REVIEW

Insights into calcium-sensing receptor trafficking and biased signalling by studies of calcium homeostasis

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Abstract

The calcium-sensing receptor (CASR) is a class C G-protein-coupled receptor (GPCR) that detects extracellular calcium concentrations, and modulates parathyroid hormone secretion and urinary calcium excretion to maintain calcium homeostasis. The CASR utilises multiple heterotrimeric G-proteins to mediate signalling effects including activation of intracellular calcium release; mitogen-activated protein kinase (MAPK) pathways; membrane ruffling; and inhibition of cAMP production. By studying germline mutations in the CASR and proteins within its signalling pathway that cause hyper- and hypocalcaemic disorders, novel mechanisms governing GPCR signalling and trafficking have been elucidated. This review focusses on two recently described pathways that provide novel insights into CASR signalling and trafficking mechanisms. The first, identified by studying a CASR gain-of-function mutation that causes autosomal dominant hypocalcaemia (ADH), demonstrated a structural motif located between the third transmembrane domain and the second extracellular loop of the CASR that mediates biased signalling by activating a novel \(\beta\)-arrestin-mediated G-protein-independent pathway. The second, in which the mechanism by which adaptor protein-2 \(\sigma\)-subunit (AP2\(\sigma\)) mutations cause familial hypocalciuric hypercalcaemia (FHH) was investigated, demonstrated that AP2\(\sigma\) mutations impair CASR internalisation and reduce multiple CASR-mediated signalling pathways. Furthermore, these studies showed that the CASR can signal from the cell surface using multiple G-protein pathways, whilst sustained signalling is mediated only by the \(G_{q/11}\) pathway. Thus, studies of FHH- and ADH-associated mutations have revealed novel steps by which CASR mediates signalling and compartmental bias, and these pathways could provide new targets for therapies for patients with calcaemic disorders.

Introduction to the CASR

Extracellular calcium (Ca\(^{2+}\)) is required for diverse biological functions ranging from blood coagulation, mineralisation of bone matrix, muscle contraction and hormone secretion (Brown 1991). Thus, calcium concentrations within the blood are tightly regulated. The parathyroid gland plays an essential role in calcium...
homeostasis by detecting Ca\(^{2+}\) in the blood, and in response to hypocalcaemia, secretes the parathyroid hormone (PTH) to normalize serum calcium concentrations. PTH achieves this by: enhancing bone resorption; activating calcium reabsorption at the kidneys; and stimulating the synthesis of 1,25-dihydroxyvitamin D\(_3\), which mobilises intestinal calcium absorption (Fig. 1A) (Riccardi & Brown 2010). The net effect of these three pathways is to increase Ca\(^{2+}\), which provides feedback inhibition to the parathyroid gland, to suppress PTH secretion (Conigrave & Ward 2013). The parathyroid is able to detect Ca\(^{2+}\) concentrations in the blood using the cell-surface-expressed calcium-sensing receptor (CASR), a class C G-protein-coupled receptor (GPCR), for which Ca\(^{2+}\) is the major ligand (Riccardi & Brown 2010, Conigrave & Ward 2013).

The CASR is a 1078-amino acid protein that exists at cell surfaces as a disulphide-linked homodimer (Ward et al. 1998), although it is capable of forming heterodimers (Gama et al. 2001, Chang et al. 2007). The CASR has a large extracellular domain, which was recently crystalized by two independent laboratories, and consists of a bilobed venus flytrap ligand binding domain (VFTD) and a cysteine-rich domain (CRD) (Geng et al. 2016, Zhang et al. 2016) (Fig. 1B). Ca\(^{2+}\) binds between the two lobes (lobe 1 and lobe 2) of the VFTD, which initiates a conformational change, facilitating lobe 2-lobe 2 dimerisation and allowing the CRDs to interact (Geng et al. 2016, Zhang et al. 2016). These conformational changes are predicted to re-orientate the seven-transmembrane (TM) domain (Geng et al. 2016), and consequently activate the associated G-proteins and initiate signal transduction (Standfuss et al. 2011). The CASR has been reported to couple to multiple G-protein subtypes, but predominantly signals by: the G\(_{i/o}\) pathway, to suppress cAMP and activate mitogen-activated protein kinase (MAPK) cascades (Kifor et al. 2001, Thomsen et al. 2012); the G\(_{q/11}\)-phospholipase C (PLC)-mediated pathway, to generate inositol 1,4,5-trisphosphate (IP\(_3\)) and diacylglycerol, that activate intracellular calcium (Ca\(^{2+}\)) mobilisations (Hofer & Brown 2003) and MAPK pathways, respectively. The CASR can also couple to a G-protein-independent mechanism involving β-arrestin proteins to also activate MAPK signalling (Thomsen et al. 2012) (Fig. 2).

