Identification and Characterization of Novel Mutations in the Human Gene Encoding the Catalytic Subunit Calpha of Protein Kinase A (PKA)

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Abstract

The genes PRKACA and PRKACB encode the principal catalytic (C) subunits of protein kinase A (PKA) Cα and Cβ, respectively. Cα is expressed in all eukaryotic tissues examined and studies of Cα knockout mice demonstrate a crucial role for Cα in normal physiology. We have sequenced exon 2 through 10 of PRKACA from the genome of 498 Norwegian donors and extracted information about PRKACA mutations from public databases. We identified four interesting nonsynonymous point mutations, Arg45Gln, Ser109Pro, Gly186Val, and Ser263Cys, in the Cα splice variant of the kinase. Cα variants harboring the different amino acid mutations were analyzed for kinase activity and regulatory (R) subunit binding. Whereas mutation of residues 45 and 263 did not alter catalytic activity or R subunit binding, mutation of Ser109 significantly reduced kinase activity but not type II holoenzyme formation. Gly186 is located in the highly conserved DFG motif of Cα and mutation of this residue to Val was predicted to result in loss of binding of ATP and Mg2+, which may explain the kinetic inactivity. We hypothesize that individuals born with mutations of Ser109 or Gly186 may be faced with abnormal development and possibly severe disease.

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Introduction

Protein Kinase A (PKA) is a cyclic AMP (cAMP)-dependent Ser/Thr kinase involved in regulating a multitude of biological processes including cell growth and division, cell differentiation, as well as metabolism and immune responsiveness [1]. In its inactive state, PKA exists as a tetrameric holoenzyme consisting of two regulatory (R) and two catalytic (C) subunits. Four different genes, PRKAR1A, PRKAR1B, PRKAR2A, and PRKAR2B, encode the R subunit proteins R1α, R1β, R2α, and R2β, respectively, with several splice variants of the R1α and R1β subunits [2,3]. Two principal genes, PRKACA and PRKACB, encode the C subunits Cα and Cβ, respectively. Also, PRKX and the retroprospons PRKY and PRKACg are identified as PKA C subunit genes. While no protein products for PRKY and PRKACg have been identified, PRKY is translated into a protein kinase which is inhibited by the R subunit in a cAMP-sensitive manner [4–7]. Both the PRKACA and PRKACB genes encode several protein products. Alternative use of two exons upstream of exon 2 in the PRKACA gene gives rise to two Cα variants with different N-termini. These proteins are designated Cα1 and Cα2 [8–11]. In the case of the PRKACB gene, several exons 5′ of exon 2 are encoding a number of Cβ splice variants designated Cβ1, Cβ2, Cβ3, and Cβ4 [12,13], as well as a number of Cβ3 and Cβ4 forms that contain N-terminal inserts from exons designated a, b, and c [14,15]. In the brain and nervous tissues of higher primates, a range of C variants lacking exon 4 encoded sequences are also identified [16].

The two major groups of the R subunits R1 and RII form two types of the PKA holoenzymes termed PKA type I and PKA type II, respectively. The C subunits are the catalytically active components, and they become activated after dissociation from the R subunits in a cAMP-dependent manner. For specificity in the cAMP-PKA signaling pathway, the PKA holoenzymes can be located to specific subcellular structures via A Kinase Anchoring Proteins (AKAPs). AKAPs typically bind RI subunits with high affinity [8,17–19]. In contrast, PKA type I holoenzymes tend to locate to the soluble fraction of the cell but can bind to dual- and RI-specific AKAPs [19]. Both the C and R subunits are differentially expressed in various tissues. Whereas Cα1 is ubiquitously expressed, Cα2 is exclusively found in sperm cells [8–10]. For the Cβ variants, Cβ1 is ubiquitously expressed, while...
the other Cβ variants are more specifically expressed in certain tissues, such as lymphoid and neuronal tissues [14,15].

Cα1 is the principal source of PKA activity in most tissues [11] and was the first protein kinase to be subjected to crystalization and 3D structure determination [20]. The Cα1 structure reveals a large C terminal lobe (C-lobe or large lobe) and a smaller N terminal lobe (N-lobe or small lobe). The large lobe is mainly composed of α-helices, and is involved in R subunit and substrate binding. It contains a number of residues involved in catalysis. The small lobe is mostly composed of β-strands and contains an ATP binding site. Between the large and small lobe is the active site cleft. The small and large lobes, as well as the active site, are known as the catalytic core and are encoded by residues 40–300 in PKA Cα1. The amino acid sequence and 3D structure of the catalytic core are conserved among all protein kinases. The catalytic core may, in addition to ATP, bind two Mg²⁺ ions that are critical for catalysis [20]. The two Mg²⁺ ions are designated activating and inhibitory Mg²⁺, or Mg1 and Mg2, respectively [21]. The binding affinity for Mg1 is higher than for Mg2 in the presence of ATP, while binding of Mg2 is believed to stabilize the protein, but also to inhibit catalysis [21–24]. Mg²⁺ is also essential for high-affinity binding to inhibitors PKI and RIα subunits [25].

For catalytic activity, Cα has to be phosphorylated on residue Thr197, located in the activation loop of the large lobe. This phosphorylation affects the conformation of the conserved DFG (Asp184-Phe-Gly) motif in the active site [26]. In the active conformation, Asp184 of the DFG motif coordinates the three phosphates of ATP via the Mg²⁺ ions, positioning the γ-phosphate of ATP optimally for catalysis. This central role of Asp184 in catalysis depends on a conserved hydrogen bond between Asp184 and Gly186 backbone amide group. This interaction orients Asp184 perfectly for coordination of the Mg²⁺ ions and efficient ATP binding [27]. The Asp184-Gly186 hydrogen bond is only established after phosphorylation of Thr197, which causes a twist of the peptide bond between Phe185 and Gly186. In this way phosphorylation of Thr197 serves as a regulatory mechanism for the kinase to become catalytically active. The C-spine is completed to be phosphorylated. Evaluation of the C- and R-spines is a regulatory (R-) spine that has to be established in order for the kinase to be active. The R-spine is composed of two hydrophobic structures called the Catalytic (C-) and Regulatory (R-) spines that have to be established in order for the kinase to become catalytically active. The C-spine is completed when the adenine nucleobase of ATP interacts with the active site. For the R-spine to be established, Thr197 in the activation loop has to be phosphorylated. Evaluation of the C- and R-spines is a helpful way of predicting whether a kinase has a catalytically active or inactive conformation.

