Phenotypic changes in mouse pancreatic stellate cell Ca$^{2+}$ signaling events following activation in culture and in a disease model of pancreatitis

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ABSTRACT The specific characteristics of intracellular Ca$^{2+}$ signaling and the downstream consequences of these events were investigated in mouse pancreatic stellate cells (PSC) in culture and in situ using multiphoton microscopy in pancreatic lobules. PSC undergo a phenotypic transformation from a quiescent state to a myofibroblast-like phenotype in culture. This is believed to parallel the induction of an activated state observed in pancreatic disease such as chronic pancreatitis and pancreatic cancer. By day 7 in culture, the complement of cell surface receptors coupled to intracellular Ca$^{2+}$ signaling was shown to be markedly altered. Specifically, protease-activated receptors (PAR) 1 and 2, responsive to thrombin and trypsin, respectively, and platelet-derived growth factor (PDGF) receptors were expressed only in activated PSC (aPSC). PAR-1, ATP, and PDGF receptor activation resulted in prominent nuclear Ca$^{2+}$ signals. Nuclear Ca$^{2+}$ signals and aPSC proliferation were abolished by expression of activated PSC (aPSC). PAR-1, ATP, and PDGF receptor activation resulted in prominent nuclear Ca$^{2+}$ signals. Nuclear Ca$^{2+}$ signals and aPSC proliferation were abolished by expression of parvalbumin targeted to the nucleus. In pancreatic lobules, PSC responded to agonists consistent with the presence of only quiescent PSC. aPSC were observed following induction of experimental pancreatitis. In contrast, in a mouse model of pancreatic disease harboring elevated K-Ras activity in acinar cells, aPSC were present under control conditions and their number greatly increased following induction of pancreatitis. These data are consistent with nuclear Ca$^{2+}$ signaling generated by agents such as trypsin and thrombin, likely present in the pancreas. PSC are present in a periacinar and periductal localization of the exocrine pancreas following a meal results in the production of a fluid rich in HCO$_3^-$ and containing a complex mixture of proteins (Williams and Yule, 2006). The proteins are predominantly inactive precursors of digestive enzymes that are subsequently activated in the lumen of the duodenum. Two epithelial cell types are primarily responsible for secretion from the gland. Acinar cells synthesize, store, and undergo regulated exocytosis of secretory granules while duct cells are responsible for the aqueous component of the secretion. Together, these cells result in the formation and delivery of pancreatic juice to the duodenum. A third, less studied cell type, pancreatic stellate cells (PSC), are also resident in the exocrine pancreas. PSC are present in a periacinar and periductal localization (Apte et al., 1998; Bachem et al., 1998; Omary et al., 2007). In common with hepatic stellate cells, they are characterized as retinol/ lipid-storing cells expressing a variety of intermediate filament proteins including desmin and glial fibrillary acid protein (GFAP) (Wake, 1980; Kordes et al., 2009). Under physiological conditions...
stellate cells appear to be benign, and relatively little is known regarding their contribution to the normal function of the gland.

PSC have, however, attracted considerable interest because of their prominent role in the etiology of pancreatic pathology (Omary et al., 2007). For example, in disease states of the pancreas including chronic pancreatitis and pancreatic cancer, stellate cells undergo a phenotypic transformation to a so-called activated form. Activated PSC (aPSC) are highly proliferative and adopt a myofibroblastic state characterized by enlargement of the endoplasmic reticulum (ER) and nucleus together with the expression of α-smooth muscle actin (α-SMA). Significantly, aPSC, again similarly to hepatic stellate cells, produce large amounts of extracellular matrix proteins (ECM) resulting in the extensive pancreatic fibrosis observed in disease states (Wake, 1980; Apte et al., 1996, 2004; Haber et al., 1999; Casini et al., 2000; Omary et al., 2007; Kordes et al., 2009).

In vivo, it is likely that the signal transduction pathways that initiate and maintain the activated phenotype are elicited by events triggered by insult and mediated by surrounding cells including infiltrating leukocytes and damaged acinar cells. Much of the current understanding of the stellate cell signal transduction has been gleaned from studies of rodent PSC, isolated and maintained in culture. Initially the cultured cells express markers consistent with a quiescent phenotype, including the presence of prominent cytoplasmic lipid droplets (Apte et al., 1998). However, following short-term culture, cells transform to an activated, proliferative phenotype expressing abundant α-SMA and ECM proteins (Haber et al., 1999). Using this paradigm, inflammatory agents, growth factors, reactive oxygen species, and autocrine/paracrine factors have been shown to act on stellate cells (Omary et al., 2007). For example, PSC proliferate in response to platelet-derived growth factor (PDGF) and transforming growth factor-β as well as proinflammatory cytokines (Apte et al., 1999; Luttenberger et al., 2000; Shek et al., 2002). These studies have implicated the MAP kinase cascade and JAK/STAT and SMAD pathways as contributing to the processes of transformation and proliferation in PSC (Masamune et al., 2004; Ochni et al., 2004; Kikuta et al., 2006). There is, however, a relative lack of data relating to other signaling systems that could potentially result in cell growth and/or transformation. In hepatic stellate cells, intracellular Ca2+ signals have been reported to induce proliferation (Soliman et al., 2009). Furthermore, agents proposed to act on PSC, including growth factors and angiotensin II, have the potential to couple to signal transduction pathways, which results in intracellular Ca2+ elevations (Luttenberger et al., 2000; Hama et al., 2006). Little information is available regarding the agonists linked to intracellular Ca2+ signaling in PSC together with the specific temporal and spatial characteristics of these signals. Moreover, data are lacking relating to the functional consequences of elevations in intracellular Ca2+.

