Supporting Information for

Construction of the Octosyl Acid Backbone Catalyzed by a Radical S-Adenosylmethionine Enzyme and a Phosphatase in the Biosynthesis of High-carbon Sugar Nucleoside Antibiotics

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5. Supplementary References

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Figure S1. Examples of nucleoside antibiotics containing high-carbon (> 6 C's) sugar scaffold.
Figure S2. In-frame deletion of polJ or polH on pJTU4774.

(A) Schematic map for the in-frame deletion of polJ on pJTU4774 (left) and the PCR confirmation (right). M, 1 kb plus ladder; 1, WT; 2 and 3, samples corresponding to the numbered gene clusters to the left.

(B) Schematic map for the in-frame deletion of polH on pJTU4774 (left) and the PCR confirmation (right).
Figure S3. Bioassay and chemical analysis of the metabolites accumulated by $\Delta$polH or $\Delta$polJ mutants.

(A) Bioassay of the metabolites produced by related strains. 1, Metabolites produced by CXR14::pJTU4774; 2, Metabolites produced by CXR14::pJTU774/$\Delta$polH; 3, Metabolites produced by CXR14::pJTU774/$\Delta$polJ; 4, Metabolites produced by CXR14::pJTU2463 as negative control. *Trichosporon cutaneum* 3071 was selected as indicator fungi for the bioassay of polyoxins and related metabolites.

(B) HPLC analysis of the metabolites produced by related strains, CXR14 was the strain with the entire polyoxin gene cluster deleted, and this strain is facilitated for the genetic analysis of the target genes involved in polyoxin biosynthesis.

(C) LC-MS analysis of the compound X. This compound indicates a [M+H]$^+$ ion at $m/z$ 329.15 whose structure was further solved by comprehensive NMR analysis.
Figure S4. $^1$H NMR of compound X in D$_2$O (500 MHz).

$^1$H NMR (500 MHz, D$_2$O) δ 7.57 (1H, s), 5.93 (1H, d, $J$ = 5.0 Hz), 4.24 (1H, d, $J$ = 5.0 Hz), 3.83 (1H, dd, $J$ = 5.0, 5.0 Hz), 4.87 (1H, s), 4.57 (2H, d, $J$ = 5.0 Hz), 1.80 (3H, s).

Figure S5. $^{13}$C NMR of compound X in D$_2$O (125 MHz).

$^{13}$C NMR (125 MHz, D$_2$O) δ 166.40, 166.38, 151.79, 149.11, 137.46, 111.70, 98.15, 89.05, 81.94, 75.46, 71.75, 60.75, 11.45.
Figure S6. DEPT NMR of compound X in D$_2$O.

Figure S7. $^1$H-$^1$H COSY of compound X in D$_2$O.
Figure S8. HMQC of compound X in D$_2$O.

Figure S9. HMBC of compound X in D$_2$O.
Proteins include PolJ (S. aureochromogenes YB172), NikL (S. tendae Tu901), scontig008-78 S. chromofuscus ATCC49982, STRAU_0240 (S. aurantiacus JA4570), and XSR1_370020 (Xenorhabdus szentirmai DSM16338). The sequences were aligned with ClustalW and merged with ESpript online program.
Proteins include PolH (*S. aureochromogenes* YB172), NikJ (*S. tendae* Tu901), sccontig008-80 *S. chromofuscus* ATCC49982, STRAU_0246 (*S. aurantiacus* JA4570), and XSR1_370015 (*Xenorhabdus szentirmaii* DSM16338). The sequences were aligned with ClustalW and merged with ESPript online program.
Figure S12. Overexpression, purification, SDS-PAGE and ESI-MS/MS analysis of purified PolJ.

