartilage, the extracellular matrix (Stockwell, 1975). They exist in a unique high-pressure, avascular environment, but chondrocytes can be enzymatically isolated from cartilage and largely retain phenotype for the first few passages in culture (Benya and Shaffer, 1982). Recent
studies have shown that chondrocytes, like other cells, express a wide variety of ion channels (Barrett-Jolley et al., 2010) including several species of potassium channel (Wilson et al., 2004; Mobasheri et al., 2005; 2007; 2010; Clark et al., 2010b; 2011) and at least two species of chloride channels (Sugimoto et al., 1996; Tsuga et al., 2002; Funabashi et al., 2010). Our own studies have also shown immunohistochemically that human chondrocytes express the epithelial sodium channel (ENaC) (Trujillo et al., 1999). We previously made a preliminary report of their functional expression in canine chondrocytes (Lewis et al., 2008), but their function has been, to date, entirely unknown. ENaC is a low conductance sodium channel consisting of various combinations of three subunits: α, β and γ (Canessa et al., 1994); and our previous immunohistochemical experiments identified each of these in human cartilage (Trujillo et al., 1999). ENaC has the key pharmacological property of sensitivity to low concentrations of benzamil and amiloride and is the prototypic member of the ENaC-degenerin (ENaC/Deg) protein family (Alvarez de la Rosa et al., 2000; Kellenberger and Schild, 2002). The most well characterized function of ENaC is transport of Na⁺ across epithelial barriers (Alvarez de la Rosa et al., 2000; Kellenberger and Schild, 2002). As such, ENaC in particular, is a regulator of sodium homeostasis, modulating sodium re-absorption in the kidney (Hager et al., 2001) and sodium absorption from the colon (Kunzelmann and Mall, 2002). However, recent evidence suggests that ENaC has more diverse functions than this, for example, sodium sensing on gustatory neurones of the anterior tongue (Staehler et al., 2008) and neurones in the paraventricular nucleus (Teruyama et al., 2011). ENaC has also been implicated in mechanotransduction in some organisms (discussed by Martinac, 2004; Chalfie, 2009). Specifically, ENaC has been shown to be ‘shrink-activated’ in certain cell types (Bohner et al., 2000; Bondarava et al., 2009). Of particular interest to us was the discovery that ENaC has an important (Wehner et al., 2000) and yet complex (Wehner et al., 2006) role in the cell volume regulation of rat hepatocytes. Active volume regulation is a well known physiological response of chondrocytes (reviewed by Lewis et al., 2011b). Under appropriate conditions, they exhibit both regulatory volume decrease (RVD) (Bush and Hall, 2001a,b) and regulatory volume increase (RVI) (Hall et al., 1996; Kerrigan et al., 2006). Typically, in eukaryotic cells, RVI occurs following exposure to hypertonic solutions (Hoffmann et al., 2009). Cells first lose water by osmosis and then shrink; RVI is the process whereby cells recover their volume by accumulating additional solutes and then osmotically imbibering water (Hoffmann et al., 2009). In rat hepatocytes, accumulation of solutes includes influx of sodium ions through ENaC (Wehner et al., 2000; 2006). Furthermore, since ENaC is a relatively sodium selective ion channel (Rossier et al., 1994), one would expect that their activation may lead to membrane depolarization, which could itself have indirect effects on cell function including volume regulation (Lewis et al., 2011a).

In the present study, we sought to characterize both protein and functional expression of ENaC/Deg channels and investigate whether they contribute to the RVI of canine chondrocytes. We show that ENaC is expressed in our canine chondrocyte model, that benzamil sensitive ion channel activity was increased by hypertonic challenge and that these ENaC/Deg-like channels contribute to the RVI response of canine chondrocytes.

Methods

Canine cartilage was obtained from stifle and elbow condyles of skeletally mature, bull terrier-type dogs killed for unrelated clinical reasons. No animals were harmed for this study. Chondrocytes were isolated as described previously (Mobasheri et al., 2010) with type II collagenase. To ensure preservation of the chondrocyte phenotype, we used freshly dissociated and up to third passage cells only.

RVI bioassay

Cells were placed in a ‘physiological saline’ (Table 1) solution including 145 mM Na⁺ and then moved to an identical solution except for the addition of 180 mM sucrose. Cells, at first, shrink as water leaves the cell due to osmosis (Figure 5A phase ‘I’) and then begin to swell as RVI takes place (Figure 5A phase ‘II’). Not all cells showed RVI, and so we used a two ‘run’ protocol. In the first ‘run’, we determined if RVI occurred by exposure to hypertonic solution. If this cell did exhibit RVI, we then continued with a second ‘run’ under either control, or treatment conditions. Live cell imaging was achieved with a Nikon Diaphot microscope equipped with a Sony ICX098QB high-sensitivity CCD, or NIKON Eclipse Ti confocal microscope (Nikon Kingston-Upon-Thames, Surrey, UK). Images were analysed offline with ImageJ (Abramoff et al., 2004). Volume was calculated from the 2D surface area (A) of the cell disc by assuming the cell is approximately spherical as described previously (Walsh and Zhang, 2005; Takeuchi et al., 2006; Lewis et al., 2011a), using the following equation:

\[
Vol = \frac{4}{3} \pi \left( \frac{A}{\pi} \right)^3
\]

