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# Ser<sup>1928</sup> phosphorylation by PKA stimulates the L-type Ca<sup>2+</sup> channel Ca<sub>v</sub>1.2 and vasoconstriction during acute hyperglycemia and diabetes

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#### Abstract

Hypercontractility of arterial myocytes and enhanced vascular tone during diabetes are, in part, attributed to the effects of increased glucose (hyperglycemia) on L-type  $Ca_V 1.2$  channels. In murine arterial myocytes, kinase-dependent mechanisms mediate the increase in  $Ca_V 1.2$  activity in response to increased extracellular glucose. We identified a subpopulation of the  $Ca_V 1.2$  channel pore-forming subunit ( $\alpha 1_C$ ) within nanometer proximity of protein kinase A (PKA) at the sarcolemma of murine and human arterial myocytes. This arrangement depended upon scaffolding of PKA by an A-kinase anchoring protein 150 (AKAP150) in mice. Glucose-mediated increases in  $Ca_V 1.2$  channel activity were associated with PKA activity, leading to  $\alpha 1_C$  phosphorylation at Ser<sup>1928</sup>. Compared to arteries from low-fat diet (LFD)–fed mice and nondiabetic patients, arteries

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from high-fat diet (HFD)–fed mice and from diabetic patients had increased Ser<sup>1928</sup> phosphorylation and Ca<sub>V</sub>1.2 activity. Arterial myocytes and arteries from mice lacking AKAP150 or expressing mutant AKAP150 unable to bind PKA did not exhibit increased Ser<sup>1928</sup> phosphorylation and Ca<sub>V</sub>1.2 current density in response to increased glucose or to HFD. Consistent with a functional role for Ser<sup>1928</sup> phosphorylation, arterial myocytes and arteries from knockin mice expressing a Ca<sub>V</sub>1.2 with Ser<sup>1928</sup> mutated to alanine (S1928A) lacked glucosemediated increases in Ca<sub>V</sub>1.2 activity and vasoconstriction. Furthermore, the HFD-induced increases in Ca<sub>V</sub>1.2 current density and myogenic tone were prevented in S1928A knockin mice. These findings reveal an essential role for  $\alpha$ 1<sub>C</sub> phosphorylation at Ser<sup>1928</sup> in stimulating Ca<sub>V</sub>1.2 channel activity and vasoconstriction by AKAP-targeted PKA upon exposure to increased glucose and in diabetes.

#### INTRODUCTION

Diabetes mellitus is a major risk factor underlying multiple pathological complications such as stroke, hypertension, heart disease, and retinal degeneration (1). These pathologies have been linked, in part, to enhanced contractility of arterial myocytes in the resistance vasculature during hyperglycemia, that is, high blood glucose that is a signature metabolic feature of diabetes (2–7). The contractile state of arterial myocytes, and thereby arterial diameter, is predominantly controlled by the amount of  $Ca^{2+}$  influx through L-type  $Ca_V 1.2$ channels. We reported that  $Ca_V 1.2$  channel activity in murine cerebral arterial myocytes is augmented in response to increased glucose and in a mouse model of diabetes through a process that is independent of changes in membrane potential (8). However, the molecular mechanisms underlying the enhanced  $Ca_V 1.2$  channel function associated with increased glucose and whether similar phenomena are engaged in the human vasculature during diabetes are unclear.

The physiological importance of  $Ca_V 1.2$  channels is underscored by their essential role in cardiac contractility, neuronal excitability, and control of vascular tone in health and disease (9, 10).  $Ca_V 1.2$  channels are composed of pore-forming  $a_{1C}$  subunits in association with several regulatory accessory subunits (11). Regulation of  $Ca_V 1.2$  function occurs through reversible phosphorylation of the channel complex by serine/threonine kinases and phosphatases, including protein kinase A (PKA) (10–12). Modulation of  $Ca_V 1.2$  channel function by PKA has been extensively studied in the context of  $\beta$ -adrenergic stimulation in the heart and brain (9, 12). Although PKA-dependent stimulation of  $Ca_V 1.2$  channels occurs in myocytes from arteries and portal veins (13–15), the molecular mechanisms and functional implications are unknown.

Regulation of Ca<sub>V</sub>1.2 channel activity by PKA in other cell types involves direct phosphorylation of one or more subunits that compose the channel (16). A highly conserved PKA consensus phosphorylation site is Ser<sup>1928</sup>, located within the intracellular C terminus of  $\alpha$ 1<sub>C</sub>. Stimulation of Ca<sub>V</sub>1.2 by PKA activity has been associated with increased phosphorylation of this residue in neurons (17–20) and cardiac cells (19), although its physiological significance in cardiac Ca<sub>V</sub>1.2 channel function is controversial (21, 22).

However, evidence for this modification and elucidation of its functional role in the vasculature are lacking.

PKA-dependent stimulation of heterologously expressed (23), as well as native cardiac (24, 25) and neuronal (20, 26, 27),  $Ca_V 1.2$  channels requires an A-kinase anchoring protein (AKAP) scaffold. Consistent with these findings, we demonstrated that potentiation of Ca<sub>V</sub>1.2 channels in arterial myocytes during acute increases in extracellular D-glucose requires an AKAP-bound PKA (8). Yet, the functional implications of PKA-dependent stimulation of  $Ca_V 1.2$  channels in response to increased glucose, as well as the identity of the AKAP, are unknown. Here, super-resolution nanoscopy and analysis by proximity ligation assay (PLA) revealed the existence of a subpopulation of Ca<sub>V</sub>1.2 and PKA catalytic subunit (PKAcat) clusters that are within nanometer range of each other in mouse and human arterial myocytes. We identified the scaffold protein AKAP150 (the murine ortholog of human AKAP79) and the binding of PKA to AKAP150 as key determinants of PKAmediated a1<sub>C</sub> phosphorylation at Ser<sup>1928</sup>. PKA-mediated phosphorylation of this residue resulted in potentiation of vascular Ca<sub>V</sub>1.2 channel activity and vasoconstriction in response to increased extracellular D-glucose. Enhanced  $Ca_V 1.2$  channel activity and intracellular  $Ca^{2+}$  concentrations ( $[Ca^{2+}]_i$ ), which augmented vasoconstriction in response to increased Dglucose and in mice fed a high-fat diet (HFD), were abolished upon prevention of Ser<sup>1928</sup> phosphorylation by selective mutation of this residue to alanine in S1928A knockin mice. We also observed PKA-dependent phosphorylation of  $\alpha 1_C$  at Ser<sup>1928</sup> and enhanced Ca<sub>V</sub>1.2 channel activity in human arterial myocytes in response to increased extracellular D-glucose and in human cells from diabetic subjects, thus highlighting the translational relevance of our findings. Together, these results revealed a fundamental role for AKAP-dependent, PKA-meditated phosphorylation of  $\alpha 1_C$  at Ser<sup>1928</sup> as a key molecular signaling event underlying potentiation of Ca<sub>V</sub>1.2 channel activity and vasoconstriction upon acute increases in extracellular D-glucose and in diabetes.

#### RESULTS

#### Super-resolution spatial maps and PLA reveal sarcolemmal Ca<sub>V</sub>1.2-PKA interactions

Enhanced Ca<sub>V</sub>1.2 channel activity in rodent cerebral arterial myocytes after an increase in extracellular D-glucose depends upon PKA activity (8). We therefore hypothesized that a subpopulation of PKA is located near Ca<sub>V</sub>1.2 channels in these cells, enabling functional regulation of channel activity. To test this, we examined the spatial relationship between Ca<sub>V</sub>1.2 and PKA in freshly dissociated cerebral arterial myocytes from wild-type mice using stochastic optical reconstruction microscopy (STORM) in the total internal reflection fluorescence (TIRF) configuration with antibodies against Ca<sub>V</sub>1.2 (28) and PKA<sub>cat</sub>. This method achieves a lateral resolution of ~30 nm with selective axial illumination of sarcolemmal and subsarcolemmal regions. Although conventional TIRF images showed diffuse and regionally varied fluorescence associated with Ca<sub>V</sub>1.2 and PKA<sub>cat</sub> (Fig. 1A, top), STORM-rendered spatial maps revealed that both proteins were broadly distributed in clusters of various sizes (Ca<sub>V</sub>1.2, 2485 ± 47 nm<sup>2</sup>; PKA<sub>cat</sub>, 3272 ± 72 nm<sup>2</sup>) at the sarcolemma of arterial myocytes [Fig. 1, A (bottom) and B]. We observed sites of close interaction between Ca<sub>V</sub>1.2 and PKA<sub>cat</sub> (Fig. 1C). Histograms of the Ca<sub>V</sub>1.2-to-nearest

 $PKA_{cat}$  distances were fit with a two-component Gaussian function with centers at ~93 and ~256 nm (Fig. 1D), suggesting that a pool of PKA is localized closer to a subpopulation (~5 to 10%) of Ca<sub>V</sub>1.2 channels.

