**SUMMARY**

NADH is a key metabolic cofactor whose sensitive and specific detection in the cytosol of live cells has been difficult. We constructed a fluorescent biosensor of the cytosolic NADH-NAD^+ redox state by combining a circularly permuted GFP T-Sapphire with a bacterial NADH-binding protein, Rex. Although the initial construct reported [NADH] × [H^+] / [NAD^+], its pH sensitivity was eliminated by mutagenesis. The engineered biosensor Peredox reports cytosolic NADH:NAD^+ ratios and can be calibrated with exogenous lactate and pyruvate. We demonstrated its utility in several cultured and primary cell types. We found that glycolysis opposed the lactate dehydrogenase equilibrium to produce a reduced cytosolic NADH-NAD^+ redox state. We also observed different redox states in primary mouse astrocytes and neurons, consistent with hypothesized metabolic differences. Furthermore, using high-content image analysis, we monitored NADH responses to PI3K pathway inhibition in hundreds of live cells. As an NADH reporter, Peredox should enable better understanding of bioenergetics.

**INTRODUCTION**

Nicotinamide adenine dinucleotide (reduced, NADH; oxidized, NAD^+) is a key cofactor for electron transfer in metabolism. Reduction-oxidation (redox) reactions catalyzed by various NAD(H)-dependent dehydrogenases are vital for biochemical processes such as glycolysis and mitochondrial metabolism. In addition, NADH-NAD^+ redox has been implicated in the regulation of embryonic development and aging (Dumollard et al., 2007; Chen et al., 2009), as well as in pathological conditions such as diabetes, cancer, and epilepsy (Eto et al., 1999; Zhang et al., 2006; Garris-Canut et al., 2006).

To assess the cellular NADH-NAD^+ redox state, there have been two general approaches. Chemical methods infer the NADH+NAD^+ ratio indirectly from the concentrations of redox couples such as lactate and pyruvate (Williamson et al., 1967). However, this requires the use of cell extracts and is, thus, incompatible with studying dynamics in intact, individual cells. A less invasive optical approach monitors cellular NAD(P)H autofluorescence. Under ultraviolet excitation, NADH and another related cofactor, NADPH, are fluorescent, whereas their oxidized counterparts NAD^+ and NADP^+ are not (Chance et al., 1962). As NADH and NADPH give identical autofluorescence signals, they are collectively denoted as NAD(P)H, although the two cofactors govern distinct metabolic reactions (Klingenberg and Bücher, 1960). Because NADH is present at higher levels than NADPH in many tissues, the NAD(P)H signal has often been interpreted as changes in NADH (Lowry et al., 1957; Klingenberg and Bücher, 1960; Chance et al., 1962). However, data from simultaneous imaging of both NAD(P)H and flavoprotein autofluorescence underscore the ambiguity of the NAD(P)H signal, suggesting that it may primarily report protein-bound NADPH instead of NADH (Rocheleau et al., 2004). In addition to the signal ambiguity, in general, cytosolic and mitochondrial NAD(P)H autofluorescence have been assumed to reflect glycolysis and oxidative metabolism, respectively (Patterson et al., 2000; Shuttleworth et al., 2003; Kasischke et al., 2004; Gordon et al., 2008). In practice, mitochondrial signals dominate the measurements, whereas cytosolic signals are small and difficult to separate from the bright mitochondrial signals. For more specific and sensitive detection of the cytosolic NADH-NAD^+ redox state, a fluorescent NADH biosensor would be valuable.

One strategy to create genetically encoded fluorescent biosensors involves circularly permuted fluorescent proteins (cpFPs), derived from green fluorescent proteins (GFPs). Circular permutation joins the original N and C termini with a peptide linker and creates new termini near the chromophore (Baird et al., 1999). A specific detector or binding protein fused to the new N and C termini creates conformational coupling between binding and fluorescence. With this strategy, fluorescent biosensors have been engineered to report calcium, hydrogen peroxide, cyclic 3',5'-guanosine monophosphate (cGMP), and ATP:ADP ratio (Nagai et al., 2001; Belousov et al., 2006; Naush et al., 2008; Berg et al., 2009). For a detector domain, we chose the redox-sensing repressor Rex, a bacterial NADH-binding protein that links metabolic state to gene expression (Brekasis and Paget, 2003). Rex is a homodimer; each subunit comprises an N-terminal domain and a C-terminal NADH-binding domain (Sickmier et al., 2005). Upon NADH binding, Rex adopts a closed instead of an open conformation (Sickmier et al., 2005; Wang et al., 2008; McLaughlin et al., 2010). Here, we found that integration of a cpFP T-Sapphire into Rex yielded a fluorescent sensor of NADH. The initial construct (and likely the native Rex protein) reported [NADH] × [H^+] / [NAD^+]. By eliminating its pH
dependence via targeted mutagenesis, we constructed a genetically encoded fluorescent biosensor of cytosolic NADH:NAD+ ratios. This reporter, Peredox, revealed cytosolic NADH-NAD+ redox dynamics in mammalian cells upon metabolic challenges. With transient or stable expression, we demonstrated the utility of Peredox in a variety of primary and cultured cell types. We found different cytosolic NADH-NAD+ redox states in primary mouse cortical astrocytes versus neurons, as well as changes in cytosolic NADH-NAD+ redox in cultured epithelial cells upon perturbation of PI3K pathway signaling.

