Distinct virulence ranges for infection of mice by *Bordetella pertussis* revealed by engineering of the sensor-kinase BvgS

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Abstract

The whooping cough agent *Bordetella pertussis* coordinately regulates the expression of its virulence factors with the two-component system BvgAS. In laboratory conditions, specific chemical modulators are used to trigger phenotypic modulation of *B. pertussis* from its default virulent Bvg\(^+\) phase to avirulent Bvg\(^-\) or intermediate Bvg\(^i\) phases, in which no virulence factors or only a subset of them are produced, respectively. Whether phenotypic modulation occurs in the host remains unknown. In this work, recombinant *B. pertussis* strains harboring BvgS variants were tested in a mouse model of infection and analyzed using transcriptomic approaches. Recombinant BP-Bvg\(^-65\), which is in the Bvg\(^i\) phase by default and can be up-modulated to the Bvg\(^+\) phase *in vitro*, could colonize the mouse nose but was rapidly cleared from the lungs, while Bvg\(^+\)-phase strains colonized both organs for up to four weeks. These results indicated that phenotypic modulation, which might have restored the full virulence capability of BP-Bvg\(^-65\), does not occur in mice or is temporally or spatially restricted and has no effect in those conditions. Transcriptomic analyses of this and other recombinant Bvg\(^i\) and Bvg\(^+\)-phase strains revealed that two distinct ranges of virulence gene expression allow colonization of the mouse nose and lungs, respectively. We also showed that a recombinant strain expressing moderately lower levels of the virulence genes than its wild type parent was as efficient at colonizing both organs. Altogether, genetic modifications of BvgS generate a range of phenotypic phases, which are useful tools to decipher host-pathogen interactions.

Introduction

*Bordetella pertussis* is the agent of an acute respiratory disease, whooping cough. Despite the current global vaccination coverage of approximately 86% of the population, this bacterium...
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Still causes 16 million cases and 200,000 deaths per year [1]. To colonize the human respiratory tract, B. pertussis produces a number of virulence factors, notably adhesins and toxins, whose expression is regulated by a two-component system called BvgAS [2]. BvgS is a sensor-kinase protein that auto-phosphorylates and transfers the phosphoryl group via a complex cascade of phosphorylation to BvgA, the response regulator. BvgA acts as a canonical transcriptional activator in its phosphorylated form [3].

The virulence status of B. pertussis is defined by three phases during which specific genes are expressed, a phenomenon referred to as phenotypic modulation [4]. At 37°C, in standard culture conditions, B. pertussis expresses all its virulence-activated genes (‘vags’), which defines the Bvg⁺ phase. In laboratory conditions, in presence of so-called negative modulators such as nicotinate or MgSO₄ at millimolar concentrations, or in conditions of nutrient restriction, the bacteria shift to the avirulent Bvg⁻ phase [5]. During this phase, the so-called virulence-repressed genes (‘vrgs’) are up-regulated, while the vags are silent. An intermediate phase (Bvgᵢ) was described at intermediate concentrations of modulators, in which a subset of vags, including bvgAS and genes coding for adhesins, are expressed [6]. Those were defined as ‘early’ vags, as they are the first genes of the Bvg regulon to be up-regulated following a shift from modulating to non-modulating conditions [7, 8]. In addition to the early vags, at least one gene specific of the Bvg’ phase, bipA, is overexpressed in the Bvg’ phase [9, 10]. Phenotypic modulation depends on the concentration of phosphorylated BvgA, itself related to the enzymatic activity of BvgS [4]. In basal conditions, BvgS functions as a kinase, while in modulated conditions it shifts to phosphatase activity [3, 11–13]. The signals that may cause the phase shift from Bvg’ to Bvg⁺ or Bvg⁻ in the host respiratory tract remain unknown.

BvgS is a large, dimeric protein that serves as a prototype for a family of poorly characterized Venus flytrap domains (VFT)-containing sensor-kinases [12, 14]. In the homodimer, each monomer is composed of two periplasmic VFT domains possibly involved in signal perception, followed by a transmembrane segment, a cytoplasmic Per-ArnT-Sim (PAS) domain, a histidine-kinase domain and two other domains, the receiver and the phosphotransfer domains, involved in a phosphorelay to BvgA. We have shown that the VFT1 domains of BvgS open and close, like typical VFT proteins, while the VFT2 domains are permanently closed [12]. Rather puzzlingly, however, negative chemical modulators such as nicotinate bind to the latter domain [13, 15]. Artificially closing the VFT1 domains strongly reduces BvgS kinase activity [12]. As VFT domains generally close upon binding their specific ligands [16, 17], this observation suggests that the VFT1 domains might perceive chemical signals that down-modulate BvgS activity at some stage of the infection.