We used PLA as an additional test of  $Ca_V 1.2$ -PKA association in arterial myocytes. PLA generates fluorescent puncta when proteins of interest are at or less than 40 nm apart (29). Mouse arterial myocytes colabeled for  $Ca_V 1.2$  and PKA<sub>cat</sub> produced fluorescent puncta, indicating the presence of the PLA reaction product (Fig. 1E). PLA signals were nearly absent when either  $Ca_V 1.2$  or PKA<sub>cat</sub> primary antibodies were omitted (Fig. 1, E and F). PLA was also performed on cells colabeled for  $Ca_V 1.2$  and transferrin receptor (TfR) or  $Ca_V 1.2$  and the  $Ca^{2+}$ -activated Cl<sup>-</sup> channel anoctamin 1 (ANO1), which are cell surface proteins that have limited to no interaction with  $Ca_V 1.2$  (30–33). Arterial myocytes were robustly stained for TfR and ANO1 when primary antibodies were included in the preparation (fig. S1A), yet no PLA signal was detected in cells colabeled for  $Ca_V 1.2$  and TfR or  $Ca_V 1.2$  and ANO1 (fig. S1, B and C). As a positive control, cells colabeled with two distinct  $Ca_V 1.2$  antibodies [rabbit FP1 (28) and monoclonal NeuroMab clone N263/31] showed surface membrane staining (fig. S1A) and produced robust PLA signals (fig. S1, B and C). Together, these data indicated close ( 40 nm) spatial proximity between a subpopulation of  $Ca_V 1.2$  channels and PKA<sub>cat</sub> in mouse arterial myocytes.

### AKAP150 facilitates sarcolemmal interactions between Ca<sub>V</sub>1.2 and PKA in murine arterial myocytes

The scaffolding protein AKAP150, which is the murine ortholog of human AKAP79, binds and coordinates a multienzyme complex that includes PKA, protein kinase C (PKC), and the protein phosphatase calcineurin to facilitate signaling to specific substrates, including  $Ca_V 1.2$  (20, 26, 27, 34–37). We found that the PKA RIIa regulatory subunit and PKCa coimmuno-precipitated with AKAP150 in arteries from wild-type mice (Fig. 2A). We tested whether the AKAP150-PKA complex facilitates the proximity between PKA<sub>cat</sub> and  $Ca_V 1.2$ in arterial myocytes by examining their distribution in cells isolated from two genetically engineered mouse strains. One expresses a truncated form of AKAP150 in which the binding site for PKA has been deleted ( A; fig. S2) (26, 38), and the other lacks the entire AKAP150 protein (*AKAP150<sup>-/-</sup>*) (37). The association between AKAP150 and PKA RIIa was either absent (*AKAP150<sup>-/-</sup>*) or markedly reduced ( A) in arteries from these mice (Fig. 2A). Coimmunoprecipitation of PKCa within the AKAP150 complex was maintained in A arteries but was absent in the arteries from the *AKAP150<sup>-/-</sup>* mice (Fig. 2A).

We used structured illumination microscopy (SIM; lateral resolution, ~114 nm) (39) in combination with an object-based analysis to compare  $Ca_V 1.2$  and  $PKA_{cat}$  proximity in arterial myocytes from wild-type, A, and  $AKAP150^{-/-}$  animals. Consistent with findings we obtained by STORM, object-based analysis identified sites of spatial overlap between  $Ca_V 1.2$  and  $PKA_{cat}$  at the sarcolemma of wild-type arterial myocytes (Fig. 2, B to D). In contrast, cells from A and  $AKAP150^{-/-}$  mice exhibited significantly fewer sites of overlap between  $Ca_V 1.2$  and  $PKA_{cat}$  (Fig. 2, B and D). Furthermore, A and  $AKAP150^{-/-}$  myocytes had a reduction in the density of  $PKA_{cat}$  but not  $Ca_V 1.2$  particles at the sarcolemma

identified by STORM/TIRF (Fig. 2E). These findings suggested that AKAP150 facilitates targeting of a pool of  $PKA_{cat}$  to a subpopulation of  $Ca_V1.2$  in arterial myocytes.

#### Enhanced vascular Ca<sub>V</sub>1.2 current in 20 mM D-glucose requires AKAP150-anchored PKA

We tested the hypothesis that increasing extracellular D-glucose from 10 to 20 mM augments Cav1.2 channel activity in an AKAP150- and PKA-dependent manner in mouse arterial myocytes. These extracellular D-glucose concentrations are within the range of observed nonfasting blood glucose concentrations reported for nondiabetic and experimental diabetic mice, respectively (3, 8, 40). To assess the effects of increased extracellular glucose on  $Ca_{\rm V}1.2$  channel activity, we used patch-clamp electrophysiology in the whole-cell configuration with barium (Ba<sup>2+</sup>) as the charge carrier before and after application of the Ltype calcium channel blocker nifedipine to determine the nifedipine-sensitive Ba<sup>2+</sup> current  $(I_{\text{Ba}})$  associated with Ca<sub>V</sub>1.2 channel activity. We found that 20 mM p-glucose produced a robust increase in  $I_{Ba}$  across a range of membrane potentials with no change in the currentvoltage (*I-V*) relationship ( $V_{\text{max}} = 8.75 \pm 1.26 \text{ mV}$  for 10 mM D-glucose and  $V_{\text{max}} = 9.45$  $\pm$  0.56 mV for 20 mM D-glucose; P = 0.6174, extra sum-of-squares F test) in arterial myocytes (Fig. 3, A and B). We also observed a similar increase in IBa in response to 20 mM D-glucose in mesenteric and femoral arterial myocytes (fig. S3, A and B), suggesting that increased extracellular D-glucose potentiates Cav1.2 current density in myocytes from different vascular beds. Incubation of arterial myocytes with the selective inhibitor of the glucose transporter GLUT4 indinavir (41) prevented the increase in Cav1.2 current density in response to 20 mM  $_D$ -glucose (fig. S4A). Furthermore,  $I_{Ba}$  was unchanged when  $_D$ glucose was substituted with equimolar concentrations of the metabolically inactive enantiomer L-glucose (10 mM D-glucose + 10 mM L-glucose; fig. S4B), suggesting that enhanced Ca<sub>V</sub>1.2 channel activity in response to increased D-glucose requires its internalization and metabolic utilization.

Contrary to findings in wild-type arterial myocytes, 20 mM G-glucose did not stimulate  $I_{Ba}$  that were elicited with a voltage pulse from -70 to +10 mV in *AKAP150<sup>-/-</sup>* and A myocytes (Fig. 3, C and D). Note that there was no change in *I-V* relationship by 20 mM D-glucose in *AKAP150<sup>-/-</sup>* cells (fig. S5) with  $V_{max}$  similar to that in wild-type cells ( $V_{max} = 8.75 \pm 1.26$  mV in wild-type and  $V_{max} = 7.98 \pm 2.06$  mV in *AKAP150<sup>-/-</sup>*; P = 0.7836, extra sum-of-squares *F* test). Consistent with the electrophysiological data, we observed an increase in [Ca<sup>2+</sup>]<sub>i</sub> in cells from wild-type mice loaded with the Ca<sup>2+</sup>-sensitive indicator Fluo-4 in response to 20 mM D-glucose (Fig. 3, E and F). This response was not observed in the presence of nifedipine or when D-glucose was substituted with L-glucose (fig. S6), and it was significantly attenuated in *AKAP150<sup>-/-</sup>* and A cells (Fig. 3, E and F). Together, these findings indicated that AKAP150-anchored PKA signaling is required for enhanced Ca<sub>V</sub>1.2 channel activity and an increase in global [Ca<sup>2+</sup>]<sub>i</sub> in response to increasing the concentration of extracellular D-glucose.

We also investigated whether AKAP150-anchored PKC participates in enhancing  $I_{Ba}$  in response to increasing extracellular D-glucose to 20 mM with a transgenic mouse expressing a form of AKAP150 that cannot bind PKC, yet retains binding to PKA (C; Fig. 4A and fig. S2). Consistent with results in arteries from *AKAP150<sup>-/-</sup>* mice (37), basal tone in

pressurized (60 mmHg) middle cerebral arteries was slightly reduced in the C mice, and receptor-mediated activation of PKC with angiotensin II evoked less constriction in arteries from C mice than in arteries from littermate controls (Fig. 4, B to D). Moreover, angiotensin II did not increase  $I_{Ba}$  in arterial myocytes from C mice, yet  $I_{Ba}$  in these cells was significantly augmented by 20 mM D-glucose (Fig. 4, E and F). These data indicated that AKAP150-anchored PKA, but not AKAP150-anchored PKC, is responsible for enhanced Ca<sub>V</sub>1.2 channel activity in response to increased D-glucose in arterial myocytes.

### Phosphorylation of $a_{1C}$ at Ser<sup>1928</sup> underlies enhanced vascular Ca<sub>V</sub>1.2 activity in response to 20 mM D-glucose

We tested whether increased Ca<sub>V</sub>1.2 channel activity was associated with an alteration in the phosphorylation state of the  $\alpha 1_{\rm C}$  subunit at Ser<sup>1928</sup>, which is a conserved PKA target site in the distal C terminus and which has been implicated in Ca<sub>V</sub>1.2 regulation (17–19). Using Western blot analysis with a specific antibody that recognizes  $\alpha 1_{\rm C}$  phosphorylation at Ser<sup>1928</sup> (42), we found increased phosphorylation of this residue in wild-type mouse arterial lysates incubated for 5 min with 20 mM D-glucose as compared with those with 10 mM D-glucose (Fig. 5, A and B). In contrast to wild-type arteries, no change in Ser<sup>1928</sup> phosphorylation in response to 20 mM D-glucose was observed in arterial lysates from A or *AKAP150<sup>-/-</sup>* mice (Fig. 5, A and B). These results suggested that AKAP150-anchored PKA is required for phosphorylation of  $\alpha 1_{\rm C}$  at Ser<sup>1928</sup> in response to increased D-glucose.

