

CACHD1 is an α2δ-like protein that modulates CaV3 voltage-gated calcium channel activity

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Research Articles: Cellular/Molecular

CACHD1 is an $\alpha 2\delta$ -like protein that modulates Ca_V3 voltage-gated calcium channel activity

Graeme S. Cottrell¹, Camille H. Soubrane¹, James A. Hounshell^{3,5}, Hong Lin¹, Venetia Owenson², Michael Rigby², Peter J. Cox², Bryan S. Barker^{3,5}, Matteo Ottolini³, Selvi Ince¹, Claudia C. Bauer¹, Edward Perez-Reyes^{4,5}, Manoj K. Patel^{3,5}, Edward B. Stevens² and Gary J. Stephens¹

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Corresponding authors: Gary J. Stephens, School of Pharmacy, University of Reading, Whiteknights, PO Box 228, Reading RG6 6AJ, UK. E-mail: g.j.stephens@reading.ac.uk Edward B Stevens, Pfizer Neuroscience and Pain Research Unit, Portway Building, Granta Park, Cambridge, CB21 6GS, UK. Email: edward.stevens@metrionbiosciences.com.

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¹University of Reading, Whiteknights Campus, Reading, RG6 6AJ.

² Pfizer Neuroscience and Pain Research Unit, Portway Building, Granta Park, Great Abington, Cambridge, CB21 6GS.

³Departments of Anesthesiology

⁴Pharmacology

⁵Neuroscience Graduate Program, University of Virginia, Charlottesville, VA, USA, 22908

1	CACHD1 is an α2δ-like protein that modulates Ca _V 3 voltage-gated calcium channel
2	activity
3	Abbreviated title: CACHD1 modulation of Ca _V 3 channels
4	
5	Graeme S. Cottrell ^{1*} , Camille H. Soubrane ^{1*} , James A. Hounshell ^{3,5} , Hong Lin ¹ , Venetia
6	Owenson ² , Michael Rigby ² , Peter J. Cox ² , Bryan S. Barker ^{3,5} , Matteo Ottolini ³ , Selvi Ince ¹ ,
7	Claudia C. Bauer ¹ , Edward Perez-Reyes ^{4,5} , Manoj K. Patel ^{3,5} , Edward B. Stevens ² , Gary J.
8	Stephens ¹
9	¹ University of Reading, Whiteknights Campus, Reading, RG6 6AJ.
10	² Pfizer Neuroscience and Pain Research Unit, Portway Building, Granta Park, Great
11	Abington, Cambridge, CB21 6GS.
12	Departments of ³ Anesthesiology, ⁴ Pharmacology, and ⁵ Neuroscience Graduate Program,
13	University of Virginia, Charlottesville, VA, USA, 22908.
14 15	*These authors contributed equally to this work.
16	Corresponding authors:
17	Gary J. Stephens, School of Pharmacy, University of Reading, Whiteknights, PO Box 228,
18	Reading RG6 6AJ, UK. E-mail: <u>g.j.stephens@reading.ac.uk</u> .
19	Edward B Stevens, Pfizer Neuroscience and Pain Research Unit, Portway Building, Granta
20	Park, Cambridge, CB21 6GS, UK. Email: edward.stevens@metrionbiosciences.com .
21	
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30	
31	Abstract
32	The putative cache (Ca ²⁺ channel and chemotaxis receptor) domain containing 1 (CACHD1)
33	protein has predicted structural similarities to members of the $\alpha2\delta$ voltage-gated Ca^{2+} channel
34	(VGCC) auxiliary subunit family. CACHD1 mRNA and protein were highly expressed in the
35	male mammalian CNS, in particular in the thalamus, hippocampus and cerebellum, with a
36	broadly similar tissue distribution to Ca_V3 subunits, in particular, $Ca_V3.1$. In expression
37	studies, CACHD1 increased cell-surface localization of Ca _V 3.1 and these proteins were in
38	close proximity at the cell surface consistent with the formation of CACHD1-Ca $_{V}3.1$
39	complexes. In functional electrophysiological studies, co-expression of human CACHD1
40	with $\text{Ca}_{\text{V}}3.1$, $\text{Ca}_{\text{V}}3.2$ and $\text{Ca}_{\text{V}}3.3$ caused a significant increase in peak current density and
41	corresponding increases in maximal conductance. By contrast, $\alpha 2\delta$ -1 had no effect on peak
42	current density or maximal conductance in either Ca _V 3.1, Ca _V 3.2 or Ca _V 3.3. Comparison of
43	CACHD1-mediated increases in Ca _V 3.1 current density and gating currents revealed an
44	increase in channel open probability. In hippocampal neurons from male and female E19 rats,
45	CACHD1 overexpression increased Ca _V 3-mediated action potential (AP) firing frequency
46	and neuronal excitability. These data suggest that CACHD1 is structurally an $\alpha2\delta\text{-like}$
47	protein that functionally modulates Ca _V 3 voltage-gated calcium channel activity.
48	
49	

50	Significance Statement
51	This is the first study to characterise the CACHD1 protein. CACHD1 is widely expressed in
52	the CNS, in particular in the thalamus, hippocampus and cerebellum. CACHD1 distribution
53	is similar to that of low-voltage-activated (Ca_V3 , T -type) calcium channels, in particular to
54	$\text{Ca}_{\text{V}}3.1$, a protein which regulates neuronal excitability and is a potential therapeutic target in
55	conditions such as epilepsy and pain. CACHD1 is structurally a $\alpha2\delta$ -like protein that
56	functionally increases $\text{Ca}_{\text{V}}3$ calcium current. CACHD1 increases the presence of $\text{Ca}_{\text{V}}3.1$ at
57	the cell surface, forms complexes with $\text{Ca}_{\text{V}}3.1$ at the cell-surface and causes an increase in
58	channel open probability. In hippocampal neurons, CACHD1 causes increases in neuronal
59	firing. Thus, CACHD1 represents a novel protein that modulates $\text{Ca}_{\text{V}}3$ activity.
60	
61	
62	

Introduction

63

64 The putative CACHD1 gene was identified following a systematic search for proteins with structural homology to $\alpha 2\delta$ VGCC auxiliary subunits. The human CACHD1 gene on 65 66 chromosome 1p31.3 encodes the putative protein CACHD1 and has many orthologs, 67 including in speciation as early as C. elegans (tag-180) and D. melanogaster (CG16868) (Anantharaman and Aravind, 2000). Despite only a 13-16% gene homology and a <21% 68 69 protein identity with the $\alpha 2\delta$ VGCC auxiliary subunits, there are several key structural 70 similarities between CACHD1 and $\alpha 2\delta$ in terms of the arrangement of protein motifs. $\alpha 2\delta$ 71 and Ca_Vβ subunits are described as auxiliary or accessory VGCC subunits that modulate cell-72 surface expression and biophysical properties of high-voltage-activated (HVA) Ca_V1 (L-type Ca²⁺ current) and Ca_V2 (P/O, N- and R-type Ca²⁺ current) VGCC major α1 subunits 73 (Dolphin, 2012; Dolphin, 2013). In particular, $\alpha 2\delta$ subunits are proposed to associate with 74 HVA channels within the secretory pathway to promote plasma membrane trafficking and, 75 76 consequentially, to contribute to synaptic abundance (Dolphin, 2012), transmitter release 77 (Hoppa et al., 2012) and to defining the extent of the active zone (Schneider et al., 2015). 78 $\alpha 2\delta$ -1 and $\alpha 2\delta$ -2 represent molecular targets of gabapentinoid drugs (Dooley et al., 2007). However, modulation of low-voltage-activated (LVA) Ca_V3 family (T-type Ca²⁺ current) by 79 80 existing $\alpha 2\delta$ and $Ca_V\beta$ auxiliary subunits has not been firmly established (Dolphin et al., 1999: Lacinová et al., 1999: Dubel et al., 2004). LVA currents are activated by small 81 82 depolarization to regulate excitability around the resting membrane potential and Cav3 83 channels have been proposed as therapeutic targets in diseases such as epilepsy and pain 84 (Perez-Reyes, 2003; Cheong and Shin, 2013; Powell et al., 2014; Snutch and Zamponi, 85 2017); therefore, knowledge of proteins that modulate Ca_V3 activity is paramount.

Here, we investigate the novel CACHD1 protein and	d test the hypothesis that
CACHD1 represents an $\alpha 2\delta$ -like protein that modulates Ca	v3 channels. We have previously
reported that, by contrast to $\alpha2\delta$, the CACHD1 subunit has	no clear effect on Ca _V 2.2
biophysical properties when co-expressed together with $\beta 2a$	a in expression system studies
(Soubrane et al., 2012). We characterise the expression of the	he CACHD1 gene in rat and
human tissue at the transcriptional and translational level, as	nd demonstrate that CACHD1,
but not $\alpha 2\delta$ -1, increases Ca_V3 (T-type) current density and	maximal conductance. CACHD1
increases Ca _V 3.1 channel levels at the plasma membrane an	d data were consistent with
CACHD1 forming complexes with Ca _V 3.1 at the cell surface	ee to increase channel open
probability. We further demonstrate that CACHD1 expressi	ion causes a functional increase in
T-type current-mediated excitability in hippocampal neuron	ns. Together, these data
demonstrate that CACHD1 is structurally an $\alpha 2\delta$ -like prote	in which functionally modulates
Ca _V 3 activity.	

Materials and Methods

RNA isolation and real-time polymerase chain reaction (PCR)

Tissue samples were dissected from 5 adult male Wistar rats (Harlan, UK) following isofluorane overdose and cervical dislocation, according to Home Office Animals (Scientific procedures) Act 1986, UK. Total RNA was extracted using an RNeasy kit (Qiagen, UK) with an on-column DNase I treatment. Additional total RNA samples from AMS Biotechnology (Abingdon, UK) originated from human male donors aged 24-65. RNA (500 ng) was reverse-transcribed and relative quantification of CACHD1 and α 28-1 transcripts was performed using SYBR green and custom-made validated primers. HPRT1 was used as housekeeping gene. Absolute quantification of CACHD1, α 28-1, -2, -3, Ca_V 2.2 and Ca_V 1, -2, -3 transcripts was evaluated using 'Best Coverage' Taqman probes (Applied Biosystems, UK) against a standard curve of plasmids containing human CACHD1 and a rat single stranded DNA standard curve.

