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Single nucleotide polymorphisms alter kinase anchoring and the subcellular targeting of A-kinase anchoring proteins

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A-kinase anchoring proteins (AKAPs) shape second-messenger signaling responses by constraining protein kinase A (PKA) at precise intracellular locations. A defining feature of AKAPs is a helical region that binds to regulatory subunits (RII) of PKA. Mining patientderived databases has identified 42 nonsynonymous SNPs in the PKA-anchoring helices of five AKAPs. Solid-phase RII binding assays confirmed that 21 of these amino acid substitutions disrupt PKA anchoring. The most deleterious side-chain modifications are situated toward C-termini of AKAP helices. More extensive analysis was conducted on a valine-to-methionine variant in the PKA-anchoring helix of AKAP18. Molecular modeling indicates that additional density provided by methionine at position 282 in the AKAP18y isoform deflects the pitch of the helical anchoring surface outward by 6.6°. Fluorescence polarization measurements show that this subtle topological change reduces RII-binding affinity 8.8-fold and impairs cAMP responsive potentiation of L-type Ca2+ currents in situ. Livecell imaging of AKAP187 V282M-GFP adducts led to the unexpected discovery that loss of PKA anchoring promotes nuclear accumulation of this polymorphic variant. Targeting proceeds via a mechanism whereby association with the PKA holoenzyme masks a polybasic nuclear localization signal on the anchoring protein. This led to the discovery of AKAP18E: an exclusively nuclear isoform that lacks a PKA-anchoring helix. Enzyme-mediated proximity-proteomics reveal that compartment-selective variants of AKAP18 associate with distinct binding partners. Thus, naturally occurring PKA-anchoringdefective AKAP variants not only perturb dissemination of local second-messenger responses, but also may influence the intracellular distribution of certain AKAP18 isoforms.

AKAPs | PKA | proximity labeling | kinase anchoring | nucleus

The recent explosion of deep-sequencing data from diverse cohorts of patients reveals the impact of genetic variation within human populations (1). Nucleotide changes that accumulate over multiple generations are considered inert or "passenger" mutations that shape us as individuals. Conversely, acute genetic lesions that alter the activity of key enzymes or structural proteins are often linked to disease. Such pathological lesions are particularly prevalent in cell-signaling enzymes, such as ubiquitin ligases and protein kinases that guide the passage of chemical signals throughout the cell (2, 3). Consequently, mining of databases that chart these mutations can be combined with high-resolution structural analysis to inform the development of drugs that combat chronic disorders, such as Parkinson's disease and various cancers (4).

Early examples of this precision-medicine approach include the identification of genetically modified forms of the BCR-Abl and B-RAF kinases that drive cancers, including chronic myelogenous leukemia, melanoma, and hepatocellular carcinoma (5, 6). The subsequent success of groundbreaking drugs, such as imatinib

(Gleevec) and sorafenib, have transformed this innovative drugdiscovery strategy into lucrative multinational ventures. The current kinase inhibitor landscape includes 243 small molecules in clinical trials for a variety of indications (7). However, manipulating complex cellular events is often more involved than simply targeting individual signaling enzymes. This is exemplified by evidence that many kinase cascades are constrained by scaffolding and anchoring proteins (8–10). These organizational elements ensure that signals are transmitted to precise sites within the cell (11). Hence, genetic lesions that disrupt protein kinase anchoring have been linked to various pathological responses and may represent untapped therapeutic targets (12–14).

A-kinase anchoring proteins (AKAPs) are prototypic kinase-targeting factors (15, 16). This structurally diverse group of proteins was initially discovered on the basis of solid-phase interaction with radiolabeled RII subunits of the cAMP-dependent protein kinase (PKA) (17). Subsequent interaction-cloning, proteomic, and bio-informatic approaches have identified 39 human genes that encode in excess of 60 AKAP isoforms (11) (Fig. 14). AKAPs were

Significance

Dissemination of chemical information throughout the cell is a fundamental biological process with clinical relevance. Pathological changes in local signaling enzyme activity are linked to diseases, including schizophrenia, Alzheimer's disease, cardiac arrhythmias, and seizures. Mining of patient datasets has uncovered genetic variation in A-kinase anchoring proteins (AKAPs) that promotes mislocalization of protein kinase A (PKA). We investigate 42 SNPs in AKAPs that interrupt association with PKA to impact local cAMP signaling. The most detrimental variants are situated within the hydrophobic face of a conserved helical region on the AKAP that is essential for kinase anchoring. An unexpected outcome is the discovery of an alternative targeting mechanism for AKAPs that utilizes the intact PKA holoenzymes as cytoplasmic "anchors."

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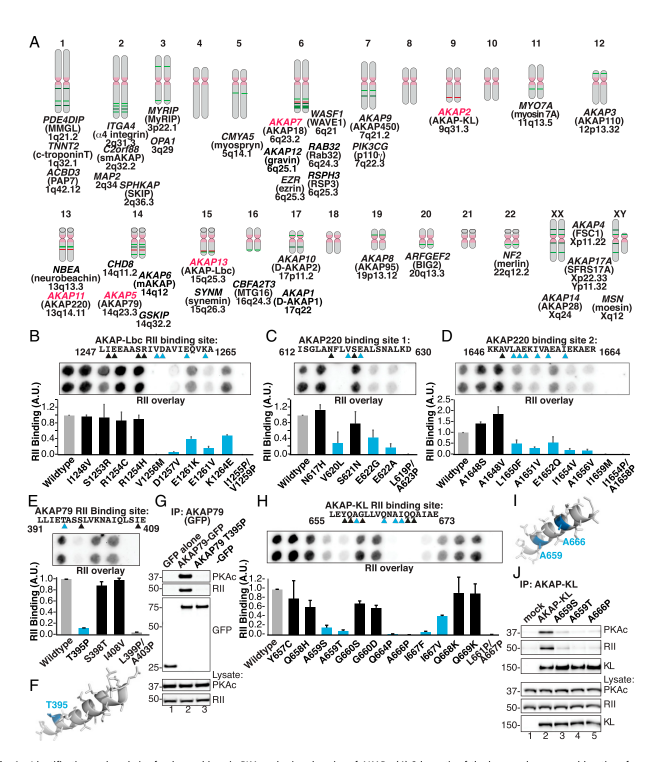