### Human disorders of the CASR provide insights into receptor signalling mechanisms

The importance of the CASR in the regulation of Ca\(^{2+}\) is highlighted in patients with germline mutations of the receptor that have disorders of calcium homeostasis. Heterozygous loss-of-function mutations in the CASR lead to the autosomal dominant condition familial hypocalciuric hypercalcaemia (FHH) (Pollak et al. 1993), which is characterised by lifelong mild-to-moderate hypercalcaemia, normal or mildly raised serum PTH levels and low urinary calcium excretion (FIREK et al. 1991, Marx 2015). FHH is considered to be a benign disorder as most patients are asymptomatic, but it can be associated with chondrocalcinosis and pancreatitis in some cases (Pearce et al. 1996a, Volpe et al. 2009, Hannan & Thakker 2013).
Homozygous and compound heterozygous loss-of-function mutations of the CASR are the common cause of neonatal severe hyperparathyroidism (NSHPT), which is characterised by marked elevations in serum calcium and PTH, failure to thrive and hyperparathyroid bone disease (Pollak et al. 1993, Chattopadhyay & Brown 2006, Hannan & Thakker 2013). On occasion, heterozygous CASR mutations may lead to an NSHPT-like phenotype, and this is largely due to the dominant-negative nature of the mutant receptor on the wild-type (WT) CASR in these cases (Pearce et al. 1995, Obermannova et al. 2009).

Gain-of-function mutations of the CASR cause autosomal dominant hypocalcaemia (ADH) characterised by mild-to-moderate hypocalcaemia and inappropriately low or normal PTH concentrations (Pearce et al. 1996b, Hannan & Thakker 2013). Up to 50% of patients present with hypocalcaemic symptoms of paraesthesia, carpopedal spasms, seizures and ectopic calcification of the kidneys and basal ganglia (Pearce et al. 1996b, Hannan & Thakker 2013). Some patients with gain-of-function mutations in CASR may have Bartter syndrome type 5, which is characterised by renal salt wasting, hypokalaemia, hyperreninaemia and hyperaldosteronaemia (Vargas-Poussou et al. 2002, Watanabe et al. 2002).

Biased signalling of the CASR

Functional studies in HEK293 cells have demonstrated that disease-causing mutations may influence CASR signalling responses in a biased manner (Leach et al. 2012). Despite the capability of the CASR to signal via multiple signalling pathways, in vitro studies in HEK293 cells have shown that it preferentially couples to the Ca\(^{2+}\) signalling pathway (Leach et al. 2012). In contrast, disease-causing CASR mutants have been shown to switch this preferential signalling, with some FHH1-causing mutations signalling equally via the Ca\(^{2+}\) and MAPK pathways, or predominantly via MAPK pathways, while many ADH1-associated mutants couple more strongly to Ca\(^{2+}\) (Leach et al. 2012). Furthermore, studies of positive and negative allosteric CASR modulating compounds, revealed they too can mediate a biased signalling response, with both classes of drugs influencing Ca\(^{2+}\) to a greater extent than ERK1/2 phosphorylation (Leach et al. 2013). Thus, these findings established that agonist-induced CASR signalling may occur in a biased manner, although the GPCR structural motifs mediating ligand-dependent bias were not described.

