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The Low-Molecular Weight Protein Arginine Phosphatase PtpB Affects Nuclease Production, Cell Wall Integrity, and Uptake Rates of Staphylococcus aureus by Polymorphonuclear Leukocytes

Mohamed Ibrahem Elhawy 1,2*, Virginie Molle 3*, Sören L. Becker 1* and Markus Bischoff 1,3

Abstract: The epidemiological success of Staphylococcus aureus as a versatile pathogen in mammals is largely attributed to its virulence factor repertoire and the sophisticated regulatory network controlling this virulon. Here we demonstrate that the low-molecular-weight protein arginine phosphatase PtpB contributes to this regulatory network by affecting the growth phase-dependent transcription of the virulence factor encoding genes/operons aur, nuc, and psma, and that of the small regulatory RNA RNAIII. Inactivation of ptpB in S. aureus SA564 also significantly decreased the capacity of the mutant to degrade extracellular DNA, to hydrolyze proteins in the extracellular milieu, and to withstand Triton X-100 induced autolysis. SA564 ΔptpB mutant cells were additionally ingested faster by polymorphonuclear leukocytes in a whole blood phagocytosis assay, suggesting that PtpB contributes by several ways positively to the ability of S. aureus to evade host innate immunity.

Keywords: Staphylococcus aureus; low-molecular-weight protein arginine phosphatase; PtpB; nuclease; whole blood phagocytosis assay; cell wall integrity; gene transcription; Triton X-100 induced autolysis; innate immunity

1. Introduction

Staphylococcus aureus is a major bacterial pathogen in humans and animals [1]. The ability of this Gram-positive bacterium to respond accurately to changing environments is one of the prerequisites for its success as a versatile pathogen in mammals [2]. The opportunistic pathogen is equipped with an armamentarium of virulence factors and a sophisticated network of regulatory molecules allowing it to control/fine-tune the expression of its virulon in a way to rapidly respond to changing conditions [3]. While virulence factor synthesis in S. aureus is mainly driven by one component systems such as the Sar family of DNA binding proteins and two-component system (TCS) response regulators [4], phosphatases are also known to contribute to this network. Earlier work demonstrated, for instance, that the serine/threonine phosphatase Stp1 directly contributes to virulence factor synthesis and infectivity of S. aureus by promoting the transcription of the α-hemolysin encoding gene hla [5]. More recent data indicate that the low-molecular-weight protein arginine phosphatase (LMW-PAP) PtpB might also contribute to staphylococcal pathophysiology by modulating the arginine phosphorylation states of regulators such as the stress response regulator CtsR and the global regulator MgrA, particularly in response to oxidative stress [6,7]. A deletion of ptpB in the S. aureus clinical isolate SA564 was additionally shown lately to reduce the capacity of the mutant to survive inside of macrophages and to cause infection in a murine-based S. aureus abscess model [8]. PtpB might thereby
form a regulatory circuit with McsB, the presumed protein arginine kinase of *S. aureus*, which is part of the *clpC* operon and known to affect the hemolytic and proteolytic activity of this bacterium [9]. The McsB homolog in *Bacillus subtilis* was shown to modulate the activity of McsB-targeted regulators by phosphorylating arginine residues within the DNA-binding domains and by marking proteins and aberrant polypeptide molecules for degradation [10,11]; similar functions are assumed for McsB in *S. aureus* [6]. Thus, by counteracting McsB-dependent arginine phosphorylation of the aforementioned global regulators, PtpB might affect the activity and stability of these transcription factors and thus the expression of stress response genes and virulence factors that are directly and/or indirectly controlled by these regulatory molecules. Additionally, PtpB might interfere with the degradation process of proteins phosphorylated at arginine residues. Notably, a functional classification of arginine phosphorylated proteins found in in vitro cultured *S. aureus* cells carried out by Junker et al. [6] revealed that nearly half (47%) of the identified arginine phosphoproteins in the *ptpB* mutant could be associated with virulence functions, supporting the idea that PtpB contributes to the regulatory network controlling the virulence factor synthesis in this bacterial pathogen.

In this study, we further strengthen this assumption by demonstrating that inactivation of *ptpB* in *S. aureus* strain SA564—a whole genome sequenced clinical human isolate of the worldwide distributed clonal complex 5 that harbors functional *agr* and *sigB* operons and a *saeS* allele [12,13]—affects the transcription of the virulence factor encoding genes *aur* (encoding the zinc-metalloprotease aureolysin), *nuc* (encoding thermonuclease) and *psm*α (encoding the phenol-soluble modulins α1-4), and of the small regulatory RNA RNAIII, one of the master regulators controlling exoprotein synthesis in this bacterium [14], which also encodes a protein, δ-hemolysin (Hld). Deletion of *ptpB* also impedes the capacity of *S. aureus* to evade phagocytosis by polymorphonuclear leukocytes (PMNs), to withstand Triton X-100 induced autolysis or lysostaphin mediated lysis, and decreases the exonuclease- and exoprotease activities of this bacterium.