Despite its central role in a multitude of processes in the body, few diseases have been linked to defects or alterations in the PKA subunit genes or proteins. Germline mutations leading to premature stop codons in PRKAR1A have been identified in patients with Carney complex, a multiple neoplasia syndrome with skin pigmentation, cardiac and other myxomas, endocrine tumors, and piasomatosus melanotic schwannomas [28,29]. To the best of our knowledge, no diseases have been linked to defects in any of the C subunit genes. Nevertheless, studies on mice that are homozygote knockout (KO) for the PRKAC1 gene reveal a severe phenotype. The majority of the Cα−/− mice die at birth or during the early postnatal period [30]. The offspring that survive beyond 2 months (<11%, [31]), all show reduced growth, and the few males, but not females, that reach puberty are all 100% infertile [30,32]. Furthermore, mice with reduced PKA C gene transcription by only expressing one functional C subunit allele of Cα or Cβ, show reduced PKA activity and neural tube defects [33].

Due to the severe phenotype of Cα KO mice we searched for mutations in the human PRKACA gene by genomic sequencing of 498 subjects. We identified two nonsynonymous point mutations in the PRKACA gene that result in amino acid switches in the Cα1 protein at residues 45 and 109. In addition, we searched for previously described nonsynonymous mutations in various human genomic DNA databases, and selected two of these, giving amino acid switches at residues 186 and 263, for further studies. These four mutations were introduced to Cα1- and Cα2-encoding plasmids and the proteins were expressed and analyzed with respect to kinase activities and R subunit binding in vitro and in vivo. Mutation of residues 109 and 186 were associated with significantly reduced and totally abrogated kinase activity, respectively. In addition, mutation of residue 186 rendered Cα incapable of forming PKA type I holoenzymes.

Materials and Methods

Analysis of Mutations

The PRKACA gene from 498 Norwegian donors deriving from three different control groups [34,35] was sequenced and analyzed for point mutations by Lark Technologies (Takeley, UK). Leukocyte DNA was isolated from thawed blood containing ethylenediaminetetraacetic acid. Using the Applied Biosystems 340A Nucleic Acid Extractor, DNA was extracted with chloroform/phenol followed by ethanol precipitation. Exons 2–10 of PRKACA were sequenced using the primers listed in table 1.

A search for mutations in the PRKACA gene was also performed in the following public databases: NCBI dbSNP [36,37], Ensembl [38], and SNPper [39].

Generation of Plasmids

The pDONR201-Cα1WT vector has previously been described [40] and was used to make plasmids containing Cα1mut using QuickChange® II Site-Directed Mutagenesis Kit (Stratagene, cat. no. 200524) according to the manufacturer’s protocol. The pDONR201-Cα1 variants were transferred to the eukaryotic

Table 1. Primers used for sequencing of PRKACA exons 2 to 10.

| Exons | Forward | Reverse |
|-------|---------|---------|
| 2     | 5′-AGGCCTTGGTGGTGAAACTGC-3′ | 5′-CTGCACTGGTGGTGAAACTGC-3′ |
| 3     | 5′-GTGGCTAAGTTGCTGCTCTCT-3′ | 5′-TGGGAAACTCCATCTCTAC-3′ |
| 4     | 5′-CCACGCGCTTGACCTCTCTGTG-3′ | 5′-TGGGAAACTCCATCTCTAC-3′ |
| 5     | 5′-CTCACGCTCCAGGACACGCCAG-3′ | 5′-TGGGAAACTCCATCTCTAC-3′ |
| 6-8   | 5′-ATGACGACGACATGTTGGAG-3′ | 5′-ATGACGACGACATGTTGGAG-3′ |
| 6-8   | 5′-TTTGACGCTGGTGGCTCTCTC-3′ | 5′-TTTGACGCTGGTGGCTCTCTC-3′ |
| 9     | 5′-ACGTTAGTAGAGGACCTTCTTCTC-3′ | 5′-ACGTTAGTAGAGGACCTTCTTCTC-3′ |
| 10    | 5′-CGGAACTCTTAAATGAGAACAACCTC-3′ | 5′-CGGAACTCTTAAATGAGAACAACCTC-3′ |
| 11    | 5′-ACGTTAGTAGAGGACCTTCTTCTC-3′ | 5′-ACGTTAGTAGAGGACCTTCTTCTC-3′ |

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expression vector pEF-DEST51 by the LR recombination reaction (Invitrogen, cat. no. 11791–019) according to the manufacturer’s protocol.

Cα1WT and Cα2WT prokaryotic expression vectors are previously described [41]. Using QuickChange™ Site-Directed Mutagenesis Kit (Stratagene, cat. no. 200518), Cα2WT was mutated into Cα2Gly186Val.

Cα1 was cloned into the mammalian expression vector pGFP-C3 (Perkin Elmer) as previously described [42], creating the BRET sensor construct GFP-Cα1 as previously described [42].

Directed Mutagenesis Kit, GFP-Cα2WT was mutated into GFP-Cα2Gly186Val according to the manufacturer’s protocol. RIα-Rluc and RIβ-Rluc constructs have previously been described [42].

For all mutagenesis reactions, the PCR reaction mixture was initially heated to 95°C for 30 s, followed by 16 cycles of 95°C for 30 s, 55°C for 1 min and 68°C for 7 min, before final elongation at 68°C for 2 min. Primers are listed in Table 2. All plasmids were verified by sequencing (Eurofins MWG Operon).