(A) Overexpression and purification of PolJ protein. A culture of *E. coli* BL21(DE3) cells harboring the pJTU2864 construct were grown in the LB medium containing 50 µg/mL of kanamycin at 37 °C with shaking (200 rpm) until the OD₆₀₀ reached ~0.5. Protein expression was then induced by addition of isopropyl β-D-1-thiogalactopyranoside (IPTG) to a final concentration of 0.5 mM, and the cells were allowed to grow at 18 °C and 200 rpm for an additional 18 h. The cells were harvested by centrifugation at 4500 × g for 15 min. All purification steps were carried out at 4 °C using Ni-NTA resin and the PolJ protein were washed and eluted by 20-250 mM imidazole in 25 mM Tris-HCl, 150 mM NaCl buffer (pH 8.0). The collected protein solution was dialyzed against 25 mM Tris-HCl, 150 mM NaCl buffer (pH 8.0) for three times. The protein solution was then flash-frozen in liquid nitrogen and stored at -80 °C until use. Protein concentration was determined by the Bradford assay using bovine serum albumin as the standard.

(B) SDS-PAGE analysis of purified PolJ.

(C) ESI-MS/MS analysis the unique peptides from the purified PolJ digested by Trypsin.
Figure S13. Purification, reconstitution, SDS-PAGE and UV-Vis analysis of the purified PolH.

(A) Purification and reconstitution of PolH protein. Following the similar procedure as the overexpression and purification of PolJ, PolH was purified anaerobically from the *E. coli* BL21(DE3)/plysE cells harboring the pJTU2877 construct in 25 mM Tris-HCl, 150 mM NaCl, 5% glycerol buffer (pH 8.0). The reconstitution was performed by incubation of as-isolated PolH with 6 equal excess mole amount of Na$_2$S, Fe(NH$_4$)$_2$(SO$_4$)$_2$ in presence of 5.0 mM DTT for 3 hr at 4°C anaerobically.

(B) SDS-PAGE of the purification of PolH.

(C) UV-Vis spectra of as-isolated, reconstituted and reduced PolH.

1. Apo: As-isolated PolH in anaerobic condition
2. RCN: Reconstituted PolH with Fe$^{2+}$ and S$^{2-}$ in presence of DTT
3. DT-: Partially reduced reconstituted PolH by diithionite
Figure S14. EPR characterization of as-isolated and reconstituted PolH.

(A) EPR spectrum of as-isolated PolH (500 µM, in 25 mM Tris-HCl, 150 mM NaCl, 10% glycerol buffer, pH 8.0).

(B) EPR spectrum of as-isolated PolH reduced by sodium dithionite.

As-isolated PolH (500 µM, in 25 mM Tris-HCl, 150 mM NaCl, 10% glycerol buffer, pH 8.0) was incubated with 5.0 mM sodium dithionite anaerobically on ice for 5.0 minutes and frozen by liquid nitrogen for EPR recording.

(C) EPR spectra of reduced as-isolated (blue line) and reconstituted (red line) PolH.

The reduced as-isolated EPR sample (blue line) is same as above (B). The reduced reconstituted (red line) PolH (400 µM) was prepared with similar condition as above B and same as Fig. 2C.
Figure S15. Analysis of NikO and LC-MS confirmation of its catalyzed-product, 3’-EUMP.

(A) SDS-PAGE analysis of the purified NikO.

(B) Schematic map of the catalyzed reaction by NikO or its homolog, PolA.

(C) HPLC analysis of the UMP reaction catalyzed by NikO. Top, NikO reaction; bottom, negative control without NikO added.

(D) LC-MS confirmation of the NikO-catalyzed product, 3’-EUMP, which indicated a [M-H]$^-$ ion at m/z 393.11, fully corresponding to its theoretical mass.
Figure S16. LC-MS/MS analysis of the reaction of 3'-EUMP catalyzed by PolJ.

(A) PolJ reaction using 3'-EUMP as substrate. The purified N-His$_6$-PoJ protein was incubated with 3'-EUMP (1.0 mM) and MgCl$_2$ (1.0 mM) in 50 mM Tris-HCl bufer (pH 8.0) at 25°C for 8 hr.

