**Ca\(^{2+}\)**-controlled competitive diacylglycerol binding of protein kinase C isoformes in living cells

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The cellular decoding of receptor-induced signaling is based in part on the spatiotemporal activation pattern of PKC isoforms. Because classical and novel PKC isoforms contain diacylglycerol (DAG)-binding C1 domains, they may compete for DAG binding. We reasoned that a Ca\(^{2+}\)-induced membrane association of classical PKCs may accelerate the DAG binding and thereby prevent translocation of novel PKCs. Simultaneous imaging of fluorescent PKC fusion proteins revealed that during receptor stimulation, PKCα accumulated in the plasma membrane with a diffusion-limited kinetic, whereas translocation of PKCe was delayed and attenuated. In BAPTA-loaded cells, however, a selective translocation of PKCe, but not of coexpressed PKCα, was evident. A membrane-permeable DAG analogue displayed a higher binding affinity for PKCe than for PKCα. Subsequent photolysis of caged Ca\(^{2+}\) immediately recruited PKCe to the membrane, and DAG-bound PKCe was displaced. At low expression levels of PKCe, PKCe concentration dependently prevented the PKCe translocation with half-maximal effects at equimolar coexpression. Furthermore, translocation of endogenous PKCs in vascular smooth muscle cells corroborated the model that a competition between PKC isoforms for DAG binding occurs at native expression levels. We conclude that Ca\(^{2+}\)-controlled competitive DAG binding contributes to the selective recruitment of PKC isoforms after receptor activation.

Introduction

PKC isoformes form a family of serine/threonine kinases that are grouped into classical (PKCα, β1, β2, and γ), novel (PKCδ, ε, η, and θ), and atypical (PKCζ and η/λ) isoformes based on their structural similarities and cofactor requirements (Nishizuka, 1992; Mellor and Parker, 1998; Ron and Kazanietz, 1999). In contrast to their highly conserved COOH-terminal catalytic core domains, NH\(_2\)-terminal regulatory domains are more variable. The diacylglycerol (DAG)*-mediated activation of classical and novel PKC isoformes is accomplished by a C1 domain that contains two cysteine-rich regions as DAG-docking motifs. DAG binding fulfills a dual role by recruiting classical and novel PKCs to the membrane compartment and by weakening the interaction of an inhibitory pseudosubstrate domain with the COOH-terminal catalytic core (Oancea and Meyer, 1998). The mode of activation, therefore, relies on the DAG-induced release from an intramolecular block. In contrast to novel PKC isoformes, classical PKCs are additionally regulated by the cytosolic Ca\(^{2+}\) concentration ([Ca\(^{2+}\)]). Their C2 domains, in conjunction with the anionic phospholipid phosphatidylinerine, bind two or three calcium ions (Shao et al., 1996; Sutton and Sprang, 1998; Verdaguer et al., 1999). The bound Ca\(^{2+}\) electrostatically facilitates the plasma membrane docking of classical PKC isoformes and may, thus, explain the diffusion-limited translocation kinetics and high collisional coupling efficiency of classical PKC isoformes observed either in vitro or in living cells (Nalefski and Newton, 2001; Schaefer et al., 2001).

In virtually each cell type and tissue, at least one member of the classical and one of the novel PKC isoformes are coexpressed (Wetsel et al., 1992; Dekker and Parker, 1994; Liu and Heckman, 1998). Stimulation of PLC-coupled receptors results in DAG formation and, via inositol-1,4,5-trisphosphate (InsP\(_3\)) receptor activation, increases the [Ca\(^{2+}\)]. One might expect that classical and novel PKC isoformes are recruited to the plasma membrane in parallel. Nonetheless, upon receptor stimulation, an isotype-selective translocation of PKCs has been observed in several cell systems (Dekker and Parker, 1994). This isotype-specific membrane translocation of PKC isoformes must be based on mechanisms that take effect downstream of receptor activation. Although a number of PKC-interacting proteins sequester activated PKCs to various...
compartments (Jaken and Parker, 2000), little is known about the mechanisms that control the selectivity of the initial membrane recruitment of distinct PKC isoforms, which is most likely based on a protein–lipid interaction.

We have recently shown that a Ca\(^{2+}\)-induced plasma membrane translocation of classical PKCs is a diffusion-driven and diffusion-limited binding process that is characterized by a high collisional coupling efficiency (Schaefer et al., 2001). Because both classical and novel PKCs share a DAG-binding C1 domain, the highly efficient Ca\(^{2+}\)-driven plasma membrane association of classical PKCs suggests that, by finally competing for the same acceptor, classical PKCs may prevent translocation of coexpressed novel PKCs. To observe the interplay of PKC translocations in living cells, we generated cyan and yellow fluorescent PKC fusion proteins and applied a multivariate regression algorithm that reliably dissects signals arising from multiple, spectrally overlapping fluorochromes. We provide evidence that an agonist-induced selective membrane recruitment of classical and/or novel PKC isoenzymes can result from a Ca\(^{2+}\)-controlled competitive binding to limiting plasma membrane concentrations of DAGs.

