Homogeneous insulin and C-peptide sensors for rapid assessment of insulin and C-peptide secretion by the islets.

Running title: Homogeneous insulin and C-peptide sensors.

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Objective: Glucose-stimulated islet insulin or C-peptide secretion experiments are a fundamental tool for studying and assessing islet function. The goal of this work was to develop antibody-based fluorescent homogenous sensors that would allow rapid, and inexpensive near-instantaneous determinations of insulin and C-peptide levels in biological samples.

Research Design and Methods: Our approach was to use molecular pincer design (Heyduk, E. et al. Anal Chem 2008, 80, 5152-5159) in which a pair of antibodies recognizing non-overlapping epitopes of the target are modified with short fluorochrome-labeled complementary oligonucleotides and are used to generate Fluorescence Energy Transfer (FRET) signal in the presence of insulin or C-peptide.

Results: The sensors were capable of detecting insulin and C-peptide with high specificity and with picomolar concentration detection limits in times as short as 20 min. Insulin and C-peptide levels determined with the FRET sensors showed outstanding correlation with determinations performed under the same conditions with ELISA. Most importantly, the sensors were capable of rapid and accurate determinations of insulin and C-peptide secreted from human or rodent islets verifying their applicability for rapid assessment of islet function.

Conclusions: Homogeneous, rapid and uncomplicated nature of insulin and C-peptide FRET sensors allows rapid assessment of β-cell function and could enable point-of-care determinations of insulin and C-peptide.

Diabetes mellitus comprises a heterogeneous group of hyperglycemic disorders. There are two major forms of diabetes: 1) type 1 diabetes which is associated with an autoimmune-mediated attack and destruction of pancreatic β-cells resulting in insulin deficiency; and 2) type 2 diabetes which is characterized by insufficient insulin production or impaired insulin action. For type 1 diabetics insulin injections are used to regulate plasma glucose levels, while type 2 diabetics are usually treated by diet, oral agents, and insulin therapy. If uncontrolled, elevated plasma levels of glucose increases the risk for the development of diabetic complications such as cardiovascular disease, kidney disease, neurological disorders and blindness. Intensive insulin therapy reduces the risk of such complications, although there is an increased risk of hypoglycemic episodes with this therapy, which if severe, can result in coma or seizures. Insulin is synthesized as a precursor protein, proinsulin, that is processed by specific proteases found in insulin granules to the active hormone (containing an A and B chain connected by two disulfide linkages). During the processing of insulin, the connecting sequence (C-peptide) is also produced, and C-peptide is released into the bloodstream with insulin at times of insulin demand. Glucose-stimulated insulin or C-peptide secretion by the islets is a fundamental tool for studying and assessing islet function. Current methodologies to determine β-cell function rely on the use of radioimmunoassay or ELISA based assays to detect either insulin or C-peptide produced by β-cells. These methods are time consuming, and in many cases require the use of radioactivity. Development of a specific methodology for detecting insulin and/or C-peptide that reduces the time required to determine β-cell function would provide a
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tremendous advantage over methods currently being employed. Such methodology could also have clinical applications, for example, as a rapid method to characterize β-cell function in the transplantation of human islets, a potential therapy for type 1 diabetics. Transplantation of human islets, isolated from cadaver donors, has been used for a number of years in an effort to gain insulin independence in type 1 diabetics; however, this procedure has been only marginally successful(1; 2). Recently, the Edmonton group has described a new protocol for the transplantation of human islets that was successful in attaining short-term insulin independence in 7 of 7 patients(3), and similar results have been reproduced by others(4-6). One key feature of the Edmonton protocol is the immediate transplantation of islets following isolation. Previous protocols cultured islets for extended periods of time in part to determine islet viability and function prior to transplantation (7). A methodology that would allow rapid assessment of glucose-stimulated insulin secretion could be thus potentially employed to enhance characterization of islets before transplantation.

In a previous study we described a new antibody-based sensor technology that allowed simple fluorescence-based homogenous detection of target proteins(7). The goal of this work was to determine if this sensor design could be adapted for rapid detection of insulin and C-peptide and utilized for very rapid determination of islet functional activity (insulin secretion). We show that our sensors allow near instantaneous determination of the insulin or C-peptide produced by isolated human or rodent islets validating their applicability for rapid assessment of islet quality. We believe that the uncomplicated and rapid characteristics of our assay will allow for the deployment of methods for insulin and C-peptide determinations in a point-of-care setting.

