Structural Investigation of Park’s Nucleotide on Bacterial Translocase MraY: Discovery of Unexpected MraY Inhibitors

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Systematic structural modifications of the muramic acid, peptide, and nucleotide moieties of Park’s nucleotide were performed to investigate the substrate specificity of *B. subtilis* MraY (MraY<sub>BS</sub>). It was found that the simplest analogue of Park’s nucleotide only bearing the first two amino acids, l-alanine-iso-d-glutamic acid, could function as a MraY<sub>BS</sub> substrate. Also, the acid group attached to the Cα of iso-d-glutamic acid was found to play an important role for substrate activity. Epimerization of the C4-hydroxyl group of muramic acid and modification at the 5-position of the uracil in Park’s nucleotide were both found to dramatically impair their substrate activity. Unexpectedly, structural modifications on the uracil moiety changed the parent molecule from a substrate to an inhibitor, blocking the MraY<sub>BS</sub> translocation. One unoptimized inhibitor was found to have a $K_i$ value of $4 \pm 1$ μM against MraY<sub>BS</sub>, more potent than tunicamycins.

Peptidoglycan is a polymer consisting of sugars and amino acids that forms the bacterial cell wall. Interrupting the biosynthesis of peptidoglycan can devastate bacterial growth and survival due to the critical role it plays in maintaining cell shape and protecting bacteria from internal osmotic pressure<sup>1,2</sup>. One of the enzymes involved in bacterial cell wall biosynthesis, MraY is an integral membrane protein that catalyzes the transfer of the monophospho-MurNAc-pentapeptide moiety from Park’s nucleotide (UDP-MurNAc-pentapeptide) onto the undecaprenyl phosphate, to give Lipid I with concomitant release of UMP (Fig. 1). MraY is an attractive antibacterial target being essential for bacterial growth; highly conserved across many bacterial species; and without a eukaryotic counterpart<sup>3–6</sup>.

One major class of MraY inhibitors, known as nucleoside antibiotics, shares a uridine nucleoside as a common moiety with Park’s nucleotide<sup>6–10</sup>. Accordingly, an understanding of the interactions between Park’s nucleotide and MraY might be useful for the design of new MraY inhibitors. Recent disclosure of an apo crystal structure of MraY from *Aquifex aeolicus* (MraY<sub>AA</sub>) shows the overall architecture of this interesting enzyme<sup>11</sup>. However, due to the lack of available complex crystal structure, detailed mechanisms or interactions between substrates or inhibitors toward MraY remain to be explored. Although some brief substrate studies of Park’s nucleotide toward MraY have been reported, their scope is limited to the structural diversity accessible by biocatalysis<sup>12</sup>. Obviously, the substrate study of MraY is hampered by difficulties to acquire the structurally complex substrates. Chemical synthesis seems to be the most straightforward approach towards the generation of pure and systematically modified samples of various desired molecules for testing against MraY.

To more thoroughly investigate how structural modification of Park’s nucleotide affects MraY substrate recognition, we first sought to identify a proper polypropyl phosphate substrate that would be conserved for all the Park’s nucleotide analogues tested. In our preliminary HPLC-based MraY activity study, NBD-Park’s nucleotide<sup>6</sup> was completely consumed in 1 h when undecaprenyl phosphate (C55P) was applied as a polypropyl phosphate substrate in our hands (Supplementary Figure 1)<sup>13</sup>. In contrast, other polypropyl phosphates with a shorter length or different configurations still can be recognized as a MraY substrate but their substrate activity is much weaker than undecaprenyl phosphate (C55P) (Supplementary Table 1). Our observation of this broad substrate specificity of MraY is consistent with previous studies in the combined Mra/Y-MurG system or membrane fractions.
containing both MraY and MurG\textsuperscript{13,14}. According to our results, C55P was chosen as the substrate coupling partner for all the Park's nucleotide analogues studies, and the substrate activity was measured after 1 h reaction for convenient purposes. Moreover, it was decided not to modify the pyrophosphate group as it is at this position that translocation occurs.

Herein, we describe the systematic preparation of Park's nucleotides with varying three parts including the peptide, \(N\)-substituted muramic acid, and uridine moieties for evaluation as MraY\textsubscript{BS} substrates (Fig. 2). This information will provide us with the essential moieties and the specificity requirements of the MraY for Park's nucleotide analogues, as an effort toward development of new inhibitors.