Fig. 1. Identification and analysis of polymorphisms in PKA-anchoring domains of AKAPs. (A) Schematic of the human chromosomal locations for genes encoding 39 experimentally validated AKAPs. AKAPs harboring nonsynonymous SNPs in the PKA-anchoring helices that are investigated in this study are highlighted in red. (B-E and H) Peptide SPOT arrays containing WT and variant PKA binding helices were constructed and probed by RII overlay. (Upper) Solidphase RII binding was assessed by far-Western blotting with biotin-RII and detected by neutravidin-HRP and enhanced chemoluminescence. (Lower) Densitometric analysis of RII binding was normalized to WT signal. (B) Peptide array and amalgamated densitometric data for analysis of the AKAP-Lbc (AKAP13) RII-binding site. Densitometric analysis of RII binding was normalized to WT signal. Variants that reduce binding to RII are highlighted in blue. (C) Screening and analysis of peptide arrays representing SNP sequences for PKA binding site 1 in AKAP220 (AKAP11). (D) Screening and analysis of peptide arrays representing SNP sequences for the PKA binding site 2 in AKAP220 (AKAP11). (E) Screening and analysis of peptide arrays representing SNP sequences for the PKA binding site in AKAP79 (AKAP5). (F) Molecular modeling of the AKAP79 amphipathic helix (PDB ID code 2H9R) showing threonine 395 at the second turn of the helix (blue). (G) Immune complexes of either WT AKAP79 or AKAP79 T395P were probed for coprecipitation of PKA RII and C subunits. Introduction of the helix-breaking proline residue disrupts the AKAP79-PKA interaction. (H) Screening and analysis of peptide arrays representing SNP sequences for the PKA binding in AKAPKL (AKAP2). (I) Modeling of the AKAP-KL amphipathic helix based on coordinates for AKAP-IS (PDB ID code 2IZX), showing the residues A659 and A666 (blue). (J) Immune complexes of either WT AKAP-KL or AKAP-KL A659S, A659T, or A666P variants were probed for coprecipitation of PKA RII and C subunits. Mutation of either position drastically impairs PKA binding. IP, immunoprecipitation

originally named colloquially or on the basis of their molecular weight, but the systematic cataloging of anchoring proteins has been complicated by a HUGO nomenclature that only recognizes 15 AKAP genes (Fig. 1A). Nonetheless, a ubiquitous and defining characteristic of the AKAP family is a 14- to 18-residue helical region that binds with high affinity to PKA holoenzymes that are composed of a regulatory (R) subunit dimer and two catalytic (C) subunits (18). X-ray crystallographic analyses have shown that the hydrophobic face of this amphipathic helix slots into a binding groove formed by the docking and dimerization domain (D/D) of R subunits (19, 20). Negative-stain electron microscopy and 3D reconstructions show that intact hetero-pentameric AKAP-RII₂-C₂ assemblies adopt a range of flexible tripartite configurations (21, 22). This is because intrinsically disordered regions between the D/ D domains and cAMP-binding sites on each regulatory subunit permit a 150- to 200-Å radius of motion to the associated catalytic subunits (21, 23). Such flexibility within anchored PKA holoenzyme complexes enables precise orientation of the catalytic subunit toward substrates (11, 24). Recent data advances this model, showing that intact, active, and anchored PKA holoenzymes are the functional signaling units inside cells (22). This "signaling island" concept radically changes the view of how AKAP complexes operate and indicates that protein phosphorylation is more regionally confined than previously appreciated (22).

As our understanding of how signaling enzyme complexes operate becomes more refined, it is evident that perturbation of AKAP-PKA interactions is linked to disease (11, 14). Scrutiny of big datasets has uncovered genetic variation in AKAPs that correlate with susceptibility to diseases, such as schizophrenia, Alzheimer's, certain cardiac arrhythmias, and the onset of febrile seizures (12, 14, 25, 26). In this report we systematically investigate the properties of polymorphic variants in AKAPs that elicit single amino acid changes in RII-anchoring helices. Unexpected outcomes of this endeavor include the discovery of an alternative AKAP-targeting mechanism that utilizes the intact PKA holoenzyme as a cytoplasmic anchor and the identification of an AKAP18 variant that does not function as an AKAP.

Results

SNPs in AKAP Helices Alter PKA Anchoring. Genome-wide association surveys have identified point mutations and SNPs in genes encoding AKAPs that are linked to a variety of neurological and cardiac diseases (reviewed in ref. 14). The National Center for Biotechnology Information (NCBI) SNP (https://www.ncbi.nlm. nih.gov/SNP/index.html) and Variation Viewer databases (https:// www.ncbi.nlm.nih.gov/variation/view/) were used to identify nonsynonymous nucleotide changes that introduce missense amino acids within the R subunit binding helices from a variety of AKAPs (Fig. 14, red). This information was used to create peptide SPOT arrays that encompass native and variant PKA-binding sequences from five different anchoring proteins (Fig. 1A and SI Appendix, Table S1). Solid-phase RII-binding (RII-overlay) assessed whether these amino acid substitutions affected PKA anchoring (Fig. 1B). Peptides containing prolines were included in the peptide array as negative controls (SI Appendix, Table S1). Prior studies have shown that substitution with proline disrupts the structural integrity of PKA-anchoring helices (18, 27).

Densitometric analyses of multiple SPOT arrays revealed that amino acid substitution within PKA-anchoring helices influences RII interaction (Fig. 1 *B–J*). Nine naturally occurring variants in AKAP-Lbc (*AKAP13*) were tested (Fig. 1*B*). Single amino acid substitutions at four positions (R1245H, I1248V, S1253R, and R1254C) had no effect on RII-binding (Fig. 1*B*). However, three side-chain variants toward the C terminus of the PKA-anchoring helix (V1256M, D1257V, and E1261V) dramatically reduced solid-phase interaction with RII (Fig. 1*B*). Two other substitutions (E1261K and K1264E) had more modest effects (Fig. 1*B*).

Next we investigated SNPs in the vesicular and membrane-associated anchoring protein AKAP220 (*AKAP11*). This anchoring protein encodes two PKA-anchoring helices with different affinities for PKA holoenzymes (28, 29). Substitutions toward the C terminus of site 1 exhibited reduced affinity for RII, whereas amino acid changes throughout helix 2 adversely affected PKA anchoring (Fig. 1 *C* and *D*). Interestingly, substitution of serine or valine at position 1648 enhanced solid-phase binding of RII in site 2 of AKAP220 (Fig. 1*D*). This suggests that side chains that constrain the initial turns of helices are positive PKA-anchoring determinants.

