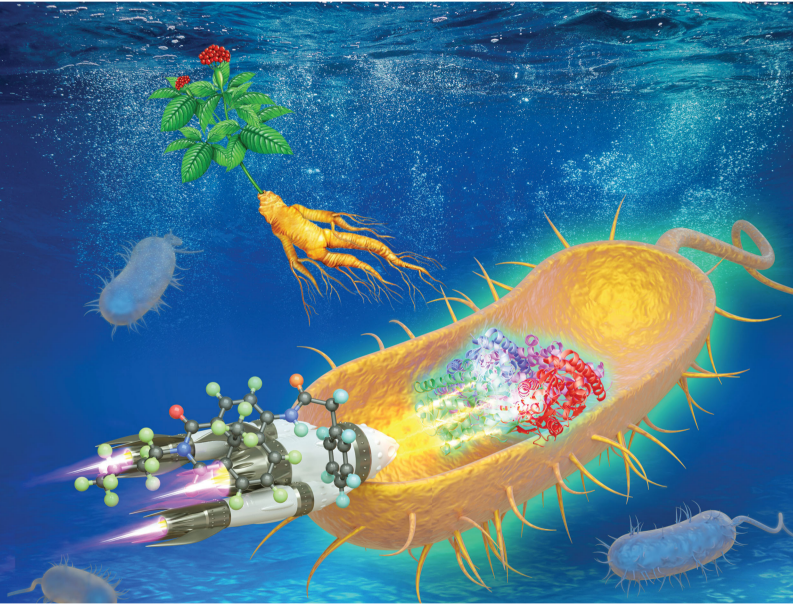
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Visualization of penicillin G acylase in bacteria and high-throughput screening of natural inhibitors using a ratiometric fluorescent probe†

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A ratiometric fluorescent probe (PNA) was developed to sense and image endogenous bacterial penicillin G acylase (PGA). Oleanolic acid was discovered as a potential natural inhibitor of PGA using high-throughput screening techniques based on PNA.

Penicillin G acylases (E.C. 3.5.1.11, PGA) mainly distribute in micro-organisms, belonging to the N-terminal hydrolase super family, which can catalyze the hydrolysis of β -lactam antibiotics producing 6-aminopenicillanic acid (6-APA), a key intermediate for the synthesis of various penicillins.^{1,2} Given the significant industrial importance of PGA, the expression of endogenous PGA in various bacteria has been evaluated using some common techniques, such as RT-PCR. In addition, penicillin acylases secreted by intestinal bacteria are one of the bile acid hydrolases, indicating the importance of penicillin acylases in the metabolism of bile acids, especially conjugated bile acids, by intestinal bacteria.³ Therefore, PGA belonging to penicillin acylases, not only mediates the production of 6-APA, but also contributes to the in vivo circulation of bile acid. Herein, on the basis of the hydrolase characteristics of PGA, we have developed an effective method for the real-time detection of endogenous bacterial PGA activity. In addition, we have investigated some potential inhibitors of PGA since such inhibitors could be used to interfere with the hydrolase activity of PGA in bacteria.

Fluorescent probes that are activated by biological enzymes with high selectivity and sensitivity, can be used for high-throughput

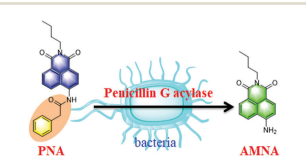
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analysis as a useful enzyme assay.⁴ 1,8-Naphthalimide derivatives have been widely used to sense biological enzymes in cells, microorganisms and even animals, due to excellent photophysicochemical properties and in many cases ratiometric fluorescence response which enables enhanced accuracy of measurement in complex systems.⁵ Recently, several fluorescent probes have been prepared to determine the PGA enzyme activity *in vitro*, based on deacylation of the phenylacetyl group from various fluorophores.⁶ However, few fluorescent probes have been developed to detect endogenous bacterial PGA activity, and image PGA in living bacterial cells.

In the present work, we designed and developed a ratiometric fluorescent probe *N*-(*n*-butyl)-4-*N*-phenylacetamide-1,8naphthalimide (**PNA**) for the real-time detection and imaging of bacterial PGA (Scheme 1). Additionally, a high-throughput screening system for PGA inhibitors has been established using **PNA**, and several inhibitors have been obtained from a natural compound library, in particular oleanolic acid with an IC₅₀ value 15.35 μ M, which could be used to block the hydrolysis of penicillins and conjugated bile acids by bacterial PGA.