Results
Excitation and emission spectra of fluorochromes form fingerprints that can be used to discriminate a number of dyes. Taking advantage of fast monochromators and linearly scaling CCD cameras, we set up a spectral evaluation method that estimates the contribution of single fluorochromes to complex signals in dynamic, multicolor labeling experiments. Image bursts were taken at five to eight different excitation wavelengths to record the spectral characteristics of the probe. These data were then compared with spectrally resolved datasets of each isolate fluorochrome (Fig. 1 A). Because fluorescence signals of a probe are additively composed of spectra of single fluorochromes, their relative contribution to the composite signal could be estimated with a pixel-by-pixel multivariate, linear regression analysis (results for two regions of interest are depicted in Fig. 1, B and C). Spectral shifts of \(~15\) nm were sufficient to achieve a reliable separation of fluorochromes such as the red-shifted green (GFP-S65T) and yellow (YFP) fluorescent proteins (unpublished data). Furthermore, by dissecting signals that arise from the spectrally shifting free and Ca\(^{2+}\)-bound forms of fura-2, a rapid calibration of cytosolic Ca\(^{2+}\) concentrations could also be achieved. Using this regression-based spectral evaluation method, we synchronously coimaged receptor-induced [Ca\(^{2+}\)]\(_i\) transients and the translocation of CFP and YFP fusion proteins of classical and novel PKC isotypes.

Our recent finding that the receptor-induced translocation of classical PKC isoenzymes is a Ca\(^{2+}\)-driven diffusion-limited binding process (Schaefer et al., 2001) prompted us to hypothesize that after initially attaching to the plasma membrane via interaction of the Ca\(^{2+}\)-binding C2 domain and phosphatidylserine, the subsequent search for DAGs may be restricted to a two-dimensional space. This two-step procedure of classical PKCs to scan for DAGs in the plasma membrane may be more efficient than the translocation mechanism of novel PKCs that have to undergo several collisions with the plasma membrane until they eventually hit a DAG molecule. Indeed, in transiently transfected human embryonic kidney (HEK) cells, YFP-tagged PKC\(\alpha\) translocated to the plasma membrane in response to maximal stimulation of a cotransfected histamine H\(_1\) receptor, whereas coexpressed CFP-fused PKC\(\varepsilon\) remained mostly cytosolic (Fig. 2, A and C). This effect was not due to massive discrepancies in the expression levels of the coexpressed PKC isoforms because the molar ratio between PKC\(\alpha\)-YFP and PKC\(\varepsilon\)-CFP was \(~1:1.5\), as detected by fluorescence intensities of the respective tags and a calibration procedure with an
intramolecularly fused CFP–YFP tandem protein (see Materials and methods). Simultaneous imaging of two classical PKC isoforms, PKCα–YFP and PKCβ1–CFP (molar ratio 1:2), however, showed coincident [Ca^{2+}] signals and simultaneous plasma membrane translocation of both PKCs (Fig. 2, B and C). Switching the fluorescent tags (PKCα–CFP co-expressed with PKCβ1–YFP) did not alter the impaired translocation of the novel isotype in coexpression experiments. Immunoblot analysis of endogenously expressed PKCα and PKCβ along with a dilution series of transiently transfected fluorescent fusion proteins demonstrates an ~10-fold overexpression of PKCα and an ~100-fold overexpression of PKCβ, as compared with endogenous expression levels (Fig. 2 D). Thus, the endogenously expressed PKC isoforms are unlikely to affect the translocation of transiently overexpressed fluorescent PKC fusion proteins. Fluorescent PKCβ fusion proteins translocated in response to histamine receptor stimulation when expressed alone (Fig. 3) or when coexpressed with the unfused CFP (unpublished data), indicat
ing that coexpressed PKCα caused the attenuation of PKCe translocation.

The initial step for the translocation of classical PKCs most likely relies on an interaction of the Ca²⁺-binding C2 domain with phosphatidylserine. Therefore, chelation of intracellular Ca²⁺ may affect both the efficiency and the temporal pattern of PKCα translocation. To evaluate the effect of intracellularly loaded BAPTA on the kinetics of PKC translocation, the localization of fluorescent PKC fusion proteins was assessed by digital videomicroscopy and subsequent analysis of fluorescence intensities within regions of interest defined over the plasma membrane and the cytosol of single cells, as shown in Fig. 3 A. The results demonstrate that the histamine-induced plasma membrane docking of PKCα–YFP is markedly affected by BAPTA-AM pretreatment. The plasma membrane association of PKCα was delayed, and the efficiency of the binding was reduced under conditions where the formation of DAGs is the only driving force for the PKCα translocation (Fig. 3 B). The half-maximal translocation of PKCα in the absence and presence of the Ca²⁺ chelator were observed 1.2 s and 7.5 s after histamine application, respectively. The agonist-induced redistribution of the DAG-sensing PKCe–YFP remained mostly unchanged irrespective of the presence or absence of [Ca²⁺], transients (Fig. 3 C). Half-maximal translocation of PKCe–YFP was observed 6–7 s after histamine stimulation in untreated or in BAPTA-loaded cells. Upon buffering of [Ca²⁺], signals, a slight, but statistically not significant, increase in the translocation efficiencies of PKCe–YFP could be observed (Fig. 3 C, top and middle). Because the endogenous expression level of PKCα in HEK293 cells is not negligibly low (Fig. 2 D), it is possible that this paradoxical inverse Ca²⁺ sensitivity of PKCe translocation could reflect a competition with endogenous classical PKCs. Applying the spectral evaluation method, [Ca²⁺], could be detected in parallel to control the Ca²⁺ clamp of intracellularly loaded BAPTA in the same cells that were used to detect the PKC translocation (Fig. 3, B and C, bottom). Thus, in the absence of [Ca²⁺], signals, the receptor-induced PKCα translocation no longer precedes the membrane recruitment of PKCe. Furthermore, the weak and delayed translocation of PKCα–YFP in BAPTA-loaded cells indicates that PKCα may bind to DAGs with a lower affinity than the novel PKC isoform.