RESULTS

A cpFP Inserted into T-Rex Reports [NADH] × [H+] / [NAD+]

To engineer an NADH biosensor, we inserted a cpFP variant of T-Sapphire (Zapata-Hommer and Griesbeck, 2003) into a tandem dimer of Rex from Thermus aquaticus (T-Rex), between the two subunits. By PCR, we created a library of peptide linkers for the cpFP insertion, expressed the protein library in bacteria, and assayed the purified proteins for fluorescence responses. We found that a cpFP inserted into T-Rex could couple conformational changes with fluorescence to report NADH (Figures 1A and S1 available online). A construct named P0 exhibited an increase in green fluorescence upon NADH application (Figure 1B). Its spectra were similar to that of T-Sapphire, with an excitation peak around 400 nm and an emission peak around 510 nm. Whereas NADH application enhanced the green fluorescence, it did not change the red fluorescence of a tandemly attached mCherry (Figure S2A; Shaner et al., 2004), which was included to normalize the signal for protein expression. The affinity of P0 for NADH was < 5 nM (Figure S2B), which was surprising given that the estimated concentration of free NADH in the cytosol is in the hundreds of nanomolar (Zhang et al., 2002). In addition to NADH, NAD+ can bind to Rex. Unlike NADH, application of NAD+ yielded only minimal change in fluorescence (Figure 1B). However, increasing NAD+ concentration effectively lowered the sensor's apparent affinity for NADH (Figure 1C), indicating that NAD+ competes with NADH for binding. According to this competitive scheme, when the concentrations of NADH and NAD+ exceed their affinity constants, the sensor's steady-state fluorescence response would report the NADH:NAD+ ratio (Figure 1D; Berg et al., 2009). Although conventionally the ratio $R = [\text{NAD}^+] / [\text{NADH}]$ is reported, we plotted the response against the alternative ratio $R' = [\text{NADH}] / [\text{NAD}^+] \times 1000$. The $R'$ at which the response is half maximal is called the “$K_{R'}$” of the sensor, analogous to the dissociation constant ($K_d$) of a receptor. At pH 7.2, P0 had ~8,000-fold higher affinity for NADH compared to NAD+.
affinity for NADH than for NAD⁺; its half-maximal response corresponded to a $K_R$ of 0.12. Because P0 was constructed with a pH-resistant GFP-T-Sapphire, its fluorescence in the unoccupied state or in the NADH-bound state was pH resistant, as expected. Nevertheless, the ability of NAD⁺ to compete with NADH for binding did vary substantially with pH (Figure 1D). With higher pH, competition by NAD⁺ became stronger, further lowering the sensor’s apparent affinity for NADH and increasing $K_R$. Remarkably, this pH dependence could be described as a strict dependence of the sensor on $[\text{NADH}] \times [\text{H}^+] / [\text{NAD}^+]$ (Figure 1E). Indeed, this expression is a component of the equilibrium constant of any NAD(H)-dependent dehydrogenase, and it is proportional to the ratio of the redox couple. For instance, when the lactate dehydrogenase (LDH) reaction is at equilibrium, the lactate:pyruvate ratio is proportional to $[\text{NADH}] \times [\text{H}^+] / [\text{NAD}^+]$.

**Optimization Eliminates the pH Sensitivity of the NADH Biosensor**

Many factors other than metabolism perturb cellular pH. Although P0 reports the interesting quantity $[\text{NADH}] \times [\text{H}^+] / [\text{NAD}^+]$, its pH sensitivity renders the signal nonspecific to cytosolic NADH-NAD⁺ redox and, thus, not highly useful. Nonetheless, this quantity $[\text{NADH}] \times [\text{H}^+] / [\text{NAD}^+]$ suggests that, similar to NADH-binding dehydrogenases, there is a single proton or charge transfer for each NAD⁺ (or NADH) binding event. We reasoned that the highly conserved residue Tyr98, located near the nicotinamide moiety of the NADH molecule in the T-Rex structure (Sickmier et al., 2005), might participate in the proton transfer upon NAD⁺ binding (Jörnvall et al., 1995). We then made a protein library with various mutations on Tyr98. In addition, to speed up kinetics (see below), we performed error-prone PCR mutagenesis and screened for improved sensor variants. The best product of the screen contained the mutations Tyr98Trp and Phe189Ile in the first subunit and Tyr98Trp in the second subunit.

This circularly permutated GFP-based sensor of NADH-NAD⁺ redox, named Peredox, demonstrated notable improvements in pH resistance and kinetics for cytosolic NADH-NAD⁺ redox sensing. Whereas Peredox retained the spectral properties of the original P0 construct (Figures S3A and S3B), it was far more resistant to pH changes in the physiological range (Figures 2A and S3C). For an increase of one pH unit, which produced a 10-fold increase of $K_R$ in P0 (Figure 1D), the $K_R$ of Peredox changed only slightly, by ~20%, with a crossover point (signal insensitive to pH) at $R^* = 3$ (Figure 2A). Upon NADH saturation, Peredox displayed an increase of ~150% in fluorescence response (i.e., a 2.5-fold increase). Due to its lower affinities for NAD⁺ and NADH, Peredox reported the cytosolic NADH level, partly compensated for the NAD⁺ level rather than strictly the NADH:NAD⁺ ratio. The NADH:NAD⁺ ratio that was required for a half-maximal response was somewhat resistant to changes in the overall $[\text{NADH} + \text{NAD}^+]$. A 3-fold change in the NAD⁺ pool size in the physiological range produced an ~2-fold change in the sensor midpoint for NADH:NAD⁺ ratio (Figure S3D). Pere- dox was specific against other metabolites structurally related to NADH, such as NADPH, NADP⁺, ADP ribose, nicotinamide, β-nicotinamide mononucleotide, AMP, and adenosine. Not only was there no change in its fluorescence response upon addition of these metabolites (Figure S3E), but the NADH:NAD⁺ titrations in their presence were also similar to that of control (Figures S3F and S3G). The main interference came from ADP and ATP, with an apparent affinity ($K$) in the millimolar range (Figure S3H), consistent with published data on B-Rex (Wang et al., 2008). In Peredox, although the specificity of binding NADH over ADP or ATP was roughly 30,000-fold, NADH is far less abundant than ADP or ATP in typical intracellular environments. Although physiological levels of ADP and ATP could compete with NADH and NAD⁺ for binding, ADP and ATP act similarly to other each other at various pHs and temperatures; also, the interference from ADP and ATP is less prominent with increasing NAD⁺ (Figures 2B and S3I–S3L). Thus, though Peredox might be slightly sensitive to changes in the total size of the adenine nucleotide pool, its response is not affected by changes in the cellular energy charge or the ATP:ADP ratio. Although metabolic challenges often lead to energy deprivation and decrease in cytosolic ATP:ADP ratio, changes in cytosolic adenine nucleotide pool size are minimal (Schwenke et al., 1981; Malaisse and Sener, 1987) and would not be expected to interfere with Peredox response. In addition, Peredox exhibited much
improved kinetics over P0. To determine the rate of NADH dissociation from the biosensor, we added the LDH enzyme with pyruvate to consume the free NADH, and we monitored the response over time. While NADH dissociation from P0 was slow, with a time constant of 25 min at 25°C, NADH dissociation from Peredox was much faster, with a time constant of 50 s at 25°C and 16 s at 35°C (Figure 2C).