Given this gap in knowledge, the goals of the present study were to characterize Ca2+ signaling events and the consequences of these signals in cultured PSC in the quiescent (qPSC) and the transformed activated state. An important consideration is that in culture these events may not be identical to those taking place during disease progression in situ. Therefore, to study these events under conditions where the architecture of the pancreas and structural relationship between cell types are relatively unperturbed, the studies are extended to investigate Ca2+ signaling in lobules of intact pancreas using multiphoton (MP) microscopy in both physiological situations and in a model of chronic pancreatitis. Data are presented that provide evidence that the complement of cell surface receptors coupled to Ca2+ signaling is altered during activation of PSC, effectively providing a diagnostic feature of the activated phenotype. Further, the specific spatial characteristics of the Ca2+ signal are important for enhancing proliferation of aPSC. In total, these data are consistent with Ca2+ signaling events triggered by factors, including those likely released following initial acinar cell damage, resulting in proliferation of PSC and contributing to the pathology of disease.

RESULTS

Characterization of Ca2+ signaling in cultured qPSC

PSC undergo morphological and phenotypic changes in culture (Apte et al., 1998). To establish the temporal sequence of these changes, PSC morphology and expression of established protein markers were monitored after isolation. On day 2 in culture, cells displayed prominent perinuclear structures, which appeared phase bright (Figure 1Aa) and stained with the lipid marker AdipoRed (Figure 1Ab). The lipid droplets could also be visualized following MP excitation at 810 nm of intrinsic fluorescence, consistent with the presence of vitamin A in the droplets known to fluoresce when excited at this wavelength (Figure 1Ac and d) (Zipfel et al., 2003). Abundant expression of the intermediate filament protein vimentin (Figure 1Ae), together with low levels of desmin and GFAP (unpublished data) were also observed in cells fixed with paraformaldehyde and processed for immunocytochemistry. In contrast, at this time point, the cells did not stain for α-SMA, the prototypical marker for aPSC (Figure 1Af). Intracellular Ca2+ signals were also monitored in fura-2-loaded qPSC using digital imaging on day 2 in culture following isolation. An increase in [Ca2+]i, was observed in qPSC in response to a high concentration of the muscarinic agonist carbachol (CCCh; >1 μM), typically consisting of oscillatory changes in the [Ca2+]i, even at concentrations as high as 10 μM (Figure 1B). qPSC also responded to angiotensin II (>100 nM; Figure 2B) and bradykinin (>100 nM; Figure 1C), and ATP (>1 μM; Figure 1C). The initial Ca2+ peak in response to these agonists was largely independent of extracellular Ca2+ and thus is likely the result of the Gq-coupled formation of InsP3 and Ca2+ release from intracellular compartments (Figure 1D). No elevation in [Ca2+]i, was evoked in qPSC when exposed to 50 mM KCl, trypsin, or thrombin, indicating that qPSC are unlikely to express voltage-gated Ca2+ channels or the protease-activated receptors (PAR) family of Gq-coupled cell surface receptors (Figure 1E). In addition, stimulation of qPSC with PDGF or cholecystokinin (CCK) did not result in an elevation of [Ca2+]i, (unpublished data).

Characterization of aPSC in culture

By day 5 in culture, phase-bright lipid droplets were not visible and the cells had adopted a more flattened myofibroblast-type morphology accompanied by the expression of α-SMA in a fraction of cells (Figure 2Aa). All cells expressed α-SMA by day 7 (Figure 2Ab) together with increased expression of vimentin (Figure 2Ac) and desmin (Figure 2Ad). The changes in morphology and expression profile are typical of the transformation of qPSC to the activated phenotype. Ca2+ signaling was also studied after day 7 in culture, when all cells express α-SMA and are assumed to represent an activated phenotype. In a similar manner to qPSC, aPSC exhibited an increase in [Ca2+]i, in response to angiotensin II (>100 nM; Figure 2Ba) and bradykinin (>100 nM; Figure 2Bb) but not 50 mM KCl (Figure 2Bc) or CCK up to 50 nM (unpublished data). Exposure of aPSC to ATP also resulted in an increase in [Ca2+]i, (Figure 2C); however, the threshold for eliciting a response was approximately threefold lower in aPSC when compared with qPSC (Figure 2D). In contrast to qPSC, aPSC were refractory to stimulation with CCK at concentrations as high as 10 μM (Figure 2Ea) but elicited robust elevations in Ca2+ in response to the PAR agonists thrombin (Figure 2Eb) and trypsin (Figure 2Ec) and to PDGF (Figure 2Ed). In a similar fashion, human stellate cells exhibited morphological markers of aPSC when monitored at day 7 in culture.
including α-SMA expression, prominent GFAP, and desmin and vimentin staining (Supplemental Figure 1A). These cells responded robustly to PAR-1 and PAR-2 agonists and bradykinin but were also refractory to stimulation with CCK up to 50 nM (Supplemental Figure 1B). The absence of CCK-stimulated Ca\(^{2+}\) signals in both mouse and human PSCs is, however, not consistent with two recent studies that have indicated that both CCK-1 and CCK-2 receptors are expressed on human (Phillips et al., 2010) and rat PSC (Berna et al., 2010). It should be noted that neither study monitored CCK-stimulated Ca\(^{2+}\) signaling events; thus the formal possibility exists that CCK receptors in PSC do not signal in a conventional manner. The human study suggested that CCK stimulation resulted in acetylcholine release without affecting PSC proliferation or the mitogen-activated protein kinase (MAPK) signaling module. In contrast, the latter study demonstrated robust proliferation in response to CCK that was mediated by the MAPK and phosphatidylinositol 3-kinase signaling cascades. Although the reasons for these discrepancies are not clear, it is possible that species differences or variations in the phenotypic state of the PSC examined might contribute to this lack of accord.