A possible Ca²⁺-controlled competition for the membrane anchorage of classical and novel PKCs was tested by coexpressing CFP- and YFP-tagged PKC fusion proteins and altering the intracellular Ca²⁺ concentrations. In HEK cells coexpressing PKCα–CFP and PKCe–YFP, maximal stimulation of a coexpressed histamine receptor induced a similar translocation of PKCα (Fig. 4 A) as in cells that expressed PKCα alone (Fig. 3 B). The histamine-induced translocation of PKCα, however, was markedly delayed, and the maximal membrane attachment was reduced in cells that coexpressed PKCα as compared with cells that were not cotransfected with PKCα (Fig. 3 C; Fig. 4 A). In PKCα/PKCe-coexpressing cells, a pretreatment with BAPTA-AM reversed the selectivity of the receptor-induced PKC translocation. The histamine-induced translocation of PKCα was further suppressed, whereas the membrane association of coexpressed PKCe now remained as efficient as in cells that only expressed the PKCe fusion protein (Fig. 4 B as compared with Fig. 3, B and C). In another experiment, we coexpressed the H₁ receptor together with PKCe–CFP and only trace amounts of PKCα–YFP (molar ratio between PKCe–CFP and PKCα–YFP ∼5:1). Under these conditions, the histamine-induced translocation of PKCe–CFP displayed identical efficiencies and kinetics as shown for PKCe–YFP without cotransfected PKCα (Fig. 3 C).

The competitive effect was not restricted to stimulation via the coexpressed H₁ histamine receptor. When PKCα–CFP/PKCe–YFP-cotransfected HEK cells were stimulated via a coexpressed human EGF receptor (EGF, 50 ng/ml), PKCα was efficiently translocated whereas PKCe remained mostly cytosolic, and BAPTA loading again reversed the selectivity of PKC translocation (unpublished data). In PKCα/PKCe-coexpressing cells that were stimulated with submaximal effective concentrations of carbachol (20 μM)
Figure 5. Differential DAG binding affinity of PKCα and PKCε in living cells. Fluorescent PKC fusion proteins were transiently transfected in HEK cells and challenged with various concentrations of the membrane-permeable DOG. (A) Confocal images of a PKCε–YFP-expressing cell taken before and 60 s after application of DOG (300 μM), and fluorescence intensity profiles were recorded over the cytoplasm and the adjacent plasma membrane of single cells. (B) The mean cytosolic and plasma membrane fluorescence intensities were recorded for either PKCα (open symbols) or PKCε (filled symbols), as shown in A, and expressed as ratios. Data represent means ± SEM of five independent experiments (each experiment comprises averaged data of three to five cells).

via an endogenously expressed M₁ family muscarinic receptor, a smaller and more short-lived [Ca²⁺], signal indicates a weaker input into the PLC signaling pathway as compared with the full stimulation of a coexpressed H₁ histamine receptor (Fig. 4, A and C). Under these conditions, the translocation of PKCα–CFP was still evident, whereas translocation of PKCε–YFP was poorly detectable. When carbachol-induced [Ca²⁺], transients were buffered by intracellular BAPTA, PKCε–YFP was translocated to the plasma membrane although the stimulation via the endogenous receptor resulted in a more delayed and weaker translocation as compared with the stimulation of the exogenously transfected H₁ receptor (Fig. 4, B and D). In the presence of [Ca²⁺], signals, the highly efficient membrane association of classical PKC isoforms may occupy the accessible DAG molecules and thereby displace novel PKC isoforms. In BAPTA-loaded cells, however, the remaining driving force for the histamine-, carbachol-, or EGF-induced PKC translocation of classical or novel PKCs is given by the formation of endogenous DAG species by PLC-β or -γ isoforms. Because PKCε

Figure 6. PKCα and PKCε compete for DAG binding in a Ca²⁺-dependent manner. (A) HEK cells were transiently cotransfected with PKCα–CFP (shown in blue) and PKCε–YFP (yellow channel), loaded with caged Ca²⁺ (o-nitrophenyl-EGTA, 10 μM), and imaged by confocal time-lapse microscopy. Images before (1) and 3 min after (2) the addition of the membrane-permeable DOG (10 μM) are shown. Subsequently, caged Ca²⁺ was photolyzed by applying a brief pulse of maximal laser energy at 364 nm (3). Bar, 10 μm. (B) Kinetic analysis of the whole experiment shown in A. Fluorescence intensities over the plasma membrane and the cytosol are expressed as ratios and SD of four cells. Comparable cell groups were allocated on the same coverslip and subjected to photolysis of caged Ca²⁺ (as indicated by the flash symbols) at later time points. (C) Similar experiment as in B, but with separately transfected PKCα–CFP and PKCε–YFP so that each cell expresses either PKCα–YFP or PKCε–CFP. (D) Equivalent experiment as in C, but without previous addition of DOG. Representatives of three independent experiments with similar results are shown.

selectively translocates to the plasma membrane under these conditions, we conclude that the novel PKC isoenzyme binds to endogenous DAG species with a higher affinity than the coexpressed PKCα.