**Peredox Reports the Cytosolic NADH-NAD⁺ Redox State in Mammalian Cells**

After characterizing Peredox-mCherry as a purified protein, we validated its utility in mammalian cells. We expressed Peredox-mCherry in cultured mouse neuroblastoma Neuro-2a cells and monitored its fluorescence response. Under confocal microscopy, the sensor fluorescence in both green and red images was fairly uniform throughout the cell (Figure 3A). The pixel-by-pixel ratio of the green image divided by the red image appeared consistent, with no apparent difference observed between cytosolic and nuclear signals. To test whether Peredox could report the cytosolic NADH-NAD⁺ redox state, we varied the concentrations of lactate and pyruvate in the extracellular solution, thereby altering the intracellular concentrations of these metabolites (Bücher et al., 1972). Interconversion between lactate and pyruvate catalyzed by endogenous LDH should lead to concomitant exchange between cytosolic NADH and NAD⁺ species. Note that cells were not permeabilized, and glucose was absent in the extracellular solution. With wide-field time-lapse microscopy, we monitored changes in the fluorescence response of Peredox. In addition, we established that cellular background autofluorescence was far weaker and did not interfere with the Peredox fluorescence signal (Figure S4). In the presence of lactate (10 mM), the green-to-red fluorescence ratio of Peredox was maximal (Figure 3B). After adding incremental amounts of pyruvate, we observed a stepwise decrease in the green-to-red fluorescence ratio, with millimolar pyruvate reducing the ratio down to 40% of the maximal value. This signal change was consistent with oxidation of NADH into NAD⁺ and a concurrent reduction of pyruvate into lactate. After each solution switch, the signal adjusted rapidly and arrived at a new steady state in a few minutes. The steady-state fluorescence response of Peredox depended systematically on the lactate:pyruvate ratio in the extracellular solution (Figure 3C); the lactate:pyruvate ratio that was required for a half-maximal response was ~40. To test the robustness of this ratio-sensing behavior, we repeated the experiments with different total concentrations of lactate and pyruvate. In each case, the lactate:pyruvate ratio, rather than their absolute concentrations, determined the response. We could also plot the Peredox response to the extracellular lactate:pyruvate ratio as a function of predicted NADH:NAD⁺ ratio (Figure 3D) by using the known equilibrium constant of the reaction.

**Figure 3. Imaging the Cytosolic NADH-NAD⁺ Redox State in Various Extracellular Lactate:Pyruvate Ratios**

(A) (Left) Confocal green and red fluorescence images of two cultured mouse neuroblastoma Neuro-2a cells expressing Peredox supplied with 10 mM lactate, 10 mM lactate plus 0.2 mM pyruvate, or 10 mM pyruvate. Scale bar, 20 μm. (Middle) Pseudocolored pixel-by-pixel green-to-red ratio images. The slight edge effect seen is likely due to the optical z shift of 1.5 μm between the green and red confocal images. (Right) Wide-field differential interference contrast (DIC), nuclear staining, and the overlay image. (B) Time course of fluorescence ratios of four Neuro-2a cells perfused with the indicated lactate:pyruvate ratios. (C) Steady-state fluorescence responses of Neuro-2a cells plotted against extracellular lactate:pyruvate ratios, with lactate of 10 mM or 20 mM (mean ± SEM, n = 12–15 cells from two independent experiments). (D) Steady state fluorescence responses in (D) plotted against the predicted $R'$, by assuming the LDH reaction was at equilibrium and a pH of 7.4. Data of purified Peredox proteins were with 80 μM NAD⁺ and 4.6 mM total adenine nucleotides at 35°C. Line fitted with a logistic function using a Hill coefficient of 1.8.
LDH reaction (Williamson et al., 1967) and assuming a constant physiological pH of 7.4. For comparison, we also show the data from purified Peredox proteins, with total adenine nucleotide of 4.6 mM and free NAD⁺ of 80 μM, consistent with published estimates (Veech et al., 1979; Zhang et al., 2002). In both the dynamic range and the relative affinities for NADH and NAD⁺, there is good agreement between the sensor responses in cells and from purified proteins. Therefore, Peredox reports cytosolic NADH:NAD⁺ ratios in mammalian cells. Furthermore, because, in the absence of glucose, we could use exogenous lactate and pyruvate to set the cytosolic NADH:NAD⁺ redox, it appears that lactate and pyruvate equilibrate readily between extracellular and intracellular environments and that the LDH reaction is approximately at equilibrium.

**Glycolysis Opposes the LDH Equilibrium to Produce a Reduced Cytosolic NADH-NAD⁺ Redox State in Cultured Cells**

In addition to LDH, glucose metabolism via glycolysis at the glyceraldehyde-3-phosphate dehydrogenase (GAPDH) reaction would be expected to affect the cytosolic NADH-NAD⁺ redox state. In cultured Neuro-2a cells supplied with glucose alone, we found the sensor signal to be maximal (Figure S5). To test whether such a strongly reduced cytosolic NADH-NAD⁺ redox state depended on glucose metabolism, we performed the following experiments. First, we varied the concentration of glucose in the extracellular solution (Figure S5A). Not only was the signal minimal in the absence of glucose, but its steady-state fluorescence response also depended on the glucose supply; the glucose concentration required for a half-maximal response was ~0.2 mM (Figure 4A). Second, when we applied iodoacetate, which irreversibly inhibits GAPDH, the signal promptly decreased (Figure S5B), indicating a decline in cytosolic NADH that was consistent with glycolytic inhibition. Therefore, the reduced cytosolic NADH-NAD⁺ redox state in cultured Neuro-2a cells supplied with glucose depended on both the presence of glucose and the GAPDH reaction. Nevertheless, given that lactate and pyruvate could readily equilibrate across the cell membrane, intracellular lactate and pyruvate might be washed away in cells that were perfused with glucose alone. If availability of intracellular pyruvate were to become limiting for the LDH reaction, this could account for the reduced NADH-NAD⁺ redox state in these cultured cells. To address this concern, we systematically varied the total concentrations of extracellular lactate and pyruvate (while keeping a constant lactate:pyruvate ratio of 10), with or without supplying glucose (Figure 4B). On one hand, when glucose was absent, Peredox indicated a cytosolic NAD⁺:NADH ratio of ~300 across a wide range of total lactate and pyruvate concentrations. On the other hand, when glucose was present, the cytosolic NADH-NAD⁺ redox state became significantly more reduced. Though this observation appeared more pronounced at lower concentrations of lactate and pyruvate, it held across a wide concentration range. With physiological amounts of lactate and pyruvate (totaling 0.8–4 mM; Williamson et al., 1967), metabolic redox state differed from the NAD⁺:NADH ratios by 2- to 4-fold to 70–130, when availability of intracellular pyruvate should not be limiting. We obtained similar results in another cultured tumor cell line, rat glioma C6 (Figure S6). In both cases, the redox status of cytosolic NADH-