Sample preparation for in situ hybridization and immunohistochemistry

Rat tissue was kindly donated by Dr Emilio Russo, University Magna Grecia of Catanzaro, Italy. Briefly, 6-month-old male rats were sacrificed by i.p. injection of pentobarbital (200 mg/kg) according to ARRIVE guidelines and local ethical approval committee of the University of Catanzaro and perfused-fixed with 4% PFA in RNAse-free PBS, pH 7.3. Brain tissue was extracted, post-fixed overnight in 4% PFA in RNAse-free PBS and then cryoprotected in 30% sucrose. After being processed to wax (Tissue-tek VIP), 5 μ m horizontal plane brain slices were cut using a microtome (Leica, UK).

124	In situ hybridization
125	A CACHD1 probe consisting of a cocktail of short 10-20bp oligonucleotides spanning ~1kb
126	was designed by ACDBio (USA) and $in situ$ hybridization was performed on 5 μm rat brain
127	sections using a RNAscope 2.0 FFPE-Red kit. Positive (POLR2A) and negative (DapB)
128	probes were run in parallel.
129	
130	Immunohistochemistry
131	Chromogenic immunohistochemistry was performed using antigen retrieval in citrate buffer
132	(Thermo, UK) for 10 min and 3,3'-diaminobenzidine (DAB) staining (ImmPACT, Vector
133	Labs, UK), dehydrated and mounted with DPX. Rabbit anti-CACHD1 (1:500) (Abcam, UK
134	Cat #AB75141, RRID: AB_1310016) with horseradish peroxidase-coupled anti-rabbit IgG
135	(ImmPRESS, Vector Labs, UK) was used to detect CACHD1 protein. Qualitative expression
136	of mRNA was evaluated with a brightfield microscope according to colour intensity of
137	labelled mRNA.
138	
139	Antibodies for biochemistry
140	The following antibodies were used: mouse anti-HA.11 (Cambridge Bioscience, UK; clone
141	16B12; Lot No. B220767, RRID: AB_10063630); rabbit anti-Na+/K+-ATPase (Novus
142	Biologicals, Abingdon, UK; NB100-80005, Lot No. YH02206, RRID: AB_2063297); mouse
143	anti-c-Myc (Sigma-Aldrich Cat# M4439, clone 9E10, Lot No. 087M4765V, RRID:
144	AB_439694), rabbit anti-c-Myc (Sigma-Aldrich Cat# C3956, Lot No. 016M4762V, RRID:
145	AB_439680), mouse anti-β-actin (Sigma-Aldrich Cat# A5441, Lot No. 028K4826, RRID:
146	AB_476744) and rabbit anti-CACHD1 (Sigma-Aldrich Cat# AV49592, Lot No. QC22258,
147	RRID: AB 1852421): goat anti-mouse or rabbit IgG coupled to horseradish peroxidase

148	(Stratech Scientific Limited, Newmarket, UK); donkey anti-mouse or rabbit coupled to
149	AlexaFluor488, 555 or 647 (Invitrogen, Paisley, UK). Note: We experienced vial-to-vial
150	variation with the rabbit anti-CACHD1 antibody for Western blotting during this study.
151	Although both vials were from the same Lot No. and specifically recognised CACHD1, the
152	vial used for Fig. 4D gave rise to more non-specific staining on HEK cell lysates than vial
153	used for Fig. 4A.
154	
155	Vectors and vector construction
156	The human CACHD1 construct was purchased from Origene (Rockville, MA, USA) and the
157	truncated clone completed by PCR. The subsequent open reading frame was then subcloned
158	into pcDNA5/FRT. An N-terminal Myc tag was inserted after the natural signal sequence
159	between Ala ³⁵ -Glu ³⁶ using standard PCR techniques. All constructs were sequenced to
160	confirm identity. Construction of the vector pcDNA5/FRT-HA-CLR-Myc-RAMP1 has been
161	described elsewhere (Cottrell et al., 2007).
162	
163	Cell maintenance and propagation
164	HEK293 tsA201 (HEK) cells were cultured in DMEM (Invitrogen, UK) containing 10% fetal
165	bovine serum (Biosera, UK) and maintained in 95% air, 5% CO ₂ at 37 °C.
166	
167	Cell-surface biotinylation
168	HEK cells were transiently transfected in 6 well plates using 3 μg DNA (ratio 2:1, GFP-
169	Ca _V 3.1-HA:CACHD1) using Lipofectamine ²⁰⁰⁰ (3:8, DNA:Lipofectamine ²⁰⁰⁰). HEK cells
170	transfected with empty vector (vector control, VC), VC + Myc-CACHD1, GFP-Ca _V 3.1-HA +
171	VC or GFP-Ca _v 3.1-HA + Mvc-CACHD1 were washed (3x PBS), incubated with 0.3 mg/ml

172	EZ-Link [™] -Sulfo-NHS-Biotin (Pierce, USA) in PBS (1 h, 4°C), washed (3x PBS) and cells
173	lysed in RIPA buffer (50 mMTris/HCl, pH 7.4, 150 mM NaCl, 5 mM MgCl ₂ , 1 mM EGTA,
174	10 mM NaF, 10 mM Na ₄ P ₂ O ₇ , 0.1 mM Na ₃ VO ₄ , 0.5% Nonidet P-40, peptidase inhibitor
175	cocktail (Roche, UK)), and centrifuged. Biotinylated proteins were recovered by incubation
176	with NeutrAvidin-agarose (30 μl, overnight, 4°C), pelleted, washed with RIPA buffer (3x 1
177	ml), boiled in Laemmli buffer and analyzed by SDS-PAGE and Western blotting.
178	
179	SDS-PAGE and Western blotting
180	Immunoprecipitations and whole cell lysates were separated by SDS-PAGE (6-9%
181	acrylamide), proteins transferred to PVDF membranes (Immobilon-P, Millipore, UK) and
182	blocked for 1 h at room temperature (1x PBS, 0.1% Tween ²⁰ , 5% non-fat milk powder
183	[blocking buffer]). Membranes were incubated with antibodies to HA (1:5,000), β -actin
184	(1:20,000), CACHD1 (1:1000), rabbit or mouse Myc (1:5000) or Na ⁺ -K ⁺ -ATPase (1:20,000)
185	(overnight, 4°C; blocking buffer). Membranes were washed for 30 min (1x PBS, 0.1%
186	Tween ²⁰) and incubated with appropriate secondary antibodies coupled to horseradish
187	peroxidase (1:10,000, 1 h, room temperature; blocking buffer). Immunoreactive proteins were
188	detected using enhanced chemiluminescence (BioRad, UK). Densitometric analysis was
189	performed using an ImageQuant-RT ECL imaging system (GE Healthcare, Chalfont St Giles,
190	UK) and analysed using ImageQuant TL software.
191	
192	Immunofluorescent detection of cell-surface proteins
193	HEK cells were transiently transfected in 12 well plates using 1 µg DNA (ratio 2:1, GFP-
194	Ca _V 3.1-HA:Myc-CACHD1) using polyethylenimine (PEI; 1:2, DNA:PEI). HEK cells
195	transfected with empty vector (vector control, VC), VC + Myc-CACHD1, GFP-Ca _V 3.1-HA +

VC, GFP-Ca _V 3.1-HA + Myc-CACHD1 or CLR•RAMP1 seeded onto coverslips and used for
experimentation after 48 h. Cells were washed twice with PBSCM, incubated in DMEM
containing 0.1% BSA and mouse anti-HA (1:100) and rabbit anti-c-Myc (1:500) antibodies
(1 h, 4°C), washed twice again with PBSCM and then fixed in 100 mM PBS containing 4%
paraformaldehyde (w/v), pH 7.4 (20 min, 4°C). Coverslips were incubated in blocking buffer
(1x PBS, 2% normal horse serum, 0.1% saponin) (30 min, room temperature (RT)) and then
incubated with appropriate secondary antibodies (1:2000, 2 h, RT). Coverslips were washed
(blocking buffer, 30 min, RT) and mounted using Vectashield containing DAPI.
Proximity ligation assays
HEK cells were transiently transfected in 12 well plates using 1 µg DNA (ratio 2:1, GFP-

HEK cells were transiently transfected in 12 well plates using 1 μg DNA (ratio 2:1, GFP-Ca_V3.1-HA:Myc-CACHD1) using polyethylenimine (PEI; 1:2, DNA:PEI). HEK cells transfected with empty vector (vector control, VC), VC + Myc-CACHD1, GFP-Ca_V3.1-HA + VC, GFP-Ca_V3.1-HA + Myc-CACHD1 or CLR•RAMP1 seeded onto coverslips and used for experimentation after 48 h. Cells were washed twice with PBSCM, incubated in DMEM containing 0.1% BSA and mouse anti-HA (1:100) and rabbit anti-c-Myc (1:500) antibodies (1 h, 4°C), washed twice again with PBSCM and then fixed in 100 mM PBS containing 4% paraformaldehyde (w/v), pH 7.4 (20 min, 4°C). After washing with PBSCM the proximity ligation assay was conducted according to the manufacturer's instructions (Duolink[®] In Situ Red Starter Kit Mouse/Rabbit, Cat No. DUO92101, Sigma). Briefly, cells were blocked (1 h, 37°C), washed twice (5 min, room temperature) and then incubated with appropriate secondary antibodies (1 h, 37°C). After washing (2x 5 min, room temperature), the ligation was conducted (30 min, 37°C) and the cells were washed twice more. Coverslips were then