This concept is corroborated by the finding that low concentrations (20–100 μM) of the membrane-permeable dioctanoyl-sn-glycerol (DOG) selectively recruited PKCε to the plasma membrane, whereas PKCα required ~10-fold higher DOG concentrations for a comparable plasma membrane association (Fig. 5). The concentration-dependent plasma membrane association was assessed by confocal measurement of transcellular concentration profiles of PKC isoforms 60 s after the addition of DOG (Fig. 5 A). The EC₅₀ for the DOG-induced membrane association of PKCε–YFP was ~90 μM, and saturation was observed at concentrations >300 μM. The fluorescent PKCα fusion protein accumulated in the plasma membrane only at DOG concentrations >200 μM (Fig. 5 B). Thus, in living cells, the affinity of PKCε to bind either a membrane-permeable DAG analogue or the endogenous DAG species formed by PLC-β or -γ is indeed higher than that of the classical PKCα.
When PKCe–YFP/PKCa–CFP-coexpressing cells were incubated with 20 μM DOG for >1 min, PKCe slowly accumulated in the plasma membrane, indicating that the number of accessible DOG molecules was just sufficient to anchor the available PKCe–YFP molecules. The subsequent photolysis of caged Ca\(^{2+}\) (intracellularly loaded α-nitrophenyl-EGTA) resulted in an immediate membrane translocation of PKCa. Coincident with the accumulation of the classical PKC in the plasma membrane, the DOG-bound PKCe–YFP was displaced by >50% (Fig. 6, A and B). The observed displacement of PKCe became less efficient after extended coincubation with DOG, although PKCa–CFP was efficiently translocated in response to Ca\(^{2+}\) pulses (Fig. 6 B, middle and right). Similarly, a displacement of PKCe by Ca\(^{2+}\)-driven membrane association of PKCa did not occur when high DOG concentrations (100–200 μM) were applied (unpublished data). We conclude that the Ca\(^{2+}\)-driven displacement of DAG-bound PKCe is restricted to situations in which the number of accessible DAG molecules in the inner leaflet of the plasma membrane is not in vast excess over the number of PKC molecules that compete for the DAG binding. Because a pretreatment with bisindolylmaleimide I (1 μM for 15 min) did not prevent the displacement of DOG-bound PKCe by coexpressed PKCa (unpublished data), the competitive effect does not rely on the catalytic activity of coexpressed PKC isoforms.

To monitor the localization of individually expressed PKC isoforms during photolysis of caged Ca\(^{2+}\), PKCe–CFP and PKCa–YFP were separately transfected into different cell populations and were then mixed and seeded on one coverslip. In cells that expressed only PKCe–CFP, the novel isoform again slowly translocated upon addition of DOG (20 μM), but a photolysis of Ca\(^{2+}\) failed to displace the DOG-bound PKCe (Fig. 6 C). The Ca\(^{2+}\) pulse, however, was sufficient because a robust translocation could be observed in PKCa–YFP-expressing cells located within the same visual field (Fig. 6 C). Thus, the competitive effect depends on a Ca\(^{2+}\)-driven membrane translocation of PKCa within the same cell. In the absence of exogenous DOG, the Ca\(^{2+}\) pulse selectively induced a translocation of PKCa–CFP and did not affect the localization of PKCe–YFP (Fig. 6 D).

Up to this point, the competition for DAG binding was observed at high PKC expression levels. This competitive mechanism may also operate at more physiological conditions with lower PKC expression levels and weaker PLC stimulation. We therefore generated a stably transfected HEKPKCe–YFP cell line that stably expresses low levels of PKCe–YFP (HEKPKCe–YFP) was generated and transfected with the H1 histamine receptor. Cells were stimulated with either histamine (100 μM) or carbachol (100 μM; acting via an endogenous muscarinic receptor) as indicated. The cell membrane association of PKCe–YFP was assessed before (open bars) and 20 s after agonist application (black bars) by confocal line-scan microscopy as described in Fig. 5. Data of three to six cells per experiment were averaged and expressed as means and SEM of four independent experiments. (B) Carbachol (100 μM)-induced [Ca\(^{2+}\)], signals were determined in fura-2–labeled single HEKPKCe–YFP cells (gray lines). The mean [Ca\(^{2+}\)] is superimposed (black line). (C) HEKPKCe–YFP cells were again transfected with PKCa–CFP. An endogenous muscarinic receptor was stimulated with carbachol (CCb; 100 μM) and PKC translocation was assessed as described in Fig. 3. The maximal translocation of PKCe–YFP in response to the agonist was normalized to cells that did not express additional PKCa–CFP (open bar). Molar ratios of PKCa and PKCe expression were calculated from the fluorescence intensities of the differently tagged PKC fusion proteins in single cells, and cells were grouped as indicated. Data represent means and SEM of seven independent transfection and imaging experiments.

