1	Sublethal β -lactam antibiotics induce PhpP phosphatase expression and
2	StkP kinase phosphorylation in PBP-independent β -lactam antibiotic
3	resistance of Streptococcus pneumoniae
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19	Running title: Pneumococcal PhpP as a PP2C type protein phosphatase
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Abstract 35

StkP and PhpP of *Streptococcus pneumoniae* have been confirmed to compose a signaling 36 37 couple, in which the former is a serine/threonine (Ser/Thr) kinase while the latter was annotated as a phosphotase. StkP has been reported to be involved in penicillin-binding 38 39 protein (PBP)-independent penicillin resistance of S. pneumoniae. However, the enzymatic characterization of PhpP and the role of PhpP in StkP-PhpP couple remain poorly understood. 40 Here we showed that 1/4 minimal inhibitory concentration (MIC) of penicillin (PCN) or 41 cefotaxime (CTX), the representatives of β -lactam antibiotics, could induce the expression of 42 43 stkP and phpP genes and phosphorylation of StkP in PCN/CTX-sensitive strain ATCC6306 and three isolates of S. pneumoniae (MICs: 0.02-0.5 µg/ml). The product of phpP gene 44 hydrolyzed PP2C type Ser/Thr phosphotase-specific RRA(pT)VA phosphopeptide substrate 45 with the Km and Kcat values of 277.35 µmol/L and 0.71 S⁻¹, and the hydrolytic activity was 46 blocked by sodium fluoride, a PP2C type Ser/Thr phosphatase inhibitor. The phosphorylation 47 levels of StkP in the four *phpP* gene-knockout ($\Delta phpP$) mutants were significantly higher 48 49 than that in the wild-type strains. In particular, the MICs of PCN and CTX against the $\Delta phpP$ mutants were significantly elevated as 4-16 µg/ml. Therefore, our findings confirmed that 50 51 sublethal PCN and CTX act as environmental inducers to cause the increase of *phpP* and *stkP* gene expression and StkP phosphorylation. PhpP is a PP2C type Ser/Thr protein phosphatase 52 53 responsible for dephosphorylation of StkP. Knockout of the *phpP* gene results in a high level of StkP phosphorylation and PBP-independent PCN/CTX resistance of S. pneumoniae. 54

Importance 55



Streptococcus pneumoniae is a common pathogen in human populations in many countries

and areas due to the prevalence of β -lactam antibiotic-resistant pneumococcal strains. 57 Production of β-lactamases and mutation of penicillin-binding proteins (PBP) have been 58 59 considered as the major β -lactam antibiotic-resistant mechanisms in bacteria, but S. pneumoniae has not been confirmed to produce any β-lactamases and many pneumococcal 60 61 strains present PBP mutation-independent β-lactam antibiotic resistance. StkP is a Ser/Thr kinase of S. pneumoniae to compose a signal-couple with PhpP protein. The present study 62 demonstrated that the PhpP is a PP2C-type phosphotase for dephosphorylation of StkP and 63 the sublethal penicillin (PCN) or cefotaxime (CTX) acted as environmental signal molecules 64 65 to induce the expression of PhpP. The knockout of PhpP-encoding gene caused the PCN/CTX resistance generation of PCN/CTX-sensitive pneumococcal strains. All the data indicate that 66 StkP-PhpP couple of *S. pneumoniae* is involved in PBP mutation-independent β-lactam 67 68 antibiotic resistance by phosphorylation of StkP.

Key words: *Streptococcus pneumoniae*; PhpP; PP2C type protein phosphatase;
 StkP-PhpP signaling couple; β-lactam antibiotics; Drug resistance

72 Introduction

Streptococcus pneumoniae is a major causative agent of bacterial pneumonia and tympanitis
in children [1-3]. More importantly, in the recent years, *S. pneumoniae*-infected meningitis
cases with high fatality have been frequently reported in many countries and areas [4-8].
Therefore, *S. pneumoniae* is a common pathogen for human beings with global importance.

β-lactam antibiotics are the first choice in clinic to cure S. pneumoniae-infected patients 77 [9]. However, in the recent years, β -lactam antibiotic-resistance of S. pneumoniae isolates 78 from patients is continuously increased and the antimicrobial-resistant S. pneumoniae strains 79 80 became more epidemic in many countries and areas [10-15], which has been considered as the major reason for increased incidence of S. pneumoniae-infected diseases [15,16]. 81 Bacterial β -lactamases have been confirmed to play a key role in generation of β -lactam 82 83 antibiotic resistance in many bacteria including S. pneumoniae [17]. Mutation of penicillin-binding proteins (PBP), the receptors of β -lactam antibiotics located on surface of 84 bacteria, has been reported as the major β -lactam antibiotic resistant mechanism in bacteria 85 [18,19]. However, recent studies found that some of the S. pneumoniae strains had no PBP 86 mutation but presented β-lactam antibiotic resistance [20-22], indicating that S. pneumoniae 87 88 may have a PBP mutation-independent mechanism of β -lactam antibiotic resistance.