2. Results and Discussion

2.1. PtpB Affects the Ability of *S. aureus* to Evade Phagocytosis by PMNs

Recent work demonstrated that PtpB contributes to the ability of *S. aureus* to survive inside macrophages [8]. However, it remains unknown whether PtpB also supports the capacity of the pathogen to evade innate immunity. In order to address this issue, we first investigated the impact of a *ptpB* deletion on attachment/phagocytosis of *S. aureus* SA564 by PMNs in whole blood (Figure 1).

When human blood was supplemented with fluorescent-labeled bacteria at a multiplicity of infection (MOI) of 50 per PMNs, a clear difference in uptake rates of *ptpB* negative and positive *S. aureus* SA564 cells by human PMNs was observed (Figure 1). After 30 min of coincubation in human whole blood, CFSE-stained SA564 isolates harboring a functional *ptpB* (wild-type and *cis*-complemented derivative) demonstrated significantly decreased rates of attachment/phagocytosis by PMNs compared to the CFSE-stained SA564 derivative lacking *ptpB* (SA564 Δ*ptpB*). These findings suggest that PtpB mediates a relevant protective effect against phagocytosis by PMNs in blood.
2.1. PtpB Affects the Ability of S. aureus to Evade Phagocytosis by PMNs

withstand Triton X-100 induced autolysis or lysostaphin mediated lysis, and decreases the RNAIII, which also encodes a protein,

Figure 1. Effect of PtpB on phagocytosis of S. aureus SA564 by polymorphonuclear leukocytes (PMNs). Carboxy fluorescein diacetate succinimidyl ester (CFSE)-stained cells of S. aureus SA564 (black boxes), SA564 ΔptpB (white boxes), and SA564 ΔptpB::ptpB (gray boxes) were added to lithium heparin anticoagulated fresh human blood at a multiplicity of infection of 50 per PMN, and incubated for 30 min at 37 °C and 1000 rpm. Attachment/uptake of bacteria by PMNs was analyzed by flow cytometry as outlined in Material and Methods. The data are representative of three biological replicates carried out in triplicate. Data are presented as box and whisker plots showing the interquartile range (25–75%; box), median (horizontal line), and whiskers (bars; min/max). ** p < 0.01 (Kruskal–Wallis test followed by Dunn’s post hoc test).

2.2. PtpB Promotes the Transcription and Secretion of Nuclease in S. aureus

Neutrophils are the main pathogen-fighting immune cells in our blood system [15]. Besides their capacity to ingest circulating pathogens, activated neutrophils exert a number of cytotoxic functions such as the production of reactive oxygen and nitrogen species, the release of antimicrobial peptides, and the formation of extracellular DNA nets called neutrophil extracellular traps (NETs) [15]. The latter bacteria-capturing and killing mechanism is counteracted by S. aureus, among others, via the secretion of nucleases and the extracellular adherence protein (Eap), which degrade and aggregate NETs, respectively [16,17]. We have recently shown that PtpB supports the capacity of S. aureus to cope with oxidative and nitrosative stress [8].

When cell suspensions of SA564 and the cis-complemented ΔptpB::ptpB derivative were spotted on DNase test agar plates and incubated at 37 °C for 24 h, clearly visible lytic areas around the bacterial growth areas were observed that were in comparable ranges (Figure 2a,b). However, when equal volumes of SA564 ΔptpB cell suspensions were spotted onto the DNase test agar plates, significantly smaller lytic areas surrounding the bacterial growth zones were observed, suggesting that the ΔptpB mutant produces and/or secretes lower amounts of nucleases into the extracellular milieu.

To elucidate whether the reduced nuclease activity observed with the ΔptpB mutant might be due to PtpB-induced changes in nuc transcription, we next assayed the transcription rates of nuc in SA564 and its ΔptpB derivative upon growth in tryptic soy broth (TSB) over time. Specifically, total RNAs were obtained from cell populations grown for 3 h (exponential growth phase), 5 h (transition phase), and 8 h (early stationary phase), respectively (Figure 2c). In line with our DNase activity findings, we observed a significantly reduced transcription of nuc in SA564 ΔptpB cells at two of the three growth stages analyzed, if compared to the wild-type transcript level, while cells of the cis-complemented derivative produced nuc transcript level that were comparable to the wild-type. Taken together, these findings suggest that PtpB contributes positively to the exonuclease activity of S. aureus by supporting nuc transcription.
extracellular adherence protein (Eap), which degrade and aggregate NETs, respectively [16,17]. We have recently shown that PtpB supports the capacity of neutrophil extracellular traps (NETs) [15]. The latter bacteria-capturing and killing mechanism is counteracted by the release of antimicrobial peptides, and the formation of extracellular DNA nets called oxidative and nitrosative stress [8]. To test whether PtpB might also interfere with other functional activities, we next tested whether the LMW-PAP might modulate the capacity of neutrophil-derived cytotoxic activities, we next tested whether the LMW-PAP might