**Figure 4. Cultured Mouse Neuroblastoma Neuro-2a Cells Supplied with Glucose Show a More Reduced Cytosolic NADH-NAD⁺ Redox State**

(A) Steady state fluorescence responses of Neuro-2a cells plotted against concentrations of extracellular glucose (mean ± SEM, n = 19 cells from two independent experiments). (B) Steady state fluorescence responses of Neuro-2a cells plotted against total concentrations of extracellular lactate and pyruvate, with a constant lactate:pyruvate ratio of 10 and glucose of 10 mM or 0 mM (mean ± SEM, n = 7 cells from three independent experiments). For the alternate y axis, the predicted NAD⁺:NADH ratio was calculated from purified protein measurements. p < 0.001 (paired t test) for all conditions in 10 mM versus 0 mM glucose.

**Primary Cultured Cortical Astrocytes and Neurons Differ in Their Cytosolic NADH-NAD⁺ Redox States**

Glucose metabolism has been proposed to differ in astrocytes and neurons (Pellerin and Magistretti, 1994; Herrero-Mendez et al., 2009). To investigate whether their cytosolic NADH-NAD⁺ redox states differ, we expressed Peredox-mCherry in primary cultured mouse cortical astrocytes and neurons. However, we observed bright red puncta that appeared to be lysosomal aggregates, due to accumulation of mCherry (Katayama et al., 2008); these red puncta rendered the normalized signal unreliable. To circumvent this problem, we used a nuclear-targeted...
Given that glucose metabolism is regulated by growth factor activation (Jones et al., 2008; Sengupta et al., 2010), we explored the effects of PI3K pathway signaling on glucose metabolism, as indicated (mean ± SEM, n = 4–5 cells from four independent experiments). For the alternate y axis, the predicted NAD⁺:NADH ratio was calculated from purified protein measurements. p < 0.01 (paired t test) for astrocytes versus neurons prior to the 10 mM lactate condition.

Stably Expressed in MCF-10A Cells, Peroxidex Reports Cytosolic NADH Decrease upon PI3K Pathway Inhibition

Given that glucose metabolism is regulated by growth factor signaling, including the PI3K/Akt/mTOR pathway (Vander Heiden et al., 2009; Sengupta et al., 2010), we explored the effects of PI3K pathway signaling on glucose metabolism, as indicated (mean ± SEM, n = 4–5 cells from four independent experiments). For the alternate y axis, the predicted NAD⁺:NADH ratio was calculated from purified protein measurements. p < 0.01 (paired t test) for astrocytes versus neurons prior to the 10 mM lactate condition.

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detects free NADH and reports the biologically relevant fluorescence signal (Ogikubo et al., 2011). In contrast, Peredox pH could affect the NAD(P)H-protein interaction and the auto-
NAD(P)H species, changes in intracellular environment such as the autofluorescence is contributed mostly by protein-bound (Avi-Dor et al., 1962; Rocheleau et al., 2004). Also, because NADH and NADPH and, therefore, is not specific for NADH alone NAD(P)H autofluorescence reflects the combined signal of (Shuttleworth et al., 2003; Kasischke et al., 2004). Nonetheless, endogenous NAD(P)H autofluorescence observed in brain slices to monitor faster events, such as the dip and the overshoot of a time resolution of a few seconds. It may be difficult to use it to monitor faster events, such as the dip and the overshoot of endogenous NAD(P)H autofluorescence observed in brain slices (Shuttleworth et al., 2003; Kasischke et al., 2004). Nonetheless, NAD(P)H autofluorescence reflects the combined signal of NADH and NADPH and, therefore, is not specific for NADH alone (Avi-Dor et al., 1962; Rocheleau et al., 2004). Also, because the autofluorescence is contributed mostly by protein-bound NAD(P)H species, changes in intracellular environment such as pH could affect the NAD(P)H-protein interaction and the auto-
fluorescence signal (Ogikubo et al., 2011). In contrast, Peredox detects free NADH and reports the biologically relevant NADH:NAD⁺ ratio, and its response is highly specific for NADH over NADPH. Furthermore, Peredox is roughly 100-fold brighter than NAD(P)H autofluorescence (Table S1). With Peredox, we could reliably monitor cytosolic NADH, which is difficult to measure with autofluorescence imaging, particularly in the face of bright mitochondrial autofluorescence (nuclear autofluores-
cence has been used as a proxy for cytosolic autofluorescence; Patterson et al., 2000). Lastly, because NAD(P)H autofluorescence requires ultraviolet (one-photon) excitation, which could induce cell toxicity, two-photon microscopy is usually performed instead. Conversely, with a red-shifted excitation wavelength and brighter fluorescence, Peredox can be used in conventional one-photon microscopy, thereby simplifying the technical requirements for imaging experiments.