Figure 7. PKCa concentration dependently suppresses the receptor-induced translocation of stably expressed PKCe. (A) A HEK cell line that stably expresses low levels of PKCe–YFP (HEKPKCe–YFP) was generated and transfected with the H1 histamine receptor. Cells were stimulated with either histamine (100 μM) or carbachol (100 μM; acting via an endogenous muscarinic receptor) as indicated. The plasma membrane association of PKCe–YFP was assessed before (open bars) and 20 s after agonist application (black bars) by confocal line-scan microscopy as described in Fig. 5. Data of three to six cells per experiment were averaged and expressed as means and SEM of four independent experiments. (B) Carbachol (100 μM)-induced [Ca\(^{2+}\)], signals were determined in fura-2–labeled single HEKPKCe–YFP cells (gray lines). The mean [Ca\(^{2+}\)] is superimposed (black line). (C) HEKPKCe–YFP cells were again transfected with PKCa–CFP. An endogenous muscarinic receptor was stimulated with carbachol (CCb; 100 μM) and PKC translocation was assessed as described in Fig. 3. The maximal translocation of PKCe–YFP in response to the agonist was normalized to cells that did not express additional PKCa–CFP (open bar). Molar ratios of PKCa and PKCe expression were calculated from the fluorescence intensities of the differently tagged PKC fusion proteins in single cells, and cells were grouped as indicated. Data represent means and SEM of seven independent transfection and imaging experiments.

From the positive local feedback of InsP\(_3\) and Ca\(^{2+}\) at InsP\(_3\) receptors, the reduced input into the PLC signaling cascade was still sufficient to generate large [Ca\(^{2+}\)]\(_i\) signals with peak values of 300–700 nM (Fig. 7 B), which clearly exceed threshold values for the Ca\(^{2+}\)-driven translocation of PKCa. HEKPKCe–YFP cells were again transfected with various amounts of a PKCa–CFP-encoding cDNA plasmid. Furthermore, the molar ratios between stably expressed PKCe–YFP and transiently expressed PKCa–CFP were determined at the single-cell level. The transient coexpression of PKCa–CFP concentration-dependently suppressed the carbachol-induced plasma membrane docking of PKCe–YFP. This reduction was ~70% at an ~1:1 molar ratio, and saturation of the inhibitory effect was evident at a threefold molar excess of PKCa–CFP over PKCe–YFP (Fig. 7 C).
Cells were lysed in either Ca\textsuperscript{2+}-activator PMA (10 μM) or ionomycin (iolo; 10 μM, 2 min), carbachol (CCh; 100 μM, 15 s) or PMA (10 μM, 5 min). Cells were lysed in either Ca\textsuperscript{2+}-free (nominally Ca\textsuperscript{2+}-free buffer supplemented with 0.5 mM EGTA) or Ca\textsuperscript{2+}-containing (1 mM) extraction buffers, however, the classical PKC isoform was mainly found in the particulate fraction, even in Ca\textsuperscript{2+}-free buffer. Although imaging experiments were instrumental in defining the competitive translocation mechanism, translocation of endogenous PKCs is commonly quantified by Western blot analysis of PKCs in soluble and particulate cell fractions. The moderate appearance of PKC\textsubscript{α} when extracts were prepared in Ca\textsuperscript{2+}-free buffer resulted in a weak redistribution to the particulate fraction. The local increase of PKC\textsubscript{α} in vascular smooth muscle cells is controlled in a Ca\textsuperscript{2+}-dependent manner. The translocation of endogenously expressed PKC\textsubscript{α} and PKC\textsubscript{ε} as indicated. PMA stimulation (10 μM, 5 min) was included as a control.

**Figure 8.** Extraction of PKC\textsubscript{α}–YFP in the absence or presence of Ca\textsuperscript{2+}. HEK cells were stably transfected with YFP-fused PKC\textsubscript{α}. After treating cells with either buffer (+), ionomycin (iolo; 10 μM, 2 min), carbachol (CCh; 100 μM, 15 s) or PMA (10 μM, 5 min). Mean fluorescence and SEM of four independent experiments. Data represent means and SEM of four independent extraction experiments.

Table 1

|          | AVP | PMA | AVP |
|----------|-----|-----|-----|
| 0″       | -   | -   | -   |
| 10″      | -   | +   | +   |
| 30″      | +   | +   | +   |

**Figure 9.** The translocation of PKC\textsubscript{α} and PKC\textsubscript{ε} in vascular smooth muscle cells is controlled in a Ca\textsuperscript{2+}-dependent manner. Neonatal rat AoSMCs were pretreated with or without BAPTA-AM and subsequently stimulated with AVP (1 μM) for the indicated times. Particulate and soluble fractions were probed for endogenously expressed PKC\textsubscript{α} and PKC\textsubscript{ε} as indicated. PMA stimulation (10 μM, 5 min) was included as a control.