Except where stated, data are presented normalized for starting volume (V₀) as V/V₀, where V is the volume at time t. Figure 5A shows confocal images verifying that the chondrocytes were approximately spherical during volume experiments. (Note that the cells only round up when removed from the culture vessels for the duration of the functional experiments, in the immunohistochemistry where they remain on coverslips, they have a flattened appearance). Cells were incubated with FM1-43 dye (Invitrogen, Paisley, UK) before being subjected to the same ‘run’ treatment as described above.

Electrophysiology

Membrane potentials and whole-cell currents were measured with the whole-cell patch clamp technique using an Axopatch 200A (Molecular Devices, Inc., Sunnyvale, CA), inside-out single channel recordings were made with an Axopatch 200A (Molecular Devices), but cell-attached patch recordings were made with an Axopatch 200B (Molecular Devices, Inc.). Except where stated, data were filtered (1 kHz) using the
inbuilt amplifier filters and sampled (5 kHz) using DigiData 1200 interface cards (Molecular Devices, Inc.). Voltage protocols were run by the WinEDR and WinWCP programs (John Dempster, University of Strathclyde). Osmolality changes were affected by addition or removal of 180 mM sucrose.

Whole-cell experiments. Patch-pipettes were fabricated from thick-walled borosilicate glass (outer diameter 1.5 mm, inner diameter 0.85 mm) with resistance approximately 5 MΩ when filled. For resting membrane potential (RMP) measurements, we used the standard physiological saline (Table 1), matched with a standard high K+ physiological intracellular saline solution (Table 2). To measure the sodium permeability of the benzamil-sensitive conductance, a voltage-ramp protocol was used. This consisted of a 4.5 s linear ramp from –80 to +80 mV, repeated at 50 s intervals. Difference currents were obtained by subtraction of a ramp in the presence of benzamil from that run in vehicle control. Intracellular [Na+] was varied (see Table 2 for the solutions used), and the reversal potential (Vrev) of the difference current at each of the [Na+] was calculated to permeability by fit of a standard variation of the Goldman Hodgkin Katz equation (Equation 2) and minimizing χ2:

\[
V_{rev} = \frac{RT}{F} \ln \left( \frac{[Na]_o + p[K]_o}{[Na]_i + p[K]_i} \right)
\]

where [Na]o is 145 mM, and [K]o is 5 mM (‘ENaC Permeability’ solution, Table 1). R, T and F have the standard definitions. p is the permeability ratio Na+/K+. [Na]i and [K]i varied between 22 and 150 mM according to the intracellular solution used (Table 2). Whole-cell currents were also recorded in voltage clamp mode using physiological saline solutions (Tables 1 and 2).

Single channel experiments. Patch pipettes were again fabricated from thick-walled borosilicate 1.5 mm (outer diameter), but resistances were approximately 10 MΩ when filled. For inside-out patch experiments, Na+ was high in both the patch pipettes (extracellular: ‘inside-out patch’, Table 1) and the bath (intracellular: ‘inside-out patch’, Table 2). For single channel kinetics, we used cell-attached patch single channel recording. Sodium ions were replaced by lithium (Li+) in the patch pipette (extracellular: ‘Lithium solution’ Table 1) and to nullify the membrane potential, the bath solution (extracellular: ‘Cell-attached patch bath solution’ Table 1) contained elevated K+ ions. Since ENaC/Deg channel conductance is so low, it is normal to filter traces heavily before performing kinetic analysis, typical cut-off frequencies from previous studies vary from 50 to 500 Hz (Eyre and Canessa, 1998; Ishikawa et al., 1998). In our study, we digitally re-filtered the data to 0.5 kHz. Events were then idealized using segmental K means (SKM) methods (Qin, 2004) using QuB software (Dr Feng Qin, Dr Lorin Milescu, Fu Qiong, Chris Nicolai and John Bannen, SUNY, Buffalo, NY). This method alternates between the Viberti method and parameter re-estimation; however, we constrained unitary amplitudes and amplitude distributions to clear periods of openings and closures. Models were fit to the idealized record by optimization of maximum interval likelihood (Qin et al., 1996; 1997). We chose our final model on the basis of significant increases of log-likelihood as described by Lema and Auerbach (2006). The probability density functions (PDFs) drawn in event frequency histogram graphs, and the respective time constants quoted are therefore those calculated from the model’s fitted rate constants. Where open probability (Po) was measured from patches with more than one channel, it was calculated as the average of those channels from Po = I/(NI) where N is number of channels in the patch (estimated as the maximum seen open at one time), and I is the mean unitary amplitude. Conductance was calculated as the slope of the fit of the IV curves. Ensemble averages were computed by simply averaging together several raw single channel traces.

