Functional comparison of protein domains within aPKCs involved in nucleocytoplasmic shuttling

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Summary
The atypical protein kinases C (PKC) isoforms ι and ζ play crucial roles in regulation of signaling pathways related to proliferation, differentiation and cell survival. Over the years several interaction partners and phosphorylation targets have been identified. However, little is known about the regulation of atypical aPKC isoforms. To address this question, we performed a comparative analysis of atypical aPKCsι/ι and ζ in MDCK cells. By using green fluorescence protein (GFP) fusion proteins containing the full-length or truncated proteins, we were able to recognize differences in subcellular localization and nucleocytoplasmic shuttling of both isoforms. We show, that an earlier described nuclear localization sequence (NLS), plays a role in the regulation of atypical aPKCsι but not in aPKCsι, despite the fact that it is present in both isoforms. Leptomycin B treatment induces accumulation of GFP-fusion protein of both isoforms in the nucleus. Regardless, the loss of the NLS only decreases shuttling of aPKCsζ, while aPKCsι remains unaffected. In addition, we identified the hinge region as a potential regulator of localization of atypical PKCs. With a set of chimeric proteins we show that the hinge region of aPKCsι mediates nuclear localization. In contrast, the hinge region of aPKCsζ causes exclusion from the nucleus, indicating two different mechanisms leading to isoform specific regulation. Taken together, we show for the first time, that the atypical isoforms aPKCsι and ζ play different mechanisms regarding their regulation of subcellular localization and translocation into the nucleus in MDCK cells.

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Key words: GFP-fusion protein, HINGE domain, NLS, Atypical Protein kinase C, Nuclear translocation, Nucleocytoplasmic shuttling

Introduction
Signal transduction is an essential process that enables the cell to interpret incoming signals and interact with the surrounding environment. Many different signaling pathways and a wide variety of mechanisms are utilized to accomplish this translation. Kinase activation is a very common mechanism that alone can amplify the incoming signal through the phosphorylation of specific substrates. Kinase substrates are often themselves kinases.

Protein Kinase C (PKC) represents a serine/threonine kinase gene family that has often been described as having a major role in signal transduction events. Nine PKC members have been described in mammals which are divided into three subgroups based on their biochemical properties and sequence homology: Classical PKCs (α, β and γ), novel PKCs (δ, ε, θ and η) and atypical PKCs (ζ and τ/λ). Classical and novel PKCs have been found to bind and be activated by tumor-promoting phorbol esters (TPA) which has connected at least some of their physiological functions in vivo functions to the promotion of cancer.

Atypical PKCs were the last to be identified and characterized. Remarkably, early findings showed that both aPKC isoforms are unresponsive to TPA and are also activated independently of calcium and diacylglycerol under normal cell conditions. Thus aPKCs represent a PKC subgroup that is clearly separated from the rest by its mode of activation. Both atypical PKC isoforms have been found to be regulated through their cysteine-rich C1 domain by Par-4 activation (Diaz-Meco et al., 1996) and through their Phox/Bem1 (PB1) domain by interaction with p62 and PAR-6 (Puls et al., 1997; Joberty et al., 2000; Sánchez et al., 1998). Another striking point within the aPKC subgroup is the high content of homologous amino acids between them (72% overall) (Akimoto et al., 1994) which limits the availability of isoform-specific tools. Thus it remains difficult to biochemically analyze and distinguish between the two aPKC isoforms. In addition, many overexpression studies have so far failed to assign functional differences to PKCsζ and PKCsτ/λ. (Diaz-Meco and Moscat, 2001; Uherbal et al., 1999).

Nevertheless recent attempts utilizing the gene targeting approach in mice have revealed that highly specific in vivo functions exist for both aPKCs. For example, the phenotypical analysis of the PKCsζ deficient mouse line lead to the identification of impairment of the NFκB signalling pathway (Leitges et al., 2001) as well as changes in IL-4 and IL-6 signaling (Martin et al., 2002; Galvez et al., 2009). A conventional knock-out of PKCsτ/λ revealed a fundamental role for this kinase during embryonic development (Bandyopadhyay et al., 2004; Soloff et al., 2004; unpublished data). Tissue specific knockouts of this isoform have also shown the importance of PKCsτ/λ in the adult organism (Farese et al., 2007; Huber et al., 2009; Murray et al., 2009; Yang et al., 2009).

Whereas the embryonic lethal phenotype of PKCsτ/λ could be partly explained by the lack of PKCsζ expression at the defined
time of development (Kovac et al., 2007) all other described phenotypes of both aPKC knockouts take place with the existence of the alternate aPKC isoform. Thus these in vivo findings clearly imply that both aPKCs have an isoform-specific, non-redundant function in the cell, even when co-expressed. How these specificities are regulated remains largely unclear. Since specific localization of PKCs within a cell is believed to be one way of controlling specificity (Rosse et al., 2010) we decided to screen for domains within both aPKCs which are able to direct localization and are different between the two isoforms.