Results and Discussions
Preparation of Park's nucleotide analogues and evaluation of their substrate activity. As shown in Fig. 3, O-debenzylation of 1 followed by a phosphorylation and phosphitylation/oxidation sequence gave the phosphate 2 in 71% yield over three steps\textsuperscript{17}. Compound 4 was obtained via the debenzylation of 2. Finally, conjugation of 4 with activated UMP-morpholine-\(N,N'\)-dicyclohexyl carbamidine salt and global deprotection under basic conditions gave Park's nucleotide 9 in 69% yield. For the preparation of 5, selective deprotection of the trimethylsilyl ethyl ester (TMSE) in 2 by treatment with TBAF in THF, followed by
coupling with H-\textit{d}-\textit{iso}-Glu(OMe)-l-Lys-(TFA)-d-Ala-d-Ala(OMe) and debenzylation gave the corresponding in 67% yield over three steps. Compound 3 was then coupled with activated UMP-morpholine-\textit{N,N}\textquotesingle-dicyclohexylcarboxamidine salt, followed by global deprotection under basic conditions to give Park's nucleotide in 35% yield over two steps. A fluorescent probe was prepared from 5 by conjugating a nitronbenzoxadiazole (NBD) fluorophore at the terminal amine site of lysine on the peptide stem in 88% yield. Compounds and were similarly prepared (Fig. 3).

The substrate activity study of 5–10 toward \textit{MraY} was performed using the HPLC-based \textit{MraY} functional assay. Substrate consumption curves of 5–10 were shown in Fig. 4A. Compounds 5–8 were recognized as a \textit{MraY} substrate, but 9 and 10 were not. The similar curves of 5 and 6 suggest that the NBD-fluorophore attached to the side chain of Lys on the pentapeptide stem of Park's nucleotide does not cause any significant effect.
on its substrate activity (Supplementary Figures 1 and 2). Compound 7, lacking the terminal two amino acids (D-Ala-D-Ala), was only slightly less active than 5 (17% activity reduced after 1 h reaction, Fig. 4B), showing that the D-Ala-D-Ala moiety is not essential for MraYBS recognition. The previous study reported by Hammes and Neuhaus pointed out that 7 is a much weaker substrate than 5 when intrinsic membrane fractions are used as a source of lipophosphate and enzyme. In our conditions, only the purified enzyme and two pure substrates were utilized, and Park’s nucleotide analogue was the limiting reagent compared to the other substrate C55P. Both individual studies show different degrees of the substrate activity loss that might be attributed to several factors such as enzyme activity, substrate ratio and assay platform. Compound 8, similar to 5 but lacking the terminal three amino acids, was a weak substrate (40% activity remained after 1 h reaction, Fig. 4B). Moreover, 9 (bearing only one amino acid (l-Ala)) and UDP-GlcNAc (10) were not substrates under these assay conditions, showing that this 3-O-lactyl-tripeptide (D-Lac-l-Ala-γ-D-Gln-l-Lys) moiety in Park’s nucleotide is important for the MraYBS catalyzing process.

Next, more subtle structural changes of Park’s nucleotide 5 were proposed, and the resultant molecules conjugated with a NBD fluorophore on the peptide stem for easy monitoring (Fig. 5A). All analogues except 17 were synthesized in a manner similar to that for 5. Initial attempts to prepare 17 by coupling of 3 and morpholine-activated 5-amino-uridine-5′-monophosphate in the presence of 1H-tetrazole were unsuccessful. Most of the morpholine-activated 5′-NH2-UMP was found to degrade into 5′-NH2-UMP, and only trace amount of product was detected in the reaction mixture. To overcome this problem, the synthetic strategy was re-designed to entail activation of the sugar moiety with the carbonyl diimidazole (CDI) instead of activation of 5-amino-uridine-5′-monophosphate, followed by global deprotection and the NBD labeling. In this way, 17 was obtained in a yield of 31% over four steps (see also Supplementary Methods).

As illustrated in Fig. 5B, both N-glycolyl 12, the natural substrate for mycobacterial MraY (also called MurX), and unnatural N-glycinol 13 had similar substrate activity to 6, indicating that there are no extra interactions, such as additional hydrogen bonds, to increase the activity between the N-substituent moiety on muramic acid of Park’s nucleotide analogues and MraYBS. Analogue 14 (R4 = H) had similar activity to 6, suggesting the methyl group on the lactate moiety to be unessential. Likewise, 15 (R5 = H) was slightly less active than 6 (about 80% relative activity after 1 h reaction, Fig. 5C). Surprisingly, 16 (R6 = H) was found to be a very poor substrate compared to 6 (<10% relative activity after 1 h reaction, Fig. 5C), showing the acid group attached to the CO of iso-D-Glu moiety in Park’s nucleotide to be critical.