(B) LC-MS/MS analysis of reaction mixture.

(C) Comparative tandem MS analysis of the substrate 3'-EUMP (from left to right: 78.967, 96.967, 111.018, 305.018, 393.034).

(D) Comparative tandem MS analysis of the product 3'-EU (from left to right: 87.007, 111.018, 201.039, 313.068).
Figure S17. LC-MS/MS analysis of the reaction of 3'-EUMP catalyzed by PolJ-PolH.

(A) PolJ-PolH reaction using 3'-EUMP as substrate. The purified N-His$_6$-PolH and anaerobically reconstituted PolH proteins was incubated anaerobically with 3'-EUMP (1.0 mM), MgCl$_2$ (1.0 mM), SAM (1.0 mM), and sodium dithionite (2.0 mM) in 50 mM Tris-HCl buffer (pH 8.0) at 25°C for 8 hr.

(B) LC-MS/MS analysis of reaction mixture.

(C) Comparative tandem MS analysis of the product isomer of 3'-EUMP (from left to right: 96.967, 111.018, 295.057, 350.028, 393.034).

(D) Comparative tandem MS analysis of the product isomer of 3'-EU (from left to right: 87.007, 111.018, 201.039, 237.090, 313.068).
Figure S18. PolH activity assay.

(A) HPLC analysis of the PolH reaction with 3’-EUMP as substrate.

(H) OAP + 3’-EUMP

(G) Purified OAP

(F) SAM std.

(E) 5’-dA std.

(D) 3-EUMP std.

(C) Denatured PolH

(B) Control exp without SAM

(A) Reconstituted PolH

(A) Reconstituted PolH (20 µM) in presence of 3’-EUMP (0.5 mM), SAM (1.0 mM) and dithionite (2.0 mM) at 37 °C for 2.0 hr anaerobically; (B) the control assay without SAM; (C) the control assay with boiled denatured PolH; (D) 3’-EUMP standard; (E) 5’-dA standard; (F) SAM standard; (G) purified PolH reaction product (OAP); (H) co-injection of OAP with 3’-EUMP. **HPLC condition:** The samples were injected into a HPLC (Agilent 1260) equipped with a phenomenex Luna C18 column (5 µm, 4.6 x 250 mm) and eluted with 5%-30% MeOH (0.15% TFA) for 10 min, 30-50% MeOH (0.15% TFA) in 10-25 min with water as secondary mobile phase at a flow rate of 0.5 mL/min at 35 °C, monitored at 260 nm.
(B) LC-MS and ESI-MS/MS analysis of the substrate of 3'-EUMP catalyzed by PolH (B-1: 393.034; B-2 (from left to right): 78.957, 96.967, 111.018, 305.017, 393.033).

(C) LC-MS and ESI-MS/MS analysis of the product isomer of 3'-EUMP catalyzed by PolH (C-1: 393.034; C-2 (from left to right): 78.957, 96.967, 111.018, 295.056, 350.029, 393.034).
Figure S19. $^1$H-, $^{13}$C-, $^{31}$P-, and DEPT NMRs analysis of purified product from the reaction of 3'-EUMP catalyzed by PolH with the contamination of glycerol.

(A) $^1$H NMR (500 MHz, D$_2$O) $\delta$ 7.96 (1H, d, $J$ = 5.0 Hz), 5.84 (1H, d, $J$ = 5.0 Hz), 5.75 (1H, s), 4.98 (1H, d, $J$ = 5.0 Hz), 4.57 (1H, d, $J$ = 10.0 Hz), 4.38 (1H, d, $J$ = 5.0 Hz), 4.08 (1H, d, $J$ = 15.0 Hz), 3.96 (1H, dd, $J$ = 10.0, 5.0 Hz), 2.42 (1H, d, $J$ = 15.0 Hz), 1.87 (1H, t, $J$ = 15.0 Hz). The signals of 3.70-3.72 (1H, m), 3.58 (2H, dd, $J$ = 15.0, 5.0 Hz), and 3.48 (2H, dd, $J$ = 10.0, 5.0 Hz) were from the glycerol contamination.