**Periacinar cells in pancreatic lobules have characteristics of qPSC**

To evaluate the phenotype of PSC in situ in terms of Ca\(^{2+}\) signaling characteristics, pancreatic lobules were prepared from wild-type (WT) animals and loaded with both Fluo-4 AM and calcein AM. The latter dye was used to visualize lobule structure independently of [Ca\(^{2+}\)]. Figure 3 shows a maximum projection image of a lobule (∼120 μm thick) generated using MP microscopy of calcein fluorescence at the indicated magnifications (Figure 3A). Because the lobules were prepared without enzymatic digestion and with minimal preparative...
The morphology of these cells could also be clearly distinguished from neurons visualized in pancreatic lobules isolated from animals engineered to express green fluorescent protein specifically in neurons. These cells were relatively rare in the fields selected and characterized by long projections extending from the cell bodies (Supplemental Figure 2). In this population of periacinar cells, robust $[\text{Ca}^{2+}]_i$ changes could be evoked by ATP (>1 µM) (Figures 4, A, images, and B–E, kinetics) as well as bradykinin (Figure 4C), angiotensin (Figure 4D), and high concentrations of CCh (Figure 4E). An increase in $[\text{Ca}^{2+}]_i$ was never observed in periacinar cells in lobules exposed to thrombin (Figure 4E), trypsin (Figure 4F), CCK (Figure 4C), or PDGF (unpublished data). However, acinar cells in the intervention, the tissue retained obvious exocrine morphology, exemplified by the phase-dark apical localization of zymogen granules in the acinar clusters. In identical lobules prepared for immunocytochemistry, acinar cells could be readily visualized by the expression of $\alpha$-amylase while vimentin was distributed in periacinar/ductal cells (Figure 3B). $\alpha$-SMA was expressed only in ductal structures, indicating the scarcity/absence of aPSC in this preparation. Nonacinar cells could also be readily visualized in the live lobules, including cells with a periacinar distribution, which exhibited preferential loading of both fluorescent dyes (Figure 3A, arrows). Periacinar cells also exhibited prominent punctate intrinsic fluorescence when excited at 810 nm, again consistent with the presence of vitamin A in lipid droplets of qPSC (Figure 3A). The morphology of these cells could also be clearly distinguished from neurons visualized in pancreatic lobules isolated from animals engineered to express green fluorescent protein specifically in neurons. These cells were relatively rare in the fields selected and characterized by long projections extending from the cell bodies (Supplemental Figure 2). In this population of periacinar cells, robust $[\text{Ca}^{2+}]_i$ changes could be evoked by ATP (>1 µM) (Figures 4, A, images, and B–E, kinetics) as well as bradykinin (Figure 4C), angiotensin (Figure 4D), and high concentrations of CCh (Figure 4E). An increase in $[\text{Ca}^{2+}]_i$ was never observed in periacinar cells in lobules exposed to thrombin (Figure 4E), trypsin (Figure 4F), CCK (Figure 4C), or PDGF (unpublished data). However, acinar cells in the
performed to define in detail the characteristics of PAR-induced $[\text{Ca}^{2+}]_i$ in aPSC. PAR are activated as proteases attack a cleavage site toward the N-termini of the receptor, revealing a previously cryptic tethered peptide ligand (Soh et al., 2010). Synthetic peptides consisting of six amino acids upstream of the cleavage site can activate the receptor independently of protease cleavage. The PAR family is encoded by four genes, and although the selectivity is not exclusive, thrombin preferentially cleaves PAR-1 and -3 whereas trypsin has been shown to act on PAR-2 and -4. Activated PAR can couple to multiple heterotrimeric G protein $\alpha$-subunits including $G_{\alpha_{11}}, G_{\alpha12/13}$, and $G_{\text{reg/11}}$. Thus direct activation of PLC by $G_{\text{reg}}$ or $G_{\beta\gamma}$ subunits are linked to $\text{Ca}^{2+}$ signaling events. Proteases may activate same fields could be clearly shown to increase $[\text{Ca}^{2+}]_i$, in response to CCK (Figure 4C, red trace), trypsin (Figure 4F, red trace), and CCh (unpublished data). Together, these data demonstrate the utility of measuring $\text{Ca}^{2+}$ signals in situ in pancreatic lobules and importantly show that periacinar-localized cells from WT animals have $\text{Ca}^{2+}$ signaling characteristics consistent with those of qPSC in culture.

Characterization of PAR agonist-induced $[\text{Ca}^{2+}]_i$ signals in aPSC

Because of the probable pathophysiological significance of protease-activated signaling in pancreas (Namkung et al., 2004; Masamune et al., 2005; Laukkanen et al., 2008), experiments were performed to define in detail the characteristics of PAR-induced $[\text{Ca}^{2+}]_i$ in aPSC. PAR are activated as proteases attack a cleavage site toward the N-termini of the receptor, revealing a previously cryptic tethered peptide ligand (Soh et al., 2010). Synthetic peptides consisting of six amino acids upstream of the cleavage site can activate the receptor independently of protease cleavage. The PAR family is encoded by four genes, and although the selectivity is not exclusive, thrombin preferentially cleaves PAR-1 and -3 whereas trypsin has been shown to act on PAR-2 and -4. Activated PAR can couple to multiple heterotrimeric G protein $\alpha$-subunits including $G_{\alpha_{11}}, G_{\alpha12/13}$, and $G_{\text{reg/11}}$. Thus direct activation of PLC by $G_{\text{reg}}$ or $G_{\beta\gamma}$ subunits are linked to $\text{Ca}^{2+}$ signaling events. Proteases may activate