Novel insights into mechanisms by which CASR can mediate signalling bias

Recently, a novel ADH1-associated CASR mutation, Arg680Gly, has provided some insights into the molecular mechanisms mediating signalling bias (Gorvin et al. 2018a). In vitro analyses showed that Ca\(^{2+}\) signalling in cells expressing the Arg680Gly mutation was not different to that observed in WT cells, in contrast to previously reported ADH1-associated CASR mutants (Gorvin et al. 2018a). However, the Arg680Gly mutation did enhance MAPK signalling (Gorvin et al. 2018a). Furthermore, this type of signalling bias, in which an ADH1 mutation enhances MAPK signalling but does not affect Ca\(^{2+}\), has not previously been described and provided an opportunity to explore the role of the Arg680 residue in CASR structure-function. As MAPK acts as a convergence pathway for multiple CASR signalling pathways (Fig. 2), each was investigated using a single reporter assay measuring...
Luciferase upstream of a serum-response element (SRE), which acts as a measure for MAPK-mediated transcription (Gorvin et al. 2017b). By applying compounds that specifically block the G_{i/o} (YM-245890 and UBO-QIC), the G_{i/o} (pertussis toxin) or the β-arrestin (β-arrestin-1 and β-arrestin-2 targeting siRNA) pathways, it was shown that the Arg680Gly mutation enhanced MAPK signalling by a β-arrestin1/2-mediated pathway (Takasaki et al. 2004, Schrage et al. 2015, Gorvin et al. 2018a).

The structural location of the mutant residue within the CASR transmembrane domain (TMD) region provides some insights into the likely mechanism mediating this bias. Homology modelling of the CASR TMD, based upon the structure of the closely related human metabotropic glutamate receptor 1 (mGlur1) (Hu et al. 2005, Gorvin et al. 2018a), predicts that the Arg680 residue lies at the extracellular side of TM3, and that the residue forms salt-bridge connections with the side chain of adjacent residues in extracellular loop 2 (ECL2) (Glu767) or TM7 (Glu837) (Fig. 3). Such connections between residues within TMDs or with ECLs of GPCRs are known to be important in receptor activity. For example, TM3 and TM6 of the β2AR form ionic interactions which lock the receptor in an inactive state, and conformational changes within these helices govern G-protein coupling at the cytoplasmic face of the receptor (Ballesteros et al. 2001, Rasmussen et al. 2007). Furthermore, disruption of a salt-bridge in β2-AR allows lateral displacement of TM3 away from TM4 and TM5, facilitating β-arrestin binding (Shukla et al. 2014), and it was therefore hypothesised that the CASR Arg680Gly mutation may break a salt-bridge between TM3 and ECL2 or TM7, allowing β-arrestin to bind more readily at the CASR cytoplasmic face (Gorvin et al. 2018a). The importance of the Arg680, Glu767 and Glu837 residues in the CASR has previously been recognised. Mutation of Glu767 and Glu837 has been shown to increase signalling by the CASR (Hu et al. 2002, 2005, 2006, Uckun-Kitapci et al. 2005), and previous structural homology models that investigated the binding of allosteric modulator drugs with the CASR demonstrated that the three residues form critical contacts for drug binding (Miedlich et al. 2004, Petrel et al. 2004), indicating their importance in CASR structure-function. To investigate the hypothesis that a salt-bridge is disrupted by mutation of Arg680, a series of mutations were engineered at the Arg680, Glu767 and Glu837 residues. First, Glu767 and Glu837 were mutated to Arg residues, to introduce unfavourable electrostatic interactions. Under these conditions, the Glu767Arg engineered mutant was shown to enhance MAPK activity in a similar way to the Arg680Gly ADH1 mutant, a response that was also β-arrestin sensitive (Gorvin et al. 2018a). In contrast, responses in cells expressing the CASR Glu837Arg engineered mutant were similar to those in cells expressing WT CASR (Gorvin et al. 2018a). To confirm that a salt-bridge is required between residues 680 and 767 in the CASR, a double mutant was generated by mutating the Glu767Arg mutant receptor with an additional missense mutation of Arg680 to Glu680. This should allow the salt-bridge between the two residues to reform. Indeed, in this double mutant (Glu680-Arg767) CASR, MAPK signalling was restored to levels similar to those seen in WT cells (Gorvin et al. 2018a).

Thus, these studies have revealed some of the important structural motifs of CASR that mediate signalling bias, and discovery of this novel β-arrestin-specific pathway may help facilitate the development of targeted therapeutics for CASR. Furthermore, it demonstrates the importance of investigating multiple signalling outputs downstream of CASR, to ensure that potentially disease-causing mutations are not classified as benign polymorphisms (Gorvin et al. 2018a).

