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bioluminescence characteristics

Red-shifted D-luciferin analogues and their

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D-Luciferin (D-LH₂) is the most used substrate for beetle luciferases in various bioluminescence applications. Here, we successfully synthesized six D-LH2 analogues including 5',7'-dimethoxy-D-LH2 and 7'-methylnaphthol-p-LH2 as novel compounds. We also developed a continuous one-pot green synthesis method to improve yields of luciferins from condensation of quinone and D-Cys (63-fold greater than the previous report). The novel p-LH2 analogues were tested with five luciferases (Fluc, SLR, Eluc, Pmluc-WT, and Pmluc-N230S), and all the compounds emitted bioluminescence at wavelengths longer than that of $D-LH_2$ (>80 nm). The reaction of SLR with 5',7'-dimethoxy- $D-LH_2$ gave the longest red-shifted bioluminescence at 663 nm. Remarkably, the reactions of 5'-methyl-p-LH2 emit longer wavelengths and brighter light than those of D-LH2 in all tested luciferases, except for Eluc. Interestingly, the novel red-shifted 5',7'-dimethyl-p-LH2 also provided prolonged bioluminescence with a rate of light decay slower than that of D-LH2. We further demonstrated applications of 5'-methyl-D-LH2 and 5',7'dimethyl-p-LH2 in mammalian cell lines expressing Fluc, SLR, and Pmluc-N230S. 5'-Methyl-p-LH2 provided about 11.2-fold greater sensitivity to detect Fluc in the HEK293T crude lysate than D-LH2, achieving the detection with a lower number of cell lines. The red-shifted 5',7'-dimethyl-p-LH2 also exhibits high sensitivity when using a red light filter to monitor live cell bioluminescence. These $D-LH_2$ analogues, 5'-methyl-p-LH2 and 5',7'-dimethyl-p-LH2, are promising substrates for future cell-based assays and real-time monitoring applications.

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Introduction

Reactions of luciferases and luciferins found in various organisms can generate cold light emission or bioluminescence (BL). These reactions are efficient analytical and visualization tools which have been applied in various detection technologies.1 It has been estimated that nearly 30% of high-throughput screening (HTP) reactions used in academia and industry involve luciferase reactions.2 Beetle luciferases, especially firefly luciferases (Fluc), are among the most widely used BL systems in biochemical and cell-based assays because of their catalytic oxidation of the D-luciferin (D-LH2) substrate to oxyluciferin generates green-yellow light (~560 nm) with a high quantum yield (41%).^{2,3} Although the Fluc reaction is useful for a variety of assays, its application for in vivo imaging is limited because BL signals generated by the natural D-LH2 emits greenyellow light which does not penetrate well into animal cells or tissues. The current use of D-LH2 in animal experiments requires the usage of high doses, thus resulting in high experimental cost.4,5 D-LH2 analogues with red-shift and steady

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BL are thus valuable for improving Fluc applications because this would allow better light penetration through the cells, blood and tissues of animal models.⁶

Numerous D-LH₂ analogues particularly in the forms of 6'-aminoluciferin (6'-NH₂LH₂) and cyclic alkylaminoluciferin (CycLuc1) emit red-shifted BL.⁷ The longest wavelengths could be achieved with AkaLumine–HCl and its derivatives including seMpai (up to 675 nm)⁸ and infraluciferin (iLH₂, >700 nm).^{7,8} However, their applications are still challenging because the natural Fluc (the most widely used luciferase for general applications) does not utilize these D-LH₂ analogues well. Several enzyme engineering campaigns were carried out to evolve luciferases suitable for using these compounds and increase the BL intensity.⁸⁻¹⁰ Although the engineered luciferase, namely Akaluc, could use AkaLumine–HCl as a substrate and exhibit brighter BL than the native Fluc,^{7,11} the overall signals are still low.

This is due to the low quantum yield of AkaLumine–HCl with Fluc (4.0 \pm 0.5%), which is significantly lower than that of natural D-LH $_2$. Therefore, new red-shifted D-LH $_2$ analogues which can be used as dropped-in substrates for the native beetle luciferases would provide alternative choices for biomedical researchers to directly use the widely available beetle luciferase systems to generate red-shifted BL for cell-based screening applications.

Currently, the synthesis of D-LH₂ and most of its analogues at a scale of >50 mg can only be performed through chemical methods which typically require multi-step organic synthesis and purification (~ 4 steps). The chemical reactions need to be carried out at 120-190 °C using hazardous chemicals such as acetic anhydride (Ac2O), pyridine, sulfolane, and dimethylformamide (DMF)¹³⁻¹⁵ by highly skilled chemist experts. We previously reported the chemo-enzymatic synthesis of novel D-LH₂ analogues with methyl substituent groups at the benzothiazole ring of D-LH2 which gave a red shift in emission up to 620 nm with good BL intensity and light stability. These compounds can be used directly as substrates for the native Fluc and have been demonstrated as environmental monitoring tools without Fluc engineering.16 Our methodology was based on the condensation of benzoquinone (BQ) and D-cysteine (D-Cys) to form the D-LH2 using the HadA enzyme (HELP, HadA Enzyme for Luciferin Preparation);^{17,18} the process was optimized to obtain a high yield of D-LH2 analogues of about $\sim 51\%$ with a 7.1 mg yield at the 0.5 L scale. Although the HELP process is green, its scale up is still limited when compared to other conventional chemical synthesis methods because it utilizes the HadA and associated enzymes.

In this work, we developed a cost-effective one-pot green synthesis procedure for methyl-, methoxy-, and naphthol-p-LH₂ analogues using the condensation reaction of BQ derivatives and p-Cys without adding the HadA and auxiliary enzymes. The process was optimized to improve the product yield for the bulk scale and easy preparation of p-LH₂ analogues while providing compatibility with green chemistry principles¹⁹ (Fig. 1a). We have used the method to prepare six methyl-, methoxy and naphthol-substituted p-LH₂ analogues including natural

D-LH₂ (2a), 5'-methyl-D-LH₂ (5'-MeLH₂, 2b), 16 5',7'-dimethyl-D- LH_2 (5',7'-DiMeLH₂, 2c),²⁰ and 4',7'-dimethyl-D-LH₂ (4',7'-DiMeLH₂, 2e)²¹ with two novel luciferins: 5',7'-dimethoxy-Dluciferin (5',7'-DiOMeLH2, 2d) and 7'-methylnaphthol-D-luciferin (7'-MeNpLH2, 2f). With an appropriate substrate ratio and continuous addition of substrates under anaerobic conditions, the developed one-pot green synthesis approach can preserve the stability of BQ derivatives and p-Cys, preventing substrate deterioration, reducing byproduct formation and increasing D-luciferin analogue yield (by 63-fold relative to that of the previously reported chemical condensation method, 17 Fig. 1b and 1c). The synthesized D-LH2 analogues (2a-2f) were tested with five known beetle luciferases including Photinus pyralis luciferase (Fluc), Phrixothrix hirtus redemitting luciferase (SLR), Pyrearinus termitilluminans emerald luciferase (Eluc), Pyrocoelia miyako luciferase wild-type (Pmluc-WT), and the Pyrocoelia miyako luciferase variant (Pmluc-N230S) commonly used in BL applications²²⁻²⁶ (Fig. 1d). Their biochemical, steady-state kinetics and BL properties of these enzymatic reactions were characterized (Fig. 1e and f). The novel methyland methoxy-D-LH2 analogues demonstrated their better redshifted BL (>600 nm), increased brightness, and slow decay rate of BL emission compared to the reaction of natural D-LH2 (2a). 5'-Methyl-D-LH₂ and 5',7'-dimethyl-D-LH₂ which showed promising BL characteristics in enzymatic reactions were further explored for their in vivo BL properties in mammalian cell lines expressing Fluc, SLR, and Pmluc-N230S. These newly discovered unique properties of methyl-D-LH₂ compounds provide the unique BL characteristics advantageous for future cell-based assays and real-time monitoring applications.