The dose–response curve was fit in Sigmaplot (Systat Software, Hounslow, UK) using the following equation, modified from (Black and Shankley, 1985) and (Barrett-Jolley et al., 1999). The SEM values quoted are those calculated by the fitting algorithm.

\[
R = m \left( \frac{1 - [c]^h}{[c]^h + 10^{pD_2}} \right)
\]

R is the response (defined as appropriate in the figure legend), [c] is the concentration of drug, m the maximum value, h is the Hill slope and pD2 the negative log of the midpoint parameter. Statistical tests were performed using SPSS (SPSS Inc., Chicago, IL), or Minitab (Minitab Ltd, Coventry, UK). Values in text are quoted as mean ± SEM (n), where n is the sample size.

Immunohistochemistry

Primary canine articular chondrocytes were grown in DMEM on autoclaved coverslips at a density of 3 x 10⁴ cells·mL⁻¹. They were then fixed in ice-cold methanol for 10 min, washed and permeabilized with PBS-T and blocked with 10% BSA in PBS. The cells were then incubated with the primary antibodies to the α, β and γ subunits of ENaC (kind gift from Dr Cecilia M Canessa, Yale University, New Haven, CT). These have been used extensively elsewhere (Trujillo et al., 1999; Rubera et al., 2003; Shakibaei and Mobasheri, 2003; Zhang et al., 2005). After three washes with PBS-T, the cells were incubated for 2 h with a goat polyclonal secondary antibody to rabbit IgG (Fc-specific, affinity-purified, pre-adsorbed) conjugated to DyLight® 488 (Abcam, Cambridge, MA, USA; ab98462) diluted according to the manufacturer’s recommendations (typically diluted 1:200). After extensive washes in PBS-T, the nuclei were counterstained with propidium iodide (red fluorescence). The chondrocyte surface antigen, CD44, and SOX-9 as an indicator of differentiation, were used as additional controls for chondrocyte phenotype (data not shown). The negative controls were exposed to non-immune rabbit IgG and shows red fluorescent nuclear staining only. The cells were visualized, and digital images were captured using a Leica DM 5000B epifluorescence imaging system.

Software

Software used for each analysis technique is described above, but further details are available at http://pcwww.liv.ac.uk/~rbj/RBJ/software.htm.
Solutions

Table 1
Extracellular solutions used during electrophysiology experiments

<table>
<thead>
<tr>
<th>Extracellular (mM)</th>
<th>Na</th>
<th>K</th>
<th>Ca</th>
<th>Cl</th>
<th>Mg</th>
<th>Cs</th>
<th>SO₄</th>
<th>MeSO₄</th>
<th>Li</th>
<th>Glucose</th>
<th>HEPES</th>
<th>pH</th>
<th>Vj (mV)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. Inside-out patch</td>
<td>155</td>
<td>0</td>
<td>4</td>
<td>158</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>10</td>
<td>NaOH</td>
<td>5.9</td>
</tr>
<tr>
<td>2. Lithium solution</td>
<td>0</td>
<td>9</td>
<td>2</td>
<td>151</td>
<td>1</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>140</td>
<td>0</td>
<td>10</td>
<td>KOH</td>
<td>6</td>
</tr>
<tr>
<td>3. Cell-attached patch bath solution</td>
<td>0</td>
<td>115</td>
<td>2</td>
<td>122.2</td>
<td>1.6</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>10</td>
<td>10</td>
<td>KOH</td>
<td>14.4</td>
</tr>
<tr>
<td>4. Physiological saline</td>
<td>145</td>
<td>5</td>
<td>2</td>
<td>151</td>
<td>1</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>10</td>
<td>NaOH</td>
<td>0.5</td>
<td></td>
</tr>
<tr>
<td>5. NaMeSO₄</td>
<td>155</td>
<td>0</td>
<td>4</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>150</td>
<td>0</td>
<td>0</td>
<td>10</td>
<td>NaOH</td>
<td>5</td>
<td></td>
</tr>
<tr>
<td>6. ENaC permeability</td>
<td>158</td>
<td>5</td>
<td>2</td>
<td>159</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>10</td>
<td>NaOH</td>
<td></td>
</tr>
</tbody>
</table>

Vj values quoted are when matched with the appropriate intracellular solutions (Table 2). Calculations were performed with JpCalc (Barry and Lynch, 1991). Osmolarity was adjusted in all cases by addition of sucrose as appropriate. Sucrose was chosen to adjust osmolarity so that the ion concentrations remained unchanged during any given experiment. Vj varied between 0.1 and 4 mV when paired with intracellular solutions with different sodium concentrations. NB All ion concentrations are the total number of ions in the solution after adjustment of pH with NaOH or KOH as appropriate.

