Calcium-sensing receptor residues with loss- and gain-of-function mutations are located in regions of conformational change and cause signalling bias

Gorvin, Caroline M; Frost, Morten; Malinauskas, Tomas; Cranston, Treena; Boon, Hannah; Siebold, Christian; Yvonne Jones, E; Hannan, Fadil M; Thakker, Rajesh V

Published in:
Human Molecular Genetics

DOI:
10.1093/hmg/ddy263

Publication date:
2018

Document version
Final published version

Document license
CC BY

Citation for published version (APA):

Terms of use
This work is brought to you by the University of Southern Denmark through the SDU Research Portal. Unless otherwise specified it has been shared according to the terms for self-archiving. If no other license is stated, these terms apply:

• You may download this work for personal use only.
• You may not further distribute the material or use it for any profit-making activity or commercial gain
• You may freely distribute the URL identifying this open access version

If you believe that this document breaches copyright please contact us providing details and we will investigate your claim. Please direct all enquiries to puresupport@bib.sdu.dk

Download date: 05. Jan. 2021
Calcium-sensing receptor residues with loss- and gain-of-function mutations are located in regions of conformational change and cause signalling bias

Caroline M. Gorvin1,†, Morten Frost1,2,¶, Tomas Malinauskas3, Treena Cranston4, Hannah Boon4, Christian Siebold3, E. Yvonne Jones3, Fadil M. Hannan1,5,¶ and Rajesh V. Thakker1,¶,*

1Academic Endocrine Unit, Radcliffe Department of Medicine, Oxford Centre for Diabetes, Endocrinology and Metabolism (OCDEM), University of Oxford, Oxford OX3 7LJ, UK, 2University of Southern Denmark, 5000 Odense C, Denmark, 3Division of Structural Biology, Wellcome Centre for Human Genetics, University of Oxford, Oxford OX3 7BN, UK, 4Oxford Molecular Genetics Laboratory, Churchill Hospital, Oxford OX3 7LJ, UK and 5Institute of Ageing and Chronic Disease, University of Liverpool, Liverpool L7 8TX, UK

*To whom correspondence should be addressed at: Academic Endocrine Unit, Radcliffe Department of Medicine, Oxford Centre for Diabetes, Endocrinology and Metabolism (OCDEM), University of Oxford, Oxford OX3 7LJ, UK. Tel: +44 (0)1865 857501; Fax: +44 (0)1865 875502; Email: rajesh.thakker@ndm.ox.ac.uk

Abstract

The calcium-sensing receptor (CaSR) is a homodimeric G-protein-coupled receptor that signals via intracellular calcium (Ca^{2+}i) mobilisation and phosphorylation of extracellular signal-regulated kinase 1/2 (ERK) to regulate extracellular calcium (Ca^{2+}e) homeostasis. The central importance of the CaSR in Ca^{2+}e homeostasis has been demonstrated by the identification of loss- or gain-of-function CaSR mutations that lead to familial hypocalciuric hypercalcaemia (FHH) or autosomal dominant hypocalcaemia (ADH), respectively. However, the mechanisms determining whether the CaSR signals via Ca^{2+}i or ERK have not been established, and we hypothesised that some CaSR residues, which are the site of both loss- and gain-of-function mutations, may act as molecular switches to direct signalling through these pathways. An analysis of CaSR mutations identified in >300 hypercalcaemic and hypocalcaemic probands revealed five ‘disease-switch’ residues (Gln27, Asn178, Ser657, Ser820 and Thr828) that are affected by FHH and ADH mutations. Functional expression studies using HEK293 cells showed disease-switch residue mutations to commonly display signalling bias. For example, two FHH-associated mutations (p.Asn178Asp and p.Ser820Ala) impaired Ca^{2+}i signalling without altering ERK phosphorylation. In contrast, an ADH-associated p.Ser657Cys mutation uncoupled signalling by leading to increased Ca^{2+}i mobilization while decreasing ERK phosphorylation. Structural analysis of these five CaSR disease-switch residues together with four reported disease-switch residues revealed these residues to be located at conformationally active regions of the CaSR such as the extracellular dimer interface and transmembrane domain. Thus, our findings indicate that disease-switch residues are located at sites critical for CaSR activation and play a role in mediating signalling bias.
Introduction

The calcium (Ca$^{2+}$)-sensing receptor (CaSR), a member of the class C subfamily of G-protein-coupled receptors (GPCRs), is highly expressed in the parathyroid glands and kidneys and plays an essential role in extracellular calcium (Ca$^{2+}$e) homeostasis by regulating parathyroid hormone (PTH) release and urinary Ca$^{2+}$ excretion (1). The CaSR is functionally active as a constitutive homodimer (2), with each monomer of the CaSR consisting of a large extracellular domain (ECD) that has been recently crystallized and shown to comprise a ligand-binding bilobed venus fly-trap domain (VFTD) linked to a cysteine-rich domain (3,4) and an intracellular domain (ICD), which are involved in activating downstream signalling proteins (5,6) (Fig. 1A and B). The CaSR couples to two major signal transduction cascades that comprise the G$\beta$$\gamma$-phospholipase C (PLC)-mediated generation of inositol 1,4,5-trisphosphate (IP$_3$) (7), which induces a rapid rise in intracellular calcium (Ca$^{2+}$i) concentrations (7) and the mitogen-activated protein kinase (MAPK) cascade, such as the extracellular signal-regulated kinase 1/2 (ERK) pathway (8). CaSR-mediated activation of MAPK signalling can occur by coupling to either the G$\beta$$\gamma$ or IP$_3$ pathways (9), and also by a G-protein-independent mechanism involving β-arrestin proteins (10).

