GEM-IL: A highly responsive fluorescent lactate indicator

Graphical abstract

Highlights

- Development of a genetically encoded metabolic indicator for lactate (GEM-IL)
- Mutagenesis and fluorophore alteration improve responsivity and specificity to lactate
- In vivo model generation, validation, and application of GEM-IL
- GEM-IL shows improved sensitivity over currently available lactate indicators

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In brief
Bekdash et al. report the development of a genetically encoded metabolic indicator for probing lactate (GEM-IL) based on superfolder fluorescent proteins and mutagenesis. With improvements in the design, specificity, and sensitivity, the authors demonstrate GEM-IL’s usefulness and functionality compared with previous lactate indicators, Laconic and Green Lindoblum.
GEM-IL: A highly responsive fluorescent lactate indicator

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SUMMARY

Lactate metabolism has been shown to have increasingly important implications in cellular functions as well as in the development and pathophysiology of disease. The various roles as a signaling molecule and metabolite have led to interest in establishing a new method to detect lactate changes in live cells. Here we report our development of a genetically encoded metabolic indicator specifically for probing lactate (GEM-IL) based on superfolder fluorescent proteins and mutagenesis. With improvements in its design, specificity, and sensitivity, GEM-IL allows new applications compared with the previous lactate indicators, Laconic and Green Lindoblum. We demonstrate the functionality of GEM-IL to detect differences in lactate changes in human oncogenic neural progenitor cells and mouse primary ventricular myocytes. The development and application of GEM-IL show promise for enhancing our understanding of lactate dynamics and roles.

INTRODUCTION

Lactate, once viewed simply as a waste product, has garnered increasing importance as a metabolite and signaling molecule that is shuttled between cells and organs (Adeva-Andany et al., 2014; Philp et al., 2005). As proteins, fats, and polysaccharides are broken down and generated via catabolic and anabolic pathways, lactate plays a critical metabolic role. As a major source of...
energy and signaling molecule, the importance of lactate in glycolysis, gluconeogenesis, and the citric acid cycle has been well documented (Baltazar et al., 2020; Goodwin et al., 2007; Takahashi and Yamada, 1996). Neurologically, it has been identified as a component of brain energy metabolism and may also act as an important signaling molecule that controls food intake in response to glucose levels (Bourmezeur et al., 2010; Cortes-Campos et al., 2013; Magistretti and Allaman, 2018). Neuronal energetics has focused on the metabolic relationships of glucose, pyruvate and lactate accumulation, and shutting between astrocytes and neurons (Diaz-Garcia et al., 2017; Lercundi et al., 2015; Machler et al., 2016; Vardjan et al., 2018). In arterial blood, lactate concentrations have been negatively correlated to patient survival following an ischemic event such as myocardial infarction, as well as septic shock (Hattori et al., 1985; Kawase et al., 2015; Lee and An, 2016). In oncology, cells prone to metastasizing have a higher rate of glucose uptake in an observed metabolic phenomenon called the Warburg effect. This increased glycolytic activity results in higher levels of intracellular lactate and has led to research in studying lactate transporters as a therapeutic target (Liberti and Locasale, 2016; Ovens et al., 2010; Tadogon et al., 2019). Recently, in epigenetics, lactate has garnered attention because histone lysine residues have been found to be lactylated, controlling gene expression in M1 macrophages (Zhang et al., 2019). The many roles of lactate in the cell highlight the necessity of developing optimized tools to enhance the ability to measure lactate in order to better understand its role (Chen et al., 2016; Hui et al., 2017).

Despite the increasing interest and importance of lactate metabolism, methods to measure lactate dynamics in situ and in vivo are still limited (Chen et al., 2016; Hui et al., 2017). Previously, indicators have proven to be useful in metabolic studies with multiple tools developed for metabolites such as glucose and nicotinamide adenine dinucleotide (Fehr et al., 2003; Yu et al., 2019; Zhao et al., 2015). The first genetically encoded fluorescent lactate indicator, Laconic, is based on Förster resonance energy transfer (FRET) and the bacterial lactate-binding molecule LdR (San Martin et al., 2013). Laconic has been utilized in subsequent studies to better understand neuronal energetics and to identify the function of Chaski, a Drosophila lactate/pyruvate shuttle, as well as being the basis for MitoToxy, a screening tool for drug-induced mitochondrial toxicity (Contreras-Baeza et al., 2019; Delgado et al., 2018; Diaz-Garcia et al., 2017; Lercundi et al., 2015; Machler et al., 2016). As a FRET sensor, however, it is limited in its ability to be used in tandem with multiple fluorescent indicators, a feature that would be of great utility given the dynamic and intertwined nature of metabolism levels. In addition, the in vivo studies using Laconic relied on the use of adeno-associated virus (AAV) to induce its expression, a process that is known to be immunogenic (Colella et al., 2018; Machler et al., 2016). No Laconic mouse lines are currently available. Recently, an additional lactate indicator, Green Lindoblum, was developed using an insertion of the bacterial binding molecule LdR into a loop of green fluorescent protein and is also similarlly based around LdR (Harada et al., 2020). Similar to the shortcomings accompanying Laconic, this indicator has poor intracellular expression, does not have a genetically encoded in vivo model available, and has significantly limited lactate sensitivity, rendering precise measurements of lactate flux difficult. We sought to improve upon these lactate indicators, by utilizing molecular biological approaches and mutagenesis to develop a single fluorescent protein-based indicator built with LdR. The advances in this next-generation tool expand the ability to observe the dynamics of lactate change in situ and in vivo. The use of single fluorescence indicators enables the utilization of multiple fluorescent molecules simultaneously to monitor several metabolites or other biological events, such as calcium ion (Ca(IN)) signaling (Germond et al., 2016; Lalonde et al., 2005; Zhang et al., 2013). The single fluorescent indicators have inherently low background signals as well as simpler detection methods compared with those necessitated by FRET-based indicators (Giepmans et al., 2006).

Here, we have developed and applied a genetically encoded metabolite indicator (GEM-IL) to study the dynamics and role of lactate in physiology and disease. Mouse in vivo models were developed, allowing for the cell/tissue-specific monitoring of lactate dynamics as demonstrated using isolated primary cells. In human neural progenitors, we were able to use GEM-IL to confirm the effect of c-Myc, an oncogene, which heightened lactate levels. GEM-IL can detect the effect of octylR-2HG, an onco-metabolite, on metabolic activity, while Laconic and Green Lindoblum could not. Altogether, we report the development and application of GEM-IL, showing its improvements on previous generations and its potential as a tool to enhance our understanding of the function and role of lactate.

