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Imaging endoplasmic reticulum calcium with a fluorescent biosensor in transgenic mice

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The advent of genetically encoded biosynthetic fluorescent sensors has been a major advance in the study of intracellular calcium. The use of biosynthetic fluorescent sensors has been an important new approach for measuring intracellular Ca$^{2+}$ in cells. Genetically encoded indicators based on green fluorescent protein, calmodulin, and fluorescence resonance energy transfer (FRET) have been utilized to measure Ca$^{2+}$ in nonmammalian transgenic organisms and provide information about the organization and regulation of Ca$^{2+}$ signaling events in vivo. However, expression of biosynthetic FRET-based Ca$^{2+}$ indicators in transgenic mammals has proven to be problematic. Here, we report transgenic expression of an endoplasmic reticulum (ER) Ca$^{2+}$ biosensor in mouse pancreas. We targeted expression of a yellow cameleon3.3er (YC3.3er) transgene with mouse insulin I promoter. YC3.3er protein expression was limited to pancreatic β-cells within islets of Langerhans and absent in the exocrine pancreas and other tissues. Animals developed and matured normally; sensor expression was unaffected by age. Glucose tolerance in transgenic mice was also unaffected, indicating the transgenic biosensor did not impair endocrine pancreas function. ER Ca$^{2+}$ responses after administration of thapsigargin, carbachol, and glucose were measured in individual β-cells of intact islets using confocal microscopy and confirmed the function of the biosensor. We conclude that controlling transgene transcription with a cell-specific promoter permits transgenic expression of FRET-based Ca$^{2+}$ sensors in mammals and that this approach will facilitate real-time optical imaging of signal transduction events in living tissues.

intracellular calcium; cameleon; fluorescence resonance energy transfer

QUANTITATIVE IMAGING OF SPATIAL and temporal gradients of intracellular Ca$^{2+}$ concentration is essential to furthering our knowledge of Ca$^{2+}$-mediated signal transduction and the wide range of Ca$^{2+}$-dependent cellular functions (2, 3, 17). The advent of genetically targeted biosynthetic sensor technology has significantly increased our ability to study complex signaling events in cells with unprecedented temporal and spatial resolution. Many genetically encoded Ca$^{2+}$ indicators based on Ca$^{2+}$-dependent changes in fluorescence resonance energy transfer (FRET) or sensor conformation are now available that provide quantitative visualization of ion fluxes in specific subcompartments of cells (1, 6, 13, 17). Although this technology has the potential for advancing our understanding of signaling in mammals in vivo, applications of fluorescent Ca$^{2+}$ biosensors have been confined to evaluation of mammalian cells and cell lines in vitro.

Expression of these probes in intact mammalian tissues has proven difficult. One of the major impediments has been the lack of an effective method to deliver biosensor cDNA to cells within a living mammal; this cannot be accomplished effectively and reproducibly with liposomal, electroporation, viral, or ballistic methods but conceivably could be attained using a transgenic approach. Biosynthetic fluorescent cameleon and camagaro Ca$^{2+}$ sensors based on green fluorescent protein (GFP) and the Ca$^{2+}$ binding protein, calmodulin (13, 17), have been engineered to label specific populations of cells within intact tissues in nonmammalian transgenic organisms (Caenorhabditis elegans, Drosophila melanogaster, and Danio rerio) (5, 9–11, 16, 20, 21). No mammalian transgenic expression of a FRET-based Ca$^{2+}$ biosensor, however, has been reported. In this paper we demonstrate transgenic expression of yellow cameleon3.3er (YC3.3er), a biosynthetic indicator of endoplasmic reticulum Ca$^{2+}$ (6), in endocrine cells of mouse pancreas. Our studies demonstrate the feasibility of applying FRET biosensor imaging technology to transgenic mammalian cell systems and suggest that this approach may be used for real-time visualization of cell signaling in living tissues.

MATERIALS AND METHODS

Transgene construction. The mouse insulin I gene promoter (MIP)-YC3.3-er transgenic construct was assembled using an 8.3-kb fragment of MIP, the 2-kb coding region of YC3.3er, and a 2-kb fragment of the human growth hormone (hGH) cassette gene for high-level expression (6, 7, 14, 15). The 12.3-kb MIP-YC3.3-er-hGH fragment was isolated from the vector by digestion of the plasmid construct with HindIII and SfiI and agarose gel electrophoresis. The fragment was further purified using an Elutip-D column (Schleicher & Schuell, Keene, NH).