**Discussion**

Coinaging of fluorescent fusion proteins revealed that a selective plasma membrane recruitment of classical and novel PKC isoforms can result from a Ca\textsuperscript{2+}-controlled competitive binding to the lipid second messenger DAG. Without additional [Ca\textsuperscript{2+}], signals, PKC\textsubscript{ε} displayed a higher DAG binding affinity than PKC\textsubscript{α}, resulting in a selective plasma membrane translocation of the novel PKC isoform. Contrast, in the presence of [Ca\textsuperscript{2+}], signals, classical PKCs bind to the plasma membrane with a diffusion-limited kinetic. Being redistributed to a two-dimensional plasma membrane compartment, the local increase of PKC\textsubscript{α} concentration within the target compartment is sufficient to occupy a majority of accessible DAG molecules and thereby prevents or displaces the DAG binding of novel PKCs. The Ca\textsuperscript{2+}-controlled competitive translocation of classical versus novel PKC iso-

![Graph](image-url)
forms was also detectable at physiological expression levels. This mechanism may enable a cell to decode stimuli by a directed membrane translocation and activation of distinct PKC isoforms.

Imaging of fluorescent PKC fusion proteins has provided a more realistic impression of the dynamics of these signaling molecules (Sakai et al., 1997; Feng and Hannun, 1998; Feng et al., 1998; Almholt et al., 1999; Codazzi et al., 2001). Most strikingly, the repetitive translocation of classical PKCs during oscillatory [Ca\(^{2+}\)], responses was only detectable using these techniques (Oancea and Meyer, 1998; Dale et al., 2001; Schaefer et al., 2001). Currently available data on sub-type-specific translocations of PKC isoforms predominantly rely on Western blot analyses of particulate and soluble cell fractions. This approach, however, does not necessarily reflect the distribution of PKCs at a given time point. In particular, during subcellular fractionation, classical PKCs may lose their membrane attachment (Kiley et al., 1990; Fig. 8). A secondary loss of membrane contact of classical PKCs during fractionation may result from (a) release of classical PKCs that are bound solely through a Ca\(^{2+}\)-dependent interaction with phosphatidylinositol and/or (b) a low affinity of classical PKCs for DAG binding, which is not sufficient to maintain the membrane-bound state if additional stabilization by Ca\(^{2+}\) is lost. The contribution of both mechanisms may be indirectly inferred from the capability of Ca\(^{2+}\)-bound classical PKCs to prevent the DAG binding of the novel isoforms. Indeed, when [Ca\(^{2+}\)] was clamped to resting concentrations (50–80 nM), translocation of PKC\(\alpha\) was impaired whereas PKC\(\varepsilon\) appeared to access DAGs more easily.

In living cells, several lines of evidence point to a higher DAG binding affinity of PKC\(\varepsilon\) as compared with PKC\(\alpha\). Low concentrations of membrane-permeable DOG preferentially recruited PKC\(\varepsilon\) to the plasma membrane. Second, receptor stimulation in BAPTA-loaded cells effectively translocated PKC\(\varepsilon\), but not coexpressed PKC\(\alpha\), and, in vascular smooth muscle cells, PMA treatment resulted in a complete translocation of endogenous PKC\(\varepsilon\), whereas the redistribution of PKC\(\alpha\) was only partial. The DAG binding of recombinantly expressed and purified PKC isoenzymes has been determined using phorbol esters as DAG surrogates. Both absolute and relative binding affinities of different PKC isoforms, however, strongly depend on the lipid composition and the dispersion technique to generate artificial membranes (Kazanietz et al., 1993; Dimitrijevic et al., 1995). Therefore, reconstitution experiments do not necessarily reflect the DAG binding of PKC isoforms in living cells. Despite its inability to estimate absolute affinities, simultaneous imaging of PKC isoforms in living cells can be applied to compare the DAG binding of different PKC isoenzymes within their physiological environment. Another advantage of imaging versus extraction experiments is the coverage of the entire time course of a receptor-induced PKC redistribution. Co-imaging of classical and novel PKCs now revealed that the translocation of the classical isoforms precedes the maximum of the membrane anchorage of novel isoforms. For a given cell type and agonist, the observation of selective translocations of PKC isoforms is, therefore, not only influenced by extraction techniques but also by the choice of the incubation time.

The Ca\(^{2+}\)-controlled competitive translocation of classical and novel PKCs represents a mechanism by which signal diversification may emerge downstream of receptor activation. Although stimulation of PLC-coupled receptors commonly induces both [Ca\(^{2+}\)], and DAG signals, their relative contribution to the PKC translocation underlies additional regulation. The catalytic activity of phosphoinositide-specific PLCs cleaves PIP\(_2\) to form equimolar amounts of DAGs and InsP\(_3\). In contrast to the direct interaction of DAGs with PKCs, InsP\(_3\) action additionally depends on the filling state and responsiveness of internal Ca\(^{2+}\) stores. Refractory Ca\(^{2+}\) stores have been demonstrated to result from InsP\(_3\) receptor downregulation (Wojcikiewicz and Nahorski, 1991; Bokkala and Joseph, 1997; Willars et al., 2001) or from a cGMP-dependent phosphorylation of IR\(\alpha\)G, an InsP\(_3\) receptor-associated cGMP kinase substrate (Schlossmann et al., 2000). Furthermore, owing to the biphasic effect of Ca\(^{2+}\) and ATP on InsP\(_3\) receptor gating, InsP\(_3\) can only operate at certain cytosolic conditions (Mak et al., 1998, 1999).