89 StkP is a sequence-conserved eukaryotic-type serine/threonine (Ser/Thr) kinase (STK) of 90 *S. pneumoniae* that has been confirmed to be involved in PBP mutation-independent 91 penicillin resistance [22]. In the chromosomal DNA of *S. pneumoniae*, StkP-encoding gene 92 (*stkP*) and *phpP* gene compose a *stkP-phpP* operon and the product of *phpP* gene is 93 annotated as a putative phosphatase [23,24]. A previous study demonstrated that the PhpP

94 and StkP of S. pneumoniae composed a StkP-PhpP signaling couple [25]. It has been reported that both prokaryotic and eukaryotic STKs are activated through phosphorylation at Ser/Thr 95 96 sites and some certain protein phosphatases can inactivate STKs by hydrolysis of phosphoryl groups at the Ser/Thr residual sites in STKs [26]. In particular, a previous study revealed that 97 98 penicillin (PCN) could cause the gene expression profile change of S. pneumoniae [27]. 99 Therefore, we presume that β -lactam antibiotics may act as environmental inducers to cause the change of PhpP and StkP expression and StkP dephosphorylation of S. pneumoniae as 100 well as the PhpP may be involved in the StkP-associated PBP mutation-independent 101 102 penicillin resistance by dephosphorylation of StkP.

103 In the present study, we used PCN and cefotaxime (CTX) as the representatives of β -lactam antibiotics to detect their induction of *phpP* and *stkP* gene expression and then 104 105 identified the product of *phpP* gene as a PP2C type Ser/Thr protein phosphatase by virtue of its ability to hydrolyze Ser/Thr phosphatase-specific substrates. Moreover, the *phpP* genes of 106 S. pneumoniae strains were inactivated to determine the role of PhpP in dephosphorylation of 107 StkP in vivo and the change of β -lactam antibiotic resistance. The results of this study 108 confirmed that the product of *S. pneumoniae phpP* gene is a Ser/Thr protein phosphatase that 109 involved in β-lactam antibiotic resistance-associated StkP-PhpP signaling couple by StkP 110 111 dephosphorylation during induction of sublethal PNC and CTX.

113 Materials and Methods

114 Bacterial strains and culture.

S. pneumoniae ATCC6306 and three β-lactam antibiotic-sensitive *S. pneumoniae* isolates
(No.: SP5, SP9 and SP14, belonging to serotype 3, 19F and 19A) from pneumonia children
were kindly provided by the Department of Medical Microbiology and Parasitology, Zhejiang
University School of Medicine. All the strains were cultured with Colombia blood agar
(bioMerieux, France) or 0.5% yeast extract-containing Todd-Hewitt (TH) broth (Sigma, USA)
at 37°C [28]. Besides, *Escherichia coli* EL21DE3 (Novagen, USA) was cultured in
Luria-Bertani (LB) medium (Oxoid, England) at 37°C.

122

123 Animal.

New Zealand white rabbits (3.0 to 3.5 kg per animal) were provided by the Laboratory Animal Center of Hangzhou Medical College (Certificate No.: SCXK[zhe]2012-0173). All the animal experimental protocols were approved by the Ethics Committee for Animal Experiment of Hangzhou Medical College.

128

129 Drug susceptibility test.

Susceptibility of each of the four *S. pneumoniae* strains to PCN or CTX was detected by E-test (bioMerieux) according to the manufacturer's instruction. The minimal inhibitory concentrations (MIC) against *S. pneumoniae* strains, ≤ 2 or ≥ 8 µg/ml of PCN (Sigma) and \leq 1 or ≥ 4 µg/ml of CTX (Sigma), were considered to be sensitive or resistant [29].

135 Detection of sublethal PCN- and CTX-induced expression of *phpP* and *stkP* genes.

Each of the four S. pneumoniae strains was inoculated into TH broth for a 200 rpm shaking 136 137 incubation at 37°C. When the value of optical density at 600 nm (OD_{600}) of pneumococcal culture turbidity reached 0.5, 1/4 MIC PCN or CTX was added and then incubated for 0.5, 1, 138 139 2, 4, 8 or 12 h as above. After centrifugation and washing with phosphate buffered saline (PBS), total RNA of each of S. pneumoniae strains was extracted using a TRIzol[®] Max[™] 140 Bacterial RNA Isolation kit (Invitrogen) plus a gDNA Eraser Kit (TaKaRa, China) and then 141 quantified by ultraviolet spectrophotometry [30]. Subsequently, cDNA from each of the total 142 143 RNAs was synthesized using a PrimeScript[™] RT Reagent Kit (TaKaRa). Using each of the cDNAs as template, the *phpP*- or *stkP*-mRNA level was assessed by real-time fluorescence 144 quantitative reverse transcription polymerase chain reaction (qRT-PCR) with the primers 145 P-1F/P-1R or S-1F/S-1R (Table 1) using a SYBR[®] Premix Ex-TaqTM Kit (TaKaRa) in an ABI 146 7500 Real-Time PCR System (ABI, USA). The primers used were designed using Primer 147 Premier 5.0 software according to the *phpP* or *stkP* gene sequences in GenBank (accession 148 No.: NC 003098 and NC 003028). In the qRT-PCR, 16S rRNA gene of S. pneumoniae was 149 used as the internal reference [31], while the PCN- or CTX-untreated S. pneumoniae strains 150 were used as the controls. The obtained qRT-PCR data were analyzed using the $\Delta\Delta$ Ct model 151 and randomization test in REST 2005 software [32]. 152

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154 Amplification and sequencing of *PhpP* and *stkP* gene segments.