MIP-YC3.3-er transgenic mice. The purified transgene DNA was microinjected into the pronuclei of CD-1 mice at the Transgenic Mouse/ES Core Facility of the University of Chicago Diabetes Research and Training Center. The transgene was maintained on the CD-1 background, and mice were housed under specific pathogen-free (SPF) conditions with free access to food and water. Tail DNA from potential founder mice was screened for the presence of the transgene by PCR using forward and reverse primers 5′-GACAACCACTACCTGAGC-3′ and 5′-ACTGGGCTTACATGGCGATACTC-3′, respectively. Whole pancreata of F1 progeny were visualized with an Olympus SZX12 stereomicroscope (Olympus, Melville, NY) in bright-field and fluorescence illumination modes. All the procedures involving mice were approved by the University of Chicago Institutional Animal Care and Use Committee.

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Glucose tolerance testing. Intraperitoneal glucose tolerance tests were performed after a 4-h fast. Blood was sampled from the tail vein before and 30, 60, 90, and 120 min after intraperitoneal injection of 2 mg/g body wt of dextrose. Glucose levels were measured using a Precision Q.I.D. Glucometer (MediSense, Waltham, MA).

Isolation of islets of Langerhans. Pancreatic islets were isolated as described (7). Briefly, the pancreas was inflated with a solution containing 0.3 mg/ml collagenase (Type XI; Sigma, St. Louis, MO) in Hanks’ balanced salt solution, injected via the pancreatic duct. The inflated pancreas was removed, incubated at 37°C for 10 min, and shaken vigorously to disrupt the tissue. After differential centrifugation through a Ficoll gradient to separate islets from acinar tissue, the islets were washed and placed on 25-mm glass coverslips. Islets were cultured in RPMI 1640 supplemented with 10% (vol/vol) fetal bovine serum, 100 U/ml penicillin, and 100 μg/ml streptomycin and incubated in a humidified incubator at 37°C in 95% air and 5% CO2. All imaging studies were performed 2–5 days after isolation.

Immunofluorescence microscopy. Sections (6 μm in thickness) from paraffin-embedded pancreatic tissue were incubated with monoclonal anti-GFP antibody (Sigma) to detect the expression of MIP-YC3.3-er, polyclonal anti-porcine insulin antibody to identify β-cells, and a combination of polyclonal anti-glucagon, anti-somatostatin, and anti-pancreatic polypeptide (PP) antibodies to identify α-, δ-, and PP cells, respectively (DakoCytomation, Carpentaria, CA). The endoplasmic reticulum (ER) was labeled with a polyclonal anti-glucose-regulated protein 94 (GRP94) antibody (Stressgen, Victoria, BC, Canada). Biotin-streptavidin-conjugated anti-mouse IgG and Cy2-conjugated streptavidin anti-mouse and Texas Red-conjugated anti-guinea pig/rabbit IgG secondary antibodies were used for MIP-YC3.3-er and insulin, respectively. Cy5-conjugated secondary antibody was used for non-β-cells and GRP94 (Jackson ImmunoResearch Laboratory, West Grove, PA). The stained sections were visualized with a Leica SP2 AOBS confocal microscope.

Confocal microscopy of isolated islets. Laser scanning confocal images were collected on a Leica SP2 AOBS spectral confocal microscope system using a 63× NA1.3 glycerol objective, viewing coverslip preparations held at 37°C on the DMIRE2 inverted scope. Excitation illumination was generated by a 405-nm laser, and simultaneous emission was detected at 453–505 nm and 525–600 nm for enhanced cyan fluorescent protein (ECFP; FRET donor) and citrine (FRET acceptor), respectively. Image stacks were reconstructed with Image J software (National Institutes of Health, Bethesda, MD).