Results

1. Generation of truncated aPKC/GFP fusion proteins.

We have used truncated aPKC proteins fused with a GFP tag to follow and identify domains that confer aPKC isoform specific translocation ability. Except for the full length constructs of both isoforms, we have generated mutant constructs that focus on changes in the N-terminal regulatory region of the proteins and have a truncated C-terminal, thus they are missing the kinase domain. Although we cannot say that translocation sequences do not exist in the C-terminal kinase region, we expected to see more changes in translocation potential upon alteration of the N-terminal regulatory region. Thus we initially generated the following for both isoforms: the GFP labeled full length construct; an (aPKC\textsuperscript{\textast }\textsuperscript{catalytic}) construct missing the whole kinase domain including a nuclear export sequence (NES); an N-terminal version (aPKC AA1-176(177*)) including the PB1 (Phox and Bem1) and a C1 (cystein-rich) domain but excluding the hinge region. We further generated a construct consisting of the hinge region alone (aPKCA1-176(177)) and found there was a reversion to a similar localization pattern between the two isoforms (Fig. 2E,F). However, in contrast to the full length proteins, both truncated versions located to the nucleus with a distinct punctate pattern. This pattern was different to that seen with the aPKC\textsuperscript{\textast }\textsuperscript{catalytic} form.

Based on these results further investigation focused on the hinge region alone. Analysis of the aPKC\textsuperscript{Hinge} construct showed a dominant localization to the nucleus as well as a diffuse weak staining in the cytoplasm (Fig. 2G). The corresponding PKC\textsuperscript{C}\textsuperscript{Hinge} construct showed cytoplasmatic staining with some aggregates surrounding the nucleus, no membrane staining, and most significantly, an exclusion from the nucleus (Fig. 2H).

Taken together these results identified first that the hinge region of both aPKCs harbors information important for subcellular localization and second that the N-terminal causes localization to the nucleus and does not need the hinge region for this localization to occur.

2. Isoform-specific domains cause different subcellular localization.

To test whether the truncation mutants are able to identify domains within aPKCs which convey isoform-specific localization we overexpressed each construct in MDCK cells. Both full length constructs showed a widely distributed diffuse GFP signal over the whole cell without prominent membrane localization. There was a slightly increased PKC\textsuperscript{c} nuclear localization when compared to aPKC\textsuperscript{AA1-176} full length (Fig. 2A,B).

Deletion of the C-terminal kinase domain changed the distribution. The aPKC\textsuperscript{\textast }\textsuperscript{catalytic} fusion protein localized exclusively in the nucleus with a regional concentration of the GFP signal within the nucleolus (Fig. 2C). In sharp contrast the corresponding truncated PKC\textsuperscript{\textast }\textsuperscript{catalytic} construct showed an almost exclusive and very strong membrane signal (Fig. 2D) which resembled that associated with the role of aPKC in polarity.

We proceeded by testing N-terminal truncations missing the hinge region (aPKCA1-176(177)) and found there was a reversion back to a similar localization pattern between the two isoforms (Fig. 2E,F). However, in contrast to the full length proteins, both truncated versions located to the nucleus with a distinct punctate pattern. This pattern was different to that seen with the aPKC\textsuperscript{\textast }\textsuperscript{catalytic} form.

3. Functional investigation of aPKC nuclear localization sequence (NLS).

Given the fact that nuclear localization of aPKC represents a new area of research in aPKC in vivo function, we decided to investigate the underlying mechanism. A functional nuclear localization sequence (NLS) located within the C1 domain had been defined in both aPKCs (Perander et al., 2001). To evaluate whether this motif drives the PKC\textsuperscript{C}AA1-177 fragment into the nucleus within our experimental context we mutated the amino acid sequence AKRNFRR to AKRNF in order to generate a nonfunctional NLS. Upon transfection with this construct in MDCK cells we were not able to detect any difference to that found with the parental construct (data not shown). These findings suggested that the N-terminal PKC\textsuperscript{C}AA1-177 fragment translocates into the nucleus in an NLS independent manner through a different mechanism and we assumed that this was also true for the N-terminal PKC\textsuperscript{\textast }AA1-176 fragment.

We next asked whether the NLS could have an impact on the kinetics of nuclear translocation of both aPKCs. Differences in the nuclear translocation kinetics among aPKCs had been earlier discussed by others (Perander et al., 2001). Based on the finding that N-terminal constructs localize to the nucleus we next worked with full length constructs for both aPKCs in order to analyze the
translocation kinetics upon Leptomycin B (LMB) treatment. Treatment with LMB effectively blocks all nuclear export related to CRM1/Exportin and represents an established method to study nucleo-cytoplasmic shuttling (Lischka et al., 2001; Kudo et al., 1998).

When comparing mouse aPKC cDNA sequences we observed that aPKC\(_\text{c}^{\text{f}}\) contains a G to R exchange at position 141(AKRFNR\(_{141}\); supplementary material Fig. S3). This sequence variation in the NLS had been described to affect the binding of importin\(_\alpha/\beta\) and subsequently affect the nuclear translocation...