Purification of Proteins

Cα1WT and Cα2WT were purified by affinity chromatography using PKI-peptide Affi-Gel, as previously described [41]. Cα2Gly186Val was purified using a modified method described by Hemmer et al [43]. Briefly, B.1.1/DE3 cells transformed with either Cα2Gly186Val or His-tagged RIα-G337Glu (provided by Antje Badel, University of Kassel, Germany) were cultured and protein expression induced with IPTG. After centrifugation, the bacterial pellets were resuspended and lysed, and the lysates were mixed in equimolar amounts forming PKA holoenzymes. The holoenzymes were then coupled to a Ni²⁺-resin binding the His-tagged R subunits. Following washing with 50 mM Na₂HPO₄ (pH 8.0), 5 mM β-mercaptoethanol, and 25 mM KCl, Cα2Gly186Val was eluted with the same buffer supplemented with 10 mM cAMP.

Cell Cultures and Transfection Methods

HEK 293T cells (ATCC, cat. no. CRL-11268) were cultured at 37°C in humidified air with 5% CO₂ in RPMI 1640 (Sigma-Aldrich, cat. no. P4458). The 293T cells were transfected with plasmid DNA (0.2 μg per well) using PEI (Polysciences, 25 kDa linear polyethyleneimine) [44]. Cells were transfected as previously described [42]. Briefly, 2 × 10⁴ COS-7 cells were seeded in each well of a 96-well microplate (Nunclon Surface, cat. no. 136101). After 24 h, cells were transfected with plasmid DNA (0.2 μg per well) using PEI.

Phosphotransferase Assay

20 h post transfection, the HEK 293T cells were harvested and washed 3 x in phosphate buffered saline, then lysed in a potassium-phosphate buffer (5 mM K₂HPO₄, 1 mM EDTA, 250 mM sucrose and 0.5% Triton X-100, 1 mM phenylmethylsulfonyl fluoride, 1 mM Na₃VO₄ and protease inhibitor cocktail (Sigma-Aldrich, cat. no. P8340)) by vortexing and 20 min on ice. Debris was pelleted by centrifugation at 16 000 g for 15 min at 4°C. After Bradford protein determination (Bio-Rad, cat. no. 500–0006), all samples were adjusted to equal concentrations. Phosphotransferase activities of lysates from Cα1WT, Cα2M13Gly and mock transfected cells were determined by the phosphotransferase assay described by Witt and Roskoski [45,46]. Briefly, phosphotransferase activity against the PKA-specific substrate Kempptide (Leu-Arg-Arg-Ala-Ser-Leu-Gly, Sigma-Aldrich) was measured in a reaction mixture (14.3 mM Mg-acetate, 145 μM ATP, 7.5 μCi/mL γ-32P-ATP (PerkinElmer), 50 mM Tris-HCl, pH 7.4) in the presence or absence of cAMP or PKI. After incubation at 30°C for 9 min, the reactions were stopped by spotting onto p81 phosphocellulose papers followed by 4 × 15 min washing in 75 mM phosphoric acid. The filter papers were washed in 96% ethanol for 10 min and air dried for approximately 1 h. Phosphotransferase activity was measured by liquid scintillation in 3 ml Opti-fluor (Packard BioScience, PerkinElmer). All experiments were repeated at least three times.

BRET Assay

48 h post transfection cells were washed with 1 × PBS and BRET read-out was performed by addition of 5 μM of the luciferase substrate coelenterazine 400 a (Biotrend, Cologne, Germany). After simultaneous detection of RLuc (410 nm) and GFP² (515 nm) emission by a multi-label reader (POLARstar, Omega, BMG Labtech GmbH, Ortenberg, Germany) the ratiometric BRET signal was calculated as followed [emission410 nm - (not transfected) COS-7 cells410 nm]/[emission515 nm - (not transfected) COS-7 cells515 nm] using the GraphPad Prism 4 software (La Jolla, CA, USA). Experiments were performed both in absence and presence of 50 μM forskolin (fsk) and 100 μM 3-

### Table 2. Primers for mutagenesis (mutated nucleotide in bold).

| Desired mutation | Primers |
|------------------|---------|
| Arg45Gln         | Forward: 5'-CTTGGATCAGTTGAACAAATCAAGACCCCGC-3' |
| Arg45Gln         | Reverse: 5'-GGCGGATGCTTTGATTTGGCGAATGCAAGG-3' |
| Ser109Pro        | Forward: 5'-CGTCAAACTGAGTTCCTTGAAGCCACCTC-3' |
| Ser109Pro        | Reverse: 5'-GAGGTGCTTGGTCAAGCGGAAAAGCTGAGG-3' |
| Gly186Val        | Forward: 5'-GTTGACAGACCTCGTTTGCTTCTGGCAAGCCGG-3' |
| Gly186Val        | Reverse: 5'-GGCGCTTGGCAGAAAAGAAAGTGGCTGTAC-3' |
| Ser263Cys        | Forward: 5'-CTTCTTCTCCACTCTTGGAGTCTGTATGAGGAC-3' |
| Ser263Cys        | Reverse: 5'-GCTTCTTCTCCACTCTTGGAGTCTGTATGAGGAC-3' |

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isobutyryl-1-methylxanthine (IBMX) (Sigma-Aldrich). All experiments were repeated at least three times.

**Immunoblotting**

Samples used in phosphotransferase assays were evaluated for C subunit expression. Protein concentrations of each sample within an assay were adjusted to the same levels, as described above. Proteins were separated by SDS-PAGE and transferred to PVDF membranes. The membranes were blocked by drying and rehydrated in methanol before incubation with primary antibody Purified Mouse Anti-PKA [C] [1:250 dilution, BD Transduction Laboratories, cat. no. 610981] for 1 h, followed by washing with TBST. After 30 min incubation with secondary antibody HRP conjugated goat anti-mouse (1:2000 dilution, ICN Biomed, cat. no. 55363) and washing, proteins were detected by enhanced chemiluminescence (Pierce Biotechnology, Rockford, IL, USA) and the Syngene G:BOX imaging system.