RESEARCH DESIGN AND METHODS

Sensor design. Fig. 1 illustrates general design of homogenous sensors for insulin or C-peptide that is based on previously described molecular pincer assay(7). A pair of antibodies recognizing non-overlapping epitopes of insulin (or C-peptide) are functionalized by attaching short complementary oligonucleotides via long nanometer scale flexible linkers. Oligonucleotides are modified with a pair of fluorophores (fluorescein and Cy5) that could function as a donor and an acceptor in Fluorescence Resonance Energy Transfer(10) (FRET). In the presence of the target analyte both antibodies will bind to the target molecule resulting in a large increase of the local concentration of signaling oligonucleotides. This in turn will lead to annealing of the oligonucleotides bringing the fluorophores to close proximity resulting in efficient FRET that can be used as a signal for target analyte detection.

Materials. NHS-PEO₈-maleimide and Traut’s reagent were from Pierce (Rockland, IL). Oligonucleotides were obtained either from Keck Oligonucleotide Synthesis Facility at Yale University or from IDT (Coralville, IA). The following oligonucleotides were used (names in parenthesis):

(A1) 5’ C6 amino-

TAGGTGCTCGACGCTGAC

(A2) 5’ C6 amino-

TAGGAGAGAGAGAGGA

(A3) 5’-Fluorescein-

GCTCAT

TGTCAGCGTCGAGCACCTA

(A4) 5’-Cy5-

ATGAGCTTCCTCTCTCTCTCCAT

A3 and A4 oligonucleotides contain short sequences (underlined) that are used in target-induced annealing to generate FRET signal. Fluorescein and Cy5-labeled oligonucleotides were purified by reversed-phase HPLC(8).
Concentrations of oligonucleotides were calculated from UV absorbance at 260 nm, after correction for fluorophore absorbance at 260 nm.

Monoclonal anti-human insulin antibodies (clones E6E5, D4B8, 8E2, 3A6) were purchased from Fitzgerald Industries International, Inc. (Concord, MA). Monoclonal anti-human C-peptide antibodies (clones 9101 and 9103) were from BiosPacific (Emeryville, CA). Insulin and C-peptide ELISA assays kits (EZH-14C and EZHCP-20C) were purchased from Millipore (Billerica, MA) and were used according to manufacturers instruction with the exception of using our assay buffer (20 mM Tris-HCl (pH 8.0), 100 mM NaCl and 10 mM EDTA containing 0.2 mg/ml BSA) and standards prepared by us in place of those provided in the ELISA kit.

Human, bovine and porcine insulin, glucagon, insulin-like growth factor II, human C-peptide, angiotensin I and II, and ACTH were purchased from Sigma (St. Louis, MO). Canine and porcine C-peptide were obtained from Anaspec (Fremont, CA) and human proinsulin was from R&D Systems (Minneapolis, MN).

**Antibody modification.** Antibodies were labeled with signaling oligonucleotides using a previously described procedure(7) (method “c” in Fig. 1B from ref. 7). A1 and A2 oligonucleotides were first attached to the antibodies via long linkers followed by annealing of A3 and A4 oligonucleotides to produce antibody-A1/A3 and antibody-A2/A4 conjugates, respectively. The first step of the procedure involves preparation of a thiol-reactive oligonucleotide that is subsequently used to react with thiolated antibody. 200 µl of 5’- amine containing oligonucleotides (A1 or A2) at ~250 µM in 20mM NaH₂PO₄ (pH 7.4), 150mM NaCl and 2.5mM EDTA buffer (conjugation buffer) were mixed with 5 µl of ~250mM of NHS-PEO₈-maleimide dissolved in DMF. The reaction mixtures were incubated for 1-1.5hr at room temperature. Oligonucleotide was purified from the excess of the crosslinker by ethanol precipitation in the presence of 1mg/ml of glycogen. Precipitated oligonucleotides were dried in Speed-Vac and were stored at -20º C until they were used for antibody modification. Antibody solutions (50-75 µl) containing 0.3-0.4 mg of the protein were run on a spin column (Zeba™, Pierce, Rockford, IL) equilibrated with the conjugation buffer. Antibodies were thiolated for 1.5hrs at room temperature with 40 molar excess of Traut’s Reagent added as ~14mM stock solution in DMF. The excess of Traut’s Reagent was removed on Zeba™ spin column equilibrated in the conjugation buffer. The thiolated antibody was then reacted with a 15-20 molar excess of linker-conjugated oligonucleotide (calculated assuming that ~50% of the oligonucleotides were conjugated with the crosslinker). Reaction mixtures were incubated for 4 hrs at room temperature followed by an overnight incubation at 4º C.