Fig. 2. Differences in sub-cellular localization of truncated aPKC protein. MDCK cells were transfected with different aPKC\(_\text{i}\) (A,C,E,G) or aPKC\(_\text{c}^{\text{f}}\) fusion proteins (B,D,F,H). The insert shows magnification of the marked area (square) in the overview. Visualization of the nucleus was achieved by DAPI staining. Membrane staining was performed using Alexa Fluor\(_{647}\) conjugated wheat-germ agglutinin (WGA). All pictures were taken with the ZEIJS LSM 520 microscope. The areas representing the cytosolic fraction or nuclear fraction are indicated with bars at the top of each histogram. The histogram analysis was performed with the ‘profile tool’ in the LSM Meta\(_{\text{H}}\) software. The analyzed area is indicated by the yellow straight line in the overview. Scale bar: 50 \(\mu\)m (overview), 10 \(\mu\)m (insert). All experiments were performed in triplicates.

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kinetics since the AKRNFRRR motif containing three arginine’s (as found in PKCθ) enables a direct binding of importin β instead of a sequential binding of importin α followed by importin β (Palmeri and Malim, 1999). Thus we first analyzed the translocation behavior of both full length aPKC-GFP fusion constructs by measuring the GFP signal ratio between the cytosol and the nucleus. Both aPKCs accumulated in the nucleus upon LMB treatment but it was clear that PKCθ accumulated faster and attained a higher level after 6 hours of treatment (Fig. 3).

We next introduced a construct with an R to G mutation at position 142 in PKCθ which mimics a PKCθ NLS. We found that the kinetics of nuclear accumulation upon LMB treatment did not change at all with this construct, which was opposite to our expectations. We then generated a R141/142E mutation in the PKCθ/GFP full length construct thereby destroying the NLS motif. The subsequent analysis of this mutant (aPKCθ R141/142E) again revealed no significant change in the nuclear translocation behavior of PKCθ thus the NLS present in aPKCθ appears to be nonfunctional in the context we tested here.

We then concentrated on PKCζ, first generating a mutant of the full length construct which mimics the PKCζ situation (aPKCζ G141R). As indicated in Fig. 3, this conversion lead to a faster increase and higher amount of the nuclear signal after 2 hours with no significant change after 6 hours, thus resembling the aPKCζ wildtype situation. Next we analyzed the aPKCζ R140E/G141E mutant representing a NLS deficient aPKCζ variant. In this case we were able to detect a significant decrease of the nuclear fraction after 2 and 6 hours LMB treatment implying that the NLS in PKCζ is functionally involved in the nuclear transport of this aPKC. Nevertheless it was also clear that even in the NLS deficient situation a smaller fraction is translocated to the nucleus suggesting that beside the NLS driven transport other nuclear transport mechanism(s) exist which aid in the localization of PKCζ to the nucleus (Fig. 3).

Taken together the mutational analysis of the NLS of both aPKCs revealed that PKCθ can translocate to the nucleus independent of its NLS whereas PKCζ’s nuclear translocation is mainly regulated by its NLS. In addition we observed that the sequence variant we identified in mouse for PKCζ slows the translocation kinetics and also that PKCζ retains a nuclear translocation capacity that is independent of the NLS.

We then focused attention on the analysis of the full length protein fusions to investigate the consequence of a PB1 mutation in aPKC nuclear localization. Having established that the generation of a NLS deficient N-terminal PKCζAA1-177 fragment does not abolish its nuclear localization we decided to investigate the possibility of other nuclear translocation domains existing within this fragment. Besides the C1 domain which includes the discussed NLS, another protein-protein interacting domain has been described in the N-terminus of both aPKCs; the Phox/Bem1 (PB1) domain. Several PB1 aPKC interacting proteins have been identified e.g. Par6 and p62 (Joberty et al., 2000; Hirano et al., 2004). Of particular interest in this context is a recent report describing the translocation of Par6 to the nucleus via an internal NLS (Cline and Nelson, 2007). Thus we hypothesized that aPKCs may be translocated to the nucleus through a “hitchhiking” mechanism by their binding to PB1 containing proteins, like Par6. To verify this hypothesis we first introduced a D63A mutation in the PKCζAA1-177 fragment. This D-A mutation has been shown by Hirano to disrupt interaction between the aPKC PB1 domain and the PB1 domain of Par6 and p62 (Hirano et al., 2004; Hirano et al., 2005). However, the abolishment of a functional PB1 domain did not prevent a nuclear translocation of this construct (Fig. 4A, B). In addition, the corresponding D62A mutation in the PKCζAA1-176 fragment reflected the same outcome (Fig. 4C, D). Thus we assumed that in both atypical PKC N-terminal fragments either a cryptic NLS or another sequence motif existed that was responsible for the observed nuclear localization.