Insights from human genetic mutations in components of the CASR signalling pathway

Both FHH and ADH are genetically heterozygous conditions with mutations in the CASR accounting for approximately 65% and 70% of cases, respectively (Hannan et al. 2012, Nesbit et al. 2013a). Two further
genetically distinct forms of FHH, and one further distinct form of ADH, have been described (Mannstadt et al. 2013, Nesbit et al. 2013a,b). Heterozygous loss- and gain-of-function germline mutations in the α-subunit of G-protein 11 (Gαq), a component of the CASR signalling pathway, give rise to FHH type-2 (FFH2) and ADH type-2 (ADH2), respectively (Mannstadt et al. 2013, Nesbit et al. 2013a). Only four FFH2 and six ADH2 mutations have been described to date, and therefore these mutations account for only a small number of cases of these disorders (Mannstadt et al. 2013, Nesbit et al. 2013a, Li et al. 2014, Gorvin et al. 2016, 2017a, Piret et al. 2016). FHH type-3 (FHH3) is due to mutations in the sigma subunit of the adaptor protein-2 (AP2α), which plays a fundamental role in clathrin-mediated endocytosis of transmembrane proteins, such as GPCRs (Nesbit et al. 2013b). The AP2α protein is ubiquitously expressed, and clathrin-mediated endocytosis is a critical cellular process; however, FHH is a largely benign condition, and the phenotypes observed in FHH3 patients are largely CASR-specific. Thus, by studying the AP2α mutations identified in FHH3, which have been described in a single residue (Nesbit et al. 2013b, Hannan et al. 2015), novel insights into the trafficking and signalling mechanisms of CASR have been elucidated, and indicate an important interplay between these two processes (Gorvin et al. 2018b).

**Trafficking of the CASR**

The cell surface expression of the CASR is important for detecting extracellular ligand and signalling by the receptor, and therefore the plasma membrane expression of CASR is carefully regulated. This involves multiple pathways including: receptor synthesis and secretion, trafficking to the plasma membrane and removal of the CASR from the cell surface by endocytosis.

**Regulation of cell surface expression by functional desensitisation**

In contrast to most GPCRs, which undergo agonist-dependent desensitisation by phosphorylation and/or by β-arrestin proteins, functional desensitisation appears to have only a minimal impact on CASR expression, and there are inconsistencies between findings in these studies (Bouschet et al. 2005, Pi et al. 2005, Lorenz et al. 2007, Thomsen et al. 2012). Initial studies of the CASR showed that several proteins desensitise the receptor. These include: PKC, which phosphorylates the receptor following activation of signalling pathways and recruits β-arrestin; G-protein regulatory kinase (GRK) 2, which binds to Gαq and inhibits its signalling; and GRK4, which phosphorylates CASR to facilitate desensitisation (Pi et al. 2005, Lorenz et al. 2007). Furthermore, GRKs and β-arrestin have independent functions as combined treatment of cells with these proteins enhances desensitisation (Lorenz et al. 2007). The traditional view of GPCR desensitisation and internalisation was one in which the receptor is phosphorylated and β-arrestin recruited, followed by internalisation of the receptor by clathrin-mediated endocytosis, which is facilitated by interactions between β-arrestin and AP2 (Shukla et al. 2011). While studies of CASR showed that β-arrestin can be recruited, this did not enhance receptor internalisation, which only occurred when cells were treated with high concentrations of Ca²⁺ (10mM) (Lorenz et al. 2007). β-Arrestin is also now recognised to have another function, as a scaffold protein that facilitates signalling either at the plasma membrane or for some GPCRs, at the endosome (Shukla et al. 2014). This function of β-arrestin has been recognised for CASR in some studies, providing seemingly contradictory information to that in studies of functional desensitisation. Thus, treatment of cells with dominant-negative forms of β-arrestin1 or β-arrestin2, or with siRNA targeting β-arrestin1 or β-arrestin2, reduces the pERK and membrane ruffling signals downstream of CASR (Bouschet et al. 2005, Thomsen et al. 2012, Gorvin et al. 2018b). Further studies are required to determine whether the discrepancies within these data sets are due to experimental differences, differences in cell type or if both desensitisation and enhanced signalling occur downstream of CASR, but at different spatial or temporal points.