The importance of the CaSR for the regulation of Ca$^{2+}$e has been highlighted by the identification of >320 different germline loss- and gain-of-function CaSR mutations that give rise to disorders of Ca$^{2+}$e homeostasis known as familial hypocalciuric hypercalcaemia type 1 (FHH1) and autosomal dominant hypocalcaemia type 1 (ADH1), respectively (11). Structural analysis has shown that FHH1- and ADH1-causing mutations usually affect different CaSR residues with FHH1-causing mutations being scattered throughout the VFTD and TMD regions (11), while ADH1-causing mutations cluster within the second extracellular loop (ECL) of the VFTD (residues 116–136) (12) and also within transmembrane (TM) helices 6 and 7 and the intervening third ECL of the TMD (residues 819–837) (10). However, four CaSR residues have been shown to be the location of both FHH1- and ADH1-causing mutations or variants on CaSR-mediated signalling, HEK293 cells were transiently transfected with pEGFP-N1-CaSR constructs expressing wild-type (WT) or mutant CaSR (Fig. S2), and Ca$^{2+}$i mobilisation and MAPK signalling was assessed following stimulation with Ca$^{2+}$e, as reported (19).

Identification of five CaSR disease-switch residue mutations

To evaluate the effect of the five FHH1-associated and ADH1-associated ECD and TMD disease-switch residue mutations and variants on CaSR-mediated signalling, HEK293 cells were transiently transfected with pEGFP-N1-CaSR constructs expressing wild-type (WT) or mutant CaSR (Fig. S2), and Ca$^{2+}$i mobilisation and MAPK signalling was assessed following stimulation with Ca$^{2+}$e, as reported (19).

Results

Identification of five CaSR disease-switch residues

An analysis of CaSR mutations and variants identified in >300 index cases of FHH1 or ADH1 that had been referred to our centre in Oxford (UK) since 2005 together with a review of previously reported CaSR mutations (17) identified five disease-switch residues, which are the location of both FHH1- and ADH1-associated mutations. Two of these disease-switch residues (Gln27 and Asn178) are located in the CaSR ECD. The Gln27 residue is affected by a novel FHH1-associated p.Gln27Pro variant and a reported FHH1-associated p.Gln27Arg mutation (18), and also a novel ADH1-associated p.Gln27Glu variant (Table 1 and Figs. 1 and S1); whereas the Asn178 residue is affected by a reported FHH1-associated p.Asn178Asp mutation (19) and also by a novel ADH1-associated p.Asn178Tyr variant (Table 1 and Figs. 1 and S1). Two disease-switch residues (Ser657 and Ser820) are located in the CaSR TMD. The Ser657 residue in TM3 is affected by a reported FHH1-associated p.Ser657Tyr mutation (20) and by a novel ADH1-associated p.Ser657Cys variant; whereas the Ser820 residue in TM6 is affected by a novel FHH1-associated p.Ser820Ala variant and a reported ADH1-associated p.Ser820Phe mutation (21) (Table 1 and Fig. 1). One disease-switch residue (Thr828) was shown to be located in ECL3, which links TM5 and TM7 of the CaSR, and this residue is affected by a novel FHH1-associated p.Thr828Ile variant and a novel ADH1-associated p.Thr828Asn variant (Table 1 and Fig. 1).

The importance of the CaSR for the regulation of Ca$^{2+}$e has been highlighted by the identification of >320 different germline loss- and gain-of-function CaSR mutations that give rise to disorders of Ca$^{2+}$e homeostasis known as familial hypocalciuric hypercalcaemia type 1 (FHH1) and autosomal dominant hypocalcaemia type 1 (ADH1), respectively (11). Structural analysis has shown that FHH1- and ADH1-causing mutations usually affect different CaSR residues with FHH1-causing mutations being scattered throughout the VFTD and TMD regions (11), while ADH1-causing mutations cluster within the second extracellular loop (ECL) of the VFTD (residues 116–136) (12) and also within transmembrane (TM) helices 6 and 7 and the intervening third ECL of the TMD (residues 819–837) (10). However, four CaSR residues have been shown to be the location of both germline loss- and gain-of-function mutations that cause FHH1 and ADH1, respectively, and are termed ‘switch’ or ‘toggle’ residues (4,14). Three disease-switch residues (Leu173, Pro221 and Glu297) are located within highly conserved regions of the VFTD that are involved in ligand binding (14,15), and one disease-switch residue (Asn802) is located in the third ICL of the TMD (16). The precise role of these disease-switch residues in CaSR function remains to be elucidated, and one possibility is that these residues act as molecular switches, which influence the transitioning of the CaSR between active and inactive conformations and direct CaSR signalling via the Ca$^{2+}$e or MAPK pathways. To gain further insights into the role of CaSR disease-switch residues, we sought for additional residues affected by loss- and gain-of-function mutations, by analysis of familial hypocalciuric hypercalcaemia and autosomal dominant hypocalcaemia patients who had been referred to our centre for diagnostic genetic testing of the CASR gene. Furthermore, we determined the effect of disease-switch residue mutations on CaSR-mediated Ca$^{2+}$i and ERK signalling and analysed the structural consequences of these mutations, using the recently determined crystal structures of the CaSR ECD (3,4).
Pro27 and Arg27 CaSR proteins had significantly reduced NFAT fold-change responses compared to WT cells, consistent with a loss-of-function; while the ADH1-associated Glu27 CaSR variant was associated with an elevated NFAT response, consistent with a gain-of-function (Fig. 2C). The effects of the Gln27 residue mutations on MAPK signalling were assessed by measuring fold-change phosphorylated ERK (pERK) responses following exposure to increasing Ca\textsuperscript{2+} concentrations ([Ca\textsuperscript{2+}]). Stimulation with Ca\textsuperscript{2+} induced a concentration-dependent increase in pERK responses in WT, Arg27 and Glu27 CaSR-expressing cells (Fig. 2D). These responses were significantly reduced at 5–7.5 mM Ca\textsuperscript{2+} for the FHH1-associated Arg27 mutant compared to WT, whereas cells expressing the FHH1-associated Pro27 mutant had absent pERK responses (Fig. 2D).
The ADH1-associated Glu27 mutant had significantly elevated pERK responses at 1–10 mM Ca\textsuperscript{2+}, consistent with a gain-of-function in MAPK signalling (Fig. 2D and Table S2). The effects of the Gln27 mutants on MAPK signalling were also assessed by measuring gene transcription induced by a serum-response element (SRE)-containing luciferase reporter construct, which is a downstream mediator of ERK signalling (26,27). These SRE responses were consistent with the pERK responses and indicate that the FHH1-associated Pro27 mutant abolished MAPK signalling (Fig. 2E), and thus exhibits an extreme bias towards Ca\textsuperscript{2+} signalling (Fig. 2F and Table S2); whereas the ADH1-associated Gln27 mutant had significantly elevated SRE responses and showed bias towards the MAPK pathway when compared to WT (Fig. 2F and Table S2).