Further support for this concept was provided by analysis of the neuronal anchoring protein AKAP79 (*AKAP5*) (30, 31). Three SNP variants in AKAP79 were investigated, but only substitution of Thr-395 with Pro affected RII-binding (Fig. 1*E*). Molecular modeling based upon the NMR structure of this region reveals that position 395 lies within the first turn of the anchoring helix (32) (Fig. 1*F*). Proline substitution prevents formation of the α-helix. Further validation was provided when this variant was introduced into full-length AKAP79 and expressed in HEK293 cells. Western blotting of WT AKAP79 immune complexes confirmed cofractionation of PKA regulatory and catalytic subunits (Fig. 1*G*, *Upper* three panels, lane 2). Cofractionation of PKA subunits was not detected in cells expressing the AKAP79-T395P variant (Fig. 1*G*, *Upper* three panels, lane 3).

AKAP-KLs (AKÂP2) are a family of alternately spliced AKAP isoforms that organize kidney and lung second-messenger signaling events (33). AKAP-KL also sustains PKA regulation of lens aquaporin-0 water channels (34). Disruption of AKAP-KLmediated PKA anchoring in the lens is implicated in pathological cataract formation (34). Peptide array analysis identified several missense variants of AKAP-KL that disrupted RII-binding (Fig. 1H). The most profound effects were observed in SNP variants that contained prolines at positions 664 and 666 in AKAP-KL. However, significant loss of PKA anchoring was also observed when positions 659 and 667 were replaced with bulkier side chains (Fig. 1H). We focused our biochemical analyses on two sites in the AKAP-KL helix that yielded severely impaired PKAanchoring variants (Fig. 11). Further validation of this result was provided when the A659S, A659T, and A666P variants were introduced into full-length AKAP-KL and expressed in HEK293 cells. Coprecipitation of PKA subunits was severely reduced or not detected in AKAP-KL immune complexes isolated from cells expressing each of the aforementioned SNP variants (Fig. 1J, Upper two panels, lanes 3-5). Control experiments with WT AKAP-KL immune complexes confirmed cofractionation of PKA regulatory and catalytic subunits (Fig. 1J, Upper two panels, lane 2). Collectively, the data presented in Fig. 1 show that naturally occurring variants on the hydrophobic face of RIIbinding helices, and particularly those located in C-terminal turns, can have profound effects on PKA anchoring.

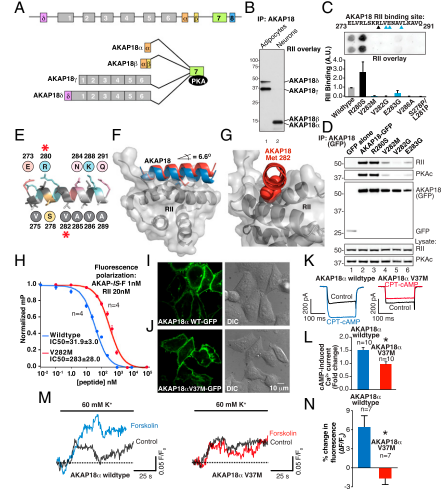
Characterization of AKAP18 Helix-Disrupting SNPs. Alternative splicing of the AKAP7 gene drives tissue-specific expression of different AKAP18 protein products that range in size from 15 to 42 kDa (35–37) (Fig. 2 A and B). The low molecular weight α and β-isoforms of AKAP18 couple PKA to a variety of ion channels in neurons, whereas the longer δ - and γ -isoforms have been implicated in modulation of excitation contraction coupling in the heart, water homeostasis in kidney collecting ducts, and roles in differentiated adipocytes (35, 38, 39) (Fig. 2B). Polymorphisms in AKAP18 have also been linked to an increased incidence of febrile seizures (26). The identification of a cluster of SNPs within the PKA-anchoring helix between residues 282 and 286 of AKAP18y provided the impetus for more indepth investigation (Fig. 2C). Peptide array analysis of SNP sequences at positions 282, 283, and 286 revealed that conservative side-chain substitutions disrupted RII-binding (Fig. 2C). Control experiments confirmed that PKA anchoring was also abolished in the context of the double-mutation AKAP18y S278/L281P (Fig. 2C). In contrast, the AKAP18γ Arg280Ser (R280S) variant appeared to strengthen the interaction with RII (Fig. 2C). More stringent cell-based validation of AKAP18y variants involved expression of the full-length proteins in HEK293 cells. Western blot analysis of AKAP18 immune complexes was used to detect cofractionation of PKA subunits (Fig. 2D, Upper panels). All variants at positions 282–286 of AKAP18y were impaired in their ability to anchor PKA (Fig. 2D, lanes 4–6). In comparison, PKA subunits remained associated with the AKAP18y R280S variant inside cells (Fig. 2D, lane 3). Secondary structure prediction algorithms place residue 280 of AKAP18y on the hydrophilic face of the AKAP helix, a region that does not directly contact the RII dimer (40) (Fig. 2E). Conversely, side chains between positions 282–286 of AKAP18y are located on the hydrophobic face of the helix (19, 40) (Fig. 2E). This substructure slots into a binding pocket formed by a four-helix bundle on the docking and dimerization domain of the R subunits (Fig. 2F, blue helix). The impaired binding in the V282M mutant provided us a unique opportunity to study this intermediate phenotype.

Molecular modeling of the AKAP18–RIIα interface using PDB coordinates from published structures predicts that the V282M substitution generates a 6.6° upward displacement of the anchoring helix in relation to the binding pocket on RIIα (Fig.

2C, red helix). More detailed examination of the V282M-RIIα interface reveals that the added bulk and length of the methionine side-chain sterically hinders association with the AKAPbinding pocket on RIIα (Fig. 2G). In silico studies using the AutoDock program (41) computed a simulated free energy of binding of -4.54 kcal/mol (estimated $K_d = 468 \mu M$) for AKAP18 V282M compared with -10.58 kcal/mol (estimated K_d = 17.4 nM) for WT. Fluorescence polarization experiments calculated an IC₅₀ of 283 \pm 28 nM (n = 4) for AKAP18V282M peptide-RIIα interaction (Fig. 2H, red) compared with an IC₅₀ of 32 ± 3 nM (n = 4) for the native sequence (Fig. 2H, blue). The differences in predicted and observed binding-affinity changes may be a consequence of AutoDock forcing all side chains into rigid conformations (41). Collectively, these data indicate that AKAP18 variants harboring the V282M substitution are severely impaired in their ability to anchor PKA holoenzymes.

