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Regulation of aldosterone secretion by Ca,1.3

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Abstract

Aldosterone-producing adenomas (APAs) vary in phenotype and genotype. Zona glomerulosa (ZG)-like APAs frequently have mutations of an L-type calcium channel (LTCC) Ca\textsubscript{V}1.3. Using a novel antagonist of Ca\textsubscript{V}1.3, compound 8, we investigated the role of Ca\textsubscript{V}1.3 on steroidogenesis in the human adrenocortical cell line, H295R, and in primary human adrenal cells. This investigational drug was compared with the common antihypertensive drug nifedipine, which has 4.5-fold selectivity for the vascular LTCC, Ca\textsubscript{V}1.2, over Ca\textsubscript{V}1.3. In H295R cells transfected with wild-type or mutant Ca\textsubscript{V}1.3 channels, the latter produced more aldosterone than wild-type, which was ameliorated by 100 µM of compound 8. In primary adrenal and non-transfected H295R cells, compound 8 decreased aldosterone production similar to high concentration of nifedipine (100 µM). Selective Ca\textsubscript{V}1.3 blockade may offer a novel way of treating primary hyperaldosteronism, which avoids the vascular side effects of Ca\textsubscript{V}1.2-blockade, and provides targeted treatment for ZG-like APAs with mutations of Ca\textsubscript{V}1.3.
Aldosterone-producing adenomas (APAs), which arise from the adrenal cortex, are one of the most common curable causes of hypertension\(^1\)-\(^3\). They account for approximately half of primary aldosteronism, which is estimated to be present in 5-13 % of all hypertensive patients, and in at least 20 % of patients with resistant hypertension\(^4\). However, it is likely that fewer than 10 % of APAs are ever diagnosed; and fewer still are removed in time to cure hypertension and prevent resistance to effective drug treatment\(^2\),\(^5\).

We previously reported somatic gain-of-function mutations in two genes that regulate Na\(^+\), K\(^+\) and Ca\(^{2+}\) transport in APAs with a zona glomerulosa (ZG)-like phenotype\(^6\). Whole exome sequencing of small-cell APAs with a ZG-like gene expression profile found five out of ten to harbour one of four different somatic mutations in the voltage dependent L-type Ca\(^{2+}\) channel, Ca\(_{\text{v}}\).1.3 (encoded by the gene \textit{CACNA1D}). These four substitution mutations, V259D, G403R, I750M, and P1336R, cluster around the Ca\(^{2+}\) pore between the S5 and S6 domains that line the inner pore surface. The mutations occur in conserved sites within functional domains such as the voltage-sensing domain to the pore (V259D and P1336R) and the channel activation gate (G403R and I750M)\(^6\). The G403R and I750M mutations were simultaneously reported as rare de novo germline mutations presenting at birth, together with several patients having somatic mutations of the same residues in sporadic APAs\(^7\). Our own replication sequencing revealed three further mutations, and sequencing of APAs in a large European consortium has now identified a total of 19 somatic mutations in or near one of the four Ca\(^{2+}\) channel pore-forming domains\(^6\),\(^8\). Patch clamping of HEK293 cells has shown that at least 6 of the 19 mutations affect the Ca\(_{\text{v}}\).1.3 channel function and allow for increased Ca\(^{2+}\) influx through either shifting voltage-dependent activation towards more negative voltages, decelerating inactivation, and/or increasing currents through a higher open channel probability\(^6\),\(^9\).
The current medical treatment of primary hyperaldosteronism is blockade of the mineralocorticoid receptor, which can lead to an increase in aldosterone secretion. Therefore, blockade of calcium entry through selective antagonism of CaV1.3 might present a valuable therapeutic target. We therefore aimed to investigate whether CaV1.3 mutations cause the postulated increase in aldosterone secretion from human adrenocortical cells, and whether blockade of calcium entry reverses this. We studied the potential value of this target using 1-(3-chlorophenethyl)-3-cyclopentylpyrimidine-2,4,6-(1H,3H,5H)-trione (compound 8), which was found to be more than 600 times more selective for CaV1.3 than CaV1.2. Nifedipine, a common antihypertensive drug, was used in comparison as a non-selective, or slightly CaV1.2 selective antagonist of L-type calcium channels (IC50=0.016μM).