(B) $^{13}$C NMR (125 MHz, D$_2$O) $\delta$ 174.95, 166.39, 151.13, 141.29, 101.47, 91.88, 76.36, 71.95, 70.89, 68.28, 32.93. The signals of 71.98 ppm and 62.39 ppm were from the glycerol contamination.

(C) $^{31}$P NMR (202 MHz, D$_2$O) $\delta$ -0.71.

(D) DEPT NMR.
Figure S20. $^1$H-NMR and $^{13}$C-NMR of the glycerol in D$_2$O solvent.

(A) $^1$H NMR (500 MHz, D$_2$O) δ 3.70-3.72 (1H, m), 3.58 (2H, dd, $J = 15.0, 5.0$ Hz), 3.48 (2H, dd, $J = 10.0, 5.0$ Hz).

(B) $^{13}$C NMR (125 MHz, D$_2$O) δ 71.94, 62.33.

Figure S21: $^1$H-$^1$H COSY of the OAP with glycerol contamination.
Figure S22: HMQC of the OAP with glycerol contamination.

Figure S23: HMBC of the OAP with glycerol contamination.
Figure S24. LC-MS/MS analysis of the reaction of OAP catalyzed by PolJ.

(A) PoIJ reaction using OAP as substrate. The purified N-His$_6$-PoHJ was incubated with OAP (0.2 mM), MgCl$_2$ (1.0 mM) in 50 mM Tris-HCl buffer (pH 8.0) at 25°C for 8 hr.

(B) LC-MS/MS analysis of reaction mixture.

(C) Comparative tandem MS analysis of the substrate OAP (from left to right, 96.967, 111.018, 295.057, 350.027, 393.034).

(D) Comparative tandem MS analysis of the product OA (from left to right, 310.902, 311.912, 312.908, 313.067, 313.908).
Figure S25: The time course of PolH activity assay with the stoichiometric production of OAP and 5'-dA.

HPLC analysis of the PolH reaction by time courses with 3’-EUMP (0.5 mM), SAM (1.0 mM), sodium dithionite (2.0 mM) at rt anaerobically. (A) 15 min, (B) 30 min, (C) 45 min, (D) 60 min, (E) 240 min. The same HPLC condition as above PolH activity assay.
(A) The scheme of PolH reaction using 3'-EUMP as substrate in 60% D$_2$O buffer solution. Reconstituted PolH (20 µM) was incubated with 3'-EUMP (0.5 mM), SAM (1.0 mM), and dithionite (2.0 mM) at 25 °C in 50 mM Tris-HCl buffer (pH 8.0) containing 60% D$_2$O anaerobically.

(B) LC-MS/MS analysis of reaction mixture (B-2 from left to right): 78.957, 96.967, 111.018, 295.057, 296.062, 350.028, 351.033, 393.034, 394.039).

(C) Comparative tandem MS analysis of the product OA (from left to right, 78.957, 96.967, 111.018, 295.057, 296.062, 350.033, 393.034).

(D) Comparative tandem MS analysis of the deuterium labeled product D-C7'-OA (from left to right, 78.957, 96.968, 111.018, 295.057, 350.028, 393.034).
Scheme S1. PolJ and PolH reaction with 3'-EUMP as substrate.
Table S1. Strains, plasmids, and cosmids used in this study.