Table 2
Intracellular solutions used during electrophysiology experiments

<table>
<thead>
<tr>
<th>Intracellular (mM)</th>
<th>Na</th>
<th>K</th>
<th>Li</th>
<th>Ca</th>
<th>Cl</th>
<th>Mg</th>
<th>Cs</th>
<th>SO₄</th>
<th>MeSO₄</th>
<th>Gluonate</th>
<th>EGTA</th>
<th>BAPTA</th>
<th>TEA</th>
<th>HEPES</th>
<th>pH</th>
<th>7.2</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. Inside-out patch</td>
<td>188</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>10</td>
<td>0</td>
<td>0</td>
<td>90</td>
<td>0</td>
<td>0</td>
<td>5</td>
<td>0</td>
<td>10</td>
<td>10</td>
<td>NaOH</td>
<td></td>
</tr>
<tr>
<td>2. Lithium solution</td>
<td>5</td>
<td>5</td>
<td>140</td>
<td>0</td>
<td>145</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>10</td>
<td>NaOH</td>
<td></td>
</tr>
<tr>
<td>3. Cell attached patch bath solution</td>
<td>0</td>
<td>150</td>
<td>0</td>
<td>0</td>
<td>28</td>
<td>1</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>115</td>
<td>5</td>
<td>0</td>
<td>0</td>
<td>10</td>
<td>KOH</td>
<td></td>
</tr>
<tr>
<td>4. Physiological saline</td>
<td>160</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>150</td>
<td>0</td>
<td>0</td>
<td>5</td>
<td>0</td>
<td>10</td>
</tr>
<tr>
<td>5. NaMeSO₄</td>
<td>160</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>150</td>
<td>0</td>
<td>0</td>
<td>5</td>
<td>0</td>
<td>10</td>
</tr>
<tr>
<td>6. ENaC Permeability</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>5</td>
<td>0</td>
<td>0</td>
<td>10</td>
<td>NaOH</td>
<td></td>
</tr>
</tbody>
</table>

aNo intracellular solutions in this configuration. bNo intracellular solutions were used for the volume recording experiments since cells were intact. c[Na] and [K] were varied between 22 and 150 mM in these experiments to determine their relative permeability (see Figure 3). All ion concentrations are the total number of ions in the solution after adjustment of pH with NaOH or KOH as appropriate.

Results

Functional expression of ENaC/Deg-like channels in canine chondrocytes

We began by showing that our isolated canine chondrocytes expressed positive immunoreactivity for all three subunits of ENaC (α-ENaC, β-ENaC and γ-ENaC; Figure 1A–F). Next, in potassium-free solutions (‘Inside-out patch’: Tables 1 and 2), we observed a small ion channel in 60% of patches with a slope conductance of 9 ± 1 pS (n = 5) (Figure 1G, 1). This reversed near to the equilibrium potential for sodium calculated for these conditions (~4.9 mV) (Figure 1I), and the open probability (Po) was significantly reduced by the ENaC/Deg blockers, 10 μM amiloride (Figure 1I) and 100 nM benzamil (inhibition of 96 ± 2%, n = 5 and 56 ± 2%, n = 3, respectively; P ≤ 0.05 in both cases, ANOVA). The kinetics of this low conductance channel was typical for ENaC channels. To quantify this, we performed a kinetic analysis. Since ENaC channels have an unusually high permeability for Li+ (Canessa et al., 1994; Schild et al., 1997; Kellenberger et al., 1999), we optimized conditions by recording events under cell-attached patch mode with Li+ included in the pipette as the charge carrier (‘Lithium solution’: Table 1). Under this configuration, and with membrane potential of -80 mV, the unitary currents were <1 pA inward, again typical of ENaC/Deg channels. Data was idealized and model fitted as described in the methods. Initially, we found events could not be satisfactorily fitted with a simple two-state model (closed–open) but was much better fitted with two open and two closed states (Figure 2). Best fit rate constants are given in full in Table 3. These rate constants were used to calculate the PDFs, mean open and mean closed times in Figure 2.