FIGURE 3: Characterization of pancreatic lobules. (A, top) Transmitted laser light images of a pancreatic lobule at magnification of 25x (left) and 50x (middle). The left panel shows intrinsic fluorescence in periacinar cells with MP excitation at 810 nm. (A, bottom) Corresponding MP-excitation images of calcein fluorescence. Arrows indicate nonacinar cells. (B) Immunocytochemical staining of fixed pancreatic lobules. Prominent duct cell staining of $\alpha$-SMA (Ba), periacinar staining for vimentin (Bb), and acinar staining with $\alpha$-amylase (Bc); overlay of Bb/Bc is shown in Bd.
During the previous series of experiments, it was noted that PAR ligands, ATP, and PDGF-stimulated Ca$^2+$ elevations were not spatially homogeneous. The spatial characteristics of the Ca$^2+$ signal were therefore next studied in Fluo-4–loaded aPSC. Following exposure to PAR-1 ligand (Figure 6, A and B), PDGF (Figure 6, C and D), or ATP (Figure 6, E and F), the Ca$^2+$ signal was initiated close to the nucleus and propagated freely into the nucleus and surrounding cytoplasm. The fluorescence signal was invariably greater within the nucleus when compared with the cytoplasm and margins of the cell (Figure 6). Ca$^2+$-sensitive dyes have been reported to behave anomalously in the nuclear milieu and often overestimate the magnitude of signals (Bootman et al., 2009). Nevertheless, these data show that multiple PAR subtypes, and therefore synthetic peptides modeled on the sequence of the ligand were next used to define the PAR subtypes expressed in aPSC. A marked elevation in [Ca$^{2+}$]$_i$ was evoked by the PAR-1 receptor peptide ligand (Figure 5, Aa and B) with an apparent $K_d$ of ~300 nM. Increasing PAR-1 peptide resulted in a recruitment of responsive aPSC; above 300 nM peptide > 80% of cells responded (Figure 5C), and a corresponding decrease in latency between ligand exposure and the elevation in [Ca$^{2+}$]$_i$ was observed (Figure 5D). The PAR-2 ligand evoked a smaller increase in [Ca$^{2+}$]$_i$ and was less potent (Figure 5Ab). In contrast, ligands directed against PAR-3 and PAR-4 were without effect (Figure 5, Ac and Ad, respectively).
Nuclear Ca\textsuperscript{2+} signals stimulate proliferation of aPSC

To investigate further the mechanism underlying the compartmentalized Ca\textsuperscript{2+} signals and to probe their physiological role, we utilized a series of adenoviruses expressing Ca\textsuperscript{2+} buffers targeted to specific cellular compartments (Rodrigues et al., 2007). The constructs consist of parvalbumin (PV) linked to the red fluorescent protein DsRed, and either a nuclear localization sequence (PV-NLS-DsRed) or nuclear exclusion signal (PV-NES-DsRed). Figure 8A shows a cell expressing a nontargeted PV-DsRed construct following virus infection and demonstrates the heterogeneous expression of the chimeric protein in aPSC. In contrast, PV-NES-DsRed was expressed in the cytoplasm and excluded totally from the nucleus (Figure 8Ab) while PV-NLS-DsRed was exclusively expressed only in the nucleus of aPSC (Figure 8Ac). Essentially 100% of cells were infected with each adenovirus, as evidenced by the presence of uniform red fluorescence. The impact on the Ca\textsuperscript{2+} signal of compartmentalized PV expression was assessed in Fluo-4–loaded aPSC stimulated with PAR-1 ligand. Expression of nontargeted PV resulted in a marked dampening of the agonist-evoked Ca\textsuperscript{2+} signals throughout the cell when Ca\textsuperscript{2+} appears to be readily released or diffuses from local release sites to maintain an elevated [Ca\textsuperscript{2+}] in the nucleoplasm of aPSC.

The spatial organization of the Ca\textsuperscript{2+} signals was further investigated using focal production of InsP\textsubscript{3} in distinct, spatially separated regions of aPSC (–1.5 μm\textsuperscript{3} focal volume) using laser photolysis of a cell-permeable form of caged InsP\textsubscript{3} (ci-InsP\textsubscript{3}) (Won et al., 2007). When the laser light was focused in the nucleus, low levels of photolysis resulted in elevated fluorescence signals that were largely confined to this compartment (Figure 7A, small arrow). Increasing energy produced larger signals in the cytoplasmic regions (Figure 7, A, large arrow; B, images). Positioning the laser spot in the margins of the cell, 20 μm distant from the nucleus resulted in an initial increase in Ca\textsuperscript{2+} signal at the photolysis site followed by a robust, maintained signal in the nucleus (Figure 7, C and D). Together, these data again support the idea that in aPSC, Ca\textsuperscript{2+} signals are freely communicated into the nuclear compartment and are consistent with InsP\textsubscript{3}-induced Ca\textsuperscript{2+} release occurring either in, or directly surrounding, the nucleus.

FIGURE 5: Characterization of PAR Ca\textsuperscript{2+} signaling in aPSC in culture. (A) Ca\textsuperscript{2+} elevations could be evoked in aPSC following exposure to peptide PAR-1 peptide ligand (SFLLRN-NH\textsubscript{2}) (Aa) and PAR-2 peptide ligand (SLIGRL-NH\textsubscript{2}) (Ab) but not PAR-3 (SFNNGGP-NH\textsubscript{2}) (Ac) or PAR-4 ligand (AYPGKF-NH\textsubscript{2}) (Ad). (B) Concentration vs. response relationship for the peak Ca\textsuperscript{2+} response evoked by PAR-1 ligand. (C) Increasing PAR-1 ligand results in progressive recruitment of cells to increase Ca\textsuperscript{2+} (increase defined as > 0.02 ratio units change from basal). (D) Mean data showing the change in latency and peak height of the Ca\textsuperscript{2+} signal initiated by increasing [PAR-1] ligand.
compared with non-PV construct–expressing cells (Figure 8Ba). In cells expressing PV-NES-DsRed, the Ca^{2+} signal in the extranuclear region was largely abolished (Figure 8Bb). However, a robust nuclear Ca^{2+} signal was still observed. Conversely, expression of PV-NLS-DsRed resulted in a markedly dampened nuclear Ca^{2+} signal, although the cytoplasmic signal was largely preserved (Figure 8Bc), indicating, not unexpectedly, that Ca^{2+} release can occur at some distance away from the nucleus.