2.2. PtpB Promotes the Transcription and Secretion of Nuclease in S. aureus

SA564 derivative lacking the ptpB gene. Quantitative transcript analyses of nuc by qRT-PCR in SA564 (black bars), SA564 ΔptpB (white bars), and SA564 ΔptpB::ptpB (gray bars) cells grown to the time points indicated. Transcript rates were quantified in reference to the transcription of gyrase B (in copies per copy of gyrB). Data are presented as mean ± SD of five biological replicates.

Figure 2. Effect of PtpB on extracellular DNase activity and nuc transcription of S. aureus SA564. (a,b) Impact of PtpB on extracellular DNase activity. Overnight cultures of S. aureus strain SA564 (black bars), its isogenic ΔptpB mutant (white bars), and the cis-complemented derivative (gray bars) were normalized with fresh tryptic soy broth (TSB) medium to an optical density at 600 nm (OD_{600}) of 12, and 5 µL of the suspensions spotted on DNase-Test-Agar plates. Inoculated plates were cultured for 24 h at 37 °C, and lytic areas were determined. (a) Representative image of one experiment. (b) Diameter of lytic areas. The data presented are the mean ± SD of three biological experiments done in duplicate. ** p < 0.01 (Kruskal–Wallis test followed by Dunn’s post hoc test). (c) Growth kinetics of S. aureus strain SA564 in TSB. Cells were cultured at 37 °C and 225 rpm at a culture to flask volume of 1:10. Data represent the mean OD_{600} readings ± SD at the time points indicated (n = 3). Time points of sampling for the transcriptional analyses are indicated by arrows. (d) Impact of PtpB on transcription of the thermonuclease encoding nuc gene. Quantitative transcript analyses of nuc by qRT-PCR in SA564 (black bars), SA564 ΔptpB (white bars), and SA564 ΔptpB::ptpB (gray bars) cells grown to the time points indicated. Transcript rates were quantified in reference to the transcription of gyrase B (in copies per copy of gyrB). Data are presented as mean ± SD of five biological replicates. * p < 0.05; ** p < 0.01 (Kruskal–Wallis test followed by Dunn’s post hoc test between wild-type and mutants at a given time point).

2.3. PtpB Promotes the Transcription of the Aureolysin Encoding Gene aur in S. aureus

Another major virulence determinant interfering with innate host immunity and supporting the survival of S. aureus in blood is the zinc-metalloprotease aureolysin, which cleaves, among others, various factors of the human complement system [18]. To test whether PtpB might aid the immune evasion of S. aureus by modulating aureolysin expression, we tested the transcription of aur during growth in TSB (Figure 3a). This transcriptional analysis revealed a significantly decreased number of aur transcripts in exponential growth phase cells of the ptpB deletion mutant when compared with cells of the wild-type and the cis-complemented derivative harvested at the same growth stage, respectively. However, aur transcript rates were rather comparable between wild-type, ΔptpB mutant, and cis-complemented cells obtained from later growth stages (i.e., 5 and 8 h of growth), suggesting that PtpB contributes to aur expression primarily in S. aureus exponential growth phase cells.
Besides its complement factors degrading functions [18], aureolysin is also known for its capacity to process and thus activate serine protease SspA (syn. V8 protease), another major exoprotease produced by *S. aureus*, which in turn cleaves the propeptide from the SspB zymogen to create the active cysteine protease SspB (syn. staphopain), a process also known as staphylococcal proteolytic cascade [19].

Wondering whether PtpB might also affect the exoproteolytic capacity of *S. aureus*, we conducted a series of skim milk agar-based protease assays (Figure 3c,d). Stationary phase cell suspensions of SA564 normalized to an OD$_{600}$ of 12 spotted onto the skim milk TSA plates produced only very small proteolytic areas during growth.

To get an idea about whether the impact of PtpB on *aur* transcription might correlate with the expression of the LMW-PAP, we determined the transcription rates of *ptpB* in SA564 during growth in TSB (Figure 3b). These analyses revealed that *ptpB* is transcribed on a rather constant level throughout growth (fold changes in relative transcription rates between growth stages < 2), suggesting that the growth phase-dependent effect of PtpB on *aur* transcription is not likely to be determined by the expression rates of the LMW-PAP during growth.