In designing Peredox, we have substantially minimized the interference of pH. In general, upon metabolic manipulations, changes in intracellular pH are common. This may interfere with biosensor measurements by altering the GFP fluorescence or the ligand-binding protein scaffold. Many cpFP-based fluo-
rescent biosensors are pH sensitive, with pKₐ’s near the physio-
logical range (Nagai et al., 2001; Belousov et al., 2006; Nausch et al., 2008; Berg et al., 2009). For some biosensors, a pH fluctu-
ation of 0.3 units could be mistaken for the entire excursion of the sensor response. Fluorescence modulation in cpFP-based sen-
sors has been attributed to a shift in the pKₐ of the chromophore that occurs upon ligand binding to the detector domain (Baird et al., 1999). Despite the apparent requirement for pH sensitivity that is inherent in this proposed mechanism, we decided to use a pH-resistant GFP. In doing so, we hoped to modulate the cpFP’s quantum yield, rather than its pKₐ, upon NADH sensing. For this reason, we chose the circularly permuted T-Sapphire (Zapata-Hommer and Griesbeck, 2003), a GFP variant that is notable for a low pKₐ of ~5. Its chromophore remains almost neutral in the ground state and ionizes only upon excitation, leading to green emission with a substantial Stokes shift of ~110 nm. We optimized the sensor response in the pH-insensi-
tive regime of the GFP (corresponding to the physiological pH range in cells). The resulting biosensor is far more pH resistant and, unlike many other cpFP-based sensors, does not need pH measurement and correction unless large pH changes are expected and high precision is needed. Moreover, the use of a pH resistant cpFP in the sensor design allowed us to uncover

Figure 6. Stably Expressed in Mammary Epithelial MCF-10A Cells, Peredox Reports Cytosolic NADH Decrease upon PI3K Pathway Inhibition

(A) Heat maps of normalized fluorescence ratios of MCF-10A cells stably expressing Peredox-NLS plotted against time. After 38 min of baseline, cells were treated with 1 μM NVP-BEZ235 (left) or DMSO (right). As a control, 20 mM lactate, 20 mM lactate and 1 mM pyruvate, and 20 mM pyruvate and 0.4 mM iodoacetate were applied at red, blue, and black arrows, respectively. For each group, ~700 cells from six fields in two experiments were collected. Though the dynamic range was less than the usual 2.5-fold, this was likely due to inadequate control of extracellular lactate and pyruvate concentrations, as these cells were imaged in the 24-well plate formats and solutions were changed without rinsing, as opposed to imaging in a chamber under continuous perfusion of fresh solutions. Fluorescence ratios binned in increments of 0.012.

(B) Histograms of normalized fluorescence ratios after 1 hr treatment with DMSO (black) or NVP-BEZ235 (blue), binned in increments of 0.027.

(C) Histograms of normalized fluorescence ratios from the indicated groups after calibration with 20 mM lactate (L), 20 mM lactate and 1 mM pyruvate (L+P), and 20 mM pyruvate and 0.4 mM iodoacetate (P+i).
the inherent pH sensitivity of the NADH-binding protein scaffold. We initially conceived T-Rex as a sensor of the NADH:NAD+ ratio, so we were surprised to find that it depended on [NADH] × [H+] / [NAD+]. Given the tight hydrophobic nicotinamide-binding pocket, the implication of a single proton transfer upon NAD(H) binding in Rex was unexpected. We were nevertheless able to eliminate its intrinsic pH dependence via directed mutagenesis of the conserved residue Tyr98.

In using a genetically encoded biosensor, a potential concern is whether it will perturb the biological systems that it is used to measure; however, Peredox seems unlikely to perturb cellular-free NADH by either catalysis or buffering. The dependence of Rex on [NADH] × [H+] / [NAD+] is reminiscent of the equilibrium constant of NADH-binding dehydrogenases, raising the question of whether Rex is not just an NADH-binding protein but also a redox enzyme. If the Rex-based biosensor were to have catalytic activity, it could alter the level of cellular NADH, and binding or release of NAD(H) might depend on the presence of substrate (Kumar et al., 2002). Yet, structural and biochemical evidence seems to suggest that Rex does not possess any catalytic activity. The T-Rex structure reveals a compact hydrophobic pocket enclosing the nicotinamide moiety of the NADH molecule, without a cavity for substrate accommodation or polar functional groups characteristic of redox enzymes (Sickmier et al., 2005). Also, most NADH-binding dehydrogenases exhibit some catalysis even with nonoptimal substrates. We, however, observed no oxidation of NADH in Rex upon pyruvate application (50 mM for > 20 hr), unlike the case for the eukaryotic transcriptional co-repressor C-terminal binding protein (Kumar et al., 2002). Even if Rex were a redox enzyme, the mutation at the highly conserved Tyr98 residue would likely spoil any potential catalytic activity in Peredox. In addition, another concern was that Peredox might perturb the cytosolic-free NADH pool by direct binding and buffering. However, this effect is likely to be negligible, as cytosolic NADH is probably buffered by the endogenous protein-bound pool, constituting ~95% of the total cytosolic NADH (Zhang et al., 2002). With a total cytosolic NADH concentration of 3 μM, biosensor expression (at 1–10 μM; Akerboom et al., 2009) is expected to change cellular NADH buffering capacity by roughly 4-fold or less.

To normalize Peredox measurements, a second FP mCherry was attached in tandem. In primary cultured cells, we found that expression of RFP, such as mCherry, led to lysosomal aggregates as previously reported (Katayama et al., 2008). As these puncta preclude reliable comparison of the green-to-red fluorescence ratios, we instead used a nuclear-targeted version of Peredox-mCherry. Possible alternative approaches to avoid puncta include the use of another FP, such as mCitrine (Griesbeck et al., 2001), for normalization or fluorescence lifetime imaging to obtain self-normalized measurements without using a second FP (Tantama et al., 2011).

Being genetically encoded, Peredox may be targeted to various cell types and compartments to monitor cellular and subcellular variations in NADH metabolism. However, Peredox, with its current affinity, would not be expected to work well in mitochondria, as mitochondrial NADH:NAD+ ratio has been estimated to be 100- to 1,000-fold higher than the cytosolic NADH:NAD+ ratio (Williamson et al., 1967). Consistent with this estimate, when we targeted Peredox to mitochondrial matrix, we did not observe any signal change upon metabolic challenges. Nevertheless, it may be possible to re-engineer its NADH-binding sites for mitochondrial redox sensing.