219	incubated with the amplification reaction mixture (100 min, 37°C), washed and coverslips
220	mounted in medium containing DAPI.
221	
222	Confocal microscopy
223	Cells were observed with a Nikon Eclipse Ti laser-scanning confocal microscope using a
224	100x/1.45 Oil DIC N2 objective. Images were collected at a zoom of 1-2 and at least five
225	optical sections were taken at intervals of $0.5~\mu m$. Single sections are shown. Images were
226	processed using Adobe Photoshop and the NIS-Elements AR software.
227	
228	Transformed human embryonic kidney cell culture and transfection for
229	electrophysiology
230	For electrophysiology experiments, HEK cells were transfected using 4 μ l Fugene6
231	(Promega, UK) with total 2 µg pcDNA3 at 50:1:25 for Ca _V 3.1/pmaxGFP, Ca _V 3.2/pmaxGFP
232	or $\text{Ca}_{\text{V}}3.3\text{/pmaxGFP}$ with or without $\alpha2\delta\text{-}1$ or CACHD1. Empty vector was used to
233	compensate when $\alpha2\delta$ or CACHD1 was omitted. Cells were maintained at 95% air, 5% CO ₂
234	at 37 °C and used for experimentation 24-48 h post transfection.
235	
236	Hippocampal neuron culture and transfection
237	Low-density hippocampal cultures were prepared from male and female E19 rat embryos as
238	described previously (Zhang et al., 2003). All experiments were carried out in compliance
239	with the Guide for the Care and Use of Laboratory Animals of the National Institutes of
240	Health and approved by the University of Virginia Animal Care and Use Committee and
241	adhered to ARRIVE guidelines. Neurons were plated onto poly-L-lysine coated glass
242	coverslips at a density of ~70 cells/mm ² and were transfected using lipofectamine 2000 at a

243 ratio of 2 µl lipofectamine 2000 per 1 µg DNA. Neurons were transfected with either 244 CACHD1 or pcDNA3.1 at a ratio of 10:1 excess to mVenus and moved 24 h after 245 transfection to a new glia-feeder layer. 246 247 Electrophysiology 248 Recordings from HEK cells were made as described previously (Vogl et al., 2015). Current-249 voltage (I-V) relationships from individual cells were fitted with a modified Boltzmann 250 equation: $I = G_{\text{max}} \times (V - V_{\text{rev}}) / (1 + \exp(-(V - V_{1/2})/k))$ where, G_{max} is the maximal 251 conductance (nS/pF), V_{1/2} is the midpoint of activation i.e. the voltage at which 50% of the 252 channels are open, V_{rev} is the null potential and k is the slope factor. Tail currents (measured 253 at -120 mV) were normalised to the maximal and minimal conductance and the resultant 254 curves were fitted with following Boltzmann function: $I=I_0+((I_{max}-I_0)/(1+\exp(V_{1/2}-V)/k)))$. 255 Throughout, all comparative electrophysiological experiments were performed in 256 transfection-matched cultures. 257 Recordings from hippocampal neurons were performed as described previously 258 (Jones et al., 2007). Throughout, data are expressed as mean S.E.M. Methods to estimate the 259 probability of channel opening, Po have been previously described by us (Shcheglovitov et 260 al., 2008), which assumes no change in single channel current, reducing the relationship 261 between whole-cell current (I) to $I \approx NPo$, where N is the number of channels in a cell and Po 262 is the probability of channel opening. N is estimated by measuring the channel gating current 263 at the reversal potential for ionic current. The peak current represents the maximal gating 264 charge Q_{max}, and is proportional to N. Peak ionic current conductance, G_{max}, was determined

by fitting the I-V curve, obtained from the same cell, with a Boltzmann-Ohm equation as

266	described earlier. G_{max} is used as a proxy for I since it is not affected by changes in driving
267	force. Therefore, the $G_{\text{max}}/Q_{\text{max}}$ ratio can be used to estimate Po.
268	
269	Experimental Design and Statistical Analysis
270	Throughout, all animal studies comply to appropriate ARRIVE and NIH guidelines and
271	comply to country and institute guidelines (as specified in Methods section for each animal
272	study). Details of animal strain, sex and method of sacrifice and use of anaesthetics are also
273	stated in Methods section for each animal study.
274	
275	Throughout, all comparative biochemical and electrophysiological experiments were
276	performed against transfection-matched culture controls. For electrophysiological
277	experiments in recombinant cells, a minimum of 5 separate transfections were performed and
278	numbers of individual replications are specified in appropriate Table. In all cases, sample size
279	is stated in text, Figure legend or appropriate Table. Data subjected to statistical comparisons
280	were assessed for assumptions of normality using a D'Agostino-Pearson omnibus test and
281	expressed as mean \pm standard error of the mean (SEM) throughout. Groups were compared
282	by two-tailed paired or unpaired Student's t-test, Mann-Whitney test, one- or two-way
283	ANOVA tests followed by Bonferroni post-hoc tests, Kruskal-Wallis test and Dunn's
284	multiple comparison test or least squares fits compared using extra sum of squares F test as
285	appropriate, using GraphPad Prism. In all cases, the statistical test used is stated in text,
286	Figure legend or appropriate Table. Throughout, P<0.05 was taken as statistically significant
287	and where appropriate values of P<0.01 and P<0.001 are specified.
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291 Results

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The novel CACHD1 protein is an α2δ paralog

We first investigated the predicted protein domain structure of CACHD1. Figure 1A illustrates that, like α2δ-1, CACHD1 has a predicted exofacial N-terminus according to its signal sequence, a von Willebrand factor A (VWA) domain, two bacterial chemosensory-like cache domains and a short hydrophobic transmembrane domain followed by an intracellular C-terminus. Although CACHD1 and α2δ share limited amino acid sequence homology (<21%), the similarities in modular domain content and arrangement between the proteins suggested the possibility that CACHD1 represents an $\alpha 2\delta$ -like protein. However, there are also a number of differences between CACHD1 and $\alpha 2\delta$ -1; these include: (i) $\alpha 2\delta$ proteins are a single gene product which is post-translationally cleaved by proteases into α2 and δ components and then associate via disulphide bonding (Calderon-Rivera et al., 2012; Segura et al., 2017); an important 6 amino acid motif for proteolytic cleavage has been identified (Andrade et al., 2007) which is absent in CACHD1. (ii) CACHD1 has a single predicted post-translational N-glycosylation site, whilst α2δ-1 is heavily glycosylated at multiple potential sites (Douglas et al., 2006). (iii) CACHD1 has a variant RSR amino acid sequence at the binding site for gabapentinoids. (iv) Despite expressing a VWA domain, the functionally important MIDAS motif in CACHD1 (DxGxS) is different from that of α2δ-1 (DxSxS). (v) $\alpha 2\delta s$ have a predicted GPI-anchoring site (Davies et al., 2010) which is absent in CACHD1, which instead has a predicted transmembrane domain and a larger intracellular C-terminus domain.

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CACHD1 is highly expressed in brain hippocampal and thalamic regions

To obtain comparative and quantitative data on CACHD1 mRNA expression, real-time PCR
was performed on rat and human mRNA from different regions of the brain and peripheral
tissue. Relative expression profiles of CACHD1 and $\alpha2\delta$ -1 transcripts in rat tissue showed
high CACHD1 expression in thalamus, hippocampus and cerebellum, whilst $\alpha2\delta$ -1 transcript
expression was prominent in cortex, hippocampus and also, superior cervical ganglia (Fig.
1 <i>B</i>). We further investigated the anatomical distribution of CACHD1 at the transcriptional
and protein levels using in situ hybridization and immunohistochemistry in adult mammalian
brain. Rat brain regions displaying high mRNA include the hippocampus, anterodorsal
thalamic nucleus, reticular thalamic nucleus, cerebellum, subiculum, medial entorhinal cortex
and zona incerta (Fig. 1-1; Fig. 1-2). Hippocampal CACHD1 mRNA staining was strong in
the dentate gyrus, as well as the CA1 pyramidal cell layer; mRNA staining was less strong in
CA3. There was strong correlation between the levels of expression of CACHD1 mRNA and
protein in rat brain (Fig. 1-2). In the thalamus, CACHD1 protein showed differential
expression between major thalamic nuclei, in particular with prominent staining in the
anterodorsal and reticular nuclei (Fig. 2). In human tissue, CACHD1 transcripts were
similarly high in hippocampus, thalamus, and cerebellum (Fig. 2-1). CACHD1 transcript
distribution was broadly similar to certain $\text{Ca}_{\text{V}}3$ subtypes, in particular to $\text{Ca}_{\text{V}}3.1$ (Fig. 2-1,
Talley et al., 1999). CACHD1 transcript expression showed a differential distribution to $\alpha 2\delta$ -
1 and $\alpha2\delta$ -2 subtypes and was most similar to $\alpha2\delta$ -3 (Fig. 2-1, Cole et al., 2005). In human
tissue, CACHD1 protein levels were most abundant in dentate gyrus granule cells and
pyramidal cells of the hippocampus cornus ammonis, cortical regions and thalamus, in both
large diameter and small diameter cells (Fig. 3).