Besides its complement factors degrading functions [18], aureolysin is also known for its capacity to process and thus activate serine protease SspA (syn. V8 protease), another major exoprotease produced by *S. aureus*, which in turn cleaves the propeptide from the SspB zymogen to create the active cysteine protease SspB (syn. staphopain), a process also known as staphylococcal proteolytic cascade [19].
milk supplemented tryptic soy agar (TSA) plates produced only very small proteolytic areas surrounding the bacterial growth zones after 48 h of incubation at 37 °C (Figure 3c). However, when normalized stationary phase cell suspensions of SA564 ΔptpB were spotted onto the skim milk TSA plates, significantly smaller proteolytic areas were observed, while cell suspensions of the cis-complemented SA564 ΔptpB::ptpB produced proteolytic areas on skimmed milk TSA plates that were in a comparable range to those seen with the wild-type cultures (Figure 3c,d). These findings suggest that PtpB affects the proteolytic capacity of S. aureus, potentially via the modulation of aur transcription.

2.4. PtpB Reduces the Autolytic Activity of S. aureus

Activated neutrophils secrete, among others, antimicrobial peptides (e.g., defensins) which exert potent in vitro microbicidal activity against S. aureus through mechanisms involving cell membrane destabilization, interference with intracellular targets, and activation of autolysins [20]. To test whether and how PtpB might influence the autolytic behavior of S. aureus, we next studied the impact of the ptpB deletion on the autolytic behavior of SA564 in Triton X-100- and lysostaphin-induced lysis assays, respectively (Figure 4).

![Figure 4. Effect of a ptpB deletion on the Triton X-100 induced autolysis and lysostaphin-mediated lysis of S. aureus SA564.](image)

(a) Cells of S. aureus strain SA564 (black symbols) and its isogenic ΔptpB mutant (white symbols) were cultured in TSB at 37 °C and 225 rpm to an OD₆₀₀ of 0.7, washed twice in ice-cold water, and resuspended in 0.05 M Tris-HCl (pH 7.2) containing 0.01% (vol/vol) Triton X-100. Triton X-100 induced autolysis was measured as the decline of optical density at 600 nm (OD₆₀₀) versus time and is expressed as the percent of the initial optical density (n = 6 biological replicates). (b) Effect of a ptpB deletion on the lysostaphin-mediated lysis. Cells of S. aureus strain SA564 (black symbols) and its isogenic ΔptpB mutant (white symbols) were cultured in TSB at 37 °C and 225 rpm to an OD₆₀₀ of 3, washed twice in PBS, and resuspended in PBS to an OD₆₀₀ of 1. The PBS cell suspensions were supplemented with 250 ng/mL lysostaphin and incubated under static conditions at 30 °C for 60 min. Lysostaphin-mediated lysis was measured as the decline of OD₆₀₀ versus time and is expressed as the percent of the initial optical density (n = 5 biological replicates). Data are presented as box and whisker plots showing the interquartile range (25–75%; box), median (horizontal line), and whiskers (bars; min/max). *p < 0.05; **p < 0.01 (Ordinary two-way ANOVA followed by Holm–Sidak’s multiple comparison test).

When washed exponential growth phase cells of SA564 and its ΔptpB mutant were challenged with a low dose of Triton X-100, respectively, cells of the ptpB deletion mutant autolyzed to a significantly larger extent after 3 h of coincubation with the detergent than wild-type cells (Figure 4a). A similar behavior was observed when SA564 and its ΔptpB mutant were challenged with the glycyl-glycyl endopeptidase lysostaphin, a S. aureus pentaglycine cross-bridges cleaving exoenzyme from Staphylococcus simulans bv. staphylolyticus [21]. Washed and PBS resuspended late exponential growth phase cells of the ptpB deletion mutant lysed again significantly faster than the wild-type cells (Figure 4b), indicating that PtpB is likely to affect the cell wall composition of S. aureus. One possible explanation for both observations is that PtpB might interfere with the production and/or...
activation of autolysins (e.g., endogenous murein hydrolases), as has been suggested for the lysostaphin resistance of some glycopeptide-intermediate-resistant \textit{S. aureus} (GISA) [22].