Considering an apparent relationship in molecular and functional data described above, we postulated that Ser<sup>1928</sup> serves as a functional PKA regulatory site in vascular Ca<sub>V</sub>1.2 channels. This hypothesis was tested using patch-clamp electrophysiology to record  $I_{Ba}$  in arterial myocytes isolated from a genetically engineered mouse expressing Ca<sub>V</sub>1.2 in which Ser<sup>1928</sup> has been mutated to an alanine residue (S1928A) (21). Our data revealed that  $I_{Ba}$  was not altered by an increase in extracellular D-glucose in S1928A myocytes (Fig. 5, C and D). Pretreatment of wild-type myocytes with the PKA inhibitor Rp-cyclic 3',5'-hydrogen phosphorothioate adenosine triethylammonium salt (rpcAMPS) also blocked the D-glucose–induced stimulation of  $I_{Ba}$  (Fig. 5, C and D). In contrast to the response of wild-type cells, we found no significant change in global  $[Ca^{2+}]_i$  in response to 20 mM D-glucose in cells from S1928A mice (Fig. 5E). We confirmed that protein abundance of K<sub>V</sub>2.1, BK<sub>Ca</sub>  $\alpha$ , and BK<sub>Ca</sub>  $\beta$ 1 subunits, which regulate arterial myocyte excitability and are glucose-sensitive (43–46), is similar between wild-type and S1928A arteries (fig. S7), thus suggesting that loss of glucose-induced stimulation of global Ca<sup>2+</sup> in S1928A myocytes is not due to changes in basal expression of these K<sup>+</sup> channel subunits.

Increases in extracellular D-glucose could modulate protein function by promoting O-linked N-acetylglucosamine (O-GlcNAc) (47, 48). Therefore, we examined whether posttranslational O-GlcNAc modification modulates Ca<sub>V</sub>1.2 channel activity in response to increased D-glucose by treating wild-type cells with the inhibitor of glutaminefructose amidotransferase, 6-diazo-5-oxo-L-norleucine (DON), and measuring the effect of glucose on  $I_{Ba}$ . Inhibition of this enzyme prevents production of the substrate for O-GlcNAc transferase. Perfusion of 20 mM D-glucose enhanced  $I_{Ba}$  in wild-type mouse arterial

myocytes in the presence of DON, indicating that the effect of glucose on  $Ca_V 1.2$  channel activity was not mediated by O-GlcNAcylation (fig. S8).

To explore the functional significance of the glucose-induced Ser<sup>1928</sup> phosphorylation, we performed measurements of arterial diameter in pressurized (60 mmHg) middle cerebral arteries in the presence of 10 and 20 mM D-glucose. Arteries isolated from wild-type mice developed stable myogenic tone and constricted in response to 20 mM D-glucose (Fig. 5F). This constriction response was independent of endothelial function, because we observed a similar response in wild-type endotheliumdenuded arteries (fig. S9). Although basal myogenic tone was not significantly different in arteries from S1928A and wild-type mice, the response to 20 mM D-glucose was abolished in arteries from S1928A mice (Fig. 5F). Together, data indicated that phosphorylation of  $\alpha 1_C$  at Ser<sup>1928</sup> by AKAP150-anchored PKA is necessary for increased Ca<sub>V</sub>1.2 channel activity, global [Ca<sup>2+</sup>]<sub>i</sub>, and vasoconstriction in response to 20 mM D-glucose.

### The single point mutation S1928A restores normal $Ca_V 1.2$ activity and vascular tone in HFD animals

We tested whether phosphorylation of  $a_{1c}$  at Ser<sup>1928</sup> is enhanced in mice on an HFD. Wildtype, AKAP150<sup>-/-</sup>, A, and S1928A animals were maintained on either a low-fat diet (LFD) (10% kcal fat) or HFD (60% kcal fat) for a period of 12 to 16 weeks. This HFD model recapitulates clinical features observed in patients with type 2 diabetes (49, 50), including enhanced arterial tone and increased blood pressure (3, 40). HFD animals exhibited significantly higher body weight and increased blood glucose concentration (table S1). Arteries and arterial myocytes from LFD and HFD mice were isolated, maintained, and immediately used in experiments with extracellular solutions containing 10 mM D-glucose. We found that  $\alpha 1_{C}$  at Ser<sup>1928</sup> was significantly more phosphorylated in arteries from wildtype HFD mice compared with those from wild-type LFD mice (Fig. 6, A and B). Additionally, arteries from the LFD and HFD A mice had similar amounts of Ser<sup>1928</sup> phosphorylation, which were not different compared with arteries from wild-type LFD mice (Fig. 6, A and B). Arterial myocytes from wild-type HFD mice exhibited significant enhancement of  $I_{Ba}$  over a range of membrane potentials with no change in *I*-V relationship  $(V_{max} = 7.66 \pm 0.87 \text{ mV} \text{ for LFD} \text{ and } V_{max} = 6.94 \pm 1.21 \text{ mV} \text{ for HFD}; P = 0.7075, \text{ extra}$ sum-of-squares F test) (Fig. 6C and fig. S10). Yet, enhanced IBa was absent in arterial myocytes from AKAP150<sup>-/-</sup>, A, or S1928A HFD mice (Fig. 6C and fig. S10).

We also examined the relevance of Ser<sup>1928</sup> phosphorylation on the development of myogenic tone in diabetic mice. Vasoconstriction in response to 60 mM K<sup>+</sup>, in which membrane potential closely follows the equilibrium potential for K<sup>+</sup> (~-20 mV) (51), was significantly higher in arteries from wild-type HFD mice, but not in arteries from S1928A HFD mice, compared with those from corresponding LFD mice (table S2). In line with other reports (3, 40), arteries from wild-type HFD mice consistently developed greater myogenic tone compared to arteries from wild-type LFD mice over a range of intravascular pressures (Fig. 6, D and E). Arterial tone development was not different between arteries from HFD or LFD wild-type mice in the presence of the L-type calcium channel blocker nifedipine (fig. S11), suggesting that the greater myogenic response in HFD mice requires Ca<sub>V</sub>1.2 activity. In

contrast to the findings in arteries from wild-type mice, arterial tone was similar between arteries from S1928A mice on LFD and HFD at all intravascular pressures tested (Fig. 6, D and E). Together, these findings indicated a key role for AKAP150-anchored PKA, leading to phosphorylation of  $\alpha 1_C$  at Ser<sup>1928</sup> in stimulation of Ca<sub>V</sub>1.2 channel activity in HFD mice. Our results suggested that substitution of a single amino acid (Ser to Ala at residue 1928) to block Ser<sup>1928</sup> phosphorylation in the  $\alpha 1_C$  is sufficient to prevent the increase in Ca<sub>V</sub>1.2 channel activity and vascular tone in HFD mice.

### PKA mediates phosphorylation of $\alpha 1_C$ at Ser<sup>1928</sup> and increased Ca<sub>V</sub>1.2 activity in native human arterial myocytes from diabetic subjects

To test the translational significance of our observations, we analyzed native human arterial myocytes and arteries from nondiabetic and diabetic patients (table S3). In native contractile human arterial myocytes from nondiabetic patients, STORM/TIRF and PLA imaging revealed a subpopulation of Ca<sub>V</sub>1.2 within close proximity to PKA<sub>cat</sub> (Fig. 7, A and B, and fig. S12), similar to our results in murine arterial myocytes (Fig. 1). We tested a role for PKA in Ser<sup>1928</sup> phosphorylation and Ca<sub>V</sub>1.2 channel activity in response to increasing extracellular D-glucose from 5 to 15 mM in native human arterial myocytes from nondiabetic patients. We selected these concentrations because they are comparable to the blood glucose concentrations observed in nondiabetic and diabetic subjects. We found that 15 mM Dglucose increased IBa in cells from nondiabetic human subjects (Fig. 7, C and D). Global [Ca<sup>2+</sup>]<sub>i</sub> also significantly increased in response to 15 mM D-glucose in these cells, and this effect was attenuated by inhibition of PKA with rpcAMPS (Fig. 7, E and F). Moreover, incubation with 15 mM D-glucose for 5 min stimulated a significant increase in phosphorylation of  $\alpha 1_C$  at Ser<sup>1928</sup> in intact arteries from the non-diabetic subjects (Fig. 7, G and H). Preincubating the arteries in rpcAMPS abolished this increase, indicating that Ser<sup>1928</sup> phosphorylation during increased D-glucose depends upon PKA activity.