Functional assessment of mutations affecting the Asn178 disease-switch residue showed the reported FHH1-associated Asp178 mutant (19) to cause a rightward shift of the Ca\textsuperscript{2+} concentration-response curve and lead to significantly increased EC\textsubscript{50} and significantly reduced maximal Ca\textsuperscript{2+} responses (Fig. 3A and B) compared to WT cells (Fig. 3A and B), consistent with the p.Asn178Tyr variant representing a pathogenic gain-of-function CaSR mutation. In keeping with these findings, NFAT reporter responses were also significantly reduced for the FHH1-associated Asp178 mutant and significantly increased for the ADH1-associated Tyr178 mutant (Fig. 3C), thereby indicating that these mutations cause a loss and gain of Ca\textsuperscript{2+} mobilisation, respectively. Evaluation of MAPK signalling showed the ADH1-associated Tyr178 mutant to cause a small but significant increase in pERK responses, whereas the FHH1-associated Asn178 mutant did not impair pERK or SRE reporter responses (Fig. 3D and E). Thus, the loss-of-function Asp178 mutant displayed a bias towards activation of pERK signalling when compared to WT CaSR (Fig. 3F and Table S2).

**TMD disease-switch residues.** An assessment of Ca\textsuperscript{2+} mobilisation mediated by Ser657 disease-switch residue mutations showed the reported FHH1-associated Tyr657 mutant (20) to cause significantly decreased Ca\textsuperscript{2+}-mediated Ca\textsuperscript{2+} maximal responses, and a significantly elevated Ca\textsuperscript{2+} EC\textsubscript{50} value compared to WT (Fig. 4A and B and Table S2). The novel ADH1-associated Cys657 variant led to significantly increased maximal Ca\textsuperscript{2+} responses, although the Ca\textsuperscript{2+} EC\textsubscript{50} response of the Cys657 protein was not significantly altered compared to WT (Fig. 4A and B and Table S2). To further characterise the functional consequences of the Ser657 residue mutations and variants, Ca\textsuperscript{2+} responses were determined using the NFAT reporter assay. This showed the FHH1-associated Tyr657 mutant to have abolished NFAT reporter responses, whereas the novel ADH1-associated Cys657 variant displayed significantly increased NFAT reporter activity (Fig. 4C), consistent with this representing a pathogenic gain-of-function CaSR mutation. An assessment of MAPK signalling showed the FHH1-associated Tyr657 mutant to have significantly reduced pERK and SRE reporter responses (Fig. 4D and E) consistent with this mutation leading to CaSR inactivation. In contrast, the ADH1-associated Cys657 mutant did not alter pERK or SRE reporter responses when stimulated with supraphysiological (≥7.5 mM) Ca\textsuperscript{2+}, which caused this mutant to have significantly impaired MAPK signalling (Fig. 4D and E). Thus, the FHH1-associated Tyr657 and ADH1-associated Cys657 mutant CaSR proteins result in biased signalling towards the Ca\textsuperscript{2+} pathway (Fig. 4F and Table S2).

**ECL3 disease-switch residue.** Functional assessment of mutations affecting the Thr828 disease-switch residue showed the novel FHH1-associated Ala820 variant to cause a rightward shift of the concentration-response curve and significantly increased EC\textsubscript{50} value (Fig. 5A and B and Table S2), which indicated that this represents a pathogenic loss-of-function CaSR mutation. Whereas, the reported ADH1-associated Phe820 mutant (21) showed a significantly reduced EC\textsubscript{50} value (Fig. 5A and B) and Table S2), consistent with a gain of CaSR function. In keeping with these findings, the ADH1-associated Phe820 mutant showed significantly elevated NFAT reporter responses at 1–2.5 mM Ca\textsuperscript{2+}, while the FHH1-associated Ala820 mutant reduced NFAT reporter responses at 7.5 mM Ca\textsuperscript{2+} (Fig. 5C). Assessment of MAPK responses revealed that the FHH1-associated Ala820 mutant induced a modest reduction in pERK responses following stimulation with 7.5 mM Ca\textsuperscript{2+} and had no effect on SRE reporter responses when compared to WT (Fig. 5D and E). The ADH1-associated Phe820 mutant showed a bimodal effect on MAPK signalling, with a significant increase in pERK and SRE responses at low [Ca\textsuperscript{2+}] (1–2.5 mM), but reduced responses at high [Ca\textsuperscript{2+}] (5–10 mM) compared to WT (Fig. 5D and E). Bias plots demonstrated that the FHH1-associated Ala820 mutant was biased towards MAPK signalling, while the Phe820 cells was more biased towards Ca\textsuperscript{2+} signalling (Fig. 5F and Table S2).