RESULTS

Development of a genetically encoded metabolic indicator for lactate

A genetically encoded metabolite indicator for lactate, GEM-IL, was developed with the goal of developing a fluorescent lactate indicator that can directly measure the metabolite both in situ and in vivo. With this in mind, we aimed to select a reporter system that would be compatible with these different applications. As previously reported, superfolder fluorescent variants show improved stability, kinetics, and resistance to chemical denaturants, making them ideal fluorescent reporters for imaging lactate compared with non-superfolder fluorescent probes (Pedelaq et al., 2006; Stepanenko et al., 2014). The split version of a superfolder fluorescent protein has been used in other tools, separating the 11th β strand of sfGFP (sfGFP11) from the first 10 β strands (sfGFP1-10) (Cabantous et al., 2005; Feng et al., 2017; Kamiyama et al., 2016). Using these strategies and designs, we tested three fluorescent proteins: superfolder cyan (sfCFP), superfolder green (sfGFP), and superfolder yellow (sfYFP). The GFP (G), YFP (Y), and CFP (C) GEM-ILs were constructed by fusing the bacterial lactate binding domain, LdR, between the 10th and the 11th β strands (named GEM-IL prototype). We hypothesized that the mature fluorescent molecule would undergo a conformational shift upon lactate binding to LdR, resulting in a detectable and reversible fluorescence change (Figure 1A). This is supported by the previous development of a single fluorescent indicator using conformationally sensitive GFP (csGFP) that can be conformationally manipulated by inducing
**Figure 1. Development of a genetically encoded metabolic indicator for lactate (GEM-IL)**

(A) Overview of the lactate indicator GEM-IL binding upon introduction of lactate.

(B) Relative fluorescence change of the cyan, green, and yellow fluorescent protein-based GEM-IL constructs after 10 mM lactate addition in NIH3T3 cells with the superfolder CFP (sfCFP), full-length cyan fluorescent protein (CFP), Laconic, and Green Lindoblum signal changes as points of comparison. Error bars represent standard deviations (n = 23, 16, 12, 18, 29, 26, and 30 samples, respectively). One-way ANOVA with multiple comparison test (Laconic versus cyan, *p = 0.0368; Laconic versus Green Lindoblum, ***p = 2.76 \times 10^{-11}; Green Lindoblum versus cyan, ***p = 0.0008; cyan versus sfCFP, ***p = 1.58 \times 10^{-13}; cyan versus CFP, ***p = 2.25 \times 10^{-11}).

(C) Representative fluorescence change traces of cyan fluorescent protein-based GEM-IL (C-GEM-IL) in NIH3T3 cells.

(D) Construct maps of GEM-IL prototype, C-GEM-IL 1.0, and C-GEM-IL 1.1.

(E) Fluorescence change of G-GEM-IL with various linker types after addition of 10 mM lactate (i.e., [Lactate] = ~545 μM according to [B]) to the imaging solution. Error bars represent standard deviations (n = 12, 8, 9, 10, 7, 9, 6, and 15, respectively). One-way ANOVA with multiple comparison tests (*p < 0.05, **p < 0.01) was used.

(F) Cyan fluorescence and bright-field images of C-GEM-IL full-length and C-GEM-IL ΔDBD in NIH3T3 cells. Arrowheads indicate the puncta seen when overexpressing C-GEM-IL. Scale bar, 10 μm.
strain on the molecule (Bonnot et al., 2014; Ganim and Rief, 2017). As expected through the known excitation and emission spectra of each fluorescent protein, we observed cyan and green fluorescence when expressing C-GEM-IL in mammalian (HEK 293T and NIH3T3) cells, as well as green and yellow fluorescence from G-GEM-IL and Y-GEM-IL (Figure S1A) (Shaner et al., 2003). The ability of GEM-IL to detect a change in lactate levels in situ was then assessed through the exogenous administration of 10 mM sodium lactate. This concentration is physiologically relevant, as circulating levels can range from 0.5 to 2.0 mM and even up to 40 mM after exercise or in tumor micro-environments (Goodwin et al., 2007; Walenta et al., 2000). This exogenous 10 mM administration led to an increase in cellular cytoplasmic levels from 33 ± 12 to 545 ± 149 mM (Figure S1B).

Importantly, this increase in cytoplasmic lactate has no effect on the innate fluorescence of cyan fluorescent protein, suggesting that any fluorescence changes we see moving forward are the result of our lactate indicator (Figures 1B and S1C) (Proia et al., 2016). Looking to now test our indicator’s response to this exogenous lactate addition, we observed that C-GEM-IL had a greater absolute fluorescence change compared with G-GEM-IL and Y-GEM-IL, with signal changes of 11.1% ± 2.5%, 4.6% ± 2.1%, and 10.2% ± 3.1%, respectively. Compared with previously developed indicators, these fluorescence changes were less than those observed using the FRET type, Laconic, but greater than Green Lindobulum (Figures 1B and 1C). We also used a full-length sfCFP molecule with no lactate binding domain as an additional control and found a signal change of only 0.38% ± 2.75%, suggesting minimal non-specific effects of the lactate addition on the fluorescent signal in live mammalian cells (Figures 1B and S1C). In addition, the binding of lactate to C-GEM-IL resulted in an increase in the fluorescent signal, which was the inverse of what is seen with G- and Y-GEM-IL. Although the exact mechanism behind this inversion is unknown, it is consistent with previous studies using csGFP (Bonnot et al., 2014). In parallel with the fluorescent protein comparisons, we tested the use of various linkers—flexible, rigid, and hinge—to better optimize our indicator given the importance of spatial constraints with superfolded fluorescent proteins. Using the green fluorescent type, G-GEM-IL, as the template, which as previously stated originally had a fluorescence change of 4.6%, we observed that adding a hinge linker between LidR and sfGFP11 increased the absolute fluorescence change of G-GEM-IL to 9.4% ± 4.3% (Figures 1D and 1E). We integrated this improvement into our most promising fluorophore, cyan, and added a hinge linker to develop C-GEM-IL 1.0 (Figure 1D).

Furthermore, we found that C-GEM-IL 1.0 formed prominent puncta when expressed in NIH3T3 cells (Figure 1F). This may be evidence of protein aggregation brought on by poor indicator stability and folding. To alleviate this, we deleted the DNA binding domain of the LidR portion of our indicator (ΔDBD), and designated it as C-GEM-IL 1.1 (Figure 1D). The deletion eradicated the previously seen puncta, resulting in a uniform distribution of the indicator in the cytoplasm, and increased the response of this indicator to lactate perfusion to 18.3% ± 3.5%, significantly greater than that of the C-GEM-IL 1.0’s 4.5% change (Figures 1F and S1D).

Fluorophore mutagenesis in C-GEM-IL
Investigating the cause behind the aforementioned inversion of fluorescence changes between C-GEM-IL and Y/G-GEM-IL, we aligned the amino acid sequences of the fluorescent proteins and found that sfCFP alone had an isoleucine at the 146th position compared with an asparagine in the other proteins (Figures 1B, 2A, and 2). Suspecting this amino acid to be the cause of the positive C-GEM-IL response to lactate, we generated an I146N mutant of C-GEM-IL 1.1 (Figure 2B). Interestingly, this did not lead to an inversion of the fluorescence change. Instead, the I146N mutation significantly enhanced the fluorescence change of C-GEM-IL 1.1 in response to lactate to 24.4% ± 10.1% (Figure 2C), leading to an improved version: C-GEM-IL 2.0. This observed change may be due to differences in the folding of mature superfolder fluorescence proteins affecting how I146 acts in relation to the chromophore compared with regular CFP. This finding is noteworthy, as in regular CFP it was previously found that the I146N mutation leads to a decrease in fluorescence (Heim and Tsien, 1996; Kubala et al., 2010; Lelimonis et al., 2009).