In order to evaluate the effects of peptide moiety of Park’s nucleotide, investigation of the binding affinity of 5, 6, 9 and 16 was performed using a biolayer interferometry-based binding (BLI) assay. Initial attempts to perform the MraYBS binding assay in the presence of both substrates (C55P and Park’s nucleotide analog) didn’t work properly because a strong non-specific binding signal was observed; presumably, the hydrophobic part of C55P might mainly contribute this non-specific interaction. To simplify the assay conditions, only Park’s nucleotides were utilized to measure the binding affinity with MraYBS. As shown in Fig. 6, compounds 5 and 6 exhibited similar binding affinity with Kd values of 120 and 127 μM, respectively. This suggests the NBD tag in 6 does not affect the binding affinity with MraYBS and this observation is consistent with the substrate activity result in Fig. 4. Structurally, 9 is the truncated form of 5 (lacking the outermost four amino acids, including iso-D-Glu); and 16 is very similar to 6 – the only difference being removal of the acid group attached to the CO of iso-D-glutamic acid (R6 = H). However, 9 and 16 showed no proper binding affinity with MraYBS – only a very low binding signal was detected, even at concentrations up to 500 μM. Our results indicate that the acid moiety (R6 = COOH) on iso-D-Glu of Park’s nucleotide plays an important role for both binding affinity and substrate activity. In addition, 11 (R1 = OH/R2 = H) and 17 (R2 = NH2) did not function as substrates, even under extreme reaction conditions, showing the equatorial hydroxyl group at R2 position to be critical and the modifications at R2 position not tolerated.

Figure 4. Evaluation of the substrate activity of Park’s nucleotide analogues 5–10 toward MraYBS. (A) The reactions were analysed in the HPLC-based MraY functional assay as described in Methods and the progresses were measured at 0, 10, 20, 30, 60 and 120 min. (B) The substrate activity of 5–10 was determined by the decrease in substrate consumption after 1 h reaction time. All experiments were repeated in triplicate (Supplementary Figure 3).
Construction of the molecular model of the Park's nucleotide- MraYBS complex. Based on these preceding results, as well as our mutagenesis (Table 1) and computational modeling studies, a putative Park's nucleotide binding site on MraYBS is proposed (Fig. 7)23,24. As illustrated in Fig. 7A, Park's nucleotide could specifically interact with the W297, K102, and Q271 of MraY, and the phosphate group of C55P. As shown in Table 1, the enzyme activities of four MraYBS mutants, including T53A, K102A, Q271A, and W297A, were significantly decreased, suggesting these residues to be important for enzyme activity. All four mutants had higher $K_M$ values compared with the wild-type MraYBS ($K_M = 18\mu M$ and NBD-Park's nucleotide applied as a substrate), showing that the mutations caused a loss of binding affinity. The highly conserved threonine (T53) located on loop A is close to the proposed catalytic pocket (Fig. 7B), and may participate in the enzyme process. In addition, the uracil moiety is embedded in a deep groove, which may interact with W297 on loop E of MraYBS. Our substrate specificity and site-directed mutagenesis study strongly suggest that Q271 on MraYBS might interact with the iso-d-glutamic acid of Park's nucleotide through a hydrogen bond to stabilize the peptide chain.

Discovery of Park's nucleotide analogues bearing modifications at the uracil 5-position as MraYBS inhibitors. We were curious whether analogues 11 (modified at R1/R2) and 17 (modified at R7) – neither of which were active substrates – could inhibit the function of MraYBS. To further evaluate the role of positions at R1, R2 and R7, we re-designed and synthesized 18–21 with a truncated peptide (Fig. 8). The inhibitory activity of each of these was determined using a fluorescent enhancement assay against MraYBS, with tunicamycin as reference (Supplementary Figure 5). As shown in Table 2, 18 ($R^7 = NH_2$) had no inhibition activity, but 19 ($R^7 = NHAc$) showed weak inhibitory activity ($K_i = 764\mu M$) against MraYBS; and 20, bearing a p-tolylacetamide moiety at R7, became a more potent inhibitor ($K_i = 11\mu M$) – strongly indicating that appropriate N-substitution ($R^7$) can enhance inhibitory activity. However, 21, the C4-hydroxyl epimer of 20, was a very poor inhibitor (30% inhibition at 1 mM). This finding implied the C4-hydroxyl epimerization of Park's nucleotide dramatically impairs both the substrate and inhibitory activities. To improve the inhibitory activity, we reinstalled the tetrapeptide