Expression of endogenous C and GFP-C constructs in transfected COS-7 cells used for BRET assays were also assessed with immunoblotting. Proteins were detected similarly to the method described above, with the following modifications: blocking was performed with 5% milk powder for 1 h; primary antibody was rabbit anti-Cα (Santa Cruz, PKA cat (c-20) cat. no. SC-903) and secondary antibody anti-rabbit IgG (peroxidase-linked species-specific whole antibody (ECL) NA934, GE Healthcare, Freiburg, Germany).

**Spectrophotometric Kinase Activity Assay**

Kinetic activities of purified CαWT and CαGly186Val were evaluated by the continuous enzyme-linked spectrophotometric method described by Cook et al. [22]. Activities were reported in U/mg, defined as µmolmg⁻¹min⁻¹. All experiments were repeated at least three times.

**Molecular Representation and Simulation**

Selected motifs, including the DFG motif and its interactions with ATP and divalent cations were presented and analyzed with PyMOL [47], using the experimental structure described by Thompson et al. [48] (PDB identifier 3FJQ). Simulated mutagenesis in PyMOL was performed on Gly 186 in the DFG motif. Relevant distances between the Gly 186/Val186 residue and surrounding atoms were calculated.

**Results**

We performed two independent studies of the **PRKAC4** gene: 1) by sequencing genomic DNA from 498 Norwegian subjects and 2) by a bioinformatics analysis of DNA sequences from various populations submitted to publicly available databases (see Material and Methods). By the first approach, exons 2 through 10 of **PRKAC4** were sequenced using exon-specific primers and genomic DNA extracted from leukocytes. In this way we detected five mutations in the **PRKAC4** gene; one in each of exons 2, 3, 4, 5, and 8. Only two of the mutations translated into an amino acid switch. The corresponding nucleotides were located in exons 3 and 4, and are affecting Arg45 and Ser109 in the Cα1 sequence, while the three silent mutations were at Pro353, Gly126, and Pro236 (Table 3, Fig. 1A). With the recent submission of a large amount of new mutations from several thousand individuals by exome and whole genome sequencing projects, four of these five mutations are now also present in dbSNP [36,49]. Our study thus confirms the presence of the mutations referred to as rs56105247 (Arg45), rs78098302 (Gly126), and rs137911238 (Pro236) in the Norwegian population.

By the second approach, we detected another twelve non-synonymous mutations (Table 4), of which the majority has been submitted within the last year. Among the mutations that were known at the initiation of this study, two were selected for further investigations. Firstly, the mutation rs53635531, resulting in a switch of Ser263 to Cys, is the only mutation that has been submitted by three independent projects. Although the frequency of this mutation has not been determined in detail, it is likely to be relatively common in at least some human populations. Secondly, the mutation causing the Gly186Val mutation was considered particularly intriguing since this residue is located in the Mg²⁺ positioning loop close to ATP and Mg²⁺ in the active site cleft of Cα (Fig. 1B) and is part of the highly conserved DFG motif [26,50,51]. The positions of the four investigated mutations resulting in amino acid switches are shown relative to exon encoded sequence in Fig. 1A and in the 3D structure of Cα1 in Fig. 1B.

**Mutation of Residues Ser109 and Gly186 but not Arg45 and Ser263 in Cα1 Influences Catalytic Activity**

We first tested if any of the mutations influenced the catalytic activity of Cα. Site-directed mutagenesis was used to introduce the required amino acid switches Arg45Gln, Ser109Pro, Gly186Val, and Ser263Cys to Cα encoding expression vectors. Wild type CαWT and the four mutated Cα1 products (collectively termed Cα1Mut) were expressed in HEK-293T cells, and 2 h post transfection the cells were lysed. Expression of protein was first assessed by immunoblotting with a pan C antibody, revealing that both Cα1WT (Fig. 1C, lane 1) and Cα1Mut (Fig. 1C, lanes 2 to 5) were expressed at comparable levels and with similar apparent size as endogenous Cα1 (Fig. 1C, lane 6). We further determined their catalytic activity in the presence of the PKA-specific substrate Kemptide and γ³⁻P-ATP (Fig. 1C). In these experiments, lysates from untransfected cells (Fig. 1C, lane 6) were included to assess endogenous kinase activity. This demonstrated high catalytic activity of expressed Cα1WT which was set to 100% (Fig. 1C, lane 1). According to this assay, the catalytic activities of Cα1Arg45Gln and Cα1Ser263Cys were shown not to be significantly different from Cα1WT. On the other hand, the activity of Cα1Ser109Pro was significantly lower compared to Cα1WT (P <0.05), and the activity of Cα1Gly186Val was comparable to background levels (Fig. 1C, lanes 4 and 6). This suggested that mutation of Gly186 rendered Cα inactive and prompted us to focus further on revealing the mechanism for this inactivation. Accordingly, we transfected cells with Cα1Gly186Val and Cα1WT and compared catalytic activity in the absence and presence of cAMP and the PKA-specific inhibitor PKI and compared phosphotransferase activities to mock transfected cells (Fig. 2A). The activity of Cα1WT was reduced to background level in the presence of PKI while activities of Cα1Gly186Val were comparable to background levels independently of stimulation with cAMP or inhibition with PKI (Fig. 2A, Mock compared to Cα1Gly186Val).

To determine if Cα1Gly186Val was completely inactive we next expressed and purified recombinant Cα1WT and Cα1Gly186Val. We used recombinant WT and mutated Cα2 since Cα2 is more easily produced than Cα1 due to higher solubility [41]. Cα2 will therefore give a higher protein yield. Cα1 and Cα2 may also be interchanged in these experiments because studies suggest that they are kinetically indistinguishable [41]. WT and mutated recombinant Cα2 (Cα2WT and Cα2Gly186Val) were produced in BL21(DE3) cells, purified and visualized by SDS-PAGE (Fig. 2B).