<table>
<thead>
<tr>
<th>Strain, plasmid, or cosmid</th>
<th>Relevant genotype or description</th>
<th>Source/Reference</th>
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<tr>
<td><strong>S. aureochromogens</strong></td>
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<td></td>
</tr>
<tr>
<td>YB172</td>
<td>Industrial polyoxin producer</td>
<td>Lab collection</td>
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<td>CXR14</td>
<td>YB172 derivative with the entire polyoxin gene cluster deleted</td>
<td>This study</td>
</tr>
<tr>
<td>CXR14::4774</td>
<td>CXR14 derivative containing pJTU4774</td>
<td>This study</td>
</tr>
<tr>
<td>CXR14::4774./JpolH</td>
<td>CXR14 derivative containing pJTU4774./JpolH</td>
<td>This study</td>
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<tr>
<td>CXR14::4774./JpolJ</td>
<td>CXR14 derivative containing pJTU4774./JpolJ</td>
<td>This study</td>
</tr>
<tr>
<td>CXR14::2463</td>
<td>CXR14 derivative containing pJTU2463</td>
<td>This study</td>
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<td><strong>E. coli</strong></td>
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<td>DH10B</td>
<td>F⁻ mcrA Δ(mrr-hsdRMS-mcrBC) φ80d lacZΔ M15 ΔlacZ474 deoR recA1 endA1 ara Δ139 Δ(ara leu)1697 galU galK ΔrpsL ΔnupG</td>
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<td>BW25113</td>
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<td><strong>Fungi</strong></td>
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<td>Trichosporon cutaneum 3071</td>
<td>Indicator fungi for the bioassay of polyoxins as well as an opportunistic human fungal pathogen¹</td>
<td>CGMCC¹</td>
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<td><strong>Plasmids/cosmids</strong></td>
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<td>pJ790</td>
<td>λ-RED (gam bet exo) Cml8 araC rep101 ts</td>
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<td>pQ446</td>
<td>aac (3)IV, SCP2, rep MB1* attPφC31 oriT</td>
<td>Stratagene</td>
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<tr>
<td>pBlueScriptII SK(+)</td>
<td>bia lacZΔ oriF1</td>
<td></td>
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<td>pJTU2463</td>
<td>9 kb cosmid vector containing 3 λ cos sites; pQ446 derivative, SCP2 replicon (3.5 kb EcoRI-XbaI) replaced by pSET152 Int attPφC31</td>
<td>(Yao et al. unpublished)</td>
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<td>pJTU2178</td>
<td>nikO structural gene was cloned into Ndel-EcoRI sites of pET28a</td>
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<td>pJTU2876</td>
<td>pBlueScriptII SK(+) derivative with EcoRI-engineered polH structural gene amplified from SADF cloned into its SmaI-EcoRI sites</td>
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<td>pJTU2863</td>
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<td>pJTU4774</td>
<td>The cosmid with XbaI and SpeI sites infilled containing the entire polyoxin gene cluster</td>
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<td>pJTU5701</td>
<td>22F7 with XbaI and Nhel sites infilled and religated separately; no nik genes deleted</td>
<td>This study</td>
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oriT, origin of transfer of plasmid RK2; tsr, thiostrepton resistance gene; aac(3) IV, apramycin resistance gene; CmlR, chloramphenicol resistance gene; KmR, kanamycin resistance gene; ^CGMCC, China General Microbiological Culture Collection Center.; rep^pMB1*, mutated rep^pMB1.

Table S2. PCR primers used in this study.

<table>
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<th>Primers</th>
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<td>polHexR</td>
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<td>polJexR</td>
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<tr>
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<td>polJtgtR</td>
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Table S3. Comparative analysis of the target pol genes and their homogs involved in potential AHA biosynthetic pathway