To further characterize our lactate indicator, we used a bacterial pCold expression system to obtain and characterize purified recombinant C-GEM-IL 2.0 protein. All recombinant protein experiments were conducted in solutions calibrated by a pH meter and confirmed stable at pH 7 unless otherwise indicated. Characteristic of CFP, we found that when excited at 425 nm, the emission spectrum remained as a single peak with a maximum emission of 490 nm (Figure 2D). The dissociation constant of the indicator (Kd) was found to be 2.33 ± 0.61 mM (Figure 2E). Measuring the response of the in vitro recombinant protein to various metabolites, C-GEM-IL 2.0 showed a specificity for lactate reflected in a 41.1% ± 17.5% fluorescence change, with no significant changes observed in the presence of other metabolites (Figure 2F). Although non-significant, the indicator’s in vitro fluorescence changes in response to glucose, pyruvate, and acetate were at levels of 15.3% ± 4.0%, 16.0% ± 8.9%, and 17.2% ± 6.1%, respectively (Figure 2F), suggesting that the substrate specificity may be a concern with the current iteration of our C-GEM-IL 2.0 construct. C-GEM-IL 2.0 had comparable substrate specificity in comparison to Green Lindobulum and Laconic, which both showed minimal non-specific signaling (Harada et al., 2020; San Martin et al., 2013).

Lactate binding domain characterization and mutagenesis in C-GEM-IL
To improve upon the specificity of our indicator, we sought to identify the conserved regions or amino acids critical for lactate binding. To do so, we aligned the amino acid sequences of the lactate binding domains from three organisms: Escherichia coli, Pseudomonas aeruginosa, and Corynebacterium glutamicum (Figure S3A). From this alignment and computational prediction tools, SWISS-MODEL and I-TASSER, several conserved residues were targeted for mutagenesis to improve binding affinity and further optimize C-GEM-IL 2.0 (Figures 3A, 3B, and 3S). Of the candidate mutants tested, the E103D mutation resulted in the largest increase in fluorescence of 42.9% ± 9.3% in response to lactate (Figure 3B). This improvement from the 24.4% ± 10.1% change observed with C-GEM-IL 2.0 led to the development of C-GEM-IL 3.0 (Figure 3C). In addition, the mutation studies
In NIH3T3 cells, AR-C155858 is an MCT1 inhibitor, which inhibits the influx of lactate into the cell (Figure 3D) (Ovens et al., 2010). Following exposure of NIH3T3 cells to AR-C155858 and observed a similar increase in fluorescence change (Figure 3B). This result suggests that the H151 residue is important for LdrR protein activity, possibly affecting oligomerization and/or lactate binding. For our studies, this construct was useful as a negative control of fluorophore point mutation improving response to lactate and could be used to further characterize our most optimized indicator, the bacterial recombinant protein was purified and imaged using a Coomassie brilliant blue stain evidencing a single band at ~50 kDa representing C-GEM-IL 3.0 (Figures 3H and S3C). The recombinant protein has a single emission peak with a maximum intensity at 480 nm (Figure 3I). To investigate whether C-GEM-IL 3.0 had multiple oligomerization states, size-exclusion chromatography-based high-performance liquid chromatography (HPLC) was conducted using the purified protein, and we found that the recombinant proteins showed a single peak (Figure S3D). Of note, there is a noticeable shoulder peak in the HPLC data that accumulates after multiple freeze/thaw cycles, suggesting possible aggregation of the purified recombinant proteins. In comparison to C-GEM-IL 2.0, whose $K_d$ was 2.33 ± 0.61 mM, C-GEM-IL 3.0 had an improved $K_d$ of 661 ± 19.2% change in response to lactate, with a significant 87.5% ± 4.6% change in response to 2DG (Figures 3F and 3G). Confirming that this response is the result of glycolysis, we perfused glucose after the cells were exposed to AR-C155858 and observed a similar increase in endogenous lactate (Figure 3F). NIH3T3 cells expressing C-GEM-IL 3.0 under these conditions could detect the change in lactate levels presumably produced through glycolysis, with a fluorescence change of 4.0% ± 1.4%, while the fluorescence change in response to 2DG was not different from baseline (Figures 3F and 3G).
Figure 3. Mutation in lactate binding domain improves response and specificity to lactate

(A) Model of the LdR structure with the predicted lactate binding site (purple) and E103 (red).
(B) Effects of targeted amino acid substitutions E103D and H151M in the lactate binding domain on lactate-induced fluorescence of C-GEM-IL 2.0 in NIH3T3 cells. Error bars represent standard deviations (n = 63 and 37). One-way ANOVA using Dunnett’s multiple comparison tests was used with wild-type (WT; ***p < 0.001).
(C) C-GEM-IL 3.0 construct map.
(D) Overview of C-GEM-IL binding upon introduction of lactate and the effect of AR-C155858 (MCT1 inhibitor).
(E) Fluorescence intensity changes of C-GEM-IL 3.0 and its response to 10 mM lactate addition with (black) and without treatment with AR-C155858 (100 nM, blue). Student’s t test was used (***p < 0.001).
(F) Schematic representation of C-GEM-IL lactate sensing upon introduction of glucose after NIH3T3 cells are glucose -starved.
(G) Average fluorescence intensity changes of C-GEM-IL 3.0 and its response to glucose (10 mM) in the absence and presence of 100 nM AR-C155858 (red and blue, respectively) and a non-metabolizable glucose analog, 2DG (10 mM; green). Before glucose or 2DG was added, the cells were kept in glucose-free Tyrode’s solution. Error bars represent standard deviations. One-way ANOVA with multiple comparisons test (***p < 0.001).
(H) Image of SDS-PAGE gel of purified recombinant C-GEM-IL 3.0 proteins (arrow).
(I) Emission spectrum of the purified recombinant proteins of C-GEM-IL 3.0.
(J) Lactate binding curve of the purified recombinant C-GEM-IL 3.0 proteins. The non-linear curve fit for total binding used to calculate the dissociation constant is shown (red).
(K) Substrate binding test of the recombinant C-GEM-IL 3.0 with each substrate at 10 mM. Error bars represent standard deviations (n = 3, using one-way ANOVA with multiple comparisons test, ***p < 0.001; n.s., no other significant differences among the groups).
As previously mentioned, the amount of lactate to which C-GEM-IL 3.0 is exposed in the cells is not the same as the lactate concentration in the external bath solution (Figure S1B). Taking this into account, the internal lactate concentration was measured in non-transfected NIH3T3 cells after exposure to 0–100 mM lactate using a biochemical assay (Figure S3E and S3F). The internal concentrations were then used to determine the in situ response and $K_D$ of C-GEM-IL 3.0. The in situ $K_D$ of C-GEM-IL 3.0 was found to be $515 \pm 167 \mu$M, similar to what was found in vitro using the recombinant proteins (Figures 3J and S3G).