(To simplify the nomenclature we refer to the mutations according to their Cα1 numbering also when introduced to Cα2. The actual position of the mutated residues in the Cα2 protein is...
achieved by subtracting the number 8). Cz2_{Gly186Val} was purified by running protein extracts over an RIIC_{Gly126Val} affinity column and Cz2 bound to RIIC was eluted with 10 mM cAMP (see Material and Methods). The purification was evaluated after each step by SDS-PAGE analysis (Fig. 2B, lanes 1 to 14). Purified Cz2_{Gly186Val} and Cz2_{WT} were tested for catalytic activity employing the Cook assay [22] that showed a specific activity for Cz2_{WT} at 20.9 ± 2.1 U/mg (95% confidence interval) (Fig. 2C, Cz2_{WT}). In comparison, catalytic activity by Cz2_{Gly186Val} was undetectable demonstrating that a Gly186 to Val mutation renders the kinase catalytically inactive (95% confidence interval of Cz2_{Gly186Val}: 0.55 ± 0.96 U/mg) (Fig. 2C, Cz2_{Gly186Val}).

**Mutation Gly186Val Alters the Catalytic Core of Cz1**

We next investigated the Gly186Val mutation in a 3D structure model of Cz1 in an attempt to understand the molecular nature of the kinase inactivity (Fig. 3). Figure 3A shows selected conserved motifs surrounding the DFG motif where residue 186 is located. The chain of interactions leading into the DFG motif from phosphorylated Thr197 is depicted. In the active conformation of Cz1_{WT}, the C- and R-spines are assembled and the structure is optimized for catalysis (Fig. 3B). Simulated mutagenesis of Gly186Val revealed three possible rotamers. For representation of the mutations would influence R subunit binding and cAMP activation. However, ATP is required for association between C and RI [33,34] and given the fact that Cz2_{Gly186Val} probably does not bind ATP efficiently, it may be suggested that Cz2_{Gly186Val} will only associate with RI and not RI. To investigate this we determined the binding of Cz1_{Gly186Val} to RI and RI using a Biosilence resonance energy transfer (BRET) assay. In these experiments, holoenzyme formation was investigated for all four Cz2 proteins to test if any of the mutations would influence R subunit binding and cAMP sensitivity. COS-7 cells were cotransfected with either Cz1_{WT} or Cz1_{Mut}, N-terminally coupled to Green Fluorescent Protein (GFP), GFP-Cz1_{WT} and GFP-Cz1_{Mut}, respectively, and RI and RI using Rluc, Renilla luciferase (Rluc), Rluc-Rluc and Rluc-Rluc, respectively. Immunoblotting shows the expression of both GFP-Cz1_{WT} and GFP-Cz1_{Mut} (Fig. 4A, lanes 2 to 6). Figure 4, lane 1 depicts the expression of endogenous C subunit in cells transfected with Rluc alone serving as background. In the case of holoenzyme formation with Rluc subunits, measurements were normalized by setting the BRET signal in holoenzyme with Rluc to 100% (Fig. 4B, WT ‘‘’’). Increasing the intracellular concentration of cAMP with forskolin (fsk) and IBMX (+‘‘’) leads to the dissociation of the holoenzyme complex and the reduction of the BRET signal by approximately 50% as reported previously [55]. The same was true for holoenzymes formed with Rluc-Rluc and GFP-Cz1_{Arg45Gln}, GFP-Cz1_{Ser263Cys} and GFP-Cz1_{Ser263Pro} and GFP-Cz1_{Ser263Cys}, respectively (Fig. 4B, ‘‘+’’). In contrast, the GFP-Cz1_{Gly186Val} showed only residual binding to RI in resting cells (Fig. 4B, Gly186Val ‘‘–’’), and only marginal activation was detected after stimulation with fsk and IBMX (Fig. 4B, Gly186Val ‘‘+’’).

**Table 3. Mutations in the PRKACA gene discovered by sequencing of 498 Norwegian subjects.**

| Exon no. | Nucleotide (nt) no. | Amino acid (aa) | WT/mut nt | WT/mut aa | No. affected individuals | Functional consequence prediction |
|----------|-------------------|----------------|----------|----------|--------------------------|----------------------------------|
| 2        | 102               | Pro33          | C/T      | Silent   | 10                       | Likely neutral                   |
| 3        | 137               | Arg45          | G/A      | Arg/Gln  | 1                        | Arg45 is highly conserved in metazoan Ca2+/Cj homologs – likely functional consequences |
| 4        | 328               | Ser109         | T/C      | Ser/Pro  | 2                        | Ser109 in conserved in Ca2+, but not in Cjβ – effect unknown |
| 5        | 381               | Gly126         | G/A      | Silent   | 3                        | Likely neutral                   |
| 8        | 711               | Pro236         | G/A      | Silent   | 1                        | Likely neutral                   |

Nucleotide enumeration is counted from the PKA Cα1 transcript start codon. Amino acid numbering is given for the mature protein with N-terminal Gly1 corresponding to codon 2. Sequence conservation is illustrated in Supplementary Fig. S1.
Figure 1. Sequences of human wild type PKA Ca1/Ca2, location of point mutations, and activities of mutant proteins. A. The human Ca wild type and mutated amino acid sequence. The N-terminal methionine of Ca1 is removed post-translationally, and glycine residue 1 in mature Ca1 corresponds to codon 2. Alternating black and blue coloring shows the contribution of the eleven exons to the mature translated protein. The codons for Val15 and Ser139 (red) are encoded by nucleotides from neighboring exons. The PRKACA gene encodes two splice variants, Ca1 and Ca2 (UniProt [59] accession number P17612), which differs by alternative use of exons 1–1 or 1–2, respectively. The four point mutations are highlighted and denoted with the amino acid change. The Gly186Val mutation is located in the DFG motif (boxed) [26].

B. The location of the four mutations are indicated (red) in a model of Ca1 based on the PDB structure 3FJQ [48]. Residues are designated according to the WT Ca1 sequence. ATP and two divalent cations are shown in yellow.

C. Immunoblotting and phosphotransferase assays of WT and mutated Ca1 subunits. Ca1 was expressed in HEK 293T cells and immunoreactive C subunit detected with anti-C (Anti-mouse PKA[C], cat. no. 610981) after separation of cell extracts by SDS-PAGE in 10% gels and immunoblotting. Expression of Ca1WT is shown in lane 1 (WT), mutated Ca1 in lanes 2 to 5, and endogenous Ca1 in lane 6 (Mock). Activities are normalized relative to Ca1WT activity. Data represents mean values ± standard deviation (SD) of triplicate experiments (*, P < 0.05. **, P < 0.005).