A competition for DAG binding of classical and novel PKCs requires that the number of accessible DAG molecules is limiting. Our data demonstrate that the Ca\(^{2+}\)-induced displacement of DOG-bound PKC\(\varepsilon\) by PKC\(\alpha\) is restricted to low concentrations of the membrane-permeable DAG analogue. The number of DAG molecules that are formed upon receptor stimulation can be roughly estimated. The InsP\(_3\) receptor is fully activated in the presence of 10\(^{-8}\) to 10\(^{-7}\) M InsP\(_3\) (Mak et al., 1998), which corresponds to a number of 12,000–120,000 molecules in a given cell of 2 picoliter volume. In various cell types, phorbol ester binding experiments have revealed a number of 40,000–800,000 binding sites per cell (Trilivas and Brown, 1989; Obeid et al., 1990; Combadière et al., 1993). Although these binding sites presumably include other DAG receptors, such as PKD, chimaerin, Munc13-1, or Ras-GRP, the total number of DAG acceptors, including classical and novel PKCs, appears not to be in vast excess over the number of DAGs formed during receptor stimulation.

From measurements of the total cellular DAG, one can calculate that a single cell contains 5 \times 10^3 to 10^8 DAG molecules (Kiley et al., 1991; Baldassare et al., 1992). The high basal values and the modest increases upon receptor stimulation indicate that a major fraction of the total DAG content cannot be accessible for PKCs. A more detailed analysis has revealed that in resting cells, DAGs with saturated or monounsaturated fatty acids are predominant, whereas PLC activity mainly forms polyunsaturated DAG species, and the hypothesis that distinct DAG species, rather than changes in the total DAG mass, regulate PKC has been raised (Eskildsen-Helmond et al., 1998; Ivanova et al., 2001). A delayed formation of DAGs in the absence of [Ca\(^{2+}\)], signals has been attributed to the catalytic activities of phosphatidylincholine-specific PLCs or PLDs. Whether this second wave of DAG formation leads to PKC activation is still under debate (Pettitt et al., 1997). Furthermore, PKC, via a noncatalytic pathway, is regarded as an activator of PLD activity rather than a target of PLD-induced DAG formation (Singer et al., 1996; Zhang et al., 1999). Consistent with the idea that second messengers formed by PLC activity may be scavenged by heterologously expressed
proteins, a modified InsP₃ receptor fragment, the “InsP₃ sponge,” has only recently been introduced as a transfectable tool that intracellularly captures InsP₃ and prevents the agonist-induced Ca²⁺ mobilization from internal stores (Uchiyama et al., 2002). Kinase-dead PKC isoforms have been considered as dominant negative modulators of endogenous pathways (Ohba et al., 1998; Matsumoto et al., 2001; Braz et al., 2002). Although competition of kinase-dead and wild-type PKCe for receptor for activated C-kinases binding has been demonstrated (Pass et al., 2001), the dominant negative mechanism is far from being clarified. Our data suggest that dominant negative effects of overexpressed PKCs may also result from scavenging the accessible DAG molecules.

In contrast to experimental settings, physiological responses to hormones or paracrine stimuli have to decode slight changes in agonist concentrations that cause only sub-maximal receptor occupancy and weaker DAG formation. Therefore, competitive processes between DAG-binding signaling molecules are likely to occur in vivo. The highly efficient Ca²⁺-driven plasma membrane association of classical PKCs and the superior DAG binding affinity of the novel PKCe represent a finely tuned system where [Ca²⁺], serves as a switch for the subtype-selective activation of PKCs. As a consequence, the competitive and/or sequential isotype-specific PKC activation is one of the mechanisms through which specific physiological responses may emerge from the signaling network of PKCs.

Materials and methods

Cell culture and transfections

HEK293 and primary cultures of rat embryonic AoSMCs (Reusch et al., 2001) were grown in MEM (Biochrom) supplemented with 10% FCS, penicillin (50 U/ml), and streptomycin (50 U/ml). Cells were seeded on 6-cm dishes and transfected the following day at 50–70% confluence with a FuGene 6 transfection reagent (Roche Molecular Biochemicals) and 4 μg total plasmid cDNA per well according to the manufacturer’s recommendations. After 18–24 h, cells were trypsinized and seeded on glass coverslips. All experiments were conducted 48–72 h after transfection. For selection and maintenance of the stably transfected HEK293-GFP cell line, the culture medium was supplemented with G418 (750 μg/ml). Vascular smooth muscle cells were starved in serum-free culture medium for 1 d before the experiments.

Generation of fluorescent PKC fusion proteins and fluorescence imaging

PKCs, β₁, and ε were COOH-terminally fused to YFP in a custom-made pcDNA3-YFP fusion plasmid as described earlier (Schafer et al., 2001). CFP fusions were generated analogous to the YFP-fused PKCs but using a custom-made pcDNA3-CFP vector that contains the open reading frame of enhanced CFP (Invitrogen) instead of enhanced YFP. Fluorescence imaging was performed with a monochromator and a cooled CCD camera ( Till-Photronics) connected to an inverted epifluorescence microscope (Axiovert 100; Carl Zeiss MicroImaging, Inc.). A 505-nm dichroic mirror (inflection point) with extended reflectivity (300–500 nm) was combined (TILL-Photonics) connected to an inverted epifluorescence microscope and a 63x/1.4 Plan-Neofluor objective (Carl Zeiss MicroImaging, Inc.). CFP was excited with the 458-nm line of an argon laser, and emitted light was collected with a 470–500-nm band pass filter. YFP was excited with the 488-nm laser line, and emission was recorded with a 530–560-nm band pass filter. For photolysis of caged Ca²⁺, cells were loaded with 10 μM α-nitrophenyl-EGTA-AM (Molecular Probes; 40 min at 25°C), washed, and incubated for another 30 min in HBS before the experiment. Caged Ca²⁺ was photolyzed in defined areas by briefly switching the 364-nm line of an argon laser to maximal intensity. Pinholes were adjusted to yield optical sections of 0.7–1.4 μm.