As seen with other fluorescence-based systems, the overall fluorescence intensity of recombinant C-GEM-IL 3.0 proteins was found to shift in relation to the pH of the solution (Figure S3H) (San Martin et al., 2013). To examine whether this feature would affect our indicator’s change in response to lactate, C-GEM-IL 3.0 was exposed to 0 or 10 mM lactate at pH 6.5. At this altered pH, C-GEM-IL fluorescence increased by 51.5% ± 31.7% in response to lactate (Figure S3I). To assess whether the fluorescence change seen in situ in NIH3T3 cells was affected by a change in intracellular pH caused by the lactate infusion, the response of the recombinant form of the indicator was measured after exposure to 30 or 500 µM lactate, levels meant to mimic the increase in cytoplasmic lactate observed with a 10 mM exogenous addition of lactate in situ (Figure S1B). This led to a 37.8% ± 11.8% and 40.7% ± 17.0% increase in the recombinant protein fluorescence purified from bacterial and mammalian cells (HEK 293T), respectively (Figure S3J). In vitro change is consistent with what is observed in NIH3T3 live cells in situ (a 42.9% increase), evidencing that the fluorescence increase can reliably be attributed to lactate binding to C-GEM-IL 3.0. Following a previous study identifying a pH-stable GFP variant (Roberts et al., 2016), a pH-insensitive mutant of C-GEM-IL 3.0, named C-GEM-IL 3.0p (N149Y/Q204H), was developed. In response to 10 mM exogenous lactate addition, this mutation led to a fluorescence change of 21.7% ± 4.2% in situ, a significant decrease compared with C-GEM-IL 3.0 (Figure S3K). Importantly, however, the purified recombinant form of C-GEM-IL 3.0p still showed a specific significant response to lactate with no significant change in the fluorescence of the sensor in response to the other metabolites (Figure S3L).

Comparison of C-GEM-IL 3.0, Laconic, and Green Lindoblum

The aim of this study is to develop an improved lactate indicator to enable more researchers to better measure lactate changes. The previously developed genetically encoded lactate sensors, Laconic and Green Lindoblum, have been utilized in various studies but have certain characteristics that may limit their use. C-GEM-IL 3.0 improves upon these indicators in several significant ways. The first is that C-GEM-IL 3.0 does not have low signal expression or form puncta when genetically overexpressed in NIH3T3 cells, which was observed with Laconic, Green Lindoblum, and C-GEM-IL 1.0 (Figures 1F and 4A). The next is that C-GEM-IL 3.0 has significantly improved response to lactate compared with Green Lindoblum (42.9% versus 6.2%, respectively, Figures 1B and 3E). Furthermore, C-GEM-IL 2.0 and 3.0 are capable of measuring the production of lactate generated via glycolysis, while Laconic and Green Lindoblum could not (responses of 4.5 ± 1.2, 12.0 ± 4.1, −1.1 ± 3.8, and 1.8 ± 1.2%, respectively, Figure 4B). In looking more at the protein expression of the different indicators, the C-GEM-IL series shows a single band on Western blotting compared with the dual unidentified bands seen with Laconic, suggesting that the Laconic construct may be degraded/truncated and/or cleaved, and the fluorescence change of Laconic might not be reliable (Figure 4C). Lastly, in an attempt to simply improve upon the currently available indicators, we looked to see whether any of the alterations conducted in C-GEM-IL development would be applicable for Laconic, and prepared E103D, ΔDBD/E103D, and H151D versions of the Laconic indicator. None of these changes improved the fluorescence response observed, and each of these changes led to a significantly less responsive form for the sensor, with FRET changes of 6.5% ± 1.1%, 6.2% ± 0.7%, and 5.5% ± 2.0% in response to lactate in situ, respectively (Figure 4D). In addition, in comparing the non-functional “dead” version of our indicator, H151M C-GEM-IL, to H151D Laconic, we found that H151M C-GEM-IL had a signal closer to baseline, with observed responses of 2.2% and 5.5%, respectively (Figures 3B and 4D). Overall, C-GEM-IL 3.0 improves on the currently available indicators, and offers a more precise and reliable method for measuring lactate changes in live cells compared with Laconic and Green Lindoblum (Figure 4D; Box 1C).

Ratiometric versions of C-GEM-IL 3.0

Although C-GEM-IL 3.0 is useful in the detection of lactate as a single-fluorophore-based indicator (Figure S4A), it does not allow for the exact quantification of lactate due to factors extrinsic to the stimulant (Grynkwicz et al., 1985; Paredes et al., 2008). Adding a ratiometric version of GEM-IL improves sensitivity through an internal reference and allows for quantitative analysis, should users of this tool prefer such capabilities or have experimental setups that support this version (Lee et al., 2015). Therefore, we developed a ratiometric version of C-GEM-IL 3.0 by adding a secondary fluorescent molecule (either YFP or red fluorescent protein, RFP) with the fluorescent indicator after a linker (Figure S4B). Interrogating the expression of appending YFP to C-GEM-IL 3.0 through Western blotting showed a single band at 75 kDa (Figures S4B–S4D). Linking a YFP to C-GEM-IL 3.0 through either an Ala-Ser (AS) or a rigid linker rendered the mature CFP unresponsive to lactate, while the YFP signal decreased, possibly a FRET change of 29.7% ± 5.7% and 16.0% ± 4.4%, respectively (Figures S4E and S4F). Examining alternative fluorophores to append, we used a rigid linker to add an RFP, mKate2 or mCherry, to the carboxyl terminus of C-GEM-IL 3.0. Expression of these constructs led to two bands between 50 and 75 kDa on western blotting due to an unknown degradation/truncation mechanism as observed in Laconic (Figures 4C, S4C, and S4D). This possible degradation/truncation suggests a potentially unstable biosensor, so as an alternative approach to address this concern, we applied a self-cleaving peptide P2A sequence, generating C-GEM-IL 3.0r (Figure S4G) (Kawano et al., 2016; Kim et al., 2011). The P2A...
cleavage was not fully efficient, with bands at 75 kDa (non-cleaved, ~10%) and 50 kDa (cleaved, ~90%) (Figure S4H). Of note, there was a band evident between 50 and 60 kDa potentially due to the truncation of the non-cleaved proteins. NIH3T3 cells transfected with C-GEM-IL 3.0 responded to lactate exposure in situ, with the cyan signal changing in reference to the mKate2 fluorophore. We observed the fluorescence change of this indicator at increasing levels of lactate to generate a linear-regression-based model for lactate quantitation (Figure S4I). Using this model, we calculated that the observed change in fluorescent signal, which had a ratiometric response of 30.0% ± 6.5%, could be equilibrated to an internal lactate increase of 622 μM (Figure S4J). Also, the mKate2 signal remained constant during lactate addition and binding, again evidencing the resistance of fluorescent molecules to cellular changes due to lactate influx into the cells, as also observed in CFP and sCFP (Figures 1B and S1C). This version of our indicator can be valuable in future applications to determine the amount of lactate generated from other metabolites and/or biological processes.

**Applications of C-GEM-IL for lactate imaging with octyl-R-2HG and c-Myc**

We then sought to examine the utility of our most optimized indicator, C-GEM-IL 3.0, in observing lactate changes in other applications and settings. Octyl-R-2HG, a membrane-permeative precursor of D-2-hydroxyglutarate, is an onco-metabolite that inhibits ATP synthase and mTOR signaling (Fu et al., 2015). Increasing levels of D-2-hydroxyglutarate were found to increase the levels of lactate (Bottcher et al., 2018). To determine whether C-GEM-IL 3.0 could detect a change in lactate levels in response to this substrate, NIH3T3 cells were treated with octyl-R-2HG. It was found that addition of octyl-R-2HG led to a detectable fluorescence increase of C-GEM-IL 3.0 by lactate of 27.9% ± 8.0% in these cells (Figure 5A), demonstrating the indicator’s ability to respond to changes in cellular metabolism elicited by chemical stimulants. Laconic and Green Lindoblum, on the other hand, were unable to detect this change in metabolism, with fluorescence changes of 2.8% ± 2.0% and 5.0% ± 3.0%, respectively (Figure 5A).
Box 1. Summary of GEM-IL development and comparison of Laconic, Green Lindoblum, and GEM-IL lactate indicators

(A) Construct maps of each GEM-IL iteration, detailing the changes made in each and in which particular domains the modifications were made.