ER Ca2+ measurements. We measured biosensor function using real-time spining disk optical confocal microscopy. Islets were placed into a microperfusion chamber mounted on an inverted epifluorescence microscope (TE-2000U, Nikon) equipped with a CARV spinning disk confocal system (Atto). Individual islets were visualized with a ×20 or ×40 fluorescent objective. Biosensor fluorescence excitation light was 440 nm and attenuated 50–90% using neutral density filters. Emitted fluorescence at 535 nm (citrine, FRET acceptor) and 485 nm (ECFP, FRET donor) was measured using a computer-controlled high-speed filter wheel (Lambda 10-2 Optical Filter Changer, Sutter Instruments, Novato, CA); the time for changing emission filters was 60 ms. Images (100- to 250-ms exposure) were captured with a 16-bit Cascade 650 digital camera (Roper Imaging) at 10-s intervals. Imaging data acquisition and analysis were accomplished using MetaMorph/MetaFluor software (Universal Imaging). Data were expressed as background subtracted intensities of the FRET acceptor and donor fluorophores and ratio of the FRET acceptor to FRET donor emission (Ratio 535/485). In addition, data were normalized to the average baseline value of Ratio 535/485 (Relative Ratio) to facilitate comparisons between responses of different cells. In all experiments, islets were superfused with warmed (37°C) buffered salt solutions consisting of (in mM) 119 NaCl, 4.7 KCl, 2.5 CaCl2, 1 MgCl2, 1 KH2PO4, 25 NaHCO3 or 10 HEPES-NaOH (pH 7.40), and 2–20 glucose.

Statistical analysis. Comparisons between groups were analyzed by ANOVA (StatView Software, SAS Institute, Cary, NC), and differences were considered to be significant at P < 0.05.

RESULTS

We generated transgenic mice expressing YC3.3-er specifically in pancreatic β-cells with a transgene construct similar to one used for transgenic expression of GFP (7). Specific cell targeting of the transgene was determined by the MIP; inclusion of a 2-kb fragment of the hGH cassette gene produced high-level expression (7, 14, 15). The MIP/hGH construct has been shown to generate transgenic mice with GFP expressed exclusively in pancreatic β-cells and not in brain, fat, skeletal or smooth muscle, gastrointestinal cells, kidney, liver, heart, spleen, or cells of the reproductive system (7). The 12.3-kb MIP-YC3.3. er transgene DNA (Fig. 1A) was microinjected into pronuclei of CD-1 mice. We obtained four male founders (MIP-YC3.3-er-5, -21, -27, and -40) that were genotypically positive for the transgene. After crossing the founders with wild-type CD-1 mice, only the F1 progeny from the MIP-YC3.3-er-21 founder demonstrated detectible fluorescence.

Using stereomicroscopy, we found protein expression of MIP-YC3.3-er in the pancreata of F1 neonates (as early as 1 day after birth) and adult mice. At the time of writing this report, our colony contains MIP-YC3.3-er-positive mice 17 wk of age, indicating that transgene expression is retained as the animals age. Stereoscopic bright-field and fluorescence imaging (Fig. 1, B and C) and laser scanning confocal microscopy (data not shown) demonstrated that expression of the fluorescent biosensor was restricted to islets of Langerhans. We did not observe MIP-YC3.3-er fluorescence in stomach, spleen, liver, or duodenum. The islet-specific pattern of MIP-YC3.3-er expression was indistinguishable from our previous transgenic mouse line, Tg(MIP-GFP)6729Hara, generated by using a similar MIP/hGH construct to drive expression of GFP specifically in insulin-secreting β-cells (7).

The MIP-YC3.3-er mice developed normally. Body weight and blood glucose levels of 6- and 8-wk-old transgenic animals were not significantly different from age-matched CD-1 control mice (P > 0.05; n = 6 mice from each group). The average ± SE body weight of 6-wk-old mice was 19.6 ± 2.0 g and 20.5 ± 1.8 g in the transgenic and wild-type mice, respectively. The average ± SE fasting blood glucose concentration in transgenic and wild-type mice was 158.0 ± 9.1 mg/dl and 173.0 ± 4.9 mg/dl, respectively. At 8 wk, the average ± SE fasting blood glucose in the transgenic mice was 166.6 ± 11.2 mg/dl. Intraperitoneal glucose tolerance tests performed in 6-wk-old (Fig. 1D) and 8-wk-old (data not shown) mice also demonstrated no significant differences (P > 0.05).

We used laser scanning confocal microscopy to illustrate the expression of the biosensor in paraffin-embedded pancreas sections (Fig. 2) and in single intact live islets of Langerhans (Fig. 3) from MIP-YC3.3-er mice. Immunofluorescence confocal microscopy revealed overlap between the ER Ca2+ indicator and insulin immunoreactivity (Fig. 2, A–C). There was no evidence of MIP-YC3.3-er expression in islet α-, δ-, and PP cells (Fig. 2, D–F). These results suggest that MIP-YC3.3-er is expressed only in the insulin-secreting β-cells. Double immunostaining showed that the subcellular distribution of MIP-YC3.3-er in β-cells (Fig. 3)
2G) colocalized with GRP94, a marker of ER (Fig. 2, H–I). This finding indicates that the transgenic biosensor is expressed within the lumen of the ER.