**Table 1.** FHH1- and ADH1-associated mutations affecting five CaSR disease-switch residues

<table>
<thead>
<tr>
<th>Wild-type</th>
<th>FHH1 mutation</th>
<th>ADH1 mutation</th>
</tr>
</thead>
<tbody>
<tr>
<td>Codon\textsuperscript{a}</td>
<td>Nucleotide</td>
<td>Amino acid</td>
</tr>
<tr>
<td>Gln27</td>
<td>c.80A &gt; G</td>
<td>Pro</td>
</tr>
<tr>
<td>Asn178</td>
<td>c.562A &gt; T</td>
<td>Asp</td>
</tr>
<tr>
<td>Ser657</td>
<td>c.1970C &gt; A</td>
<td>Tyr</td>
</tr>
<tr>
<td>Thr828</td>
<td>c.2483C &gt; T</td>
<td>Ile</td>
</tr>
</tbody>
</table>

\textsuperscript{a}Codon number according to full-length 1078 amino acid CaSR protein. All variants were observed in the heterozygous state.
functional consequences of the Thr828 residue variants, Ca\textsuperscript{2+}i responses were determined using the NFAT reporter assay. This showed the FHH1-associated Ile828 variant to have significantly reduced NFAT responses, whereas the ADH1-associated Asn828 variant had significantly increased NFAT responses (Fig. 6C), consistent with these variants representing pathogenic loss- and gain-of-function CaSR mutations, respectively. In keeping with the findings of the Ca\textsuperscript{2+}i signalling studies (Fig. 6A–C), an assessment of MAPK signalling showed the FHH1-associated Ile828 mutation to have significantly reduced pERK and SRE responses, whereas the ADH1-associated Asn828 mutation had significantly increased MAPK responses (Fig. 6D and E). Bias plots showed that both FHH1 and ADH1 mutants at the 828 residue had similar bias towards the Ca\textsuperscript{2+}i pathway to WT expressing cells (Fig. 6F).

Cellular expression of CaSR disease-switch residue mutations

Western blot analysis of whole cell lysates and plasma membrane fractions of HEK293 cells transiently transfected with WT or mutant CaSR disease-switch residue proteins was undertaken to determine whether alterations in cellular expression may have contributed to the loss- and gain-of-function caused by these mutant proteins. Untransfected HEK293 cells were shown not to endogenously express CaSR protein (Fig. S3A), whereas transfected cells expressed the monomeric CaSR as two bands at 140 and 160 kDa, and the dimeric CaSR at 250 kDa (Fig. S3A), consistent with previous reports (28). An assessment of the FHH1-associated and ADH1-associated ECD Gln27 and Asn178 disease-switch residue mutations showed no alterations in the
Figure 3. Effect of the ECD Asn178 disease-switch residue mutations on Ca$$^{2+}$$ and pERK signalling. (A) Concentration-response curves showing Ca$$^{2+}$$ responses following stimulation with Ca$$^{2+}$$e in HEK293 cells expressing WT (Asn178, black), or FHH1-associated (Asp178, blue) or ADH1-associated (Tyr178, red) mutant (m) CaSR proteins. Responses are expressed relative to the WT maximal responses with mean ± SEM of 4–10 biological replicates. (B) EC50 values obtained from the Ca$$^{2+}$$ concentration-response curves shown in panel A. (C–E) Concentration-response curves of Ca$$^{2+}$$e-induced (C) NFAT luciferase reporter responses, (D) pERK responses expressed as the ratio of pERK to total ERK concentrations and (E) SRE reporter responses of HEK293 cells expressing WT (Asn178) or FHH1-associated (Asp178) or ADH1-associated (Tyr178) mutant CaSR proteins. Responses at each [Ca$$^{2+}$$]e are expressed as a fold-change of basal [Ca$$^{2+}$$]e responses and shown as mean ± SEM of 6–12 biological replicates. (F) Bias plots for Ca$$^{2+}$$ and pERK signalling responses of mutations affecting the Asn178 CaSR disease-switch residue. Curves located above grey dotted line indicate signalling biased towards Ca$$^{2+}$$, while curves below grey dotted line indicate signalling biased towards pERK. Statistical analyses comparing WT versus Asp178 (blue asterisk), and WT versus Tyr178 (red asterisk) ***P-value < 0.0001, **P-value < 0.01, *P-value < 0.05 compared to WT, by a two-way ANOVA with Tukey’s multiple-comparisons test.

Structural characterisation of the CaSR disease-switch residues

The reported crystal structures of the ligand- and non-ligand-bound forms of the homodimeric form of the bilobed CaSR ECD (3,4) were analysed to determine whether the ECD disease-switch residues (Gln27 and Asn178) may act as molecular switches that could influence the adoption of the CaSR into an active or inactive conformation (Fig. 7A–E). Such crystal structures for the CaSR TMD are not established, and the structural properties of the TMD and ECL3 disease-switch residues (Ser657, Ser820 and Thr828) were therefore characterised using a homology model of the CaSR TMD (Fig. 7F–H), that is based on the crystal structure of human metabotropic glutamate receptor type 1 (mGluR1) in the inactive state, as previously described (10, 30). The location of the four previously reported CaSR disease-switch residues (Leu173, Pro221, Glu297 and Asn802) (14–16) were also mapped onto the ECD and TMD of the CaSR (Figs. 1A and B, 7F and S1).