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Figure 5. Evaluation of the substrate activity of Park's nucleotide analogues 6 and 11–17 toward MraYBS. 
(A) The chemical structures of synthetic Park's nucleotide analogues (11–17) are shown. (B) The reactions were analyzed in the HPLC-based MraY functional assay as described in Methods, and the progresses were measured at 0, 10, 20, 30, 60 and 120 min. (C) The substrate activity of 6 and 11–17 was determined by the substrate consumption after 1 h reaction time. All experiments were repeated in triplicate (Supplementary Figure 4).
moiety on 20 to give 22, behaving as a competitive inhibitor with the \( K_i \) value of 4 µM toward MraYBS, approximately two-fold more potent than tunicamycins (\( K_i = 9 \) µM). Our results indicated that (1) analogues 19, 20, and 22 all function as MraY BS inhibitors, even though they all contain a pyrophosphate moiety; and (2) inhibitory potency can be significantly improved through modification of the substituent at the 5-position of the uracil; modification of the oligopeptide moiety can moderately increase the inhibition activity.

To understand the binding contribution derived from structural modifications, 19–22 were used to evaluate their binding affinity toward MraYBS in the BLI assay (Table 2 and Supplementary Figure 6). The results are interesting. For example, moderate binding signals were revealed after acetylation of 18, and the binding affinity of 20 (\( R = \text{p-tolylacetamido group}, K_D = 197 \) µM) was approximately 1.5 fold stronger than that of 19 (\( R = \text{acetamido group}, K_D = 281 \) µM). From the modeled complex structure, both inhibitors could exert an additional H-bond interaction with the K226 in the hydrophobic cleft formed by P52, K226, F228, and W297 (refer to Fig. 7B) to compensate the losing interaction with Q271. Although the binding affinity of 19 and 20 is subject to the modification at the 5-position of the uracil, there is no obvious reason to explain why their inhibition abilities (\( K_i \) values in Table 2) vary by 70 fold. Presumably, the p-tolylacetamido substituent at the 5-position of the uracil might further affect the interaction with MraYBS by inducing an enzyme conformation change – a detailed molecular dynamic simulation of Park's nucleotide and inhibitors toward MraYBS remains to be performed. Compound 21 had no sufficient binding interactions with MraYBS even at the concentration up to 250 µM, emphasizing that the

| Proteins | \( K_D \) (µM) | \( V_{max} \) (FLU/min) | Relative activity (%) |
|----------|----------------|-------------------------|----------------------|
| WT       | 18 ± 1         | 18 ± 0.8                | 100                  |
| T51A     | 24 ± 2         | 5 ± 0.2                 | 60                   |
| T53A     | 70 ± 11        | 7 ± 0.8                 | 18                   |
| K102A    | 49 ± 9         | 5 ± 0.2                 | 33                   |
| K226A    | 24 ± 3         | 23 ± 4.7                | 100                  |
| F228A    | 18 ± 2         | 10 ± 1.5                | 98                   |
| Q271A    | 72 ± 11        | 6 ± 1.7                 | 16                   |
| R281A    | 20 ± 1         | 10 ± 1.4                | 81                   |
| W297A    | 63 ± 9         | 4 ± 0.4                 | 17                   |

Table 1. Evaluation of kinetic parameters of mutant MraYs. *Kinetic parameters were determined by fluorescent enhancement assay with using 6 as the substrate. FLU: fluorescent units. The assay was determined by HPLC-based MraY functional assay with using 6 as the substrate. Relative activity was determined by monitoring the consumption of 6 in one-hour reaction.
Figure 7. Docked pose of Park’s nucleotide 5 in the MraY binding site. (A) Surface view of the substrate binding site of MraY and the docked ligand poses. (B) Cartoon view of the MraY activity site and the docked ligand poses. TM refers to transmembrane domain. The carbons of ligand (Park’s nucleotide 5) are colored in light blue and the carbons of protein are colored in brown. C55P is shown by a ball style representation. Amino acid residues of the protein are labeled in one-letter code; residues of the ligand are labeled by three-letter code.