Results and discussion

Synthesis of D-LH₂ analogues and their yield optimizations

We investigated the condensation of various BQ (1a) derivatives and D-Cys to synthesize D-LH₂ analogues with more red-shifted light emission (Fig. 2a). High-resolution quadrupole time of flight mass spectrometry (QTOF-MS) was used for identifying potential products. The results (Table S1) revealed that the BQ derivatives (1a–1f) can serve as electrophilic substrates (Fig. 2b) for the D-Cys condensation reaction, producing the D-LH₂ analogues (2a–2f) shown in Fig. 2c. Structures of the novel D-LH₂ analogues were elucidated by LC/QTOF-MS and NMR (Tables S1–S3) as 5′,7′-DiOMeLH₂ (2d) and 7′-MeNpLH₂ (2f), as shown in Fig. 2c. These compounds were not previously found in nature nor reported.

Previously, the yield of D-LH₂ (2a) obtained using BQ (1a) and D-Cys condensation alone was poor (0.3%).¹⁷ We hypothesized that this was due to the instability of 1a. We thus screened for buffers and reaction conditions which might be suitable for providing better yields. Results in Fig. S1 indicate that the organic buffer, 100 mM HEPES, gave the highest D-LH₂ yield compared to other systems. We further explored co-solvent systems to stabilize 1a and enhance the yield of D-LH₂ synthesis of 2a. We found that the addition of 20% (v/v) methanol

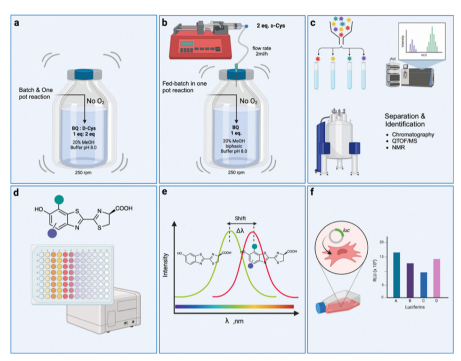


Fig. 1 Graphical representation of the overall workflow. (a) Batch and one-pot reaction for $D-LH_2$ synthesis. (b) Fed-batch synthesis of $D-LH_2$ analogues in one-pot reaction. (c) $D-LH_2$ analogues were isolated and characterized and their structures were elucidated by 1H and ^{13}C NMR and high-resolution mass spectrometry. (d) BL reactions of new $D-LH_2$ analogues with five types of beetle luciferases. (e) BL characteristics of novel $D-LH_2$ analogues. (f) Demonstration of applications for new $D-LH_2$ analogues in mammalian cell line assays and real-time bioluminescence monitoring of live cells.

increased the yield of 2a formation by about 2.8-fold compared to the system without any solvent addition (Fig. S2). We further identified a bottleneck of 1a and D-Cys condensation by varying the percentage of oxygen in the reaction. The reaction buffers were prepared by equilibration with various concentrations of O_2 (<0.0005%, 10%, and 20%) and used in the reactions to monitor the product 2a formed. The results in Fig. 3a reveal that low concentrations of O_2 tend to improve the yield of 2a formation, possibly by preventing the decay of 1a to another form. With <0.0005% O_2 (anaerobic condition), the reaction gave the highest yield of 2a formation which was greater than that of the 20% O_2 (air-saturation) condition by about 1.8-fold with a D-LH₂ formation rate of 3.26 μ M h⁻¹ (Fig. S5 and Table S4).

After finding that the instability of the **1a** derivatives and D-Cys substrates is a limiting factor for the generation of D-LH₂ analogues, we designed a flow chemistry strategy by continuously mixing one substrate with another under various conditions. Fig. 3b illustrates the reactions to generate **2a** in batch and continuous feeding (fed-batch) systems. Due to the O₂ vulnerability of the substrates, only low percentages of **2a** formation (up to 18%) could be obtained from the batch reactions within 24 h. For the continuous feeding of **1a** into the anaerobic buffer containing 2 (orange) and 4 (old rose) equivalents (eq.) of D-Cys, the yield of **2a** formation was about twice that of the batch reactions (Fig. 3b, orange bar at 24 h). Remarkably, when 2 equivalents (eq.) of D-Cys were pumped into 1 eq. of **1a** under anaerobic conditions, the highest yield of **2a** formation (about 86 μM) could be obtained (Fig. 3b, pink bar

at 24 h). This yield was 16.4-fold greater than that of the batch reaction after 12 h (Fig. 3b). Therefore, by pumping 2 eq. of D-Cys into a 1 eq. solution of **1a** under anaerobic reactions (Fig. 3c), the stability of the substrates could be greatly enhanced and the reaction could be finished after 8 h (Fig. 3d).

To synthesize D-LH2 analogues at a semi-large scale, our method described above and the recently published protocol²⁷ of using BQ, L-Cys methyl ester and D-Cys to synthesize D-LH2 were tested. We found that 2a and 2c compounds could be prepared with 50 and 66% yield, respectively, when using the BQ derivatives (1a and 1c) and the L-Cys methyl ester and D-Cys condensation protocol. However, this protocol could not be used to prepare other D-LH₂ analogues (compounds 2b and 2d) because it gave a mixture of products (non-specific reactions in the case of 2b synthesis) and the electron-donating group of 1d affected the cyclization of the benzothiazole ring (in the case of 2d synthesis). These compounds could only be prepared using a fed-batch reaction shown in Fig. 3 and described above. The fed-batch protocol developed in this work is thus useful for synthesizing D-LH2 analogues without utilizing harmful catalysts or toxic chemicals. The reaction also requires less usage of organic solvent and thus is compatible with green chemistry principles. These findings set an important foundation for future large-scale production of the compounds.