We then focused attention on the analysis of the full length protein fusions to investigate the consequence of a PB1 mutation in aPKC nuclear localization in this context. Thus we introduced the D63A or D62A mutations respectively in each full length aPKC construct which we subsequently analyzed in an LMB experiment.

In order to verify the functionality of the introduced mutation we first checked whether the PB1 mutation caused a loss of membrane association due to the protein’s inability to form the polarity complex with Par6/aPKC/Par3 (Lin et al., 2000; Wodarz...
subcellular localization of these constructs we observed that both aPKC isoforms harboring the D to A mutation showed a significant increase in the pool of nuclear located fusion protein in the non-LMB treated samples (Fig. 4E,F, control values).

With regard to PKCζ, LMB treatment caused a similar kinetic rate of nuclear accumulation for both wild type and mutated constructs whereby the D62A mutant translocation started at a significantly higher level, but both reached maximum accumulation levels after 4 hours of LMB treatment (Fig. 4F).

In contrast, PKCζ gave a different picture of nuclear accumulation. Whereas the wild type construct increased its amount in the nucleus constantly over the 6 hour time line, the D63A mutation of PKCζ increased its nuclear level only slightly (but significantly) within the first 2 hours. We were unable to detect any further significant increase of the mutant construct in the nucleus beyond this time-point (Fig. 4E).

Taken together we have shown that the PB1 domain within both aPKCs is not involved in the nuclear translocation of the N-terminal constructs used here indicating the existence of an alternate nuclear translocation mechanism for these two fusion proteins. However, this analysis also revealed that in the context of the full length protein, the PB1 domain does influence the localization of aPKCs in an isofrom specific manner: whereas PKCζ appears to translocate into the nucleus independent of its PB1 domain, PKCζ translocation seemed unable to proceed beyond a certain level of nuclear content (reached at 2 hours after addition of LMB) implying that at least a fraction of the transported GFP fusion protein depends on a functional PB1 domain.

5. Atypical PKCs hinge region mediates different localization of aPKCζ and aPKCζ.

Our initial screen revealed that the hinge region of both aPKCs appeared to be involved in the unique sub-cellular localization of each isoform (Fig. 2). To further elucidate these findings we generated mutants with exchanged hinge regions that either harbored the aPKCζ specific hinge sequence within the aPKCζ specific background or vice versa (Fig. 1B). Upon transfection of the N-terminal chimeric aPKCζ N-HINGE/GFP fusion protein into MDCK cells it became apparent that the aPKCζ specific hinge sequence is able to keep the aPKCζ protein outside of the nucleus (Fig. 5A,C). Interestingly this chimeric fusion protein did not reflect the same pattern as the aPKCζ N-HINGE/GFP fusion protein where the primary target of the hinge mutation is the N-terminal hinge region of the aPKCζ protein (Fig. 4F). This suggests that the hinge region of aPKCζ is not involved in the nuclear translocation of the N-terminal chimeric aPKCζ N-HINGE construct.

These findings clearly identified that the hinge region of atypical PKCs can be distinguished as harboring specific sub-cellular localization information.

We then proceeded to ask the same question in the context of the full length protein. We used either wildtype or chimeric fusion proteins which contained the corresponding hinge region of the alternate aPKC isoform (Fig. 1B). We first analyzed the subcellular localization of these constructs by measuring the nuclear to cytosol ratio of the GFP signal. As indicated in Fig. 5E.
(control lane) an aPKCζ protein harboring a ψ-hinge region showed a significant increase in the nuclear fraction of the GFP signal whereas the chimeric aPKCζ protein did not appear to be affected by the ψ-hinge region with regard to its nuclear localization (Fig. 5F, control lane). After 6 hours of LMB treatment the chimeric aPKCζψHINGE construct showed a higher accumulation of the GFP signal in the nucleus (at a similar level to that of the aPKCζ wildtype construct) (Fig. 5E). In contrast, the chimeric aPKCζψHINGE protein showed a significant decrease of its nuclear signal after 6 hours of LMB treatment thus mimicking the levels reached by the aPKCζ wild type construct (Fig. 5F).

In summary, both approaches identify the hinge region of both aPKCs as a domain that contains isoform specific sub-cellular localization information.

Discussion

Atypical PKCs represent a subgroup of the PKC family distinguished by their mode of activation and their remarkably high homology content to each other (Steinberg, 2008). Most published work on aPKCs has not distinguished between the two isoforms due to the lack of specificity of available tools such as reliably specific antibodies and peptide inhibitors. In addition, due to the high homology of the amino acid sequence, there is a great potential for redundancy between these two isoforms.

Nevertheless studies using either RNA knockdown (Durgan et al., 2011) or gene targeting approaches in mice (Leitges et al., 2001; Farese et al., 2007) clearly indicate the existence of isoform-specific functions. Phenotypical analysis has revealed quite opposite in vivo functions even in cells where both isoforms coexist, for example in the intestinal epithelial cell, (Oster and Leitges, 2006; Murray et al., 2009). Thus the question of how these different functions are regulated given the high degree of homology becomes poignant.