(B) Flow chart highlighting the major modification in each construct. Mouse model is available for C-GEM-IL 3.

(C) Comparison table of Laconic, Green Lindoblum, and GEM-IL indicators.

|                          | Laconic | Green Lindoblum | GEM-IL |
|--------------------------|---------|-----------------|--------|
| Indicator Size           | 83.8 kDa| 60 kDa          | 48.5 kDa|
| Available Fluophores     | FRET (mTFP/Venus) | GFP | various fluorescent protein (sfCFP, sfGFP & sfYFP) |
| Response to 10 mM Lactate (in vitro, NIH 3T3 cells) | ~15% | ~6% | ~40% |
| Substrate Specificity    | O       | O               | O      |
| Detects Metabolic Activity (glucose, drug metabolism & cancer metabolism) | - | - | O |
| in vivo Mouse Model Availability | - | - | O |
| Expression Pattern in Live Mammalian Cells (ex. NIH 3T3) | Aggregated | Aggregated & low expression | Uniform in the cytoplasm |
| Kd                       | 8 µM / 830 µM (San Martin, 2013) | 3mM (Harada, 2020) | 661 ± 142µM |
| pH-insensitive Construct Availability | - | - | O |
| Quantification Reliability | High (FRET) | Medium (single fluorescent) | Medium (single fluorescent) |
| Dual Imaging Capability  | -       | O               | O      |

(A) Construct maps of each GEM-IL iteration, detailing the changes made in each and in which particular domains the modifications were made.

(B) Flow chart highlighting the major modification in each construct. Mouse model is available for C-GEM-IL 3.

(C) Comparison table of Laconic, Green Lindoblum, and GEM-IL indicators.
Overexpression of c-Myc, an oncogene, has been shown to alter the metabolic state of cells with an increased need to support cellular proliferation accompanied by an increase in lactate production (Miller et al., 2012). To assess if our indicator can detect this shift in cellular profiles, we virally overexpressed c-Myc in NIH3T3 cells and human neuronal progenitors derived from induced pluripotent stem cells (iPSCs) as a model of the central nervous system primitive neuro-ectodermal tumor (CNS-PNET) (Behdad and Perry, 2010). In this model, c-Myc overexpression results in a heightened level of lactate production in the cellular environment. Here we found that the addition of lactate to these c-Myc-infected oncogenic cells resulted in a smaller fluorescence change compared to non-infected cells, showing that the indicator can distinguish between the two metabolic states (Figures S5B–S5E). The lower response seen in the oncogenic cells could be due to the heightened metabolic state induced by increased c-Myc expression, resulting in significantly higher endogenous lactate concentrations compared with non-infected controls, which was supported by a lactate biochemical assay of the cell lysate and culture medium (Figures S5A–S5C). The higher basal lactate levels would result in a higher baseline fluorescence of our indicator, so any further lactate addition would not result in as large of an increase in fluorescence.

**Simultaneous imaging of lactate and calcium ion**

To further demonstrate the possibility of simultaneous fluorescence imaging using C-GEM-IL 3.0, we conducted live cell imaging for lactate and Ca²⁺ using C-GEM-IL 3.0 and R-GECO1, a genetically encoded red fluorescent Ca²⁺ indicator (Zhao et al., 2011). We found that the cytoplasmic Ca²⁺ levels significantly decreased with the increase in lactate in the cells (Figures S5D and S5E). This finding further evidences that GEM-IL can be used in tandem with other indicators, even those that monitor metabolites from traditionally unassociated metabolic pathways. This will allow for the examination of the effects of various chemicals and metabolites and how they relate to varying levels in lactate.

**Generation of C-GEM-IL mouse lines**

Rather critically, the measurement of lactate in organs and tissue relies on indirect measurements, including the use of electrodes measuring extracellular lactate or radioisotope-labeled metabolites coupled with mass spectrometry, effectively...
allowing for only snapshots of lactate handling (Marzouk et al., 2002; Sugiura et al., 2016). An improved in vivo model would be invaluable to study lactate dynamics in healthy and disease models. In cardiac physiology, for example, an increase in the serum level of lactate correlates negatively with patient survival after heart failure (Hattori et al., 1985; Kawase et al., 2015). To enable the monitoring of lactate levels during normal physiology and disease, in vivo mouse models expressing either C-GEM-IL 2.0 or C-GEM-IL 3.0 were generated by inserting either version of the indicator into the Rosa26 locus (Rosa26-Cag-lox-stop-lox-C-GEM-IL, Figures 5F and S5F). The C-GEM-IL mouse models were used to induce the expression of either indicator in cardiac tissue by crossing the C-GEM-IL 2.0 and 3.0 mouse lines with either a tamoxifen-inducible α-myosin heavy chain promoter-driven Cre-expression mouse (αMHC/Myh6-MerCreMer/CreERT2), which restricts the indicator expression to the heart in response to tamoxifen, or a non-inducible-type, αMHC-Cre, mouse, respectively (Sohal et al., 2001).

The expression was confirmed through PCR showing Cre-loxP recombination specifically in cardiac tissue, and the expression of C-GEM-IL 2.0 and 3.0 was further confirmed by Western blotting; furthermore, Cre transient expression could also induce C-GEM-IL 3.0 protein expression in other tissues, such as liver (Figures S5F–S5K). The overexpression of the indicator did not deleteriously affect the architecture of the mouse hearts as evidenced by the echocardiograms (Figure S5L). To confirm the functionality, we isolated adult primary cardiomyocytes from C-GEM-IL 2.0 and 3.0 mice and observed the changes in fluorescence after lactate exposure. C-GEM-IL 3.0 cardiomyocytes had a significantly higher response of 68.8% ± 4.3% compared with the 12.9% ± 1.4% change observed with the 2.0 version, confirming the successful generation and validation of our mouse lines (Figures 5G, 5H, and S5M).

DISCUSSION

This work describes the development and applications of C-GEM-IL 3.0, a single superfolder fluorescent protein-based indicator with a mutation, E103D, which improves upon the previously available lactate indicators, Laconic and Green Lindoblum. The novel design of GEM-IL series with varying kinetics and expression to targeted subcellular organelles such as the nucleus, plasma membrane, mitochondria, and endoplasmic reticulum. To this point, the pH-stable and ratiometric versions, C-GEM-IL 3.0p and 3.0r, can be useful to increase potential applications of fluorescent lactate imaging. Unfortunately, the pH-insensitive version, 3.0p, had significantly lower fluorescent response to lactate. The two point mutations (N149Y and Q204H) used to generate a pH-insensitive version appear to have also decreased the sensor’s overall sensitivity to lactate. This is supported by the recombinant protein data, which were collected at a stable pH 7 and showed that 3.0p still has a significant sensitivity to lactate but lower than that of the original C-GEM-IL 3.0. Control perfusion experiments using just the fluorophores to examine whether lactate addition can change the fluorescent signals in live mammalian cells showed that neither CFP nor sfCFP had any significant/obvious changes in signal when 10 mM lactate was added to the imaging solution (Figures 1B and S1C). While fluorescent proteins are known to be pH sensitive, the lactate perfusion to live cells does not affect the fluorescent signal of CFP and sfCFP in the live cells. A benefit of developing a single fluorescence indicator is the ability to use multiple fluorescent sensors at the same time to measure different probes and reporters. As evidence of this, C-GEM-IL 3.0 was successfully used in tandem with R-GECO1, a Ca2+ red fluorescent indicator, showing a negative correlation between lactate administration and the R-GECO1 Ca2+ signal. The molecular mechanism underlying the effects of lactate on intracellular Ca2+ handling remains unclear. Investigating the association of lactate with Ca2+ handling is an interesting direction as a future study.