Modified antibodies were purified from the excess of the oligonucleotides by size exclusion FPLC chromatography using 10/30GL Superdex™ 200 column (Pharmacia) equilibrated with 10 fold-diluted 20 mM Tris (pH 8.0), 100 mM NaCl, 10 µM EDTA buffer. Fractions containing modified antibodies were pooled and concentrated 10-fold in the Speed-Vac. The protein concentration was estimated using Bradford assay. Labeling of the antibodies with oligonucleotides was confirmed (and the extent of the labeling estimated) by analyzing the UV spectra of purified final product. Observed spectra were fitted by a linear combination of the spectra of free antibody and free oligonucleotide to determine relative amounts of the protein and oligonucleotide in the sample.

**Islet Isolation.** Human islets were isolated from cadaver donors using protocols approved by the IRB at University of
Alabama in Birmingham. Rodent islets were isolated from male Sprague Dawley rats (250-300 g) by collagenase digestion as previously described(9).

Islets were cultured overnight in CMRL-1066 (containing 2 mM L-glutamine, 10% heat-inactivated fetal calf serum, 100 U/ml penicillin, and 100 µg/ml streptomycin) at 37°C under an atmosphere of 95% air and 5% CO₂ prior to experimentation.

**Glucose-stimulated Insulin Secretion**: The islets were washed with KRB buffer (25 mM Hepes, 115 mM NaCl, 24 mM NaHCO₃, 5 mM KCl, 1 mM MgCl₂, 2.5 mM CaCl₂, and 0.1% BSA, pH 7.4) containing 3.3 mM glucose followed by preincubation for 30 min at 37°C in KRB containing 3.3 mM glucose. The islets (~ 10 and ~15 in case of human or rodent, respectively) were aliquoted to vials containing either KRB with 3.3 mM glucose or KRB with 16.7 mM glucose and were incubated for 60 min (human islets) (KRB with 20 mM glucose and 30 min incubation in the case of rodent islets) at an atmosphere of 95% air, 5% CO₂ at 37°C. After the incubation, the supernatant was removed and the samples were stored frozen at -70 deg until analyzed. In experiments where FRET sensors were compared with ELISA, the samples were thawed and analyzed in parallel by sensors and ELISA to eliminate any variability due to differences in sample treatment.

**Insulin and C-peptide determinations.** All insulin and C-peptide measurements were performed in 20 mM Tris-HCl (pH 8.0), 100 mM NaCl and 10 μM EDTA containing 0.2 mg/ml BSA (assay buffer). Standard solutions of insulin and C-peptide were prepared from stock solutions obtained by dissolving human insulin and C-peptide (from Sigma (St. Louis, MO)) in assay buffer. Concentrations of stock solutions were assigned based on information provided by the supplier. The standards were not calibrated to WHO reference material. Stock solutions were stored frozen in small aliquots at -70 deg. No significant differences were observed between the results obtained with different samples of stock insulin or C-peptide solutions. When comparing FRET sensors with ELISA, to eliminate variability of the results due to potential instability of calibrators included in commercial ELISA kits, FRET sensors and ELISA experiments were done in parallel using the same solutions of insulin or C-peptide standards for preparation of FRET sensor or ELISA calibration curves. Thus, ELISA assays were not performed exactly according to manufacturer’s instructions. While this may have affected somewhat the performance of ELISA, it was important or the purpose of validating sensor assay to perform sensor and ELISA assays under the same conditions. Typically for a standard curve 8 samples of insulin (or C-peptide) at concentrations in 0 – 15 nM range were prepared by diluting stock solutions in assay buffer. Standard curves for quantitative determinations performed by ELISA were obtained by assaying 2.5-5 μl of standard solutions diluted to 60 μl with the assay buffer. Standard curves using the fluorescent sensor methodology were obtained by assaying 5 μl of standard solutions diluted into 20 μl of assay mix. Concentrations of insulin and C-peptide in unknown samples were determined from the described above calibration curves using absorbance values (ELISA) obtained with 2.5-5 μl of the samples diluted to 60 μl with the assay buffer or using fluorescence values (sensors) obtained with 5 μl of the samples diluted into 20 μl of assay mix. Since the samples from islet secretion experiments were in modified Kreb’s Ringer bicarbonate (MKBr) buffer, 5 μl of MKBr buffer were included in 20 μl assay mix used to measure calibration standards in this case. Absorbance at 450 nm of ELISA assay samples was measured in 96-well plates using SpectrofluorPLus plate reader (Tecan). Fluorescence of sensor mixes were measured...
in 20 µl in 384-well low-volume black microplates (Corning cat #3676) at 25 ºC. Typically, sensor assay mixtures contained (final concentrations in 20 µl) 20 nM and 25 nM or 10 nM and 12.5 nM of fluorescein-labeled antibody and Cy5-labeled antibody, respectively. Prior to fluorescence measurements, the A1 and A2-labeled antibodies were annealed with fluorescent A3 and A4 oligonucleotides by incubating 100 nM antibodies with equimolar amounts of A3 or A4 for 30 min at room temperature. These 100 nM stocks of labeled antibodies were then used to prepare 2x sensor mix solutions in the assay buffer. Appropriate volumes of samples and assay buffer were then added to 10 µl of 2x assay solutions to obtain final assay volume of 20 µl that was used for fluorescence measurements.