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WT (‘−’). PKA type II holoenzymes formed with GFP-CαWT and GFP-CαMut dissociate almost completely resulting in a stronger BRET signal reduction (Fig. 4C, ‘+’). Interestingly, whereas GFP-CαGly186Val does not stably interact with RII-Rluc, the holoenzymes formed with RIIz-Rluc shows a comparable magnitude to GFP-CαWT (Fig. 4C, Gly186Val ‘−’ and WT ‘−’).

Discussion

Here we have examined the human PKA\(_\alpha\) gene for mutations by sequencing genomic DNA from 498 Norwegian individuals and by searching for earlier reported mutations in publicly available databases. By genomic sequencing, we identified two interesting mutations that would lead to residue

| dbSNP  | Exon no. | Nucleotide no. | Amino acid | WT/mut nt | WT/mut aa | Functional consequence prediction |
|--------|----------|----------------|------------|-----------|-----------|-----------------------------------|
| rs142045517 | 2       | 103           | Ala34      | G/A       | Ala/Thr    | Ala34 is Ser in rodent C\(_\alpha\) and Thr in human C\(_\beta\) – likely few functional consequences |
| rs56029020 | 3       | 121           | Leu40      | T/G       | Leu/Val    | Leu40 is absolutely conserved in metazoan C\(_\alpha\)/C\(_\beta\) homologs – likely deleterious |
| rs112360106 | 3      | 149           | Leu49      | T/A       | Leu/His    | Conserved in metazoan C\(_\alpha\)/C\(_\beta\) homologs – likely deleterious |
| rs148280386 | 5      | 409           | Gly136     | G/A       | Gly/Arg    | Conserved in metazoan C\(_\alpha\)/C\(_\beta\) homologs – possibly functional effects |
| rs142007512 | 6      | 541           | Ile180     | A/G       | Ile/Val    | Highly conserved in vertebrate C\(_\alpha\)/C\(_\beta\) – possibly functional effects |
| rs11541563 | 7       | 560           | Gly186     | G/T       | Gly/Val    | Conserved in metazoan C\(_\alpha\)/C\(_\beta\) homologs – likely deleterious |
| rs34988264 | 8       | 673           | Leu224     | -/G, 1 nt insertion | Frame shift in exon B – very likely results in nonfunctional C\(_\alpha\) |
| rs35635531 | 9       | 791           | Ser263     | C/G       | Ser/Cys    | Highly conserved in metazoan C\(_\alpha\)/C\(_\beta\) homologs – possibly functional effects |
| rs146946205 | 9     | 842           | Arg280     | G/A       | Arg/His    | Conserved in metazoan C\(_\alpha\)/C\(_\beta\) homologs – likely deleterious |
| rs187770246 | 9    | 926           | Arg308     | G/A       | Arg/Lys    | Conserved as Arg or Lys in metazoa – likely few consequences |
| rs149832080 | 10    | 955           | Phe318     | T/C       | Phe/Leu    | Not conserved in vertebrates – likely few consequences |
| rs141087932 | 10   | 1045          | Ser348     | T/C       | Ser/Pro    | Not conserved in mammals – likely few consequences |

Enumeration is given as in Table 3, and sequence conservation is illustrated in Supplementary Fig. S1. Reference identifiers from dbSNP [36,49] are included in the left column.

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C\(_\alpha\)Gly186Val was successfully purified by RIIz-affinity chromatography followed by elution with cAMP. This, together with the fact that C\(_\alpha\)Gly186Val forms cAMP-sensitive holoenzymes with RIIz in vitro suggests that lack of enzymatic activity is not caused by an overall misfolding of the protein. We rather suggest that lack of kinase activity is due to less extensive molecular changes, affecting only parts of the protein structure. The introduction of a bulky residue in place of Gly\(_{186}\) is likely to affect many critical factors necessary for kinase activity. The most apparent explanation is that the aliphatic Val\(_{186}\) side chain leads to less efficient binding of Mg\(_1\)/Mg\(_1\) due to displacement of the cation itself or water molecules that are solvating Mg\(_1\)/Mg\(_1\). Loss of Mg\(_1\)/Mg\(_1\) would in turn disable ATP binding, thus explaining the kinase inactivity. This hypothesis is supported by our BRET results, which show exclusive binding to RII but not RI subunits. Moreover, we were unable to purify C\(_\alpha\)Gly186Val using PKI affinity chromatography (results not shown). It has previously been shown that the R\(_1\) subunits, as well as PKI, need ATP to bind C with high affinity [53,54].

According to our model, it is not unlikely that the hydrogen bond between residues 184 and 186 fails to form in C\(_\alpha\)Gly186Val, leading to displacement of Asp\(_{184}\) and inability to position Mg\(_1\) or Mn\(_1\) and thereby ATP in the active site. The inability to bind ATP implicates that the C-spine is not established, which is necessary for the active conformation. An alternative explanation for the inactivity is that Val\(_{186}\), instead of suppressing binding of Mg\(_1\)/Mg\(_1\) and ATP, is rather displaced, leading to a conformational change in the DFG motif itself. This would lead to a malformed R-spine, which is also thought to be incompatible with kinase activity [26].
Reduced kinase activity was also observed for Ca1Ser109Pro. Ser109 is located in the middle of β-strand 4 in the small lobe and this β-strand stabilizes the N-terminal end through targeting Ser109 to Thr37 [48]. It is well known that the N-terminal tail is important for Ca subunit stability, and deletion of the N-terminus has previously been demonstrated to lead to a significant reduction in thermal stability [56]. We speculate that mutation of Ser109 to Pro leads to partial loss of N terminal structure and destabilization of the kinase, and that this may in part provide an explanation for the reduced kinase activity. A second consequence of mutating Ser109 may be associated with our recent findings that Ser109 belongs to a series of signature residues that can be used to distinguish the Ca1 from Ca2 ortholog (unpublished results). Due to this, the Ser109Pro mutation may result in alteration of Ca1-specific functions which do not include holoenzyme formation since our BRET results showed that Ca1Ser109Pro formed holoenzymes with both RIα and RIIα with comparable affinities as the Ca1WT. Full comprehension of the reduced kinase activity and other features associated with mutation at Ser 109 merits further investigation.