NMR, mass spectra, absorption, and fluorescence characteristics of D-LH₂ analogues

The method described in the previous section was used to synthesize D-LH₂ analogues (2a-2f) from the BQ derivative

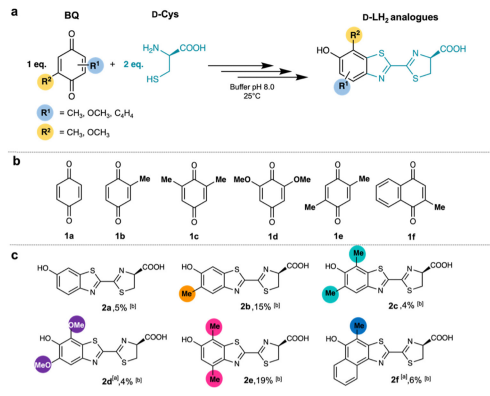


Fig. 2 Synthesis of D-LH₂ and its analogues by condensation of D-Cys and BQ derivatives. (a) D-LH₂ and its derivatives could be obtained under batch conditions. (b) BQ derivatives (1a-1f) serve as substrates for synthesizing p-LH2 analogues. (c) The p-LH2 analogues (2a-2f) obtained from condensation of p-Cys and BQ derivatives (1a-1f). [a] Novel p-LH₂ analogues are 5',7'-DiOMeLH₂ (2d) and 7'-MeNpLH₂ (2f). [b] Isolated yields of p-LH₂ analogues after purification.

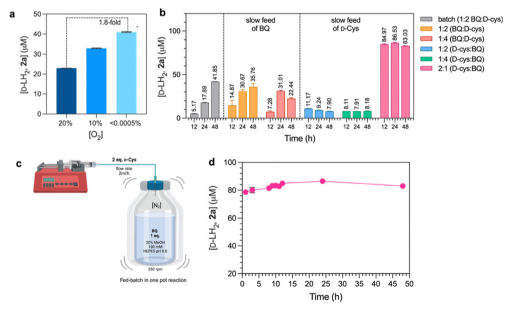


Fig. 3 Optimization of 2a production yield. (a) Effects of O₂ concentrations on 2a formation. (b) Synthesis of 2a by pumping a solution of one substrate into another substrate solution with different ratios under anaerobic conditions. (c) The synthesis of 2a could be set up by continuously feeding 2 eg. of D-Cys (2 mL h⁻¹) into 1 eq. of **1a** under anaerobic conditions. (d) Kinetics of continuous formation of **2a** by feeding 2 eq. of D-Cys (2 mL h⁻¹) into 1 eq. of **1a** under anaerobic conditions.

substrates (1a-1f, Fig. 1b). The products were isolated (purity up to 97%), and their structures were characterized using

QTOF-MS and NMR spectroscopy (Fig. 1c and Fig. S3 and S4 and Tables S3 and S4). The observed m/z with low mass error values of D-LH₂ analogues (2a–2f) from the QTOF-MS data (Table S1) confirmed that the BQ derivatives (1a–1f) can react with D-Cys to yield the products in one-pot synthesis. The structures of 2a–2f were then elucidated using ¹H NMR (Fig. S3 and Table S2) and ¹³C NMR (Fig. S4 and Table S3) to identify the molecular structures of the D-LH₂ analogues. The NMR results indicate that the benzothiazole rings of the D-LH₂ analogues were substituted with methyl and methoxy groups at 4′, 5′, and 7′-positions (Fig. 2c) and demonstrated that we could obtain novel compounds including 5′,7′-DiOMeLH₂ (2d) and 7′-MeNpLH₂ (2f). All of the D-LH₂ analogues (2b–2f) were further investigated for their absorption and fluorescence characteristics in comparison with the native D-LH₂ (2a) as described below.

Absorption and fluorescence characteristics of D-LH2 analogues (2a-2f) in 100 mM HEPES pH 8.0 at 25 °C were recorded. The natural D-LH₂ (2a) displayed a maximum adsorption peak (λ_{max}) at 327 nm, while **2b** which has a methyl group addition at the 5'-position of the benzothiazole moiety also gave a similar λ_{max} at 323 nm (Table S6). However, for 2c and 2e which have two methyl substitutions at the benzothiazole skeleton, their absorption spectra shifted to longer wavelengths of λ_{max} at 334 and 330 nm, respectively (Table S5). The addition of dimethoxy (2d) and naphthol groups (2f) resulted in longer wavelength shifts (λ_{max} at 412 and 437 nm, respectively) (Table S5). Similarly, the fluorescence properties of natural D-LH₂ (2a) and methyl-D-LH2 analogues (2b, 2c, and 2e) showed similar fluorescence emission spectra with peaks around 523-560 nm. For 2d and 2f, their emission spectra showed a greater bathochromic shift with the peaks around 620 and 580 nm, respectively (Table S5). These absorption and fluorescence characteristics of the D-LH₂ analogues (2a-2f) provide a means by which BL can be red shifted (shown later). It should be noted that their uniqueness in fluorescence is also valuable for future use in florescencebased sensor applications. 28,29

Bioluminescence characteristics of D-LH2 analogues

The purified D-LH2 analogues (2a-2f) were investigated for their BL characteristics using known luciferases, i.e., Fluc, SLR, Eluc, Pmluc-WT, and Pmluc-N230S. Results in Fig. 4a showed that 2b-2f could serve as substrates for various luciferases (except 2f, which could not serve as a substrate for Pmluc-WT and Pmluc-N230S) and generate light emission with longer wavelengths (red-shifted) compared to the natural D-LH₂ (2a). Interestingly, the pairing of novel D-LH2 analogues (2d and 2f) with SRL and Fluc gave red light emissions at longer than 630 nm which could be recorded by a digital single-lens reflex (DSLR) camera (Fig. 4b-e). We found that a methyl substituent in the benzothiazole ring of D-LH2 could also modulate absorption, fluorescence and BL characteristics of luciferases (Table S5). Particularly, the BL emission was altered from yellow-green (\sim 560 nm) of the natural D-LH₂ (2a) to be in the red-shifted region (>600 nm).

We also compared the photon emission of methyl-substituted D-LH₂ analogues 2b and 2c to that of the natural compound (2a) using an integrating sphere-based multichannel spectrometer

with a charge-coupled device (CCD) detector (details are given in SI, Section S1.5). The photon/pixel emission spectra of the luciferase reaction showed that red-shifted methyl-substituted D-LH₂ analogues (**2b** and **2c**) provided greater photon emission than that of the natural compound (**2a**). Interestingly, **2b** showed greater photon emission than those of **2a** in Fluc, SLR, and Pmluc-N230S reactions (Fig. 4f-i). The spectra of **2b** and **2c** also showed the emission wavelengths longer than 600 nm. BL properties of D-LH₂ analogues (**2a–2c**) correlate well with their quantum yields in the reactions of Fluc, SLR, and Pmluc-N230S measured later.

Importantly, the D-LH $_2$ analogues (2b–2f) in this study allowed the reactions of Fluc and other luciferases to gain a maximum BL emission band shift ($\Delta^{\rm BL}\lambda$) of >40 nm and up to 88 nm compared to 2a (Fig. 4j). The largest BL wavelength shift in this study could be obtained from the 2d-SLR orthogonal pair that emitted a maximum light peak at 663 nm and a broad BL spectrum beyond 750 nm, which could be beneficial for bioimaging and real-time deep tissue *in vivo* monitoring applications in the future.