Genomic DNA of each of the four *S. pneumoniae* strains was extracted using a Bacterial Genomic DNA Preparation Kit (Axygen). By using a High Fidelity PCR Kit (TaKaRa), the

entire *phpP* or *stkP* gene segments were amplified from the DNA templates by PCR using the
primers P-2F/P-2R or S-2F/S-2R (Table 1). The PCR products were examined by 1.5%
ethidium bromide-stained agarose gel electrophoresis and then cloned into pMD19-T plasmid
using a T-A Cloning Kit (TaKaRa) to form recombinant pMD19-T^{*phpP*} and pMD19-T^{*stkp*}
plasmids for sequencing by Invitrogen Co. in Shanghai of China.

162

163 **Bioinformatic analysis of** *phpP* and *stkP* genes.

Since the nucleotide and amino acid sequence identities of *phpP* and *stkP* genes from the four *S. pneumoniae* strains were as high as 98.7%-100%, the *phpP* and *stkP* genes of *S. pneumoniae* strain ATCC6306 were analyzed using TMHMM and NCBI Database Conserved
Domain Database (CDD) software [33].

168

169 Generation of prokaryotic expression systems of *phpP* and *stkP* genes.

The pMD19-T^{phpP} or pMD19-T^{stkP} plasmid from S. pneumoniae strain ATCC6306 and 170 171 pET42a vector (Novagen) were digested with both Nde I and Xho I or Nde I and Hind III (TaKaRa). The recovered *phpP* or *stkP* gene segment was linked with the linearized pET42a 172 using T4 DNA ligase (TaKaRa) and then transformed into E. coli BL21DE3 by CaCl₂ 173 transformation method to form *E. coli* BL21DE3^{pET42a-phpP} or *E. coli* BL21DE3^{pET42a-stkP}. The 174 engineered strains were cultured in LB medium containing 50 µg/ml kanamycin (Sigma) and 175 the pET42a-phpP and pET42a-stkP were extracted from the strains using a Plasmid 176 177 Extraction Kit (Axygen) for sequencing again.

179 Expression of *phpP* and *stkP* genes and extraction of expressed products.

The engineered strains, E. coli BL21DE3 pET42a-phpP and E. coli BL21DE3 PET42a-stkP, were 180 181 cultured in kanamycin-containing LB liquid medium to express the target recombinant proteins (rPhpP and rStkP) under induction of 0.5 mM isopropy-β-D-thiogalactoside (IPTG, 182 183 Sigma). After ultrasonic breakage on ice and a $13,800 \times g$ centrifugation for 10 min (4°C), the 184 supernatants of cultures were collected to extract soluble rPhpP and rStkP using a Ni-NTA affinity chromatographic column (BioColor, China). The extracted rPhpP or rStkP was 185 quantified using a BCA Protein Assay Kit (Thermo Scientific, USA). Both the expressed and 186 187 extracted rPhpP and rStkP were examined by sodium dodecyl sulfate polyacrylamide gel electropheresis (SDS-PAGE) plus a Gel Image Analyzer (Bio-Rad, USA). 188

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190 Preparation of rPhpP-IgG and rStkP-IgG

New Zealand rabbits were intradermally immunized on days 1, 14, 21 and 28 with Freund's adjuvant-mixed 2 mg rPhpP or rStkP per animal. Fifteen days after the last immunization, the sera were collected to separate rStkP-IgG or rPhpP-IgG by ammonium sulfate precipitation plus a DEAE-52 column chromatography using 10 mM phosphate buffer (pH 7.4) for elution [34]. The titer of rPhpP-IgG or rStkP-IgG binding to rStkP or rPhpP was detected by immunodiffusion test.

197

198 **Phosphatase activity assays**.

The enzymatic activity of rPhpP was detected using a pNPP-Phosphate Assay Kit (BioAssay
Systems, USA) and a PP2C-specific Ser/Thr Phosphatase Assay Kit (Promega, USA) [35,36].

201	Briefly, 0.5, 1, 2.5, 5, 10 or 20 µg rPhpP was mixed with 500 nM para-nitrophenyl phosphate
202	(p-NPP), an universal Ser/Thr phosphatase substrate, in 100 µl reaction buffer. After a
203	30-min incubation at 37°C, the OD ₄₀₅ values reflecting p-NPP hydrolysis were detected using
204	type M5 spectrophotometer (Bio-Rad). On the other hand, 0.5, 1, 2.5, 5, 10 or 20 μ g rPhpP
205	was mixed with 100 μ M RRA(pT)VA phosphopeptide, a PP2C type Ser/Thr protein
206	phosphatase-specific substrate, in 100 μ l reaction buffer. After incubation as above, the OD ₆₀₀
207	values reflecting RRA(pT)VA hydrolysis were detected by spectrophotometry.