The intention of this study was to identify how specific domains within both isoforms transfer specificity to the individual protein. As described for other PKCs (Parker and Murray-Rust, 2004) we assumed that one method to achieve specificity is through protein localization within the cell. Thus in an unbiased approach we tested aPKC/GFP fusion constructs for their location after their transient transfection in MDCK cells. In accordance with others (Horikoshi et al., 2009), we found both full length constructs mainly located in the cytoplasm but we also observed that while the aPKCζ construct was found exclusively in the nucleus the aPKCζψHINGE construct showed weak nuclear staining implying a tendency for its nuclear localization compared to aPKCζ.

A clear difference between the two isoforms became obvious when analyzing the N-terminal constructs missing the catalytic kinase domain (aPKCζψcatalytic and aPKCζψcatalytic). It is worth stating here that this truncation eliminates a nuclear export sequence (NES) from both constructs (aPKCζ AA 247–255, aPKCζ AA 239–247) and leaves the truncated protein in a somewhat open conformation. Whereas the aPKCζψ specific construct showed prominent membrane staining, aPKCζψ was found exclusively in the nucleus. These observations per se
indicate a clear difference between both aPKC isoforms. The robust membrane localization of aPKCζ could be explained by its previously reported association with the signaling cascade of the ternary polarity par3/aPKC/par6 complex which is localized at the cell membrane (Suzuki et al., 2001), while the accumulation of the aPKCι construct might be a consequence of the missing NES. The explanation for this obvious difference in localization remains elusive.

Further investigation identified the hinge region of both isoforms to be essential in this localization process. We found that in both isoforms the deletion of the hinge domain caused a nuclear localization that was different in appearance to that observed in aPKCιAcatalytic construct expression where the hinge region is intact. We also found that expression of hinge/GFP fusion proteins showed an isoform-specific difference.

Whereas the aPKCιHINGE predominantly localized in the nucleus (showing a homogenous distribution within) the aPKCζHINGE was excluded. Thus, given the fact that the hinge region within aPKCs contains the most diverse nucleotide sequence it is interesting to observe that both isoforms contain information related to isoform-specific, distinct, subcellular localization in this region. This was finally confirmed by the expression of chimeric proteins that were generated containing the hinge fragment of one isoform sub-cloned to the corresponding N-terminus (or full length) of the alternate aPKC isoform (Fig. 5). In both cases τ-hinge drives the protein into the nucleus whereas ζ-hinge either keeps it outside (N-terminal construct) or significantly reduces (in the context of the full length protein) the nuclear pool upon LMB treatment.

Until now, the only verified functional motif within the atypical hinge sequence had been a caspase cleavage site which exists only in aPKCζ and is involved in the regulation of the protein’s enzymatic activity rather than its localization (Frutos et al., 1999). Any other functional derivation based on the crude amino acid sequence is highly speculative without experimental verification. Current approaches in our laboratory employed mutagenesis to establish hinge region mutants with abolished localization behavior, thereby enabling the identification of the underlying sequence motifs (and mechanisms) that cause these observed differences between hinge regions in different aPKC isoforms.

The N-terminal construct (depleted of the hinge region) of both aPKCs was found exclusively in the nucleus which could be explained by the NLS motif located in the C1 domain, functionally described by others (Perander et al., 2001; Pu et al., 2006). Nevertheless we investigated the NLS in more detail and showed that mutations destroying the functionality of the NLS did not completely abolish the nuclear translocation of either the N-terminal or the full length GFP fusion proteins in either aPKC. This was a surprising result since the NLS had been shown to cause localization to the nucleus in the context of both a smaller fragment and a full length construct containing a kinase defective mutation in a different cellular context (Perander et al., 2001). We found that neither an N-terminal nor a full length aPKCζ/GFP construct containing an R141/142 mutation appeared to be affected in its nuclear translocation behavior. However, a full-length aPKCζ/GFP construct harboring an R140/141 mutation showed a significant reduction in the amount of nuclear accumulation after 2 and 6 hours LMB treatment. Although reduced, the accumulation was significant, thus indicating the possible existence of alternate methods of translocation.

During our analysis we identified a sequence variant in the NLS sequence which appears to be mouse specific and converts the PKCζ specific NLS to AKRFNR (Uniprot.org entry Q02956). This variation could explain the slower nuclear translocation of PKCζ observed upon LMB treatment compared with PKCι based on the described mechanism of the importin α/β system. Also the G141R mutation of PKCζ which converts the ζ-NLS into a τ-NLS, supports this notion since the construct mimicked the PKCι nuclear translocation kinetics exactly (higher maximum, reached after 2 hours LMB treatment). However, this is not a valid argument for the observed differences regarding the nuclear translocation kinetics between both aPKC isoforms made previously by others (Perander et al., 2001) since the aPKC cDNAs from other species that do not contain this sequence variation were used. Thus, our observation may be either a cell type or species specific phenomena or a combination of both. In any case our data show a clear difference between both aPKCs and their NLS dependent nuclear translocation in MDCK cells upon LMB treatment when using mouse specific constructs. PKCι translocates to the nucleus independently of its NLS whereas PKCζ mainly depends on a functional NLS.