Kinetics and thermodynamic properties of new D-LH₂ analogues with different luciferases

To investigate the substrate specificity and the influence of substituents of D-LH2 analogues on BL characteristics, steadystate kinetics of luciferase (Fluc, SLR, Eluc, Pmluc-WT, and Pmluc-N230S) reactions using various D-LH2 analogues (2b-2f) as substrates were carried out at 25 °C and pH 8.0 to measure their kinetic constant, $K_{\rm m}$ and $k_{\rm cat}$, values compared to those of the natural D-LH₂ (2a). The results in Table S6 show that, for all luciferases studied, the $K_{\rm m}$ values of 2b were comparable to those of 2a, suggesting that a methyl group substituent at the 5'-position of luciferin's benzothiazole ring (2b) does not interfere with the ability of the enzyme to interact and react with a substrate. It should be noted that while the natural D-LH₂ mostly exhibits strong substrate inhibition with a range of > 1-10 μM against various luciferases, p-LH₂ analogues (2b-2f) showed lower severity of substrate inhibition (Table S6). For the compounds with methoxy substituents at 5' and 7'-positions or methyl substituents at the 4' and 7'-positions, such as 2d, 2e and 2f, their $K_{\rm m}$ values were mostly greater than those of 2a, implying that these substituents may disrupt the interactions of these luciferins to serve as substrates for luciferases.

The $k_{\rm cat}$ values of the BL reactions indicate that the reactions of ${\bf 2b}$ with various luciferases, Fluc, SLR, Pmluc-WT, and Pmluc-N230S, yielded $k_{\rm cat}$ values greater than those of the native substrate (${\bf 2a}$). In particular, the reaction with Pmluc-N230S resulted in a 5.2-fold increase in the $k_{\rm cat}$ value. However, the stability of the light emission ($t_{1/2}$) of ${\bf 2b}$ is very similar to that of ${\bf 2a}$ in Fluc and Pmluc-WT, and light emission by SLR and Pmluc-N230S could be prolonged longer than those of other systems by about 1.5 and 1.7-fold, respectively (Fig. 5b). This result suggests that ${\bf 2b}$ with a methyl substituent at the 5'-position of the benzothiazole skeleton does not display any adverse effects on the activities of luciferases or their BL properties. The reactions of ${\bf 2c}$ exhibited $k_{\rm cat}$ values of

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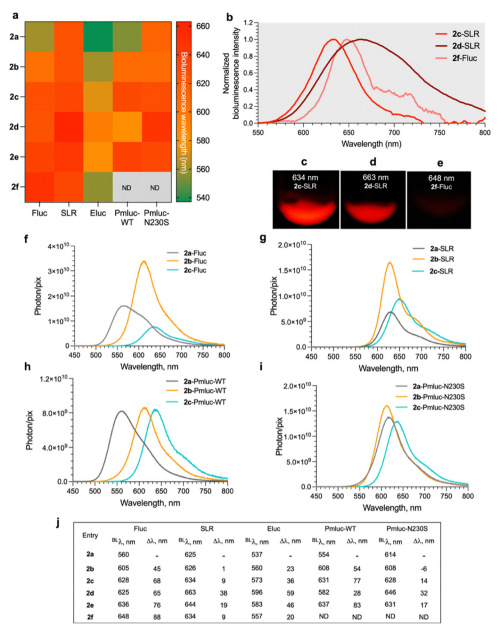


Fig. 4 BL characteristics of p-LH2 analogues (2a-2f) reacted with various luciferases. Reactions of various luciferases with p-LH2 analogues (2b-2f) were carried out at 25 °C in 100 mM HEPES buffer at pH 8.0 and compared to those of natural D-LH₂ (2a). (a) Heat maps describing BL emission of D-LH₂ analogues (2a-2f) generated from their reactions with Fluc, SLR, Eluc, Pmluc-WT, and Pmluc-N230S luciferases. (b) BL emission spectra of novel p-LH₂ analogues (2d and 2f) obtained from their reactions with their orthogonal luciferases. (c-e) BL signals obtained from the reactions of novel D-LH₂ analogues (2d and 2f) and their orthogonal luciferases were recorded by a DSLR camera. (f-i) The BL emission spectra of the luciferase (5 μM)-luciferin (2 μM) reactions were measured using the integrating sphere-based multichannel spectrometer. (j) The difference in emission wavelengths of D-LH₂ analogues (2b-2f) compared to those of the natural D-LH₂. ND denotes the non-detectable signals.

approximately half that of 2a (except in the reaction of SLR, in which 2c gave a slightly higher k_{cat} than 2a by approximately 1.3-fold). 2d-2f provided very low bioluminescence intensity (BLI) in all luciferases tested (Fig. 5a and Table S7). The results indicate that 2d-2f showed larger $K_{\rm m}$ and lower BLI values than the native substrate (2a) in most of the enzyme reactions investigated. The data indicate that reactions of 2d-2f were not suitable with currently available luciferases and would require future work to engineer luciferases to obtain the best orthogonal pair of luciferase-D-LH2 analogues with high BLI. We noted that Eluc prefers to use 2a as a substrate rather than other analogues (Fig. 5a and Table S6) based on the $k_{\rm cat}$ and $K_{\rm m}$ values, but other enzymes such as Fluc, SLR, Pmluc-WT, and Pmluc-N230S could use 2b as an alternative substrate well. We also measured the stability of light emissions of the D-LH₂ analogues (2a-2c). The results in Fig. 5b clearly indicate that 2c gave longer BL half-lives $(t_{1/2})$ than the natural D-LH₂ when using Fluc and Pmluc-N230S, by about 1.7- and 5.6-fold,

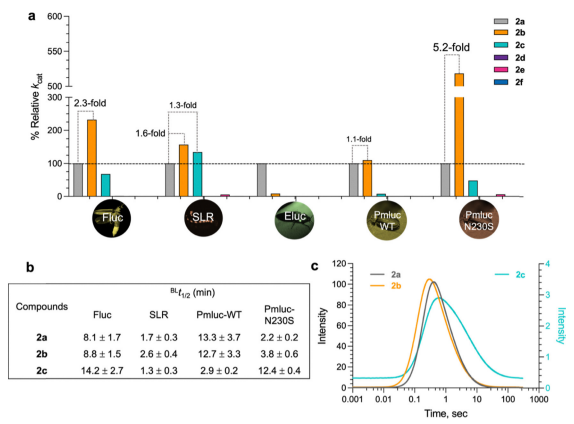


Fig. 5 BL characteristics of D-LH₂ analogues and luciferases. (a) The relative k_{Cat}, which represents the BL intensity (BLI) of each D-LH₂ analogue (2a-2f) when it reacts with luciferases compared to those of 2a reactions. (b) The half-time ($t_{1/2}$) values of BL emission of p-LH₂ analogues (2a-2c) which react well with Fluc, SLR, Pmluc-WT, and Pmluc-N230S luciferases. (c) Kinetics of light formation and light decay of Fluc reactions with D-LH2 analogues (2a-2c) monitored by stopped-flow luminometry.

respectively, as compared to those of 2a. Without the addition of coenzyme A (CoA), the ability of 2c in prolonging BL in Fluc and Pmluc-N230S should be useful for future live-cell imaging applications. Thus, the current best substrate analogues of luciferases such as Fluc, SLR, Pmluc-WT, and Pmluc-N230S are 2b and 2c which are also promising substrates for redshifted BL and steady light emission in the reaction of SLR. Although 2d and 2f are novel compounds with potentially useful long wavelength emission, their usage with native luciferases is not very efficient due to their low BL signals. This is due to non-compatibility of these compounds with native enzymes. Engineering campaign of beetle luciferases is required for improving their BL signals in the future.