As previous studies have already demonstrated that the global regulator MgrA acts as a repressor of autolysins in \textit{S. aureus} [23,24], and that MgrA serves as a substrate for PtpB [6,7], it was tempting to speculate that PtpB might affect the autolytic behavior of \textit{S. aureus} via modulation of MgrA activity. In order to address this hypothesis, we first studied the impact of the \textit{ptpB} deletion on the transcription of MgrA regulated genes [25,26], focusing on genes whose products are involved in the autolytic behavior of \textit{S. aureus} (Figure 5). Specifically, the transcription of \textit{atlR} (encoding a MarR family transcriptional regulator repressing \textit{atl} transcription), \textit{fmtB} (encoding a cell wall-anchored protein involved in methicillin resistance and cell wall biosynthesis), \textit{lrgA} (encoding the holin-like murein hydrolase regulator LrgA), and \textit{lytN} (encoding the cell-wall hydrolase LytN) was analyzed, all of which were reported to directly or indirectly affect the autolytic behavior of \textit{S. aureus} [27–29].

![Figure 5. Effect of a \textit{ptpB} deletion on the transcription of MgrA-regulated genes affecting autolysis of \textit{S. aureus}. Quantitative transcript analyses of MgrA regulated genes by qRT-PCR in SA564 (black bars) and SA564 \textit{ΔptpB} (white bars) cells grown in TSB at 37 °C and 225 rpm to the time points indicated. Growth-phase-dependent transcript rates of MgrA-regulated genes \textit{atlR} (a), \textit{fmtB} (b), \textit{lrgA} (c), and \textit{lytN} (d). Transcript rates were quantified in reference to the transcription of gyrase B (in copies per copy of \textit{gyrB}). Data are presented as mean + SD of five biological replicates. * \textit{p} < 0.05; ** \textit{p} < 0.01 (Mann–Whitney-U test between wild-type and mutant at a given time point).

Notably, all four MgrA-regulated genes were transcribed in a significantly different manner in SA564 \textit{ΔptpB} cells than in wild-type cells. Importantly, genes reported to be repressed by MgrA (i.e., \textit{atlR}, \textit{fmtB}, and \textit{lytN}) were all transcribed on higher levels in the \textit{ptpB} deletion mutant, while transcription of \textit{lrgA}, which is positively affected by MgrA activity [25], was increased in wild-type cells throughout growth. However, while transcription of \textit{atlR}, \textit{fmtB}, and \textit{lrgA} was affected by the \textit{ptpB} deletion basically at all three growth stages (Figure 5a–c), this was not the case with \textit{lytN}. Transcription of the cell-wall hydrolase encoding gene was only affected by the \textit{ptpB} deletion during exponential growth phase and transition phase, but not in the early stationary phase (Figure 5d), suggesting that \textit{lytN} expression in the stationary phase cells is dominated by PtpB/MgrA-independent mechanisms. Taken together, these data indicate that the PtpB-driven dephosphorylation...
of MgrA phosphoarginine residues observed by Junker et al. [6] is likely to affect MgrA activity, and supports our hypothesis that PtpB interferes with autolysis of S. aureus via modulation of MgrA activity.

2.5. PtpB Does Not Alter MgrA Activity per Se

Given that MgrA also promotes the transcription of aur and nuc [25], which were also found to be transcribed in significantly higher levels in wild-type cells when compared to ΔptpB mutant cells (Figures 2c and 3), we wondered whether PtpB might affect the transcription of the whole MgrA regulon. In order to address this hypothesis, we additionally tested the transcription of two MgrA-regulated surface factor encoding genes, ehb (encoding extracellular matrix-binding protein Ebh) and spa (encoding immunoglobulin G binding protein A), both of which are known to be strongly repressed by MgrA activity on the transcriptional level [26]. In contrast to findings obtained for the MgrA-regulated genes affecting autolysis (Figure 5), we surprisingly observed no clear differences in transcription rates of ehb and spa between the wild-type and the ptpB deletion mutant at any time point analyzed (Figure 6).

As MgrA directly represses ehb transcription [26], our findings presented here indicate that PtpB-driven dephosphorylation of MgrA does not alter the activity of the transcription factor towards its regulon per se, but might be important for the expression of a subset of MgrA-regulated factors, particularly of those involved in autolysis. One explanation for our observations might be that PtpB-mediated dephosphorylation of MgrA arginine residues influences the activity of the regulator by differentiating the binding specificity among target gene promoters, depending on the phosphorylation status at arginine residues. In such a scenario, target genes with a high binding affinity for MgrA would not be affected in their expression by the arginine phosphorylation status of the regulator and thus would not display a change in transcription depending on the PtpB status of the cell.