Consistent with our findings in HFD animals, we found that phosphorylation of  $\alpha 1_C$  at Ser<sup>1928</sup> was significantly increased in arterial lysates prepared from diabetic compared to those from non-diabetic human subjects, without any change in total Ca<sub>V</sub>1.2 protein abundance (Fig. 8A). We recorded single L-type Ca<sup>2+</sup> channel activity in the cell-attached configuration from freshly isolated arterial myocytes from nondiabetic and diabetic human subjects. Cells from diabetic subjects had significantly enhanced L-type Ca<sup>2+</sup> channel open probability (n*P*<sub>o</sub>) (Fig. 8, B and C). Preincubation of the cells from diabetic subjects with rpcAMPS significantly reduced L-type Ca<sup>2+</sup> channel n*P*<sub>o</sub> to values similar to those observed in cells from nondiabetic patients (Fig. 8, B and C), suggesting that PKA plays a key role in stimulation of L-type Ca<sup>2+</sup> channel activity during diabetes in human subjects. Together, our results are consistent with a mechanistic model whereby PKA activity determines phosphorylation of  $\alpha 1_C$  at Ser<sup>1928</sup>, leading to functional enhancement of L-type Ca<sup>2+</sup> channel activity in human arterial myocytes in response to increased D-glucose and in diabetes.

#### DISCUSSION

Many studies have revealed impaired myogenic autoregulation that may contribute to increased vascular reactivity in the diabetic state (7). This has been associated, at least in part, with enhanced contractility of arterial myocytes in response to increased blood D-glucose (hyperglycemia) during diabetes (2–6). The mechanisms underlying this hypercontractility remain the subject of intense investigation, but links have been made to changes in PKC and Rho kinase signaling (52, 53), transcriptional remodeling due to activation of the prohypertensive transcription factor NFATc3 (3, 54), and reduced K<sup>+</sup> channel abundance and channel activity (3, 40, 55, 56). The results of the present study provide strong support for an additional model in which direct phosphorylation of  $\alpha_1_C$  at Ser<sup>1928</sup> by an AKAP150-anchored PKA plays a key role in stimulating Ca<sup>2+</sup> influx through Ca<sub>V</sub>1.2 channels, which leads to vasoconstriction during increases in extracellular glucose and in diabetes (Fig. 8D).

A central observation in this study was that increasing the concentration of glucose triggered an increase in global  $[Ca^{2+}]_i$  in arterial myocytes, leading to vasoconstriction through a PKA-dependent pathway. These findings are paradoxical, because PKA activation plays a well-established role in arterial myocyte relaxation in response to endogenous and exogenous vasodilatory agents (57-61). Data reported here provide insight into this paradox by supporting a role for precise subsarcolemmal compartmentalization of a subpopulation of PKA in  $Ca_V 1.2$  channel regulation and vasoconstriction. Accordingly, targeting PKA to  $Ca_V 1.2$  channels by scaffolding proteins, such as AKAPs, may provide a means for selective phosphorylation of the channel to promote increased global Ca<sup>2+</sup> and vasoconstriction rather than vasodilation. When PKA is tethered by an AKAP, dissociation of the catalytic subunit may not be required for substrate phosphorylation (62). Hence, association with the AKAP complex may be indispensable for spatial limitation of kinase activity within discrete cellular compartments. To extend the range of activity of an AKAP-anchored PKA, intrinsically disordered regions within RIIa subunits afford a high degree of flexibility within the anchored PKA holoenzyme, giving the catalytic subunits ~20-nm radius of motion. Considering the length of antibodies used to label proteins (~10 to 15 nm), the intermolecular distances between Cav1.2 and PKAcat reported in the current study are conceivably within the limits required for selective modulation of  $Ca_V 1.2$  channel activity by PKA-mediated phosphorylation to promote contraction. In contrast, stimuli that induce cell-wide activation of PKA, such as forskolin, or activate a different PKA pool to stimulate other substrates, such as the phosphorylation of  $K^+$  channels in response to  $\beta$ -adrenergic agonists (61, 63), may cause relaxation. Therefore, we predict that the spatial and temporal characteristics of the activated PKA signal determine the functional effects of PKA activity on vascular reactivity.

Another fundamental finding was that enhanced Ca<sub>V</sub>1.2 channel activity in arterial myocytes in response to increases in extracellular D-glucose and in diabetes relies on phosphorylation of  $\alpha 1_C$  at Ser<sup>1928</sup>. This is a highly conserved residue among  $\alpha 1_C$  subunits and is phosphorylated by PKA in response to various stimuli, such as  $\beta$ -adrenergic signals (17–21). Here, we showed that, in murine arterial myocytes and arteries, PKA-dependent phosphorylation of  $\alpha 1_C$  at Ser<sup>1928</sup> enhances vascular Ca<sub>V</sub>1.2 channel activity and

vasoconstriction in response to increased extracellular D-glucose and in diabetic mice. Similar increases in PKA-mediated Ser<sup>1928</sup> phosphorylation, Ca<sub>V</sub>1.2 channel activity, and global Ca<sup>2+</sup> signals were found in nondiabetic human arteries and myocytes in response to increasing the concentration of glucose and in arteries and myocytes from diabetic human subjects, thus highlighting the consistency of the results from the murine model with human pathology. In agreement with an important role for Ser<sup>1928</sup>, we found that this single Ser-to-Ala substitution, which precludes phosphorylation of this site in S1928A knockin mice (21), prevented the enhancement in Cav1.2 channel activity and vasoconstriction in response to increased p-glucose. Moreover, enhanced  $Ca_V 1.2$  channel activity and myogenic tone in an HFD mouse model of diabetes were blocked in S1928A knockin mice on HFD, with no effect on basal Ca<sub>V</sub>1.2 channel function. A reduction in K<sup>+</sup> channel function has also been proposed to promote membrane depolarization and vasoconstriction in response to acute increases in extracellular D-glucose (55, 56). Our data showing no functional impact of acute high glucose on arterial diameter in arteries from S1928A mice argue that direct phosphorylation of  $\alpha 1_{C}$  at Ser<sup>1928</sup>, leading to membrane potential-independent increase in  $[Ca^{2+}]_i$ , is required for enhanced vasoconstriction. However, this Ca<sub>V</sub>1.2-dependent response could also act in concert with mechanisms of membrane potential dysregulation to exacerbate vascular tone in diabetes.

The essential influence of Ser<sup>1928</sup> on Ca<sub>V</sub>1.2 channel activity during PKA signaling in arterial myocytes is conserved in neurons, as revealed by Qian *et al.* (64). However, this phosphorylation event does not seem to be important for PKA-dependent regulation of Ca<sub>V</sub>1.2 channel activity in cardiac cells (21).  $\beta$ -Adrenergic stimulation of Ca<sub>V</sub>1.2 channels in ventricular myocytes from S1928A mice was normal, suggesting that Ser<sup>1928</sup> phosphorylation by PKA is not necessary for sympathetic stimulation of channel activity and contraction in murine hearts (21). In contrast, Qian *et al.* report that S1928A mice have impaired long-term potentiation and compromised  $\beta$ -adrenergic–mediated stimulation of Ca<sub>V</sub>1.2 activity by PKA in neurons, demonstrating a key role for Ser<sup>1928</sup> in  $\beta$ -adrenergic regulation of Ca<sub>V</sub>1.2 channel activity and neuronal function (64). The molecular mechanisms underlying these marked tissue-specific differences are unclear but may involve functional heterogeneity among distinct tissue-specific channel splice variants, a unique phosphorylation profile of  $\alpha$ 1<sub>C</sub>, posttranslational regulation of other interacting proteins at distinct sites, or differential interaction with or regulation by different scaffold proteins and accessory subunits.

Ser<sup>1928</sup> can also be phosphorylated by PKC (65), and activation of PKC increases  $Ca_V 1.2$  channel activity in arterial myocytes (66, 67). Enhanced activity of this kinase has also been associated with increased  $Ca_V 1.2$  channel activity and vascular reactivity during hypertension (68). We speculate that increased  $Ca_V 1.2$  channel activity during hypertension could be attributed to PKC-dependent phosphorylation of the same target residue. Thus,  $Ser^{1928}$  could represent a point of convergence to regulate  $Ca_V 1.2$  channel function by diverse signaling pathways. Furthermore, this phosphorylation site may be a distinct critical site of action for signal transduction in arterial myocytes, as well as a potential therapeutic target to treat vascular complications in diabetes and perhaps other pathological conditions.

Our previous studies have demonstrated that AKAP150 is necessary for regulation of vascular  $Ca_V 1.2$  channel by receptor-stimulated signals, such as angiotensin II (37, 68). This potent vasoconstrictor acts on G<sub>q</sub>-coupled AT1 receptors to stimulate phospholipase C, generate diacylglycerol, and activate PKC. AKAP150 associates with AT1 receptors (69), and binding of PKC to AKAP150 is necessary for both acute and chronic angiotensin II stimulation of vascular Ca<sub>V</sub>1.2 channels (37). Here, we confirmed the importance of AKAP150-bound PKC in modulating Cav1.2 channel activity and vascular reactivity during angiotensin II signaling using the C knockin mouse. We also discovered that anchored PKC was not necessary for enhanced  $Ca_V 1.2$  channel activity in response to increased Dglucose. In contrast, the AKAP150 mutant lacking the PKA-binding site ( A) failed to enhance D-glucose-mediated phosphorylation of  $a_{1C}$  at Ser<sup>1928</sup> to promote Ca<sub>V</sub>1.2 channel activity. Although the upstream mechanisms by which an increase in extracellular D-glucose leads to activation of AKAP150-anchored PKA in arterial myocytes are unclear, research from several groups suggested that glucose might stimulate a  $Ga_s$  signal (70, 71). This would activate adenylyl cyclase to produce cyclic adenosine 3',5'-monophosphate (cAMP), which leads to cAMP-dependent activation of PKA. The incorporation of AKAP150 in this pathway may coordinate the clustering of specialized G protein (heterotrimeric guanine nucleotide-binding protein)-coupled receptors, G proteins, and the downstream effectors to enable the decoding of different stimuli.