To comprehensively characterize the bioluminescence properties of the D-LH₂ analogues (2a-2d), we measured their bioluminescence quantum yields from their reactions with Fluc, SLR, and Pmluc-N230S. The reactions were performed using the conditions obtained from steady-state kinetics studies (Fig. 5a and Table S6). Results shown in Table 1 and Table S8 indicate that the reaction of Fluc with 2b gave the greatest quantum yield (0.52) and the value was higher than those of the natural D-LH2 and other D-LH2 analogues. The data clearly support potential applications of Compound 2b. For the reactions with SLR and Pmluc-N230S, compounds 2a-2c showed the same range of quantum yields. However, the

Table 1 Quantum yields of luciferases and D-LH₂ analogues

	Quantum yield with different luciferases				
Luciferins	Fluc	SLR	Pmluc-N230S		
2a	0.44 ± 0.04	0.160 ± 0.005	0.160 ± 0.018		
2b	0.52 ± 0.01	0.190 ± 0.005	0.170 ± 0.034		
2c	0.200 ± 0.007	0.160 ± 0.003	0.150 ± 0.014		
2d	ND^a	ND^a	ND^a		

^a ND denotes "not determined" because 2d gave low bioluminescence signals.

quantum yield of Compound 2d with all luciferases could not be measured due to low bioluminescence signals. Altogether, the results of quantum yield measurements indicate that the Fluc-2b pair gives the greatest value of BL signals.

Furthermore, the equilibrium binding constant (K_d) values of D-LH₂ analogues (2b-2d) with Fluc were measured using isothermal titration calorimetry (ITC) and compared with the values of natural D-LH2 (2a). The results in Table 2, Table S9 and Fig. S7 reveal that the K_d value of 2b was about half that of 2a, indicating that 2b has a greater binding affinity to Fluc than 2a. The data suggest that the 5'-methyl substituent in 2b does not interfere with the binding interactions between the substrate and the Fluc pocket site but rather promotes a tighter binding. For other D-LH2 analogues, the results indicate that

Table 2 Thermodynamics of ligand binding and kinetics constants for light formation and decay of the reactions of $D-LH_2$ analogues (**2a–2c**) with Fluc

	Dissociation constar	nt (<i>K</i> _d , μ M)	Observed rate constant (k_{obs})		
				Light decay (s ⁻¹)	
Compound	Fluc: Luciferin ^a	Fluc-ApCPP: Luciferin b	Light formation (s^{-1})	Faster phase	Slower phase
2a	21.1 ± 7.4	4.2 ± 0.1	6.64 ± 0.05	1.57 ± 0.02	0.29 ± 0.00
2b	13.4 ± 2.5	4.00 ± 0.02	9.63 ± 0.04	1.55 ± 0.15	0.27 ± 0.00
2c	35.1 ± 3.1	25.3 ± 1.1	7.28 ± 0.02	0.27 ± 0.00	0.05 ± 0.00
2d	30.4 ± 8.2	<u></u> c	8.13 ± 0.08	0.50 ± 0.00	ND^d
2e	NC^e	864 ± 7	<u></u> c	<u></u> c	<u></u> c

^a The K_d values were measured by titration between Fluc and luciferins. ^b The K_d values were measured by titration between Fluc-ApCPP complex and luciferins. To probe interactions between D-LH₂ analogues and the Fluc-ATP complex, we used the non-hydrolyzable ATP analogue (ApCPP, Adenosine-5'-[(α,β)-methyleno]triphosphate) instead of ATP. ^c—Denotes "not measured". ^d ND denotes "not determined" because **2d** kinetics gave only one decay phase. ^e NC denotes "not calculated" by ITC because **2e** gave a very high K_d value with Fluc.

modification of the luciferin benzothiazole ring at the 5'- and 7'-positions with methyl (2c) and methoxy groups (2d) yielded $K_{\rm d}$ values comparable to that of 2a, indicating that these substituents also do not interfere with the binding behavior. To investigate whether the pre-binding of ATP can enhance the interactions of D-LH₂ analogues (2b-2d) with Fluc, we measured $K_{\rm d}$ values for binding of D-LH₂ analogues to the preformed complex of Fluc and adenosine-5'- $[(\alpha,\beta)$ -methyleno]triphosphate (ApCPP) (Table 2 and S9). ApCPP is an ATP analogue mimicking the adenylated-ATP intermediate structure which does not allow Fluc to proceed with catalysis. This Fluc-ApCPP complex thus allowed us to probe interactions between D-LH₂ analogues and the Fluc-ATP complex. Results in Table S9 indicate that all binding of D-LH2 analogues exhibited lower $K_{\rm d}$ values in the Fluc-ApCPP complex, implying that the presence of ATP enhances stronger interactions of D-LH2 analogues with Fluc.30

We also performed stopped-flow experiments to determine the observed rate constants $(k_{\rm obs})$ of single turnover reactions of light formation and decay of the Fluc reactions with ${\bf 2a-2c}$ (Fig. 5c and Table 2). The BL kinetics revealed three exponential phases (Fig. S6). The first phase was light formation, while the second and third phases were light decay with faster and slower rate constants, respectively. The kinetic analysis showed that all compounds displayed comparable $k_{\rm obs}$ values of light formation, suggesting that the substituents of D-LH2 analogues synthesized in this work do not affect the light formation kinetics. Interestingly, the $k_{\rm obs}$ values of light decay in both faster and slower phases of the ${\bf 2c}$ reaction were about 5–6-fold slower than those of ${\bf 2a}$ and ${\bf 2b}$, indicating that the overall light emission yield in the reaction with ${\bf 2c}$ would be much greater than those of ${\bf 2a}$ and ${\bf 2b}$.

We then used molecular docking and molecular dynamics (MD) simulations to explain the binding and kinetic behaviors of the D-LH₂ analogues and luciferases as shown in Fig. S8–S19. The data revealed that all D-LH₂ analogues (**2b–2e**) can bind to the active site of Fluc (Fig. S8a–e). To further investigate the stability of the enzyme and D-LH₂ analogue complexes, MD simulations were performed across a range of temperatures (300–360 K). Results in Fig. S9–S18 showed that the structure of Fluc with **2a** and **2b** bound remained stable across all temperatures tested, particularly at the elevated temperature of 360 K

compared to that of Fluc with 2c bound. These findings suggest that 2a and 2b exhibit stronger binding to Fluc compared to 2c; these data correlate well with K_d values obtained from the ITC experiments (Table 2).