aPSC are highly proliferative, and expansion of cell numbers followed by secretion of ECM is thought to play a major role in disease progression in chronic pancreatitis. Experiments were subsequently performed using targeted buffers to evaluate whether the prominent compartmentalized increases in [Ca^{2+}] play any role in promoting cell proliferation. aPSC were serum deprived for 16 h before addition of media replete with 1% fetal bovine serum (FBS). Proliferation was assessed by monitoring incorporation of 5-bromo-2′-deoxyuridine (BrdU) 24 h later. Addition of FBS resulted in a significant increase in BrdU incorporation. Basal proliferation in the absence of FBS was unaffected regardless of the expression and localization of targeted PV, while incorporation of BrdU was significantly increased in cells not expressing PV constructs exposed to FBS (Figure 9). Global uniform expression of PV (PV-DsRed) or buffering of cytoplasmic Ca^{2+} specifically with PV-NES-DsRed did not alter the proliferation of aPSC. In contrast, expression of PV-NLS-DsRed and thus buffering of nuclear Ca^{2+} completely eliminated proliferation (Figure 9). These data highlight the general importance of nuclear Ca^{2+} changes for proliferation and strongly suggest that agents that increase nuclear [Ca^{2+}], including PAR agonists, PDGF, and ATP, likely present in or around the gland following insult and play an important role in aPSC proliferation.

**FIGURE 6:** Spatial characteristics of Ca^{2+} changes. Exposure to PAR agonist evokes prominent, sustained fluorescence changes in the nucleus of aPSC (A, black trace) vs. a cytoplasmic region of interest (A, red trace). (B) Corresponding images acquired at time points corresponding to the arrows in (A). (C, images; D, kinetics) PDGF stimulation results in similar prominent nuclear Ca^{2+} signals. (E, images; F, kinetics) ATP stimulation results in prominent nuclear Ca^{2+} signals.

**PSC in an animal model of chronic experimental pancreatitis**

LSL-K-Ras^{G12D} mice express an activating mutation of the oncogene K-Ras driven by its endogenous promoter in acinar cells and thus exhibit moderately elevated Ras activity in acinar cells (Ji et al., 2002; Guerra et al., 2007). The animals develop extensive pancreatic fibrosis only following induction of experimental pancreatitis with serial intraperitoneal (IP) cerulein injections. These mice therefore provide an excellent model system to study Ca^{2+} signaling in PSC during the development of fibrosis. Studies were performed in lobules prepared from LSL-K-Ras^{G12D} mice under control conditions and subsequently following cerulein injection as detailed in Materials...
Figure 3). Periacinar cells responded to angiotensin II (Figure 10E), and a proportion of these cells also increased [Ca$^{2+}$] when exposed to thrombin (Figure 10B; pooled data in Figure 12G) or trypsin (Figure 10C; pooled data in Figure 12H) and low concentrations of ATP (Figure 10D; pooled data in Figure 12I), suggesting that some cells exhibit the phenotype of aPSC. Consistent with this idea, in a limited number of experiments cells also responded to PDGF. Furthermore, when stellate cells were isolated from these animals, α-SMA staining could be observed in 21% of cells on day 2 in culture (Figure 10F, 24/114 cells; 17 fields prepared from two mice), which was much earlier than cells prepared from WT animals (Figure 2), indicating the presence of aPSC in the pancreata of LSL-K-Ras G12D mice (Figure 10F). 

Pancreatic lobules from WT and LSL-K-Ras G12D animals were isolated 3 d after induction of experimental pancreatitis as detailed in Materials and Methods following IP injection of the CCK analogue cerulein. The structure of lobules from WT animals injected with cerulein was again not markedly different from noninjected WT animals, with sparse localization of α-SMA confined to ductal structures (Supplemental Figure 3). Patent acinar structure and occasional periacinar cells were clearly visualized by MP imaging of calcein fluorescence (Figure 11A, top). In ∼50% of these lobules, cells responded to thrombin (9/20 lobules examined) and trypsin (7/13) (Figure 12, A and B; pooled data in Figure 12, G and H), indicative of the presence of aPSC following treatment. This number was similar to noninjected LSL-K-Ras G12D lobules. Similar morphology and lack of proliferation of nonacinar cells were also seen in animals in which the injection protocol was repeated and the animals killed at day 28 (Figure 11C). In contrast, identical treatment of LSL-K-Ras G12D resulted in a severe disruption of exocrine pancreatic structure, a striking increase in the expression of α-SMA (Supplemental Figure 3), and a marked increase in nonacinar cells, presumably PSC (Figure 11, A, bottom, and C). Specifically, in numerous foci there was a loss of acinar cell structure, with the polarized cells replaced by cells with cuboidal morphology (Figure 11A, bottom). Foci were surrounded by numerous elongated cells that preferentially loaded with calcein AM. Paraformaldehyde-fixed lobules from these animals showed extensive expression of α-SMA in rings of cells surrounding cells expressing amylase (Figure 11B). The cells surrounding the remnants of acinar cells are therefore likely aPSC. Consistent with this hypothesis, in ∼95% of these lobules, cells were present that increased [Ca$^{2+}$], in response to trypsin, thrombin, and low concentrations of ATP (Figure 12, D–F; pooled data in Figure 12, G–I). This loss
of acinar structure and increase in PSC numbers, as monitored by calcein fluorescence in cells surrounding the foci, were even more marked after 28 d (Figure 11C). However, lobules prepared from these animals loaded very poorly with Fluo-4, precluding extensive investigation of Ca\textsuperscript{2+} signaling events.