PNA as a ratiometric fluorescent probe displayed fluorescence emission at 470 nm (Fig. S1, ESI⁺), after deacylation of the phenylacetyl moiety by PGA the *N*-(*n*-butyl)-4-amino-1,8-naphthalimide (**AMNA**) produced displays fluorescence emission at 540 nm (Fig. S1, ESI⁺). The hydrolysis of **PNA** mediated by PGA was determined by HPLC-DAD, and indicated the production of **AMNA** in a co-incubated solution of **PNA** with PGA (Fig. S2,



Scheme 1 Hydrolysis of **PNA** mediated by penicillin G acylase.

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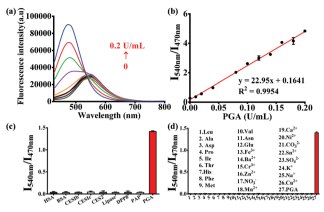


Fig. 1 (a) Fluorescence spectra of **PNA** (10 μ M) upon addition of increasing concentrations of PGA (0–0.2 U mL⁻¹). (b) Linear relationship between the fluorescence ratio (l_{540}/l_{470}) and PGA concentration. Fluorescence intensity ratio (l_{540}/l_{470}) of **PNA** upon addition of various enzymes and proteins (c), ions and amino acids (d). (PGA 0.05 U mL⁻¹, PBS : acetonitrile 2:1, v/v, λ_{ex} 400 nm).

ESI[†]). In addition, the interaction and hydrolysis mechanism between **PNA** and PGA was investigated using *in silico* docking analysis. As shown in Fig. S3 and S4 (ESI[†]), hydrogen bonds were observed between 4-NH and GLN B: 23 amino acid residue, the carbonyl group of the phenylacetyl moiety and the ALA B: 69 amino acid residue, which plays a key role in the hydrolysis of the amide bond of **PNA**. Importantly, PGA could efficiently catalyze the hydrolysis of **PNA** to produce **AMNA**.

The optimal incubation temperature for the enzymatic hydrolysis of PNA by PGA, was found to be 37 °C (Fig. S5, ESI†). We then evaluated the fluorescence responses of PNA towards PGA as a function of PGA concentration (Fig. 1a). A good linear relationship was observed for the fluorescence intensity ratio at different PGA concentrations $(0-0.2 \text{ U mL}^{-1})$ (Fig. 1b), indicating that the intensity ratio was closely related to the PGA activity. Furthermore, the selectivity of PNA toward PGA was evaluated in the presence of various species. As shown in Fig. 1c, general proteins and enzymes including human serum albumin (HSA), bovine serum albumin (BSA), carboxylesterases (1b, 1c, and 2), lipase, dipeptidyl peptidase 8 (DPP8) and proline aminopeptidase (PAP) could not catalyze the hydrolysis of PNA. In addition, various amino acids and ions displayed no interference of the fluorescence intensity ratio (Fig. 1d). As an important parameter for this enzymatic reaction, the kinetic behavior of the hydrolysis of PNA mediated by PGA was investigated (Fig. S7, ESI[†]). A Michaelis-Menten kinetic model was observed for the hydrolysis experiment, with V_{max} 6.44 nmol min⁻¹ mg⁻¹ and K_{m} 9.12 μ M, indicating a high affinity of PNA towards PGA.

As a bacterial hydrolase, PGA has been reported to be expressed in several bacteria, such as lactobacillus.⁷ On the basis of hydrolase activity of endogenous bacterial PGA, **PNA** as a substrate of PGA was used to sense bacterial PGA and image bacteria. After screening several bacteria, *Acinetobacter baumannii* LMG 994 HAM, *Bacillus cereus* 994000168 LBK, *Brevibacillus parabrevis* 090915_03 LBK, and *Staphylococcus aureus* ssp. aureus DSM 799 DSM all expressed PGA. As shown in Fig. 2, the blue channel and green channel were monitored corresponding to the

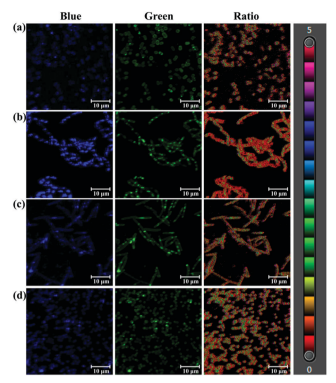


Fig. 2 Fluorescence images of bacteria incubated with **PNA** (50 μM). (a) *Acinetobacter baumannii* LMG 994 HAM (ratio 1.995). (b) *Bacillus cereus* 994000168 LBK (ratio 0.41). (c) *Brevibacillus parabrevis* 090915_03 LBK (ratio 1.135). (d) *Staphylococcus aureus* ssp. aureus DSM 799 DSM (ratio 0.842). Blue channel: λ_{ex} 405/ λ_{em} 425–459 nm, green channel: λ_{ex} 405/ λ_{em} 525–575 nm. Scale bar 10 μm.

existence of **PNA** and **AMNA** in these bacteria. Importantly, the ratiometric nature of **PNA**, facilitates the precise determination of endogenous PGA activity for various bacterial species. According to the fluorescence intensity ratio values, it was found that *Acinetobacter baumannii* (ratio 1.995) expressed more PGA than other bacteria. Therefore, **PNA** as a ratiometric fluorescent probe could be used to sense and image endogenous bacterial PGA, which would be helpful for the characterisation of bacterial PGA and the investigation of the biological function of bacterial PGA.

