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Review Review **TRPV channels as temperature sensors Christopher D. Benham*, Martin J. Gunthorpe, John B. Davis** *Neurology and GI Centre of Excellence for Drug Discovery, GlaxoSmithKline Research and Development Ltd., New Frontiers Science Park (North), Third Avenue, Harlow, Essex CM19 5AW, UK* Received 5 February 2003; accepted 10 February 2003

9 Abstract

10 The past year has seen a doubling in the number of heat-sensitive ion channels to six, and four of these channels are from the TRPV family. These channels characteristically have Q_{10} values of >10 above the thermal threshold, very different from the Q_{10} values of 1.5–2.0 11 seen in most ion channels. Cells expressing TRPV1 show similar temperature sensitivity to small capsaicin-sensitive nociceptor neurons, 12 consistent with these neurons expressing homomers of TRPV1. A- δ fibres exhibit properties that may be explained by TRPV2 containing 13 channels which is present in large diameter sensory neurons that do not express TRPV1. TRPV3 has a lower temperature threshold and 14 may contribute to warm-sensitive channels together with TRPV1. Warm sensation may also be transduced by TRPV4 expressing sensory 15 neurons and hypothalamic neurons. We can now look forward to further work defining the properties of the recombinant channels in more 16 17 detail and a re-analysis of endogenous i_{heat} currents in thermosensitive neurons and other cells. Data from the study of mice in which TRPV2, TRPV3 or TRPV4 have been deleted are also eagerly awaited. 18

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20 Keywords: TRPV channels; Sensor; Neuron

21 1. Introduction

Temperature sensing is important in all animals. Mam-22 mals require precise assessment of body temperature for set-23 ting internal thermoregulation, while cold-blooded animals 24 need to sense internal body temperature and to sense warm 25 and cool surroundings to regulate their behaviour in seeking 26 warming or cooling environments. In addition, all animals 27 depend on the rapid sensation of noxious heat to activate 28 rapid avoidance reflexes. 29

In principle, sensitivity to small temperature changes 30 could be conferred by incorporating any biological reac-31 tion with high entropy into a signal transduction cascade. 32 In Caenorhabditis elegans there is evidence that levels of 33 cGMP in a sensory neuron confer thermosensation through 34 their gating of tax4 the C. elegans analogue of the mam-35 36 malian cyclic nucleotide gated channel [1]. In mammals 37 also, ion channels seem to be the main signal transduction mechanism for thermosensation, but at least some of these 38 mammalian channels are directly gated by heat. The prop-39 erties of all ion channels are affected by temperature but ef-40 fects are modest, usually resulting in small linear increases 41

in current flow with Q_{10} values of around 2, where Q_{10} is 42 the change in rate of the reaction resulting from a 10 °C 43 rise in temperature. However, the heat-sensitive channels 44 are characterised by having gating mechanisms that show a 45 much greater sensitivity for heat than standard biochemical 46 reactions and have Q_{10} values much greater than 2. 47

Of the six molecularly defined heat-sensitive channels, 48 one is a two-pore domain potassium channel, TREK-1 49 [2] while the other five are all cation channels from the 50 TRP family. The TRPV or vanilloid sub-family, currently 51 comprising six members, four of which sense temper-52 atures around and above body temperature. The fifth 53 temperature-sensitive TRP channel is TRPM8 [3,4] from the 54 melastatin sub-family which functions as a cold sensor, re-55 sponding to decreases in temperature below 22 °C. This re-56 view summarises recent work on the temperature sensitivity 57 of the TRPV family of ion channels and attempts to corre-58 late these properties with endogenous temperature-sensitive 59 currents in native tissues. 60

2. Properties of recombinant heat-sensitive61TRPV channels62

The TRPV channels are a sub-group of the TRP family 63 of cation channels [5,6]. Structurally, these channels share 64

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homology with potassium channels. Each protein sub-unit 65 has six trans-membrane domains and recent experimen-66 tal work confirms the expectation based on analogy to 67 potassium channels that the functional channels are likely 68 composed of tetramers [7]. Homologous genes have been 69 identified in C. elegans, osm-9 is the most closely related 70 gene to mammalian TRPV1. Interestingly, osm-9 gene 71 product functions in C. elegans as an osmosensor and plays 72 no part in thermosensation which is performed by a single 73 thermosensitive sensory neuron, the AFD cell [1]. A recent 74 functional characterisation of two of four further TRPV ho-75 mologues in C. elegans indicated that they too functioned 76 as part of the osmosensing pathway [8]. Two open reading 77 frames of genes that fall within the TRPV family have been 78 identified in the Drosophila genome [8] but their function 79 is not known. The exquisite temperature sensitivity of the 80 mammalian TRPVs reviewed here may thus be an adap-81 tation during the more recent evolution of warm-blooded 82 animals. 83