DISCUSSION

The central role of Ca\textsuperscript{2+} signaling in pancreatic acinar cells in both health and disease is firmly established (Sutton et al., 2003; Kiselyov et al., 2006; Williams and Yule, 2006; Petersen and Tepikin, 2008). There are, however, relatively few reports characterizing the dynamics and impact of these potentially important signaling events in other exocrine pancreatic cell types. In this study, we have characterized Ca\textsuperscript{2+} signaling events in PSC both in culture and in situ, in acutely isolated lobules. The lobule preparation retains gland architecture that is largely comparable to the native pancreas and therefore provides an excellent platform to study signaling events in the various cell types resident in the exocrine pancreas. Significantly, we demonstrate that the phenotypic changes observed in PSC during activation in culture are mirrored by changes that occur in PSC in situ in WT animals and in an animal model of chronic pancreatitis. In both qPSC and aPSC, Ca\textsuperscript{2+} signaling is initiated via the canonical pathway.
which results in InsP$_3$R-mediated Ca$^{2+}$ release from intracellular stores. Notably, however, the complement of PSC cell-surface receptors coupled to Ca$^{2+}$ signaling events was dramatically altered during activation, and these changes are paralleled whether in culture or in situ. Although the "route" of activation may be different in the case of the animal model versus cultures of acutely isolated PSC, the observation that aPSC in each situation share fundamental similarities provides important validation for studies of cultured PSC.

Ca$^{2+}$ signaling events in cultured aPSC

Agonist-stimulated Ca$^{2+}$ changes are ubiquitous signals controlling a diverse array of downstream effectors that ultimately direct various cellular endpoints as disparate as proliferation and apoptotic cell death (Berridge et al., 2000). It is clear that cells must use mechanisms to ensure fidelity and specificity in order to guarantee the appropriate activation of effectors. This is widely believed to occur as a consequence of the specific temporal and spatial characteristics of the Ca$^{2+}$ change. Stimulation of aPSC in culture resulted in Ca$^{2+}$ signals with distinct spatial features. The signals appear to initiate close to the nuclear envelope, subsequently rapidly invade, and are maintained in the nucleoplasm relative to the cytoplasm. Ca$^{2+}$ signals with similar spatial characteristics were also initiated following focal liberation of InsP$_3$ from a caged precursor some 20 µm distant from the nucleus, suggesting that the generation of nucleoplasmic signals is favored in aPSC. In addition, uncaging directly in the nucleus produced signals that were maintained in this compartment longer than the corresponding cytoplasmic signals. A question arises as to the mechanism that underlies the striking spatial attributes of the signal. A simple hypothesis is that InsP$_3$ is produced at the plasma membrane, diffuses through the cytoplasm, and engages InsP$_3$R on perinuclear ER to initiate Ca$^{2+}$ release (Allbritton et al., 1994; Lipp et al., 1997; Eder and Bading, 2007; Bootman et al., 2009). The nucleoplasmic Ca$^{2+}$ rise then passively follows the Ca$^{2+}$ rise from the ER by diffusion through the nuclear pore complex. The signal remains elevated because of the relatively low buffering capacity and paucity of Ca$^{2+}$ ATPase activity within the nucleus (Fox et al., 1997).

Studies using targeted PV indicate, however, that simple diffusion between compartments may not be the complete story. Nuclear but not cytoplasmic Ca$^{2+}$ elevations were markedly attenuated in cells expressing nuclear-targeted PV, indicating, not unexpectedly, that Ca$^{2+}$ signals can be initiated away from the nucleus and are consistent with a contribution of passive diffusion into the nucleus. To the contrary, nuclear Ca$^{2+}$ signals were readily observed in the presence of PV targeted to the cytoplasm where Ca$^{2+}$ was shown to be effectively buffered. This observation is consistent with the notion that the Ca$^{2+}$ signal observed in the nucleoplasm can result from Ca$^{2+}$ release through InsP$_3$R localized in the close vicinity of the nucleus. Potential release sites with precedent established in other cell types include InsP$_3$R localized to the outer nuclear envelope, localized to the inner nuclear membrane, or alternatively directly from ER invaginations projecting within the nucleus (Hennager et al., 1995; Echevarria et al., 2003; Cardenas et al., 2010).

Do these nuclear Ca$^{2+}$ changes play any specific role in cultured aPSC? Nucleoplasmic Ca$^{2+}$ changes have been shown to alter the activity of numerous downstream targets including transcription factors, kinases, and phosphatases (Dolmetsch et al., 2001; Pusl et al., 2002; Mellstrom et al., 2008). An elegant demonstration of the profound impact of nuclear Ca$^{2+}$ on cellular function has been reported in a series of studies using targeted PV constructs developed by Nathanson, Leite, and colleagues (Rodrigues et al., 2007). A major finding of these studies was that PV targeted to the nucleus markedly reduced cell proliferation in hepatocytes and the size of hepatic tumors (Rodrigues et al., 2007). The predominant effect of nuclear PV was to block progression through the cell cycle by controlling mitosis. Significantly, cytosolic-targeted PV did not alter cell proliferation but attenuated the activity of cytosolic effectors. The current data indicate that nuclear Ca$^{2+}$ plays an analogous role in aPSC because in a similar fashion simple global attenuation of the Ca$^{2+}$ signal, or specific cytosolic reduction, did not alter proliferation while nuclear-targeted PV selectively abrogated aPSC proliferation. That the specific expression of PV in the nucleus reduced proliferation while a general reduction of [Ca$^{2+}$] with the nontargeted PV did not, may be related to the fact the NLS construct, as a function of specific targeting and presumably increased relative "residence" in the nucleus, is better able to tightly control the local Ca$^{2+}$ environment in this compartment. Interestingly, these data are in contrast to a report on hepatic stellate cells that indicated that while Ca$^{2+}$ changes were important for proliferation, the specific compartment was not (Soliman et al., 2009). In total, the experiments point to an important role of compartmentalized nuclear Ca$^{2+}$ signals in the proliferation of aPSC. This may contribute to the mechanism’s underpinning expansion of aPSC cell numbers in disease states of the pancreas.