Most kits for the activity assay and inhibitor screening of biological enzymes work on the basis of OD values at specific wavelengths for the designed chromogenic reagents, which show low sensitivity. Herein we used **PNA** with higher sensitivity to assay PGA activity and establish a high-throughput screening system to rapidly discover inhibitors for **PNA**.

In the present work, 36 natural compounds with a variety of chemical skeletons (*e.g.* flavones, coumarins, monoterpenoids, diterpenoids, triterpenoids, steroids, alkaloids, and polyphenols) from a natural compound library have been evaluated for their inhibitory effects on PGA (Fig. S12 and Table S1, ESI[†]). Compared with the control group (Con.), weak fluorescence intensity was observed for the wells corresponding to compounds **13** and **21**, indicating potential PGA inhibition (Fig. 3a). In comparison with the control group, the relative fluorescence intensity and remaining PGA activity of each well were determined as shown in Fig. 3b. As a result, compound **13** (oleanolic acid) was tested as a PGA inhibitor, and found to have an

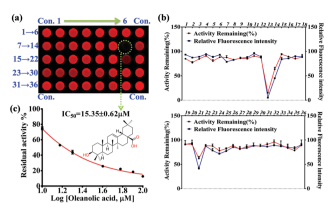


Fig. 3 Discovery of an inhibitor for PGA from a natural compound library using an established high-throughput screening system. (a) Fluorescence image for the inhibitor screening in plates against PGA measured by Amersham Typhoon RGB. (b) The fluorescence intensity and remaining activity of PGA for each plate corresponding to different compounds (1–36). (c) The inhibitor effect of compound 13 (oleanolic acid) against PGA.

 IC_{50} value of 15.35 \pm 0.63 μ M (Fig. 3c). Thus, using our high-throughput screening system, oleanolic acid was determined to be a potential PGA inhibitor from the natural compound library.

The selectivity and specificity of sensing endogenous bacterial PGA by **PNA** has been elucidated by fluorescence imaging experiments using PGA expression regulation and the inhibitor oleanolic acid. In the imaging experiment of bacteria, the PGA plasmid was constructed and transfected into *E. coli*, which led to the preparation of an *E. coli* strain with high expression of PGA (*Escherichia coli*-PGA⁺). As shown in Fig. 4, a strong fluorescence intensity in the green channel was observed for *E. coli*-PGA⁺, which suggests that the fluorescence imaging of *E. coli* was PGA activity dependent.

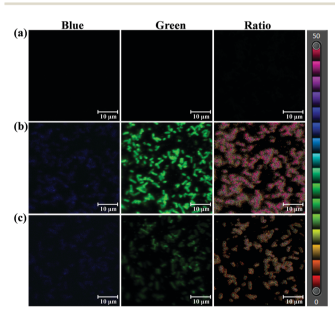


Fig. 4 Confocal microscopic images of *Escherichia coli* incubated with **PNA** (50 μ M). (a) Blank group. (b) *Escherichia coli*-PGA⁺ with PGA high expression. (c) *Escherichia coli*-PGA⁺ incubated with the inhibitor oleanolic acid (50 μ M). Blue channel: λ_{ex} 405/ λ_{em} 425–459 nm, green channel: λ_{ex} 405/ λ_{em} 525–575 nm. Scale bar 10 μ m.

Additionally, the co-incubation of oleanolic acid (50 μ M) and **PNA** in *E. coli*–PGA⁺ culture inhibited the production of **AMNA** according to the weak fluorescence intensity from the green channel. The ratio images confirmed high expression of PGA in *E. coli*–PGA⁺ and the strong inhibitory effect of oleanolic acid towards PGA.

In summary, based on the hydrolase properties of PGA, a ratiometric fluorescent probe **PNA** has been successfully designed and developed for sensing and imaging bacterial PGA. Using **PNA** for PGA activity assay, a high-throughput screening system has been established for the discovery of new PGA inhibitors, and oleanolic acid was discovered as a potential PGA inhibitor from a natural compound library, capable of interfering in the hydrolysis of penicillins and conjugated bile acids by bacterial PGA.

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Conflicts of interest

There are no conflicts to declare.

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