In order to explain the equilibrium binding constant (K_d) values of D-LH₂ analogues (2b-2d) with Fluc, the crystal structure of the luciferase enzyme (Fluc) from *Photinus pyralis* with adenylate analogue (DLSA) bound (PDB: 4G36) was analyzed. Residues within 6 Å of the 7'-position of the bound DLSA are shown in Fig. S19. The 7'-position of DLSA is surrounded by the helix (F247 and T251), the lower beta strand (L286), the middle beta strand (A313 and S314), the middle loop (G315 and G316), and two upper beta strands (Q338, G339, Y340 and A348, L350, I351) which are linked by the top loop (S347). The introduction of a methyl substituent at the 5'-position could interact with T251, leading to the lower K_d of 2b compared to 2a. In contrast, the introduction of a methyl substituent at the 7'-position could clash with the middle beta strand (A313 and S314) and the upper beta strand (Q338, G339, Y340) leading to the higher $K_{\rm d}$ of **2c** and **2d** compared to **2a**.

The overall data reported here indicate that **2b** and **2c** can be used well as dropped-in substrates for the currently available luciferases without requiring any enzyme engineering to provide red- shifted, high BL signals, and good light stability which could be useful for live-cell and animal BL applications in the future.

Demonstration of the use of D-LH₂ analogues in Mammalian cell assays

We tested the ability of the purified D-LH₂ analogues (2a–2f) to serve as substrates for Fluc expressed in mammalian cell lines including HepG2, HEK293T, HeLa, and RAW264.7. These cells are commonly used as hosts for monitoring the gene expression, cellular activities and molecular reactions of diseases. Si,31–34 BL signals generated by expression of Fluc in these cells have been used as tools to address various research topics in biomedical research. We transfected mammalian cells with the plasmid pGL4.13 expressing Fluc (details are given in SI, Section S1.6). Cells were collected after 24 h of growth for lysis, and the crude lysates were then assayed using the D-LH₂ analogue cocktail reagents (2a–2f). The results showed that 2b provided light intensities similar to those of 2a with the crude lysates of HepG2, HeLa, and RAW264.7 (Fig. 6a). Interestingly, 2b

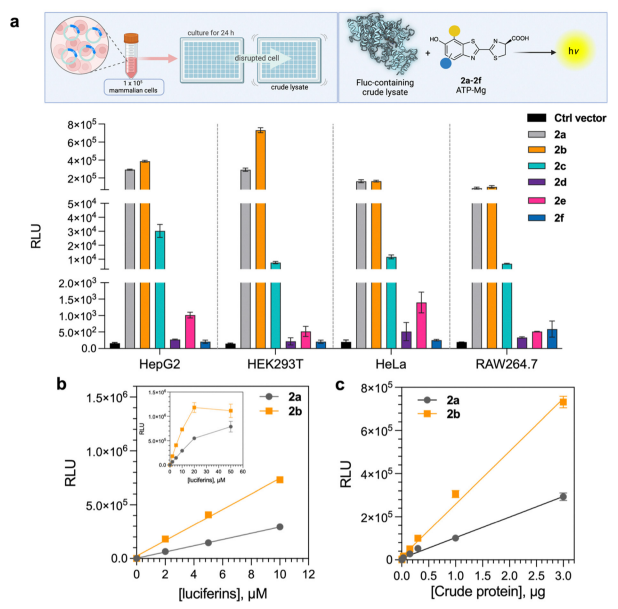


Fig. 6 Applications of $D-LH_2$ analogues (2a-2f) in mammalian cell line BL assays. (a) The crude lysate of each cell line was tested with $D-LH_2$ analogues (2a-2f). (b) Comparison of light intensity from reactions of the Fluc-containing HEK293T crude lysate with 5'-MeLH₂ (2b) and natural $D-LH_2$ (2a) at various concentrations. (c) Dependence of light intensity on various concentrations of natural $D-LH_2$ (2b).

showed a higher light intensity in the HEK293T crude lysate than **2a**. In addition, **2c** exhibited good light intensity with HepG2, HEK293T, HeLa, and RAW264.7 cell lysates, whereas other analogues (**2d–2f**) gave poor light intensity (Fig. 6a). All results suggested that **2b** has good potential to be used as a substrate to generate redshifted signals in BL applications because it can be used directly in commonly used cell-based assays with the wild-type Fluc.

We then further investigated the BL properties of **2b** by varying its concentration in the reaction with a fixed amount of HEK293T crude lysate and compared the light intensity with that produced by the native substrate **2a**. The results in Fig. 6b clearly demonstrated that **2b** gave greater light intensity than **2a** by about 2.5-fold. We then varied the concentrations of **2a** and **2b** with a fixed amount of HEK293T lysate and then varied the

cell lysate amount with a fixed concentration of both compounds. The results as shown in Fig. 6b and c indicate that **2b** provided greater sensitivity than **2a** by about 2-fold.

The limit of detection (LOD) of expressed Fluc in crude cell lysates using **2a** and **2b** was identified to be 51.6 and 4.6 ng, respectively, indicating that **2b** can give a better sensitivity than the natural p-LH₂ (**2a**) by about 11.2-fold. Altogether, these results also confirmed that 5'-MeLH₂ (**2b**) can serve as a promising substrate for future cell-based assay applications.

Bioluminescence characteristics of D-LH₂ analogues in live cell applications

To demonstrate the use of D-LH₂ analogues (2a-2c) in live cells, we measured the BL spectra of 2a-2c in the HepG2 cell line

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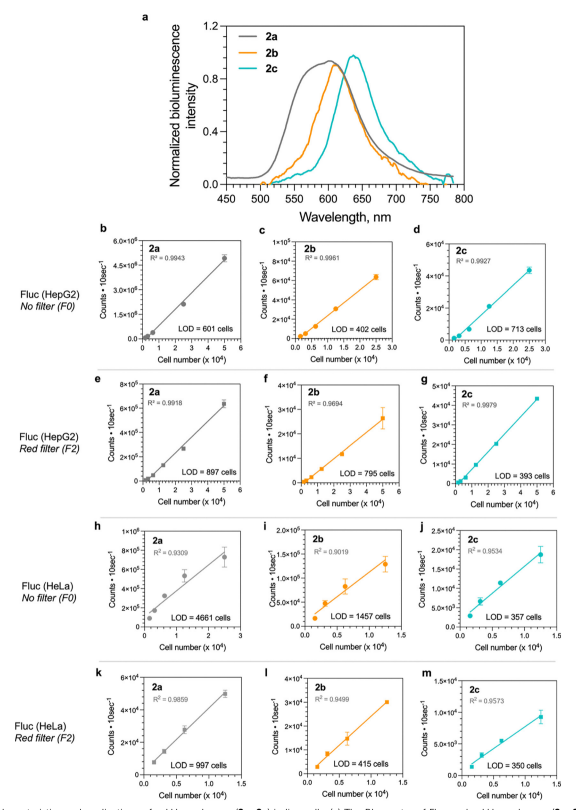