We also undertook immunohistochemistry of normal human adrenals, and APAs, in order to determine whether CaV1.3 is a ZG-selective LTCC and whether blockade may have greater expected effect on aldosterone secretion from APAs than from normal adrenal.

To study the role of CaV1.3 on aldosterone secretion, we first investigated the substitution mutations near the voltage-sensing domain, P1336R and V259D, on 24-h aldosterone production in transiently transfected H295R cells to find if the different changes seen in our electrophysiology data translated to changes in aldosterone secretion. We then contrasted the aldosterone secretion of cells transfected with mutant CaV1.3 channels to those transfected with wild-type CaV1.3 channel in the presence of compound 8 or nifedipine to study if blockade of calcium entry affects APAs with a CaV1.3 mutation differently.

Transfection of H295R cells with exogenous CaV1.3 was performed with β3 and α2δ accessory subunits, the subunits we used previously to show gain-of-function effects of the mutations on Ca2+ currents. As transfected channels and subunits do not necessarily emulate in vivo expression, we also tested the effect of compound 8 and nifedipine on endogenous CaV1.3
Results

**CaV1.3 mutations and compound 8 alter aldosterone production**

Transfection of H295R cells with CaV1.3 mutants P1336R and V259D caused a 2.4±0.2 (P=0.0004) and 2.1±0.2 (P=0.002) fold increase, respectively, compared to wild-type transfected H295R cells (Fig. 1a) and similarly in angiotensin II stimulated aldosterone production (Supplementary Fig. 1).

Exposure of H295R cells transfected with wild-type CaV1.3 to low concentration (1 µM) of compound 8 almost doubled aldosterone secretion (P=0.007), whereas high concentration (100 µM) of compound 8 decreased aldosterone production to 35±0.1% of basal level (P=0.003, Fig. 1b).

**Effect of calcium blockade on CaV1.3 genotypes**

In cells transiently transfected with wild-type, P1336R, or V259D CaV1.3, there was a similar biphasic effect of compound 8 on aldosterone secretion from the mutant P1336R cells, as that seen in wild-type CaV1.3. In mutant V259D cells, compound 8 was inhibitory only at 100 µM (Fig. 2a). Using the same protocol as for compound 8, the inhibitory effect of nifedipine on aldosterone secretion from CaV1.3 transfected H295R cells was determined. In wild-type CaV1.3 transfected cells, after treatment with 1 µM, 10 µM or 100 µM nifedipine, a 35±12 % decrease of aldosterone secretion was observed only at the highest concentration of nifedipine interrogated (100 µM)(P=0.0001, Fig. 2b) a considerable excess of its Kᵢ for CaV1.2 (IC₅₀=0.016 µM)¹²⁻¹³. In P1336R and V259D transfected cells, despite the increased
aldosterone secretion compared to that of wild-type cells (as seen at 0 µM), the presence of
high concentration of nifedipine (100 µM) decreased aldosterone production similarly across
all genotypes (Fig. 2b).

In non-transfected H295R cells (with only endogenous CaV1.3 and endogenous CaV1.3
accessory subunits present), compound 8 and nifedipine, 1-100 µM, decreased basal
aldosterone secretion (Fig. 2c).

**Compound 8 decreases aldosterone production in primary human adrenal cells**

In primary human adrenal cells cultured from the normal adjacent adrenals of patients with
an APA, 10 and 100 µM of compound 8 inhibited aldosterone production by 35±10 and
43±11 %, respectively (P<0.05; Fig. 3a). Cortisol secretion was also decreased to 72±1 and
50±4 % of basal level, respectively (P<0.05; Supplementary Fig. 2). As for nifedipine, effect
on aldosterone production was varied - not all cell cultures from the different patients
showed a reduction, even at the high concentration of 100 µM (Fig. 3b).