Employing C-GEM-IL 3.0 allowed us to detect the difference in lactate response of metabolically disparate cells driven by c-Myc overexpression in NIH3T3 cells and human iPSC-derived neural progenitor cells. Octyl-R-2HG led to a detectable increase in lactate in live cells, revealing the ability of C-GEM-IL 3.0 to respond to changes in cellular metabolism induced by such onco-metabolites. The other currently available indicators, Laconic and Green Lindoblum, were unable to achieve such results. Our GEM-IL can be beneficial in chemical biology study to determine the metabolic effects of various small molecules on the levels of lactate as well as being used as a screening tool to assess the utility of these compounds in biomedical research.

To enable the investigation of lactate levels in vivo, we developed genetically encoded mouse strains that contain a Cre-inducible cassette for the expression of either C-GEM-IL...
2.0 or C-GEM-IL 3.0 by inserting the sequences into the Rosa 26 locus. When crossed with cardiac Cre drivers such as αMHC-Cre, isolated adult primary cardiomyocytes responded to lactate perfusion, demonstrating that the indicator was expressed and functioned with C-GEM-IL 3.0, showing a significantly improved response over C-GEM-IL 2.0.

Overall, GEM-IL is a powerful tool set that allows us to obtain new insights into the roles of lactate as an essential metabolite in live cells with single-cell resolution. As this is the initial set of iterations on the indicator, future work should focus on refining the kinetics and subcellular and extracellular expression of the GEM-IL series. We hope that further development of genetically encoded fluorescent indicators, similar to C-GEM-IL, for other metabolites will be useful in biomedical applications that probe metabolic changes in both normal and disease states with single-cell resolution to better understand the importance of various metabolites in physiology and disease.

**Limitations of the study**

Our tool, GEM-IL, provides an additional method for monitoring lactate influx and efflux in cells. Capitalizing on the compact nature and innate characteristics of superfolder fluorescent proteins, we were able to generate a tool with stable expression, high specificity, better sensitivity, and wider-spread applicability compatible with other lactate indicators. In addition, we achieved and demonstrated in vivo expression and functionality in a cardiac-specific murine model. Given that this is an indicator based on a fluorescent protein, we recognize some of the limitations that accompany this feedback system, such as pH sensitivity, photobleaching, and quantitative analysis. We sought to address these issues through different iterations of GEM-IL: C-GEM-IL3.0p and C-GEM-IL3.0r. In addition, we recognize that our indicator requires an experimental setup capable of monitoring changes in cyan fluorescence. Although we believe this to be a reasonable requirement, our indicator can also be visible with green and yellow fluorescent protein filters, although the signal will not be as strong due to the close spectral properties of these proteins. Last, we acknowledge that in this study we focused on proof of principle and benchmark experiments to validate our new lactate indicators, not on biological applications, as a method report. However, the plasmid DNA, viral construct, and mouse line of GEM-IL will be widely applicable for various researchers as a new resource to investigate lactate metabolism and provide new insights into the role of lactate.

**STAR+METHODS**

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**QUANTIFICATION AND STATISTICAL ANALYSIS**

**SUPPLEMENTAL INFORMATION**

Supplemental information can be found online at https://doi.org/10.1016/j.crmeth.2021.100092.

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**AUTHOR CONTRIBUTIONS**

M.Y. conceived of and designed this project. R.B., J.R.Q., S.U., F.K., K.M., A.D.K., T.C.L., K.N., T.M.L., R.L., M.V.Y., and M.Y. designed and/or performed the experiments. S.U., F.K., and M.Y. designed constructs and performed the mutagenesis to improve the indicator constructs. C-S.L. and M.Y. conducted mouse embryonic stem cell targeting and the mouse generation. R.B., J.R.Q., S.U., F.K., K.M., K.K., K.N., T.L., A.C., and M.Y. analyzed the data. R.B., J.R.Q., and M.Y. interpreted the data and wrote the manuscript. F.K., S.H., A.L.S., and K.G. helped M.Y. supervise this interdisciplinary team and proofread the manuscript. All co-authors approved the manuscript.

**DECLARATION OF INTERESTS**

The authors declare no competing interests.

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**STAR★METHODS**

**KEY RESOURCES TABLE**

| REAGENT OR RESOURCE | SOURCE | IDENTIFIER |
|---------------------|--------|------------|
| Antibodies          |        |            |
| β-tubulin           | Sigma-Aldrich | #5201 ; RRID:AB_609915 |
| GFP                 | MBL    | #598 ; RRID:AB_591819 |
| anti-mouse secondary| Thermo-Fisher | #31430 ; RRID:AB_228307 |
| anti-rabbit secondary| Thermo-Fisher | #31460 ; RRID:AB_228341 |
| Bacterial and virus strains |        |            |
| pCMV-dR7.2 dvpr     | Addgene | #8455 |
| pCMV-VSV-G          | Addgene | #8454 |
| LV-Cre-SD           | Addgene | #12105 (no longer available) |
| pcDNA3.1            | Life Technologies | cat no. V79020 |
| pCold II            | TaKaRa Bio | #3362 |
| A38                 | Addgene | #34883 |
| Rosetta DE3 Competent Cells | Novagen | #70956 |
| XL10-Gold Cells     | Agilent | #200315 |
| Chemicals, buffer reagents, and recombinant proteins |        |            |
| Tamoxifen           | Sigma-Aldrich | T5648 |
| Lenti-X concentrator| TaKaRa Bio | #631232 |
| poly-ornithine      | Sigma-Aldrich | #P0421 |
| Lipofectamine 2000  | Life Technologies | #11668027 |
| Lipofectamine 3000  | Life Technologies | #L3000008 |
| OptiMEM             | Thermo-Fisher | #31985070 |
| Ni-sepharose Beads  | GE Healthcare | #17-5318-01 |
| IPTG                | Sigma-Aldrich | #6758 |
| Cell Lysis Buffer   | Cell Signaling Technologies | #9803 |
| Pierce ECL western blotting substrate | Thermo-Fisher | #32209 |
| Superblock          | Thermo-Fisher | #37515 |
| Critical Commercial Assays |        |            |
| Lactate Assay Kit   | Sigma Aldrich | #MAK064 |
| Experimental Models: Cell Lines |        |            |
| Human: HEK 293T     | ATCC   | #CRL-3216 |
| Human: NIH 3T3      | ATCC   | #CRL-1658 |
| Experimental Models: Mouse Lines |        |            |
| αMHC-MerCreMer      | JAX    | #005657 |
| αMHC-Cre            | JAX    | #011038 |
| Oligonucleotides    |        |            |
| Oligonucleotides designed in this study | Full sequence maps available upon request | N/A |
| Software and algorithms |        |            |
| Vevo770             | VisualSonics | N/A |
| e33 Ultrasound      | Philips | N/A |
| CueMol2             | N/A    | http://www.cuemol.org/ |
| I-TASSER            | Zhang Lab | https://zhanglab.ccb.med.umich.edu/I-TASSER/ |
| MetaFluor           | Molecular Devices | Version 7.8.1.0 |
| Prism v9.0          | GraphPad | https://www.graphpad.com/ |
RESOURCE AVAILABILITY

Lead contact
Further information and requests for resources and reagents should be directed to and will be fulfilled by the Lead Contact Dr. Masayuki Yazawa (my2387@columbia.edu)

Materials availability
All materials and constructs used in this study are maintained by Dr. Yazawa’s laboratory and are available upon request. In the future, the constructs and mouse line will be deposited and available to order through Addgene and JAX, respectively.