The donor (fluorescein; excitation at 485 nm, emission at 535 nm) and sensitized acceptor emission (Cy5; excitation at 485 nm, emission at 665 nm) signals were read with Analyst AD plate reader (LJL Biosystems, Sunnyvale, CA). Reaction mixtures were incubated for at least 20 min before fluorescence measurements were made (with the exception of kinetic experiments where incubation time varied).

Results of fluorescence measurements were expressed as normalized FRET signal change (ΔFRET):

\[
\Delta FRET = \frac{F_{SA}/F_D}{F_{SA0}/F_{D0}} - 1
\]

where \( F_{SA} \), \( F_D \), \( F_{SA0} \) and \( F_{D0} \) are sensitized acceptor and donor emission intensities in the presence and absence of insulin (or C-peptide), respectively. Buffer background was subtracted from the measured fluorescence intensities before FRET values were calculated. Using the ratio of sensitized acceptor and donor emission intensities for calculating FRET according to eq. 1 reduces the variability of FRET measurements due to dilution and instrumentation errors.

Radioimmunoassay (RIA) determinations of insulin in rat islet secretion experiments were performed by Washington University DRTC (Saint Louis, MO).

**RESULTS AND DISCUSSION**

We have previously demonstrated feasibility of the sensor design illustrated in Fig. 1 using thrombin, cardiac troponin and C-reactive proteins as models and using antibodies(7) or aptamers(11) as the binders. The assay depicted in Fig. 1 requires only adding the sample to the sensor mix followed by a simple fluorescence intensity measurement. We hypothesized thus that if the sensors with adequate sensitivity and specificity could be developed for insulin or C-peptide, they could be an extremely useful tool in rapid assessment of islet function.

We first prepared an insulin sensor using anti-insulin monoclonal antibodies clones E6E5 and D4B8 (Fitzgerald Industries International). When titrated with increasing concentrations of insulin, a robust and highly reproducible insulin concentration-dependent FRET signal was observed (Fig. 2A, grey symbols). While robust, the signal was lower when compared to what we have previously observed in case of troponin or C-reactive protein (7). We wondered if the antibody pair we selected for the sensors was indeed the most optimal. We tested another monoclonal antibody pair (clones 8E2 and 3A6 from Fitzgerald Industries International, Inc.) that according to manufacture’s specifications had higher insulin binding affinity. The sensors prepared from this antibody pair exhibited a much better (~ 5 fold higher) FRET signal in the presence of insulin (Fig. 2A, black symbols). Limit of detection was also improved substantially (from ~ 1 nM (grey symbols) to ~ 100 pM (black symbols)). Upper limit of detection is function of sensor concentration used and thus can be tailored to desired value by changing sensor concentration (data nor shown). Analytical sensitivity (defined as the mean of the measured concentrations to the actual
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concentrations) was 1.13+/−0.13. Comparison of these two data sets obtained with two pairs of monoclonal antibodies shows that the binding affinity of the antibodies that are used for preparing the sensors is a key determinant of the performance of the sensors.