We next investigated whether constitutive activity of these channels influenced the cellular membrane potential (Vm). We used standard physiological saline solutions (extracellular: Table 1, intracellular: Table 2) and measured Vm in current clamp mode. Control Vm was ~13.2 ± 3.8 mV, similar
Immunohistochemistry and patch-clamp recording reveal the presence of ENaC protein expression and a benzamil and amiloride sensitive cation conductance. Immunofluorescence staining of αENaC (A), βENaC (B) and γENaC (C) in primary cultures of canine articular chondrocytes (first passage cells, 100× objective). Positive immunoreactivity for all subunits was observed although strongest for α- and β-ENaC. The concentration of the primary antibodies was 1 μg·mL⁻¹ (dilution 1:200). The secondary antibody was a goat polyclonal to rabbit IgG (Fc-specific, affinity purified, pre-adsorbed) conjugated to DyLight® 488 (Abcam ab98462). After extensive washes in PBS-T, the nuclei were counterstained with propidium iodide (red fluorescence). Green staining shows the ENaC protein, orange staining is the nuclei and yellow staining is the overlap of the two. (D–F) The negative control was treated identically, but with the omission of primary IgG and shows the red fluorescent nuclear staining with very little background green fluorescence (scale bars for A–F: 10 μm). (G–J) Inside-out patch clamp recordings from canine chondrocytes using inside-out solutions: Tables 1 and 2. (G) Traces of inside-out low conductance single channel activity at the given membrane potentials. The scale bar horizontal line is 100 ms; vertical line is 500 fA. (H) All-points amplitude histogram for the low conductance channel at −40 mV. (I) Single channel current–voltage curve for the low conductance channel. \( V_{rev} \) was −1 ± 5 mV (\( n = 5 \)), slope conductance 9 ± 1 pS (\( n = 5 \)). (J) Open probability \( (P_o) \) versus time, calculated over successive 0.4 s windows before and during the addition of the ENaC channel inhibitor, amiloride (10 μM). Low conductance single channel \( P_o \) was reduced by 96 ± 2% (\( n = 5 \)).
to previously reported resting membrane potentials in chondrocytes (Wright et al., 1992, Lewis et al., 2011a). Application of 10 mM amiloride caused the \( V_m \) to become significantly more negative (Figure 3A, 9.5/±0.8 mV, \( n = 5 \)).

Addition of 100 nM benzamil (Figure 3B), an even more specific inhibitor of ENaC (Kellenberger and Schild, 2002; Alexander et al., 2011), also significantly changed \( V_m \) (7.5/±1.7 mV, \( n = 5 \) hyperpolarization, \( P < 0.05 \) ANOVA). These effects were reversible and dose-related (Figure 3A–C).

To determine the sodium permeability of the benzamil-sensitive current (relative to \( K^+ \)), we switched to whole-cell voltage clamp. With the standard physiological solutions and at a holding potential of \(-15\) mV, application of 1 \( \mu \)M benzamil blocked a small constitutive inward current and resulted in an outward current deflection (mean conductance \( 1.5 \pm 0.4 \) nS, \( n = 9 \), Figure 3). Whole-cell voltage ramps were then run in the presence and absence of benzamil, using four different intracellular sodium concentrations optimized for

### Table 3

<table>
<thead>
<tr>
<th>From State 1</th>
<th>To State 2</th>
<th>( K (s^{-1}) )</th>
<th>Mean</th>
<th>SEM</th>
</tr>
</thead>
<tbody>
<tr>
<td>Closed (2)</td>
<td>Open (3)</td>
<td>183.4</td>
<td>21.8</td>
<td></td>
</tr>
<tr>
<td>Open (3)</td>
<td>Closed (2)</td>
<td>203.8</td>
<td>5.1</td>
<td></td>
</tr>
<tr>
<td>Closed (2)</td>
<td>Closed (1)</td>
<td>19.3</td>
<td>3.0</td>
<td></td>
</tr>
<tr>
<td>Closed (1)</td>
<td>Closed (2)</td>
<td>14.6</td>
<td>3.7</td>
<td></td>
</tr>
<tr>
<td>Open (3)</td>
<td>Open (4)</td>
<td>41.7</td>
<td>17.0</td>
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</tr>
<tr>
<td>Open (4)</td>
<td>Open (3)</td>
<td>38.6</td>
<td>9.8</td>
<td></td>
</tr>
</tbody>
</table>

State numbers are in parenthesis (see model in Figure 2).

Figure 2

Kinetic analysis of the low conductance cation channel. Kinetic analysis of the ENaC/Deg-like low conductance was performed under cell-attached patch conditions. \( \text{Li}^+ \) was included in the patch-pipette and served as the charge carrier (Solutions: Table 1). (A) Upper panel, the SKM idealized record from this trace (see Methods). Lower panel, raw single channel trace at \(-80\) mV, baseline corrected. (B) An example patch closed-time distribution superimposed with a bi-exponential PDF calculated from panel D: \((\tau_1, \text{area})\): 5.9 ms, 0.83, \((\tau_2, \text{area})\): 94.5 ms, 0.17. Mean: 21.0 ms.