Phenotypic changes in cultured PSC on activation: implications for pancreatic disease

A major change during PSC transformation to an activated phenotype is the expression of PAR-1 and PAR-2 receptors (Masamune et al., 2005) and increased sensitivity to ATP. The demonstration of Ca$^{2+}$ signaling events generated following thrombin and trypsin exposure exclusively in aPSC is consistent with these proteins initiating Ca$^{2+}$ signaling events that contribute to pancreatic disease. How might aPSC be exposed to agents that subsequently activate Ca$^{2+}$ signaling events? In experimental models of acute pancreatitis, there is considerable evidence that trypsinogen, normally inactive until cleaved by enteropeptidases in the small intestine, is converted to active trypsin within the acinar cell (Saluja et al., 1999; Kruger et al., 2000; Ratary et al., 2000). Trypsin, as well as other cytoplasmic contents including adenine nucleotides, is likely released into the extracellular space following acinar cell damage. In addition, during episodes of pancreatitis there is increased permeability of the vasculature, exposing the pancreas to the contents of plasma (Turjanski et al., 2007). It is therefore probable that during episodes of pancreatic insult the gland is exposed to PAR ligands, PDGF, and adenine

**FIGURE 9:** aPSC proliferation is attenuated by nuclear-targeted PV. Proliferation induced by serum was attenuated only by PV-NLS-DsRed.
damage to acinar cells and induction of the inflammatory response (Omary et al., 2007). Of note, however, despite evidence of “activation,” is that no apparent increase in stellate cell numbers was observed in WT animals following cerulein treatment. Presumably, in this experimental paradigm the unchanged aPSC numbers reflect the need for an additional signal to either enhance the basal proliferation rate or overcome efficient removal of aPSC from the pancreas. This is consistent with little evidence of pancreatic fibrosis or severe morphological disruption in this model of pancreatitis (Lampel and Kern, 1977).

In contrast, in lobules isolated from LSL-K-RasG12D mice subjected to the same protocol, an approximately twofold increase in the numbers of stellate cells was observed. A majority of these cells exhibited an activated phenotype as assessed by response to PAR agonists and low [ATP], and this was accompanied by extensive expression of α-SMA and secretion of ECM, as reported previously. Interestingly, although LSL-K-RasG12D animals have been reported to have no overt phenotype until pancreatitis is induced (Guerra et al., 2007), our studies provide clear evidence that PSC are primed in an activated state before insult. The primary evidence for this is that a proportion of the PSC responds to PAR agonist, and aPSC can be prepared from these animals that do not require culture for transformation. By virtue of the targeting strategy used to generate the LSL-K-RasG12D mouse (Guerra et al., 2007), activation of PSC must occur as a result of the diverse signaling pathways downstream of K-Ras activation in acinar cells (Schubbert et al., 2007; Kholodenko et al., 2010). This “priming” of the PSC in an activated state likely promotes the proliferation of aPSC following induction of pancreatitis in this animal, resulting in a more severe manifestation of disease. It is tempting to speculate that nuclear Ca²⁺ signals initiated by agents including trypsin, thrombin, and ATP may contribute to this phenomenon. Consistent with this idea, camostat, an orally active serine protease inhibitor, has been reported to decrease pancreatic fibrosis and PSC proliferation in animal models of chronic pancreatitis (Gibo et al., 2004; Emori et al., 2005).

A clinical parallel may be relevant in regard to the data generated in the LSL-K-RasG12D mice. Patients with chronic pancreatitis have an increased risk of developing pancreatic ductal adenocarcinomas (PDAC) (Lowenfels et al., 1988; Hruban et al., 1993; Luttges et al., 2000). Additional strong evidence to support this proposal has come from several genetic mouse models.
tent with the relatively low, but nevertheless increased, level of K-Ras activity in LSL-K-Ras\textsuperscript{G12D} animals effectively functioning as a "coincidence detector" that increases the likelihood of pancreatic disease; the increased K-Ras activity leads to primed aPSC that can respond to further insult, including those that lead to Ca\textsuperscript{2+} signaling events and aPSC proliferation. It is envisioned that further studies using a similar combination of in situ imaging and cultured PSC from these mouse models will provide additional insight into the role of PSC in pancreatic disease.

MATERIALS AND METHODS

Fluo-4 AM and Fura-2 AM were purchased from Teflabs (Austin, TX). Calcein AM was from Invitrogen (Carlsbad, CA). Ci-InsP\textsubscript{3} was from Axxora (San Diego, CA). \(\alpha\)-SMA antiserum and \(\alpha\)-amylose were purchased from Sigma (St. Louis, MO); \(\alpha\)-vimentin, \(\alpha\)-GFAP, and \(\alpha\)-desmin antiserum were from Abcam (Cambridge, MA). Alexa Fluor 488/546–coupled secondary anti-bodies were from Invitrogen. PAR ligand peptides were from AnaSpec (San Jose, CA). Adenovirus encoding targeted parvalbumin constructs were a kind gift from Michael Nathanson (Yale University, New Haven, CT) and Fatima Leite (Federal University of Minas Gerais, Brazil).