In the next set of experiments a sensor of similar quality to that obtained for insulin was developed for C-peptide. Fig. 2B shows that, as in the case of insulin, a C-peptide sensor exhibited a robust and highly reproducible C-peptide concentration-dependent response. The estimated limit of detection (~ 50 pM) was slightly better than the limits for the insulin sensor. Analytical sensitivity was 0.99+/−0.06.

The data shown in Fig. 3A demonstrates specificity of insulin sensor. While a robust response in the presence of insulin is observed, no FRET signal was detected in the presence of the negative control (C-peptide). More extensive data regarding insulin sensor specificity is summarized in Table S9 (Supplemental Data in the online appendix available at http://diabetes.diabetesjournals.org). No significant crossreactivity was observed for any of the C-peptides or various unrelated polypeptides tested. The sensor could detect bovine, porcine and rat insulin as well (in some case better) as human insulin. Partial crossreactivity with human proinsulin was observed. Specificity of the sensors reflects specificity of the antibodies that were used to prepare them so it could be altered by using different set of antibodies for sensor preparation, if available. We expected that our sensors, due to their homogenous nature, would allow quick, uncomplicated measurement of insulin in a sample. Fig 3B shows the time dependence of insulin sensor response (FRET signal) upon addition of indicated amounts of insulin to sensor mix. Full response was obtained within ~ 20 min with ~ 50% of the maximum signal observed 5 min after addition of the test sample. Data in Fig. 3B demonstrate a key feature of our sensors, the ability to detect insulin in a very rapid manner using a simple homogenous assay format that does not require specialized instrumentation (all the data were collected using a standard fluorescence plate reader). The C-peptide sensor was also highly specific as the FRET signal was not observed in the presence of human insulin (Fig. 3C). More extensive data regarding C-peptide sensor specificity is summarized in Table S10 (Supplemental Data). No significant crossreactivity was observed for any of the C-peptides from other species, any insulins or various unrelated polypeptides tested. Partial crossreactivity with human proinsulin was observed. The time course of the response of the C-peptide sensor (Fig. 3D) was very similar to the insulin sensor requiring ~ 20 min for the maximal FRET signal and exhibiting >50% of maximal signal change after 5 min incubation.

The data presented in Figs. 2-3 established the feasibility of homogenous FRET sensors for insulin and C-peptide. To examine the performance of these sensors for quantitative determination of insulin and C-peptide we compared our sensors with standard and established methodologies employed in commercially available ELISA kits (Fig. 4). Thirty-two samples containing randomly chosen concentrations of insulin or C-peptide (in 0 to 14 nM range) were prepared in the assay buffer. The levels of insulin (Fig. 4A) and C-peptide (Fig. 4B) in these samples were quantified in parallel using our sensors and ELISA. There was outstanding correlation between the two methods (R² ≥ 0.98; Fig. 4) demonstrating the utility of FRET sensors for quantitative determination of insulin and C-peptide.

To confirm the functionality and to examine the practical utility of our sensors the levels of insulin and C-peptide were determined in randomly chosen samples from glucose-stimulated secretion experiments
performed with human islets. Thirty two samples from secretion experiments at low and high glucose were analyzed in parallel by FRET sensors and ELISA assays (Fig. 5). In every sample tested it was possible to determine the level of insulin (or C-peptide) using FRET sensors showing that the sensors could be used for measuring insulin and C-peptide secreted at low and high glucose. There was a cluster of data points in Fig. 5 at low insulin (or C-peptide) corresponding to low glucose samples and a more diffuse cluster at higher insulin (or C-peptide) corresponding to high glucose samples. For both insulin and C-peptide, the correlation between the results obtained with the sensors and ELISA was outstanding ($R^2 = 0.98$ and $0.99$, respectively) demonstrating the utility of the sensors for rapid assessment of islet function. Similarly, in randomly chosen samples from rodent islets secretion experiments, insulin levels were determined using our sensors and RIA assay (Fig. 6) and excellent correlation between the two techniques was observed.

Results of experiments used to obtain basic performance characteristics of FRET sensor assays are presented in Supplemental Data. Between and intra assay precision (Table S1-S4), spike and recovery (Table S5-S6), assay linearity (Table S7-S8), and assay crossreactivity (Tables S9-S10) were determined and were found to be similar to what’s typically observed for ELISA assays.