2.6. PtpB Alternates the Transcription of Some but Not All SarA Regulated Genes

Transcription of aur, lrgA, and nuc is also reported to be affected by SarA [30–32], suggesting that PtpB might also interfere with the transcription of the aforementioned genes via modulation of SarA activity, although SarA was not identified as a direct substrate for McsB/PtpB [6,7]. To test whether and how PtpB affects the transcription of the SarA regulon in SA564, we assayed the transcription rates of three additional SarA target genes, hla (encoding α-hemolysin), hld (encoding δ-hemolysin), and splB (encoding serine protease SplB; Figure 7) [32–34], which all contain a putative Sar box within the gene/operon’s promoter region [32]. Similar to the situation seen with MgrA-regulated genes, we found
significant changes in transcription rates in ptpB lacking cells for some but not all SarA-regulated genes. While transcription of the hld encoding RNAIII was affected by the ptpB deletion during the exponential growth phase and early stationary phase (Figure 7b), this was not the case with hla nor splB (Figure 7a,c). Notably, exponential growth phase cells of the ptpB mutant produced lower RNAIII transcript rates than wild-type cells, while the opposite was encountered with early stationary phase cells, suggesting PtpB to modulate the transcription of the agr locus in SA564 by several means.

![Figure 7](image-url) Effect of a ptpB deletion on the transcription of SarA- and/or AgrA-regulated genes in S. aureus SA564. (a-d) Quantitative transcript analyses of SarA- and/or AgrA-regulated genes by qRT-PCR in SA564 (black bars) and SA564 ΔptpB (white bars) cells grown in TSB at 37 °C and 225 rpm to the time points indicated. Growth phase-dependent transcript rates of Sar box containing genes hla (a), hld (b), and splB (e), and of the AgrA-driven psma cluster (d). Transcript rates were quantified in reference to the transcription of gyrB in copies per copy of gyrB. Data are presented as mean + SD of five biological replicates. * p < 0.05; ** p < 0.01; ns, not significant (Mann–Whitney-U test between wild-type and mutant at a given time point). (e,f) Overnight cultures of S. aureus strain SA564 (black bar), its isogenic ΔptpB mutant (white bar), and the cis-complemented SA564 ΔptpB:ptpB derivative (gray bar) were normalized with fresh TSB medium to an OD600 of 12, and 5 µL of the suspensions spotted on sheep blood agar plates. Inoculated plates were cultured for 24 h at 37 °C, and hemolytic areas were determined. (e) Representative image of one experiment. (f) Diameter of hemolytic areas. The data presented are the mean + SD of three biological experiments done in duplicate. ns, not significant (Kruskal–Wallis test followed by Dunn’s post hoc test).

One potential factor contributing to this phenomenon could be AgrA, the response regulator of the agr locus, and the main transcription factor driving RNAIII transcription [35]. Since AgrA is also known to control the expression of the phenol-soluble modulin operons psma and psmb [36], we additionally tested whether psma transcription in SA564 is altered by the ptpB deletion (Figure 7d). In line with our RNAIII transcription data (Figure 7b), we observed significantly increased psma transcript rates in early stationary phase cells of the ΔptpB mutant, suggesting that PtpB might modulate RNAIII and psma transcription in stationary phase cells via AgrA.

To support our transcriptional findings indicating that PtpB does not markedly alter the expression of α-hemolysin in SA564, we tested the hemolytic activity of our strain triplet on sheep blood agar plates (Figure 7e,f). In line with our hla transcription findings,
we did not encounter clear differences in the hemolytic areas surrounding the growth zones for all three SA564 derivatives.

Together with the observations that SarA was not identified to be phosphorylated at arginine residues [6,7], our transcriptional findings made for the Sar box-containing genes *hla*, *spa*, and *splB* suggest that PtpB is not likely to directly modulate SarA activity, albeit of the fact that the transcription rates of other Sar box containing genes such as *aur*, *hld*, and *nuc* (Figures 2c, 3 and 7b) were markedly affected by the *ptpB* deletion. The differences in transcription rates observed for the latter genes between the wild-type and the *ptpB* deletion mutant might be attributed to other direct PtpB mediated effects on regulatory factors such as MgrA and AgrA, and/or indirect PtpB effects on regulators such as ArlR, CodY, SaeR, SarR, SarV, SarX, and SarZ, which all were found to be expressed to different extents in wild-type and *ptpB* deletion mutant cells of *S. aureus* strain COL under non-stress conditions [6].