Assessing the temperature sensitivity of an ion channel 84 may be approached in an analogous manner to charac-85 terising its voltage dependent properties. The steady state 86 properties of the current at different test potentials can be 87 examined using an instantaneous voltage clamp or alterna-88 tively, a voltage ramp may be applied to rapidly determine 89 the current flow at a range of potentials. Clearly, the lat-90 ter is most useful when channels show little in the way 91 of time dependent changes in gating at different poten-92 tials. Both temperature ramps and temperature jumps have 93 been used to measure thermal sensitivity (cf. [9,10]) but 94 not usually both techniques in the same laboratory. As the 95 TRPV channels show slow time dependent temperature 96 responses these properties will affect the data generated 97 by the two stimulus protocols. This should be borne in 98 mind when comparing properties of the channels listed in qc Table 1. 100

3. TRPV1

The expression cloning of the capsaicin-sensitive vanil-102 loid receptor was a ground breaking landmark [11] from 103 which subsequent work on the molecular basis of the other 104 temperature-sensitive TRP channels followed. TRPV1 is 105 a Ca^{2+} permeable cation channel activated by exoge-106 nous vanilloids such as capsaicin, but also by endogenous 107 lipid signalling molecules such as anandamide [12] and 108 eicosanoids [13]. As suspected, given the co-location of 109 vanilloid sensitivity and noxious heat gated currents in 110 small sensory neurons and the knowledge that capsaicin 111 evokes a "hot" sensation in humans, VR1 or TRPV1 can be 112 activated by noxious heat with a threshold of about 43 °C in 113 rat [11,14] human [15] or by inference in the mouse knock-114 out [16,17]. Rapid temperature jumps show that TRPV1 is 115 activated relatively rapidly with currents reaching a plateau 116 after less than 500 ms [15]. Thus, temperature ramps of 117 <0.5 °C s⁻¹ will report currents close to the steady state 118 values. This may explain the consistent responses reported 119 above, obtained from ramps or temperature jumps. 120

The temperature threshold for TRPV1 is not fixed but 121 modulated by chemical ligands and the phosphorylation state 122 of the channel. The various activating ligands have syner-123 gistic effects, so that any specific chemical ligand concen-124 tration will result in a unique setting of the temperature sen-125 sitivity of the channel. Thus, changes in endogenous lipid 126 ligand concentration might be expected to vary the thermal 127 sensitivity of the channels. The phosphorylation state of the 128 channels is also important. So, for example, phosphoryla-129 tion of TRPV1 by protein kinase C results in activation of 130 the channel at normal body temperature [18]. This plasticity 131 potentially confers a broad range of temperature sensitivity 132 to cells expressing TRPV1. Thus, while responses in recom-133 binant expression systems such as HEK293 cells in standard 134 culture conditions may be quite consistent, there is much

Table 1

Properties of ihe	_{eat} in	recombinant	systems	expressing	TRPV	subunits
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Channel expressed	TRPV1	TRPV2	TRPV3	TRPV4 VRL-2, VR-OAC, OTRPC4, trp12	
Pseudonyms	VR1	VRL-1	VRL-3		
High expression	DRG, TG	DRG, TG	DRG, TG, skin	TG	
TRPV heteromers	1 and 3	Not 2 and 1	3 and 1	?	
$p_{\rm Ca}/p_{\rm Na}$	9.6	2.9	12.1	6.3, 4-αPPD	
i _{heat}			2.6		
Heat threshold (°C)	>43	>53	>31	>24	
			>35	>33	
			>39		
Q_{10}	21	?	25 or 6.6	10 or 19	
Effect of prior heating	Sensitises/desensitises	Strongly sensitises	Strongly sensitises	Desensitises	
Threshold shift after pre-heating (°C)	~-7	-13	-4	+6	
<i>i</i> _{heat} in isolated patches	Yes	Not tested	Yes	No	
$[Ca^{2+}]_i$	Desensitises	?	Desensitises	Blocks IC ₅₀ 0.4 µM	
Ruthenium red	Blocks	IC ₅₀ 0.6 μM	$IC_{50} < 1 \mu M$	Voltage dep block	
Capsazepine	Blocks	Inactive	Inactive	Inactive	

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more potential for a range of responses in native cells. This
must be borne in mind when attempting to explain native
currents in molecular terms.