Fluorescence from individual cells within isolated live islets was easily detected (Fig. 3). The intensities of the fluorescent emission from the FRET donor and acceptor channels were very bright, ranging between 3- and 10-fold higher than the background autofluorescence observed in islets from control mice. YC3.3-er fluorescence was distributed throughout the cytoplasm of individual cells in a reticulated pattern (Fig. 3A) and analogous to the distribution of ER-targeted cameleon, YC4-er, in transiently transfected insulin-secreting cell lines (19). Intranuclear expression of the biosensor was evident in the H9252-cells, characterized by membrane-bound lamellar and branching tubular structures contiguous with the ER and nuclear envelope (Fig. 3, B–C). The morphology of these structures in primary β-cells was strikingly similar to a nucleoplasmic reticulum found in SKHep1 epithelial cells that is thought to regulate highly localized intranuclear Ca\(^{2+}\) signaling (4).

Similar to what we observed in paraffin-embedded sections of transgenic mouse pancreas (Fig. 2), not all cells within the isolated islets expressed YC3.3-er. Nonfluorescent cells appeared as dark voids or cavities within the islets (Fig. 3, B–D) and likely represent α-cells, δ-cells, and PP cells of the endocrine pancreas. This conclusion is consistent with immunohistochemical studies showing an absence of transgenic GFP expression in other endocrine cell types within islets from MIP-GFP mice (7).

Our confocal imaging studies demonstrated intercellular and intracellular heterogeneity of the FRET emission intensity ratio in unstimulated islets (Fig. 3D). In some cells, the perinuclear ER appeared to exhibit a higher FRET ratio than the ER in the cytoplasm (Fig. 3D, arrow). These observations raise the possibility that the FRET ratio heterogeneity might reflect regional variations in ER Ca\(^{2+}\) levels and...
suggest that β-cell ER Ca\(^{2+}\) is regulated by different mechanisms in distinct subcellular regions. More work will need to be done to explore these hypotheses.

We next determined whether the transgenic biosensor was functional. Cameleon indicators rely on Ca\(^{2+}\)-dependent FRET between chromophores of two GFP mutants as the basis for optical measurements of Ca\(^{2+}\) gradients (1, 6, 13, 17). The GFP mutants in YC3.3-er are ECFP and citrine, a mutant of enhanced yellow fluorescent protein (6). As Ca\(^{2+}\) concentration ([Ca\(^{2+}\)]) rises, FRET between ECFP and citrine increases, causing fluorescence emission from ECFP to decrease and citrine to increase. In contrast, a fall in [Ca\(^{2+}\)] reduces FRET, and consequently ECFP emission intensity increases and citrine decreases. The ratio of citrine (FRET acceptor) and ECFP (FRET donor) fluorescence emission is indicative of [Ca\(^{2+}\)]. We measured FRET between ECFP and citrine in individual β-cells within intact transgenic islets visualized (at 10-s intervals) with a real-time spinning disk confocal microscope. Approximately 50–60 s after application of thapsigargin, an inhibitor of sarco(endo)plasmic reticulum Ca\(^{2+}\)-ATPases that causes irreversible depletion of Ca\(^{2+}\) sequestered within the ER (18), biosensor FRET (Fig. 4A) and the ratio of FRET acceptor-to-donor emission decreased (Fig. 4B). This effect was observed in every cell (n = 20) we examined within transgenic islets expressing YC3.3-er (Fig. 4C) and indicated that MIP-YC3.3-er was providing a readout of [Ca\(^{2+}\)]\(_{\text{er}}\) ([Ca\(^{2+}\)]\(_{\text{er}}\)). This conclusion was further substantiated by exposing islets to carbachol, a muscarinic agonist that discharges ER Ca\(^{2+}\) stores by phospholipase C-mediated production of inositol 1,4,5-trisphosphate (IP\(_3\)) and activation of IP\(_3\) receptor Ca\(^{2+}\) channels located in the ER membrane. Administration of carbachol transiently lowered [Ca\(^{2+}\)]\(_{\text{er}}\), but not to the same extent as thapsigargin (Fig. 4D). The nadir of the carbachol-induced decrease in [Ca\(^{2+}\)]\(_{\text{er}}\) occurred within 1 min and then ER Ca\(^{2+}\) increased and, after 5–6 min, returned to baseline levels. This is consistent with ER store refilling and likely reflects activation of thapsigargin-sensitive sarco(endo)plasmic reticulum Ca\(^{2+}\)-ATPases. However, we were surprised to observe that after depletion of ER Ca\(^{2+}\) stores by prolonged exposure of islets to thapsigargin (5 µM for 8 min) in solutions containing 2 mM glucose, administration of 20 mM glucose increased the average FRET ratio by 79% (n =
6 cells). This unexpected finding suggests that primary mouse β-cells possess a glucose-stimulated, thapsigargin-insensitive mechanism capable of refilling ER Ca$^{2+}$ stores. The underlying mechanism remains to be determined.