In APA cells, with increasing concentrations of 1, 10 and 100 µM, both compound 8 and
nifedipine showed a dose dependent decrease in aldosterone production, to a minimum
average of 54±2 and 43±13 % of basal level, respectively (P<0.005; Fig. 3c & d).

**Localization of CaV1.3 in adrenals containing an APA**

In sections of adjacent normal adrenal, that were adjacent to an APA or pheochromocytoma,
CaV1.3 was detected in the ZG and the zona reticularis (ZR) (Fig. 4a). *Only in ZG were*
juxtanuclear accumulation seen (as shown in the zoomed image), as ZR staining was mainly cytoplasmic (Fig. 4a). Exogenous CaV1.3 in H295R transfected cells had mainly membranous expression (Supplementary Fig. 3).

In APAs, different patterns of CaV1.3 expression were observed. CaV1.3 was expressed at the cell membrane, cytoplasmic, at the edge of cell clusters, or sparsely, or not at all (Fig. 4b and Supplementary Fig. 4).

Discussion

We previously reported that somatic mutation of CaV1.3 is present in a subset of APAs, distinguished by several features resembling normal ZG. In a large multi-centre study of 474 APAs, the frequency of CaV1.3 mutation was estimated to be 9.3%8. Although no particular histological phenotype was found in the multi-centre study8, one centre within the study did subsequently report that of their 71 APAs, CaV1.3 mutant APAs (3 of 71) were composed mainly of zona glomerulosa-like cells15. Thus the current approximation could be a substantial underestimation since (a) half of our selected ZG-like APAs that were exome sequenced had a CaV1.3 mutation6; and (b) our experience that such tumours are frequently too small to be detected by conventional adrenal imaging. We therefore wished to show whether the mutations are likely to increase aldosterone production, rather than trigger development of the adenoma, and whether this increase could be reversed by blockade of calcium entry. As few APAs are diagnosed in time to offer high likelihood of surgical cure from hypertension, and the increased recognition of aldosterone morbidity, a need arises for novel therapies that suppress aldosterone production, and lack the adverse effects of aldosterone receptor blockade and other less specific therapies for hypertension.
Herein we report that the two CaV1.3 mutations studied, selected for having different electrophysiological effects\(^6\), do increase aldosterone secretion of transfected human adrenocortical cells (Fig. 1). Furthermore, calcium blockade using compound 8, an investigational CaV1.3 inhibitor, and nifedipine, a non-selective L-type calcium channel inhibitor, reversed the increase (Fig. 2). The inhibition of aldosterone secretion was seen in the presence of the highest concentration of compound 8 interrogated in this study (100 \(\mu\)M) in H295R cells transfected with CaV1.3 mutants (Fig. 2); whereas in non-transfected H295R cells and primary adrenal cells, inhibition of aldosterone secretion could be seen at lower concentrations (1 and 10 \(\mu\)M) (Fig. 2c & Fig. 3d). We also postulate that regardless of whether a given APA has a somatic mutation of CaV1.3, the channel is often more active than in normal ZG cells, where immunohistochemistry suggests CaV1.3 is mainly internalised (Fig. 4).

Compound 8 was interrogated in this study as it was found to be the most selective CaV1.3 antagonist among 60,480 commercial compounds and a few hundred non-commercial compounds (Silverman lab) tested for efficacy in blocking CaV1.3 or CaV1.2 in stably transfected HEK293 cells. Compound 8 was reported to inhibit CaV1.3 >600-fold more potently than CaV1.2\(^{11}\). Subsequent studies have questioned this degree of selectivity, and even whether compound 8 is an agonist or antagonist\(^{16, 17}\). Nevertheless, it is well known that the effects of L-type Ca\(^{2+}\) channel blockade can differ among tissues depending on factors such as resting membrane potential\(^{18}\). Consequently, the hyperpolarisation of adrenocortical cells may have enhanced our ability to detect an antagonist effect of compound 8. Further, we may have fortuitously selected the \(\text{CACNB}\) isoform which maximises compound 8 selectivity, namely \(\text{CACNB3}\) (encoding for the \(\beta_3\) subunit). In subsequent analysis, however, we found \(\text{CACNB2}\) to be the predominant isoform in human
adrenal, indeed being one of the genes most up-regulated in ZG compared to zona
fasciculata (ZF). Thus, for the pharmacological responses of the different mutations to be
legitimately compared, a better CaV1.3 antagonist than Compound 8 is needed. Future
antagonists should be developed not only based on its selectivity for CaV1.3 but also on its
functionality with the prevalent accessory subunits in the human adrenal.