138 4. TRPV2

An initial search of genomic databases for TRPV1 homo-139 logues yielded vanilloid receptor like protein 1 or VRL1, 140 now called TRPV2, that is expressed in a sub-population of 141 medium to large sensory neurons but also at lower levels 142 in other tissues. This family member was not activated by 143 any TRPV1 chemical ligand, but was activated by noxious 144 heat >53 °C resulting in a cation current that was Ca^{2+} 145 permeable. The current showed outward rectification at 146 positive potentials, like other TRP channels, but also in-147 ward rectification at very negative membrane potentials. 148 The temperature-evoked currents were specific to TRPV2 149 transfected cells whether oocytes or HEK293 cells, and po-150 tently blocked by ruthenium red (IC₅₀ 0.6μ M) consistent 151 152 with current flow through a specific ion channel pathway. Repeated heating resulted in sensitisation such that the cur-153 rent threshold moved to much lower temperature at around 154 40°C [19]. 155

This initial description provides a clear role for TRPV2 156 channels in sensing high threshold noxious heat. The 157 broader distribution suggests other possible functions. Inter-158 estingly, the murine form of TRPV2 was shown to be con-159 stitutively active at room temperature following treatment 160 of TRPV2 transfected CHO cells with insulin-like growth 161 162 factor (IGF-1) for a few minutes. The development of a 163 functional cation current correlated with the IGF-1 stimulated translocation of the channel protein to the cell mem-164 brane. The mouse isoform (79% homology to rat TRPV2 165 was also sensitive to noxious heat, currents being increased 166 by 140% compared to currents at room temperature [20]. 167

Attempts by several laboratories including our own to 168 measure temperature gated currents in TRPV2 transfected 169 cells have been unsuccessful, indicating that important as-170 pects of the functional expression of this channel remain to 171 be determined. The recent suggestion that soluble co-factors 172 are required to mediate heat responsiveness of TRPV4 (see 173 below) may be a clue to the variable success in evoking heat 174 gated currents from TRPV2 expressing cells. 175

176 5. TRPV3

Analysis of thermosensation in the TRPV1 null mouse 177 demonstrated virtually normal thermal nociception in the 178 absence of inflammation [16,17]. Sensing hot temperatures 179 could be ascribed to TRPV2 but there was clearly a need for 180 further molecular candidates for warm sensation in addition 181 to TRPV1. The virtual completion of the human genome 182 project provided genomic sequence in which to search for 183 any outstanding TRPV homologues. This search yielded 184

one further member, TRPV3, which is expressed mainly in 185 the CNS and sensory neurons in humans [9,10], but also 186 in skin and in particular in keratinocytes found at the in-187 ner boundary of the epidermis [21]. Applying temperature 188 ramps to CHO or HEK293 cells expressing TRPV3, evoked 189 a temperature-sensitive cation current with moderate perme-190 ability to Ca^{2+} and a high Q_{10} [10,21]. To date there is no 191 evidence that TRPV3 can be activated by any chemical lig-192 and. Heating isolated outside out patches from TRPV3 ex-193 pressing cells activated a cation channel of 172 pS unitary 194 conductance [22]. This suggests that direct heating of the 195 channel protein or at least a membrane delimited pathway 196 mediates channel opening. 197

Thermal sensitivity depended on the thermal history of the 198 cell and this may be one reason why there is some variation 199 in the reported threshold of activation of TRPV3, ranging 200 from 23 °C [10] through 35 °C [21] to 39 °C [9]. Repeated 201 warming sensitises the channel to heating, both increasing 202 the maximum current at the end of the temperature ramp 203 and shifting the temperature threshold to lower temperatures. 204 The effect of repeated heating to 48 °C on currents recorded 205 from TRPV1 and -3 expressing HEK 293 cells is shown in 206 Fig. 1. 207