CACHD1 promotes cell-surface expression of Cav3.1

Our expression data indicated high levels of CACHD1 expression in the thalamus,
hippocampus and cerebellum. As expression levels of $\text{Ca}_{\text{V}}3$ subunits are also high in the
thalamus and hippocampus, we hypothesized that CACHD1 may modulate $\text{Ca}_{\text{V}}3$ subunits in
a recombinant HEK cell system. As a first step, we expressed CACHD1 in HEK cells and
confirmed the specificity of the CACHD1 antibody (Fig. 4A). Immunoreactive CACHD1 was
detected at approximately 170 kDa. We also confirmed that CACHD1 is present at the cell-
surface of HEK cells (Fig. 4B). Next, we determined if expression of CACHD1 affected the
subcellular localization of Ca _v 3.1 using a cell-surface biotinylation assay. Cell-surface
proteins from HEK cells expressing empty vector, empty vector + CACHD1, GFP-Ca _V 3.1-
HA + empty vector and GFP-Ca _V 3.1-HA + CACHD1 were extracted and levels of GFP-
Ca _V 3.1-HA analysed by Western blotting. Our data show that co-expression of CACHD1
increased cell-surface localization of GFP-Ca _V 3.1-HA (2.65 \pm 0.40 fold over control P<0.05
two-tailed paired Student's <i>t</i> -test; Fig. 4 <i>C</i>). We also quantified the whole-cell expression of
GFP-Ca _V 3.1-HA in the same HEK cells, normalising to levels to β -actin (Fig. 4 <i>D</i>).
Importantly, our data shows that CACHD1 increases levels of GFP-Ca _V 3.1-HA at the cell-
surface without affecting the total cellular level.
CACHD1 and Ca _V 3.1 are in close proximity at the cell-surface
To determine if Ca _V 3.1 and CACHD1 are present in a complex at the cell-surface, an epitope-
tagged CACHD1 (Myc-CACHD1) was used to aid cell-surface precipitation and detection.
First, we tested the expression of the tagged protein and examined the ability of an anti-Myc
antibody to bind to CACHD1 at the cell-surface. Myc-CACHD1 was expressed in HEK cell
with a similar molecular mass (~170 kDa) to untagged CACHD1 (Fig. 5-1). Furthermore, we
could detect Myc-CACHD1 at the cell-surface using immunofluorescence and confocal

362	microscopy (Fig. 5-1). Proximity ligation assays are commonly used to predict the likelihood
363	that two proteins are sufficiently close enough to be present in the same complex. First, we
364	determined if we could simultaneously detect Myc-CACHD1 and GFP-Ca _V 3.1-HA at the
365	cell-surface by confocal microscopy. Live HEK cells expressing empty vector, empty vector
366	+ CACHD1, GFP-Ca $_{V}$ 3.1-HA + empty vector and GFP-Ca $_{V}$ 3.1-HA + CACHD1 were
367	incubated with antibodies to the Myc and HA epitope tags of CACHD1 and $\text{Ca}_{\text{V}}3.1$,
368	respectively and immunoreactive proteins visualized by immunofluorescence (Fig. 5A). No
369	immunoreactive signals were detected in cells expressing empty vector, indicating antibody
370	specificity. We were able to detect immunoreactive Myc signals only in cells expressing
371	Myc-CACHD1. Similarly, we were able to detect signals for the HA antibody only in cells
372	expressing GFP, indicating expression of GFP-Ca _V 3.1-HA. We were also able to
373	simultaneously detect CLR and RAMP1 at the cell-surface of transfected cells (Fig. 5A).
374	Next, we labelled cells from the same transfections and performed a proximity ligation assay
375	and visualized the cells using confocal microscopy. No PLA signals were detected in cells
376	transfected with empty vector, empty vector + CACHD1, GFP-Ca _V 3.1-HA + empty vector
377	(Fig. 5B). By contrast, we could readily detect PLA signals in our positive control
378	(CLR•RAMP1) and in transfected with GFP-Ca _V 3.1-HA + CACHD1. Importantly, we could
379	only detect PLA signals in cells expressing GFP (Fig. 5B). Thus, CACHD1 and Ca _V 3.1 are in
380	close proximity (<40 nm) at the cell-surface of HEK cells, indicating that they are likely in
381	the same protein complex. As discussed more fully below, together, these data are consistent
382	with CACHD1 increasing the cell-surface localization of $\text{Ca}_{\text{V}}3.1$ and with formation of
383	CACHD1-Ca _V 3.1 complexes at the cell surface.

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CACHD1 modulates recombinant Cav3 family VGCCs

386	We next tested the hypothesis that CACHD1 modulates T-type Ca ²⁺ current. Co-expression
387	of CACHD1 with $\text{Ca}_{\text{V}}3.1$ caused an increase in current density around peak values (Fig.
388	6A,B) and a corresponding increase in maximal conductance (Fig. 6B inset; Table 1). By
389	contrast, in our hands, $\text{Ca}_{\text{V}}3.1$ peak current and conductance was not modulated by $\alpha2\delta\text{-}1$ in
390	transfection-matched experiments (Fig. 6A,C; Table 1). CACHD1 effects were not
391	accompanied by any overall change in the midpoint of activation or slope factor k (Table 1)
392	and CACHD1 had no effect on Ca _V 3.1 steady-state inactivation (data not shown). Neither
393	CACHD1 nor $\alpha 2\delta$ -1 affected Ca _V 3.1 recovery from inactivation, as measured by lack of
394	effect on mid-time of recovery from inactivation or $\tau_{recovery}$ (p>0.1 for both, one-way
395	ANOVA with Bonferroni post-hoc test, data not shown).
396	We next investigated potential modulation of Ca_{V} 3.2 and Ca_{V} 3.3 by CACHD1. Peak
397	current density of Ca _V 3.2 (Fig. 7 <i>A</i> , <i>C</i>) and Ca _V 3.3 (Fig. 7 <i>B</i> , <i>D</i>) was increased by CACHD1
398	with corresponding increases in maximal conductance (Table 1). CACHD1 had no significant
399	effect on midpoint of activation or slope factor k for either $\text{Ca}_{\text{V}}3.2$ or $\text{Ca}_{\text{V}}3.3$ (Table 1) or
400	steady-state inactivation (p>0.1, Kruskal-Wallis test with Dunn's multiple comparison test,
401	data not shown). CACHD1 was without effect on Ca _V 3 activation or inactivation kinetics
402	(Fig. 7-1; Table 1). In our hands, $\alpha 2\delta$ -1 was without effect on current density in Cav3.2 (Fig.
403	7 <i>E</i>) or Ca _V 3.3 (Fig. 7 <i>F</i>). α 2 δ -1 was without effect on Ca _V 3.2 activation kinetics or on Ca _V 3.2
404	and $Ca_V 3.3$ inactivation kinetics (Fig. 7-1; Table 1). $\alpha 2\delta$ -1 had subtle effects on $Ca_V 3.1$
405	activation and inactivation kinetics and $\text{Ca}_{\text{V}}3.3$ activation kinetics (Fig. 7-1; Table 1).
406	Overall, these data suggest that CACHD1, but not $\alpha 2\delta$ -1, has a major effect on recombinant
407	Ca _V 3 VGCCs in terms of increased Ca ²⁺ current density and maximal conductance.
408	To determine the mechanism by which CACHD1 increased T-type channel currents,
409	we estimated channel opening probability by measuring Ca _V 3.1 gating currents at the reversal

potential for the ionic current (Fig. 8). In these experiments, the CACHD1-mediated increase in current density was recapitulated; thus, $Ca_V3.1$ maximal conductance 280 ± 30 pS/pF was significantly increased to 860 ± 15 pA/pF (n = 12 for each condition from 3 separate transfections; P<0.001 Mann-Whitney test) (data not shown). Measurement of area under the gating current provides a measure of the maximal gating charge Q_{max} . A plot of conductance versus gating current amplitude of the ionic current of the same cell provides a measure of open probability (Po) (Agler et al., 2005). Under these conditions, there was a ~1.4 fold increase in $Ca_V3.1$ Po in CACHD1 expressing cells (P<0.001, Fig. 8). These findings are consistent with CACHD1 interaction with $Ca_V3.1$ at the cell surface causing a functional increase in Po as a major contribution to CACHD1-mediated increases in Ca^{2+} current density.

CACHD1 increase Ca_V3-mediated excitability in hippocampal neurons

Ca_V3 channels are predicted to affect neuronal excitability around the resting membrane potential (Perez-Reyes, 2003; Cheong and Shin, 2013). To investigate the role of CACHD1 in controlling neuronal excitability, we expressed CACHD1 (vs. empty vector controls) in hippocampal neurons. Transfected neurons were identified by co-expression of the biomarker mVenus (Fig. 9*A*). At a depolarizing current injection step of 220 pA, CACHD1 expressing neurons fired at a higher frequency than control neurons (Fig. 9*B*, *C*,*D*; Table 2). To further determine the role of T-type currents in establishing the increase in neuronal firing frequencies, we used the selective Ca_V3 channel blocker, TTA-P2 (Dreyfus et al., 2010). TTA-P2 (1 μM) reversed the firing frequency in CACHD1 expressing neurons back to control levels, but was without effect on control neurons (Fig. 9*D*; Table 2). To increase the

contribution of T-type current to neuronal excitability, a hyperpolarizing prepulse was used to recover LVA Ca²⁺ channels from inactivation, followed by a short depolarizing pulse to evoke an AP (Eckle et al., 2014). Under these conditions, CACHD1 expression caused a more profound increase in rebound firing frequency in CACHD1-transfected, but not control, neurons (Fig. 9*E*,*F*,*G*; Table 2). TTA-P2 (1 µM) reversed the increase in rebound AP firing in CACHD1 expressing neurons back to control levels, but was without effect on control neurons (Fig. 9*G*; Table 2). Throughout these experiments, CACHD1 had no significant effects on AP waveform properties (Fig. 9-1). These data support a CACHD1-mediated selective increase in T-type Ca²⁺ current, which leads to an increase in AP firing frequency and excitability in native neurons.