Figure 8. Structures of Park’s nucleotide analogues 18–22.

| Entry | Compounds | $K_i$ ($\mu$M)$^a$ | Inhibition Mode$^a$ | $K_D$ ($\mu$M)$^b$ |
|-------|-----------|--------------------|-------------------|------------------|
| 1     | 18        | —$^d$              | —$^d$             | Not determined$^e$ |
| 2     | 19        | 764 ± 127          | Competitive       | 281 ± 97         |
| 3     | 20        | 11 ± 3             | Competitive       | 197 ± 47         |
| 4     | 21$^*$    | —$^d$              | —$^d$             | NB$^f$           |
| 5     | 22$^g$    | 4 ± 1              | Competitive       | 86 ± 12          |
| 6     | Tunicamycins | 9 ± 1              | Competitive       | 93 ± 16          |

Table 2. Evaluation of the inhibitory activity and the binding affinity of Park’s nucleotide derivatives (18–22) toward MraY$_{bs}$. $^a$The $K_i$ and inhibition mode were determined by fluorescent enhancement assay. $^b$The dissociation constants ($K_D$) were determined by BLI assay. $^c$No inhibitory activity was observed at 1 mM for 18. The $K_D$ value of 18 is not determined. $^d$The $K_i$ and inhibition mode cannot be measured due to the low inhibitory activity. $^e$Only 30% inhibition was observed at 1 mM for 21. NB refers to no significant binding signal at 250 $\mu$M. $^g$Compound 22 was not a MraY$_{bs}$ substrate even under extreme conditions.
orientation of the C4-hydroxyl group of muramic acid plays an essential role in molecule-enzyme recognition. Notably, the binding affinity of 22 \( (K_i = 86 \mu M) \) was approximately 2.3 fold stronger than that of 20, which can be attributed to the contribution of the tetrapeptide moiety.

The antibacterial activities of 20 and 22 were also investigated and the minimal inhibitory concentrations (MIC) against S. aureus and B. subtilis were determined using standard-broth dilution methods\(^\text{25}\). Unfortunately, both 20 and 22 showed no antibacterial activity, even at a high concentration of 200 \( \mu \text{M} \). It may be because the compounds containing the highly charged pyrophosphate moiety were difficult to penetrate the bacterial cytoplasmic membrane\(^\text{26}\). In order to improve the antibacterial activity, finding a surrogate to replace the pyrophosphate moiety on 22 remains to be explored.

**Conclusions**

A series of Park’s nucleotide analogues with modifications at the peptide, muramic acid, and nucleotide moieties has been designed and synthesized, and their MrAY\(_{BS}\) substrate activity and specificity were evaluated. Our results led to several important findings: (1) the first two amino acids (L-alanine-\( \text{iso-}d\)-glutamic acid) of the oligopeptide chain are essential for MrAY\(_{BS}\) recognition; (2) the configuration of the C4-OH on muramic acid is important for MrAY\(_{BS}\) substrate specificity; and (3) modifications at the 5-position of the uracil dramatically impair the substrate activity (Fig. 9). Also, the substrate specificity data together with mutagenesis and computational modeling studies allowed us to infer a putative Park’s nucleotide binding site on MrAY\(_{BS}\).

Unexpectedly, analogues bearing modifications at the 5-position of the uracil were found to be MrAY\(_{BS}\) inhibitors though these molecules contain a pyrophosphate moiety. Of these, an unoptimized inhibitor 22 \( (K_i = 4 \mu \text{M}) \) was found to be roughly twice as potent as tunicamycins \( (K_i = 9 \mu \text{M}) \) against MrAY\(_{BS}\), the first example of Park’s nucleotide-based inhibitors. However, our results not only allow us to infer the minimal structure requirements of Park’s nucleotide as a MrAY substrate, but also illuminate a new direction for MrAY inhibitor design. And more generally, the concrete nature of these conclusions validate our strategy of systematic substrate structure modifications for the elucidation of enzyme binding site mapping, for membrane proteins without available co-crystal structures. Investigation of the MrAY mechanisms, and development of more potent inhibitors with \textit{in vivo} antibacterial activity are currently ongoing in our laboratory.

**Methods**

**General.** All chemicals were obtained from commercial suppliers and used without further purification. All solvents were anhydrous grade unless indicated otherwise. All non-aqueous reactions were performed in oven-dried glassware under a slight positive pressure of argon unless otherwise noted. Reactions were magnetically stirred and monitored by thin-layer chromatography on silica gel. Flash chromatography was performed on silica gel of 40–63 μm particle size. Concentration refers to rotary evaporation. Yields are reported for spectrscopicily pure compounds. NMR spectra were recorded on dilute solutions in D\(_2\)O, CDCl\(_3\), and CD\(_3\)OD on Bruker AVANCE 600 at ambient temperature. Chemical shifts are given in δ values and coupling constants \( J \) are given in Hz. The splitting patterns are reported as \( s \) (singlet), \( d \) (doublet), \( t \) (triplet), \( q \) (quartet), \( m \) (multiplet), and \( dd \) (double of doublets). High resolution ESI mass spectra were recorded on a Bruker Daltonics spectrometer. Compound 7–22 were prepared as described in supplementary Methods. Tunicamycins were purchased from Sigma-Aldrich (St. Louis, MO).