All three groups used electrophysiological or Ca²⁺ fluo-208 rescence recording based on a baseline resting temperature 209 at room temperature. While, all cells will have been incu-210 bated at 37 °C post transfection, the duration held at room 211 temperature before commencing recording probably varied 212 and might be expected to affect the subsequent tempera-213 ture sensitivity. Prolonged incubation at 37 °C might also 214 select against cells expressing channels with low tempera-215 ture thresholds. Whatever the precise thermal sensitivity of 216 the channel, from a parallel comparison of TRPV1 and -3, it 217 appears that TRPV3 has a lower temperature threshold than 218 TRPV1 [9]. 219

A further complexity is the demonstration that TRPV3 220 can heteromise with TRPV1 when expressed in HEK293 221 cells. These heteromers may function with many of the poly-222 modal properties of TRPV1 including capsaicin and proton 223 sensitivity. Co-localisation in native DRG cell bodies sug-224 gests that this may happen in native cells [9]. Further care-225 ful comparison of TRPV1, -3 and -1/3 expressing cells will 226 be needed to confirm that TRPV1/3 heteromers are func-227 tional and identify any differences from TRPV1 homomers. 228 The profound differences between TRPV1 and -3 behaviour 229 (Fig. 1) may allow for heteromeric channels to be robustly 230 identified and characterised. 231

6. TRPV4

TRPV4 is the final member of the TRPV family with 233 reasonably close homology to TRPV1. TRPV5 and 6, the 234 ECAC channels are more distant cousins [6]. This TRPV4 235 channel was originally described as an osmosensor, opening 236 in response to hypotonic swelling of the cell [23,24]. While

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Fig. 1. Heat activation of TRPV1 and -3. (A) Expression of either TRPV1 or TRPV3 receptors alone in HEK293 cells generates heat-activatable ion channels with heat thresholds in the warm (TRPV3) or noxious (TRPV1) temperature ranges. Whole-cell patch clamp recordings of membrane currents in response to heat application (48 °C, or at room temperature, RT), for the duration of the bar, are shown. These traces typify the slower kinetics of TRPV3 receptor activation relative to TRPV1 [9]. It is also noteworthy that the inward heat-gated currents are also accompanied by an increase in current noise (variance) which is consistent with the stochastic activation of an ion channel of relatively high single channel conductance in both cases. (B) TRPV1 receptors typically desensitise in response to agonist stimulation such as capsaicin or acid. Heat activation of TRPV1 appears to cause similar effects. Repetitive stimulation of TRPV1 with supra-threshold heat stimuli (48-51 °C) led to pronounced receptor desensitisation (even in the nominal absence of external Ca²⁺) such that the magnitude of inward current responses after four test stimuli at approximately 1 min intervals were only 51% of the original control response. The behaviour of TRPV3 is quite different since TRPV3 responses actually increase with repeated stimulation at supra-threshold temperatures (43-47 °C), indicating a marked sensitization of this receptor by heat. TRPV3 responses increased by approximately 100% at each subsequent stimulus challenge such that current responses increased by about 10-fold (1246%) over the course of the experiment. The pooled datasets for TRPV1 (n = 3-6) and TRPV3 (n = 4) are shown in panel (C) and experimental conditions are as described previously [15,9] for TRPV1 and -3, respectively). The scale bars used are calibrated as follows: vertical, 100 pA; horizontal, 500 ms.

the initial reports suggested that the channel could not be 237

238 activated solely by a rise in temperature, it was reported that

responses to osmotic stress increased significantly at body 239 temperature compared to room temperature [24].

More detailed examination of the properties of TRPV4 241 expressing cells has shown that TRPV4 also acts as a ther-242 mosensor. Application of temperature ramps from 22 °C to 243 oocytes or HEK293 cells expressing rat TRPV4 resulted in 244



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rises in intracellular Ca²⁺ and inward currents with thresh-245 olds around 34°C. [25]. Starting from a lower holding 246 temperature of 14 °C, inward currents were activated with 247 a threshold of 24 °C in HEK293 cells expressing mouse 248 TRPV4. Careful comparison of the heat-evoked current 249 250 with that activated in the same cells by the TRPV4 agonist, $4-\alpha$ PDD [26], supported the conclusion that the heat acti-251 vated current was through TRPV4 channels [27]. In contrast 252 to TRPV1 [14] and TRPV3 [22] (Table 1), isolated patches 253 that contained functional TRPV4 channels showed no cur-254 rent activation when exposed to increasing temperature. 255 This suggests that additional soluble factors are required to 256 mediate thermosensation. Either heating produces a soluble 257 ligand or a soluble ligand is required as a co-activator [27]. 258