<table>
<thead>
<tr>
<th>Protein (aa)</th>
<th>Proposed Function</th>
<th>S. tendae Tü 901, Homolog (aa), Identity, Similarity, (%)</th>
<th>S. chromofuscus ATCC49982 Homolog (aa), Identity, Similarity, (%)</th>
<th>S. aurantiacus JA4570 Homolog (aa), Identity, Similarity, (%)</th>
<th>Xenorhabdus szeentirmai DSM16338, Homolog (aa), Identity, Similarity, (%)</th>
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</thead>
<tbody>
<tr>
<td>PolA (445)</td>
<td>UMP enolpyruvyl transferase</td>
<td>NikO (473) (63.73)</td>
<td>sccontig008-75 (495) (40.53)</td>
<td>STRAU_0242 (423) (33.47)</td>
<td>XSR1_370018 (436) (35.57)</td>
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<td>PolD (215)</td>
<td>Hydroxylase</td>
<td>NikI (218) (50, 66)</td>
<td>sccontig008-76 (201) (28,46)</td>
<td>STRAU_0246 (216) (53,67)</td>
<td>XSR1_370026 (221) (31,48)</td>
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<td>PolH, (469)</td>
<td>Radical SAM protein</td>
<td>NikJ (454) (70, 81)</td>
<td>sccontig008-80 (465) (52,63)</td>
<td>STRAU_0239 (436) (51,63)</td>
<td>XSR1_370015 (447) (43,60)</td>
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<td>PolI (380)</td>
<td>Aminotransferase</td>
<td>NikK (368) (62, 70)</td>
<td>sccontig008-81 (392) (35,47)</td>
<td>STRAU_0238 (374) (32,44)</td>
<td>XSR1_370019 (371) (27,47)</td>
</tr>
<tr>
<td>PolJ (273)</td>
<td>Phosphatase</td>
<td>NikL (240) (61, 69)</td>
<td>sccontig008-78 (242) (31,48)</td>
<td>STRAU_0240 (223) (35,49)</td>
<td>XSR1_370020 (231) (30,46)</td>
</tr>
<tr>
<td>PolK (213)</td>
<td>Hydroxylase</td>
<td>NikM (213) (58, 70)</td>
<td>sccontig008-74 (240) (47,59)</td>
<td>STRAU_0236 (207) (36,53)</td>
<td>XSR1_370017 (206) (35,48)</td>
</tr>
</tbody>
</table>
4. Methods and Experimental section

Method S1. Materials, instruments and general methods.

All chemicals and reagents were purchased from Sigma-Aldrich (USA) or Aladdin (China) unless otherwise stated. Strains, plasmids and primers used in this study were independently listed in Table S1 and Table S2. General methods were according to the standard methods of Sambrook et al.\cite{7} or Kieser et al.\cite{8} The final antibiotic concentrations used in this study were as follows: 100 μg/ml ampicillin, 30 μg/ml apramycin, 50 μg/ml kanamycin, 25 μg/ml chloramphenicol and 12.5 μg/ml thiostrepton. NMR spectra were recorded on a 500 MHz instrument (Bruker). The tandem MS of activity assay and digested protein was detected on the Q Exactive Orbitrap Mass Spectrometry (Thermo) in the core facility center of SIPPE. The LC condition of LC-MS/MS was conducted using an analytical Phenomenex Luna NH2 column (3 µM, 4 × 100 mm). The sample was eluted with a gradient of solvent A (50 mM NH₄OAc, pH 9.5) and solvent B (acetonitrile). The gradient was run from 30% A over 1 min, 30 to 95% A over 8 min, 95% A over 3 min, then 0.1 min at 95–30% A, and then re-equilibrated at 30% A for 8 min. The flow rate was 0.3 mL/min at 15 °C. The peaks corresponding to the enzymatic reaction products were subjected to ESI-MS or MS/MS analyses. EPR spectra were recorded using a Bruker EMX Plus 10/12 spectrometer (Bruker Co., Ltd., Germany) equipped with an Oxford ESR910 liquid helium continuous flow cryostat (Oxford Instrument Co., Ltd., UK) at the National High Magnetic Field Laboratory of the Chinese Academy of Sciences, Hefei, China. Acquisition conditions for the Fe-S cluster analysis are: microwave frequency, 9.39 GHz; microwave power, 1 mW; field modulation amplitude, 5 Gauss; modulation frequency, 100 kHz; temperature, 30 K.

Method S2. In frame deletion of \textit{polH} or \textit{polJ} in cosmid pJTU4774 by PCR-targeting technology.