The augmented Ca<sup>2+</sup> influx observed in cells exposed to prolonged increases in extracellular D-glucose or cells from animals or humans with diabetes may also activate a prohypertensive transcriptional cascade involving calcineurin and NFATc3 (Fig. 8D) (54, 68, 72, 73). Consistent with this, NFATc3 signaling is activated in an HFD mouse model of diabetes through a process that requires AKAP150-anchored calcineurin and Ca<sup>2+</sup> influx through Ca<sub>V</sub>1.2 channels (3). However, the abundance of Ca<sub>V</sub>1.2 in arteries from wild-type HFD mice (3) and humans with diabetes (Fig. 8A) is similar to that in arteries from the corresponding LFD mice and nondiabetic subjects. Yet, NFATc3 activation leads to selective reduction in BK<sub>Ca</sub> β1 and K<sub>V</sub>2.1 subunit abundance and function, which may further contribute to enhanced vascular tone during diabetes (3, 40). Thus, we propose that AKAP150-anchored PKA and phosphorylation of  $\alpha 1_{\rm C}$  at Ser<sup>1928</sup>, leading to increased Ca<sub>V</sub>1.2 channel activity during chronic increases in extracellular D-glucose and diabetes, may activate AKAP150-anchored calcineurin to promote NFATc3 nuclear accumulation, transcriptional changes, and vascular dysfunction. Findings of restored myogenic tone in S1928A HFD animals may reflect the absence of an effect of diabetes on PKA-mediated Ca<sub>V</sub>1.2 channel stimulation and a normalization of the steady-state global [Ca<sup>2+</sup>]<sub>i</sub> as well as minimization of K<sup>+</sup> channel remodeling and normalization of membrane potential regulation through prevention of downstream NFATc3-dependent signaling. Therefore, regulatory events that are independent of membrane potential (PKA-mediated potentiation of  $Ca_V 1.2$ channel activity) and those that are dependent on membrane potential (down-regulation of  $K^+$  channel activity, leading to depolarization and further Ca<sub>V</sub>1.2 channel activation) may both modulate arterial myocyte  $[Ca^{2+}]_i$  and contractility during diabetes.

We found that  $Ca_V 1.2$  channel activity is higher in cells and arteries from HFD mice and human subjects with diabetes even when measurements were performed in the presence of normal low concentrations of p-glucose. These results are similar to those observed in

arterial myocytes from a dB/dB mouse model of diabetes (8). Both Navedo *et al.* (8) and we here found that these differences were eliminated by PKA inhibition, suggesting that increased  $Ca_V 1.2$  channel function is the result of higher PKA activity in cells from diabetic animals and human subjects, thus providing evidence for "molecular memory." At present, it is unclear whether this increase in PKA activity results from altered upstream signaling (for example, activity of adenylyl cyclase), impaired activity of a phosphodiesterase, or changes in PKA subunit abundance or regulation.

In summary, our data indicated that, upon an increase in extracellular glucose, phosphorylation of  $\alpha 1_C$  at Ser<sup>1928</sup> by AKAP-anchored PKA is essential for enhanced Ca<sub>V</sub>1.2 channel activity and increased global [Ca<sup>2+</sup>]<sub>i</sub>. Because contractility of arterial myocytes depends upon changes in global [Ca<sup>2+</sup>]<sub>i</sub>, higher Ca<sub>V</sub>1.2 channel open probability in response to increased glucose and during diabetes may stimulate vasoconstriction in the resistance vasculature. Thus, activation of this pathway may have profound implications for tissue perfusion and blood pressure regulation during chronic hyperglycemia and diabetes.

#### MATERIALS AND METHODS

#### Animals

Male mice (5 to 8 weeks of age) were used for this study in strict accordance with protocols approved by the Animal Care and Use Committee of the University of California, Davis. In some experiments, mice were maintained on either an LFD (10% kcal) or HFD (60% kcal) (Research Diets) starting at 5 weeks of age for 12 to 16 weeks. The composition of these diets and the propensity of mice maintained on this HFD to develop type 2 diabetes and induce vascular dysfunction of small resistance arteries have been well documented in other studies (3, 50, 74). Wild-type (C57BL/6J), AKAP150<sup>-/-</sup> (C57BL/6J), and knockin mice expressing AKAP150 lacking its PKA RII binding site (A; C57BL/6J) or PKC (C; C57BL/6J) or in which Ser<sup>1928</sup> of  $a_{1C}$  was mutated to Ala (S1928A) were euthanized by intraperitoneal injection of sodium pentobarbital (250 mg/kg). Pial cerebral arteries were acutely dissected, and arterial myocytes were isolated using enzymatic digestion techniques as described (3, 75). For some experiments, third-order mesenteric and femoral arteries were isolated, and arterial myocytes were dissociated following the procedure above. Arteries and arterial myocytes were maintained in ice-cold nominally  $Ca^{2+}$ -free Ringer's solution (5 mM KCl, 140 mM NaCl, 2 mM MgCl<sub>2</sub>, 10 mM Hepes, and 10 mM D-glucose adjusted to pH 7.4 with NaOH) containing normal 10 mM D-glucose until use.

#### Human arteries and cells

Excised human arteries from surgical patients undergoing gastric bypass procedure were obtained after institutional review board (IRB) approval and written consent and in accordance with the guidelines of the Declaration of Helsinki. Because our IRB exemption requires the use of noncoded tissue, a detailed therapeutic profile of the patient is unknown. Available patient information is included in table S3. Tissue was collected and placed in cold phosphate-buffered saline (PBS) solution containing normal D-glucose [138 mM NaCl, 3 mM KCl, 10 mM Na<sub>2</sub>HPO<sub>4</sub>, 2 mM NaH<sub>2</sub>PO<sub>4</sub>, 5 mM D-glucose, 0.1 mM CaCl<sub>2</sub>, and 0.1 mM MgSO<sub>4</sub> (pH adjusted to 7.4)] and transferred to the laboratory. Small-diameter (~150- to

250- $\mu$ m-diameter) arteries were carefully dissected out of surrounding tissue and used for experimentation. Arterial myocytes were enzymatically isolated using a two-step digestion process as described (3, 75). Cells were maintained in ice-cold nominally Ca<sup>2+</sup>-free Ringer's solution with normal D-glucose concentration (5 mM) until use.

#### Immunolabeling and STORM

Isolated arterial myocytes were allowed to adhere to a coverslip before fixing with PBS containing 3% paraformaldehyde + 0.1% glutaraldehyde followed by 0.1% sodium borohydride. After washing with PBS, cells were permeabilized and blocked with 0.5% Triton X-100 and 3% bovine serum albumin and exposed overnight to custom antibody against Ca<sub>V</sub>1.2 (28, 76) and antibody against PKA<sub>cat</sub>  $\alpha$ ,  $\beta$ ,  $\gamma$  (Santa Cruz Biotechnology). For secondary antibodies, we used a goat FLIP 565-conjugated antibody recognizing rabbit immunoglobulin G (IgG) and goat Alexa Fluor 647-conjugated antibody recognizing mouse IgG. The specificity of secondary antibodies to detect corresponding primary antibodies was tested in control experiments in which cells were only treated with secondary antibodies before imaging (-1° antibody control). We used a Nikon N-STORM super-resolution microscope system to capture images after TIRF illumination with 561- and 640-nm laser light. During imaging, cells were bathed in PBS (pH 7.4) containing glucose oxidase (0.56 mg/ml), catalase (0.3 mg/ml), and cysteamine (10mM). Aseries of 20,000 images were acquired using a Plan Apochromat TIRF 100× oil immersion lens [numerical aperture (NA), 1.49] and an iXon3 DU-897E electron-multiplying charge-coupled device (EMCCD) camera (Andor Technology). STORM images were rendered by plotting molecule localization as a Gaussian that was proportional to the number of photons released per photoswitching event. The full width at half maximum measured for single-molecule signals was ~31 nm. Reconstructions were filtered to reject localizations generated from less than 600 photons or from events that were detected on eight or greater consecutive frames. Half-maximal intensity for single molecules in reconstructed images was used to set a threshold for binary analysis, and the shortest intermolecular distance was determined for Cav1.2 and PKA<sub>cat</sub> using the JACoP plug-in of ImageJ software [National Institutes of Health (NIH)] as before (77, 78). Intermolecular distance histograms were fit with a sum of two Gaussian functions of the following equation:  $Y = A \times \exp(-0.5((X - \text{Mean}_1)/\sigma_1)^2) + B \times \exp(-0.5((X - \text{Mean}_1)/\sigma_1)^2)$ Mean<sub>2</sub>)/ $\sigma_2$ )<sup>2</sup>), where A and B are the heights of the center of the distribution of distances expressed as a percentage of total points, Mean<sub>1</sub> and Mean<sub>2</sub> are the x values of distance at the center of the two distributions, and  $\sigma_1$  and  $\sigma_2$  are the widths of each distribution in nanometers.