Over the range 24–36 $^{\circ}$ C, current increased with a Q_{10} of 259 19.1 and showed dramatic desensitisation towards the peak 260 of the ramp. A further manifestation of this property was that 261 repeated heat ramps evoked smaller responses with higher 262 thresholds [27] in contrast to the properties of the other TR-263 PVs (Table 1). Temperature sensitivity that spans normal 264 body temperature suggests that TRPV4 can respond to small 265 changes in body temperature around 37 °C but the rapid de-266 sensitisation properties make extrapolating these results to a 267 steady state 37 °C hazardous. However, acclimatising cells 268 to mammalian body temperature and then increasing tem-269 perature with a ramp produced further increases in intracel-270 lular Ca²⁺ indicating that TRPV4 expressing cells can sense 271 a change in temperature as seen in pyrexia [25]. 272

It will be interesting to generate comparative data using
the same protocols as in Fig. 1 for TRPV4 and for TRPV2.
Data summarised in Table 1 suggests that TRPV4 shows
strong desensitisation of responses after prior heating and

during a single heat ramp [27], while TRPV2 shows strong sensitisation [19]. Detailed information on the activation kinetics of these two channels is awaited. This should permit a more comprehensive comparison with the properties of native currents and aid identification of further functional heteromers. 282

7. Properties of heat-sensitive currents in sensory neurons

7.1. In vivo single unit recording

In vivo recording of the activity of single nerve fibres 286 in response to heating the skin has identified four types of 287 thermosensor with specific temperature sensitivities (Fig. 2) 288 (reviewed by [28]). Warming skin from around 30°C, the 289 skin temperature if the ambient temperature is 20 °C, first 290 excites a population of unmyelinated c-fibres which convey 291 a sensation of innocuous warmth (Fig. 2). This temperature 292 sensitivity indicates a role for TRPV3 as suggested by Peier 293 et al. [21] who believe that the localisation of this channel 294 in keratinocytes at the inner surface of the epidermis where 295 sensory nerve terminals terminate, is important to warm tem-296 perature signal transduction. They propose a signalling path-297 way that involves temperature sensation by the keratinocytes 298 that then excite nerve terminals by the release of a transmit-299 ter such as ATP. This neatly explains the lack of warm tem-300 perature responsive sensory neurons in DRG isolated from 301 TRPV $1^{-/-}$ mice [16,17]. However, this observation might 302 be expected given the low number of warm-sensitive primary 303 afferent fibres and the relatively small sample size in these



Fig. 2. Fibre firing rates plotted against temperature of the four types of thermosensors in somatic afferent nerves. For native fibres, temperatures reflect ambient temperature at the tissue surface, so that temperatures at nerve endings are likely to be a few degrees lower. C-polymodal nociceptors show leftward shifts in temperature threshold in response to inflammatory mediators (red line) as do TRPV1 expressing cells. Activation ranges of recombinant TRPV channels are indicated by the horizontal bars below *X*-axis. Note that there is some uncertainty over the activation thresholds for TRPV3 and -4 (see Table 1).

Table 2 Properties of native *i*_{heat} recorded from rat dorsal root ganglion cells in culture

Classification (references)	Tissue	Cell diameter mean (µm)	Threshold (°C)	Q_{10}	$P_{\rm Ca}/P_{\rm Na}~i_{\rm heat}$	Desensitisat-ion of heat response	Capsazepine block <i>i</i> _{heat}	Ruthenium red block IC ₅₀ (µM)
Low threshold capsaicin [30]	3-5 days rat DRG	<25	42	_	1.3	No		
Sensitive cells [37]			43	17.8		Yes, decrease threshold		
[31], [33]	Adult rat DRG	18	45		1.2, i _{capsaicin} , 2.4		No effect	
[32]		27.5				No	IC ₅₀ 13 μM 25% insensitive	>5
Low threshold capsaicin insensitive [38]	3 days rat DRG	<20	40	>10		No, decrease threshold, biphasic, respectively		
LT capsaicin sensitivity n.d. [29]	Adult rat DRG	<30	40				No effect at 10 µM	No effect at 100 μM
High threshold [31]	Adult rat	25	51					
Capsaicin insensitive [34]	DRG	30	51.6		3.5		$5\mu M$ blocks 55% only	0.3
						イ		