For the targeted inactivation of \textit{polH}, a kanamycin resistance cassette (neo) from SuperCos1 amplified using the primers polHtgtF and polHtgtR (Table S2), was recombined into pJTU5701 by λ-Red-mediated recombination\cite{4} to give pJTU4774/\textit{polH-neo}. The \textit{neo} cassette was then deleted \textit{in vitro} by XbaI and Spel digestion (unique sites) and religated to produce pJTU4774/Δ\textit{polH}. The unmarked deletion was confirmed by PCR using primers polHexpF/R. Likewise, \textit{polJ} in pJTU4774 was inactivated using the primers polJtgtF/R for the the targeted gene replacement, and polJexpF/R to confirm the unmarked deletion in pJTU4774/Δ\textit{polJ}.

Method S3. Cultivation and metabolites analysis of the \textit{S. aureochromogenes} and its derivatives.

\textit{S. aureochromogenes} and its derivatives were cultivated in liquid fermentation medium.\cite{9} After the fermentation, the broth was adjusted to pH 3.0 with oxalic acid and filtered for LC-MS analysis and targeted metabolites preparation. Bioassay of the metabolites was based on the procedures by Zhai et al.\cite{6}
For the detection of the target metabolites accumulated by related *S. aureochromogenes* and its derivatives, the HPLC (Shimadzu LC-20A) analysis were performed on a ZORBAX SB-C18 column (Agilent, 3.5 mm, 4.6×250 mm) with an elution gradient of 5%-55% methanol: 0.15% aqueous TFA (HPLC grade) over 30 min at a flow rate of 0.5 mL/min, and then keep the ratio of 55% methanol : 0.15% aqueous TFA for another 30 min at 0.5 mL/min. The elution was monitored at UV 263 nm with DAD detector.

For the purification of the target metabolite (Compound X), the conditions are the identical to the above except the preparation was performed on a ZORBAX SB-C18 Column (Agilent, 5 µm, 9.4×250 mm) at flow rate of 3 mL/min. The purified metabolite was subsequently concentrated by vacumm, after that, the identity of the metabolite was preliminarily confirmed by LC-MS analysis (Thermo LTQ-Orbitrap) with positive ion mode, and the detailed parameters are as follows: drying gas temperature: 275 °C, drying gas flow: 10 liters/mL, and nebulizer pressure: 30 p.s.i.

**Method S4. Preparation of the NikO-catalyzed product, 3'-EUMP.**

NikO was overexpressed and purified based on the procedures described previously. After the reactions, the NikO protein was inactivated by adding equal volume of methanol, and removed by centrifuging at 12,000 r/m for 10 min. Subsequently, the 100 mL supernatant of the reactions was concentrated by vacumm to volume of 5 ml, then 3'-EUMP was prepared and its identity was confirmed by LC-MS (in negative ion mode) and NMR analysis. The HPLC preparation was conducted on a ZORBA SB-C18 Column (Agilent, 5 µm, 9.4×250 mm) with an elution gradient of 5%-30% methanol: 0.3% aqueous TFA (HPLC grade) over 20 min at a flow rate of 3.0 mL/min, and then keep the ratio 30% methanol: 0.3% aqueous TFA for another 20 min at 3.0 mL/min. The elution was monitored at UV 263 nm with a DAD detector.

**Method S5. Cloning of PolH and PolJ in *E. coli.***

For the construction of the *polH* and *polJ* expression vector, the structural gene of them were amplified with individual pairs of the primers polHexF/polHexR and polJexF/polJexR. The EcoRI-engineered PCR products were then cloned into SmaI-EcoRI sites of pBlueScriptII SK(+) to generate pJTU2876/pJTU2863. After confirmation by DNA sequencing, the Ndel-EcoRI engineered fragments containing both structural genes from the plasmids were cloned into the counterpart sites of pET28a to give pJTU2877 and pJTU2864, respectively.