(C) An example open-time distribution, again fit with a bi-exponential PDF calculated from panel D: \((\tau_1, \text{area})\): 3.1 ms, 0.5, \((\tau_2, \text{area})\): 26.5 ms, 0.46. Mean: 13.8 ms. (D) Schematic of our best fit kinetic model, full data in text (Table 3). Throughout this figure, red is used to denote closed states, and blue denotes open state.
recording ENaC currents (22, 66, 110 and 150 mM). Subtraction of the whole-cell currents in the presence of benzamil from those recorded in vehicle gave the benzamil difference currents shown in Figure 3F. Difference currents at four different intracellular sodium concentrations were obtained (Figure 3F, G), and the reversal potentials for these followed the calculated changes in Na⁺ equilibrium potential (Figure 3G).

Finally, we investigated whether ENaC/Deg channel activity contributed to canine chondrocyte RVI. In cell-attached
**Discussion**

In this work, we show that canine chondrocytes express ENaC subunit protein and exhibit single channel activity electrophysiologically and pharmacologically, characteristic of ENaC/Deg channels. Block of these channels significantly alters membrane potential and reduces RVI following osmotic shrinkage.

**Identification of ENaC protein and ENaC/Deg-like single channel activity**

Several previous studies have demonstrated that isolated chondrocytes exhibit a characteristically differentiated phenotype for the first few passages in culture (Benya and Shaffer, 1982). ENaC protein expression (Trujillo et al., 1999) and RNA transcription (Karlsson et al., 2010) have previously been shown in human chondrocytes. We used reliable and well-tested antibodies that have been used in numerous studies of ENaC (Trujillo et al., 1999; Rubera et al., 2003; Shakhibaei and Mobasheri, 2003; Zhang et al., 2005) and found positive immunoreactivity for all three ENaC subunits in canine chondrocytes. Immunostaining did not appear limited to the membrane. There could be several technical reasons for this without need to infer that the majority of protein was cytoplasmic. For immunohistochemical analysis, cells were grown on coverslips and thus morphologically very flat and photographed with epifluorescence (not confocal). These conditions make it difficult to distinguish between membrane and cytoplasmic staining. Second, the standard methods we adopted for fixation/permeabilization vigorously disrupted the membrane with both methanol and detergent. Similar patterns of whole-cell staining have been observed with many membrane ion channels antibodies including those directed against ENaC (Trujillo et al., 1999; Mobasheri et al., 2004; Sauter et al., 2006; Teruyama et al., 2012). Our patch-clamp data show ion channels with physiology and pharmacology consistent with that of ENaC/Deg channels. First, at the single channel level, our measured reversal potential was suggestive of selective Na+ permeability (Smith and Benos, 1991; Garty and Palmer, 1997), and this was supported by explicit calculation of the permeability ratio $P_{Na}/P_{K}$ value. In our calculation of permeability ratio, we find that the change in $V_{rev}$ is rather flat at intracellular Na+ ($[Na]_i$) concentrations between 60 and 150 mM, but $V_{rev}$ rises steeply as $[Na]_i$ falls below 60 mM. This suggests that at the benzamil concentrations used here (1 μM, much greater than our calculated benzamil $EC_{50}$) there may be a contamination of the ENaC/Deg conductance with a non-specific cation conductance. This would lead to an underestimate of the permeability ratio. Despite this, we calculate a high selectivity for sodium (24:1), very similar to that expected for ENaC (Alexander et al., 2011). Furthermore, these channels had the low conductance that is a key feature of ENaC/Deg channels and ENaC in particular (Kellenberger and Schild, 2002). The value of ENaC single channel conductance depends very much on the subunit composition and the conditions (Kellenberger and Schild, 2002), but our measured value of 9pS lies well within the previously reported range (Canessa et al., 1994; Waldmann et al., 1995; Awayda et al., 1996; Fyfe and Canessa, 1998; Ishikawa et al., 1998; Caldwell et al., 2005). In any set of...
RVI is retained in lithium, but sensitive to low concentrations of benzamil. (A) Confocal images showing changes of cell volume with time. The top panels show the chondrocyte from above (X–Y plane), and the lower panels show the reconstructed cell view from the side (Z–X plane). In order to capture relatively rapid volume changes, scan resolution was set to a low value (31 Z layers per time slice). These images were necessary to verify that cells were approximately spherical (please see Methods: Equation 1). 60× objective scale bar 20 μm. (B) Under control conditions, when cells are incubated in 309mOsm physiological saline (Table 1) then exposed to hypertonic solutions (489mOsm), they first shrink passively as water leaves the cell via osmosis (marked ‘I’). They then swell through the process of RVI, eventually returning to the starting volume (RVI indicated by ‘II’). Osmolarity was increased with addition of sucrose. Not all cells underwent RVI. We use a two-exposure protocol. Only those cells that exhibited RVI on the first exposure to hypertonic challenge were then used for these experiments. The second exposure either contained benzamil or further control vehicle solution, mean of eight cells. (C) The same protocol as that used in panel B, but with the presence of 100 nM benzamil; RVI is inhibited. Mean of five cells. (D) RVI protocol with extracellular Na$^+$ replaced by Li$^+$. For (B) to (D) solid filled markers indicate data points and the solid line (and y-axis on the right) indicates the osmolarity. (E) Concentration inhibition curve for inhibition of RVI by benzamil. Five cells per concentration. Data are represented as fractional recovery from the theoretical maximum shrinkage (0.63 = 309/489) and fit with Equation 3, where $R$ is fractional recovery, $m$ was 0.84 ± 0.2, $h$ is 3.5 ± 0.9 and $pD_2$ = 7.5 ± 0.01. (F) Schematic of our working model for RVI, with phases ‘I’ and ‘II’ indicated.
electrophysiological experiments on native cells (unlike a recombinant expression system), there is the possibility of mixed populations of channels. In this study, however, the single channel properties, together with the pharmacological profile of these channels (high sensitivity to amiloride and benzamil), are strongly suggestive of the principle component of the low conductance cation channel being ENaC. Acid-sensing channels, specifically ASIC1a (Yuan et al., 2010b), are the only other members of the ENaC/Deg family we are aware of, which have been shown to be present in mammalian chondrocytes. Whilst amiloride (Waldmann et al., 1997; Yuan et al., 2010a) and benzamil (Waldmann et al., 1997) can inhibit these channels at sufficient concentration, benzamil IC50 for ASIC1a (Waldmann et al., 1997) is approximately 1000x that of the value we calculated for inhibition of the ENaC/Deg channel in this study (benzamil dVm EC50 calculated from the fitted pD2 is equivalent to approximately 4 nM). We found that the inhibition of inside-out patch clamp recorded events were apparently less sensitive to benzamil inhibition than the membrane potential, which would be consistent with an external binding site for ENaC inhibition (Schlied et al., 1997). The identity of the low conductance cation channel studied in the present study is therefore highly likely to be ENaC.