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studies, which may also explain why there is no detailed de-304 scription of warm responsive sensory neurons in dissociated 305 culture. Alternatively, culture conditions, age of neurons or 306 the absence of satellite cells might also explain this obser-307 vation. Study of TRPV3^{-/-} mice will be useful in further 308 309 exploring the role of TRPV3 as will direct recordings of heat activated currents from isolated keratinocytes. The temper-310 ature range in which TRPV4 is activated also fits well with 311 sensing warm temperatures, particularly the desensitisation 312 properties as noxious temperatures are approached (Fig. 2). 313 It is tempting to speculate that both channels may contribute 314 in some sensory pathways. 315

As skin temperature is elevated beyond 40 °C, the firing 316 rate of these c-fibres declines and a separate population of 317 polymodal c-fibres are excited that are also responsive to 318 noxious chemical and mechanical stimuli. This fibre phe-319 notype correlates well with the behaviour of isolated small 320 diameter sensory neurons that are responsive to capsaicin 321 (see below and Table 2), suggesting a role for TRPV1 in 322 thermosensation in these nerve endings. Further increases 323 324 in temperature successively recruit myelinated A- δ fibres at thresholds of about 46 and 53 °C, the latter having a similar 325 326 threshold to TRPV2. These types I and -II AMH fibres are also mechanosensitive resulting in the AMH (A, mechano 327 and heat sensitive) nomenclature. In vitro correlates of these 328 fibre phenotypes are seen in large diameter, capsaicin insen-329 sitive, sensory neurons, although clear distinction into two 330 types is less obvious (see Table 2). Thus, these precise ther-331 mal sensitivities suggest that multiple thermosensitive sen-332 sory transduction elements are involved rather than for ex-333 334 ample, graded expression of a single temperature-sensitive 335 element that might result in different temperature thresholds for firing, dependent on expression level. 336

337 7.2. In vitro single cell studies

Studies on sensory neurons in the late 1990s had suggested that there were multiple heat-sensitive channels contributing to noxious thermosensation. While small, capsaicin-sensitive, sensory neurons responded to temperatures >45 °C [29,30], larger diameter temperature-sensitive neurons, also distinguished by being capsaicin insensitive had temperature thresholds of around 51 °C [31].

The cloning of TRPV1 [11] and TRPV2 [19] rapidly provided a molecular basis for this diversity and immediately stimulated more detailed analysis of the properties of sensory neurons.

This recent work is summarised in Table 2. A number 349 of studies have now provided many details of the prop-350 erties of currents with activation threshold 40-45 °C that 351 can be evoked in dorsal root ganglion neurons that also re-352 spond to capsaicin (see citations in Table 2). The loss of 353 such currents in sensory neurons dissociated from TRPV1 354 null mice [16,17], provides strong evidence for an obliga-355 tory role of TRPV1 in the composition of channels carrying 356 these currents. However, there are inconsistencies in the de-357

tailed properties of the native currents which could be ex-358 plained if all the current is not carried by TRPV1 homo-359 mers. The pharmacology of native i_{heat} currents is variable 360 in capsaicin-sensitive neurons although the ligands used are 361 not ideal for performing definitive studies. So, for example, 362 capsazepine has been reported as a weak blocker or to have 363 no effect (Table 2). Similarly, ruthenium red block was poor 364 [32] or had no effect up to 100 µM [29]. Further, in isolated 365 patches at the single channel level, there is a poor correlation 366 between numbers of capsaicin activated and heat activated 367 channels consistent with some heterogeneity in the native 368 signalling units [33]. 369

In larger capsaicin insensitive DRG neurons, inward cur-370 rents activated by temperatures above 51 °C, with higher 371 Ca^{2+} permeability, are seen [31,34]. This sub-type of larger 372 sensory neurons express TRPV2 protein but are not im-373 munoreactive for TRPV1 [34]. These authors demonstrate 374 that the high threshold response is a specific current and not 375 due to non-selective membrane or protein destruction, be-376 cause the current is reversible, specifically blocked by ruthe-377 nium red and only observed in a sub-population of neurons. 378 Comparing the properties of these currents with those evoked 379 in HEK293 cells expressing VRL-1 (TRPV2) there are clear 380 similarities. In addition to general similarities in the inward 381 current properties, the temperature threshold, Ca²⁺ perme-382 ability and block by ruthenium are quantitatively almost 383 identical in the native and recombinant current (Tables 1 384 and 2). Taken with the localisation of TRPV2 to these large 385 diameter cell bodies, the evidence suggests that TRPV2 is 386 a major component of the high threshold current in DRG. 387 Clearly, more work is needed to understand the functional 388 expression of TRPV2-mediated currents as a number of 389 laboratories including our own have failed to replicate the 390 findings of Caterina et al. [19]. Some unidentified accessory 391 protein, which was endogenously expressed in the Julius 392 lab, is the most likely explanation. Alternatively, if as it 393 seems possible for TRPV4 (see above and [27]), TRPV2 394 thermosensitivity might depend on some endogenous intra-395 cellular ligand acting as a co-agonist. If so, this might also 396 explain the lack of response when tested in some expression 397 systems. 398