**Chemistry.** Compound 2. A mixture of 1 (200 mg, 0.31 mmol) and 20% \( \text{Pd(OH)}_2/C \) (20 mg) in THF (10 mL) was stirred at RT for 24 h under a hydrogen atmosphere. The reaction mixture was filtered through a pad of celite and the filtrate was concentrated to give a crude intermediate as white solid. The intermediate was used directly without further purification. The solution of intermediate, 1H-tetrazole (43 mg, 0.62 mmol) and dibenzyl (\( N,N\)-diisopropyl) phosphoramidite (213 mg, 0.62 mmol) in CH\(_2\)Cl\(_2\)/ACN (10 mL, v/v = 1:1) was stirred at 0°C for 2 h. The reaction was then cooled to −40°C and treated with \textit{tert}-butylhydroperoxide (0.6 mL, 6.0 mmol). The reaction was allowed to warm to RT over a period of 1 h. The mixture was diluted with CH\(_2\)Cl\(_2\) (50 mL), and extract with water (20 mL \times 2). The organic layers were collected, dried over MgSO\(_4\), concentrated, and purified.
by cc (EtOAc/CH2Cl2 = 1:2, silica gel) to afford 2 as a colorless oil (177 mg, 0.22 mmol, 71% over three steps). 1H NMR (600 MHz, CDCl3): δ 7.26–7.31 (m, 10 H), 6.72 (d, 1 H, J = 7.2 Hz), 6.58 (d, 1 H, J = 9.0 Hz), 5.61 (dd, 1 H, J = 5.4 and 6.0 Hz), 4.96–5.05 (m, 5 H), 4.31–4.34 (m, 1 H), 4.23–4.26 (m, 1 H), 4.11–4.14 (m, 2 H), 4.02 (dd, 1 H, J = 4.8 and 13.2 Hz), 3.84–3.92 (m, 3 H), 3.49 (dd, 1 H, J = 9.0 and 10.2 Hz), 2.01 (s, 3 H), 1.94 (s, 3 H), 1.71 (s, 3 H), 1.35 (d, 3 H, J = 7.2 Hz), 1.25 (d, 3 H, J = 6.6 Hz), 0.91–0.93 (m, 2 H), −0.03 (s, 9 H). 13C NMR (150 MHz, CDCl3): δ 172.4, 172.0, 170.7, 170.6, 169.0, 135.2, 135.0, 128.9 (×2), 128.7 (×4), 129.0 (×2), 128.0 (×2), 96.7, 78.1, 76.6, 70.1, 69.9 (×2), 68.8, 63.8, 61.4, 53.0, 45.3, 48.1, 22.8, 20.7, 20.6, 18.8, 17.6, 17.1, −1.59 (×3); HRMS calcd for [C37H53N2O14PSiH+]+ 809.3081, found 809.3045.

Compound 3. A mixture of 2 (100.0 mg, 0.12 mmol) and 1.0 M TBAF in THF (0.24 mL, 0.24 mmol) was stirred at RT for 2 h. The reaction mixture was concentrated, and the residue was extracted with EtOAc (50 mL), washed with 1.0 N HCl(aq) (50 mL × 2) and water (50 mL × 2). The combined organic layers were dried over MgSO4, and concentrated to afford the monosaccharide intermediate as a colorless oil. The sugar intermediate was purified by cc (CH2Cl2/EtOAc/Water = 50:50:50, silica gel) to give a fluorescent product. The compound was subjected to 2D NMR analysis, the reaction mixture containing MraY BS (10 μg/mL) and NBD-X-OSu (3.3 mg, 8.6 μmol) in DMF/sat. NaHCO3(aq) (2 mL, v/v = 1:1) was stirred at RT for 4 h. The solvent was removed and the reaction mixture was purified by cc (iPrOH/NH4OH = 1:2, silica gel) to afford 3 as a reddish brown solid (11.0 mg, 7.7 μmol, 88%). The purity of 3 is higher than 95% by analytical anion-exchange HPLC (Supplementary Figure 7). HRMS calcd for [C25H32N3O8PSi−H+]+ 627.1981, found 627.1996.