Because the kinase-anchoring helix is conserved in most splice variants of AKAP18, we introduced the corresponding mutation (V37M) into the α -isoform. This membrane-associated anchoring protein mediates PKA-dependent modulation of ion channels in excitable cells (42). Fluorescent imaging confirmed that GFP-tagged AKAP18 α V37M is retained at the plasma membrane when expressed in HEK293 cells (Fig. 2 *I* and *J*). Coexpression of AKAP18 α V37M with L-type Ca²⁺ channel subunits permitted whole-cell electrophysiological analysis of local PKA modulation

Fig. 2. Polymorphisms in AKAP18 isoforms (AKAP7) alter PKA binding and regulation of ion channels. (A) Schematic of AKAP7 gene organization depicting exon-intron structure and alternatively spliced isoforms. (B) IP-RII overlay analysis show differential expression of distinct AKAP18 isoforms in 3T3-L1 adipocytes and cultured hippocampal neurons. (C) Peptide array and amalgamated densitometric data for analysis of the AKAP18 (AKAP7) RII-binding site. Densitometric analysis of RII binding was normalized to WT signal. Variants that reduce binding to RII are highlighted in blue. (D) AKAP18 SNP variants (denoted above each lane) were immunoprecipitated and copurification of PKA regulatory (Upper) and catalytic (Upper-mid) identified by Western blotting. Expression levels of each protein are indicated in the Lower three panels. (E) Model of AKAP18 PKA binding helix highlighting selected residues (red asterisk) on either face of the amphipathic helix. (F) Structural model shows WT AKAP18 helix (blue) and AKAP18 V282M helix (red) in complex with the RII D/D domain (gray). The V282M helix (red) is oriented at a 6.6° angle from WT. (G) En face and expanded view of the AKAP18 V282M helix shows displacement of the mutant peptide by bulky methionine side chain (red). (H) Fluorescence polarization data shows AKAP18 WT and V282M peptide displacement of a fluorescent AKAP-IS peptide from RII. The IC50 values for each are indicated. (/) Fluorescent and DIC images of HeLa cells expressing WT AKAP18α-GFP. (J) Fluorescent and DIC images of HeLa cells expressing AKAP18α V37M-GFP. (Scale bar: I-J, 10 μm.) (K) Whole-cell electrophysiological recording of cAMP-potentiated Ca²⁺ current in HEK293 cells expressing the L-type Ca²⁺ channel and either AKAP18 α WT or the AKAP18 α V37M variant. (L) Quantification of peak current potentiation. Data are presented as mean + SEM. *P < 0.001 by unpaired Student's t test. (M) Fluo4-AM measurement of cAMP-potentiated Ca2+ current in mouse aortic smooth muscle cells expressing either WT AKAP18 α (blue) or the AKAP18 α V37M variant (red). Calcium accumulation in response to de-



polarization was measured before and after brief forskolin stimulation. (N) Quantification of the change in fluorescence response to K^+ -induced depolarization. Data are presented as mean \pm SEM. *P < 0.001 by unpaired Student's t test.

of calcium currents (36, 43). Stepwise depolarization of cells was used to generate a pulse of Ca^{2+} current. In cells expressing WT AKAP18 α , application of cell-soluble cAMP elicits a 1.5-fold increase in Ca^{2+} current (n=10) (Fig. 2 K and L, blue). This second-messenger response was abolished in cells expressing the AKAP18 α V37M variant (Fig. 2 K and L, red).

Further substantiation in a more physiologically relevant system was possible by measuring depolarization-induced changes in Ca^{2+} flux in unpassaged cultured mouse aortic smooth-muscle cells expressing AKAP18 α or AKAP18 α V37M. Global Ca^{2+} influx through calcium channels was measured using the fluorescent calcium indicator dye acetoxymethyl-ester (AM) Fluo-4 (Fluo4-AM) upon depolarization with 60 mM potassium. In cells expressing AKAP18 α , a brief pulse of forskolin to activate PKA enhanced Fluo4-AM fluorescence twofold (n=7) (Fig. 2 M and N, blue). Importantly, this response was absent in cells expressing the AKAP18 α V37M variant (Fig. 2 M and N, red). Taken together, these experiments suggest that naturally occurring variants in the RII-binding helix of AKAP18 isoforms uncouple cAMP-responsive events at the plasma membrane.

AKAP18 Long Isoforms Transit to the Nucleus in a PKA-Dependent Manner. Long isoforms of AKAP18 encode targeting sequences that direct the anchoring protein to a variety of other subcellular locations (37, 44). For example, AKAP18γ and -δ are sequestered at the sarcoplasmic reticulum through protein-protein interactions with SERCA2 calcium reuptake channels, whereas in kidney collecting ducts, AKAP188 is a component of intracellular vesicles (38, 39). However, these static targeting interactions do not fully reflect the complexity of AKAP18 retention in the cytoplasm. Surprisingly, application of a cell-soluble PKA-anchoring disruptor peptide (stHt-31) elicited the nuclear accumulation of AKAP18y over a time course of 40 min (Fig. 3 A and B). In contrast, AKAP187 remained cytoplasmic upon application of the control peptide stHt-31-P (Fig. 3C). These results raised the intriguing possibility of whether or not association with the PKA holoenzyme influences the cytoplasmic retention of AKAP18γ or -δ.