Glycolysis opposes the lactate dehydrogenase equilibrium to produce a reduced cytosolic NADH:NAD+ redox state in cultured Neuro2a and C6 cancer cells. From a biological standpoint, the measured value of the cytosolic NADH:NAD+ redox is surprisingly reduced, corresponding to an NAD+:NADH ratio of < 150, compared with a typical range of 200–700 from tissue estimates obtained under a range of conditions (Williamson et al., 1967). Qualitatively, this is not unexpected: cancer cells are known to undergo aerobic glycolysis (Warburg, 1956), whereby most of the glucose is metabolized into lactate, and the production of lactate from pyruvate is presumably driven by elevated NADH:NAD+ ratios. However, at these extremely reduced redox values, it seems surprising that the GAPDH reaction can proceed forward to establish net glycolysis and a net production of NADH (Cerdán et al., 2006). The reduced redox level is dependent on the presence of glucose, arguing against net gluconeogenic flux. Perhaps the net forward flux through glycolysis is driven by rapid consumption of intermediate metabolites via pathways downstream of the GAPDH reaction, in accordance with the high biosynthetic requirements of cancer cells (Vander Heiden et al., 2009). In addition, considering that the GAPDH reaction has been proposed to be coupled to the subsequent step, the 3-phosphoglycerate kinase (PGK) reaction (Veech et al., 1979), future imaging experiments of both cytosolic NADH:NAD+ ratio and ATP:ADP ratio (Berg et al., 2009) could yield insights on how these parameters and the potential coupling between these two reactions regulate the state of glycolysis.

We found a difference in cytosolic NADH:NAD+ redox between primary cultures of cortical astrocytes and neurons, measured under identical conditions of extracellular glucose, lactate, and pyruvate. This redox difference, as observed in separately cultured cells, is consistent with the proposed redox gradient between astrocytes and neurons (Cerdán et al., 2006). More generally, this redox difference is consistent with the hypothesis that glucose metabolism in the brain is compartmentalized, with different cell types performing distinct metabolic functions. Whereas cytosolic NADH:NAD+ redox states in cultured cells are likely to be different than those that occur in vivo, Peredox, being genetically encoded, should in the future allow us to monitor NADH metabolism in astrocytes and neurons more physiologically, such as in brain slice preparations and in vivo.

By expressing Peredox stably in a cultured cell line, we could monitor changes in cytosolic NADH:NAD+ redox states in hundreds of individual cells upon perturbation of PI3K pathway signaling. We demonstrated that Peredox can be used in a high-content imaging format to obtain kinetically detailed metabolic data from individual cells. In the future, Peredox should facilitate studies on how NADH:NAD+ redox metabolism is regulated by growth factor signaling pathways in live cells, as well as how individual cell responses correlate with the status of other signaling systems. As a reporter of the cellular NADH:NAD+ redox state, Peredox should open the door to a better understanding of such potentially elaborate organization of metabolism and bioenergetics, as well as the consequences of such metabolic regulation for cellular signaling and physiology.
Cell Metabolism

Fluorescent Biosensor of Cytosolic NADH-NAD+ Redox

EXPERIMENTAL PROCEDURES

Gene Construction and Protein Characterization
Gene construction and characterization of the sensor are described in Supplemental Information. Sequences and plasmids for expression of Peredox have been deposited with Addgene.

Cell Culture, Transfection, and Transduction
Cells were maintained in 95% air and 5% CO2 at 37°C. Mouse neuroblastoma Neuro-2a cells (CCL-131, American Type Culture Collection) were cultured in Minimal Essential Medium (MEM; all media and serum from Invitrogen) with 10% fetal bovine serum (FBS), 24 mM NaHCO3, 2 mM HEPES, and 1 mM sodium pyruvate. Rat glioma C6 cells (CCL-107, ATCC) were cultured in Dulbecco’s modified Eagle’s medium-nutrient mixture Ham’s F-12 (DMEM/ F-12) with 2.5% FBS and 14 mM NaHCO3 (pH 7.1) (NaOH). Neuro-2a and C6 cells were plated onto protamine-coated coverslips 4–8 hr prior to transfection using Effectene (Qiagen). The next day, Neuro-2a and C6 cells were rinsed with phosphate-buffered saline and maintained in MEM with 1% FBS, 24 mM NaHCO3, 2 mM HEPES, 1 mM sodium pyruvate, and 100 units/ml penicillin with 100 μg/ml streptomycin (pen-strep; Lonza). For Neuro-2a differentiation, 10–20 μM retinoic acid was added. Primary dissociated mouse cortical astrocytes (Po-P2) and neurons (E16-E18) were prepared as described (Bartlett and Banker, 1984) and seeded onto poly-D-lysine-coated plates in MEM supplemented with 10% horse serum, 33 mM glucose, 2 mM glutamine (Lonza), 1 mM sodium pyruvate, and pen-strep for 3 hr. Then, primary astrocytes were maintained in DMEM/F-12 with 10% FBS, 24 mM NaHCO3, and pen-strep, and they were passed twice before transfection using Effectene on DIV (days in vitro) 21. Primary neurons were maintained in Neurobasal medium with B27 serum-free supplements, 2 mM glutamine, and pen-strep before transfection using calcium phosphate on DIV 4. For stable expression of Peredox-NLS, mammary epithelial MCF-10A cells (CRL-10317, ATCC) were transduced using the retroviral pMSCV vector, followed by puromycin selection. MCF-10A cells were cultured in DMEM/F-12 with 5% horse serum, 20 ng/ml epidermal growth factor (EGF), 0.5 μg/ml hydrocortisone, 100 ng/ml cholera toxin, 10 μg/ml insulin, and pen-strep, as described (Debnath et al., 2003).

Confocal and Wide-Field Time-Lapse Microscopy
Cells were imaged 2–5 days after transient transfection. The extracellular solution contained (in mM): 121.5 NaCl, 25 NaHCO3, 2.5 KCl, 1.25 CaCl2, 1.25 NaH2PO4, and 1 MgCl2, bubbled with 95% air and 5% CO2 delivered at 24 mM NaHCO3, 2 mM HEPES, 1 mM sodium pyruvate, and 100 units/ml penicillin with 100 μg/ml streptomycin (pen-strep; Lonza) before transfection using calcium phosphate on DIV 4. For stable expression of Peredox-NLS, mammary epithelial MCF-10A cells (CRL-10317, ATCC) were transduced using the retroviral pMSCV vector, followed by puromycin selection. MCF-10A cells were cultured in DMEM/F-12 with 5% horse serum, 20 ng/ml epidermal growth factor (EGF), 0.5 μg/ml hydrocortisone, 100 ng/ml cholera toxin, 10 μg/ml insulin, and pen-strep, as described (Debnath et al., 2003).