ECD disease-switch residues. The Gln27 disease-switch residue is located within lobe 1 of the VFTD and forms hydrogen bonds
Figure 4. Effect of the TMD Ser657 disease-switch residue mutations on Ca\textsuperscript{2+} and pERK signalling. (A) Concentration-response curves showing Ca\textsuperscript{2+} responses following stimulation with Ca\textsuperscript{2+} in HEK293 cells expressing WT (Ser657, black), or FHH1-associated (Tyr657, blue) or ADH1-associated (Cys657, red) mutant (m) CaSR proteins. Responses are expressed relative to the WT maximal responses with mean ± SEM of 4–10 biological replicates. (B) EC\textsubscript{50} values obtained from the Ca\textsuperscript{2+} concentration-response curves shown in panel A. (C-E) Concentration-response curves of Ca\textsuperscript{2+}-induced (C) NFAT luciferase reporter responses, (D) pERK responses expressed as the ratio of pERK to total ERK concentrations and (E) SRE reporter responses of HEK293 cells expressing WT (Ser657) or FHH1-associated (Tyr657) or ADH1-associated (Cys657) mutant CaSR proteins. Responses at each [Ca\textsuperscript{2+}] are expressed as a fold-change of basal [Ca\textsuperscript{2+}] responses and shown as mean ± SEM of 6–12 biological replicates. (F) Bias plots for Ca\textsuperscript{2+} and pERK signalling responses of mutations affecting the Ser657 CaSR disease-switch residue. Curves located above grey dotted line indicate signalling biased towards Ca\textsuperscript{2+}, while curves below grey dotted line indicate signalling biased towards pERK. Statistical analyses comparing WT versus Tyr657 (blue asterisk), and WT versus Cys657 (red asterisk) ****P-value < 0.0001, ***P-value < 0.001, **P-value < 0.01, *P-value < 0.05 compared to WT, by a two-way ANOVA with Tukey’s multiple-comparisons test.

with the adjacent His344 (Fig. 7A–C). The side chain of Gln27 adopts similar conformations in all crystal structures (Fig. 7B), and the FHH1-associated p.Gln27Pro (Fig. 7B and C) mutation is predicted to abolish a hydrogen bond between an amino group of Gln27 and carbonyl oxygen of His344 (Fig. 7B and C). A similar analysis of the FHH1-associated p.Gln27Arg and ADH1-associated p.Gln27Glu mutations did not reveal the occurrence of any structural alterations involving the VFTD, using the available crystal structures (3,4). The Asn178 disease-switch residue is located at the lobe 1–lobe 1 VFTD dimer interface (Figs. 1, 7D and E) and, in the inactive conformation (Fig. 7D), forms an intramolecular hydrogen bond with the neighbouring Asn176 residue. However, in the active (ligand-bound) conformation (Fig. 7D), Asn178 could participate in hydrophilic interactions with the neighbouring Asn176 residue and form hydrogen bond interactions with Tyr246 from the opposing molecule in the CaSR dimer, thereby stabilising an active conformation of CaSR (3). The FHH1-associated p.Asn178Asp mutation is predicted to introduce a negative charge at the dimer interface, which could lead to repulsion between Asp178 and Asp48 from the opposing molecule in the active conformation (Fig. 7E). Thus, the p.Asn178Asp mutation may impair dimerisation and destabilise the active CaSR conformation. In contrast, the ADH1-associated p. Asn178Tyr mutation is predicted to introduce a bulky residue at the dimer interface that may potentially form pi–pi interactions with Tyr246 and stabilize the CaSR dimer (Fig. 7E), thereby leading to CaSR activation. Structural analysis also showed that the reported Leu173 (lobe 1) and Pro221 (lobe 2) disease-switch residues (14) are located at the hinge region of the VFTD dimer interface, while
Figure 5. Effect of the TMD Ser820 disease-switch residue mutations on Ca\(^{2+}\)\(_e\) and pERK signalling. Concentration-response curves showing Ca\(^{2+}\)\(_e\) responses following stimulation with Ca\(^{2+}\)\(_e\) in HEK293 cells expressing WT (Ser820, black) or FHH1-associated (Ala820, blue) or ADH1-associated (Phe820, red) mutant (m) CaSR proteins. Responses are expressed relative to the WT maximal responses with mean ± SEM of 4–10 biological replicates. (B) EC\(_{50}\) values obtained from the Ca\(^{2+}\)\(_e\) concentration-response curves shown in panel A. (C–E) Concentration-response curves of Ca\(^{2+}\)\(_e\)-induced (C) NFAT luciferase reporter responses, (D) pERK responses expressed as the ratio of pERK to total ERK concentrations and (E) SRE reporter responses of HEK293 cells expressing WT (Ser820), or FHH1-associated (Ala820) or ADH1-associated (Phe820) mutant CaSR proteins. Responses at each [Ca\(^{2+}\)\(_e\)] are expressed as a fold-change of basal [Ca\(^{2+}\)\(_e\)] responses and shown as mean ± SEM of 6–12 biological replicates. (F) Bias plots for Ca\(^{2+}\)\(_e\) and pERK signalling responses of mutations affecting the Ser820 CaSR disease-switch residue. Curves located above grey dotted line indicate signalling biased towards Ca\(^{2+}\)\(_e\), while curves below grey dotted line indicate signalling biased towards pERK. Statistical analyses comparing WT versus Ala820 (blue asterisk), and WT versus Phe820 (red asterisk) ***P-value < 0.0001, **P-value < 0.01, *P-value < 0.05 compared to WT, by a two-way ANOVA with Tukey’s multiple-comparisons test.

Discussion

Our studies have identified five CaSR disease-switch residues, and these findings together with the four previously reported...
disease-switch residues (14–16) indicate that the CaSR has at least nine disease-switch residues, which are the location of both germline loss- and gain-of-function mutations that lead to FHH1 and ADH1, respectively (Fig. 1). The finding that different mutations affecting the same residue may lead to distinct genetic disorders agrees with previous studies of the human vasopressin type 2 receptor (V2R), which is a class A GPCR involved in the regulation of renal water excretion (33,34). Thus, a loss-of-function germline mutation affecting the highly conserved Arg137 residue of the V2R, which is located in the TMD, has been shown to cause nephrogenic diabetes insipidus, which is characterized by excess renal water loss (33), whereas gain-of-function mutations of the V2R Arg137 residue give rise to inappropriate renal water retention that causes the nephrogenic syndrome of inappropriate antidiuresis (NSIAD) (34).