Leptomycin B (LMB) is a bacterial metabolite that blocks CRM1-dependent nuclear export in mammalian cells. This compound traps proteins that have the capacity to translocate into the nucleus (45). Hence, we reasoned that the diminished capacity of AKAP18y V282M to anchor PKA would render this variant more prone to nuclear translocation. Live-cell imaging of HeLa cells treated with LMB (20 nM) monitored nuclear accumulation of GFP-tagged AKAP18y adducts (Fig. 3 D and E). Nuclear accumulation rates were calculated by quantifying fluorescent intensity ratios between the nucleus and cytoplasm over a time course of 40 min (Fig. 3F). Minimal movement of the WT anchoring protein was detected (Fig. 3 D and F, blue) (n =33 cells). In contrast, nuclear accumulation of AKAP18y V282M was pronounced (Fig. 3 G and H, red) (n = 58 cells) and occurred at a rate that is 3.75 times faster than the WT anchoring protein (Fig. 3F). Considering these data in light of the experimental evidence presented in Fig. 2, we propose that a reduced affinity for PKA contributes to the transit of AKAP18y V282M into the nucleus. Independent support for this hypothesis was provided by evidence that AKAP187 S278P/L281P, a PKA binding-null mutant, was exclusively detected in the nucleus (Fig. 3 G and H). Control experiments confirmed that WT AKAP18y associates with PKA, and is retained in the cytoplasm under these conditions (Fig. 3 G and H, Upper).

The long isoforms of AKAP18 (γ and δ) encode a putative nuclear localization signal (NLS) within exon 3 of the *AKAP7* gene (46). Therefore, a potential explanation for the results in Fig. 3 is that PKA anchoring obscures this NLS. To test this hypothesis, a family of deletion constructs was generated where regions of the AKAP18 δ were fused to YFP (Fig. 4*A*). In situ interaction with PKA was confirmed by coprecipitation of RII (Fig. 4*B*). The subcellular location of each YFP-tagged fragment was established by fluorescence microscopy (Fig. 4 *C*–*F*). Differential interference contrast (DIC) microscopy outlined cell contours (Fig. 4 *C*–*F*). These studies reveal that residues 1–98 of the anchoring protein are necessary and sufficient for nuclear targeting (Fig. 4*E*). To further demonstrate that the NLS is sufficient to direct AKAP18 into the nucleus in in the absence of

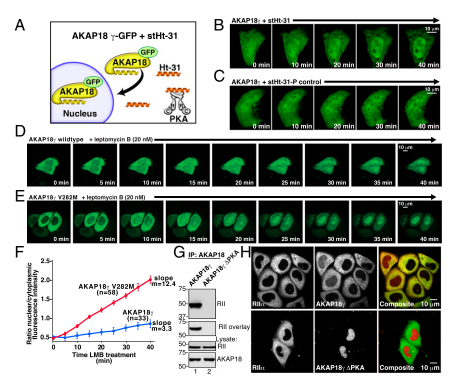


Fig. 3. Association with the PKA holoenzyme is necessary for cytoplasmic retention of AKAP18 long forms. (A) Schematic depicting displacement of PKA from AKAP18y by the cell-soluble PKA-anchoring disruptor peptide (stHt-31, orange) and the effect on subcellular targeting of AKAP18 long isoforms. Selected time points are shown from 0 to 40 min. (B and C) Live-cell imaging of cells expressing AKAP18y-YFP following application of (B) stHt-31 or (C) the proline control peptide, stHt-31-P. (D and E) Live-cell imaging of cells expressing (D) WT AKAP18γ-GFP or (E) V282M AKAP18γ-GFP. Cells were treated with LMB and subcellular distribution was recorded over 40 min. (F) Amalgamated data from multiple experiments in D and E. Changes in the ratio of nuclear/ cytoplasmic fluorescent signal WT AKAP18 γ -GFP (n =33 cells) and V282M AKAP18 γ -GFP (n = 58 cells). (G) The PKA-binding deficient mutant AKAP18y S278P/L281P (AKAP18 γ Δ PKA) is unable coprecipitate RII or interact with it by solid-phase overlay. (H) Fluorescent confocal images of HeLa cells expressing $\mbox{RII}\alpha\mbox{-YFP}$ (green) and (Lower) mCherry-AKAP18y (red) or (Bottom) the PKAdeficient AKAP18y mutant.

PKA, we deleted the 8-amino acid KKRKKKRK sequence and introduced proline mutations that perturb the RII-binding helix (AKAP18 γ Δ PKA Δ NLS) (Fig. 4 A and G, Top and Middle, lane 2). Fluorescent imaging revealed that both AKAP18γ ΔPKA ΔNLS and RII α are retained within the cytoplasm (Fig. 4 H and I). Within an enlarged region of interest, it was apparent that the AKAP18γ ΔPKA ΔNLS (red) and RIIα (green) signals exhibit distinct but overlapping cytoplasmic distributions (Fig. 4*J*, *Inset*). This is consistent with the ability of RII α to be incorporated into a spectrum of AKAP-PKA holoenzyme complexes dispersed throughout the cytoplasm (38, 47-50). Taken together, the results in Fig. 4 indicate that cytoplasmic retention of native AKAP18γ or -δ proceeds through a mechanism in which binding to the PKA holoenzyme masks a nuclear localization signal.

Proximity Labeling and Proteomics Define Pools of AKAP18 Complexes.

A corollary of these findings is that cytoplasmic and nuclear pools of AKAP18 interact with compartment-specific binding partners. To address this question, we tagged nuclear and cytoplasmic AKAP18 variants with miniTurbo, an engineered promiscuous ligase that permits local and efficient biotinylation of proteins within a radius of $\sim 5-10$ nm (51). Tagged proteins are captured on NeutrAvidin beads and identified by MS (Fig. 5A). The intracellular location of nuclear and cytoplasmic AKAP18-miniTurbo was determined by immunofluorescence detection of a V5 tag (Fig. 5 B and C). In situ detection of biotinylated target proteins was assessed by immunofluorescence using NeutrAvidin-Rhodamine Red X dye (Fig. 5 B and C). Immunoblot analysis using streptavidin-HRP revealed distinct staining patterns for the nuclear and cytoplasmic pools of AKAP18-binding partners (Fig. 5D, Upper, lanes 1 and 2). Immunoblot detection of the AKAP18 miniTurbo constructs served as a loading control (Fig. 5D, Lower). MS identified 131 binding partners of AKAP18 (Fig. 5 E-G). Forty of these proteins preferentially interacted with the nuclear variant (Fig. 5G, blue), whereas 67 proteins bound only to the cytoplasmic form (Fig. 5G, orange). Network analysis suggests that nuclear AKAP18 isoforms associate with RNA splicing machinery (Fig. 5H). Cytoplasmic forms of AKAP18 interact with proteins involved in translational control and cell cycle progression (Fig. 51). Taken together, these proximity-labeling results imply that AKAP18 isoforms associate with compartment-selective biding partners to participate in different cellular processes.