Fig. 7 BL characteristics and applications of $p-LH_2$ analogues (2a-2c) in live cells. (a) The BL spectra of Fluc and $p-LH_2$ analogues (2a-2c) in stable reporter HepG2 cell lines harboring the fluc gene. The spectra were measured in live cells in 25 mM HEPES-NaOH pH 7.0 containing DMEM culture medium at 37 °C. The use of $p-LH_2$ analogues (2a-2c) in living HepG2 cells. $p-LH_2$ analogues were tested for their detection sensitivities by varying the number of HepG2 reporter cells using different BL filters including F0 (no filter) and F2 (red filter, >620 nm long-pass filter). (p-d) Linearity ranges of HepG2 cell number detection using p-d2 cell number detection using p-d3 cell number detection using p-d4 cells expressing fluc gene by transient transfection were tested with $p-LH_2$ analogues using different BL filters including F0 (no filter) and F2 (red filter, p-d4 cell number detection using p-d4 cell number detection using p-d5 no filter). (p-d4 cell number detection using p-d6 no filter). (p-d6 no filter). (p-d6 no filter). (p-d7 no filter). (p-d8 no filter). (p-d8 no filter). (p-d9 no filter).

stably expressing a Fluc reporter (details are given in SI, Section S1.6). The BL spectra in intact cells were captured using a highly sensitive CCD spectrophotometer with 2 min exposure time. The spectra in Fig. 7a and Fig. S21 showed a broad spectrum of **2a**-Fluc in live cells with a peak at 604 nm, which differed from those of the purified Fluc reactions (Fig. 4f). However, the spectra of **2b** and **2c** in living cells were identical to those of the purified Fluc reactions (refer to Fig. 7a and 4f) with red-shifted wavelengths at 609 and 638 nm, respectively. HepG2 and HeLa cell lines were chosen for further investigation because they are among the most commonly used cell lines in biomedical research (41 200 and 46 000 publications in 2023, respectively, searched by Google Scholar, Fig. S20).

We further investigated the sensitivity of reporter HepG2 and HeLa cell line detection when 2a-2c were used as substrates. The number of HepG2 and HeLa cell lines was varied by 2-fold dilution and the cells were seeded in a 96-well plate. 100 μM (final concentration) of 2a-2c in culture medium was then added to the cells and incubated for 30 min. The BL signals (photon counts) after a 30 min incubation period were measured with different filters to increase the selectivity for the BL wavelength-especially towards red-shifted light. In this work, we used optical filters including F0 (no filter) and F2 (red filter, > 620 nm long-pass filter) to enhance the BL signals at specific wavelength regions to reduce background signals. The detection of live HepG2 and HeLa cell lines using an F0 filter (no filter) in Fig. 7b-m showed a good linearity ($R^2 > 0.99$) with all compounds (2a-2c). The sensitivity for detection in live Fluc-expressing HepG2 cells using 2b was better than those of 2a and 2c, with an LOD of approximately 402 cells in the assay, using the F0 filter (no filter). The results in Fig. 7e-g showed linearity ranges for detection by 2a-2b measured using the F2 filter, which collected light signals >620 nm. 2c showed a prominent ability of red emitted signals for live cell detection in red BL regions, displaying high sensitivity when measured in HepG2 cells with a red filter, which required only 393 cells. In contrast, LODs for the similar systems with 2a and 2b are 897 and 795 cells, respectively (Table 3).

Interestingly, the sensitivity for detection in live Flucexpressing HeLa cells treated with **2c** gave an LOD of approximately 357 cells which had higher sensitivity than those of **2b** and **2a** about 4.1 and 13.0-fold, respectively, using the F0 filter (no filter) (Fig. 7h–j and Table 3). Detection using the F2 filter for HeLa reporter cells also showed the LODs with **2c**, **2b**, and **2a** as 350, 415, and 997 cells, respectively (Fig. 7k–m and Table 3). The HeLa cells with **2c** gave higher sensitivity than that of **2a** about 2.8-fold using the F2 filter.

We further tested the usage of p-LH₂ analogues (2a-2c) with SLR and Pmluc-N230S expressing HepG2 and HeLa cells. Results in Fig. S22 and S23 and Table 3 indicate that most of the red-shifted p-LH₂ (2b and 2c) provided higher sensitivity with lower LOD values than that of 2a, except for Pmluc-N230S-expressing HepG2 and HeLa cells treated with 2c.

The results suggested that methyl-substituted luciferins (2b and 2c) can permeate through the cell membrane of live cells and serve as substrates for Fluc inside the cytosol well. The high

Table 3 Detection sensitivities of luciferase-expressing cells treated with $D-LH_2$ analogues using different optical filters

		Limit of detection (LOD), cells		
Reporter cell	Filter	2a	2b	2c
Fluc (HepG2)	F0	601	402	713
SLR (HepG2)		802	643	500
Pmluc-N230S (HepG2)		2010	752	ND ^a
Fluc (HepG2)	F2	897	795	393
SLR (HepG2)		1071	783	613
Pmluc-N230S (HepG2)		1880	247	ND ^a
Fluc (HeLa)	F0	4661	1457	357
SLR (HeLa)		1352	973	1437
Pmluc-N230S (HeLa)		1428	630	ND ^a
Fluc (HeLa)	F2	997	415	350
SLR (HeLa)		1268	826	707
Pmluc-N230S (HeLa)		1185	812	ND ^a

^a ND denotes "not determined" because 2c gave weak BL signals.

light intensity of **2b** in this red BL region also demonstrated its ability to serve as a new substrate (in replacement of **2a**) for luciferases. **2c** also showed high BL signals at longer red-shifted wavelengths which was suitable for detecting luciferase-expressing cells in low numbers because it provides high sensitivity for red light emission.

The reporter HepG2 cell line expressing Fluc, SLR, and Pmluc-N230S was also real-time monitored after adding 100 μM (final concentration) of D-LH₂ analogues (2a-2c) in Fig. 8a-i. The BL characteristics of all D-LH2 analogues in live cells were measured for 3 days. The results showed that the light intensity of 2b in stable Fluc-expressing reporter cells was slightly higher than that of 2a for 24 h when measured using both F0 and F2 filters and then gradually decreased over time after 30 h. 2c showed greater red BL signals (highest F2/F0 ratio) than other compounds but gave lower BL intensity than 2a and 2b when using the F0 filter or no filter was used (Fig. 8a-c). The highest BL signals of 2a-2c could be achieved at 44, 35, and 6 h, respectively. The data indicate that the highest signals of 2c and 2b BL could be obtained at 7.3 and 1.2-fold faster rates than that of 2a, respectively. This may be due to the ability of 2b and 2c to penetrate the cells faster than 2a.