**Regulation of cell surface expression by the secretory pathway and agonist-driven insertional signalling (ADIS)**

The cell surface expression of CASR and mechanisms by which the receptor is synthesised and exported to the plasma membrane are generally well understood. The CASR is synthesised at the endoplasmic reticulum (ER) where it forms homodimers (Pidasheva et al. 2006) and undergoes quality control and immature glycosylation steps prior to progression to the Golgi (Fan et al. 1997). At the Golgi, the CASR undergoes further maturation. This forward trafficking through the secretory pathway involves a number of regulatory proteins that have been well described in previous reviews (Huang et al. 2011, Breitwieser 2013). The core-glycosylated CASR is then retained within pre-plasma membrane compartments (Fan et al. 1997).
Therefore, CASR is retained intracellularly within two large intracellular reserves: one at the ER and one at pre-plasma membrane compartments (Breitwieser 2012).

The large pool of fully mature CASR plays a unique role in regulation of \( \text{Ca}^{2+} \). Unlike many GPCRs, the CASR is chronically exposed to its ligand at baseline concentrations, and thus the receptor undergoes very little functional desensitisation (Gama & Breitwieser 1998). Furthermore, CASR is able to elicit signalling responses for as long as elevated \( \text{Ca}^{2+} \) is available (Pi et al. 2005, Lorenz et al. 2007, Grant et al. 2011). Experimental studies have sought to explain this apparent paradox and using a combination of total internal reflection fluorescence microscopy (TIRFm) and a construct that allows the simultaneous measurement of CASR insertion within plasma membranes and endocytosis (known as BSEP-CASR), a new model for CASR cell surface expression has emerged (Fig. 4). This model proposes that in the basal state, CASR is only weakly expressed at the plasma membrane, but on exposure to increases in \( \text{Ca}^{2+} \), there is an increase in anterograde trafficking through the secretory pathway (Grant et al. 2011) (Fig. 4). This increase in the secretory pathway involves both mature CASR located in pre-plasma membrane regions, and newly synthesised CASR from the ER (Grant et al. 2011, Breitwieser 2013), and at present, the mechanisms that trigger the ADIS events are not fully elucidated. However, studies in which ADIS and CASR signalling were measured simultaneously revealed that elevated \( \text{Ca}^{2+} \), and the \( \text{G}_{q/11} \) pathway are likely involved (Grant et al. 2012, Gorvin et al. 2018b). Furthermore, the 14-3-3 proteins, which bind CASR at an Arg-rich site within the C-terminus, limit the release of CASR from the ER, and therefore regulate CASR cell surface expression by reducing the ADIS-mobilisable pool of the receptor (Grant et al. 2015). The presence of the ADIS system and constitutive endocytosis may also explain some of the inconsistencies between studies of CASR trafficking, as endocytosis of the receptor is balanced by the continual insertion of new CASR from the secretory pathway (Breitwieser 2013). Those studies in which a tagged construct and cell surface labelling (e.g. FLAG) was used in isolation, without also measuring insertion of new CASR within the plasma membrane, may not reflect the physiological state. Similarly, measurement of total cell surface expression of receptor cannot be used as a surrogate for determining the rate of internalisation.

These studies have demonstrated that trafficking of the CASR plays a critical role in receptor signalling and calcium homeostasis. More recent studies, building upon these initial findings, have identified further insights

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**Figure 4**

Mechanisms by which calcium-sensing receptor (CASR) expression is regulated at plasma membranes. Schematic showing trafficking pathways regulating the expression of CASR at the plasma membrane. The CASR is continuously synthesised at the endoplasmic reticulum (ER) and undergoes post-translational modifications at the ER and Golgi, before its export to the plasma membrane surface. In addition to this constitutive anterograde trafficking of the CASR, an additional export pathway for the CASR has been described. This pathway is activated by high [\( \text{Ca}^{2+} \)], and has been named the agonist-driven insertion signalling, or ADIS, pathway. Removal of the CASR from the cell surface is regulated by clathrin-mediated endocytosis that requires the heterotetrameric adaptor protein-2 (AP2). AP2 binds directly to transmembrane proteins using its \( \mu \)-subunit or \( \alpha \)-subunit. The CASR has a putative dileucine motif within its C-terminus with which it is predicted to bind to AP2\( \alpha \). The \( \alpha \)-subunit and \( \beta \)-subunit have large appendages that are important for binding to clathrin coat proteins, plasma membrane phospholipids and endocytic accessory proteins. Once internalised within clathrin-coated vesicles, CASR is targeted to the endosomal-lysosomal system. Experimental evidence suggests that CASR is degraded at the lysosome, and very little, if any, recycling of the receptor occurs.
into CASR cell surface expression and endocytosis, and recognised further connections between the trafficking and signalling of this receptor.