SUPPLEMENTAL INFORMATION

Supplemental Information includes Supplemental Experimental Procedures, six figures, and one table and can be found with this article online at doi:10.1016/j.cmet.2011.08.012.

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Imaging Cytosolic NADH-NAD$^+$ Redox State
with a Genetically Encoded Fluorescent Biosensor

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SUPPLEMENTAL EXPERIMENTAL PROCEDURES

Materials
Custom gene synthesis was performed by GenScript. Standard oligonucleotides were from Integrated DNA Technologies. Custom doped oligonucleotides were from Alpha DNA. Error prone PCR was performed using GeneMorph II Random Mutagenesis Kit from Stratagene. Chemicals and reagents were from Sigma unless otherwise noted. Lactate dehydrogenase (EC 1.1.1.27, rabbit-muscle, NAD+-dependent) was from Worthington.

Gene Construction
A synthetic gene encoding two tandem subunits of Rex from *Thermus aquaticus* (T-Rex) was designed with mammalian codon bias and selected restriction sites. The first subunit was from Met1 to Trp205 (wild type T-Rex numbering), and the second subunit was from Lys2 to Gly211. To minimize potential interactions with DNA or NADPH, both subunits contained the mutations S30A, R46D, K58D, E116G, and K117M. A synthetic gene encoding T-Sapphire was designed with restriction sites to facilitate circular permutaion. Circularly permuted monomeric T-Sapphire (cpmTS) had its N terminus at Phe145 (standard GFP numbering); the mutation A206K; and a linker Gly-Gly-Thr-Gly-Gly-Ser between the original N and C termini. Using nested PCR, cpmTS was inserted between the two Rex subunits. The P0 construct contained the mutation K204T in the first Rex subunit; a linker Ser-Ala-Ala-Gly-Gly-His-Gly between the first Rex subunit and the N terminus of cpmTS; mutations N146T S147A in T-Sapphire; an amino acid Thr inserted between the C terminus of cpmTS and the second Rex subunit; and the mutation K47E in the second Rex subunit. A linker Ser-Gly-Thr-Gly-Gly-Asn-Ala-Ser-Asp-Gly-Gly-Ser-Gly-Gly connected the Rex-cpmTS chimera to mCherry (Figure S1A). To create Peredox, three additional mutations (Y98W and F189I in the first Rex subunit and Y98W in the second Rex subunit) were introduced. The complete protein and DNA coding sequence of Peredox has been submitted to Addgene. For purified protein measurements, Peredox was subcloned with an N-terminal 7His tag into the bacterial expression vector pRSET B (Invitrogen). For cell imaging experiments, Peredox was subcloned into the mammalian expression vector GW1 (British Biotechnology). Peredox-NLS was created by adding the nuclear localization signal Pro-Lys-Lys-Lys-Arg-Lys-Val-Glu-Asp-Ala to the sensor C terminus (Nagai et al., 2001).

Protein Purification and Characterization
Standard DH5α bacteria expressing the constructs were grown in YT media in 96-well deep well plates at 37°C for 24 hr, then at room temperature for 1–4 days. Bacteria were then centrifuged, lysed with CellLytic B, and incubated in 96-well HIS-Select high capacity nickel-coated plates at room temperature overnight. Wells were washed with Tris buffered saline with Tween 20 at pH 8.0, and with
the MOPS buffer containing (in mM): 100 MOPS, 50 KCl, 5 NaCl, and 0.5 MgCl2, pH 7.3 (KOH). For spectroscopic and kinetic measurements, proteins were eluted using the MOPS buffer supplemented with 20 mM EGTA. Fluorescence was measured with a Fluorolog-3 spectrofluorometer (Horiba Jobin Yvon). Rapid mixing was done with a stopped-flow unit RX.2000 (Applied Photophysics). For microplate experiments, 0.1% bovine serum albumin (BSA) was added to the elution buffer. Eluted samples were transferred to 96-well microplates pre-blocked with the BSA-containing elution buffer. Fluorescence was measured with a Synergy 4 microplate reader (BioTek) using the following excitation and emission filters (BioTek): 400/30 nm and 528/20 nm for green; 575/15 nm and 635/32 nm for red. After background subtraction, all green to red fluorescence ratios were normalized by a common factor to yield a signal of 1 in the absence of pyridine nucleotides at pH 7.2. Solutions of ATP and ADP were added with Mg2+ to maintain a concentration of free Mg2+ of 0.5 mM at pH 7.3. For experiments where the chloride concentration was modified (Fig. S3F), potassium gluconate was added to balance the osmolarity. For determination of extinction coefficient and quantum yield, proteins were purified using Ni-NTA spin columns (QIAGEN) and dialyzed into the MOPS buffer with three buffer changes at 4°C. Protein concentrations were determined independently with both the BCA assay (Pierce) and the alkali denaturation method, using the absorbance of alkali denatured Aequorea chromophores (44,100 M–1 cm–1 at 447 nm; Ward, 1981). Extinction coefficients were estimated using absorbance measurements and Beer’s law. Quantum yields were determined using as the reference standards both fluorescein in 0.1 M NaOH (Lakowicz, 1999) and 9-aminoacridine in water (Weber and Teale, 1957). For comparison, the photophysical properties of NAD(P)H autofluorescence was taken from published values (Scott et al., 1970). To determine Peredox specificity against compounds structurally related to NADH, before NADH-NAD+ titration other metabolites were applied at concentrations at or above the published estimates of physiological concentrations listed below: adenosine, 200 μM (Smolenski et al., 1991); ADP-ribose, 10 μM (Perraud et al., 2005); AMP, 200 μM (Ballard, 1970); free NADP+ and NADPH, <1 μM (Zhang et al., 2002); nicotinamide mononucleotide, 100 μM (Revollo et al., 2007); and nicotinamide, 400 μM (Bitterman et al., 2002).
Supplemental References


Figure S1. Design of a Fluorescent NADH Reporter

(A) Schematic representation of P0, encoding a cpFP variant of T-Sapphire (cpmTS) inserted into a tandem dimer of Rex from *Thermus aquaticus* (T-Rex). The first subunit of T-Rex contained residues Met1 to Trp205 (wild type T-Rex numbering), and the second subunit contained residues Lys2 to Gly211. To minimize potential interactions with DNA or NADPH, both subunits contained the mutations S30A, R46D, K58D, E116G, K117M. T-Sapphire was circularly permuted, beginning at Phe145 (standard GFP numbering) and ending at Asn144; the mutation A206K was introduced to minimize potential aggregation to yield cpmTS. To normalize for sensor concentration, mCherry was attached with a linker to the C-terminus of the Rex-cpmTS chimera. Three additional mutations (Y98W and F189I in the first Rex subunit and Y98W in the second Rex subunit) were introduced in P0 to create Peredox.