Data and code availability
- All data reported in this paper will be shared by the lead contact upon request.
- This paper does not report original code.
- Any additional information required to reanalyze the data reported in this paper is available from the lead contact upon request.

EXPERIMENTAL MODEL AND SUBJECT DETAILS

Mice
This study was carried out in strict accordance with the guidelines of the National Institutes of Health and the Institutional Animal Care and Use Committee at Columbia University, New York. The protocol was approved by the IACUC at Columbia University (protocol #AC-AAAU2453/AABK3550). All mice are housed under a 12 hour light/dark cycle and provided with food and water ad libitum. The synthesized C-GEM-IL 2.0/3.0 DNA fragment (Integrated DNA Technologies) was inserted into the Ai38 targeting construct (Addgene, #34883) utilizing the FseI restriction site (NEB, R0588S). The construct was then linearized using PvuI-HF digestion enzyme (NEB, R3150S). The linearized construct was electroporated into home-made mouse embryonic stem cells (C-S. L generated) using a standard protocol. The germ-line transmitted C-GEM-IL 2.0/3.0 mice were crossed with the αMHC-MerCreMer line (JAX #005657) or a MHC-Cre line (JAX #011038). In case of αMHC-MerCreMer use, the subsequent progeny was injected intraperitoneally with 50mg/kg body weight of tamoxifen (Sigma-Aldrich, dissolved in sunflower oil, 1 dose/day, 3 days) to induce the cardiac specific expression of C-GEM-IL. Mouse primary adult ventricular myocytes were isolated from C-GEM-IL 2.0/3.0 mice using our established protocol. These cells were then calcium ion-adapted and imaged as described in the below perfusion section. Genomic DNA was extracted from the cardiac and tail samples from C-GEM-IL 2.0 and 3.0 positive mice using standard phenol/chloroform-ethanol precipitation method. The target sequence was then amplified evidencing the cardiac-specific Cre-lox recombination utilizing Ex Taq DNA polymerase (TaKaRa/Clontech, RR0018).

Cell culture
Human embryonic kidney (HEK) 293T cells (ATCC, #CRL-3216) and NIH 3T3 cells (ATCC, #CRL-1658) were cultured in Dulbecco’s Modified Eagle Media (DMEM, Thermo-Fisher/Gibco #10313021) supplemented with GlutaMax-I and penicillin, streptomycin (PS) and 10% fetal bovine serum (FBS, not heat-inactivated, HyClone, SH30071.03, Thermo-Fisher) under normoxia (20% O2, 5% CO2, at 37°C).

Human normal induced pluripotent stem cells (iPSCs) were cultured using a standard feeder-free protocol with Essential 8 medium and Geltrex (Thermo-Fisher/Gibco #A1517001) following the manufacturer’s instructions. Human iPSC lines were differentiated using Neural Induction Media (NIM, Thermo-Fisher/Gibco #A1647801) to generate neural progenitor cells. The progenitors were cultured using NIM supplement, Neurobasal and advanced DMEM media (Thermo-Fisher/Gibco) following the NIM manufacturer’s instruction.

METHOD DETAILS

Lactate biochemical assay
The lactate content of the cells and the media was measured using a lactate assay kit (Sigma-Aldrich/Millipore-Sigma, #MAK064) following the manufacturer’s instructions. In brief, the cells were infected with the c-Myc virus for 24 hours. The cell media was then replaced by phenol red-free DMEM (Thermo-Fisher/Gibco, #31053028) supplemented with GlutaMax-I, PS and 10% FBS. Twenty-four hours later, the media and cells were harvested and filtered through a 10kDa filter (Amicon, UFC801024). The samples were then plated on a flat bottom 96-well plate (Falcon, #353072) and read using a plate reader Spectramax ID3 (Molecular Devices).

Plasmid DNA construction
Plasmid DNA constructs were generated using standard methods using restriction enzymes (New England BioLabs), DNA ligase (MightMix, TaKaRa) and polymerase chain reaction (PCR) with Phusion polymerase chain reaction (Thermo-Fisher, 2× master
solution three times (normal Tyrode’s solution: 1.8 mM CaCl₂, 1 mM MgCl₂, 140 mM NaCl, 5.4 mM KCl, 10 mM glucose, and 10 mM

Lentiviral production and infection of NPC and NIH 3T3 cells
Lentiviral production and infection were conducted using standard method. Briefly, HEK 293T cells were transfected with lentiviral human c-Myc vectors together with pCMV-dR7.2 dvpr and pCMV-VSV-G (Addgene #8455 & 8454). Lentivirus c-Myc plasmid was generated using standard PCR and LV-SD with EcoRI and Xhol cloning sites. The lentivirus was then concentrated using a Lenti-X concentrator (TaKaRa/Clontech, #631232) following manufacturer’s instructions to infect either neural progenitor cells or NIH 3T3 cells.

Plasmid DNA transfection
Plasmid transfection of HEK 293T and NIH 3T3 cells was conducted using standard lipofection method. Briefly, HEK 293T and NIH 3T3 cells were plated at 0.5 x 10⁵ cells/well in a 24-well plate format on poly-ornithine coated (Sigma-Aldrich/Millipore-Sigma, #P0421) glass cover slips (Warner instruments). HEK 293T cells were transfected with 0.9 μg of DNA in 2 μL of Lipofectamine 2000 (Invitrogen/Life Technologies #11668027) into 100 μL of OptiMEM (Thermo-Fisher/Gibco #31985070) in 400 μL of DMEM 10% FBS (PS-free). NIH 3T3 cells were transfected with 0.5 μg DNA, 1 μL P3000 and 1.5 μL Lipofectamine 3000 (Invitrogen/Thermo-Fisher #L3000008) into 50 μL OptiMEM in 400 μL of DMEM 10% FBS (PS-free).