The transient SLR-expressing HepG2 cells with **2b** showed faster light generation kinetics, which gradually decreased after 10 h. The real-time BL signal patterns of **2a** and **2c** were also similar to that of **2b**, with **2c** exhibiting the highest F2/F0 ratio (red BL characteristics) than those of **2a** and **2b** (Fig. 8d–f). The transient Pmluc-N230S-expressing HepG2 cells showed similar kinetic patterns with all compounds **2a–2c** (Fig. 8g–i). The cells with **2b** and **2c** appeared to prolong the BL signals after 40 h better than that of **2a**.

Moreover, when the HeLa cell lines transiently expressing Fluc, SLR, and Pmluc-N230S were treated with 100 μ M (as a final concentration) of D-LH₂ analogues (2a–2c), their real-time monitored kinetic characteristics were different from those of HepG2 cells (Fig. S24). The highest BL signals could be achieved approximately at 510 h. After a maximum peak at

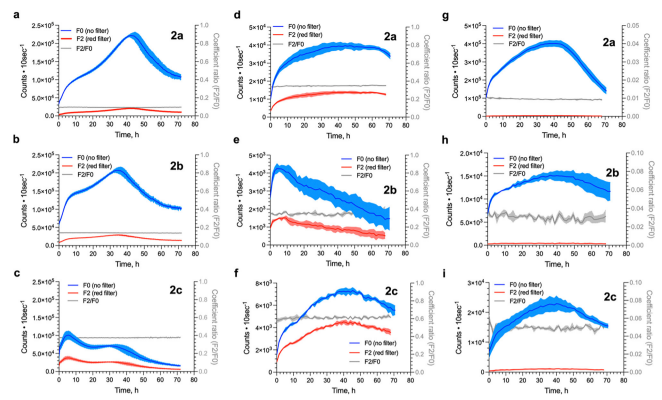


Fig. 8 Real-time monitoring of BL signals of HepG2 cell lines with luciferase reporters and p-LH2 analogues (2a-2c). (a-c) BL real-time monitoring signals of stably Fluc-expressing HepG2 cells with natural p-LH₂ (2a), 5'-MeLH₂ (2b), and 5',7'-DiMeLH₂ (2c), respectively. (d-f) BL real-time monitoring signals of transient transfection of SLR reporter in HepG2 cells with natural D-LH2 (2a), 5'-MeLH2 (2b), and 5',7'-DiMeLH2 (2c), respectively. (g-i) BL realtime monitoring signals of transient transfection of Pmluc-N230S reporter in HepG2 cells with natural D-LH₂ (2a), 5'-MeLH₂ (2b), and 5',7'-DiMeLH₂ (2c),

5 h, the BL signal of the 2a compound (a native p-LH₂) with Fluc-expressing HeLa cells gradually decreased. BL signals of 2b and 2c could be prolonged much better, up to 13 h before decreasing (Fig. S24b and c). Similarly, the kinetics patterns in SLR-expressing HeLa cells (Fig. S23d-f) showed more prolonged BL signals (up to 18 h) with 2c than those of 2a and 2b compounds (up to 10 h) before signal decreasing. The Pmluc-N230S-expressing HeLa cells exhibited low BL signals with all 2a-2c compounds, particularly with the 2c compound (Fig. S23g and h). Kinetics of these BL signals indicate the ability of these D-LH2 analogues (2a-2c) to penetrate the cell membrane and maintain their activities inside the cells.

All results suggest that D-LH₂ analogues with a methyl substituent at the benzothiazole ring, including 2c and 2b, can penetrate the cells faster than natural D-LH₂ (2a). Therefore, 2b and 2c showed great potential to be used as substrates for cell-based assays and real-time monitoring experiments.

Conclusions

Here, we developed a new setup for cost-effective one-pot green synthesis of six D-LH2 analogues using mild chemical reagents. We discovered that by establishing a continuous feeding of D-cys (2 eq.) into an anaerobic one-pot reaction containing

anaerobic 1a (1 eq.), the substrate instability (1a derivatives and D-Cys) and yield of D-LH2 analogues could be significantly improved. The process could improve the production yield by 63-fold compared to the previously reported chemical condensation method.¹⁷ The synthesis method here could synthesize two new D-LH2 analogues including 5',7'-DiMeOLH2 (2d) and 7'-MeNpLH₂ (2f). Substituents such as methyl, methoxy, and naphthol groups at the luciferin's benzothiazole ring (2b-2f) generated longer red-shifted BL wavelength than that of the natural D-LH₂ (2a) by up to 88 nm when reacting with five types of beetle luciferases. The longest red-shifted BL could be obtained from the 2d-SLR orthogonal pair reaction with a maximum BL at 663 nm. However, the reactions of luciferases with 2d and 2f gave low signals; this requires enzyme engineering to improve the BL signals of 2d and 2f in the future.

We demonstrated the use of D-LH2 analogues (2a-2f) in mammalian cell lines. The crude lysate assays exhibited good light intensity in the reactions of 2b and 2c in HepG2, HEK293T, HeLa, and RAW264.7 cells, especially 2b provided greater detection sensitivity of the Fluc enzyme in crude lysates than 2a by about 11.2-fold; this is beneficial for bio-reporter and cell-based assay applications. Real-time BL measurements in live cells showed that 2b and 2c provide greater BL redshifted signals than 2a. Altogether, our results indicate that a small methyl group substituent in the luciferin's benzothiazole ring does not disrupt the activity of wild-type beetle luciferases but enhances light intensity, red-shifted BL, and provided greater light stability—features that are highly advantageous for *in vitro*, *ex vivo*, and *in vivo* BL detection applications in the future.

Author contributions

P. W. and P. C. designed experiments; P. W., I. K., P. K., C. K., and D. S. performed experiments. I. K., W. J. and P. W. elucidated the structures of synthesized D-LH₂ analogues by QTOF-MS and NMR. P. W. and C. K. optimized the conditions of bioluminescence assays. P. W. characterized the synthesized D-LH₂ analogues. N. L. and A. P. performed molecular docking and MD simulations. P. W., I. K., P. K., W. J., C. K., D. S., A. P., N. L., S. V., R. T., S. M., J. S., Y. N. and P. C. performed the data analysis. P. C., R. T., S. M., J. S., R. N., K. N., Y. N., and Y. O. provided advice. P. W. and P. C. wrote the manuscript with suggestions and comments from all authors.

Conflicts of interest

A patent application related to synthesis of red-shifted luciferin analogues has been submitted to the Department of Intellectual Property, Thailand (TH 2301002330).

Data availability

The authors declare that all data supporting the findings of this study are available within the paper and its supplementary information (SI). Supplementary information: experimental procedures, tables and figures showing D-LH₂ analogue synthesis, yield optimization, structural elucidation and bioluminescence characterization, and molecular docking, and applications of D-LH₂ analogues in crude lysates and live mammalian cell lines. See DOI: https://doi.org/10.1039/d5cb00287g.

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