**Regulation of cell surface expression by endocytosis**

The CASR has been described to have two types of internalisation: a constitutive pathway (Grant et al. 2011), and an agonist-driven pathway (Lorenz et al. 2007, Gorvin et al. 2018b). The CASR was shown to internalise using clathrin-mediated endocytosis in early studies (Holstein et al. 2004). In addition, CASR has been described to associate with several proteins that facilitate clustering at the plasma membrane and therefore increase the efficiency of internalisation, or act as scaffolds to enable signalling to occur. One such protein is caveolin-1, with which CASR has been shown to coimmunoprecipitate (Kifor et al. 1998). CASR has been described to be enriched in caveoleae structures in parathyroid chief cells and osteosarcoma cells (Kifor et al. 1998, Jung et al. 2005, Sun & Murphy 2010). CASR-mediated signalling is impaired in cells treated with caveolin-1 targeting siRNA, and it is likely that this signalling function is facilitated by caveoleae acting as signalling hubs, allowing the CASR, G-proteins and PKC to cluster together (Kifor et al. 1998, Jung et al. 2005, Sun & Murphy 2010). However, it is currently unknown whether caveolin facilitates CASR clustering within caveoleae structures to enhance signalling, or whether the signalling itself drives this clustering to encourage endocytosis (Breitwieser 2013). Filamin is an actin-binding cytoskeleton protein that is important for protein scaffolding (Hjalm et al. 2001) and may also facilitate clustering that aids in receptor endocytosis. Filamin binds to the CASR C-terminus and increases total cellular content of CASR by preventing its proteosomal degradation (Hjalm et al. 2001), with some studies showing CASR expression is reduced in cells treated with siRNA targeted against filamin. However, other reports show no such changes in CASR expression, and this likely requires further investigation (Huang et al. 2006, Mingione et al. 2017). The net result of filamin A binding to CASR is increased MAPK signalling by the receptor (Hjalm et al. 2001, Pi et al. 2002).

**Effect of AP2σ mutations on CASR signalling and trafficking**

Mutations in AP2σ have been demonstrated to cause FHH3, and are associated with impaired CASR-mediated Ca^{2+}_i signalling (Nesbit et al. 2013b). The AP2 complex is a ubiquitously expressed heterotetrameric protein that plays a fundamental role in the clathrin-mediated endocytosis of transmembrane proteins, such as GPCRs. The two larger subunits, α and β, have appendages that bind to the clathrin coat proteins, plasma membrane phospholipids and endocytic accessory proteins (e.g. β-arrestin); while the two smaller subunits, μ and σ, bind to endocytic motifs of cargo proteins (Collins et al. 2002, Kelly et al. 2008, Jackson et al. 2010, Kirchhausen et al. 2014) (Fig. 4). The AP2µ subunit recognises tyrosine-based motifs and the AP2σ subunit recognises dileucine-based motifs (Haucke & De Camilli 1999, Kelly et al. 2008) (Fig. 4). FHH3-associated mutations in the AP2σ protein have been reported to affect the Arg15 residue, and structural modelling studies using a published structure of the AP2 complex have shown these missense mutations (to Cys15, His15 and Leu15) likely disrupt interactions with a putative dileucine motif in the CASR C-terminus (Nesbit et al. 2013b), and thus affect CASR endocytosis.