444	Discussion
445	This study characterises the protein CACHD1, encoded by the cache domain containing 1
446	gene, and presents evidence that it represents a novel protein that modulates Ca _V 3 VGCC
447	activity. These data also provide further evidence that the major $\alpha 2\delta$ -1 auxiliary calcium
448	channel subunit does not fulfil a similar role for Ca _V 3 channels. Detailed examination of
449	Ca _V 3.1 channels suggests an underlying mechanism whereby CACHD1 promotes increased
450	$\text{Ca}_{\text{V}}3.1$ levels at the plasma membrane. In addition, data were consistent with CACHD1
451	forming a complex with the channel at the cell surface to increase open probability and
452	potentiate T-type current.
453	
454	CACHD1 protein modulates Ca _V 3 VGCCs
455	At a cellular level, CACHD1 transcripts were localised to granule and pyramidal cells of the
456	hippocampus, and specific thalamic nuclei, notably the anterodorsal thalamic nucleus and
457	reticular nucleus. Compared to the gene expression of the major $\alpha2\delta$ -1 and $\alpha2\delta$ -2 subunits,
458	CACHD1 protein displayed a unique expression signature with, in particular, high expression
459	in the thalamus and hippocampus and also in some regions of the cerebellum and cortex.
460	CACHD1 was largely co-incident with the expression pattern of the Ca _V 3.1 channel in the
461	CNS (Talley et al., 1999). CACHD1 co-transfection with Ca _V 3.1 in recombinant cells
462	increased cell surface expression and Ca ²⁺ current levels and maximal conductance.
463	CACHD1 similarly modulated $Ca_V3.2$ and $Ca_V3.3$ current levels. Under equivalent
464	conditions, $\alpha 2\delta$ -1 was without significant effect on current levels in any Ca_V3 subtype.
465	Proximity ligation assays were consistent with CACHD1 being able to form complexes with
466	$\text{Ca}_{\text{V}}3.1$ at the cell surface. Mechanistically, CACHD1 effects on $\text{Ca}_{\text{V}}3.1$ were associated with
467	increases in channel Po. A similar role has been reported for α2δ auxiliary subunit

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interactions with Ca_V1 channels; thus, $\alpha 2\delta$ -1 increased channel Po and channel number as well as allosterically regulating drug binding (Shistik et al., 1995; Wei et al., 1995). Other studies have reported either an α2δ-mediated reduction in Po (Wakamori et al., 1999) or a lack of effect on Po (Brodbeck et al., 2002). The latter study suggested that $\alpha 2\delta$ predominantly performs a VGCC trafficking function to increase the number of active channels at the membrane (reviewed by Dolphin, 2012). The demonstrated CACHD1mediated increase in Ca_V3.1 cell surface expression is proposed to contribute to increase in cell Ca²⁺ current levels and maximal conductance. Here, the ~1.4-fold increase in Po is insufficient to fully account for the ~3 fold increase in current density seen in this set of experiments; channel number is predicted to increase (according to I= iNPo, where I is the whole-cell current, i is the single channel current (predicted to be constant) and N is the number of functional channels). Thus, increase in channel number may be attributable to either CACHD1-mediated increases in forward trafficking or reduced endocytosis of Cay3.1. With respect to $\alpha 2\delta$ auxiliary subunits, HVA Ca_V $\alpha 1$ - $\alpha 2\delta$ interactions are reported to occur during early maturation at an intracellular site to drive forward trafficking to the plasma membrane (Cantí et al., 2005). Whilst Ca_V2.2 proteomic data have reported only a low appreciable amount of co-purified $\alpha 2\delta$, with detection dependent on solubilising agent used (Müller et al., 2010), recent work using exofacial tags and antigen stripping techniques has supported $\alpha 2\delta$ also remaining associated with Ca_V2.2 at the plasma membrane (Cassidy et al., 2014). In the present study, clear indication of CACHD1 and Cav3.1 complex formation at the cell surface was obtained using proximity ligand assays. Moreover, $\alpha 2\delta$ has the propensity to sequester into lipid raft compartments, as reported by us (Ronzitti et al., 2015) and others; this may also limit efficient detection of $\alpha 2\delta$ -Ca_V $\alpha 1$ complexes and it will be of interest to determine if CACHD1 similarly localizes to lipid rafts. Overall, we propose that

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CACHD1 acts to increase Ca_V3 expression at the plasma membrane, at the cell surface

CACHD1 can form a complex with the channel to increase Po and, consequentially, increase

T-type current.

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Potential functional impact of CACHD1 on Cav3 VGCCs

T-type Ca²⁺ currents are active around the resting membrane potential, where noninactivating channels generate low threshold Ca²⁺ spikes and the consequential triggering of Na⁺-dependant APs (Llinás 1988; Cheong and Shin, 2013). Of further interest here is that multiple mechanisms and proteins involved in folding and trafficking are reported to be involved in Ca_V3 expression at the cell surface. For example, proteins such the actin binding protein kelch-like 1 (Aromomolaran et al., 2010), stac1 (Rzhepetskyy et al., 2016) and calnexin (Proft et al., 2017) have a proposed role in Ca_V3 expression. Moreover, the glycosylated form of Ca_V3 represents the mature, correctly folded protein that is associated with higher Po (Weiss et al., 2013; Ondacova et al., 2016). T-type current has also been implicated in regulating presynaptic transmitter release in hippocampal and nociceptive circuitry (Huang et al., 2011; Jacus et al., 2012). Increases in Ca_V3 current are predicted to have profound effects on neuronal firing (McCormick and Huguenard, 1992). Correspondingly, over-expression of CACHD1 caused a pronounced increase in T-type current-mediated spike firing in hippocampal neurons. This activity was enhanced using a protocol to trigger recovery of Cay3 channels from their inactivated states, thereby increasing contribution of T-type current to neuronal excitability. Ca_V3 subtypes have been suggested as targets for anti-epileptic drugs (Powell et al., 2014). In models of temporal lobe epilepsy (TLE), selective up-regulation of T-type current in hippocampal neurons causes intrinsic bursting activity (Sanabria et al., 2001; Su et al., 2002). Ca_V3.2 transcripts were upregulated

516	in TLE models and intrinsic burst firing was reduced in Ca _V 3.2 knock-out mice (Becker et
517	al., 2008). Moreover, the deubiquitinating enzyme USP5 (Garcia-Callero et al., 2014), and
518	preventing $\text{Ca}_{\text{V}}3.2$ deubiquitination was suggested to be beneficial in neuropathic and
519	inflammatory pain. Our data suggest CACHD1 as a potential future target in
520	hyperexcitability disorders associated with $\text{Ca}_{\text{V}}3$ dysfunction, such as epilepsy and pain.
521	Moreover, CACHD1 gene expression has been shown to be modulated in patients with Type
522	1 diabetes (Rassi et al., 2008) and Parkinson's disease (Aguiar and Severino, 2010).
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524	CACHD1 protein structure dictates α2δ-like function
525	There are clear similarities in protein structural motifs between CACHD1 and $\alpha2\delta$, namely,
526	the presence of an N-terminal signal sequence, VWA and two downstream cache domains,
527	these similarities suggest a conserved evolution (Anantharaman and Aravind, 2000).
528	However, a number of important differences are also present. CACHD1 has a RSR variant at
529	the gabapentin binding motif; whilst $\alpha 2\delta$ -1 and $\alpha 2\delta$ -2 were found to bind to gabapentinoids
530	via their RRR binding motif, $\alpha 2\delta$ -3 and $\alpha 2\delta$ -4 have variant RNR sites which do not bind
531	gabapentin (Wang et al., 1999; Marais et al., 2001). Earlier studies also identified porcine
532	$\alpha 2\delta$ -1 residues 516 to 537 within the first cache domain and residues 583 to 603 as also
533	contributing to gabapentin binding (Wang et al., 1999). It will be of interest to determine if
534	CACHD1 binds gabapentanoids. Despite sharing a common VWA domain, CACHD1 has a
535	variant MIDAS motif. The $\alpha 2\delta$ -1 MIDAS motif is functionally important in Ca^{2^+} channel
536	trafficking and synaptic function (Cantí et al., 2005; Hoppa et al., 2012). However, it has
537	been suggested that MIDAS is unlikely to represent a key $\text{Ca}_{V}2.2/\alpha 2\delta$ -1 interaction site,
538	rather other regions are more likely involved (Cassidy et al., 2014); such regions may include
539	cache domains, for example, rat $\alpha 2\delta$ -1 residues 751-755, which are within a modelled cache

region, were implicated in $Ca_V 2.2/\alpha 2\delta$ -1 interaction (Cassidy et al., 2014). By contrast, comparative data investigating $\alpha 2\delta$ effects on $Ca_V 1.2$ point to aspartate and the first serine residue within the DxSxS MIDAS site as molecular determinants for interaction and correct modulation of $Ca_V 1.2$ (Briot et al., 2018). Of interest here is that CACHD1 contains a variant MIDAS with a glycine residue at the equivalent position of the critical serine residue identified by Briot et al. (2018). It has also been proposed that the $\alpha 2\delta$ amino terminal (amino acids 26-230, termed the R-domain) contains all the machinery required to support $\alpha 2\delta$ -1-mediated current enhancement in $Ca_V 2.2$ channels (Song et al., 2015). This study identified a tryptophan residue (W205), which is conserved across all four $\alpha 2\delta$ isoforms, as an important molecular determinant for these R-domain effects; it is of note that CACHD1 also contains a conserved tryptophan residue at the equivalent position.

In bacteria, the cache domain is proposed to arise from bacterial small molecule binding domains PAS and GAF (Anantharaman et al., 2001) and to play a key role in chemotaxis by acting as an extracellular receptor (Anantharaman and Aravind, 2000). Recent computational work has suggested that cache domains represent the dominant extracellular sensor in prokaryotes; by contrast, cache domains are largely limited to only $\alpha 2\delta$ subunits in metazoa (Upadhyay et al., 2016). The present study adds CACHD1 to this classification. Whilst the functional relevance of mammalian cache domains remains to be fully established, deletions within the cache domain of $\alpha 2\delta$ -4 have been associated with familial bipolar disorder (Van Den Bossche et al., 2010). Roles for 'free' $\alpha 2\delta$ (not associated with VGCCs) have also been extended to functions including synaptogenesis and neurodegeneration via interaction with alternative ligands such as thrombospondins and prion proteins, respectively (Eroglu et al., 2009, Lana *et al.*, 2016; Senatore et al., 2012); it will also be of interest to see if CACHD1 possesses similar functionality.