In BvgS and the majority of its homologs, the transmembrane domain is followed by a two-helix linker called linker 1 that leads to the PAS domain. This is followed by a second two-helix linker, linker 2, that leads to the α-helical Dimerization and Histidine phosphorylation moiety (DHp) of the kinase domain [18, 19]. However, a sizeable proportion of BvgS homologs are devoid of a PAS domain and flanking linkers. Instead, a two-helix linker called the linker X directly connects the transmembrane segment to the DHp domain [20]. The two-helix linkers form coiled coils that regulate the BvgS enzymatic activity [18, 19]. We have built several chimeras by replacing the region between the TM and DHp domains of BvgS with those from homologs devoid of PAS domain [18, 20]. Among the various regulation phenotypes of those chimeras, that of the so-called BvgSΔ65 is inverted relative to that of BvgS [20]. Thus, BvgSΔ65 shows low kinase activity at the basal state, most likely corresponding to the Bvg⁺ phase, while chemical modulation or closure of the VFT1 domains triggers a large increase of kinase activity, shifting the bacteria to the Bvg⁻ phase [20]. In this study, we characterized this and other BvgS chimeras using transcriptomic analyses and a mouse model of infection. We obtained no evidence for a shift of BvgSΔ65 to the kinase mode of activity in the mouse during the time.
course of infection. In addition, we identified different ranges of virulence factor expression for colonization of, and persistence in, distinct sites of the mouse respiratory tract.

Materials and methods

Strains and plasmids

The strains used to perform animal experiments, BPSM, BPSM$_{SS1}$ (previously called BPSM$_{BvgS-E113C+N177C}$), BP-BvgS$_{Δ65}$ and BP-BvgS$_{Δ65-SS1}$ (previously called BvgS$_{Δ65-VFT1-SS}$) were described previously [12, 20]. In all of them, expression of the $bvgS$ variants from the natural chromosomal locus was achieved by allelic exchange as described in [12]. Similarly, to construct BP-BvgS$_{T733M}$ the mutation of interest was introduced by mutagenesis on a pUC19 derivative containing the appropriate region of the $bvgS$ gene, followed by cassette exchange in pUC19mint, transfer of the EcorI-HindIII fragment into pSORTP1 and homologous recombination into BvgS$_{new}$ as in [12].

The BvgS$_{Δ65}$ R572L variant was expressed from a plasmid for activity measurement. It was constructed by mutagenesis on a pUC19 derivative containing the region of interest, followed by cassette exchange in pUC19mpla and then in pBBRmpla [12]. The recombinant $B$. pertussis strains were obtained by introducing the pBBRmpla variant by conjugation in $B$. pertussis BPSM$_{new}$ carrying the chromosomal ptx-lacZ transcriptional fusion [12]. $bvgS_{A65}$ was cloned on a plasmid in a similar manner for comparison.

Animal experiments

After 36 h of growth on standard Bordet-Gengou (BG)-blood plates or plates containing 50 mM MgSO$_4$ to modulate the various strains, bacteria were resuspended in sterile PBS to $10^6$ bacteria per 20 $\mu$L. After intraperitoneal anesthesia with a mixture of ketamine, atropine and valium, female 6-weeks-old JAX BALB/cByJ mice from Charles River were infected by intranasal inoculation with $10^6$ bacteria. Groups of 5 animals per bacterial strain were sacrificed by cervical dislocation after 3 h and 3, 7, 14 or 28 days post-inoculation in the first experiment, and groups of 3 animals after 3 h and 7, 14 or 21 days in the second. Noses and lungs were collected and homogenized using an Ultra Turrax apparatus. Serial dilutions were performed in PBS and plated on BG agar plates to count the bacteria. All the experiments were carried out in accordance with the guidelines of the French Ministry of Research regarding animal experiments, and the protocols were approved by the Ethical Committees of the Region Nord Pas de Calais and the Ministry of Research (agreement number APAFIS#9107–201603311654342 V3).