#### Structured illumination microscopy

Arterial myocytes were fixed, permeabilized, blocked, and incubated with primary antibodies against  $Ca_V 1.2$  and  $PKA_{cat}$  as described above. Goat Alexa Fluor 488–conjugated antibody against rabbit and goat Alexa Fluor 568–conjugated antibody against mouse were used as secondary antibodies. Images were captured using a Nikon N-SIM microscopy system coupled to a Nikon ECLIPSE Ti-E inverted microscope. The specimen was illuminated with 488- and 561-nm laser light by high-frequency patterned illumination. Images were captured at a focal plan at the *z*-axis center of an arterial myocyte using a CFI SR Apochromat TIRF 100× oil immersion lens and an iXon3 DU-897E EMCCD camera

(Andor Technology). Cellular protein ultrastructure was determined by reconstruction using a computer algorithm in Nikon Elements AR software. The percent of total  $Ca_V 1.2$ -associated centroids within 100 nm to the nearest PKA<sub>cat</sub>-associated centroids was determined using the JACoP plug-in of ImageJ software (NIH) as described above.

#### Immunofluorescence

Immunofluorescence labeling of freshly isolated arterial myocytes was performed as described (37) using a monoclonal antibody against Ca<sub>V</sub>1.2 (NeuroMab, clone N263/31; 1:1000) and TfR (Thermo Scientific, clone H68.4; 1:1000) and goat antibody against ANO1 (Santa Cruz Biotechnology, clone S-20; 1:1000). The secondary antibody was either a donkey Alexa Fluor 568–conjugated antibody against mouse (5 mg/ml) or donkey Alexa Fluor 568–conjugated antibody against goat (5 mg/ml) from Molecular Probes. Cells were imaged (512 × 512–pixel images) using an Olympus FV1000 confocal microscope coupled with an Olympus 60× water immersion lens (NA, 1.2) and a zoom of 3.0 (pixel size, 0.138  $\mu$ m). Images were collected at multiple optical planes (*z*-axis step size, 0.35  $\mu$ m). The specificity of the primary antibody was tested in negative control experiments in which the primary antibody was substituted with PBS. Ca<sub>V</sub>1.2-, TfR-, and ANO1-associated fluorescence was not detected under this experimental condition. The same laser power, gain settings, and pinhole were used to image arterial myocytes under different experimental conditions.

#### Proximity ligation assay

A Duolink In Situ PLA kit was used to detect complexes consisting of Ca<sub>V</sub>1.2 and PKA<sub>cat</sub> in freshly dissociated arterial myocytes. Cells were plated on glass coverslips and allowed to adhere (30 min, room temperature) before fixing with 4% paraformaldehyde (20 min), quenching in 100 mM glycine (15 min), and washing in PBS ( $2 \times 3$  min). Cells were then permeabilized with 0.1% Triton X-100 (20 min) and blocked (1 hour, 37°C) in 50% Odyssey blocking solution (LI-COR Biosciences). Cells were incubated overnight with a specific combination of primary antibodies [mouse antibody against PKAcat (Santa Cruz Biotechnology, sc-365615, 1:1000), custom rabbit antibody specific for  $Ca_V 1.2$  (1:1000) (28, 76), monoclonal antibody specific for  $Ca_V 1.2$  (NeuroMab, clone N263/31 AB\_11000167, 1:1000) and TfR (Thermo Scientific, clone H68.4, #13-6800, 1:1000), and goat antibody against ANO1 (Santa Cruz Biotechnology, clone S-20, sc-69343, 1:1000)] in Duolink antibody diluent solution. As a control, cells were incubated with only one primary antibody. Secondary antibodies conjugated with oligonucleotides (PLA probes, antibody against rabbit PLUS and antibody against mouse MINUS) were used to detect  $Ca_V 1.2$  and PKA, Ca<sub>V</sub>1.2 and TfR, and Ca<sub>V</sub>1.2 and ANO1 (1 hour, 37°C). After incubation with probes, a ligation solution consisting of two distinct oligonucleotides and ligase was added and incubated for 30 min at 37°C to allow hybridization and formation of a circular DNA template at sites of dual labeling. The ligation step is followed by a rolling circle amplification reaction (100 min,  $37^{\circ}$ C) using the ligated circle as a template. The ligation reaction was followed by washing in Duolink buffer B ( $2 \times 10$  min) and in 1% buffer B ( $1 \times$ 1 min). Coverslips were allowed to dry and subsequently mounted on a microscope slide with Duolink mounting medium. The fluorescence signal was visualized using an Olympus FV1000 confocal microscopy system on an Olympus IX81 microscope with a 60× oil

immersion lens (NA, 1.54). Images were acquired at different optical planes (*z*-axis step size,  $0.5 \mu m$ ). The stack of images for each sample was then combined into a single-intensity projection image that was subsequently used for analysis of number of puncta per square micrometer per cell. For presentation, representative PLA images at the center of the *z* axis are shown. Before labeled specimens were imaged, photomultiplier gain and laser power parameters were determined using control cells in which primary antibodies were omitted. Images for all conditions were obtained using the same acquisition parameters.

#### Electrophysiology

Freshly isolated arterial myocytes were allowed to adhere to a glass coverslip in a recording chamber for 10 min. Ca<sub>V</sub>1.2 channel activity was examined using the perforated whole-cell patch-clamp technique to record macroscopic currents with Ba<sup>2+</sup> as a charge carrier. The pipette solution contained 120 mM CsCl, 20 mM tetraethylammonium chloride (TEA-Cl), 1 mM EGTA, and 20 mM Hepes with amphotericin B (250 µg/ml; pH adjusted to 7.2 with CsOH). Cells were bathed in a solution containing 115 mM NaCl, 10 mM TEA-Cl, 0.5 mM MgCl<sub>2</sub>, 10 mM D-glucose, 5 mM CsCl, 20 mM BaCl<sub>2</sub>, and 20 mM Hepes (pH adjusted to 7.4 with CsOH). All electrophysiological experiments were performed at 21° to 23°C. Experiments were terminated by application of the experimental solution containing 1 µM nifedipine to determine the nifedipine-sensitive component, which is produced by Ca<sub>V</sub>1.2 channels containing a 1<sub>C</sub> subunits. To obtain the *I*-*V* relationship of nifedipine-sensitive Ba<sup>2+</sup> currents, we depolarized cells for 200 ms from the holding potential of -70 mV to voltages ranging from -60 to +60 mV. The *I*-*V* relationship for averaged data sets was fit with a peak Gaussian function:  $I(V) = I_{max} \times \exp(-0.5((V - V_{max})/b)^2)$ , where  $I_{max}$  is peak *I*,  $V_{max}$  is *V* at  $I_{max}$ , and *b* is the slope of the distribution, as described previously (79).

Single-channel Ca<sup>2+</sup> currents were recorded from arterial myocytes in the cell-attached configuration of the patch-clamp technique as described (80-82). Briefly, data were acquired at a sampling rate of 50 µs and low pass-filtered at 2 kHz with an Axopatch 200B amplifier and Digidata 1440 digitizer (Molecular Devices). Data were subsequently filtered with a Gaussian filter (400 Hz) during analysis. Pipettes were pulled from borosilicate capillary glass using a micropipette puller (model P-97, Sutter Instruments). Pipettes were polished to achieve resistances that range from 3.5 to 6.5 megohms. The pipette solution contained 120 mM TEA-Cl, 110 mM CaCl<sub>2</sub>, and 10 mM Hepes. BayK-8644 (500 nM) was included in the pipette solution to promote longer open times, thereby facilitating measurement of small unitary L-type Ca<sup>2+</sup> currents and increasing the probability of sweeps with channel activity. Bath solution contained 145 mM KCl, 10 mM NaCl, and 10 mM Hepes [pH 7.4 (NaOH)]. This high  $K^+$  solution was used to fix the membrane resting potential to ~0 mV (83, 84). Ltype  $Ca^{2+}$  currents were evoked by a 2-s step depolarization to -30 mV from the holding potential of -70 mV. Capacitive currents were compensated by subtraction of blank sweeps. Single-channel events were recorded using the Axopatch module of the pCLAMP 10 software. Cav1.2 channel activity was negligible when nifedipine was included in the patch pipette. The single-channel event half-amplitude detection algorithm from pCLAMP 10 was used to analyze single-channel openings and  $nP_0$ . The unitary channel amplitude in our single-channel recordings is ~0.53 pA, which corresponds to the expected single-channel

activity for  $Ca_V 1.2$  channels under our experimental conditions (81, 85). All experiments were performed at room temperature.

#### Arterial diameter measurements

Freshly isolated posterior cerebral arteries were cannulated on glass micropipettes mounted in a 5-ml myograph chamber (University of Vermont Instrumentation and Model Facility) as described (86). For some experiments, endothelial function was ablated by passing an air bubble through the lumen during the cannulation procedure. To allow for equilibration, arteries were pressurized to 20 mmHg and continuously perfused (37°C, 30 min, 3 to 5 ml/ min) with physiological saline solution consisting of 119 mM NaCl, 4.7 mM KCl, 2 mM CaCl<sub>2</sub>, 24 mM NaHCO<sub>3</sub>, 1.2 mM KH<sub>2</sub>PO<sub>4</sub>, 1.2 mM MgSO<sub>4</sub>, 0.023 mM EDTA, and 10 mM D-glucose aerated with 5% CO<sub>2</sub>/95% O<sub>2</sub>. Bath pH was closely monitored and maintained at 7.35 to 7.4. After equilibration period, intravascular pressure was increased and arteries were allowed to develop myogenic tone. Arteries not exhibiting stable myogenic tone after ~1 hour were discarded. To assess the response of arterial diameter to hyperglycemic conditions in pressurized vessels, D-glucose was increased in the perfusion solution from 10 to 20 mM. Arterial tone data are presented as a percent decrease in diameter relative to the maximum passive diameter at a given intravascular pressure obtained at the end of each experiment using Ca<sup>2+</sup>-free saline solution containing 1  $\mu$ M nifedipine.