One reason for the NLS independent translocation of PKCι might be hidden in its hinge region which we have shown to be able to direct nuclear translocation. Other mechanisms must also exist since N-terminal constructs of both aPKC isoforms (missing each hinge region and containing NLS mutations) retain the capacity to translocate to the nucleus.

We have further analyzed the PB1 domain which remained intact in our N-terminal constructs and which has been implicated as a mediator of protein-protein interactions between PB1 containing proteins (Hirano et al., 2004; Hirano et al., 2005). To our knowledge, the PB1 interacting proteins include at least 13 mammalian proteins (Lamark et al., 2003).

The background notion for this attempt was that it was possible that interacting partners of aPKCs were able to translocate to the nucleus and could serve as a shuttle vehicle for aPKCs. However, when introducing a D-A mutation which had been described to abolish PB1 specific interactions (Hirano et al., 2004) we were not able to detect any effect on the N-terminal constructs with regard on the nuclear/cyttoplasm ratio. Upon the introduction of the same mutation into full length proteins we were readily able to detect an increased nuclear fraction in non-treated cells. We interpreted this result as an indication that there is reduced export rather than increased import.

One likely candidate for the export shuttle might be p62. This protein was originally identified as a binding partner for aPKC (Puls et al., 1997; Sánchez et al., 1998) and has received some attention recently based on its involvement in starvation induced autophagy (Pankiv et al., 2010; Wooten et al., 2008; Lamark et al., 2009). More important for us was the identification of p62 in the nuclear compartment (Bjørkøy et al., 2005) and the finding that the shuttling of p62 between cytoplasm and nucleus appears to be a very dynamic process (Pankiv et al., 2010) similar to that we have observed for aPKC. In addition, most recently it has been shown that p62 is involved in the nuclear export of ALFY (autophagy-linked FYVE) proteins through protein-protein interaction (Clausen et al., 2010). Thus, it could well be that p62 is also involved in the nuclear export of aPKC. Nevertheless, we do not exclude other PB1 containing proteins, like Par6, also
recently described as a nuclear protein (Cline and Nelson, 2007), being involved in the nuclear translocation of aPKC.

With regard to nuclear import, the PB1 mutation did not influence PKCζ (similar kinetics and amounts of nuclear accumulation were seen) but it did reduce the total amount of PKCζ in the nucleus upon LMB treatment. Thus a certain fraction of PKCζ translocation to the nucleus depends on an as yet undefined PB1 domain protein-protein interaction.

Our study revealed clear differences between the two aPKC isoforms which have not been analyzed in such a comparative structure to function study before. The hinge region, representing the most heterologous amino acid stretch between the two aPKCs, contains isoform-specific, sub-cellular localization information. In particular, with regard to the nuclear localization of aPKCs the functionality of the NLS, the PB1 domain and in the case of PKCζ the hinge region, are shown to influence the nuclear import to a different degree when both isoforms are compared directly.

We are not the first to show that aPKC is present in the nucleus but our data suggests that there are multiple modes of nuclear transport that differ between aPKC isoforms. This could mean that the physiological conditions in which each aPKC translocates to the nucleus are distinguishable. What exactly drives PKCζ and PKCζ into the nucleus is not clear. Our attempts to locate aPKC/GFP proteins to a more precise sub-nuclear domain or to the nucleus during specific cell cycle periods have thus far been unsuccessful (data not shown).

Another option is the activation of aPKC in the context of NFκB signaling leading to a translocation of NFκB to the nucleus. Besides this translocation, specific phosphorylations are required to regulate NFκB specific gene expression. Ser 311 on the RelA subunit has been identified as a PKCζ target (Durand et al., 2003) but where this phosphorylation takes place remains unknown. Thus it could well be that PKCζ must be translocated into the nucleus to phosphorylate RelA on Ser 311.

Both aPKC isoforms have been described as forming complexes with Par6 and Par3 (Noda et al., 2003; Henrique and Schweiguth, 2003; Joberty et al., 2000) to build a ternary complex (also called polarity complex (Suzuki and Ohno, 2006). Recent work has also identified nuclear accumulation of Par6 in MDCK cells after LMB treatment indicating nucleocytoplasmic shuttling similar to that we observed for aPKC (Cline and Nelson, 2007). Cline could not identify a specific nuclear function for Par6, but a participation in gene transcription, RNA splicing and mRNA transport was excluded.

Par3 has also been found to be located in the nucleus (Fang et al., 2007). In this case an association with Ku70/Ku80 proteins was reported and functional analysis revealed that Par3 regulates DNA-PK activity through this interaction which is needed in the repair of double-strand breaks. Neither Par6 nor Par3 has been established polarity complex (Par6/aPKC/Par3) exists in the nucleus independently and fulfills as yet uncharacterized functions.