RNA extractions

$B$. pertussis strains were grown on BG agar plates for 2 days at 37°C and then cultured in modified Stainer Scholte (SS) liquid medium supplemented when indicated with 50 mM MgSO$_4$ at 37°C under agitation. The bacterial cultures were stopped at mid exponential phase ($OD_{600} = 2$) by adding 1 mL of a mixture of 5:95 phenol:ethanol (v:v) to 4 mL of bacterial suspensions. Bacteria were pelleted, and total RNA was extracted using TriReagent (Invitrogen) following the manufacturer’s instructions. Genomic DNA was removed by DNase I treatment (Sigma Aldrich).

Illumina RNA sequencing

RNA-seq experiments were performed with two independent cultures of BPSM and BPSM$_{SS1}$, BP-BvgS$_{A65}$,
BP-BvgS\textsubscript{A65-SS1} and BP-BvgS\textsubscript{T733M}. For each RNA-seq sample, DNA-depleted total RNA was treated with the Ribo-Zero rRNA Removal Kit (Illumina) following the manufacturer’s recommendations. The RNA-depleted RNA was then used to make the Illumina library using the TruSeq RNA Library Preparation Kit, following sequencing on an Illumina NextSeq 500 benchtop sequencer on SR150 high output run mode. The RNA-seq data of each sample were analyzed using Rockhopper v2.0.3 with the default parameters to calculate the RPKM value for each CDS using the *Bordetella pertussis* Tohama I BX470248 genome annotation [21]. The RNA-seq data reported in this paper have been deposited in the Sequence Read Archive, [www.ncbi.nlm.nih.gov/sra](http://www.ncbi.nlm.nih.gov/sra) (submission SUB4097406; NCBI BioProj PRJNA474836; BioSample accessions: SAMN09374659, SAMN09374660, SAMN09374661, SAMN09374662, SAMN09374663, SAMN09374664, SAMN09374665, SAMN09374666).

**Generation of cDNA and quantitative real-time polymerase chain reaction (qRT-PCR)**

Fifteen micrograms of total RNA were treated with DNase I, and then 500 ng of total RNA was reverse-transcribed using the Verso cDNA synthesis kit (Thermo Scientific). Polymerase chain reaction (PCR) was performed on 30 ng of cDNA using a LightCycler 480 SYBR Green I Master kit (Roche) and a Roche LightCycler 480 Instrument II. The efficiency for each primer pair was determined by serial dilutions. The experiments were performed three times for BPSM, MgSO\textsubscript{4}-treated BPSM, BP-BvgS\textsubscript{A65}, MgSO\textsubscript{4}-treated BP-BvgS\textsubscript{A65}, BPSM\textsubscript{SS1}, BPSM\textsubscript{T733M} and BP-BvgS\textsubscript{A65}.rev 79, twice for BP-BvgS\textsubscript{A65}.rev 80 and from single biological sampling for the other BP-BvgS\textsubscript{A65}.rev strains. For each sample, at least three technical replicates were performed. The results were analyzed with the Light Cycler 480 software. The expression of the housekeeping gene *bp3416* was used as reference to normalize the expression of the genes of interest.

**Extraction of genomic DNA and sequence determination**

The various segments of the *bvgS* gene were amplified from clarified lysates obtained by heating bacteria resuspended in H\textsubscript{2}O at 95°C for 30 min, using the pairs of primers described in [22], and the amplicons were sequenced by the Sanger method. DNAseq experiments were performed on genomic DNA extracted using the Genomic tip 100/G kit (Qiagen). The sequencing libraries were prepared with the Nextera XT sample prep kit (Illumina) following the manufacturer’s instruction and sequenced on HiSeq 2500 with 2x250 bp reads. The read mapping was performed with CLC Genomics Workbench V11, and variants were searched using the Basic Variant Detection module.

**β-galactosidase assay**

β-galactosidase assays were performed as described previously [12] with 3 different clones at different times, and the means and standard errors of the means were determined. TCEP was added to the cultures for 6 or 16 hours prior to harvesting the bacteria.

**Statistical analyses**

Statistical analyses were performed using an unpaired t test for β-galactosidase results, and using the one-way ANOVA followed by Bonferroni’s multiple comparison post-test for the animal experiments and qRT-PCR results. For the RNA-seq data, the adjusted p-values were determined using the Benjamini-Hochberg procedure in Rockhopper.
Results

Enhancement of animal colonization by reversion of the BvgSΔ65 regulation phenotype