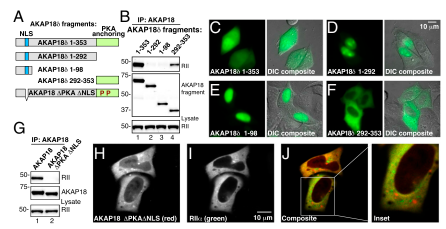
Discovery of the Non-PKA-Anchoring AKAP18 Epsilon. As outlined in Fig. 2A, numerous transcripts emanate from the AKAP7 gene. Scrutiny of the TCGA SpliceSeq database exposed a previously unidentified splicing event that bypasses the PKA-anchoring helix and replaces this region with a cryptic exon from the 3'UTR (Fig. 6A, blue). To amplify this transcript, we performed PCR on pooled human reverse-transcribed cDNA using specific primers (Fig. 6B, lane 1). PCR was also conducted with a 3' primer that bridged sequences in exon 9 and exon 12.2 (p3R) (Fig. 6B, lane 2). Nucleotide sequencing of the cloned product revealed an ORF that encodes a non-PKA-anchoring isoform of 329 amino acids with an alternate C terminus (Fig. 6D, blue shading). We have named this variant AKAP18ε. When expressed in HEK293 cells, AKAP18ε-GFP is only detected in the nucleus (Fig. 6D). In contrast RII α and α -tubulin are exclusively cytoplasmic (Fig. 6D). Biochemical characterization of AKAP18E immune complexes isolated from HEK293 cells demonstrate that this isoform does not interact with either RII α or C α subunits (Fig. 6E, Top and Middle-mid panels, lane 3). Control experiments confirmed both PKA subunits were detected in AKAP18y immune complexes (Fig. 6E, Top and Middle-mid panels, lane 2). Immunoblot detection of both PKA subunits in cell lysates served as loading controls (Fig. 6E, Bottom). Taken together these studies indicate that at least one nuclear isoform of AKAP18 does not function as a PKA-anchoring protein.

Discussion

Genome-wide association metaanalyses have identified SNPs and point mutations in AKAPs from patients with neurological disorders, hypertension, cardiac arrhythmias, and certain cancers (14, 52, 53). In this report we investigate SNPs within exons that encode the PKA-binding domains of several AKAPs. These functionally conserved regions form amphipathic helices that bind PKA holoenzymes with high affinity (11). In total we screened 42 missense variants gleaned from a variety of patient-derived datasets. In vitro binding studies confirmed that half of these single amino acid substitutions disrupted PKA anchoring. Further scrutiny of AKAP-Lbc, AKAP220, and AKAP-KL sequences reveal that a majority of detrimental substitutions occur within the penultimate and final turns of the anchoring helix (Fig. 1). Structural studies corroborate that these hydrophobic side-chains interface with the D/D of the R subunits of PKA (19, 54).

Nonsynonymous SNPs in the AKAP7 gene have been indirectly linked to an increased incidence of febrile seizures, cardiovascular events, and autism spectrum disorders (14, 26, 55, 56). This provided an impetus for more extensive analysis of the valine-to-methionine variant in the PKA-anchoring helix. Molecular models presented in Fig. 2 F and G show that the extra

Fig. 4. Long isoforms of AKAP18 contain a nuclear localization signal that may be masked by interaction with PKA. (A) Schematic showing fragments of YFPtagged AKAP188 isoforms that were used to map targeting regions within the anchoring protein. The first and last residue of each AKAP18 fragment is indicated. A polybasic NLS (blue) and PKA-anchoring regions (green) are identified. (B) IP of YFP-tagged AKAP188 fragments. (Top) Western blot analysis of immune complexes shows PKA RII coprecipitating with full-length protein and the 292-353 fragment. (Middle) Loading control for AKAP18 fragments and (Bottom) loading controls for RIIα. (C-F) Fluorescent and DIC imaging detecting the differential subcellular distribution YFP-tagged AKAP188 fragments. (G) Characterization of WT AKAP18γ and AKAP18γ ΔPKA ΔNLS mutant immune complexes. (Top) RII coprecipitates with the WT only. (Middle) Loading control for AKAP18. (Bottom) loading control for RIIa. (H) Fluorescent con-



focal image of the mCherry-tagged AKAP18γ ΔPKA ΔNLS mutant shows cytoplasmic retention. (/) Rllα-YFP is also cytoplasmic. (/) Single channel and composite images show both an overlapping and distinct cytoplasmic distribution of RII (green) and AKAP18 APKA ΔNLS (red). (Inset) Magnified image showing the regional distribution of RII and AKAP18.

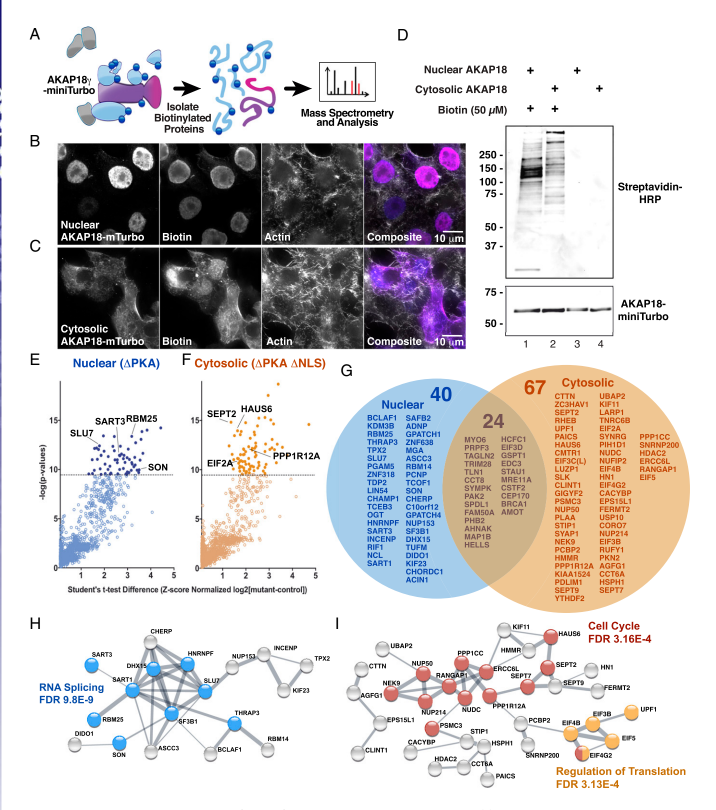
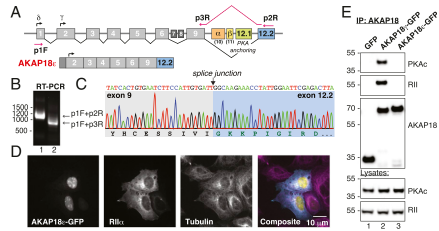