Overall, our data are consistent with CACHD1 structurally representing an $\alpha 2\delta$ -like
protein that act to increase Ca _V 3 cell surface expression and current. Identification of the
CACHD1 protein as a modulator of Ca _v 3 activity expands the range of VGCC associated
proteins and may provide an additional target itself, or via its modulation of T-type current, in
different disease states.

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824	

825	Figure Legends
826	Figure 1. Predicted protein sequence homology and relative expression profile of
827	CACHD1 and α2δ-1.
828	CACHD1 and $\alpha 2\delta$ -1 subunits both contain a N-terminus signal peptide, a VWA domain, two
829	cache domains, and transmembrane and intracellular domains. GBP: gabapentin binding
830	domain (RRR). GBP*: gabapentin binding domain variant (RSR). MIDAS: metal-ion-
831	dependent adhesion site (DxSxS). MIDAS*: metal-ion-dependent adhesion site variant
832	(DxGxS). VWA: von Willebrand factor A. Cache: Ca ²⁺ channel and chemotaxis receptor.
833	TM: transmembrane domain. Cys: cysteine. His: histidine (locations of domains are
834	approximate and from data from www.Uniprot.org, figure drawn using DOG: Domain
835	Graphics). (B) Relative expression profile of CACHD1 and $\alpha 2\delta$ -1 mRNA in rat tissue
836	determined using SYBR green real-time quantitative PCR and HPRT1 as housekeeping gene.
837	DRG: dorsal root ganglion. SCG: superior cervical ganglion. (Data normalised to lowest
838	tissue expression; n=3 experiments using 3 animals each). Figure 1 is supported by <i>in situ</i>
839	hydridization data in different rat brain regions (Fig. 1-1) and qualitative expression profile of
840	CACHD1 mRNA and protein in the adult rat brain (Fig. 1-2).
841	
842	Figure 2. CACHD1 protein expression in adult rat brain.
843	Immunoreactive protein was detected using rabbit anti-CACHD1 with peroxidase anti-rabbit
844	secondary antibody and DAB staining (brown). AD: anterodorsal thalamic nucleus; AVDM:
845	anteroventral thalamic nucleus (dorsomedial); AVVL: anteroventral thalamic nucleus
846	(ventro-lateral); fi: fimbria; MD: mediodorsal thalamic nucleus; Po: posterior thalamic
847	nucleus; sm: strai medullaris; Rt: reticular thalamus nucleus; RtSt: reticular VL: ventrolateral
848	thalamic nucleus: VPI: ventro-nosterior lateral thalamus; or oranule cell layer; m: molecular

849	layer; p: Purkinje cell; wm: white matter. Figure 2 is supported by expression profiling of
850	CACHD1 and different voltage-gated calcium channel subunit mRNA in human tissue (Fig.
851	2-1).
852	
853	Figure 3. CACHD1 protein expression in human brain.
854	Immunohistochemistry of adult human brain using rabbit anti-CACHD1 with peroxidase
855	anti-rabbit secondary antibody with (brown) DAB stain. CA1-3: cornus ammonis 1-3; DG:
856	dentate gyrus.
857	
858	Figure 4. Characterisation of CACHD1 and its effects on Ca _V 3.1 channel expression.
859	HEK cells were transfected with empty vector (vector control, VC), CACHD1, Myc-
860	CACHD1, GFP-Ca _V 3.1-HA alone or in combination, as shown in each panel. (A) HEK cell
861	lysates were analysed by Western blotting (WB). An antibody to CACHD1 recognised a
862	single protein similar to the predicted size for CACHD1, but also recognized a non-specific
863	protein in all lysates. (B) Cell-surface proteins were biotinylated and pull downs analysed for
864	CACHD1 and Na ⁺ /K ⁺ -ATPase (loading control). In control cells, no immunoreactive
865	CACHD1 was detected, confirming antibody specificity. In CACHD1 expressing cells,
866	immunoreactive CACHD1 was detected. In both cell types, immunoreactive $\text{Na}^+\!/\text{K}^+\text{-ATPase}$
867	was detected. (C) Cell-surface proteins were biotinylated and pull downs analysed for GFP-
868	Ca_{V} 3.1-HA (HA) and $\text{Na}^{\text{+}}/\text{K}^{\text{+}}$ -ATPase (loading control). In control cells and cells only
869	expressing CACHD1, no HA signals were detected, confirming antibody specificity. In cells
870	expressing GFP-Ca _V 3.1-HA, HA signals were readily detected. Quantification of the HA
871	signals (normalised to Na ⁺ /K ⁺ -ATPase) revealed expression of CACHD1 increased signals
872	for GFP-Cav3 1-HA at the cell-surface *n<0.05 Na ⁺ /K ⁺ -ATPase signals were detected in all

cell types (D) Inputs of the biotin pull down assays were analysed by WB. Signals for HA
were only detected in cells expressing GFP-Ca _V 3.1-HA, signals for CACHD1 were only
detected in cells expressing Myc-CACHD1 and signals for $\beta\text{-actin}$ were detected in all cell
types. All blots are representative of n≥3 experiments.
Figure 5. Ca_{V} 3.1 and CACHD1 are present at the cell-surface and are in close
proximity. Live HEK cells expressing empty vector (vector control, VC), VC + Myc-
CACHD1, GFP-Ca _V 3.1-HA + VC, Myc-CACHD1 + GFP-Ca _V 3.1-HA or CLR•RAMP1
(positive control) were incubated with antibodies to HA and Myc, washed and fixed. (A)
Cells were then incubated with appropriate secondary antibodies and immunoreactive
proteins localised by immunofluorescence and confocal microscopy. In HEK-VC cells, no
signals for GFP, HA or Myc were detected indicating specificity of detection. HA signals
(arrowheads) were only detected in cells expressing GFP-Ca $_{V}$ 3.1-HA (as determined by the
GFP signal) and CLR•RAMP1. Similarly, Myc signals (yellow arrowheads) were only
detected in cells expressing Myc-CACHD1 and CLR•RAMP1. Scale bar, 10 μm (B) After
the proximity ligation assay, no signals were detected in cells expressing empty vector or in
cells expressing only Myc-CACHD1 or GFP-Ca $_{V}$ 3.1-HA. In contrast, PLA signals were
detected in cells expressing Myc-CACHD1 + GFP-Ca _V 3.1-HA (arrows) and CLR•RAMP1
(arrows). Single optical sections are shown except for the PLA panel (CLR•RAMP1
excluded) where 5 optical sections are merged, two above and two below (0.5 μm
increments) from the optical sections shown in the GFP/DAPI panel. Scale bar, 20 $\mu m.\ All$
images are representative of n=3 experiments. Figure 5 is supported by analysis of cell-
surface CACHD1 construct expression studies (Fig. 5-1).

897	Figure 6. Effects of CACHD1 and \$\alpha 20-1 on Cay5.1 channels
898	CACHD1 significantly increased current density as shown by (A) representative current
899	density traces at -25 mV and (B) I-V relationships, V_H -90 mV (*p<0.05, **p<0.01,
900	*** p <0.001, two-way ANOVA with Bonferroni post-hoc test). α 2 δ -1 had no significant
901	effect on current density as shown by (A) representative current density traces at -25 mV and
902	(C) I-V relationships, $V_{\rm H}$ -90 mV. CACHD1, but not $\alpha 2\delta$ -1, significantly increased maximal
903	conductance (inset, p<0.05, one-way ANOVA with Bonferroni post-hoc test).
904	
905	Figure 7. Effects of CACHD1 and $\alpha2\delta1$ on Cav3.2 and Cav3.3 channels
906	CACHD1 significantly increased current density as shown by representative current density
907	traces at -20 mV for (A) $Ca_V3.2$ and (C) $Ca_V3.3$, and I-V relationships for (B) $Ca_V3.2$ and (D)
908	$Ca_{V}3.3;\ V_{H}\ -90\ mV\ (*p<0.05,\ **p<0.01,\ ***p<0.001,\ two-way\ ANOVA\ with\ Bonferroni$
909	post-hoc test). $\alpha2\delta$ -1 had no effect on (E) Ca _V 3.2 and (F) Ca _V 3.3 I-V relationships, V _H -90
910	mV. Figure 7 is supported by analysis of effects of CACHD1 and $$ α 2 δ -1 on Ca _V 3 channel
911	kinetic properties (Fig. 7-1).
912	
913	Figure 8. CACHD1 expression increases Ca_{V} 3.1 gating currents and open probability
914	(Po).
915	Representative gating currents recorded from $\text{Ca}_{V}3.1$ (Aa) and $\text{Ca}_{V}3.1 + \text{CACHD1}$ (Ab) at
916	the observed reversal potential. Expanded time scale illustrates the increase in area under the
917	gating current for CACHD1 expressed cells. B) Conductance vs gating current plot for
918	multiple cells. Line represents linear regression to data points. The slopes $(G_{\text{max}}/Q_{\text{max}})$ were
919	significantly different (P=0.0004, least squares fits compared using extra sum of squares F

942

920	test; $Ca_V 3.1$: 0.09 ± 0.003 , $n=10$, and $Ca_V 3.1 + CACHD1$: 0.14 ± 0.090 , $n=11$). C) Plot
921	showing the slopes (i.e relative Po) and S.E.M. for fits shown in B (***p<0.001).
922	
923	Figure 9. Effects of CACHD1 in hippocampal neurons
924	(A) Co-labelling of hippocampal neurons with CACHD1 and mVenus. (B) CACHD1
925	increased firing frequency of hippocampal neurons. (C) Example traces in response to
926	depolarizing current injections steps of -20, 70 and 140 pA. (D) Summary data from separate
927	experiments confirming CACHD1-mediated increased firing frequency and also showing that
928	TTA-P2 (1 μM) reduced firing rates in CACHD1-expressing neurons, but not in controls. (E)
929	Rebound APs were evoked using a -50 pA hyperpolarizing prepulse followed by a
930	depolarizing step from 0 pA to 200 pA in steps of 10 pA for 200 ms, CACHD1 expressing
931	neurons displayed a significantly greater number of rebound APs compared to controls. (F)
932	Example traces representing depolarizing current injection steps of 40, 90 and 140 pA. (G)
933	Summary data from separate experiments confirming CACHD1-mediated increased in
934	rebound APs and also showing that TTA-P2 (1 $\mu M)$ reduced firing rates in CACHD1-
935	expressing neurons, but not in controls. *P<0.05 throughout, two-tailed paired Student's t-test
936	or one-way ANOVA with Bonferroni post-hoc test. Figure 9 is supported by analysis of
937	effects of CACHD1 and TTA-P2 on biophysical properties of hippocampal neurons (Fig. 9-
938	1).
939 940	Extended Data Figure Legends
941	

Figure 1-1: CACHD1 mRNA expression in adult rat brain.