Structural analysis of the CaSR demonstrated that four extracellular disease-switch residues are located in key sites within the VFTD (Figs. S1 and 7). Thus, the reported Glu297 disease-switch residue (15) is located within the cleft formed by lobes 1 and 2 of the VFTD (Fig. 1) and has been shown to be involved in ligand binding which in turn leads to CaSR activation by mediating closure of the two lobes of the VFTD (3). Whereas, the Leu173, Asn178 and Pro221 disease-switch residues are located in the proximity of the hinge region (Figs. 1A and S1) between lobes 1 and 2, and also at the extended dimer interface that is formed by the two protomers of the VFTD upon agonist-mediated CaSR activation (Fig. 1) (3). Moreover, three of the TMD disease-switch residues clustered around TM6 (Figs. 1 and 7), which has been shown in structural studies of class A and B GPCRs to undergo a substantial outward movement in order
Figure 7. Structural analysis of CaSR disease-switch residues. (A) The Gln27 residue is located in the β1-strand of lobe 1 of the CaSR ECD, shown in the active conformation (PDB: 5K5S) (3) (Figs. 1 and S1). The peptide backbone amino group hydrogen on nitrogen (blue) of Gln27 forms a hydrogen bond (H-bond) with the peptide carbonyl group of His344 located in the α10 and α11 loop. (B) Superposition of all known crystal structures of CaSR: active conformation (carbons are in dark red as in panels A and C; PDB: 5K5T) (3),inactive conformation (cyan, PDB: 5K5T) (3), cyclomethyltryptophan/magnesium-bound form (green, PDB: 5FBK) (4), cyclomethyltryptophan/magnesium/gadolinium-bound form (blue, PDB: 5FBH) (4). The Gln27-His344 interaction is not altered in the active and inactive conformations of the CaSR. (C) The FHH1-associated p.Gln27Pro mutation abolishes the Gln27-His344 hydrogen bond (circled). (D) The Asn178 is located at the lobe 1-lob e1 dimer interface (Figs. 1 and S1). The side chain of Asn178 is exposed to solvent in the inactive conformation (cyan, PDB: 5K5T) (3) and is positioned at ≥7.0 Å from neighbouring residues (Asp48, Tyr246) on the opposing CaSR molecule. (E) The side-chain of Asn178 is located within a tightly packed environment between two CaSR ECD protomers in the active conformation (PDB: 5K5S). Mutation to Asp178 or Tyr178 could alter the relative orientation between the two protomers. (F) Homology model of the CaSR TMD viewed parallel to the cell membrane and based on the structure of mGluR1 (PDB: 4OR2) (30). Side chains of Ser657, Asn802, Ser820 and Thr828 are shown as spheres (dashed circles). Seven TM helices are numbered. (G) The Ser657 residue is located within TM2 and its side chain projects towards Cys691 located on TM3. (H) The Ser820 residue is located within TM6 and its side chain points towards the transmembrane bilayer. The Thr828 residue is located within ECL3, which connects TM6 and TM7.

Thus, our finding that ECD and TMD CaSR disease-switch residues are located within regions that undergo large conformational changes upon ligand binding is consistent with an analysis of molecular switch residues identified in class A GPCRs, which has shown these to be located between the GPCR ligand-binding pocket and the G-protein interaction site (38). In this region of the TMD, class A GPCR molecular switch residues were shown to particularly cluster around the kink region of TM6, which undergoes the greatest structural alteration during receptor activation (38). Moreover, large conformational changes upon ligand binding involving residues within TM3 and TM6 have also been shown to be important in class C GPCR activation and G-protein coupling (39,40).

We used CaSR crystal structures (3,4) to obtain detailed insights into how the two VFTD disease-switch residues identified in this study may be influencing CaSR activation. Our analysis indicated that the Asn178 residue adopts distinct conformations and forms different interactions in the ligand bound and unbound forms of the CaSR (Fig. 7D and E). In the inactive CaSR, Asn178 forms weak intramolecular hydrogen bonds with Asn176, whereas upon ligand binding Asn178 interacts with the opposing molecule and could facilitate agonist-induced dimerisation of the VFTD, which in turn triggers CaSR-mediated signal transduction (3). It is therefore possible that the Asn178 residue and its mutations may switch the CaSR between active and inactive conformations. This finding that residues affected by loss- and gain-of-function mutations can act as molecular

...
switches is consistent with that previously reported for the Arg137 molecular switch residue of the human V2R, which participates in an ionic lock mechanism common to class A GPCRs and determines whether these receptors adopt active or inactive conformations (33,34). However, our analysis of the Gln27 residue within the context of the available crystal structures revealed that this residue was unlikely to be acting as a molecular switch as its interactions with other residues did not change upon CaSR activation (Fig. 7B). Moreover, Gln27 is not located in a region directly involved in ligand-binding or in mediating large conformational changes. Instead, the finding that Gln27 mutations can lead to loss- or gain-of-function is most likely a consequence of the specific mutations affecting this residue rather than due to the properties of the WT residue. Detailed structural analysis of the TMD disease-switch residues was not possible because the crystal structure of the CaSR TMD has not been established. However, homology modelling of the CaSR TMD based on the crystal structure of the related mGluR1 (10) provided insights into the role of the Ser657 disease-switch residue, which may mediate interactions between TM2 and TM3 (Fig. 7C). It should be noted that all of the TMD disease-switch residues detected to-date (Ser657, Asn802, Ser820 and Thr828) have polar side chains. Previous studies of class A GPCRs have shown that conserved TMD polar residues contribute to a water-mediated hydrogen-bonded network within the TMD that plays a key role in regulating receptor activity and that engineered alanine substitutions of these residues can cause loss or gain of receptor function (41). Thus, the polar TMD disease-switch residues may potentially play a complex role in mediating interactions between the different helices within the CaSR TMD.