In our cells transfected with exogenous CaV1.3, the stimulatory effect of apparent low dose
calcium blockade on aldosterone secretion was observed only for Compound 8, but not
nifedipine. This increase in aldosterone secretion could have been due to low dose
compound 8 behaving as a channel activator; but toxicity (and hence leakage of
aldosterone) due to high calcium influx in transfected H295R cells cannot be dismissed, since
no stimulation of aldosterone secretion was seen in untransfected cells (Fig 2c). The
limitation of our expression CaV1.3 model, however, was that the cell line we used, H295R
cells, express a mixture of endogenous CaV1.2 and CaV1.3 whereas primary human ZG cells
express mainly CaV1.3. Moreover, the immortalised H295R cells were not a perfect
model for primary aldosteronism as other adrenal corticosteroids are secreted. This cell
line was used mainly due to the ease of transfecting exogenous mutant CaV1.3. Hence, to
supplement our transfection experiments, not only was compound 8 also studied in un-
transfected H295R cells, but also in primary adrenal cells (of which we have a limited
supply), to support endogenous CaV1.3 role in aldosterone regulation. To note, as we did not
find a linear relationship between increase in aldosterone production and amount of
transfected constructs, no correction for transfection efficiency whether by Western blots or
qPCR was performed. Transfection rates of exogenous CaV1.3 were confirmed as similar
visually, using its GFP-tag.
Previous studies have shown a number of dihydropyridines to reduce aldosterone secretion from adrenocortical cells\textsuperscript{22}. We chose nifedipine as a comparator because of experience with its use in patients, in whom it was the first dihydropyridine to be used\textsuperscript{23-25}, and also because of its modest Ca\textsubscript{v}1.2 selectivity. Nifedipine is expected to exert its Ca\textsubscript{v}1.2 blockade at concentrations around 4.45nM\textsuperscript{14}. At the lowest concentration of nifedipine that we had interrogated (1 µM), a concentration which should have easily blocked Ca\textsubscript{v}1.2, only some inhibition of aldosterone could be seen in non-transfected H295R and primary adrenal cells and none at all in H295R cells transfected with Ca\textsubscript{v}1.3 mutants (Fig. 2 & Fig. 3). The shallow concentration-response curves are consistent with blockade of different sites at low and high concentrations (Fig. 2 & 3). Dihydropyridines sometimes cause substantial reductions in plasma aldosterone in patients with primary aldosteronism\textsuperscript{26}. However this is not the predominant response at usual clinical doses, and increasing the dose to the presumed Ca\textsubscript{v}1.3-blocking range is precluded by the vascular side effects, particularly peripheral edema\textsuperscript{25,27}.

The potential attraction of selective Ca\textsubscript{v}1.3 blockade is that such a drug can be used at a dose which achieves substantial suppression of aldosterone secretion, without the vascular side effects of currently used L-type Ca\textsuperscript{2+} blockers\textsuperscript{25,27}. Previously, a T-type Ca\textsuperscript{2+} channel blocker, mibefradil, was introduced whose reduction in aldosterone secretion was among the theoretical advantages over L-type Ca\textsuperscript{2+} blockade\textsuperscript{28}; however the drug was withdrawn due to reports of dangerous and even fatal interactions with other drugs and was later found to cause serious effects on QTc\textsuperscript{29}. In vitro studies at least, have shown that single blockade of either L-type or T-type Ca\textsuperscript{2+} channels can decrease aldosterone production, even though the influx of Ca\textsuperscript{2+} in the ZG is thought to be mediated by both channels\textsuperscript{28,30-32}. While there has also been considerable attempt to develop inhibitors of aldosterone synthase as a
therapeutic class\textsuperscript{33}, these have foundered on the challenge of developing a drug, which
inhibits aldosterone synthase, without effect on the 95% homologous enzyme catalysing
cortisol synthesis (encoded by the gene \textit{CYP11B1}). By contrast, the homology between
\textit{Ca}_\text{v}1.2 and \textit{Ca}_\text{v}1.3 is only 75\%\textsuperscript{34}. Thus, even though compound 8 itself may not be the ideal
drug candidate to progress for treatment of hyperaldosteronism, there are a number of sites
outside the dihydropyridine-binding site where \textit{Ca}_\text{v}1.2 and \textit{Ca}_\text{v}1.3 differ sufficiently to
suggest that selective blockade is achievable.