943	In situ hybridization of adult rat brain. CACHD1 mRNA was labelled pink with blue
944	counterstain (Gill's I Haematoxylin). CA1-3: cornus ammonis 1-3; DG: dentate gyrus; g:
945	granule cell layer; m: molecular layer; p: Purkinje cell; wm: white matter.
946	
947	Figure 1-2: Qualitative expression profile of CACHD1 mRNA and protein in the adult
948	rat brain.
949	+ labelling similar to background; ++ weak labelling; +++ moderate labelling, ++++ strong
950	labelling; +++++ very strong labelling.
951	
952	Figure 2-1: Expression profile of CACHD1 and voltage-gated calcium channel subunit
953	mRNA in human tissue.
954	Absolute quantification of CACHD1, $\alpha 2\delta$ -1, -2, -3, $Ca_V 2.2$ and $Ca_V 1$, -2, -3 transcripts was
955	assessed in triplicate by TaqMan® qPCR using 'Best Coverage' Taqman probes (Applied
956	Biosystems, UK) against a 5-point standard curve of plasmids consisting of 10-fold dilution
957	of a known copy number of plasmid containing cDNA of the gene of interest. Total RNA was
958	extracted using an RNeasy kit (Qiagen, UK) with an on-column DNase I treatment.
959	Additional total RNA samples from AMS Biotechnology (Abingdon, UK) originated from
960	human male donors aged 24-65.
961	
962	Figure 5-1: Analysis of cell-surface CACHD1 construct expression.
963	(A, B) HEK cells were transfected with empty vector (vector control, VC) or Myc-CACHD1
964	and cell lysates analysed by (A) Western blotting (WB) and (B) immunofluorescence and
965	confocal microscopy. (A) Immunoreactive signals for Myc (mouse Myc, mMyc) were
966	detected at a similar molecular mass to that predicted for CACHD1 only in cells expressing

967	CACHD1. (B, upper panel) Cells were incubated with antibody to Myc (rabbit Myc, rMyc),
968	washed, fixed and then incubated with appropriate secondary antibodies. Myc signals
969	(arrowheads) were only detected in cells expressing Myc-CACHD1. (B, lower panel) Cells
970	were fixed, incubated with antibody to Myc (rMyc), washed and then incubated with
971	appropriate secondary antibodies. Myc signals were detected at the cell-surface (arrowheads)
972	and in intracellular vesicles only in cells expressing Myc-CACHD1. Scale bar, 10 $\mu \text{m}.$
973	
974	Figure 7-1: Effects of CACHD1 and α2δ-1 on Ca _V 3 channel kinetic properties
975	CACHD1 co-expression had no significant effect on $t_{activation}$ in (Aa) $Ca_V3.1$, (Ba) $Ca_V3.2$ and
976	(Ca) $Ca_V 3.3$. $\alpha 2\delta$ -1 significantly increased $Ca_V 3.1$ $t_{activation}$ at all voltages tested (Aa)
977	(*p<0.05, **p<0.01, ***p<0.001, two-way ANOVA with Bonferroni post-hoc test); α2δ-1
978	had no effect on $Ca_V 3.2\ t_{activation}$ (Ba); $\alpha 2\delta$ -1 significantly decreased $Ca_V 3.3\ t_{activation}$ at -35
979	and -30 mV (Ca) (*p<0.05, ***p<0.001, two-way ANOVA with Bonferroni post-hoc test).
980	CACHD1 co-expression had no significant effect on $t_{inactivation}$ in (Ab) $Ca_V3.1$, (Bb) $Ca_V3.2$
981	and (Cb) Cav3.3. α 2 δ -1 co-expression with Cav3.1 (Ab) resulted in significantly faster
982	inactivation kinetics (*p<0.05, one-way ANOVA with Bonferroni post-hoc test), but had no
983	effect on $t_{inactivation}$ in (Bb) $Ca_V 3.2$ and (Cb) $Ca_V 3.3$. Inactivation traces at -20 mV or -30 mV
984	were fitted with a single exponential function.
985	
986	Figure 9-1: Effects of CACHD1 and TTA-P2 on biophysical properties of hippocampal
987	neurons.
988	Extended Data Fig. 9-1 supports Figure 9.
989	
990	

991 Table 1. Effects of CACHD1 and α2δ-1 on biophysical properties of Ca_V3 subtypes.

Table 1. Effects of CACHD1 and 0.20-1 on biophysical properties of Ca _V 5 subtypes.						
	G _{max} (pS/pF)	V _{1/2} (mV)	k (mV)	τ activation (ms)*	τ inactivation (ms)**	
Ca _V 3.1 (18)	628 ± 70	-34.5 ± 0.8 (30)	5.4 ± 0.1 (30)	2.0 ± 0.1	25.8 ± 2.0	
Ca _V 3.1/ CACHD1 (19)	944 ± 90*	-36.3 ± 0.9 (29)	5.6 ± 0.2 (29)	2.0 ± 0.2	22.2 ± 4.8	
Ca _V 3.1/ α2δ-1 (13)	672 ± 90	-35.7 ± 1.4	5.6 ± 0.3	$3.3 \pm 0.2^{\Delta\Delta}$	$18.9 \pm 0.86^{\Delta}$	
*=p<0.05 vs. Ca _V 3.1 (one-way ANOVA with Bonferroni post-hoc test) ^Δ =p<0.05, ^{ΔΔ} =p<0.05 vs. Ca _V 3.1 (two-way ANOVA with Bonferroni post-hoc test)						
Ca _v 3.2 (13)	596 ± 120	-34.4 ± 2.4	5.7 ± 0.2	7.1 ± 0.40	33.3 ± 0.97	
Ca _v 3.2/ CACHD1 (15)	1060 ± 140*	-33.4 ± 0.8	5.9 ± 0.2	5.9 ± 0.38	32.0 ± 1.6	
*=p<0.05 vs. Ca _V .	3.2 (two-tailed u	inpaired Student	t's t-test)			
Ca _V 3.3 (12)	573 ± 88	-36.1 ± 1.2	4.3 ± 0.2	24.4 ± 1.9	134 ± 12	
Ca _V 3.3/ CACHD1 (10)	849 ± 78*	-38.9 ± 1.6	4.0 ± 0.3	28.5 ± 3.4	126 ± 8.3	
*=p<0.05 vs. Ca _V 3.3 (two-tailed unpaired Student's <i>t</i> -test)						

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996

997 998 In all cases, comparisons were performed in culture-matched experiments. Numbers in parenthesis represents number of cells each from a minimum of 5 separate transfections.

^{*} τ activation was measured at -25 mV in all cases.

^{**} τ inactivation was measured at -20 mV for Ca_V3.1 and Ca_V3.2 and at -30 mV for Ca_V3.3.

999 Table 2. Effects of CACHD1 and TTA-P2 on hippocampal neuronal firing

	Firing frequency (Hz)	Rebound firing frequency (Hz)
Control	$6.0 \pm 1.2 \ (41/6)$	$7.2 \pm 1.2 (32/5)$
CACHD1	9.8 ± 1.1* (29/5)	12.1 ± 0.9* (28/5)
Control	$8.5 \pm 1.4 (6/3)$	$10.0 \pm 1.8 \ (6/3)$
Control + TTA-P2	6.5 ± 1.2 (6/3)	9.2 ± 1.5 (6/3)
CACHD1	14.1 ± 1.7 (7/3)	$16.7 \pm 0.8 \ (10/3)$
CACHD1 + TTA-P2	6.9 ± 1.4* (7/3)	10.0 ± 1.2* (10/3)

^{*=} p<0.05 vs control two-tailed paired Student's t-test

Values represent means \pm S.E.M; number in parenthesis = number of neurons/number of separate transfections.