CONCLUSIONS

Insulin and C-peptide sensors that we have developed allow rapid determination of insulin and C-peptide with high specificity and sensitivity by a procedure that requires only the addition of the sample to the sensor mix. The sensors exhibit sensitivity that is sufficient for rapid assessment of islet functionality by near-instantaneous measurement of insulin or C-peptide secreted by the islets. The use of the sensors does not require any complicated specialized instrumentation (any standard fluorescence plate reader will be sufficient).

Limits-of-detection that we observe with these sensors are near the concentrations of insulin and C-peptide observed in serum (for example, normal insulin concentrations in plasma are 40-160 pM). However, in its current formulation the sensors are not yet suitable for serum measurements. Although they do work with serum samples (data not shown), the significant background fluorescence produced by serum requires significant sample dilution. In order to use these sensors for serum measurements the sensors will need to be further improved to increase their sensitivity (~5-10 fold) and to reduce the background of the serum (by employing fluorescence probes with excitation at higher wavelength or by using time-resolved fluorescence signal detection). We believe these improvements will be possible and if successful, they will widen significantly the applicability of our sensors. With these improvements we envision that the sensors could be adopted for use with a simple hand-held fluorescence reader allowing point-of-care insulin and C-peptide determination.

Author Contributions. E.H. researched data, contributed to discussion, reviewed/edited manuscript, wrote manuscript. M.A.M. contributed to discussion, reviewed/edited manuscript. A.S. researched data, reviewed/edited manuscript. J.A.C. contributed to discussion, wrote manuscript, reviewed/edited manuscript. T.H. contributed to discussion, reviewed/edited manuscript, wrote manuscript.

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REFERENCES


FIGURE LEGENDS

Fig. 1. Design of homogenous sensors for detecting insulin and C-peptide. “T” denotes a target (insulin or C-peptide).
Fig. 2. (A) Response of the insulin sensor to indicated amounts of human insulin. Experiments were performed with a sensor mixture containing 20 nM fluorescein-labeled antibody and 25 nM of Cy5-labeled antibody. Black and grey symbols correspond to experiments performed with a sensors utilizing 8E2/3A6 and E6E5/D4B8 antibody pairs, respectively. The average and standard deviation of three measurements are shown. (B) Response of the human C-peptide sensor to indicated amounts of C-peptide. Experiments were performed with a sensor mixture containing 20 nM fluorescein-labeled antibody and 25 nM of Cy5-labeled antibody. The average and standard deviation of three measurements are shown. FRET signals were read after a minimum of 20 min. incubation after adding sample to the sensor mix.

Fig. 3. (A) Specificity of insulin sensor. Response of insulin sensor to human insulin and human C-peptide is compared. Sensor mixture containing 20 nM fluorescein-labeled 8E2 antibody and 25 nM of Cy5-labeled 3A6 antibody was used. FRET signals were read after a minimum of 20 min incubation after adding sample to the sensor mix. (B) Kinetics of insulin sensor response at indicated human insulin concentrations. Sensor mixture containing 20 nM fluorescein-labeled E6E5 antibody and 25 nM of Cy5-labeled D4B8 antibody was used. (C) Specificity of C-peptide sensor. Response of C-peptide sensor to human C-peptide and human insulin is compared. FRET signals were read after a minimum of 20 min incubation after adding sample to the sensor mix. (D) Kinetics of C-peptide sensor response at indicated C-peptide concentrations. Sensor mixtures containing 20 nM fluorescein-labeled antibody and 25 nM of Cy5-labeled antibody were used.

Fig. 4. Correlation between measurements using the sensors and ELISA assay in 32 samples containing randomly selected concentrations of human insulin (A) and human C-peptide (B) dissolved in a buffer.

Fig. 5. Correlation between measurements of human insulin (A) and human C-peptide (B) using the sensors and ELISA assays in human islet secretion experiments. The average and standard deviation of three measurements of 32 samples are shown.

Fig. 6. Correlation between measurements of rat insulin using the sensors and RIA assay in 15 samples from rodent islet secretion experiments.
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Figure 1

Figure 2

A

B

Normalized ΔFRET

Normalized ΔFRET

insulin (nM)

c-peptide (nM)
Figure 5

A

\[ R^2 = 0.98 \]
\[ y = 1.15x + 0.12 \]

B

\[ R^2 = 0.99 \]
\[ y = 0.88x - 0.16 \]

Figure 6

\[ R^2 = 0.97 \]
\[ y = 0.72x + 0.18 \]