208

209 **Phosphatase inhibition test.**

Okadaic acid (OA) is an inhibitor of PP1, PP2A and PP2B type Ser/Thr phosphatases while
sodium fluoride (NaF) is a PP2C type Ser/Thr phosphatase inhibitor [37,38]. Briefly, 5 μg
rPhpP was mixed with 0.1, 0.5, 1, 5 or 10 μM OA (Sigma) or 0.1, 0.5, 1, 5 or 10 mM NaF

213 (Sigma) in 100 μl reaction buffer and then incubated at 37°C for 30 min. The activity of OA-

or NaF-treated rPhpP to hydrolyze RRA(pT)VA substrate was detected as described above.

215

216 **Determination of Km and Kcat values.**

According to the results of phosphatase activity assays, 5 μ g rPhpP was mixed with 50, 100, 150, 200 or 250 μ M RRA(pT)VA substrate and then detected by spectrophotometry as described above. According to the OD₆₀₀ values reflecting RRA(pT)VA hydrolysis and free phosphate concentration standard curve, the Km and Kcat values of rPhpP hydrolyzing the substrate were calculated by double reciprocal Lineweaver-Burk plot [39].

223 Generation and identification of *phpP* gene-knockout mutant.

pEVP3 plasmid has been used to generate the target gene-knockout mutant of S. pneumoniae 224 225 [40,41]. Briefly, a 495-bp phpP gene segment (phpP-495) from S. pneumoniae strain ATCC6306 was amplified using a High Fidelity PCR Kit (TaKaRa) with the primers 226 P-3F/P-3R and then cloned into pMD19-T to form pMD19-T^{*phpP-495*} for sequencing as above. 227 The pMD19-T^{phpP-495} and pEVP3 plasmid were digested with both Xho I and BamH I 228 (TaKaRa). The recovered *phpP*-495 segment was linked with the linearized pEVP3 using T4 229 DNA ligase (TaKaRa) to form suicide plasmid pEVP3^{phpP495} for sequencing again. Each of 230 the four S. pneumoniae strains was inoculated in TH broth containing 0.01% CaCl₂ and 0.2% 231 BSA (Sigma) for a 250-rpm incubation at 37°C to the OD_{600} value as 0.25 and then collected 232 by centrifugation. The competent pneumococcal cells were suspended in 200 µl TH broth 233 234 containing 0.01% CaCl₂, 0.2% BSA and 200 ng/ml competence stimulation peptide (CSP, AnaSpec, USA) and then added with 200 ng pEVP3^{*phpP-495*} for transformation. The mixtures 235 were smeared on 5% sheet blood Columbia agar (bioMerieux) plates containing 2.5 µg/ml 236 chloromycin (CM, Sigma) for a 48-h incubation at 37°C in a 5% CO₂ atmosphere to obtain 237 *phpP* gene-knockout colonies [42]. The strategy for generating *phpP* gene-knockout mutants 238 $(\Delta phpP-6306, \Delta phpP-SP5, \Delta phpP-SP9 and \Delta phpP-SP14)$ is summarized in Fig. 1. 239

240

241 Identification of *phpP* gene-knockout mutants.

Growth of the $\Delta phpP$ -6306, $\Delta phpP$ -SP5, $\Delta phpP$ -SP9 or $\Delta phpP$ -SP14 mutant was assessed by

243 spectrophotometry. The *phpP* gene knockout in the $\Delta phpP$ mutants was determined by PCR

using the primers P-4F/P-4R and P-5F/P-5R (Table 1) and sequencing of the 5'-phpP-pEVP3

and pEVP3-3'-*phpP* segment amplification products. Using 1:200 diluted rabbit anti-rPhpP-IgG as the primary antibody and HRP-conjugated goat anti-rabbit-IgG (Abcam, USA) as the secondary antibody, Western blot assay was performed to detect the PhpP form the $\Delta phpP$ mutants, in which the wild-type strains were used as the controls.

249

250 **Detection of sublethal PCN- or CTX-induced StkP phosphorylation.**

The $\Delta phpP$ mutants and their wild-type strains were treated with 1/4 MIC PCN or CTX for 1, 251 2, 4 or 8 h at 37°C. After centrifugation and washing with PBS, the pneumococcal pellets 252 253 were suspended in deionized water and then ultrasonically broken on ice. The lysates were centrifuged at 14,000×g for 10 min (4°C) and then the supernatants were collected to detect 254 255 protein concentrations using a BCA Protein Assay Kit (Thermo Scientific). 200 µg of each of 256 the total pneumococcal proteins was mixed with 20 µg rabbit anti-rStkP-IgG for a 2-h incubation in a 90-rpm rotator (4°C). The mixture was added with 600 µg protein-A-coated 257 agarose beads (Millipore, USA), followed by a 60-min incubation as above. After a 14,000×g 258 259 centrifugation for 5 min and washing thoroughly with PBS, the precipitated beads were suspended in 200 µl 50 mM Tris-HCl buffer (TB, pH7.5) and then mixed with 200 µl 2M 260 261 NaOH solution for a 30-min water-bath at 65°C to release phosphate groups from the captured StkP according to the instruction of Phosphoprotein Phosphate Detection Kit 262 (Sangon Biotech, Canada). The mixture was neutralized with 200 µl 4.7 M HCl solution and 263 then added with 200 µl detection buffer for a 30-min incubation at room temperature. The 264 OD₆₂₀ value reflecting phosphate group concentration released from IgG-captured StkP were 265 detected using a spectrophotometer (Molecular Devices, USA) [43]. In the detection, the 266

267 PCN- or CTX-untreated $\Delta phpP$ mutants and their wild-type strains were used as the controls.

268

269 **Detection of** β **-lactam antibiotic resistance of** Δ *phpP* **mutants.**

- 270 Susceptibility of the $\Delta phpP$ -6306, $\Delta phpP$ -SP5, $\Delta phpP$ -SP9 and $\Delta phpP$ -SP14 mutants to PCN
- 271 or CTX was detected by E-test as described above. In the detection, the wild-type S.
- 272 *pneumoniae* strains were used as the controls.