The single channel kinetic properties we measured were similar, but not identical, to those of ENaC in other cell types; resting open probability was somewhat lower than reported for ENaC in Xenopus laevis oocytes (Fyfe and Canessa, 1998), but similar to that seen in rat cortical collecting tubules (Palmer and Frindt, 1986). Some previous studies found a satisfactory fit with a single open and closed state (Palmer and Frindt, 1986), however, we achieved a much better fit with two open and two closed states. Palmer and Frindt (1986), for example, saw virtually no short events at all, whereas we saw many. That these differences represent differences in filtering/fitting algorithms is unlikely, but more likely to reflect different subunit expression between systems (Fyfe and Canessa, 1998). Kellenberger et al. (2002) used similar filtering conditions to us and also adopted a four-state model to describe their β5S18C point mutant ENaC channels. They did not calculate model rate constants, but their mean open and closed states were both considerably shorter than those we observed. The absolute rate constants presented in our schema (Figure 2D) should be viewed with some caution, since single channel kinetic studies with a 9pS channel are likely to be more greatly affected by noise than larger channels. In order to have sufficient signal to noise ratio to perform the study of these very small channels, we had to digitally re-filter the data to 0.5 kHz, rather lower than would usually be desirable for a single channel kinetic analysis study. Whilst the Qin algorithms (Qin, 2004) we used take this into account, it seems logical that the accuracy is less than for a larger channel.

**Role of benzamil-sensitive ion channels in volume control**

When cells capable of RVI are exposed to hypertonic solutions, three processes take place: first, cells shrink as water leaves the cell along its osmotic gradient; second, the shrinkage in some way activates an influx of ions; and third, water re-enters the cell along this new osmotic gradient until equilibrium is restored (reviewed by Hoffmann et al., 2009). The second and third processes occur in parallel and comprise RVI (‘II’ in Figure 5A). Whilst the water fluxes are generally accepted to be conducted by aquaporin ‘water channels’, there is little consensus as to the nature of the shrink-activated influx of ions. It seems likely that it is different from cell type to cell type. ENaC/Deg has previously been shown to be an important component of RVI in rat hepatocytes (Wehner et al., 2000; 2006).