(B) In this model, the green cpFP (PDB: 3evp) interposed between the two Rex subunits (blue and orange) changed its fluorescence upon NADH binding (balls and sticks). The open (left) and closed (right) conformations of Rex were from the crystal structures of B-Rex (PDB: 2vt2) and T-Rex (PDB: 1xcb), respectively. The green cpFP signal was normalized by the red fluorescence of mCherry (PDB: 2h5q).
Figure S2. Additional Characterization of Purified P0

(A) Excitation and emission spectra of the mCherry tandemly attached to Rex-cpmTS in the control condition (solid black), after addition of 100 µM NAD⁺ (dash purple), or 100 µM NAD⁺ and 0.2 µM NADH (solid green), normalized to the peak intensity in the control condition. For excitation spectra, emission was measured at 610 ± 5 nm; for emission spectra, excitation was at 580 ± 2.5 nm.

(B) NADH affinity. Fluorescence response to NADH at 25°C, normalized to initial and final values. The affinity of P0 for NADH was under 5 nM.
Figure S3. Additional Characterization of Purified Peroxidox

(A) and (B) Excitation and emission spectra of Peroxidox in the control condition (solid black), after addition of 100 µM NAD⁺ (dash purple), or 100 µM NAD⁺ and 1.5 µM NADH (solid green), normalized to the peak intensity in the control condition. In (A), for excitation spectra, emission was measured at 510 ± 5 nm; for emission spectra, excitation was at 400 ± 2.5 nm. In (B), for excitation spectra, emission was measured at 610 ± 5 nm; for emission spectra, excitation was at 580 ± 2.5 nm.

(C) Fluorescence ratios in the control condition, after addition of 170 µM NAD⁺, 4 µM NADH, or 170 µM NAD⁺ and 4 µM NADH in the pH range between 4.0 and 8.4.

(D) Fluorescence ratios at indicated pH plotted against $R'$ or $R$, with NAD⁺ of 0.08 mM or 0.23 mM.

(E) Fluorescence ratios plotted against concentrations of metabolite applied to the sensor alone.

(F) Fluorescence ratios in the presence of ADP ribose (10 µM), NADP⁺ (4 µM), NADPH (4 µM), nicotinamide (400 µM), β-nicotinamide mononucleotide NMN (400 µM), or chloride (5 mM, 55 mM, and 105 mM) plotted against $R'$ or $R$, with 0.08 mM NAD⁺.

(G) Fluorescence ratios in the presence of AMP (200 µM) or adenosine (1 mM) plotted against $R'$ or $R$, with 0.08 mM NAD⁺.

(H) Fluorescence ratios in the presence of indicated ADP and ATP plotted against $R'$ or $R$, with 0.08 mM NAD⁺ at 25°C. Fluorescence ratios (mean ± SEM, n = 3) were normalized to the control condition in the absence of pyridine nucleotides at pH 7.2 at 25°C.
Figure S3 (continued). Additional Characterization of Purified Peredox

(I-L) Fluorescence ratios at the indicated ATP:ADP ratios (low, 0.3; high, 3.6), NAD\(^+\) and total adenine nucleotide concentrations, pH, and temperature plotted against \( R' \). Fluorescence ratios (mean ± SEM, \( n = 3 \)) were normalized to the control condition in the absence of pyridine nucleotides at pH 7.2.
Figure S4. Comparison of Fluorescence Intensity between Transfected and Control Cells

Cumulative fraction plotted against green and red fluorescence pixel intensity for Neuro-2a cells transfected with Peredox (solid) or non-transfected control (dotted). For each set, multiple images were collected, and 5,000 of the brightest pixels were plotted. Data had not been background subtracted, with the region below the threshold for ratio analysis highlighted in grey. Compared to autofluorescence in the control, both green and red fluorescence signals in transfected cells were far higher.
Figure S5. Imaging Cytosolic NADH-NAD⁺ Redox Dynamics by Varying Glucose Supply or during Metabolic Challenge

(A) Time course of fluorescence responses of four Neuro-2a cells (top) or average response (bottom; mean ± SEM, n = 9) perfused with glucose concentration as indicated.

(B) Time course of fluorescence responses of four Neuro-2a cells (top) or average response (bottom; mean ± SEM, n = 9) challenged with 0.5 mM iodoacetate (IAA) as indicated.
Figure S6. Cultured Rat Glioma C6 Cells Supplied with Glucose Showed a More Reduced Cytosolic NADH-NAD⁺ Redox State

Steady state fluorescence responses of cultured rat glioma C6 cells, plotted against total concentrations of extracellular lactate and pyruvate, with a constant lactate:pyruvate ratio of 10, and glucose of 10 mM or 0 mM (mean ± SEM, n = 5 cells from four independent experiments). For the alternate y axis, the predicted NAD⁺:NADH ratio was calculated from purified protein measurements. * p < 0.05 and ** p < 0.01 (paired t-test) vs. corresponding condition with 0 mM glucose.
Table S1. Summary of the Photophysical Properties of Peredox as Compared to the NAD(P)H Autofluorescence

Excitation wavelength maximum (Ex), emission wavelength maximum (Em), extinction coefficient (EC), quantum yield (QY), and brightness (percentage as compared to EGFP) for Peredox in the apo state, or saturated with NADH, as well as for free NAD(P)H autofluorescence (Scott et al., 1970). In this case, mCherry was not attached to the NADH biosensor to minimize the interference of RFP on the measurements of the green signal.

<table>
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<th>Ex (nm)</th>
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