**DISCUSSION**

Elucidating precisely how Ca$^{2+}$ and other signaling molecules are organized and integrated dynamically to influence mammalian cell function in tissues and organ systems will be accomplished by measuring signal transduction events in situ. Problems with loading cells in thick biological tissues, and the lack of cell and subcellular compartment specificity of synthetic polycarboxylate Ca$^{2+}$ dyes such as fura-2, preclude their use for ex vivo or in vivo imaging of Ca$^{2+}$ signaling within intact multicellular mammalian organ systems. Although transgenically expressed fluorescent Ca$^{2+}$ biosensors are potential candidates for quantitative real-time visualization of signaling in mammalian tissues, transgenic expression of GFP- and calmodulin-based cameleon and camgaroo indicators has been restricted to nonmammalian organisms (5, 9–11, 16, 20, 21).

The reasons for the difficulty in development of mammalian transgenic biosensor models are unclear. It has been proposed that a high level of expression is necessary for the detection of cameleon fluorescence and that biosensor overexpression interferes with endogenous calmodulin-dependent signal transduction (17). On the other hand, interactions between yellow cameleon biosensors and endogenous calmodulin or calmodulin-binding proteins are unlikely: the major effect of overexpression is to increase Ca$^{2+}$ buffering (12). An intracellular concentration of 20 μM is sufficient to detect cameleon fluorescence, and in vitro studies suggest that as much as 1 mM cameleon does not perturb calmodulin-dependent signaling (12). An alternate explanation for the failure of transgenic expression of protein-based Ca$^{2+}$ biosensors in mammals might be the strategy employed to create the transgenic model. Our initial efforts to generate transgenic Ca$^{2+}$ biosensor mice using a pancellular approach, driving transgene expression by β-actin or cytomegalovirus promoters, failed. Attempts to produce transgenic YC2.1 mice, a cameleon that measures cytoplasmic Ca$^{2+}$, resulted in embryonic lethality or insufficient levels of expression for detection (M. W. Roe and M. Rincon, unpublished observations). Tissue-specific targeting of YC3.3er resulted in the generation of transgenic mice with brightly fluorescent cells that were easily visualized (Fig. 1, B and C). Our studies indicate that mice tolerate expression of a GFP/calmodulin-based Ca$^{2+}$ biosensor throughout maturation from neonate to adult without impairing the physiology of the transgenically targeted cells. The absence of a
detectible effect of the MIP-YC3.3-er transgene on whole animal carbohydrate regulation is similar to transgenic mice expressing GFP in H9252-cells (7). The primary physiological function of H9252-cells is blood glucose sensing and insulin secretion, and it is well known that these processes are critically dependent on precise regulation of intracellular Ca2\(^{+}\) signaling (8). The findings suggest that MIP-YC3.3-er expression levels in the transgenic mice do not affect endogenous calmodulin-dependent physiological processes in vivo. This hypothesis is in close agreement with in vitro evidence from previous studies of the cameleons (12).

In the present study, we have established that a genetically encoded functional fluorescent Ca2\(^{+}\) biosensor can be expressed transgenically in intact mammalian tissues and engineered to label specific cells within a multicellular organ system. Our results suggest that cell-specific targeting of GFP-based biosensors is a feasible approach to generate transgenic expression in mammals. It is conceivable that this approach could be adapted to exploit new biosynthetic fluorescent sensor designs as they become available and advance real-time optical analysis of signal transduction and cellular biochemistry in complex organ systems of living animals.

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