Initially, this hypothesis was tested using an ELISA assay of total CASR at cell surfaces in HEK293 cells stably overexpressing CASR (HEK-CASR), and transiently transfected with AP2σ-mutant proteins. This showed that AP2σ-mutant expressing cells had increased CASR cell surface expression when compared to AP2σ-WT cells following stimulation with 5 mM Ca^{2+}_i (Nesbit et al. 2013b). Thus, it was concluded that CASR endocytosis is impaired in these cells, resulting in increased total CASR cell surface expression (Nesbit et al. 2013b). More detailed studies characterising the ADIS and endocytosis components of CASR regulation in AP2σ mutant cells using TIRFm and the BSEP-CASR construct showed both ADIS and CASR endocytosis were impaired, resulting in the net effect of an increased total CASR cell surface expression in cells expressing AP2σ mutant protein compared to WT cells (Gorvin et al. 2018b). Additional TIRFm studies focussing on CASR and clathrin demonstrated that CASR and clathrin colocalise at plasma membranes, and that the duration of this colocalisation is prolonged in AP2σ-mutant expressing cells (Gorvin et al. 2018b). Furthermore, the vesicles containing both CASR and clathrin in AP2σ-mutant cells, when compared to WT cells, were less motile, which is an indication that the vesicles are less likely to result in viable endocytic events (Rapportor & Simon 2003, Gorvin et al. 2018b). Thus, cells expressing AP2σ-mutant proteins have delayed recruitment of clathrin, and colocalisation with CASR is prolonged, resulting in impaired CASR endocytosis (Gorvin et al. 2018b).

The effect of the FHH3-associated AP2σ-mutant proteins on multiple CASR-mediated signalling pathways has been characterised in a number of cellular assays
in HEK293 cells stable overexpressing AP2σ WT and mutant proteins and lymphoblastoid cell lines derived from blood samples from FHH3 patients with the AP2σ-Cys15 mutation (Nesbit et al. 2013b, Gorvin et al. 2018b). These studies confirmed that AP2σ mutations of the Arg15 residue impair Ca\(^{2+}\), mobilisation, phosphorylated ERK1/2 (pERK1/2) MAPK signalling, membrane ruffling and suppression of cAMP, and that all of these pathways occur downstream of G\(_{\alpha q/11}\) and G\(_{\alpha o}\) (Nesbit et al. 2013b, Gorvin et al. 2018b). Therefore, these studies have shown that AP2σ mutations reduce endocytosis, resulting in increased CASR cell surface expression, but paradoxically decrease CASR-mediated signalling (Gorvin et al. 2018b). To explain this paradox, we hypothesised that CASR may be able to continue signalling from within the cell (i.e. sustained signalling) (Gorvin et al. 2018b). Such sustained signalling has been previously reported for some class A (e.g. β2-adrenergic receptor (β2AR), dopamine receptor D1 (DRD1), thyroid-stimulating hormone receptor, vasopressin receptor 2 (V2R) and luteinizing hormone receptor (LHR)), and class B (e.g. parathyroid hormone 1 receptor (PTH1R)) GPCRs (Calebiro et al. 2009, Ferrandon et al. 2009, Kotowski et al. 2011, Feinstein et al. 2013, Irannejad et al. 2013, Jean-Alphonse et al. 2014). Thus, in cells with the AP2σ mutation, in which there is impaired endocytosis, the availability of internalised receptors from which sustained signals could emanate would be reduced and thus the net effect would be impaired overall CASR-mediated signalling. To test this hypothesis, a combination of imaging and biochemical analyses, along with chemical inhibitors, were used in HEK-CASR, HEK-AP2σ and CRISPR-Cas generated β-arrestin knockout cells to assess CASR-mediated MAPK signalling (Gorvin et al. 2018b). To assess sustained signalling, two primary assays were used: (1) assessment of pERK1/2 over 60 min following treatment of cells with a 5 min pulse of 5 mM Ca\(^{2+}\); and (2) analysis of SRE luciferase reporter responses over 12 h, following a 5 min pulse of 5 mM Ca\(^{2+}\). Using these methods, MAPK-sustained signals were demonstrated in HEK-CASR cells, and evidence for an internal, likely endosomal, source was shown in three ways (Gorvin et al. 2018b) (Fig. 5). First, addition of Dyngo, a chemical inhibitor of dynamin, which is required for vesicle scission during clathrin-mediated endocytosis, abolishes sustained signals, whilst rapid plasma membrane-mediated endocytosis, abolishes sustained signals, and therefore slows clathrin-mediated endocytosis. Third, reduction or loss of these sustained signals was observed in cells expressing AP2σ mutant proteins (Fig. 5).