273

274 Statistical analysis

275 Data from a minimum of three experiments were averaged and presented as mean \pm standard

276 deviation (SD). One-way analysis of variance (ANOVA) followed by Dunnett's multiple

- 277 comparisons test were used to determine significant differences. Statistical significance was
- 278 defined as p < 0.05.

280 **Results**

281 Increase of *stkP*- and *phpP*-mRNA levels induced by sublethal PCN and CTX.

The *phpP*- and *stkP*-mRNA levels of each of the four tested *S. pneumoniae* strains were relatively lower. When the strains were treated with 1/4 MIC PCN or CTX, the *phpP*- and *stkP*-mRNA levels were rapidly elevated (Fig. 2A-2D). The data suggested that sublethal PCN and CTX can act as the stimulators to induce the expression of *S. pneumoniae phpP* and *stkP* genes.

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288 Characterization of *phpP* and *stkP* genes.

The product of *S. pneumoniae phpP* or *stkP* gene was predicted as a secretary cytosolic or a transmembrane protein (Fig. 3A and 3B). The *phpP* gene contains a PP2Cc type protein phosphatase domain (6-238 aa) with five enzymatic active sites (Fig. 3C). The *stkP* gene contains a C type STK domain belonging to STKc_PknB superfamilay containing twelve ATP-binding sites, six dimer interface sites, two activation loops, eight polypeptide substrate-binding sites and sixteen enzymatic active sites as well as four penicillin-binding protein and STK-associated (PASTA) superfamily domains (Fig. 3D).

296

297 Effects of expression and extraction of *stkP* and *phpP* genes.

The nucleotide and amino acid sequence identities of *phpP* or *stkP* genes from the four *S*. *pneumoniae* strains were 99.3%-100% and 99.1%-100% or 99.2%-99.7% and 98.7%-99.9%, compared to the same two genes in GenBank (accession No.: NC003098) (data not shown). The two engineered strains expressed the target recombinant proteins rPhpP and rStkP, respectively, and the extracted rPhpP or rStkP was showed as a single band in gels (Fig.4A-4D).

304

305 **Powerful Ser/Thr protein phosphatase activity of rPhpP.**

The rPhpP from S. pneumoniae ATCC6306 hydrolyzed p-NPP, an universal Ser/Thr 306 phosphatase substrate, and RRA(pT)VA, a PP2C type Ser/Thr protein phosphatase-specific 307 308 substrate, with concentration-dependent manners (Fig. 5A). The Km and Kcat values of rPhpP hydrolyzing RRA(pT)VA substrate were 277.35 µmol/L and 0.71 S⁻¹, respectively (Fig. 309 5B). Moreover, the PP2C type Ser/Thr protein phosphatase inhibitor NaF, but not the PP1, 310 PP2A or PP2B type Ser/Thr phosphatase inhibitor OA, inhibited the hydrolytic activity of 311 rPhpP (Fig. 5C). The data suggested that the product of S. pneumoniae phpP gene is a PP2C 312 type Ser/Thr protein phosphatase. 313

314

315 PCN/CTX-induced StkP phosphorylation and PhpP-caused StkP dephosphorylation.

316 The $\Delta phpP$ mutants could grow persistently in the TH broth similarly to the wild-type strains (Fig. 6A). The PCR plus sequencing and Western Blot assay confirmed the phpP gene 317 knockout and no PhpP expression in all the four $\Delta phpP$ mutants (Fig. 6B and 6C). 1/4 MIC 318 PCN or CTX could cause the increase of StkP phosphorylation levels in the $\Delta phpP$ mutants 319 and their wild-type strains (Fig. 6D and 6E). However, the $\Delta phpP$ mutants presented 320 significantly higher PCN- or CTX-induced StkP phosphorylation levels than the wild-type 321 322 strains (Fig. 6D and 6E). The data suggested that sublethal PCN and CTX can induce phosphorylation of StkP and PhpP can cause dephosphorylation of StkP in vivo. 323

324

325 Increased PCN and CTX resistance of Δ*phpP* mutants.

- 326 The MICs of PCN and CTX against wild-type ATCC6306, SP5, SP9 and SP14 strains of *S*.
- 327 pneumoniae were 0.02-0.5 µg/ml, indicating all the strains were sensitive to both PCN and
- 328 CTX. However, the MICs of PCN and CTX against the four $\Delta phpP$ mutants were
- 329 significantly elevated as 4-16 μg/ml (Table 2). The data suggested that PhpP is involved in
- 330 β -lactam antibiotic resistance of *S. pneumoniae*.