Three drugs do have clinical efficacy in patients with primary aldosteronism: spironolactone,
eplerenone and amiloride\textsuperscript{35,36}. However, the efficacy of the latter two is modest, and the use
of spironolactone is severely limited in men by the anti-androgenic effects of higher
doses\textsuperscript{37,38}. All three drugs cause substantial increases in plasma aldosterone secretion,
probably secondary to the rise in plasma $K^+$, and there is some concern whether aldosterone
could have adverse vascular effects through a non-canonical aldosterone receptor\textsuperscript{39,40}.
Although no evidence exists in humans, there is an additional theoretical benefit from
blocking aldosterone synthesis rather than response – that such a drug could cause
involution of aldosterone-producing cells. This is suggested by the observation that genetic
deletion of the enzyme aldosterone synthase leads to apoptosis of the normal ZG cells\textsuperscript{41}.

In summary, we previously reported ZG-like APAs to have \textit{Ca}_\text{v}1.3 mutations. In this study, we
confirmed that \textit{Ca}_\text{v}1.3 is localized to the human adrenal ZG. By blocking endogenous \textit{Ca}_\text{v}1.3
in primary human adrenal and transfecting mutant \textit{Ca}_\text{v}1.3 in the human adrenocortical cell
line, H295R, we have also confirmed that \textit{Ca}_\text{v}1.3 plays a role in human adrenal
steroidogenesis. Taken together, these discoveries suggest that \textit{Ca}_\text{v}1.3 can provide a novel
mechanism and target for regulating excess aldosterone secretion and may be a novel way
of treating hyperaldosteronism, especially those caused by ZG-like APAs with a Ca\textsubscript{v}1.3 mutation. Since non-selective or Ca\textsubscript{v}1.2 selective dihydropyridines are dose-limited clinically by vascular effects, a selective Ca\textsubscript{v}1.3 antagonist may be valuable for suppressing aldosterone secretion in some patients with aldosterone-dependent hypertension.

Methods

Cell culture experimentation

H295R cells, were cultured in growth medium consisting of DMEM/Nutrient F-12 Ham supplemented with 10 % foetal bovine serum, 100 U of penicillin, 0.1 mg/mL streptomycin, 0.4 mM L-glutamine and insulin-transferrin-sodium selenite medium (ITS) at 37°C in 5% CO\textsubscript{2}. For transient transfection, wild-type or mutant P1336R or V259D Ca\textsubscript{v}1.3 GFP-tagged constructs were co-transfected together with constructs for \(\beta_3\) and \(\alpha_2\delta\) auxiliary subunits of Ca\textsubscript{v}1.3 into H295R cells using Amaxa Nucleofector kit R (Lonza, Germany) with electroporation program P20. The GFP-Ca\textsubscript{v}1.3 wild-type and mutant vectors were obtained from our collaborators; Dr. Jöerg Striessnig’s group at University of Innsbruck Center for Chemistry and Biomedicine, Austria. These constructs were derived from the ‘long’ isoform of the Ca\textsubscript{v}1.3 \(\alpha_1\) pore-forming subunit, with isoform A of the alternatively spliced exon 8. Transfected cells were seeded into 24-well plates at 100,000 cells per well in 0.5 mL of growth medium. At 24-h post-transfection, H295R cells were serum deprived in unsupplemented DMEM/Nutrient F-12 Ham medium for 24-h. At 48-h post-transfection, the transfection efficiency was visualised and qualitatively quantified by fluorescence microscopy. Further experiments were performed on cells with 60-80% transfection efficiency.
For primary cell culture, adrenocortical cells were obtained from the adrenals of patients with Conn’s syndrome that had undergone adrenalectomy at Addenbrooke’s Hospital, Cambridge, UK (Supplementary Table 1). Local ethical approval and informed consent were obtained for each patient and the procedures followed were in accordance with institutional guidelines. After macroscopic identification of APA and adjacent normal adrenal by a trained histopathologist, tissue samples were placed in growth medium within 45 minutes of surgical excision. The APA and adjacent normal adrenal was then digested separately in 3.3 mg/ml collagenase at 37 °C for 2h. Within a week of procurement, the primary human adrenocortical cells were then randomly seeded into 24-well plates at 50,000 cells per well in 0.5 mL of growth medium and allowed to settle for a further 48-h before drug treatments were performed.