Figure 5
The calcium-sensing receptor (CASR) can activate plasma membrane and endosomal signalling pathways. Schematic diagram showing the signalling pathways that occur downstream of the CASR. As described in Fig. 2, CASR predominantly signals via: the G\(_{\alpha q}\) pathway, leading to activation of the intracellular calcium (Ca\(^{2+}\)) and mitogen-activated protein kinase (MAPK) pathways; and the G\(_{\alpha o}\) pathway, leading to activation of MAPK and reductions in the cAMP pathways. These pathways are activated rapidly (within 2–5 min), and originate at the plasma membrane. The CASR can also signal from an internal location using a G\(_{\alpha q}\) pathway. This sustained signal that occurs later than the plasma membrane signal (from 30 min) is sensitive to: global clathrin-mediated endocytosis blockade (shown experimentally using Dyngo); inhibition of CASR-mediated internalisation (demonstrated in cells expressing FHH3-associated AP2σ mutants); and maturation of internalised vesicles to the early endosome (demonstrated experimentally using a dominant-negative (DN) form of Rab5). Thus, this sustained signal likely arises within endosomes.
The G-proteins involved in these sustained signals were also explored as MAPK signalling is a convergence pathway for multiple CASR-mediated signalling pathways (Fig. 2). The ability of the Gαq/11 and Gαi/o pathways to activate CASR sustained signals was investigated using G-protein-specific inhibitors (UBO-QIC and PTx) and the SRE luciferase reporter assay, which showed that inhibition of both Gαq/11 and Gαi/o impaired the early, plasma-membrane-mediated CASR signal, while only Gαq/11 was important for the later sustained signal. Furthermore, confocal microscopy confirmed that Gαq and phosphatidylinositol 4,5-bisphosphate (PIP2), the lipid hydrolysed by PLC, colocalise with CASR at plasma membranes and a subpopulation of endosomes (Gorvin et al. 2018b). In addition, inhibitors of the PLC-DAG-IP3 pathway (U73122, GF-109203X and 2-aminoethoxydiphenyl borate (2-APB), which inhibit PLC, PKC and the IP3-Receptor, respectively) were shown to reduce sustained pERK1/2 signals, indicating that Gαq/11 and its signalling pathway are important for CASR-mediated signalling (Gorvin et al. 2018b). Finally, the effects of the β-arrestin scaffold proteins, which are important for the sustained endosomal signalling of some GPCRs such as V2R and PTH1R (Feinstein et al. 2013, Wehbi et al. 2013), were assessed in HEK-CASR cells with deletion of the β-arrestin1 and β-arrestin2 genes. In these cells, both pERK and SRE reporter responses were unaffected by deletion of the β-arrestin proteins (Gorvin et al. 2018b). Thus, the CASR mediates some MAPK signals from endosomes using Gαq and PLC, but does not require β-arrestin for this pathway.

Conclusions and important lessons for the future (or unanswered questions)

Since the cloning of the CASR gene 25 years ago (Brown et al. 1993), many insights have been gained into the role of the CASR in calcium homeostasis, and by studying patients with mutations within this gene, we have learnt much about CASR signalling and trafficking mechanisms. The discovery that CASR can signal from within the cell, most likely from an endosomal source, and that disruption of a specific structural motif can mediate G-protein independent signalling bias, opens up many new avenues of investigation. How do mutations in the CASR affect these endosomal pathways? Can drugs that target the CASR affect these pathways? How are trafficking and signalling of the CASR so intimately linked? Future studies investigating these questions will undoubtedly reveal further complexities into the regulation of CASR, and could provide mechanisms relevant to other GPCRs.

Declaration of interest

The author declares that there is no conflict of interest that could be perceived as prejudicing the impartiality of this review.

Funding

This research did not receive any specific grant from any funding agency in the public, commercial or not-for-profit sector.

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Received in final form 16 March 2018
Accepted 29 March 2018
Accepted Preprint published online 29 March 2018

http://jme.endocrinology-journals.org
Published by Bioscientifica Ltd.
Printed in Great Britain