**Method S6. Over-expression and purification of PolJ.**

A culture of *E. coli* BL21(DE3) cells harboring the PolJ/pET28a construct were grown in the LB medium containing 50 µg/mL of kanamycin at 37 °C with shaking (200 rpm) until the OD$_{600}$ reached ~0.5. Protein expression was then induced by addition of isopropyl β-D-1-thiogalactopyranoside (IPTG) to a final concentration of 0.5 mM, and the cells were allowed to grow at 18 °C and 200 rpm for an additional 18 h. The cells were harvested by centrifugation at 4500 x g for 15 min. All purification steps were carried out
at 4 °C using Ni-NTA resin and the PolJ protein were washed and eluted by 20-250 mM imidazole in 25 mM Tris-HCl, 150 mM NaCl buffer (pH 8.0). The collected protein solution was dialyzed against 25 mM Tris-HCl, 150 mM NaCl buffer (pH 8.0) for three times. The protein solution was then flash-frozen in liquid nitrogen and stored at -80 °C until use. Protein concentration was determined by the Bradford assay using bovine serum albumin as the standard.

**Method S7. PolJ activity assays.**

The PolJ activity assays were performed with the purified N-His\textsubscript{6}-PoHJ incubated with OAP (0.2 mM) or 3'-EUMP (1.0 mM), MgCl\textsubscript{2} (1.0 mM) in 50 mM Tris-HCl buffer (pH 8.0) at 25 °C for 8 hr. The reaction mixture was quenched with equal volume amount of MeOH and centrifuged to remove protein precipitation then filtrated with 0.22 µM of filter before injecting to HPLC. The assay was monitored by LC-MS/MS with the general method by the Q Exactive Orbitrap Mass Spectrometry.

**Method S8. Purification and reconstitution of PolH protein.**

Following the similar procedure as the over-expression and purification of PolJ, PolH was purified anaerobically in the Coy-Chamber glove box from the *E. coli* BL21(DE3)/plysE cells harboring the PolH/pET28a construct in 4 °C.

The reconstitution was performed by incubation of as-isolated PolH with 6 equal excess mole amount of Na\textsubscript{2}S, Fe(NH\textsubscript{4})\textsubscript{2}(SO\textsubscript{4})\textsubscript{2} in presence of 5.0 mM DTT for 3 hr at 4 °C anaerobically. Then the reconstituted solution was desalted by G-10 column then washed with the buffer (25 mM Tris-HCl, 150 mM NaCl, 5% glycerol buffer, pH 8.0). The fractions contained PolH were collected and concentrated by ultrafiltration.

**Method S9. PolH activity assays**

The PolH activity assays were performed at 37°C or room temperature under anaerobic conditions for a proper time in a Coy-chamber glove box. The reaction mixture contained the following: reconstituted PolH (20 µM), 3'-EUMP (0.5 mM), SAM (1.0 mM) and sodium dithionite (2.0 mM) in 50 mM Tris-HCl (pH 8.0) anaerobically. The reaction mixture was quenched with equal volume amount of MeOH and centrifuged to remove protein precipitation then filtrated with 0.22 µM of filter before injecting to HPLC.

The reaction conversion was monitored in the Agilent 1260 HPLC equipped the phenomenex luna C18 column (5.0 µM, 250 mm x 4.6 mm) with a gradient condition of solvent A (MeOH with 0.15% TFA) and solvent B (0.15% TFA in H\textsubscript{2}O) below. The gradient was run from 5% to 30% A over 10 min, 30% to 50% A over 15 min, 30% to 50% A over 1 min, then 4.0 min at 50–5% A, and then re-equilibrated at 5% A for 5 min. The flow rate was 0.5 mL/min at 35 °C.

The product was purified with the HPLC condition above. The LC-MS/MS was detected with the general method by the Q Exactive Orbitrap Mass Spectrometry.
5. Supplementary References.

Reference S1. Supplementary References.