331 Discussion

Kinases play key roles in signal transduction in both eukaryotes and prokaryotes and are 332 333 activated by either phosphorylation or dephosphorylation [44]. Unlike eukaryotes, histidine kinases but not serine/threonine/tyrosine (Ser/Thr/Tyr) kinases act as the major signal sensors 334 and transducers in prokaryotic bacteria [45]. Previous studies confirmed that StkP of S. 335 pneumoniae is an eukaryotic-type Ser/Thr protein kinase containing PBP-like domains in its 336 extracellular region and Ser/Thr phosphorylation sites in its intracellular region [46], and 337 PhpP and StkP of S. pneumoniae compose a StkP-PhpP signaling couple [25]. Operon is a 338 339 genic unit for transcription in prokaryotic genome that composed of promoter, operator and function-associated structural genes [47]. Our bioinformatic analysis showed that PhpP- and 340 StkP-encoding genes of S. pneumoniae are located in the same one operon for co-expression, 341 342 implying the close functional association between the two genes. As described in the introduction, StkP of S. pneumoniae is involved in PBP mutation-independent penicillin 343 resistance [22]. Therefore, the product of *phpP* gene, PhpP, may act as a protein phosphotase 344 to down-regulate StkP phosphorylation level by dephosphorylation to participate in 345 StkP-involved β -lactam antibiotic resistance. 346

According to the differences of amino acid sequences and structures, Ser/Thr protein phosphatases are classified into MPP and PPP families as well as the former contains Mg^{2+} or Mn^{2+} -dependent PP2C type phosphatases while the latter includes PP1, PP2A and PP2B type phosphatases [48]. PP2C type phosphatases has an extensive distribution in bacteria, yeasts, plants and mammalian cells to play various and complicated functions such as signaling transduction by dephosphorylation, cellular generation cycle regulation, monitoring DNA

353 damage and RNA splicing [48-50]. Our bioinformatic prediction showed that *phpP* gene of S. pneumoniae contains a PP2C type protein phosphatase domain. The recombinant expression 354 355 product of *phpP* gene (rPhpP) hydrolyzed both the general Ser/Thr phosphatase substrate p-NPP and PP2C type Ser/Thr protein phosphatase-specific substrate RRA(pT)VA in a 356 357 concentration-dependent manner but its RRA(pT)VA-hydrolyzed ability was blocked by the 358 PP2C type Ser/Thr protein phosphatase-specific inhibitor NaF. Previous studies confirmed that StkP and PhpP of S. pneumoniae compose a StkP/PhpP signaling couple in which StkP is 359 activated by Ser/Thr residual phosphorylation and PhpP could hydrolyze phosphoryl groups 360 361 in StkP in vitro [25,51]. However, we confirmed that the sublethal PCN and CTX induced the phosphorylation of StkP and the StkP phosphorylation levels in the $\Delta phpP$ mutants were 362 363 significantly higher than that in the wild-type strains. All the data suggested that the product 364 of S. pneumoniae phpP gene is a PP2C type Ser/Thr protein phosphatase to play a reverse regulation role in StkP/PhpP signaling couple by dephosphorylation of StkP. 365

Previous studies reported that penicillin-binding protein and STK-associated (PASTA) 366 367 superfamily domains in some proteins from many Gram-positive bacteria have a potential ability to bind to β -lactam antibiotics [52,53], and our bioinformatic analysis also revealed 368 that StkP of S. pneumoniae possesses four PASTA domains located its carboxyl terminal, 369 implying that the StkP may served as the β -lactam antibiotic receptor and β -lactam antibiotics 370 may cat as environmental activators of StkP-PhpP signaling couple. In the present study, the 371 sublethal PCN or CTX (1/4 MIC) caused the significant increase of stkP- and phpP-mRNA 372 levels and StkP phosphorylation. The data suggested that StkP-PhpP couple is a β -lactam 373 antibiotic-associated signaling pathway of S. pneumoniae. 374

375 Bacterial PBP is a group of peptidoglycan biosynthesis-associated transpeptidases, carboxypeptidases and endopeptidases that located on surface of bacteria [54]. β-lactam 376 377 antibiotics can bind to the peptidases to cause them inactivation by due to enzymatic molecular allosterism but PBP mutation can cause β-lactam antibiotic resistance due to the 378 decrease of β -lactam antibiotic-binding ability of PBP [55,56]. However, some S. 379 pneumoniae strains presented PBP mutation-independent β-lactam antibiotic resistance 380 [20-22]. In the present study, all the tested S. pneumoniae strains were sensitive to both PCN 381 and CTX (MICs: 0.02-0.5 μ g/ml). When the *phpP* genes were knockout, the $\Delta phpP$ mutants 382 became resistant to the two antibiotics (MICs: 4-16 µg/ml). The data imply that StkP 383 phosphorylation promotes but PhpP-based StkP dephosphorylation inhibits the generation of 384 PBP mutation-independent β -lactam antibiotic resistance. 385

386

387 Conclusions

Sublethal PCN and CTX can act as environmental inducers to cause the increase of *stkP* and *phpP* gene expression and StkP phosphorylation of *S. pneumoniae*. The product of *phpP* gene is a PP2C type Ser/Thr protein phosphatase to cause dephosphorylation of StkP that plays a reverse regulating role in StkP/PhpP signaling couple. Sublethal β -lactam antibiotics can act as environmental inducers of *phpP* and *stkP* gene expression and knockout of *phpP* gene caused the significant increase of β -lactam antibiotic resistance of *S. pneumoniae*.