Fig. 5. Proximity labeling and proteomic identification of compartment-selective AKAP18 interactors. (A) Schematic depicting the miniTurbo-based biotinylation/MS experiments. Biotin is shown as dark-blue circles. (Left) Upon biotin addition, miniTurbo biotinylates only proximal proteins. (Center) Biotinylated proteins are isolated with streptavidin pulldown and (Right) identified by MS. (B and C) Fluorescence microscopy depicting the subcellular location of (B) nuclear and (C) cytoplasmic AKAP18-miniTurbo variants and detection of compartment specific biotinylated proteins. (D) Western blot analysis of biotinylated proteins in cell lysates shows distinct and compartment-specific labeling patterns. (E and F) MS results plotted as log difference of experimental intensity minus control intensity plotted against the negative log of the P value. Selected nuclear (dark-blue) and cytoplasmic (tan) AKAP18 interacting partners are indicated. (G) Lists of statistically significant target proteins in nuclear and cytosolic preparations. Of 131 identified proteins, 40 were selectively associated with the nuclear variant, whereas 67 preferentially bound to the cytoplasmic variant. (H and I) STRING database-generated network depictions of compartment-specific proteins. Gene ontology process annotation and false-discovery rates (FDR) are listed for the highlighted nodes. Putative biological functions for each subnetwork are indicated.

Fig. 6. Identification of an AKAP18 isoform that lacks a PKA binding helix. (A) AKAP7 intron-exon map as adapted from TCGA SpliceSeq. The start sites for the δ - and γ -isoforms are noted, as are the α - and β-isoform-specific exons. The RII anchoring site (green) is encoded in exon 12.1 and the alternate 12.2 exon (blue) are presented. The locations of the primers used for RT-PCR are marked. (B) RT-PCR from pooled human cDNA samples using primers against predicted transcripts and novel exon junctions produces the expected products where intervening exons are spliced out. (C) Sequencing chromatogram of the exon 9-12.2 splice junction shows an inframe protein sequence corresponding to a distinct C terminus. (D) Fluorescent confocal images showing nuclear localization of EmGFP-tagged AKAP18ε and cytoplasmic labeling of RII α -V5 and α -tubulin. (E) Immune complexes of either WT AKAP187 or AKAP18ε were immunoprecipitated and copur-



ification of PKA catalytic (*Upper*) and RII (*Upper*-mid) subunits were identified by Western blotting. Expression levels of both AKAP18 forms and GFP are indicated in the *Lower* three panels.

molecular density of a methionine at position 282 in AKAP18y deflects the helical pitch of the hydrophobic anchoring surface upward by 6.6°. This relatively modest topological flaw decreases the binding affinity for RII by 8.8-fold and impairs PKAmediated potentiation of L-type Ca²⁺ currents, as shown in Fig. 2H. Because alternative splicing of the AKAP7 gene generates multiple isoforms, Met-282 polymorphic variants might negatively impact a certain local PKA signaling events. These include modulation of neuronal sodium channels, regulation of SERCA2-mediated calcium reuptake in the heart, and aquaporin-2-mediated control of water homeostasis in the kidney (38, 39, 57). The physiological significance to this concept is underscored by recent evidence that intact and anchored PKA holoenzymes are the predominant cytoplasmic signaling units and are sequestered within 200-400 Å of their preferred substrates (21, 22). Hence, AKAP variants that impact the intracellular targeting of PKA holoenzymes have the capacity to alter the dissemination of second-messenger signals.

Although polymorphic variation within AKAPs has the potential to uncouple local cAMP signaling, there is little information about the allelic frequency or prevalence in specific populations. In addition, scrutiny of the cBioPortal cancer genomics data interface does not reveal any clear pattern of AKAP-anchoring helix mutations that correlate with cancers. Perhaps the best-documented example of genetic variation comes from analysis of a nonsynonymous SNP that promotes an isoleucine to valine substitution at position 646 within the PKAanchoring domain of D-AKAP-2 (AKAP10) (12, 58). This variant is associated with cardiac dysfunction in aging populations of European-American individuals that display a shortened electrocardiogram PR interval (12). The D-AKAP2 I646V variant is also associated with increased basal heart rate (58). Because basal heart rate correlates inversely with lifespan, the altered PKA-anchoring properties of D-AKAP2 may be linked to adverse pathological outcomes (59). However, the downstream signaling pathways affected by this modified kinase-anchoring event remain obscure. In vitro biochemical studies show that the D-AKAP2 646V variant binds to type I PKA holoenzyme with threefold higher affinity than D-AKAP2 646I, yet this variation has no effect on type II PKA anchoring (12). The physiological ramifications of these relatively small changes in affinity may be limited as D-AKAP2 exhibits a strong binding preference for type II PKA holoenzymes (60, 61). Furthermore, the porcine, murine, and chicken orthologs of D-AKAP2 have a valine at the corresponding position in the PKA-anchoring helix (12). While structural analyses of these D-AKAP2 orthologs are scant, it seems that the Ile to Val substitution can be tolerated with minimal effects on PKA anchoring. This raises the issue of whether or not silent variation in AKAP

helices is a more general phenomenon. Data presented in Fig. 1F indicate that a silent change in the final turn of the AKAP79 helix (Ile-406 to Val) has no apparent effect on RII-binding. This may represent another example of evolutionary drift, as valine naturally occurs at this position in the mouse, rat, and macaque orthologs of AKAP79. Thus, malleability within amphipathic helices can accommodate structurally conserved variants or passenger mutations without any substantial impact on PKA anchoring.