Neither mutations of Arg45Gln nor Ser263Cys influenced apparent kinase activity. Arg45 is located near a recently identified conserved pocket in the N lobe known as the N lobe cap, which is above the crucial amino acids Ala70 and Lys72 [48]. Ala70 is part of the C-spine while Lys72 is directly involved in ATP binding. It might have been expected that mutation of Arg 45 would affect kinase activity, also because this residue is highly conserved in metazoan PKA Ca/Cβ homologs (Supplementary Fig. S1) and is clearly under strong purifying selection. The fact that we did not observe any change in phosphotransferase activity suggests that the mutation does not influence the positioning of Ala70 and Lys72. Despite of this, it may be that mutation of residue 45 affects other features necessary for function of the kinase. It could be speculated that the N lobe cap may also be a docking site for proteins inhibiting kinase activity by disturbing either the C-spine or Lys72. This may indicate that the Arg45Gln mutation could be involved in deregulated PKA activity due to altered interactions with so far unidentified interaction partners docking to the N lobe cap. This binding partner does not include the R subunit since the BRET experiments of Ca1Arg45Gln showed no differences in binding and

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**Figure 2.** Activity of wild type and Gly186Val mutated Ca1 and Ca2. A. Phosphotransferase activity of Ca1WT and Ca1Gly186Val expressed in HEK-293T cells was measured in the presence of Kemptide and γ-32P-ATP, and in the presence (+) and absence (−) of cAMP or PKI kinase inhibitor. Enzyme activity was normalized to activity of Ca1WT which was set to 100%. The data are presented as mean values ± SD of triplicate experiments. B. Recombinant Ca2WT (not shown) and Ca2Gly186Val were expressed in BL21 (DE3) cells. For Ca2Gly186Val expressed protein was purified by running cell extracts over a Ni-resin loaded with His6RIIαGly337Glu affinity column and eluted with 10 mM cAMP. The purification steps of Ca2Gly186Val are shown after separation of the various fractions by SDS-PAGE. The crude cell extract (lane 1, crude) was centrifuged and divided into a soluble (lane 2) and an insoluble fraction (lane 3, pellet). Proteins not retained on the column is shown (lane 5, Flowthrough). W1, W3, W4 and W5 (lanes 6 and 8–10) depict protein contents of successive washing steps using buffer containing 50 mM NaH2PO4 (pH 8.0), 5 mM β-mercaptoethanol, and 25 mM KCl. E1-E4 (lanes 11–14) depict protein content in consecutive elution fractions using buffer containing 50 mM NaH2PO4 (pH 8.0), 10 mM cAMP, 5 mM β-mercaptoethanol, and 25 mM KCl. To assure equal amounts of RIIαGly337Glu and Ca2Gly186Val cell extracts of RIIαGly337Glu and Ca2Gly186Val expressed in separate bacteria cultures were mixed 1:1 (lane 7, 1:1 mix). Protein with the correct molecular mass (arrow to the right) was obtained in fractions E1–E3. Ten μl was applied in lanes 6, 8, 9 and 10, seven μl in lane 1, 2, 3, 5 and 7, and five μl in lane 4. (Molecular Weight Marker (MWM)). C. Kinase activity of recombinant Ca2WT and Ca2Gly186Val was determined by employing the spectrophotometric Cook assay. The data are presented as mean values ± SD of triplicate experiments. doi:10.1371/journal.pone.0034838.g002

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Novel Mutations in the Gene Encoding Calpha of PKA

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release upon fsk/IBMX stimulation of RIα and RIIα subunits compared to Ca1WT. This is also consistent with the localization of Arg45 far from the R subunit docking site.

Ser263 is highly conserved among metazoa (Supplementary Fig. S1) and is part of the H helix at the very lower end of the large lobe. This site is close to the R subunit docking sites, and the

![Figure 3. Mutation of Gly186 in Ca1 prevents Ca from binding ATP and divalent cations. A. The 3D structure of the catalytic site of Ca1. Selected conserved motifs and their relations to divalent cations Mn1 and Mn2 and ATP are shown. Residues connecting phospho-Thr197 (pThr197) to the DFG motif are represented as stick models [50]. Mn2ATP (green), the DFG motif (teal), Gly-rich loop (salmon), catalytic loop (yellow), and activation loop (purple) are also highlighted. B. Overall structure of Ca1WT with the conserved structural motifs the C- and R-spine structural motifs highlighted. The boxed segments depict spatial relations between residue 186 (Gly or Val) and ATP, divalent cations, and the C- and R-spines. C. DFG motif in Ca1WT (left) and Ca1Gly186Val (right) and its relations to Mn1 and ATP. Residues are represented as stick models with carbon (orange), oxygen (red) and nitrogen (blue) atoms. The hydrogen bond between the side chain of Asp184 and the amide group of Gly186 (dashed line) is predicted to be broken in Ca1Gly186Val due to the Val side chain. The models are based on the structure with PDB identifier 3FJQ [48]. doi:10.1371/journal.pone.0034838.g003

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A. Co-expression of GFP-Cx1wt or GFP-Cx1mut with wt Rluc and Rluc subunits fused to Renilla luciferase (Rluc-Rluc, Rluc-Rluc, as described in Diskar et al [55] and Material and Methods). Western blot analysis of COS-7 cells expressing either Renilla luciferase alone (lane 1), or Rluc plus GFP-Cx1wt (lane 6) and GFP-Cx1mut (lanes 2–5). Molecular mass and C subunit identity is shown at the left and right, respectively. B. Analysis of PKA type I holoenzyme formation (Rluc-Rluc) with GFP-Cx1wt or GFP-Cx1mut. Expression of Rluc alone defines the background signal (lane 1). 48 h post transfection the luciferase substrate was added and light emission from GFP and luciferase was monitored in the absence (−) or presence (+) of 50 μM forskolin (fsk) and 100 μM IBMX (+). The results are shown as mean values ± SEM of 3 experiments. C. Analysis of PKA type II holoenzyme formation (Rluc-Rluc) was done as described for PKA type I in panel B. doi:10.1371/journal.pone.0034838.g004