Compound 4. A mixture of 2 (100 mg, 0.12 mmol) and 20% Pd(OH)2/C (10 mg) in MeOH (8 mL) was stirred for 1 h under a hydrogen atmosphere. The reaction mixture was filtered through a pad of celite, and the filtrate was concentrated and purified by cc (CHCl3/MeOH/H2O = 60:25:4, silica gel) to give 4 as white solid (57 mg, 0.09 mmol, 75%). 1H NMR (600 MHz, CDCl3): δ 5.08 (t, 1 H, J = 9.6 Hz), 4.16–4.32 (m, 8 H), 4.10 (d, 1 H, J = 12.6 Hz), 3.86 (t, 1 H, J = 9.6 Hz), 2.11 (s, 3 H), 2.06 (s, 3 H), 1.95 (s, 3 H), 1.40 (d, 3 H, J = 7.2 Hz), 1.31 (d, 3 H, J = 6.6 Hz), 1.01 (m, 2 H), 0.06 (s, 9 H); 13C NMR (150 MHz, CDCl3): δ 173.6, 173.5, 173.4, 173.1 (×2), 172.1, 171.9, 171.2, 170.8, 170.3, 157.5, 115.2, 93.9, 78.0, 77.8, 69.2, 68.5, 61.8, 54.4, 52.3, 53.8, 51.5, 51.4, 49.4, 49.8, 39.0, 30.8, 30.6, 28.1, 26.7, 22.7, 21.7, 19.6, 19.3, 17.8, 16.6, 16.3, 15.9; HRMS calcd for [C39H61F3N7O21P+Na]+ 1074.3502, found 1074.3518.

Compound 5. A mixture of 3 (84.0 mg, 0.08 mmol), UMP-morpholine-N,N′-dicyclohexyl-carboxamide salt (80.0 mg, 0.12 mmol), 1H-tetrazole (8.4 mg, 0.12 mmol) and 4 Å molecular sieves in anhydrous pyridine was stirred at RT for 24 h under argon. The reaction mixture was concentrated and purified by cc (CHCl3/MeOH/H2O = 60:25:4, silica gel) to afford a crude mixture of 5. The intermediate was dissolved in a solution of 1.0 M LiOH(aq)/MeOH (2 mL, v/v = 1:1) and stirred at RT for the global deprotection. After stirring for 4 h, the reaction mixture was neutralized by 1.0 N HCl(aq) and concentrated, and purified by cc (iPrOH/NH4HCO3 = 2/1, silica gel) to give 5 as white solid (32 mg, 0.027 mmol, 35% over two steps). 1H NMR (600 MHz, CDCl3): δ 7.94 (d, 1 H, J = 8.4 Hz), 5.94–5.96 (m, 2 H), 5.45 (dd, 1 H, J = 3.6 and 7.2 Hz), 4.03–4.36 (m, 15 H), 3.91–3.94 (m, 1 H), 3.80–3.87 (m, 2 H), 3.77 (t, 1 H, J = 10.2 Hz), 3.62 (dd, 1 H, J = 9.0 and 10.2 Hz), 2.98 (m, 2 H), 2.28 (t, 1 H, J = 7.8 Hz), 2.12–2.14 (m, 1 H), 1.98 (s, 3 H), 1.65–1.89 (m, 5 H), 1.31–1.44 (m, 12 H). HRMS calcd for [C50H53N3O22P2+H]+ 1150.3589, found 1150.3644.

Compound 6. A mixture of Park’s nucleotide 5 (10.0 mg, 8.7 μmol) and NBD-X-OSu (3.3 mg, 8.6 μmol) in a solution of DMF/sat. NaHCO3(aq) (2 mL, v/v = 1:1) was stirred at RT for 2 h. The solvent was removed and the reaction mixture was purified by cc (iPrOH/NH4HCO3 = 2/1, silica gel) to give a fluorescent product. The compound was further purified by a semi-preparative reverse-phase HPLC (ZORBAX RX-C18 column, 5 μm, 9.4 × 250 mm) with gradient elution of 0.25 M NH4HCO3/15% MeOH in water (100:0 to 100) at a flow rate of 1 mL/min over 45 min, to give compound 6 as a reddish brown solid (11.0 mg, 7.7 μmol, 88%). The purity of 6 is higher than 95% by analytical anion-exchange HPLC (Supplementary Figure 7). HRMS calcd for [C50H37N3O22P2+H]+ 1426.4447, found 1426.3308.