 $\begin{array}{c} 1001 \\ 1002 \end{array}$

1000

Figure 1. Cottrell et al

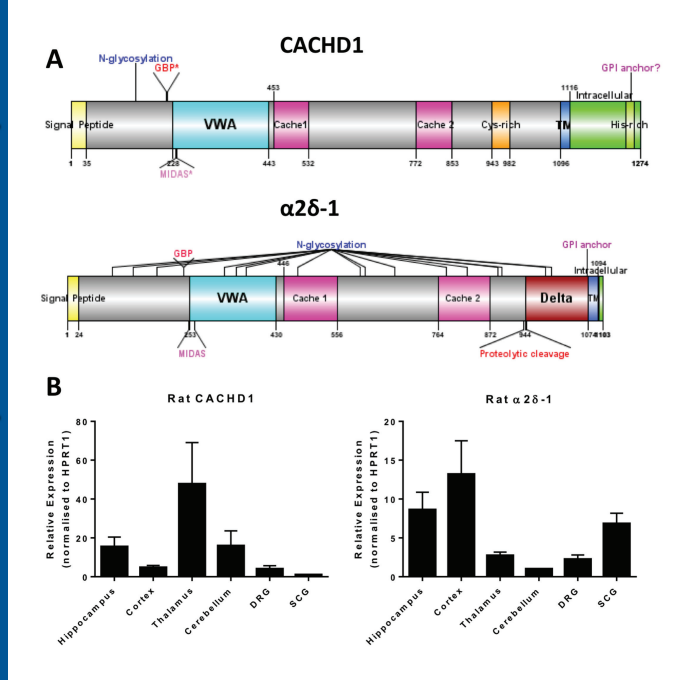


Figure 2. Cottrell et al

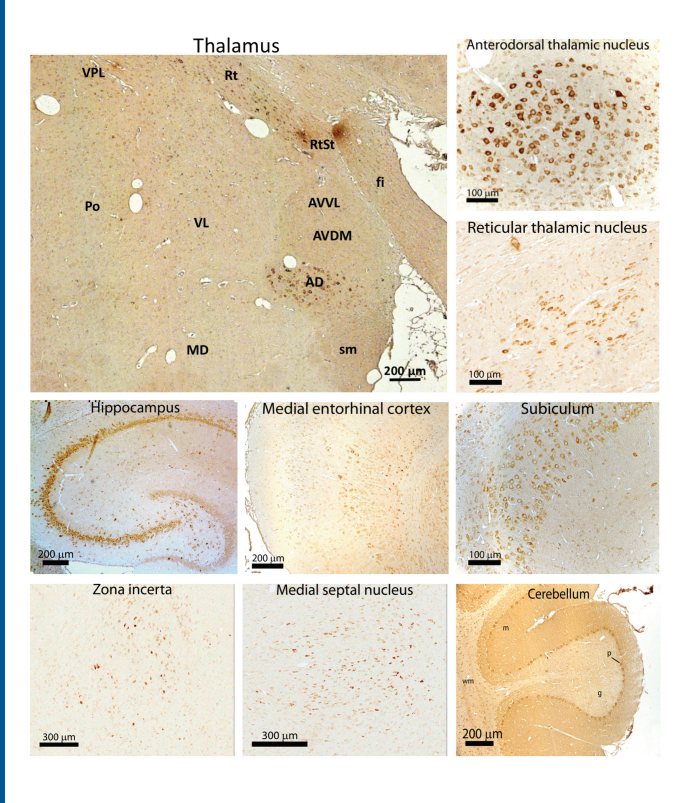
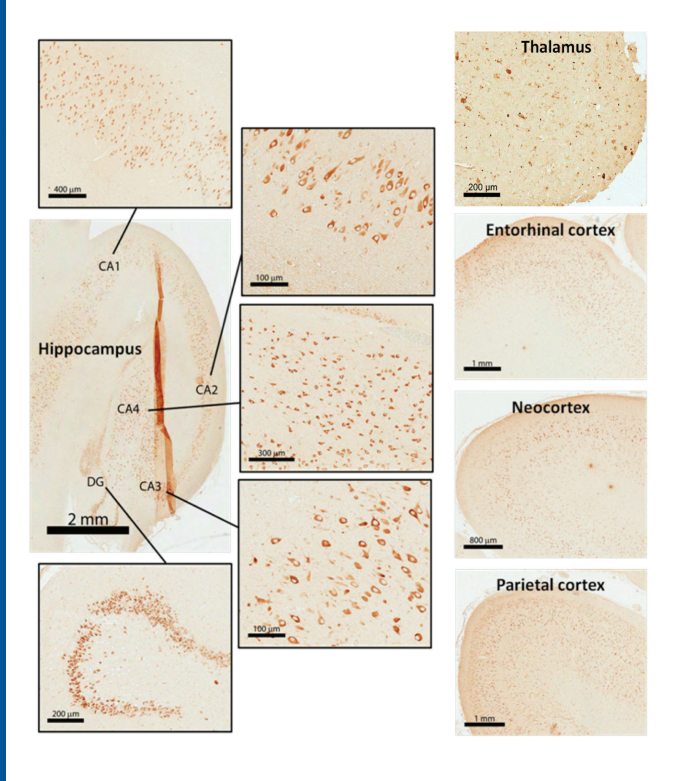
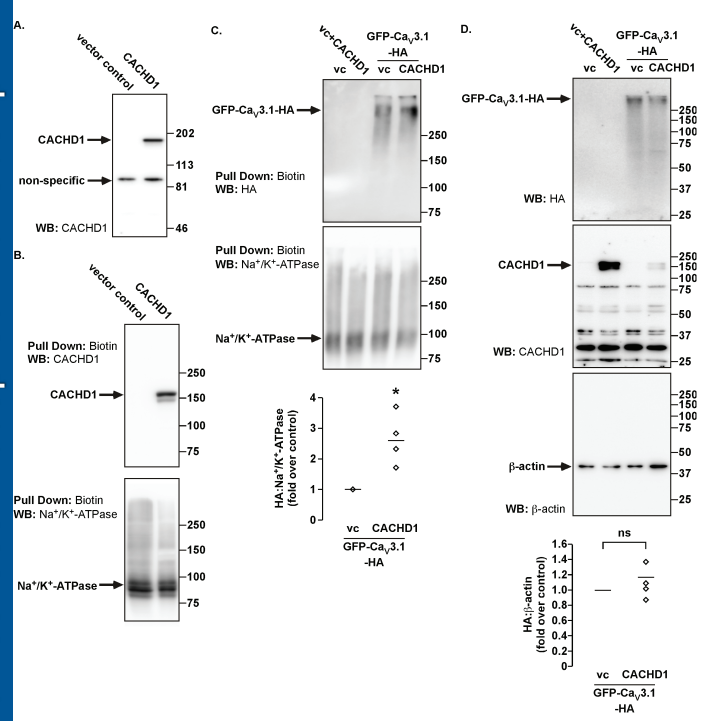


Figure 3. Cottrell et al





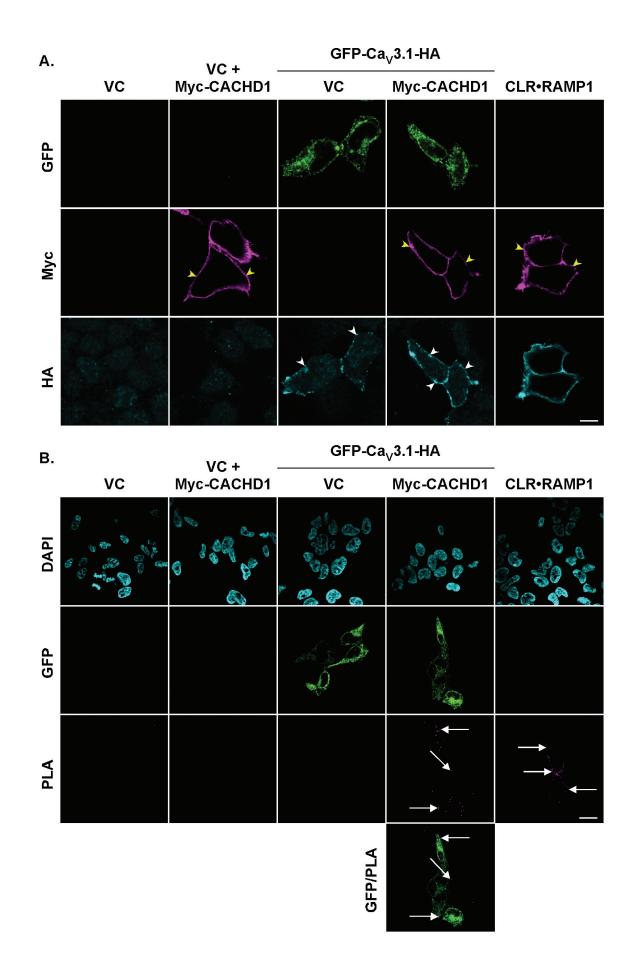


Figure 6. Cottrell et al

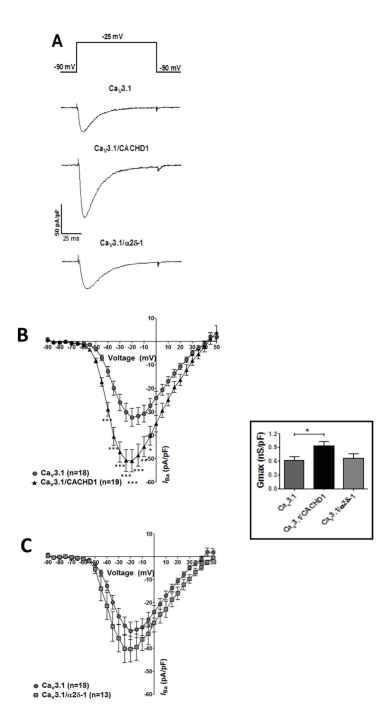


Figure 7. Cottrell et al

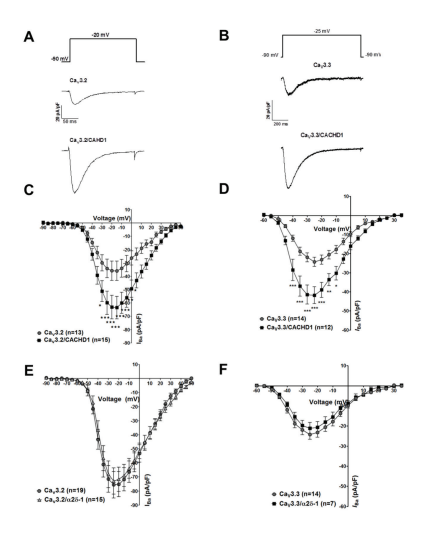


Figure 8. Cottrell et al

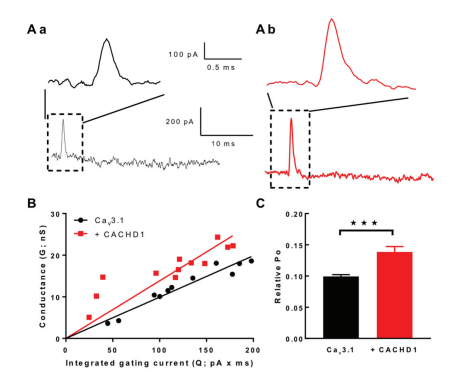


Figure 9. Cottrell et al