3. Conclusions

Protein posttranslational modifications such as reversible phosphorylation by kinases/phosphatases is a common mechanism employed by bacteria and eukaryotes to modulate the activity of enzymes and regulatory molecules, which is also utilized by *S. aureus* to adjust central metabolic pathways and virulence factor synthesis [37]. The serine/threonine protein kinase-phosphatase pair Stk1-Stp1 is, for instance, known for its ability to modulate the activities of the SarA family transcription factors MgrA, SarA, and SarZ [38], with Stp1 promoting infectivity, while Stk1 attenuates infectivity [5,38]. Stk1-driven phosphorylation of the catabolite control protein A was additionally shown to inhibit the DNA-binding capacity of the master regulator of carbon catabolite repression in *S. aureus*, thereby modulating the expression of metabolic and virulence genes [39], and infectivity [40]. Our findings, presented here and elsewhere [8], demonstrate that the deletion of *ptpB* in the clinical *S. aureus* isolate SA564 alters the transcription of various genes/operons whose products are involved in stress adaptation and infectivity, suggesting that PtpB-driven removal of phosphates from arginine phosphosites is another posttranscriptional mechanism utilized by this pathogen to fine-tune the expression and activity of its virulon, in order to successfully adapt to the diverse host environmental conditions encountered by the bacterium during infection. Given the clear impact of PtpB on the transcription of specific virulence determinants shown here, its impact on *S. aureus* to cause disease in mice [8], and the fact that several low molecular weight protein tyrosine phosphatase inhibitors are currently in development to combat diseases such as cancer, diabetes/obesity, and bacterial infections [41–45], PtpB might constitute an additional interesting target for drug development against this notorious human nosocomial pathogen.

4. Materials and Methods

4.1. Bacterial Strains, Media, and Growth Conditions

The bacterial strains used in this study are listed in Table 1. *S. aureus* isolates were plated on Tryptic Soy Agar (TSA; BD, Heidelberg, Germany), or grown in Tryptic Soy Broth (TSB; BD) medium at 37 °C and 225 rpm with a culture to flask volume of 1:10.

Table 1. Strains used in this study.

| Strain         | Description                           | Reference |
|---------------|---------------------------------------|-----------|
| SA564         | *S. aureus* clinical isolate, wild type | [12]      |
| SA564 Δ*ptpB* | SA564 Δ*ptpB*:lox66-erm(B)-lox71; ErmR | [8]       |
| SA564 Δ*ptpB*:ptpB | *cis*-complemented SA564 Δ*ptpB* derivative | [8]       |

1 ErmR, erythromycin-resistant.
4.2. Human Whole Blood Phagocytosis Assay

The uptake of *S. aureus* cells by PMNs in whole blood was performed essentially as described in [46]. Overnight cultures of *S. aureus* strains SA564, SA564 ΔptpB, and SA564 ΔptpB::ptpB were inoculated into fresh TSB to an optical density at 600 nm (OD_{600}) of 0.05 and grown at 37 °C and 225 rpm to mid-exponential growth-phase (i.e., 2 h). Cultures were centrifuged at 10,000 × g for 5 min, bacterial pellets washed three times with phosphate buffered saline (PBS; Thermo Fisher, Dreieich, Germany), and subsequently stained with a 50 µM carboxy fluorescein diacetate succinimidyl ester (CFSE; Invitrogen, Darmstadt, Germany)-PBS solution for 15 min at 37 °C and 1000 rpm. CFSE-stained bacterial cells were afterwards washed again three times with PBS to remove unbound dye, and adjusted to an OD_{600} of 1. Fresh human whole blood was withdrawn from healthy donors and anticoagulated with lithium heparin (S-Monovette; Sarstedt, Nümbrecht, Germany). The PMN contents of the blood samples were determined using the RAL DIFF-QUICK kit (RAL Diagnostics, Martillac, France) according to the manufacturer’s recommendations, and blood samples were substituted with fluorescent-labeled bacteria at an MOI of 50 per PMN. Infected blood samples were cultured in the dark at 37 °C and 1000 rpm for 30 min and subsequently placed into 5 mL round bottom polystyrene tubes (BD). Erythrocytes were lysed by adding FACS lysis solution (BD), and lysed cell debris was removed by centrifugation at 450 × g for 5 min. The cell pellets were resuspended in PBS supplemented with 2% fetal calf serum (PAA, Pasching, Germany) and 0.05% sodium azide, and subjected to flow cytometry using a FACSCalibur (BD) system. PMNs were gated using the CellQuest Pro Software version 4.02 (BD), and the mean fluorescence intensity (MFI) per PMN was recorded, indicating the number of bacteria that were attached to or ingested by the leukocyte.

4.3. Extracellular DNase-, Hemolytic- and Proteolytic Activity Assays

Overnight cultures of the *S. aureus* SA564 strain triplet were adjusted for all three assays with fresh TSB to an OD_{600} of 12. For extracellular DNase activity measurements, 5 µL of the adjusted bacterial suspensions were spotted on DNase-Test-Agar plates (Carl Roth, Karlsruhe, Germany) and incubated for 24 h at 37 °C. Lytic zones were visualized by overlaying the agar with 1N HCl to precipitate undigested DNA. The hemolytic activities of the bacterial cell suspensions were tested by spotting 5 µL aliquots of the bacterial suspensions on TSA II plates supplemented with 5% Sheep Blood (BD), and diameters of hemolytic zones were determined after 24 h of incubation at 37 °C. The proteolytic activity of the bacterial cell suspensions was determined by spotting 2 µL aliquots of the bacterial suspensions on TSA plates supplemented with 10% skim milk (BD). Variations in zones of proteolysis were evaluated after incubating the plates for 48 h at 37 °C.