#### [Ca<sup>2+</sup>]<sub>i</sub> measurements

Global cell-wide  $[Ca^{2+}]_i$  was recorded as described (8). Briefly, global cytosolic  $[Ca^{2+}]_i$  was imaged in isolated arterial smooth muscle cells loaded with the membrane-permeable acetoxymethyl ester Fluo-4 (15 min, room temperature). Cells were bathed in a Hepesbuffered saline solution containing 140 mM NaCl, 5 mM KCl, 1 mM MgCl<sub>2</sub>, 2 mM CaCl<sub>2</sub>, 10 mM Hepes, and 10 or 20 mM D-glucose. Images were collected using an Andor spinning disk confocal system coupled to an Olympus iX-81 inverted microscope equipped with an Olympus 60× oil immersion lens (NA, 1.49). Background-subtracted images were normalized by dividing the fluorescence intensity of each pixel (*F*) by the average resting fluorescence intensity (*F*<sub>0</sub>) of a confocal image to determine the *F*/*F*<sub>0</sub>-time relationship.

#### Immunoprecipitation and immunoblotting

Whole lysates were obtained from cerebral and mesenteric arteries by sonication (20 min,  $4^{\circ}$ C) in lysis buffer solution containing 150 mM NaCl, 10 mM Na<sub>2</sub>HPO<sub>4</sub>, and 1 mM EDTA with 1% deoxycholic acid, 0.1% SDS, and protease inhibitors (cOmplete Mini Protease Inhibitor Cocktail, Roche) with 2 µM microcystin and 1 mM *p*-nitrophenyl phosphate to inhibit endogenous phosphatase activity. Samples were cleared from nonsolubilized material by ultracentrifugation (250,000*g* for 30 min) before immunoprecipitation (4 hours, 4°C) with an antibody against Ca<sub>V</sub>1.2 [4 µg of FP1 (28)] using protein A–Sepharose. An equal amount of nonspecific rabbit IgG was used as control for immunoprecipitation. Beads were washed three times with washing buffer [150 mM NaCl, 10 mM EDTA, 10 mM EGTA, 10 mM tris-HCl, 0.1% Triton X-100 (pH 7.4)]. Samples were boiled in Laemmli Sample Buffer (Bio-Rad) for 5 min at 95°C and run on an ~8% acrylamide gel in SDS–polyacrylamide gel electrophoresis before they were transferred onto polyvinylidene difluoride membranes. Membranes were blocked in blocking solution [5% dry milk in tris-buffered saline (TBS)

containing 0.05% Tween 20] before incubation with primary antibodies (blocking solution for 1 hour) followed by three 15-min washes in TBS with 0.05% Tween 20, incubation with horseradish peroxidase (HRP)–protein A (1:10,000 in blocking solution for 1 hour), several washes for 2 hours, and detection of HRP with enhanced chemiluminescence (ECL), ECL Plus, or Femto chemiluminescence reagents. Primary antibodies were from Santa Cruz Biotechnology ( $\alpha$ -tubulin, sc-32293), BD Transduction Laboratories (PKA-RII $\alpha$ , #612242), EMD Millipore (AKAP150, #07210), NeuroMab (K<sub>V</sub>2.1, clone K89/34), Alomone (BK<sub>Ca</sub>  $\alpha$ , APC-021), Abcam (BK<sub>Ca</sub>  $\beta$ 1, ab3587), and Cell Signaling Technology (PKC $\alpha$ ). Multiple exposures with increasing time periods were obtained to ensure that signals were in the linear range, as described (17, 34).

#### Chemicals and statistical analyses

All chemical reagents were from Sigma-Aldrich unless otherwise stated. Data are expressed as means  $\pm$  SEM. Data were analyzed using GraphPad Prism software. Statistical significance was determined using appropriate paired or unpaired Student's *t* test, ANOVA, or nonparametric tests. Statistical significance, denoted by asterisk in figures, was considered at *P*<0.05.

#### Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

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#### Fig. 1. Spatial organization of Ca<sub>V</sub>1.2 and PKA in murine arterial myocytes

(A) Representative conventional TIRF images (top) and corresponding STORM reconstruction maps (bottom) from a murine arterial myocyte labeled for Ca<sub>V</sub>1.2 and PKA<sub>cat</sub> (scale bar, 2 µm). Lower panels are enhanced magnification of areas shown in yellow boxes (scale bar, 100 nm). (B) Histograms showing the area of clusters of Ca<sub>V</sub>1.2 channels and PKA<sub>cat</sub> in isolated arterial myocytes. (C) Expanded merged images (top) and associated *x-y* fluorescence intensity profiles (bottom) of sites of close interaction between Ca<sub>V</sub>1.2 (green) and PKA<sub>cat</sub> (red; scale bars, 100 nm). (D) Histogram of the lowest intermolecular distance to PKA<sub>cat</sub> centroids for Ca<sub>V</sub>1.2 fluorescence particles. Data were fit with a sum of two Gaussian functions with depicted  $R^2$  and centroids (n = 10,306 particles from nine cells). Differential interference contrast (DIC; top) and fluorescence PLA (red)/4',6-diamidino-2phenylin-dole (DAPI) (blue) images (E) and quantification of PLA fluorescent puncta per square micrometer cell area for isolated arterial myocytes (F) labeled for Ca<sub>V</sub>1.2 (n = 20cells), PKA<sub>cat</sub> (n = 18 cells), or Ca<sub>V</sub>1.2 + PKA<sub>cat</sub> (n = 19 cells).



**Fig. 2. AKAP150 targets PKA to a subpopulation of Cav1.2 channels in arterial myocytes** (**A**) Representative immunoprecipitation (IP) of AKAP150 followed by immunoblot detection of AKAP150, PKA RIIa, and PKCa in lysates from wild-type (WT), A, and *AKAP150<sup>-/-</sup>* mice (n = 3 lysates). (**B**) Representative SIM reconstructions of isolated WT, A, and *AKAP150<sup>-/-</sup>* arterial myocytes labeled with antibodies against Cav1.2 and PKA<sub>cat</sub>. Object-based analysis shows location of objects (red crosses) corresponding to Cav1.2 and PKA<sub>cat</sub> fluorescence overlap. Scale bars, 5 µm. (**C**) Enlarged SIM (upper) and associated object-based colocalization (lower) images for Cav1.2 (yellow centroids) and PKA<sub>cat</sub> (red particles) of regions marked by yellow boxes in image in (B). Scale bar, 200 nm. (**D**) Percentage of total Cav1.2 channels associated with PKA<sub>cat</sub> at a distance of <100 nm for WT (n = 7 cells), A (n = 6 cells), and *AKAP150<sup>-/-</sup>* (n = 8 cells) arterial myocytes (\*P < 0.05, Kruskal-Wallis test). (**E**) Summary bar plot showing the density of sarcolemma-associated particles of Cav1.2 and PKA<sub>cat</sub> in TIRF/STORM images for arterial myocytes from WT (n = 6 cells) and A (n = 9 cells) animals (\*P < 0.05, unpaired *t* test of WT PKA<sub>cat</sub> versus A PKA<sub>cat</sub>).



Fig. 3. Enhanced Ca<sub>V</sub>1.2 channel activity and elevation of global  $[Ca^{2+}]_i$  in arterial myocytes in response to 20 mM D-glucose require anchoring of PKA by AKAP150

(A) Representative  $I_{Ba}$  in WT arterial myocytes evoked by 200-ms depolarization steps from holding potential of -70 mV to voltages ranging from -20 to +40 mV in 10 mM D-glucose (D-glu) and 5 min after application of 20 mM D-glucose. (B)  $I_{Ba}$ -voltage relationship (-60 to +60 mV) in WT arterial myocytes in the presence of 10 and 20 mM D-glucose (n = 8 cells; \*P < 0.05, paired t test). (C) Representative  $I_{Ba}$  from A and  $AKAP150^{-/-}$  arterial myocytes in response to step depolarizations from -70 to +10 mV before and after application of 20 mM D-glucose. (D) Amalgamated  $I_{Ba}$  (to +10 mV) before and after application of 20 mM Dglucose in WT, A, and  $AKAP150^{-/-}$  arterial myocytes (n = 8 to 11 cells; \*P < 0.05, paired t test). (E) Exemplary [Ca<sup>2+</sup>]<sub>i</sub> traces in isolated WT,  $AKAP150^{-/-}$ , and A arterial myocytes in the presence of 10 mM D-glucose and in response to 20 mM D-glucose in shaded time frame. (F) Plot of change in Ca<sup>2+</sup> ( $F/F_0$ ) to 20 mM D-glucose in isolated WT (n = 7 cells),  $AKAP150^{-/-}$  (n = 8 cells), and A (n = 7 cells) arterial myocytes [\*P < 0.05; one-way analysis of variance (ANOVA) with Tukey post hoc test].