394

396 Additional file

397 This manuscript has no additional files.

398

399 Abbreviations

S. pneumoniae: Streptococcus pneumoniae; PNC: penicillin; CTX: cefotaxime; quantitative 400 reverse transcription polymerase chain reaction: qRT-PCR; $\Delta phpP$: phpP gene-knockout 401 mutant; MIC: minimal inhibitory concentration; mRNA: messenger ribonucleic acid; PBP: 402 penicillin-binding proteins; STK: serine/threonine kinase; DNA: deoxyribonucleic acid; TH: 403 404 Todd-Hewitt; LB: Luria-Bertani; OD: optical density; PBS: phosphate buffered saline; RNA: ribonucleic acid; SDS-PAGE: sodium dodecyl sulfate polyacrylamide gel electropheresis; 405 CDD: conserved domain database; IPTG: isopropy-β-D-thiogalactoside; OA: Okadaic acid; 406 407 NaF: sodium fluoride; SD: standard deviation; PASTA: penicillin-binding protein and STK-associated; TB: Tris-HCl buffer. 408

409

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418

419 Availability of data and materials

- 420 The datasets analyzed during the current study are available from the corresponding author
- 421 on reasonable request.

422

423 Author's contributions

- 424 A.H.S. and J.Y. conceived and designed the experiments. Y.Y.H., Y.H.S. and P.D. performed
- 425 the experiments. Y.Y.H., Y.H.S. and X.X.L. analyzed the data. Y.Y.H., A.H.S. and J.Y. wrote
- the manuscript. A.H.S. obtained the funding. All authors have read and approved themanuscript.

428

429 Ethics approval and consent to participate

This study has no medical ethic problems. All the animal experimental protocols were
approved by the Ethics Committee for Animal Experiment of Hangzhou Medical College.

432

433 **Consent for publication**

434 Not applicable.

435

436 **Competing interests**

437 All the authors have no potential conflict of interest to declare.

438

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617 Legend

Fig. 1 Strategy for generation of *S. pneumoniae* $\Delta phpP$ mutant. See section of materials and methods in text for details.

- Fig. 2 Increase of *phpP* and *stkP*-mRNAs after treatment with sublethal PCN and CTX.
- 622 (A). Increase of *phpP*-mRNA induced by 1/4 MIC PCN, detected by qRT-PCR. Bars show
- 623 the mean \pm SD of three independent experiments. The *phpP*-mRNA levels in the
- 624 PCN-untreated S. pneumoniae strains (before treatment) were set as 1.0. *: p<0.05 vs the
- 625 *phpP*-mRNA levels in the PCN-untreated strains.
- 626 (B). Increase of *phpP*-mRNA induced by 1/4 MIC CTX, detected by qRT-PCR. The legend is
- 627 the same as shown in A but for CTX-induced *phpP*-mRNA detection.
- 628 (C). Increase of *stkP*-mRNA induced by 1/4 MIC PCN, detected by qRT-PCR. Bars show the
- 629 mean \pm SD of three independent experiments. The *stkP*-mRNA levels in the PCN-untreated *S*.
- 630 *pneumoniae* strains (before treatment) were set as 1.0. *: p < 0.05 vs the stkP-mRNA levels in
- 631 the PCN-untreated strains.
- (B). Increase of *stkP*-mRNA induced by 1/4 MIC of CTX, detected by qRT-PCR. The legend
- 633 is the same as shown in C but for CTX-induced *stkP*-mRNA detection.
- 634
- Fig. 3 Predicted characteristics of *S. pneumoniae phpP* and *stkP* genes.
- 636 (A). Structure and location of *phpP* gene product, predicted using TMHMM software.
- 637 (B). Structure and location of *stkP* gene product, predicted using TMHMM software.
- 638 (C). PP2C type phosphatase domain in phpP gene product (PhpP), predicted using NCBI

639 Database CDD software.

640 (D). STK and PASTA domains in *stkP* gene product (StkP), predicted using NCBI Database
641 CDD software.

642

- Fig. 4 Amplification and expression of *S. pneumoniae phpP* and *stkP* genes.
- (A). Amplification fragments of *phpP* genes from *S. pneumoniae* strains, determined by PCR.
- Lane M: DNA marker. Lanes 1-4: amplicoms of phpP genes from S. pneumoniae strains
- ATCC6306, SP5, SP9 and SP14 (738 bp). Lane 5: blank control.
- (B). Amplification fragments of *stkP* genes from *S. pneumoniae* strains, determined by PCR.
- ⁶⁴⁸ The legend is the same as shown in A but for *stkP* gene amplification (1977 bp).
- 649 (C). Expression and extraction effects of rPhpP, detected by SDS-PAGE. Lane M: protein
- 650 marker. Lane 1: wild-type *E. coli* BL21DE3. Lane 2: the expressed rPhpP (~28.3 kDa). Lane
- 3: the extracted rPhpP by Ni-NTA affinity chromatography.
- (D). Expression and extraction effects of rStkP, detected by SDS-PAGE. The legend is the

same as shown in C but for rStkP expression and extraction (~75.8 kDa).