**Drug treatments with Ca\textsubscript{2+}1.3 selective antagonist, compound 8, and L-type calcium blocker, nifedipine**

Compound 8 and nifedipine (Sigma-Aldrich, UK) were reconstituted in DMSO to stock concentrations of 1, 10, and 100 mM. Stock concentrations were further diluted (1:1000) in sterile un-supplemented DMEM/Nutrient F-12 Ham for treatments.

Transfected H295R cells were treated at 48-h post transfection (after 24-h of serum deprivation) with either vehicle or compound 8 or nifedipine in un-supplemented DMEM/Nutrient F-12 Ham medium in the presence of 10 nM angiotensin II. Supernatant and cells were harvested after 24-h incubation at 37°C.

For non-transfected H295R cells, cells were seeded into 24-well plates at 100,000 cells per well in 0.5 mL of growth medium, serum deprived for 24-h, and treated with either vehicle,
compound 8 or nifedipine in un-supplemented DMEM/Nutrient F-12 Ham medium.

Supernatant and H295R cells were harvested after 24-h incubation at 37°C.

For primary human adrenal cells, APA and adjacent normal adrenal cells were serum deprived for 24-h, and treated with either vehicle or compound 8 or nifedipine in un-supplemented DMEM/Nutrient F-12 Ham medium in the presence or absence of 10 nM angiotensin II. Supernatant and H295R cells were harvested after 24-h incubation at 37°C.

**Immunohistochemistry**

Immunohistochemistry was performed on formalin-fixed, paraffin-embedded adrenal sections (4 μm) using an automated immunostainer with cover tile technology (Bond-III system, Leica Biosystems). The commercial antibody of CACNA1D, clone N38/8 (UC Davis/NIH NeuroMab Facility; 1:500 dilution), was used as the primary antibody. Negative control experiments, in which the primary antibody was omitted, resulted in a complete absence of staining. Images were captured using a standard bright-field microscope, a U-TV1-X digital camera and CellID software (Olympus UK).

**Confocal Imaging**

H295R cells were cultured in complete media on sterilised and poly L-lysine coated coverslips at the density of 10^5 cells/well in 12-well cell-culture plate for 24h. Cells were serum-starved overnight before transfection. Serum-free media was replaced with antibiotic-free serum-containing media at the time of transfection with Lipofectamine 3000 transfection reagent (Life Technologies). Cells were co-transfected with GFP-tagged Ca\(_2\)1.3 WT, β\(_3\) & α\(_2\)δ constructs according to manufacturer’s instructions. 48h later plasma membranes of cells were stained with 2μg/ml Wheat Germ Agglutinin, Alexa Fluor® 633 Conjugate (W21404, Life
Technologies) in complete media for 10min at 37°C. Cells were washed twice with PBS (5min each), followed by fixing with 4% paraformaldehyde and permeabilisation with 1% trition-X100 (PBST), 10min each at room temperature. Cells were incubated with blocking buffer (3% BSA in PBS) for 1h at room temperature and overnight with α-Ca,1.3 (1:500, Clone N38/8, NeuroMab) in 3% BSA-PBST. Goat anti-mouse IgG, Alexa Fluor® 568 Conjugate (A11004, Life Technologies) was used as secondary antibody at 1:1000 dilution in 3%BSA-PBST for 1-h at room temperature. Finally cells were washed thrice in PBST and cover-slips were mounted on slides using VECTASHIELD Antifade Mounting Medium with DAPI (H-1200, Vector Laboratories). Confocal images were taken using Zeiss LSM510 Meta confocal microscope and analysed using Zen 2011 software.