Isolation and culture of PSCs

PSC were isolated by a modification of the method described by Apte et al. (1998). Briefly, pancreatic tissue from mouse (approximately 25 g) was minced with scissors and digested with 0.02% Pronase (Roche, Indianapolis, IN), 0.05% Collagenase P (Roche), and 0.1% DNAse in Gey’s balanced salt solution (GBSS; Sigma Aldrich, St. Louis, MO) for 50 min. Digested tissue was then filtered through a 100-µm nylon mesh. Cells were washed and then resuspended in GBSS containing 0.3% bovine serum albumin (BSA). The cell suspension was centrifuged into a 28.7% (wt/vol) solution of Nycodenz (Sigma Aldrich) at 1400 \(\times\) g for 20 min. The cells of interest separated into a fuzzy band just above the interface of the Nycodenz cushion and the GBSS with BSA. This band was harvested, and the cells were washed and resuspended in a mix of DMEM and F-12 medium (1:1 vol/vol) containing 10% FBS, 4 mM glutamine, and antibiotics (penicillin 100 U/ml and streptomycin 100 \(\mu\)g/ml). Cells were seeded at a density of 4 \(\times\) 10\textsuperscript{4} cells/cm\textsuperscript{2}. Viability of isolated cells was assessed by trypan blue exclusion, and cells were counted using a hemocytometer. Cells were maintained at 37°C in a humidified atmosphere of 5% CO\textsubscript{2}/95% air. The culture medium was replaced the day after initial seeding and subsequently each 3 d. After reaching 80% confluence, the cells were released expressing a range of K-Ras activity, including the mice used in this study. For example, although LSL-K-Ras\textsuperscript{G12D} mice develop pancreatic fibrosis and precancerous lesions only after further pancreatic insult, a mouse engineered to express higher levels of K-Ras in acinar cells exhibits extensive pancreatic fibrosis and a high penetrance of PDAC without further insult (Ji et al., 2002). Our data are consist
from the plastic by trypsinization and subcultured into 12-well plates at a density of $1 \times 10^4$ cells/well. All experiments were performed between passages 1 and 5.

**Induction of pancreatic fibrosis**

As previously described, mice were injected IP five times at 2-h intervals with 50 µg/kg cerulein on day 1 of the protocol followed by a single injection on days 2 through 5 (Ju et al., 2009). The mice were killed on day 8 for preparation of stellate cells and lobules. In a number of experiments the protocol detailed for the first week was repeated on days 21–25 and the animals were killed on day 28. Animal procedures were approved by the University of Rochester Institutional Animal Care and Use Committee.

**Immunocytochemistry and staining of lipid droplets**

Immunocytochemistry in pancreatic slices was performed as previously described by Warner et al. (2008) for parotid gland slices. Briefly, 200-µm-thick pancreas slices were fixed with 4% paraformaldehyde. Following permeabilization and blocking, the slices were incubated overnight with the indicated primary antisera in phosphate-buffered saline–Triton (PBS-Tx) at 4°C. The slices were rinsed with PBS and incubated with 2 µl AdipoRed per ml PBS for 10 min. Fluorescence was observed on a Nikon C1 confocal microscope following excitation with 563 nm HeNe laser with emission monitored above 600 nm.

**Cell proliferation assay**

Cell proliferation was assessed via the incorporation of BrdU according to the manufacturer’s instruction (Cell Proliferation ELISA, Roche) using a chemiluminescence-based assay detected by a plate reader equipped with a luminescence detector (Synergy 2; BioTek, Winooski, VT). Briefly, PSCs (passages 2 and 3) were seeded at a density of $3 \times 10^4$ cells/well in 96-well culture plates. After 1 d in culture, cells were infected with PV-NLS-DsRed, PV-NES-DsRed, or PV-DsRed virus at 37°C for 16 h. The cells were washed with serum-free culture medium and incubated with serum-free or 1% FBS culture.

**Ca$^{2+}$ imaging and focal uncaging in PSC**

Imaging was performed as previously described (Won and Yule, 2006; Won et al., 2007). Briefly, stellate cells grown on cover glass were loaded with 2 µM fura-2 AM at room temperature for 30 min. Imaging was performed on a Nikon TE200 Inverted microscope using a 40x oil immersion objective lens coupled to a monochromator-based illumination system (TILL Photonics, Grabeling, Germany) and digital frame transfer CCD camera controlled by Vision Suite software (Betzenhauser et al., 2008). Changes in Fura-2 fluorescence are presented as a change in ratio of the emitted fluorescence following excitation at 340/380 nm. For focal uncaging experiments, PSC were loaded with 4 µM Fluo-4. Focal uncaging was accomplished as previously described using a custom-built system consisting of a laser light source (90 mW, 405-nm diode) coupled through a single-mode fiber to an epifluorescence condenser (Won et al., 2007). The system produces an intense, focused, and restricted area (∼1.5 µm$^3$) of illumination at the focal plane. Brief exposure (<50 ns) provided sufficient energy to photolyse ci-InsP$_3$ without any obvious deleterious effects on the cells. Changes in Fluo-4 fluorescence are expressed as ΔF/F$_{00}$, where F is the fluorescence captured at a particular time and F$_{00}$ is the mean of the initial 10 fluorescence images captured.
MP imaging in pancreatic lobules and slices

Pancreatic lobules were isolated following injection of saline beneath the capsule with a 27-gauge needle. Using a binocular dissecting scope, individual lobules < 1 mm³ were removed with scissors. Some lobules were embedded in 3% low-temperature melting point agarose and equilibrated with 95% O₂ and 5% CO₂ for 1–4 h. The lobules and slices were loaded with Fluo-4 AM for intracellular Ca²⁺ measurements (5 µM) or calcein AM for visualization of structure (5 µM) by incubation for 30–60 min. Intrinsically fluorescent GECOS was excited at the indicated wavelength and visualized in lobules not loaded with dye. Pancreatic tissue was placed in a low-volume perfusion chamber (Warner Instruments, Hamden, CT) and maintained in position by using a wire grid in a weighted metal ring. MP imaging was performed using an Olympus BX61WI upright microscope coupled to an Olympus Fluoview 1000 multiphoton/confocal (FV1000MP) microscope equipped with Spectra Physics Mai Tai (DeepSee, Santa Clara, CA) mode-locked Ti:sapphire tunable laser. Imaging was performed using a 25× water-immersion objective (XL Plan N; numeric aperture = 1.05). The laser was tuned to provide excitation at 810 nm, and emitted light was separated using a 565-nm dichroic mirror followed by emission filters HG525/50 and HQ605/50 in front of the detectors to visualize green and red emitted fluorescence, respectively. An individual plane, typically at a depth of 80–150 µm into the slice, was scanned at 1 Hz for measurement of intracellular Ca²⁺. A Z-series through the preparation followed by analysis of a reconstructed maximum projection image was used to analyze lobule structure.

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