Fig. 4. Anchoring of PKC by AKAP150 is necessary for angiotensin II, but not potentiation of Ca<sub>V</sub>1.2 channel activity in response to elevated glucose in arterial myocytes (A) Immunoprecipitation of AKAP150 from arterial lysates from WT and C animals followed by immunoblot detection of AKAP150, PKCa, and PKA RIIa (n = 3 lysates). (**B**) Representative diameter recordings in pressurized arteries (60 mmHg) before and after application of 100 nM angiotensin II (Ang II) with subsequent application of 1 µM nifedipine (Nifed) from WT (n = 5 arteries from three animals) and C (n = 6 arteries from four animals) middle cerebral arteries (starting and passive diameters for representative recordings: 118 and 147 µm for WT and 109 and 136 µm for C, respectively). (C and D) Summary vascular tone data in the absence (ct) and presence of angiotensin II and angiotensin II + nifedipine (C), and angiotensin II-induced constriction (D) in arteries from WT and C mice (\*P < 0.05, Mann-Whitney U test). (E) Representative  $I_{Ba}$  from C arterial myocytes in response to step depolarizations to +10 mV from a holding potential of -70 mV before and after application of angiotensin II or 20 mM D-glucose (n = 8 cells per condition). (F) Summary  $I_{Ba}$  data from C arterial myocytes before and after application of angiotensin II or 20 mM  $_{\rm D}$ -glucose (\*P< 0.05, one-way ANOVA with Tukey post hoc test).



# Fig. 5. Phosphorylation of Ser<sup>1928</sup> by AKAP150-targeted PKA is required for increased Ca<sub>V</sub>1.2 channel activity, enhanced global $[{\rm Ca}^{2+}]_i$ , and vasoconstriction in response to 20 mM D-glucose in arterial myocytes

(A) Representative immunoblot detection of phosphorylated Ser<sup>1928</sup> (pSer<sup>1928</sup>) and total Ca<sub>V</sub>1.2 from WT, A, and AKAP150<sup>-/-</sup> arteries after incubation with D-glucose concentrations, as indicated. (B) Quantification by densitometric analysis of pSer<sup>1928</sup> normalized to total respective Ca<sub>V</sub>1.2 for WT, A, and AKAP150<sup>-/-</sup> arteries incubated with either 10 or 20 mM D-glucose (n = 4 arterial lysates per condition; \*P < 0.05, Mann-Whitney Utest). a.u., arbitrary units. (C) Characteristic  $I_{Ba}$  recordings from WT, WT + 10  $\mu$ M rpcAMPS, and S1928A arterial myocytes in response to step depolarizations to +10 mV from a holding potential of -70 mV before (10 mM D-glucose) and after application of 20 mM D-glucose. (**D**) Summary data showing  $I_{Ba}$  for WT (n = 10 cells), WT + rpcAMPS (n5 cells), and S1928A (n = 9 cells) arterial myocytes in 10 and 20 mM D-glucose (\*P < 0.05, paired t test). (E) Representative fluorescence traces showing global  $[Ca^{2+}]_i$  and plot of changes in global  $[Ca^{2+}]_i$  in response to 20 mM D-glucose (shaded area) in isolated WT (n =9 cells) and S1928A (n = 11 cells) arterial myocytes (\*P < 0.05, unpaired t test). (F) Representative diameter recordings and summary arterial tone (percent decrease in diameter from maximum passive diameter) data from pressurized (60 mmHg) WT and S1928A arteries in response to increasing extracellular D-glucose from 10 to 20 mM (n = 6 arteries per condition). Starting and passive diameters for representative recordings: 111 and 132 µm for WT and 126 and 155  $\mu$ m for S1928A, respectively (\*P < 0.05, Wilcoxon matched-pairs signed-rank test).



Fig. 6. Increased Ca<sub>V</sub>1.2 channel activity and enhanced vascular tone during diabetes require AKAP150-PKA phosphorylation of  $\alpha 1_C$  at Ser^{1928} in vivo

(A) Representative immunoblot detection of pSer<sup>1928</sup> and total Ca<sub>V</sub>1.2 in arteries isolated from WT and A LFD and HFD animals. (B) Densitometric quantification of pSer<sup>1928</sup>/  $Ca_V 1.2$  ratio in lysates from WT and S1928A LFD and HFD mice (n = 3 lysates from three mice per condition) (\*P < 0.05, Mann-Whitney Utest). (C)  $I_{Ba}$ -voltage relationship (-60 to +60 mV) in arterial myocytes from WT (LFD, n = 14 cells from five mice; HFD, n = 9 cells from five mice), AKAP150<sup>-/-</sup> (LFD, n = 5 cells from two mice; HFD, n = 5 cells from two mice), A (LFD, n = 6 cells from three mice; HFD, n = 6 cells from two mice), and S1928A (LFD, n = 8 cells from three mice; HFD, n = 9 cells from three mice) mice in LFD and HFD (\*P < 0.05, unpaired t test). (**D**) Representative diameter records obtained using arteries isolated from WT and S1928A mice in LFD and HFD over a range of intravascular pressures from 20 to 100 mmHg. Arteries were exposed to 1 µM nifedipine at 100 mmHg to assess contribution of  $Ca_V 1.2$ -mediated  $Ca^{2+}$  influx to tone generation in each group. (E) Bar plots summarizing arterial tone, expressed as a percent decrease in diameter from maximum passive diameter at a given pressure for arteries in WT and S1928A mice in LFD and HFD (n = 7 to 8 arteries from three animals per group; \*P < 0.05, Mann-Whitney U test).



## Fig. 7. A subpopulation of Ca<sub>V</sub>1.2 associates with PKA to stimulate Ca<sub>V</sub>1.2 channel activity, global Ca<sup>2+</sup>, and Ser<sup>1928</sup> phosphorylation in response to elevated glucose in native human arterial myocytes

(A) Differential interference contrast (top) and fluorescence PLA (red)/DAPI (blue) images of freshly dissociated human arterial myocytes labeled for Ca<sub>V</sub>1.2 or PKA<sub>cat</sub> or Ca<sub>V</sub>1.2 + PKA<sub>cat</sub> from nondiabetic patients. (**B**) Quantification of PLA fluorescent puncta per square micrometer cell area for human arterial myocytes labeled for Ca<sub>V</sub>1.2 (n = 14 cells), PKA<sub>cat</sub> (n = 15 cells), and Ca<sub>V</sub>1.2 + PKA<sub>cat</sub> (n = 16 cells) (\*P < 0.05, one-way ANOVA with Tukey post hoc test).  $I_{Ba}$  records (from -70 to +10 mV) (**C**) and amalgamated current density data (**D**) obtained from freshly dissociated human arterial myocytes from nondiabetic subjects exposed to 5 and 15 mM D-glucose (n = 7 cells from three patients) (\*P < 0.05, paired t test). (**E** and **F**) [Ca<sup>2+</sup>]<sub>i</sub> records and summary data of the response ( $F/F_0$ ) to increasing bath D-glucose from 5 to 15 mM in freshly dissociated arterial myocytes from nondiabetic subjects in the absence (n = 11 cells from four patients) or presence (n = 9 cells from two patients) of 10 µM rpcAMPS (\*P < 0.05, unpaired t test). (**G**) Immunoblot detection of pSer<sup>1928</sup> and total Ca<sub>V</sub>1.2 in human arteries incubated (10 min) with 5 or 15 mM D-glucose in the absence or presence of rpcAMPS. (**H**) Normalized densitometric quantification of pSer<sup>1928</sup> relative to total Ca<sub>V</sub>1.2 (n = 4 lysates from four patients for each) (\*P < 0.05, Mann-Whitney *U* test).



### Fig. 8. Arterial myocytes from diabetic patients exhibit enhanced $\mathrm{Ser}^{1928}$ phosphorylation and PKA-dependent augmentation of $\mathrm{Ca_V}1.2$ channel activity

(A) Representative immunoblot detection of pSer<sup>1928</sup>, total Ca<sub>V</sub>1.2, and α-tubulin and densitometric summary data for total Ca<sub>V</sub>1.2 (relative to α-tubulin) and pSer<sup>1928</sup> (relative to total Ca<sub>V</sub>1.2) in arterial lysates isolated from nondiabetic (n = 4) and diabetic patients (n = 4) (\*P < 0.05, Mann-Whitney U test). (**B**) Representative elementary Ca<sub>V</sub>1.2 channel records obtained during a 2-s step depolarization from -80 to -30 mV in freshly dissociated human arterial myocytes from non-diabetic (Non-db) and diabetic (db; ±10 µM rpcAMPS) patients. Channel openings (o) are represented by downward deflections from baseline (c). (**C**) Plot shows Ca<sub>V</sub>1.2 channel n $P_0$  for cells from nondiabetic patients (n = 27 cells from six patients) and diabetic patients in the absence (n = 24 from four patients) or presence (n = 14 cells from three patients) of rpcAMPS (\*P < 0.05, Kruskal-Wallis test). LTCC, L-type Ca<sup>2+</sup> channel. (**D**) Proposed model of Ca<sub>V</sub>1.2 channel stimulation upon increases in extracellular glucose and in diabetes.