AKAPs are sequestered within defined subcellular locations by specialized targeting motifs that interact with membranes or organelles (62–65). The α - and β -isoforms of AKAP18 were originally identified as membrane proximal lipid-modified PKAanchoring proteins that facilitate cAMP-responsive modulation of ion channels (36, 66). In contrast the longer γ - and δ -isoforms of AKAP18 are retained within the cytoplasm through different targeting mechanisms (37, 39). Studies presented in Figs. 3-5 argue that interaction with the PKA holoenzyme influences the cytoplasmic distribution of AKAP18 long isoforms. This unique mode of AKAP targeting is supported by time-lapse imaging experiments in Fig. 3B, showing that application of PKA anchoringdisruptor peptides facilitates nuclear accumulation of AKAP18y. This may be an AKAP18-selective phenomenon, because other anchoring proteins remain firmly attached to their targeting sites upon displacement of PKA. For example, delivery of anchoring inhibitor peptides has no effect on the mitochondrial location of D-AKAP-1 and OPA1 or the membrane tethering of AKAP79/150 and Gravin (47, 63, 67, 68). However, one discriminating feature may be the relatively small size of AKAP18, which is within the molecular dimensions of particles (30-60 kDa) that diffuse through the nuclear pore (69). With this in mind, we took advantage of the reduced PKA-anchoring affinity of the AKAP18y V282M variant to show that the rate of nuclear accumulation is enhanced 3.8-fold compared with WT in the presence of LMB. Thus, paradoxically we can propose that intact PKA holoenzymes serve as a cytoplasmic anchor for certain long isoforms of AKAP18. Molecular explanations for this alternative targeting mechanism include the dimensions of the AKAP18-RII₂-C₂ pentameric complex (~230,000 kDa), which is too large to transit the nuclear pore and that R subunit dimers are only detected in the cytoplasmic compartment (22). In contrast, free catalytic subunits only transit into the nucleus in response to pharmacological regimens that promote supraphysiological accumulation of cAMP or in adrenal adenomas where pathological Cα mutants are excluded from PKA holoenzymes (70–72).

Our demonstration of AKAP18 γ nuclear shuttling and the subsequent discovery of AKAP18 ϵ further expands our view of AKAP biology. For example, PKA anchoring contributes to the

cytoplasmic retention of AKAP18 by masking a classic NLS that is located between residues 52-59. Furthermore, data in Fig. 6 characterizing the AKAP18E isoform that lacks a PKA-anchoring helix, infer that this splice variant utilizes this NLS as its sole targeting signal. In both cases this would permit direct transport through the nuclear pore by targeting factors, such as Ran GTPase and importin (73). In addition, enzyme-catalyzed proximitydependent proteomic screens substantiate that cytoplasmic and nuclear AKAP18s coordinate distinct interactomes. Network analyses presented in Fig. 5 H and I argue that cytoplasmic AKAP18 variants interface with RNA transport and translation machinery, whereas their nuclear counterparts facilitate signaling to spliceosomal components. Although individual nuclear functions for AKAP18γ, -δ, and -ε have yet to be ascribed, X-ray crystallography identified a common 2'-3' phosphoesterase-like fold within each isoform (74, 75). One notable feature was the detection of 5'-AMP embedded deep within a grove of this shared domain. This degradation product of cAMP is coordinated by two His-X-Thr motifs that are hallmarks of the 2H-phosphoesterase-like fold (74). Postulated roles for this nucleotide-protein interaction include energy sensing or as an auxiliary nuclear localization element akin to ligand mediated targeting of steroid hormone receptors (74). Another important concept investigated in this study is that non-PKA-anchoring products can be transcribed from AKAP genes. On the surface, this notion may seem counterintuitive to the AKAP model (15, 76). However, omission of the PKA-anchoring helix could favor formation of AKAP18 macromolecular assemblies that respond to different activation signals (11, 43). This latter issue may have confounded analysis of AKAP18 knockout mice, as only an exon encoding the PKA-anchoring region was ablated from the *AKAP7* gene (77).

Methods

Sequence Analysis and Variant Identification. The variants described here were captured at several different points in time as new SNPs were deposited into the NCBI dbSNP and NCBI Variation Viewer databases.

Peptide Array Synthesis. SPOT arrays were generated as described previously (60).

Live Cell Imaging. HeLa cells expressing AKAP18-GFP constructs were washed in HBSS, mounted in a Ludin chamber (Life Imaging Services), and imaged

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using a Leica AS-MDW workstation. For nuclear trapping experiments, LMB (Calbiochem) was added to 20 nM and images were captured every 5 min for at least 90 min

Structural Modeling with AutoDock. AutoDock calculations were carried out using AutoDock Tools 1.5.6, according to the methods described in ref. 42.

Electrophysiology. HEK293 cells were transfected with cDNAs encoding the α 1c and β 2a subunits of cardiac L-type Ca²⁺ channels (36) and WT or V37M AKAP18α-GFP. Whole-cell recordings were performed using an Axopatch 200B patch-clamp amplifier (Axon Instruments). L-type calcium channel currents were evoked step depolarization (200 ms) from a holding potential of –80 mV to +10 mV.

Calcium Imaging. Male WT C57BL/6J mice of 5 to 8 wk of age were euthanized with sodium pentobarbital (250 mg/kg; intraperitoneally), as approved by the University of California, Davis Animal Care and Use Committee (protocol #20321). Mouse aorta were dissected, washed, and digested in DMEM solution containing 2.2 mg/mL of collagenase type 2 (Worthington). Unpassaged arterial myocytes were seeded on glass coverslips coated with laminin and kept in an incubator at 37 °C with 5% $\rm CO_2$ for 7–10 d before transfection. Transfected arterial myocytes were loaded with Fluo4-AM for 20 min at room temperature. Images were collected using an Andor spinning-disk confocal system coupled to an Olympus iX-81 inverted microscope equipped with an Olympus 40× oil immersion lens (NA = 0.75).

Proximity Labeling and Mass Spectrometry. The cDNA for miniTurbo (51) was fused to 3′ end of AKAP18γ cDNAs and tested for expression in HEK293T cells. Transfected cells were incubated in media containing 50 μM biotin for 1 h. After lysate preparation and biotinylated target capture, proteins were reduced and alkylated digested using Lys-C (Wako) followed by further trypsin digestion. Peptides were analyzed by nanoLC-MS on an Orbitrap Fusion Lumos Tribrid Mass Spectrometer. Protein network prediction and gene ontology analysis were performed using STRING database version 10.5 (78). Networks were visualized using Cytoscape v3.6.1 software.

Complete methods can be found in SI Appendix.

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