It is necessary at this point to mention that the N-terminal constructs (PKCζAA1-177 and PKCζAA1-176) we have used in our study have an average molecular weight of around 55 kDa (and even smaller with regard to the individual hinge constructs). Thus as a consequence we are not able to exclude that some of our immunofluorescence data are discolored by the fact that smaller proteins can enter the nucleus by passive diffusion (Nigg, 1997).

Regardless of this fact we are convinced that our major findings are not influenced by this fact since all relevant mutation have been introduced and analysed in the full length proteins as well. In addition also the hinge domains have been tested in the context of a bigger fusion protein.

In conclusion, evidence for the implication of aPKCs in nuclear function(s) are accumulating, some may be redundant and some may be assigned to a single aPKC isoform. The functions may be dependent on specific stimuli since we have found that defined domains within the aPKC protein have various influences on the nuclear translocation behavior of the protein. Future challenges include the identification of the physiological conditions in which individual aPKC isoforms translocate to the nucleus which may then suggest links to specific nuclear function.

**Materials and Methods**

**Cell cultures**

Madin Darby Canine Kidney (MDCK) cells were grown in Gibco DMEM+Glutamax®-I (Invitrogen) with 15% Gibco fetal calf serum (Invitrogen), Gibco non essential amino acids (Invitrogen), Penicillin (100 units/ml) and streptomycin (100 μg/ml) (Invitrogen). For the nuclear export experiments, Leptomycin B (Sigma-Aldrich) was added to the medium to a final concentration of 4 ng/ml.

**Plasmid constructions and site-directed mutagenesis**

The murine cDNA was amplified from mouse brain RNA after reverse transcription by PCR using HotStarTag Master Mix from QIAGEN (203,443). Following Primers were used: *f*PKCwt for: CAC CAT GCC GAC CCA GAG GGA CAG CAG C; *l*PKCwt rev: GAC ACA CTC TTC TGC AGC CAT CAA GAG GGG; *f*PKCwt for: CAC CAT GCC CAG GAC GAG CCC CAA CAA GAT G; *l*PKCwt rev: CAC GGA CTC CAC AGC AGA CAG CAG AA; *f*PKCAK rev: ACT GGA CGA CGA CCT TTT AAC ACT CTT CCG CT; *l*PKCAK rev: GCG CCA CCC TGG GAT TGT GAT CCC ATT CAC C; *l*PKCAAI-186rev: GAC CAG CTT G TG CCA CCT CTT CTG GAT G; *l*PKCAAI-178rev: GAG GAG GAC GAG CAC GCA GGG CTT ATG AGC; *f*PKCAAI-227 for: CAG GAT TGT GCC GGC GCA CAC GGC CCG GTG; *f*PKCAAI-184-246: CAG CAG CTT G CGC TGT GAA GCC GCA CTA CCG; *f*PKC Hinge for: CAC CATGAC AAT TGA GTG TGG GCA GCC CT; *f*PKC Hinge: CAC CAGGCG TCG CCA GCA GCC AFA TGG AT; All PCR products were sub-cloned into pENTR-D-TOPo by using the Gateway cloning system (Invitrogen) according to the manufacturer’s instructions. All plasmids were verified by restriction enzymes and sequencing to confirm the correct coding sequence. All expression plasmids were constructed using the LR clonase Kit (Invitrogen) according to the manufacturer’s instructions. The pDEST expression vector with a c-terminal GFP-tag was provided by Kjetil Tasken (the Biotechnology Centre of Oslo, Norway). For sequencing of the constructs we used following primer EGFP-N rev: CTAACCTGGTGGCGCTTACGEP-N for: TCCGGCCATTGAGCGAAT; for the site directed mutagenesis we used the Quickchange Site Directed Mutagenesis Kit (NO. 200518, Stratagene). The primer-design was done with the Quick Change Primer Design tool on the companies homepage. Following primers were used for the mutagenesis: *f*PKCR141/142E for: GCC AAA CGT TTC AAT GAG GAG GGC GCC CAC TGT GAC CTC; *l*PKCR141/142E rev: GAT GGC ACA GTG GGC GCC CTT ATG GAA AGC GGG GCC; *f*PKCR142G for: GCC AAA CGT TTC AAT GAG GAG GGC GCC CAC TGT GT; *l*PKCR142G rev: GCA CAG TGG GCG CCC CAC TTA TGG AAA CAGT TTC GC; *f*PKC140/141E for: AAG CCA AGC GCT TTA ACG AGG AAG CAG CTC GAT GCC GAC; *l*PKC140/141E rev: CTT CGT GGC GCA GTC GCC TCT CTT GCT TAA GAG GTG CTT GGC; *f*PKCG141R for: AGC CCT GTA AGA GAG GGC GCC AGT CCG; *l*PKCG141R rev: GCC GCA GTA GGC TCT CTT GCT TAA GAG CCG CT; *f*PKCD63A for: CAT CAC GAA ATG AGC TGA GGA AGG AGA CCC AT; *l*PKCD63A rev: ATG GGT CTC CTT CCG CAG CTA TCC ATT TCA TGG TG; *f*PKCD62A for: CTT CCT TGG GGG GCC CAC TGG CTA CCG; *l*PKCD62A rev: CTT CAC CCT CAC TGG CCC CAC ATG TGA.