However, despite the importance of volume regulation to chondrocytes, there have been only a few previous studies on RVI in chondrocytes themselves. Activity of the Na⁺-K⁺-2Cl⁻ membrane transporter (NKCC) appears to be involved both in chondrocytes (Kerrigan et al., 2006; Bush et al., 2010) and the related C-20/A4 cell line (Qous et al., 2011), but the Na⁺-Ca²⁺ exchanger (NCX) is also implicated, since its block prevents Ca²⁺ ion changes in response to hypertonic challenge (Sanchez and Wilkins, 2004). We certainly do not rule out a contribution of these transporters to volume control, but since benzamil inhibits RVI at well below the concentration required to block NCX, our data are also consistent with a central role for ENaC/Deg. The fact that the EC50 for benzamil block of RVI was 3- to 10-fold greater than for block of the ion channel conductance would support a small contribution from a non-ENaC, but benzamil-sensitive conductance. Our working hypothesis is that volume control is a complex process that requires a number of ion channels and transporters to both maintain an appropriate membrane potential and electrochemical gradient to support volume control. Since chondrocytes exhibit both RVD (K⁺ efflux) and RVI (Na⁺ influx), one would anticipate that optimal RMP would be midway between EK and ENaC. This is consistent with the values we, and others, have found previously (Wright et al., 1992; Lewis et al., 2011a) and in the present study (−13 mV). One would expect EK for a chondrocyte to be around −80 mV and ENaC to be greater than +90 mV (extracellular Na⁺ > 200 mM, Urban et al., 1993), thus driving forces for K⁺ efflux and Na⁺ influx are approximately 70 and 100 mV respectively. Therefore, interestingly, whilst the ENaC/Deg channels have a low conductance, in terms of RVI, there is a huge driving potential for Na⁺ influx.

There are therefore at least three ways in which ENaC/Deg block could affect volume regulation: (i) The level of the membrane potential has a strong influence on volume control (Lewis et al., 2011a), and so by hyperpolarizing the cell, ENaC/Deg block could result in a reduced driving force for a hypothetical anionic osmolyte channel. In our experiments, RVI was blocked by 100 nM benzamil, however, which only hyperpolarized the membrane by 8 mV so this would seem unlikely. (ii) ENaC/Deg could be essential for maintaining the electrochemical gradient, and its block could indirectly result in a change such that volume regulation is no longer possible. (iii) ENaC/Deg could open in response to mechanostimulation and allow sufficient Na⁺ influx to drive RVI. To investigate this possibility, we measured ENaC/Deg channel activity in cell-attached patch mode whilst applying hypertonic saline. Channel activity increased markedly supporting this third alternative mechanism. The mechanism of the opening of ENaC/Deg channels in response to hypertonic solution remains to be determined.
This could again be a direct ‘shrink’ activation of the channels as previously reported for the ENaC/Deg-like channels in macrophages (Gamper et al., 2000), or the opening in response to a change of intracellular concentration following the cell’s response to the hypertonic solutions. Activity of ENaC/Deg is well known to be sensitive to the ionic environment (Anantharam et al., 2006).

**Functional significance of volume control in chondrocytes**

A number of cell and molecular changes have been shown to take place in cartilage during onset of osteoarthritis (OA). These changes are complex but include changes in proteoglycan content (Venn and Maroudas, 1977; Brocklehurst et al., 1984), loss of collagen (Maroudas, 1976; Venn and Maroudas, 1977), increases in water content (Bollet and Nance, 1966; Mankin and Thrasher, 1975; Brocklehurst et al., 1984; Grushko et al., 1989; Chou et al., 2009) (decreases in osmolarity) and swelling of the chondrocytes themselves (Jones et al., 1999; Bush and Hall, 2003). The sequence of these changes is unknown. However, a decrease in osmolarity leads to increased vulnerability of chondrocytes to physical damage (Bush et al., 2005). Furthermore, chondrocytes from OA cartilage exhibit poor RVD (Jones et al., 1999), and so it has been suggested that inappropriate increases in chondrocyte volume may contribute to the progression of osteoarthritis (Bush and Hall, 2003). Transgenic deletion of TRPV4, an ion channel critical to volume control in chondrocytes, also predisposes chondrocytes to physical damage (Clark et al., 2010a). Since chondrocytes have robust systems in place to control cellular volume following changes of osmolarity, it seems unlikely that the increase in cellular volume of OA chondrocytes is a simple result of increased cartilage water content. Another possibility is that the changes which take place in cartilage include changes in ion channel expression per se. A number of such changes have been reported; a recent transcriptomic analysis showed a total of three ion channels transcript levels increased in chondrocytes from OA cartilage KCNMA1 (large Ca\(^{2+}\)-activated potassium channel), KCNN4 (small/intermediated Ca\(^{2+}\)-activated potassium channel) and TMEM16A (a Ca\(^{2+}\)-activated chloride channel) (Karlsson et al., 2010; Lewis et al., 2011b). They also observed a very significant decrease in the transcript abundance of SCNN1A (ENaC \alpha\-subunit) (ibid). This observation supports the observation of decreased \(\alpha\)-ENaC protein expression previously observed in OA chondrocytes (Trujillo et al., 1999). Whether these changes, or changes in other as yet unidentified ion channels lead to progression of OA, or results from it, remains to be determined. Future studies on models of cartilage degeneration and OA will be necessary to examine whether pharmacological intervention in the processes of chondrocyte volume regulation can influence the progression of OA.

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**References**


**Conflicts of interest**

The authors have no conflicts of interest.
Benzamil sensitive ion channels in chondrocytes