Aldosterone concentration measurements

Aldosterone concentration was quantitatively measured using three methods due to availability of the kits; Coat-A-Count® Aldosterone (Siemens Medical Solutions, USA), a¹²⁵I solid-phase radioimmunoassay and after the discontinuation of this kit, an ELISA method adapted from researchers in Gomez-Sanchez’s group and finally a commercially available Homogenous Time Resolved Fluorescence Resonance Energy Transfer (HTR-FRET) assay from Cisbio Bioassays, France (used according to manufacturer’s instructions). ELISA was carried out using a selective validated aldosterone monoclonal antibody gifted to us and produced by Gomez-Sanchez’s lab, USA⁴². The aldosterone concentrations from transfected H295R cells were normalized to total cell protein, which was determined by performing the bicinchoninic acid (BCA) protein assay (Pierce Biotechnology, USA).

Statistical analysis
Experiments were performed with vehicle/plasmid controls where appropriate. Each experiment was performed with biological replicates and the averages were calculated. Aldosterone measurements are expressed as a ratio of basal (control) for each experiment. Results are shown as mean values ± SEM of separate experiments/transfections unless stated otherwise. Statistical analysis, two-tailed Student's t-tests or analysis of variance, was performed as indicated using the standard statistical software, Prism 6 (GraphPad Software, Inc).

References


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Author contributions

C.B.X. and L.H.S. designed and performed the experiments on CaV1.3 transfected H295R cells with the help of S.G.. E.A.B.A. and L.H.S. designed and performed experiments on non-transfected H295R cells and primary adrenal cells with the help of A.E.D.T. and J.Z.. W.Z. performed the immunohistochemistry and E.A.B.A. documented the results. S.K. and R.B.S. designed and provided compound 8. G.T. preformed the confocal microscopy. L.H.S. performed the statistical analysis on the data generated. C.M. provided the clinical information of patients. C.B.X., L.H.S., E.A.B.A. and M.J.B. wrote the manuscript.

Competing financial interests: The authors declare no competing financial interests.
Fig.1: CaV1.3 mutations and compound 8 alter aldosterone production

Comparison of stimulated aldosterone production in (a) wild-type (WT), P1336R, and V259D CaV1.3 transfected H295R cells (n=5) and in (b) different concentration of compound 8 on WT H295R cells (n=3).

Student t-test was used to calculate significance. **P<0.01 and ***P<0.001, compared to baseline (Wild-type or 0 M compound 8).

The n value represents number of separate experiment/transfection performed. Each experiment/transfection had 6 biological replicates. Aldosterone results shown here were measured by RIA method and are relative to basal level (Wild-type or 0 M compound 8).
Fig. 2: Effect of compound 8 on aldosterone production of different Ca\textsubscript{v}1.3 genotype

(a) Stimulated aldosterone secretion (n=3) in the presence of compound 8 and (b) stimulated aldosterone secretion in the presence of nifedipine (n=3) on WT, P1336R and V259D Ca\textsubscript{v}1.3 transfected H295R cells. There was a similar biphasic effect of compound 8 on aldosterone secretion from the mutant P1336R cells \( (P=0.02; \) Student’s \( t \)-test), as that seen in wild-type Ca\textsubscript{v}1.3, but not so in mutant V259D cells or when transfected cells were treated with nifedipine. (c) Comparison of basal aldosterone production of non-transfected H295R cells in the presence of 0-100 µM of compound 8 or nifedipine (n=3).