To generate aPKC-NLS and aPKC-N hinge constructs we designed primers to add an additional BsfI restriction site to the N-terminus of the corresponding Hinge-region and the C-terminus of the N-terminal region without changing the amino acid sequence. The following primers were used: *f*PKCN-BSfor: GAC TAG GAG GTG CCA CCT ATG; *l*PKCHINGE-Bsfrev:
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24 hours after transfection the cells were trypsinized with 1 reagent from Invitrogen dependent of the size of the used cell culture dish. MDCK cells were transfected according to the Lipofectamine 2000 linear tool to analyze the localization of the GFP-fusion protein within the cell. To user to analyze the pixel strength of a defined area and all channels. We used the obtained data were in the arbitrary units of the 8-bit gray scale (0–256 units).

Subcellular localization analyzes and immunocytochemistry

For the sub-cellular localization studies of the different GFP fusion proteins, MDCK cells were seeded in a 4-well dish with a 14 mm microscope cover glass (Marenfeld), coated with 0.2% Gelantine at a density of 7.3 i × 104 per well 18 h before transfection. The cells were then transfected with 0.8 μg of expression vector using Lipofectamine. The cells were fixed with 4% paraformaldehyde for 15 min and permeabilized in 0.01% Triton X-100 for 10 min at room temperature, 24 hours after transfection. We used the Vectashield mounting medium with DAPI (Vector) to mount the cells on a microscope slide (VWR). The GFP fusion proteins were visualized by confocal microscopy using the Zeiss LSM 510 Meta inverted microscope equipped with a Zeiss LSM laser module and a AxioCam Mrm digital camera.

Image analysis

The color digital images of the fluorescing cells were processed by the Adobe Photoshop CS5 (Adobe Systems Inc., USA) and analyzed by the Image 1.43r (National Institute of Health, USA, http://rsb.info.nih.gov/iij/) software. The ‘Mean Gray Value’ was quantitatively assessed in the selected regions of interest (ROI “Oval Selection”) from the images using “Analyze” plugin (“Measure” option). The obtained data were analyzed by a linear tool to analyze the localization of the GFP-fusion protein within the cell. To visualize and analyze the dimension of a cell and the position of the nucleus we used Alexa Fluor® 647 (Invitrogen) labeled Wheat Germ Agglutinin (WGA). In the histogram each peak represents a cell membrane. The two outer peaks represent the outer cell membrane and the inner peaks the nuclear envelope. The position and dimensions of the nuclei were visualized and analyzed by using 4′,6-diamidino-2-phenylindole (DAPI). Channel coding: red = WGA, blue = DAPI, green = GFP.

Data analysis

The histogram analysis of the fluorescence signal was performed with the profile tool provided by the LSM Image Examiner from Zeiss. The software enables the user to analyze the pixel strength of a defined area and all channels. We used the linear tool to analyze the localization of the GFP-fusion protein within the cell. To visualize and analyze the dimension of a cell and the position of the nucleus we used Alexa Fluor® 647 (Invitrogen) labeled Wheat Germ Agglutinin (WGA). In the histogram each peak represents a cell membrane. The two outer peaks represent the outer cell membrane and the inner peaks the nuclear envelope. The position and dimensions of the nuclei were visualized and analyzed by using 4′,6-diamidino-2-phenylindole (DAPI). Channel coding: red = WGA, blue = DAPI, green = GFP.

Western Blot analysis

MDCK cells were transfected according to the Lipofectamine 2000® transfection reagent from Invitrogen dependent of the size of the used cell culture dish. 24 hours after transfection the cells were trypsinized with 1× Trypsin (Invitrogen) for 10 min at room temperature, the reaction was stopped by supplementing DMEM containing 15% FCS. The pellet was collected and incubated in 80 μl of protein extraction buffer (50 mM Tris/HCl, pH 7.5–8.0, 2 mM EDTA, pH 7.0, 10 mM EGTA, pH 7.0, 0.1% Triton-X100, 3% β-mercaptoethanol) supplemented with Proteinase Inhibitor Cocktail (Sigma, NO. P-2714), incubated on ice for 20 min. After supplement of 25 μl of protein sample loading buffer the samples were boiled for 5 min. Subsequently, the samples were run on a 10% SDSD-polyacrylamide gel and blotted onto BioRad nitrocellulose membranes. The membranes were blocked in 5% nonfat dry milk in TBST for 1 h at room temperature. Incubation with the primary antibody happened overnight in TBST at 4°C. The following antibodies were used: anti-GFP (Santa Cruz Antibodies NO. sc-8334) (1:500).

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Competing Interests

The authors would like to declare that they do not have conflicting financial interests.

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