Purification and characterization of recombinant protein
Recombinant forms of the indicator protein were purified through batch nickel purification taking advantage of the 6xHis tag (GE Healthcare #17-5318-01) in the pCold_2 vector. Briefly, Rosetta (DE3)pLysS Competent cells (Novagen #70956) were transfected with the pCold II vector with the respective inserts and grown at 37°C overnight. These primary cultures were then used for larger scale production at a 1:100 inoculation. The larger cultures were grown for 4-6 hours at 37°C (or until the O.D. was between 0.4-0.8). The cultures were then plated in ice for 1 hour, followed by addition of 0.5 mM IPTG. These cold-shocked cultures were then placed in a 16°C shaker for 24 hours at 250 rpm. Samples were then spun down and resuspended in lysis buffer (20 mM Na₂PO₄, 500 mM NaCl, 20 mM Imidazole, 1 mM EDTA, 0.1% Triton X-100, 150 mM sucrose) followed by sonication (Branson sonifier 450). The lysed samples were then then added to the beads, washed, and then eluted (elution buffer: 20 mM Na₂PO₄, 500 mM NaCl, 500 mM Imidazole, 1 mM EDTA, 150 mM sucrose). The protein was then concentrated using centrifugal filters with a 10,000 Da (Amicon Ultra-15 UFC901024) cutoff and resuspended in cytoplasmic buffer (pH 7.0, 10 mM NaCl, 130 mM KCl, 1.25 mM MgSO₄, 10 mM HEPES). HEK 293T cells were transfected with C-GEM-IL 3.0 as described above. Forty-eight hours after transfection, the cells were lysed with cell lysis buffer (CST, #9803). The protein was then isolated with nickel beads similarly to that from bacteria. The purified recombinant protein (100 nM) was added to a flat black bottom 96-well plate (Costar 3916) in the respective solutions. The plates were then read in Spectramax ID3.

Fluorescent HPLC
Autosampler SIL-20ACHT was used to inject 100-μL protein samples into the Superose 6 10/30 gel filtration chromatography column connected to the Shimadzu HPLC. The samples were monitored using the RF-20A spectrofluorometric detector at the excitation wavelength of 435 nm and the emission wavelength of 485 nm.

Cellular perfusion experiments
Cultured cells were plated at 0.5 x 10⁵ cells/well in a 24-well plate format on poly-ornithine coated glass cover slips. Cells were transfected as described above. For dual imaging experiments, cells were transfected with C-GEM-IL 3.0 and R-GECO1 simultaneously using the same transfection protocol. 24-48 hours following transfection cover slips were washed three times with normal Tyrode’s solution three times (normal Tyrode’s solution: 1.8 mM CaCl₂, 1 mM MgCl₂, 140 mM NaCl, 5.4 mM KCl, 10 mM glucose, and 10 mM HEPES (pH 7.4 adjusted with NaOH at 25°C). After wash, the coverslips were picked and placed in the imaging chamber (Warner Instruments/Harvard Apparatus, RC-20H, #64-0223). Cells were imaged and fluorescence was traced throughout the perfusions using an inverted Nikon Eclipse Ti microscope with a Nikon 20X objective and MetaFluor Software Version 7.8.1.0 (Molecular Devices). Baseline fluorescence in Tyrode’s solution was gathered for 300 seconds, followed by a 2 mL perfusion of the compound of interest dissolved in Tyrode’s solution with recording for 300 seconds, followed by a 2 mL perfusion of Tyrode’s solution as a wash. Changes in fluorescent data were quantified and analyzed relative to baseline in Excel.

Kinetic measurement
NIH 3T3 cells were plated on coverslips, transfected with C-GEM-IL 3.0, and placed in the imaging chamber set up as described above. Cells were again imaged using MetaFluor Software Version 7.8.1.0. A 150-second baseline was recorded in Tyrode’s solution, followed by 2 mL perfusions of increasing doses (1, 5, 10, 20, and 50 mM) of sodium lactate dissolved in Tyrode’s solution each recorded for 150 seconds. No pH changes were observed after adding 10-20 mM sodium lactate into the Tyrode’s solution. Changes in fluorescent data were quantified and analyzed relative to baseline in Excel.
Multi-filter fluorescent imaging
Cells were plated 1.5 x10^5 cells/dish in 35mm glass bottom polyornithine coated dishes. The cells were transfected as previously described. 24-48 hours post-transfection cells were washed three times with Tyrode’s solution and then imaged using MetaMorph Imaging Software. A Sutter Lambda DG-4 Illumination System was used with standard BFP, CFP, GFP, and YFP filters (Chroma).

Hydrodynamic tail vein injection
5-6 week-old C-GEM-IL 3.0 positive mice were injected with PAC-Cre plasmid using TransIT®-EE Delivery Solution (Mirus Bio LLC) respectively according to the manufacturer’s protocol. The amount of injected plasmid DNA was 10μg per mouse. The volume of delivery solution was 0.1mL per mouse weight (g). After the hydrodynamic injection of DNA, the mice were allowed to recover for 48 hours. The mice were then sacrificed using standard procedure approved in our animal protocol and their livers were immediately harvested for protein expression characterization using Western blotting.

Echocardiographic measurements
Echocardiography was conducted using standard method. Briefly, C-GEM-IL mice were placed under light anesthesia (2% isoflurane) and imaged using Vevo 770 with a 40-MHz transducer (for C-GEM-IL 2.0) or iE33 with a 15-MHz linear transducer (for C-GEM-IL 3.0). M-mode images and two-dimensional parasternal short-axis images at the mid-papillary level were recorded in each mouse.

Homology modelling
The LldR lactate binding domain was determined using known templates in the SWISS-MODEL (Biozentrum) that produced a high identity with the transcriptional factor CGL2914 from Corynebacterium glutamicum (2DI3a). This was used to generate a homology model that was visualized using CueMol2.

Western blotting
Western blot experiments were conducted using standard method. In brief, cardiac/liver samples harvested from C-GEM-IL 2.0/3.0 positive and wild-type mice were homogenized and lysed in 10x cell lysis buffer (Cell Signaling Technology, #9803) with 1% protease inhibitor cocktail (Sigma-Aldrich/Millipore-Sigma). SDS-polyacrylamide gel electrophoresis was performed using Tris-Glycine-based gels (Bio-Rad) containing 10% Acrylamide-Bis (Fisher Scientific) which was then transferred to polyvinylidene difluoride (PVDF) membranes. Primary bodies to GFP (MBL#598, 1/4,000 dilution) and β-tubulin (Sigma-Aldrich, #t5201, 1/4000 dilution). Secondary antibody α-mouse (Invitrogen, #31430, 1/8,000 dilution) and α-rabbit (Thermo Scientific, #31430, 1/8,000 dilution). SuperBlock blocking buffer (PBS based, Thermo Scientific #37515) was used for blocking and antibody incubations of the PVDF membranes. Pierce ECL western blotting substrate (Thermo Scientific, #32209) was used for the chemiluminescent reaction.

QUANTIFICATION AND STATISTICAL ANALYSIS
Quantification of fluorescent intensity was done using MetaMorph software (Molecular Devices). Cellular regions were manually defined and quantified during a defined imaging study. Two-tailed Student’s t-test was used when comparing two separate groups of observations. Welch’s correction was used when the variance between the two groups was statistically different. Two-tailed paired Student’s t-test was used when comparing data sets conducted on samples before and after treatment. One-way ANOVA with Bonferroni or Sidak’s comparison corrections was used for comparison between multiple sets of observations. One-way ANOVA with Dunnet’s correction was used for multiple comparisons against a specified data set. GraphPad Prism non-linear regression model of total site binding was used to generate dissociation constants. All error bars represented standard deviation. p values < 0.05 were considered significant. Unless otherwise stated, all experiments were conducted in at least three independent experiments.