The BP-BvgSΔ65 recombinant strain shows an inverted regulation phenotype relative to its wild type (wt) parent BPSM in laboratory conditions [20], as chemical modulation shifts BvgSΔ65 to a high-kinase mode of activity (Fig 1A and 1B). We characterized the full gene expression profile of BP-BvgSΔ65 in the ‘default state’ (i.e., not modulated) by performing Illumina RNA sequencing (RNA-seq) experiments and compared it with that of BPSM modulated or not (Fig 1C; S1 Table). The transcription profile of BPSM was in good agreement with previous reports [23–25], with high transcription levels of the vags and low transcription levels of the vrgs (Fig 1C; green and red ovals; S1 Table). Except for a few sets of genes, the gene expression profile of BP-BvgSΔ65 was rather close to that of modulated BPSM. Thus, expression of late vags, e.g. coding for PTX or the type III secretion system, decreased to the same levels as in modulated BPSM (Fig 1C, green oval; S1 Table). In contrast and congruent with the BvgΔ65 phase, the expression levels of early vags such as fhaB, fimBCD or fhuA in BP-BvgSΔ65, were similar to those in BPSM (Fig 1A, blue oval), and bipA (bp1112), the hallmark gene of the BvgΔ phase in B. pertussis, was expressed at higher level in BP-BvgSΔ65 than in BPSM grown in standard or modulated conditions (Fig 1C, purple circle) [23]. Additionally, some other vags, like fhuA (bp2667), fntB (bp2936), prn (bp1054), and other genes including lgmABC (bp0397, bp0398 and bp0399) and putA (bp2749) were expressed at higher levels in BP-BvgSΔ65 than in modulated BPSM (S1 Table). Most of the vrgs were expressed at levels similar to those in modulated BPSM, which was not previously reported for the BvgΔ phase (Fig 1A, red ovals) [23]. Altogether, these results indicate that the default state of BP-BvgSΔ65 is the BvgΔ phase.

As in vitro modulation shifts BP-BvgSΔ65 to the BvgΔ phase, we investigated whether animal experiments might reveal specific conditions present in the mammalian respiratory tract that induce phenotypic modulation, thus causing a shift of BP-BvgSΔ65 to the colonization-proficient, BvgΔ phase. If this occurred, the ‘up-modulated’ recombinant bacteria should be able to colonize the animal lungs, while they would be cleared quickly if they remained in the BvgΔ phase, as described earlier using other BvgΔ-phase strains [26, 27]. In contrast, we did not expect them to be rapidly cleared from the mouse noses, as B. pertussis locked in the BvgΔ phase was reported to survive in the mouse upper respiratory tract in a way similar to virulent bacteria [26, 27]. Mouse colonization experiments were thus performed with BP-BvgSΔ65 cultured in standard conditions (i.e., 37°C without modulator) prior to inoculation, or at 37°C in the presence of 50 mM MgSO4 to set the bacteria in the BvgΔ phase prior to colonization. We reasoned that if the bacteria encountered modulating conditions at a late stage of mouse colonization rather than early on, pre-modulation of BP-BvgSΔ65 might facilitate initial survival in the animals, before induction of virulence factor expression resulting from in vivo modulation. If no modulation occurred in vivo, we expected pre-modulated BP-BvgSΔ65 to progressively lose virulence factor expression and therefore to be cleared like its non-modulated counterpart. The wt control strain, BPSM, was cultured in the same two conditions before mouse inoculation. The mice were infected intranasally with 10⁶ bacteria. After 3 h, approximately 10⁶ and 10⁵ bacteria were found in the lungs and in the noses of mice infected with the BPSM control or with BP-BvgSΔ65 (Fig 1D and 1E). The colonization profiles by BPSM were as typically reported. At day 3, the bacterial loads increased in both organs, the bacteria were found in similar numbers at day 7, and then their numbers markedly decreased at days 14 and 28. Bacterial multiplication and clearance from both organs appeared to be slightly delayed for BPSM chemically down-modulated prior to inoculation, possibly because virulence factor
Fig 1. Characterization of BP-BvgS_{Δ65} and its colonization of the mouse respiratory tract by comparison with control strain BP-PSM. A. Schematic representation of wild type BvgS as found in the control strain BP-PSM and of the BvgS_{Δ65} variant.
production was initiated only when the bacteria encountered the \textit{in vivo} environment of the animals’ respiratory tract and shifted to the Bvg$^+$ phase. For BP-BvgS$\Delta 65$, in contrast, no bacterial multiplication was observed at day 3 in either organ. On the contrary, marked decreases of the bacterial loads were seen irrespective of prior up-modulation of the bacteria. Thus, \textit{in vitro} modulation before inoculation failed to restore a wt-like colonization profile. The fast decreases of the BP-BvgS$\Delta 65$ populations in both organs do not support the hypothesis that up-modulating signals were present in the respiratory tract that might have shifted BvgS$\Delta 65$ to a high-kinase mode of activity.