8. Properties of heat-sensitive currents399in mammalian thermostats400

In mammals maintenance of core body temperature is 401 achieved with the aid of multiple thermosensors present in 402 the pre-optic anterior hypothalamus (POAH), medulla ob-403 longata and spinal cord [35]. Extracellular recording from 404 POAH revealed a population of warm-sensitive neurons that 405 showed thresholds of 37 °C and firing rates above this tem-406 perature of about 5 spikes $s^{-1} \circ C^{-1}$. Other neurons either 407 showed no temperature sensitivity or were cold sensitive 408 with firing rates that declined on heating [36]. Voltage clamp 409 studies of the warm-sensitive neurons identified a cation cur-410 8

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rent reversing at 0 mV activated by increasing temperature 411 and cell-attached patch recording identified unitary inward 412 currents that were about 2 pA in amplitude at the resting 413 membrane potential [36]. As currents indicative of action 414 415 potential firing were seen in these patches, this would suggest the membrane potential was less negative than -45 mV. 416 Assuming a membrane potential of $-40 \,\mathrm{mV}$ and a reversal 417 potential of 0 mV gives a unitary conductance of \sim 50 pS for 418 this temperature-sensitive channel. 419

TRPV4 is expressed in the anterior hypothalamus [25] 420 and TRPV4 currents have similar temperature sensitivity 421 [25] to the POAH cell currents [36]. Watanabe et al. [27] 422 found a lower threshold for TRPV4 current activation but 423 interestingly the unitary currents through these recombinant 424 channels had a similar conductance (59 pS) to the native 425 POAH currents. TRPV3 also has appropriate temperature 426 sensitivity so it would be interesting to see if this channel was 427 expressed in hypothalamus. Direct measurement of the uni-428 tary conductance of this channel gave a value of 170-200 pS 429 [10] but this was at $+60 \,\mathrm{mV}$. At negative membrane poten-430 431 tials a unitary conductance consistent with the noise analysis derived estimate of 48 pS [9] is probable. Thus, data to date 432 support a role for TRPV4 in transducing temperature in 433 these hypothalamic neurons but does not exclude TRPV3. 434

9. Other heat-sensitive cells 435

The functional activation of TRPV4 expressed in vascular 436 endothelial cells [27] suggests that this channel may have 437 438 a role in local vascular responses to changes in tempera-439 ture. Elevating temperature above body temperature would be expected to activate channels, causing a rise in endothe-440 lial Ca²⁺ levels. This would stimulate release of vasorelax-441 ants resulting in local vasodilatation. Conversely, cooling 442 could lead to vasoconstriction as the basal tonic Ca^{2+} influx 443 through TRPV4 was reduced. This, then provides a theo-444 retical mechanism for mediating peripheral cardiovascular 445 responses to limb heating and cooling in mammals. It will 446 be interesting to test this hypothesis in intact tissues. 447

10. Conclusions: TRPVs and endogenous 448 heat sensation 449

The past year has added two new TRPV channels to the 450 451 collection of heat-sensitive channels. We most likely now have the complete set of cation channels with which to fully 452 explore the molecular basis for thermosensation in mam-453 malian cells. We can now look forward to further work defin-454 ing the properties of the recombinant channels in more de-455 tail, including the mechanism of heat sensation. This will 456 also guide and stimulate a re-analysis of endogenous i_{heat} 457 currents both in cells where i_{heat} has been described but also 458 in cells such as endothelial cells and keratinocytes that had 459 not, until now been thought of as thermosensors. Data from 460

the study of mice in which TRPV2, -3 or TRPV4 have been 461 deleted are also eagerly awaited. 462

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