In vitro functional expression studies of mutations affecting the five disease-switch residues identified in this study have provided insights into the role of these residues in mediating CaSR signal transduction. For example, most of the disease-switch residue mutations showed similar levels of cell-surface expression compared to WT CaSR (Fig. S3), and it is thus unlikely that the alterations in signalling caused by these mutants is a consequence of altered anterograde or retrograde receptor trafficking. Moreover, none of the disease-switch residue mutations had significant effects on CaSR function in the absence of the agonist (Ca2+)e, and therefore these disease-switch residues are not involved in regulating the basal activity of the CaSR. However, mutations affecting the Gln27, Ser657 and Thr828 disease-switch residues altered the maximal responses of the CaSR without influencing EC50 values, which may indicate that these mutations altered the intrinsic signalling efficacy of the CaSR without affecting agonist affinity (42). Another possibility is that the alterations in maximal Ca2+i responses may be due to changes in cell-surface expression of the CaSR switch residue mutants. However, an alteration in cell-surface expression was only demonstrated for the Tyr657 disease-switch residue mutant. Moreover, in contrast with other ADH1-associated disease-switch residue mutants, which showed increased maximal Ca2+i responses, the gain-of-function Phe820 disease-switch residue mutant showed reduced maximal Ca2+i responses compared to WT, which indicates that the CaSR activation caused by this mutant may have led to depletion of intracellular stores of Ca2+, as reported (29).

As most of the nine CaSR disease-switch residues are located in regions of the CaSR that are predicted to undergo conformational rearrangements upon ligand binding (Figs. 1 and 7) (3), mutations affecting these residues have the potential to stabilise the CaSR in conformations that selectively activate G-protein dependent or independent signalling pathways (43). In keeping with this, many of the CaSR disease-switch residue mutations showed distinct patterns of signalling bias compared to the WT CaSR, which preferentially activates Ca2+i, mobilisation (29). The disease-switch residue mutations were commonly found to show signalling bias towards either Ca2+i (FH11-associated p.Gln27Pro or ADH1-associated p.Asn178Tyr and p.Ser657Cys) or MAPK pathway activation (FH11-associated p.Asn178Asp and p.Ser820Ala, or ADH1-associated p.Gln27Glu). In particular, the FH11-associated p.Asn178Asp and p.Ser820Ala mutations led to reductions in Ca2+i mobilisation without any significant impairment of pERK responses. Thus, these mutations may result in the CaSR adopting conformational states that favour signalling away from Gq11-mediated PLC activation and towards Gi/o and/or β-arrestin-mediated MAPK signalling, as has been recently demonstrated for a CaSR mutation that disrupts a salt-bridge within the TMD (10). In contrast, the ADH1-associated p.Ser657Cys disease-switch residue mutation showed signalling bias towards Ca2+i mobilization, as this mutation significantly increased Ca2+i responses, yet led to a reduction in MAPK signalling as assessed by pERK and SRE reporter responses (Fig. 4). To our knowledge, this is the first disease-causing CaSR mutation reported to cause both loss- and gain-of-function, depending on which signalling pathway is evaluated, and highlights a potential uncoupling of Ca2+i and MAPK responses. This understanding of signalling bias caused by CaSR mutations may provide a precision medicine approach to the treatment of patients with symptomatic forms of FHH1 and ADH1. For example, FHH1 patients harbouring a CaSR mutation such as p.Ser820Ala, which shows a preferential impairment of Ca2+i responses, may benefit from treatment with cinacalcet, as this CaSR positive allosteric modulator (PAM) is biased towards the Ca2+i mobilisation pathway (44), whereas FHH1-causing mutations such as p.Thr828Ile that cause a preferential reduction in MAPK signalling may benefit from treatment with the AC265347 compound, as this CaSR PAM causes biased activation of pERK (45).

In conclusion, our studies have demonstrated that CaSR residues affected by disease-causing loss- and gain-of-function mutations are located within receptor domains that are pivotal for CaSR function and may undergo conformational rearrangements during GPCR activation. Moreover, these disease-switch residues likely determine whether the CaSR signals via Ca2+i or MAPK pathways.

Materials and Methods
Bioinformatic analysis of DNA sequence variants
Publicly accessible databases [dbSNP (http://www.ncbi.nlm.nih.gov/projects/SNP/) (46) and the ExAC (exac.broadinstitute.org), which contains details from exomes of 60 706 unrelated individuals (22)] were examined for the presence of CASR sequence variants.

Protein sequence alignment and structural analysis
Protein sequences of CaSR orthologs were aligned using ClustalOmega (http://www.ebi.ac.uk/Tools/msa/clustalo/) (47). Crystal structures of CaSR ECDs in both active and inactive conformations were analysed [Protein Data Bank (PDB) accession codes 5FBK, 5FBH, 5K5S, 5KST] (3,4). The HHpred homology detection server (https://toolkit.tuebingen.mpg.de/hhpred) was used to identify crystal structures with similar sequences to the CaSR TMD, and to perform amino acid sequence alignment (30,48). The CaSR sequence was threaded onto the mGluR1 (PDB ID 4OR2) (30) template coordinates using Modeller
**Cell culture and transfection**

Studies were performed in HEK293 cells maintained in DMEM-Glutamax media (ThermoFisher) with 10% fetal bovine serum (Gibco) at 37°C, 5% CO₂. Mutations were introduced into the reported WT pEGFP-N1-CaSR construct by site-directed mutagenesis using the Quikchange Lightning Kit (Agilent Technologies) and gene-specific primers (SigmaAldrich), as described. WT and mutant pEGFP-N1-CaSR constructs, and luciferase reporter constructs (pGL4.30-NFAT and pGL4.33-SRE, Promega) were transiently transfected into HEK293 cells using Lipofectamine 2000 (LifeTechnologies) 48 h before experiments, as described (50).