Two-way ANOVA was used to calculate overall significance. Table of \( P \)-values shows significance of mutation status (Mutation), concentration of treatment (Concentration), and type of treatment (Drug), on aldosterone production. The \( n \) value represents number of separate experiment/transfection performed. Each experiment/transfection had 6 biological replicates. Aldosterone was measured by RIA (a & b) or RIA and HTR-FRET (c) method. Results of both methods are relative to basal level (Wild-type or 0 M of treatment).
Fig. 3: Compound 8 decreases aldosterone production in primary human adrenal cells

Aldosterone secretion of (a & b) normal primary adrenal cells or (c & d) aldosterone-producing adenomas (APAs) in the presence of compound 8 (a & c) or nifedipine (b & d).

Dose response curve between 0-100 μM of compound 8 on (a) normal primary adrenal cells (n=4) and (c) APA cells (n=3), and nifedipine on (b) normal primary adrenal cells (n=3) and (d) APA cells (n=3).

Two-way ANOVA was used to calculate overall significance. Table of P-values shows significance of patient differences (Patient variability) and concentration of treatment (Concentration) on aldosterone production. The n value represents number of individual patient samples used for each experiment. Each concentration was replicated 2-12 times within each individual patient samples (which depended on quantity of primary cells available). Aldosterone results shown here are relative to 0 M of treatment.

Numbers 181, 182, 184, 187, 196, and 221 represent individual patient ID. Clinical data for these patients is provided in Supplementary Table 1. Primary cell cultures from patients 181, 182, and 184 were performed in the absence of angiotensin II (seen as solid bars) whereas primary cell cultures 187, 196, and 221 were stimulated with 10 nM angiotensin II (seen as hatched bars). Aldosterone was measured by RIA and ELISA method.
Figure 4: Localization of CaV1.3 in human adrenals

a) Immunohistochemistry (IHC) of CaV1.3 on formalin-fixed paraffin-embedded (FFPE) adrenal sections localized the channel to the zona glomerulosa (ZG) and zona reticularis (ZR) of the adrenals. In the ZG, cytoplasmic and juxtanuclear accumulation of CaV1.3 was observed whereas in the ZR, staining was mainly cytoplasmic. Picture is representative of 12 normal adjacent adrenal glands, 3 from patients with a phaeochromocytoma and 9 from patients with an aldosterone-producing adenoma (APA). C, capsule; G, zona glomerulosa; F, zona fasciculata; R, zona reticularis; M, adrenal medulla.

b) CaV1.3 expression in APA cells. IHC of CaV1.3 on FFPE adrenal sections were performed on three different types of APAs: (i-iii) ZG-like (low nucleus to cytoplasm ratio) APAs without a CaV1.3 mutation, (iv-vi) APAs with a CaV1.3 mutation, and (vii-ix) APAs with a KCNJ5 mutation. Immunostaining reveals a mixture of cytoplasm and membranous sublocalization in APA cells.
### a

<table>
<thead>
<tr>
<th>Compound 8 (μM)</th>
<th>WT</th>
<th>P1336R</th>
<th>V259D</th>
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</thead>
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**Two-way ANOVA**

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### b

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**Two-way ANOVA**

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### c

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**Two-way ANOVA**

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### Primary Adrenals Normal

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<tr>
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</table>

**Two-way ANOVA**

- **Patient variability**: 0.0004
- **Concentration**: < 0.0001

### Primary Adrenals Tumour

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<thead>
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<th>Aldosterone</th>
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<td>1.5</td>
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<tr>
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</table>

**Two-way ANOVA**

- **Patient variability**: 0.003
- **Concentration**: < 0.0001

### Primary Adrenals Normal

<table>
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<th>Aldosterone</th>
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<tbody>
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**Two-way ANOVA**

- **Patient variability**: < 0.0001
- **Concentration**: < 0.0001

### Primary Adrenals Tumour

<table>
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<tr>
<th>Nifedipine (μM)</th>
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**Two-way ANOVA**

- **Patient variability**: 0.003
- **Concentration**: < 0.0001