654

Fig. 5 Powerful protein phosphatase activity of rPhpP from *S. pneumoniae*.

656 (A). Ability of rPhpP hydrolyzing phosphatase substrates p-NPP and RRA(pT)VA, 657 determined by spectrophotometry. Bars show the means \pm SD of three independent 658 experiments. p-NPP is a universal Ser/Thr phosphatase substrate while RRA(pT)VA is a 659 PP2C type Ser/Thr protein phosphatase-specific substrate.

660 (B). Km and Kcat values of rPhpP hydrolyzing RRA(pT)VA substrate, determined by

spectrophotometry. 5 μg rPhpP and 50, 100, 150, 200 or 250 μM RRA(pT)VA
phosphopeptide were used.

(C). Enzymatic activity of rPhpP after treatment with phosphatase inhibitors, determined by
spectrophotometry. NaF is a PP2C type Ser/Thr protein phosphatase inhibitor while OA is an
inhibitor of PP1, PP2A or PP2B type Ser/Thr phosphatases.

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667 Fig. 6 PCN/CTX-induced StkP phosphorylation and PhpP-caused StkP-dephosphorylation.

668 (A). Growth curves of the $\Delta phpP$ mutants and wild-type strains in TH broth, determined by

spectrophotometry. Bars show the means \pm SD of three independent experiments.

670 (B). Schematic diagram of PCR and sequencing results of the $\Delta phpP$ mutants.

671 (C). PhpP absence in the $\Delta phpP$ mutants, determined by Western Blot assay. Lanes 1-4:

672 immunoblotting bands of PhpP proteins from the wild-type S. pneumoniae strain ATCC 6306

and *S. pneumoniae* isolates SP1, SP5 and SP9. Lanes 5-8: no immunoblotting bands of PhpP

674 proteins in the $\Delta phpP$ mutants.

(D) Sublethal PCN-induced increase of StkP phosphorylation and decrease of StkP phosphorylation in the $\Delta phpP$ mutants, detected by spectrophotometry. Bars show the means \pm SD of three independent experiments. *: *p*<0.05 *vs* the StkP phosphorylation levels of the PCN-untreated wild-type strains and $\Delta phpP$ mutants. [#]: *p*<0.05 *vs* the StkP phosphorylation levels of the wild-type strains.

680 (E) Sublethal CTX-induced increase of StkP phosphorylation and decrease of StkP 681 phosphorylation in the $\Delta phpP$ mutants, detected by spectrophotometry. The legend is the 682 same as shown in D but for detection of CTX-induced StkP phosphorylation.

683

Primer	Sequence (5' to 3')	Purpose	Size (bp)
P-1	F: CTTCTGCCTSCTTCTGGTGT	phpP-mRNA detection	119
	R: TATTGGTGATTCGCGTATC		
S-1	F: TCGCTCGGTTTCAGAGATGTATGT	stkP-mRNA detection	190
	R: TGGGTTTTTGATTGATTTGTGTTC		
16S	F: CGACGATACATAGCCGACCTG	16S RNA segment as	145
	R: AAAACCTTCTTCACTCACGCG	qRT-PCR inner reference	
P-2	F: CGC <u>CATATG</u> (Nde I)GAAATTTCATTATTA	phpP gene detection and	738
	R: CGC <u>CTCGAG</u> (Xho I)TTCTGCATCCTCCTCGTTC	expression	
S-2	F: GCG <u>CATATG(Nde</u> I)ATCCAAATCGGCAAG	stkP gene detection and	1977
	R: GCGAAGCTT(Hind III)AGGAGTAGCTGAAGTTGT	[expression	
P-3	F: CGC <u>CTCGAG</u> (Xho I)ATATCGCTAGTGAAATGG	phpP gene segment for	495
	R: CGC <u>GGATCC(BamH I)CACTGGTTACAATATCAC</u>	knockout	
P-4	F: ACATGAATGTAAGGATATCATG	$\Delta phpP$ mutant identification	895
	R: GGATGTGCTGCAAGGCGATTA		
P-5	F: GCTTTCTTCATTAGAATCAATCC	$\Delta phpP$ mutant identification	907
	R: TCACTGCCACTTCTTCCCCATC		

Table 1. Sequences of primers used in this study. 684

F: forward primer. R: reverse primer. Underlined areas indicate the sites of endonucleases. 685

686

Table 2 MICs of PCN and CTX against S. pneumoniae strains. 687

Strain —	MIC (µg/ml)				
Stram	PCN	CTX			
Wild-type ATCC6306	0.05	0.12			
$\Delta phpP$ -6306	8	8			
Wild-type SP5	0.05	0.25			
$\Delta phpP$ -SP5	8	16			
Wild-type SP9	0.25	0.5			
$\Delta phpP$ -SP9	16	16			
Wild-type SP14	0.02	0.03			
$\Delta phpP$ -SP14	4	4			

688

