**Western blot analyses**

Expression of WT and mutant proteins by the pEGFP-N1-CaSR constructs was confirmed by Western blot analysis, with the calnexin housekeeping protein being used as a loading control (50). For cell fractionation studies, cells were transfected with CaSR constructs and 48 h later plasma membrane and cytoplasmic fractions were isolated using a plasma membrane extraction kit (Catalog No. 65400, Abcam), as described (51). Plasma membrane calcium ATPase (PMCA1) protein was used as a loading control for plasma membrane fractions. The following primary antibodies were used for Western blot analysis: anti-CaSR (ADD, ab19347, Abcam), anti-calnexin (ab2301, Millipore) and anti-PMCA1 (ab190355, Abcam). The Western blots were visualized using an Immuno-Star WesternC kit (BioRad) on a Chemidoc XRS+ system (50).

**Ca²⁺ measurements**

Ca²⁺-induced Ca²⁺ responses were measured by Fluo-4 calcium assays as previously described (52). HEK293 cells were plated in 96-well plates (Corning) and transiently transfected with 1000 ng/mL WT or mutant pEGFP-N1-CaSR constructs. Ten wells were transfected with each construct. Cells were loaded with Fluo-4 dye, prepared according to manufacturer’s instructions (Fmriogen), for 30–60 min at 37°C. Baseline measurements were made for each well and increasing doses of CaCl₂ injected automatically into each well. One well was used for each [Ca²⁺]₀. Changes in Ca²⁺ were recorded on a PHERAstar instrument (BMG Labtech) at 37°C with an excitation filter of 485 nm and an emission filter of 520 nm. The peak mean fluorescence ratio of the transient response following each individual stimulus was measured using MARS data analysis software (BMG Labtech). Responses were normalized to the maximal response of WT expressing cells and plotted using GraphPad Prism (29). Assays were performed in 4–12 biological replicates (independently transfected wells, performed on at least 4 different days) for each of the expression constructs. Statistical analysis was performed using two-way analysis of variance (ANOVA) and the F test for EC₅₀ values (50,52).

**Functional assays**

AlphaScreen assays were undertaken to measure pERK using cells transiently transfected with 100 ng of WT or mutant pEGFP-N1-CaSR constructs. Cells were treated with 0.1–10 mM CaCl₂ for 5 min then lysed in Surefire lysis buffer and pERK total and ERK assays performed, as previously described (10,52). The fluorescence signal in AlphaScreen assays was measured using the PHERAstar FS microplate reader (BMG Labtech) (10,52). Assays were performed in 6–12 biological replicates (independently transfected wells, performed on at least 3 different days) for each of the expression constructs. Luciferase reporter assays were undertaken to measure SRE and NFAT responses, as previously described (50,52). Cells were transiently transfected with 100 ng/mL of the WT or mutant pEGFP-N1-CaSR constructs, 100 ng/mL luciferase construct (either pGL4-NFAT or pGL4-SRE) and 10 ng/mL pRL. At 48 h post-transfection, cells were then treated with 0–10 mM CaCl₂ and incubated for 4 h prior to lysis and measurement of luciferase activity using Dual-Glo Luciferase (Promega) on a Veritas Luminometer (Promega), as previously described (50). Luciferase:renilla ratios were expressed as fold-changes relative to responses at basal CaCl₂ concentrations (0.1 mM). Assays were performed in 8–12 biological replicates (independently transfected wells, performed on at least 4 different days) for each of the expression constructs. Statistical analysis was performed using two-way ANOVA. To construct the bias plots the responses at each [Ca²⁺]₀ in Ca²⁺ (x-axis) and pERK (y-axis) assays, were plotted against each other, as previously described (29). Data was normalized to WT levels, expressed between 0 and 100%. If the agonist showed no preference for one pathway over another the concentration-response curves would be coincident, and the bias plot would overlay the line of identity (dotted line in figures) (29). If the WT or mutant CaSR couples more favourably to one pathway over another, the bias plot line will fall on either side of the line towards the preferred pathway.

**Bias factor calculation**

Bias factor calculations were based on previous studies (53,54). To normalise data, responses of each mutant CaSR construct were expressed relative to WT as a percentage. Nonlinear regression of concentration-response curves was performed with GraphPad Prism (GraphPad) and the EC₅₀ and Emax calculated. The intrinsic relative activity (RA₁) value was calculated using the following equation for each mutant and WT CaSR:

\[
RA₁ = \frac{E_{\text{max}1}}{E_{\text{max}2}} \frac{EC_{50,1}}{EC_{50,2}}
\]

where ‘1’ denotes response 1 (Ca²⁺) and ‘2’ denotes response 2 (pERK). The bias factor was calculated relative to WT as follows:

\[
\text{Bias factor} = \log \frac{RA₁_{\text{mutant}}}{RA₁_{\text{WT}}}
\]

**Statistical analysis**

A minimum of four independent biological replicates (independently transfected wells, performed on at least 3 different days) were used for all statistical comparisons. All data was analysed by two-way ANOVA with Tukey’s multiple-comparisons test. Statistical analyses were undertaken using GraphPad Prism (GraphPad), and a value of P-value < 0.05 was considered significant for all analyses. Comparison of EC₅₀ values was performed using the F test (50,52).
Supplementary Materials

Supplementary Materials are available at HMG online.

Conflict of Interest statement. None declared.

Funding

Wellcome Trust Investigator Award (106995/Z/15/Z to R.V.T.); National Institute for Health Research (NIHR) Oxford Biomedical Research Centre Program (to R.V.T.); and NIHR Senior Investigator Award (NF-SI-0514-10001 to R.V.T.); Cancer Research UK (C20724/A14414 to T.M. and C.S.); the European Research Council (647278 to C.S.); a long-term postdoctoral fellowship from the Human Frontier Science Program (LT000021/2014-L to T.M.).

References