4.4. Triton X-100 Induced Autolysis Assay

Triton X-100 induced autolysis of *S. aureus* was assayed using a modified version of the protocol described in [47]. Cells of *S. aureus* SA564 and its ptpB deletion mutant were cultured in TSB at 37 °C and 225 rpm to an OD_{600} of 0.7, washed twice in ice-cold water, and suspended in 0.05 M Tris-HCl (pH 7.2) containing 0.01% (vol/vol) Triton X-100 (Merck, Darmstadt, Germany). The cell suspensions were incubated at 30 °C and 225 rpm and the OD_{600} measured every 30 min. Triton X-100 induced autolysis was determined as the decline of optical density versus time and is expressed as the percent of the initial optical density.

4.5. Lysostaphin Induced Autolysis Assay

The lysostaphin-induced autolysis of *S. aureus* was assayed as described in [48]. Cells of *S. aureus* SA564 and SA564 ΔptpB were cultured in TSB at 37 °C and 225 rpm to an OD_{600} of 3, washed twice in PBS, and resuspended in PBS to an OD_{600} of 1. Cell suspensions were supplemented with 250 ng/mL of lysostaphin (Dr. Petry Genmedics GMBH, Reutlingen, Germany) and incubated under static conditions at 30 °C for 60 min. The OD_{600} was...
measured every 10 min. Lysostaphin-induced autolysis was determined as the decline of optical density versus time and is expressed as the percent of the initial optical density.

4.6. qRT-PCR Analyses

RNA isolation, cDNA synthesis, and qRT-PCR were carried out as previously described [49], using the primer pairs listed in Table 2. Transcripts were quantified in reference to the transcription of gyrB using the $2^{-\Delta\Delta C_t}$ method [50].

Table 2. qRT-PCR primer used in this study.

| Gene Target | Primer | Sequence (5′-3′) |
|-------------|--------|-----------------|
| *atlR*      | forward reverse | AACTTATTACACTGACTAACAATG TGTCATAATTCTTACTAATAAG|
| *aur*       | forward reverse | AATAGATTTGACGTTGATTTT AATTGCTTGAATTTACCTCCTGATG |
| *ebh*       | forward reverse | GTATAATGAACAGACTGAGATG CCGTAGATGGATATGACTATT |
| *fmtB*      | forward reverse | GACTGATCCGATGTGGA AACGGTGACGTGCAATA |
| *gyrB*      | forward reverse | GACTGATCCGATGTGGA AACGGTGACGTGCAATA |
| *hla*       | forward reverse | AACCAGGATATGGCAATCACAT CTCGCTCTCTTACAGCCATT |
| *hld*       | forward reverse | AGGAGTGGTTCAATGGCACAAG TGTCGATATCCATTTACATAGTCAG |
| *lrgA*      | forward reverse | GCCGGATATCGTTGTAAATCTT AAATGGTCCTTGCAATGAC |
| *lytN*      | forward reverse | CTATGCGTTTAAATGGCATATG ATCTGAACCTTGTGCAAACCATTA |
| *nuc*       | forward reverse | TAGCTCAGCAAATGGCATACAA GAACACTTTTTGACAGCCATTATC |
| *psma*      | forward reverse | ATCAACAACACTACATGGTAAATCAAC GCCATCCGTTTTCTGTCCTTG |
| *ptpB*      | forward reverse | AGGCCATAGCCGAAATAGATTG AAATTTGATGTTGCGATAACCTCTC |
| *spa*       | forward reverse | TACCTATACGTTGTGCGATGA GGTCGTTTTAAGACTTTTG |
| *splB*      | forward reverse | AAGGTAATGTTGATTTACAT CCAGACTGATACATCTCATTT |

4.7. Statistical Analyses

The statistical significance of changes between groups was determined using the GraphPad software package Prism 6.01. $p$ values < 0.05 were considered statistically significant.

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Institutional Review Board Statement: The study was conducted according to the guidelines of the Declaration of Helsinki and approved by the Ethics Committee of the Medical Association of Saarland (code number 39/20, approved 25 February 2020).

Informed Consent Statement: Informed consent was obtained from all subjects involved in the study.

Data Availability Statement: The datasets generated during and/or analyzed during the current study are available from the corresponding author on reasonable request.

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