**Biology.** HPLC-based MraY functional assay. The purified MraY from *Bacillus subtilis* was prepared as described in our previous reports16. For 5 and 7–10, the reaction mixture containing MraY 6 (10 μg/mL), C6P (200 μM) in 40 mM reaction buffer (30 mM Tris, 10 mM MgCl2, 10 mM NaCl, 0.1 mM Tween-20, 2.5% DMSO, pH 8.0) were pre-incubated at 37 °C for 15 min. The reactions were initiated by the addition of Park’s nucleotide analogues (10 μM) and incubated at 37 °C for 1 h. The reaction mixtures were added with uridine (0.01 mg/mL) as the internal standard and heated to 100 °C to stop the transferring reaction. The samples were analyzed by RP-C18 HPLC with gradient elution of 0.25 M NH4HCO3/15% MeOH in water (100:0 to 100) at a flow rate of 1 mL/min over 45 min, and the peaks of substrates were monitored at UV 260 nm. For 6 and 11–17, the reaction
mixture containing MraYBS (10 μg/mL), C25P (200 μM), 6-(7-Nitro-2,1,3-benzoxadiazol-4-ylamino)hexanoic acid (5 μM, internal standard) in 10 μL reaction buffer (30 mM Tris, 10 mM MgCl2, 10 mM NaCl, 0.1 mM Tween-20, 2.5% DMSO, pH 8.0) were pre-incubated at 37 °C for 15 min. The reactions were initiated by the addition of NBD-Park’s nucleotide analogues (10 μM) and incubated at 37 °C for 1 h. The reaction mixtures were heated to 100 °C to stop the transferring reaction, and samples were analyzed by an anion-exchange column (SAX1, Supelco Co., 5 μm, 4.6 × 250 mm) with a linear gradient elution of NH2OAc (20 mM to 1 M in MeOH) at a flow rate of 1.0 mL/min over 30 min. The fluorescent substrates were monitored with λem 466 nm/λex 535 nm by fluorescence detector. In an extreme reaction condition, which is performed in reaction buffer (30 mM Tris, 10 mM MgCl2, 10 mM NaCl, 0.1 mM Tween 20, 2.5% DMSO, pH 8.0) containing Park’s nucleotides (10 μM), MraYBS (40 μg/mL) and C55P (1 mM) at 37 °C for 24 h, compounds 11, 17 and 22 were confirmed to not be a substrate.

**Biolayer interferometry-based (BLI) binding assay.** MraYBS binding was measured in a biolayer interferometry-based binding assay on Octet Red96® instrument (Fortebio, Inc.). The biotinylation of MraYBS was performed with using EZ-Link® Sulfo-NHS-LC-LC-Biotin Kit (Thermo Fisher Scientific, Inc.). In general, MraYBS (1.7 mg/mL) in 50 μL reaction buffer (0.01 M HEPES pH 7.4, 0.15 M NaCl, 0.005% v/v Surfactant P20), was added by the sulfo-NHS-LC-LC-biotin reagent (1 mM, 1 μL). After incubating for 30 minutes at room temperature, the biotinylated-MraYBS was purified by a Zeba™ Spin Desalting Columns, 7K MWCO (Thermo Fisher Scientific, Inc.), and the buffer was changed to the binding buffer (30 mM Tris, 10 mM MgCl2, 10 mM NaCl, 0.1 mM Tween-20, pH 8.0) during the purification. Comparing to the non-biotinylated-MraYBS, the biotinylated-MraYBS remains 62% activity measured by the HPLC-based MraY functional assay in 1 h reaction. The biotinylated MraYBS was then immobilized to Super Streptavidin (SSA) Octet tips (Fortebio, Inc.) for reaching a fixed signal of 3 nm. Before the assay starting, the MraYBS-labeled SSA sensors were pre-soaked in binding buffer (30 mM Tris, 10 mM MgCl2, 10 mM NaCl, 0.1 mM Tween-20, pH 8.0) containing Park’s nucleotides (10 μM) and MraYBS (40 μg/mL) and C55P (1 mM) at 37 °C for 24 h, compounds 11, 17 and 22 were confirmed to not be a substrate.

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Author Contributions
K.-T.C. planned the experiment, and W.-C.C. supervised the experiment. K.-T.C., P.-T.C., C.-K.L. and Y.-F.C. synthesized compounds. T.-J.R.C., L.-Y.H. and C.-M.H. prepared enzymes. Y.-T.W. and H.-T.H. constructed computational model. K.-T.C., P.-T.C. and C.-M.H. performed measurements, and analysed the data. K.-T.C. and W.-C.C. wrote the manuscript. All authors commented on the results and manuscript.

Additional Information
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