Regression-based spectral evaluation of multiple fluorochromes

At any combination of excitation and emission wavelengths λ, the background-corrected signal of the probe Fᵢ is additively composed of fluorochromes emitted from m different dyes Fᵢ,c. Because Fᵢ,c scales with both the unknown relative concentration of the dye cᵢ and its normalized fluorescence intensity at the chosen wavelength settings Xᵢ, the term Fᵢ = Cᵢ,Xᵢ,cᵢ dissect known (cᵢ) and unknown (Xᵢ,cᵢ) parts. The signal of the probe can be described as Fᵢ = ΣΣXᵢ,cᵢ,Fᵢ,cᵢ (Fig. 1 A). The determination of fluorescence intensities of the probe at n different combinations of excitation and/or emission wavelengths provides n independent solutions for Fᵢ. The resulting linear systems of equations can be solved if the number of different spectral settings n equals or exceeds the number of unknown relative dye concentrations cᵢ,cᵢ (provided that linear dependence between the solutions can be excluded). For best reliability, we solved overdetermined (n > m) linear systems of equations by means of multivariate, constrained (cᵢ,cᵢ ≥ 0), linear regression analyses:

\[
\begin{align*}
C &= \begin{bmatrix} c_1 & \cdots & c_m \end{bmatrix} \\
F &= \begin{bmatrix} F_1 \\
\vdots \\
F_n \end{bmatrix} \\
X &= \begin{bmatrix} X_{11} & \cdots & X_{1m} \\
\vdots & \ddots & \vdots \\
X_{n1} & \cdots & X_{nm} \end{bmatrix} \\
\text{As a result, the relative concentrations of free (cᵢ,cᵢ) and the Ca²⁺-bound fura-2 (cᵢ,cᵢ=2) are known, and interfering signals of CFP variants are eliminated. Therefore, the equation } [Ca²⁺] = 224 nM \times C_{Ca²⁺}\text{ is applied to calibrate } [Ca²⁺]. \text{ The regression analysis was repeated for each image pixel at any time point resulting in a spatially and/or temporally resolved calibration of the } [Ca²⁺], \text{ and in a detection of signals arising from coexpressed CFP- and YFP-fused proteins. Furthermore, the molar ratio between CFP- and YFP-tagged fusion proteins can be assessed at the single-cell level. Based on the equimolar expression of CFP and YFP in an intramolecularly fused CFP-YFP tandem protein, a compensation for the lower fluorescence intensity of CFP could be calibrated. In our optical system used for imaging of } [Ca²⁺], \text{ CFP, and YFP, the fluorescence of YFP (Fᵢ) appeared 8.1-fold brighter than for the intramolecularly coupled CFP (cᵢ,cᵢ=2) as detected after computerized disruption of fluorescence resonance energy transfer by YFP photobleaching). The term cᵢ,cᵢ=2×1/cᵢ,cᵢ corresponds to the molar ratio between coexpressed YFP- and CFP-fused proteins.}
\end{align*}
\]

Cell fractionation and immunoblotting

For detection of PKCs in cytosolic and particulate fractions, cells were homogenized by two 5-s pulses of sonification (Branson sonifier) in an ice-cold lysis buffer containing 25 mM Tris/HCl, pH 7.5, 2 mM EDTA, 0.5 mM EGTA, 1 mM DTT, 200 μM phenylmethylsulfonyl fluoride and leupeptin, aprotinin (20 μg/ml each). Particulate material was removed (30,000 × g, 20 min at 4°C), and supernatants representing the soluble PKC fractions were collected. The pellets were resuspended in lysis buffer supplemented with...
1% (wt/vol) Triton X-100, sonified (5 s), and placed on a rocking platform (20 min at 4°C). Supernatants of a second centrifugation (30,000 g, 20 min at 4°C) were assigned as particulate PKC fraction. For immunoblotting, samples were diluted in Laemmli buffer containing 10% β-mercaptoethanol and subjected to a 10% SDS-PAGE. Separated proteins were electroblotted on nitrocellulose membranes, blocked in PBS containing 5% nonfat dry milk, and probed with polyclonal rabbit anti-PKCα or anti-PKCε antisera (1:1,000 in PBS; BD Biosciences) and a secondary, peroxidase-coupled anti-rabbit IgG antibody (Sigma-Aldrich). Chemiluminescence was detected with a Lumiglo reagent (New England Biolabs, Inc.).

The expression levels of endogenously expressed PKC isozymes and of heterologously expressed PKC fusion proteins were assessed in cytosolic fractions prepared from quiescent cells. 20 μg of protein or diluted samples of the transiently overexpressing cells were loaded on the gel. The transfection efficiency in these experiments was >50%, as detected by counting the total number and number of fluorescent cells.

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