Figure 4. Assessment of PKA type I and II holoenzyme formation and dissociation with CaWT and CaMut applying BRET in living cells. Co-expression of GFP-Cx1wt or GFP-Cx1mut with wt Rluc and Rluc subunits fused to Renilla luciferase (Rluc-Rluc, Rluc-Rluc, as described in Diskar et al [55] and Material and Methods). Western blot analysis of COS-7 cells expressing either Renilla luciferase alone (lane 1), or Rluc plus GFP-Cx1wt (lane 6) and GFP-Cx1mut (lanes 2–5). Molecular mass and C subunit identity is shown at the left and right, respectively.

B. Analysis of PKA type I holoenzyme formation (Rluc-Rluc) with GFP-Cx1wt or GFP-Cx1mut. Expression of Rluc alone defines the background signal (lane 1). 48 h post transfection the luciferase substrate was added and light emission from GFP and luciferase was monitored in the absence (−) or presence (+) of 50 μM forskolin (fsk) and 100 μM IBMX (+). The results are shown as mean values ± SEM of 3 experiments.

C. Analysis of PKA type II holoenzyme formation (Rluc-Rluc) was done as described for PKA type I in panel B. doi:10.1371/journal.pone.0034838.g004

The other investigated mutations are likely more compatible with normal development and may not be associated with disease since neither of them influenced holoenzyme formation and mutation of Ser109 only partly reduced catalytic activity. The reason why Gly in the DFG motif is a relatively frequent site of disease-causing mutation in various protein kinases which is most likely due to kinase inactivation [57]. As described above, homozygote targeting mutation of the PREICA gene in mouse is associated with high pre- and postnatal lethality, most probably due to lack of kinase activity at critical steps in embryonic development [30,32]. Based on this, it is expected that any mutation affecting Cα activity may lead to a severe phenotype and possibly disease in humans. Since the Cα-Gly186Val mutation resulted in catalytic inactivation and partial lack of holoenzyme formation, homozygote mutation for Gly186 in vivo may functionally be considered a gene KO and hence may be incompatible with normal development and life. The same would most likely be the case for the frame shift mutation detected in exon 8. The fact that Cα-Gly186Val exclusively forms holoenzyme with RII subunits suggest that individuals with heterozygote mutation of Gly186 may have reduced levels of PKA type I holoenzymes, in addition to harboring type II holoenzymes occupied by inactive C subunits, which can hypothetically cause an imbalance in PKA signaling.

Among the mutations identified in the sequencing of the PREICA gene, Arg45Gln and Ser109Pro were identified in one and two samples, respectively. Of the two mutations identified in the database search only the Ser263Cys mutation was identified by two independent submitters. To what extent this indicates anything about the prevalence of the different mutations, remains to be verified. However, Gly in the DFG motif is a relatively frequent site of disease-causing mutation in various protein kinases which is most likely due to kinase inactivation [57]. As described above, homozygote targeting mutation of the PREICA gene in mouse is associated with high pre- and postnatal lethality, most probably due to lack of kinase activity at critical steps in embryonic development [30,32]. Based on this, it is expected that any mutation affecting Cα activity may lead to a severe phenotype and possibly disease in humans. Since the Cα-Gly186Val mutation resulted in catalytic inactivation and partial lack of holoenzyme formation, homozygote mutation for Gly186 in vivo may functionally be considered a gene KO and hence may be incompatible with normal development and life. The same would most likely be the case for the frame shift mutation detected in exon 8. The fact that Cα-Gly186Val exclusively forms holoenzyme with RII subunits suggest that individuals with heterozygote mutation of Gly186 may have reduced levels of PKA type I holoenzymes, in addition to harboring type II holoenzymes occupied by inactive C subunits, which can hypothetically cause an imbalance in PKA signaling.

The other investigated mutations are likely more compatible with normal development and may not be associated with disease since neither of them influenced holoenzyme formation and mutation of Ser109 only partly reduced catalytic activity. Despite this there is a possibility that homozygote mutation of Ser109 may be associated with disease since experiments on mice have demonstrated that reduced C subunit gene expression can lead to spinal neural tube defects [33]. In total, 13 nonsynonymous point mutations in the PREICA gene were identified in the present study. For example the Gly186Val mutation was only identified in a single EST sequence, and no frequency data was available. Hence, the information on the prevalence of the various mutations is therefore limited and a full comprehension of their existence in patients or patient groups remains to be elucidated. Finally, it is also worth mentioning that several thousand regions of the human genome have structural variation in large segments termed Copy Number Variation (CNV) [58]. In this paper they report that three out of 95 individuals were found to have a loss of ~160 kb which included the whole PREICA gene as well as up- and downstream genes. Due to this it may be speculated that a combination of CNV deletion at the PREICA locus and a heterozygote loss-of-function mutation of Cα could be associated with disease due to severe reduction in C subunit gene dose. To what extent this is a cause of disease remains to be determined.
Supporting Information

Figure S1  Multiple sequence alignment of human PKA Cα (top row) and homologous sequences from a number of metazoan species for the sequence segments containing the 13 mutations discussed in the present study. Enumeration is according to the human PKA Cα1 splice variant and the mutated residues are highlighted. The sequences were obtained from the NCBI (http://www.ncbi.nlm.nih.gov) and UniProt (http://www.uniprot.org) protein sequence databases with the following identifiers: P17612, P05132, P00517, NP_001003039, Q9W0N3, A3KMS9, NP_001003470, P22694, P68118, P05131, XP_867543, XP_422359, Q7ZWW0, Q3ZB92, Q7T374, XP_001175934, XP_002740161, CAG44453, XP_393285, XP_968170, NP_476977.

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