At later time points, however, BP-BvgS$\Delta 65$ appeared to survive at similar levels to BPSM. BP-BvgS$\Delta 65$ is non-hemolytic on blood agar plates in non-modulated culture conditions, due to the lack of or the very low-level expression of the hemolysin/adenylate cyclase gene \textit{cyaA}, a late \textit{vag} whose transcription requires high concentrations of phosphorylated BvgA [8]. Intriguingly, however, all colonies recovered at day 3 from the lungs of mice infected with non-modulated BP-BvgS$\Delta 65$, and 40% of those recovered from the lungs of mice infected with BP-BvgS$\Delta 65$ modulated prior to infection were hemolytic (Fig 2A). At day 7, only hemolytic colonies were recovered from the lungs of the two groups of mice. Hemolytic colonies were also recovered from the noses, in low proportions at day 3 that appeared to increase at day 7 (Fig 2B). Nevertheless, non-hemolytic clones were detected in the noses up to day 28. The hemolytic phenotype was maintained after re-streaking those clones on Bordet Gengou agar-blood plates with or without modulators, suggesting that the bacteria were locked in the Bvg$^+$ phase.

The hemolytic bacteria appear to have undergone a reversion from the Bvg$^+$ phase to the Bvg$^-$ phase in most of the animals. Possibly because of this reversion, we observed a moderate increase of the bacterial loads of BP-BvgS$\Delta 65$ in the lungs between days 3 and 7, for both the bacteria chemically modulated and those not modulated prior to inoculation (Fig 1D and 1E). Nevertheless, the bacterial loads remained 1000-fold lower than those of BPSM even at their peak at day 7. The numbers of bacteria in the lungs of those two groups of mice leveled off at day 14 and decreased thereafter. In the noses, where the proportions of hemolytic colonies were initially lower, the bacterial counts remained steady between days 3 and 7 and started to decrease from day 14. Thus, infection of mice with BP-BvgS$\Delta 65$ resulted in the appearance of hemolytic variants whose proportions in the bacterial populations recovered from both organs increased over time. This suggested that it could be due to selective pressure in mice. As only 0.1% of the initial bacterial loads were recovered from the lungs after 3 days, the remaining
99.9% bacteria, which were presumably non-revertants, most likely died in the first three days. The potential selective pressure for virulent bacteria—using the hemolytic phenotype as a proxy for virulence—appears to be less intense in the nose than in the lungs, as more than 40% of the BP-BvgSΔ65 bacteria recovered from the noses were non-hemolytic at day 14 (Fig 2B).

### Colonization profiles by distinct Bvg1-phase bacteria

Since hemolytic revertants were obtained from several animals, the most likely explanation is that they spontaneously arose at low frequency in our master stock of BP-BvgSΔ65 and were selected for by the host environment. However, when BP-BvgSΔ65 was streaked on blood-agar plates and incubated for 5 days at 37°C, no hemolytic colony was detected, indicating that hemolytic bacteria, if present, would represent a very small proportion of the stock. Furthermore, deep Illumina whole-genome sequencing of BP-BvgSΔ65 did not reveal any mutations.

We nevertheless performed a new round of single colony isolation starting from our initial BP-BvgSΔ65 stock, which yielded BP-BvgSΔ65new, and we performed another mouse colonization experiment with the newly isolated clone. In this second experiment, in addition to BPSM and BP-BvgSΔ65new, we included two other strains that express BvgS variants with different levels of enzymatic activities, BPSMSS1 [12] and BP-BvgSΔ65SS1 [20] (Fig 3A and 3B). In BPSMSS1, two selected residues at the lips of the lobes of the VFT1 domain of BvgS were replaced with Cys residues, which results in the formation of an inter-lobe disulfide (S-S) bond that...
Fig 3. Characterization of strains used in this study and colonization of the lungs and noses of mice. A. Schematic representation of the BvgS variants present in BPSMSS1 and BPSM-BvgSΔ65-SS1 (see legend of Fig 1). Disulfide bond formation between the two lobes of the VFT1 domain is indicated with 'S-S'. B. Activities of the variants as measured using the ptx-lacZ or fhaB-lacZ reporter systems. The strains were grown in standard conditions (0), and where indicated 10 mM Tris(2-carboxyethyl)phosphine hydrochloride (TCEP 10) was added to reduce the S-S bond in VFT1. The measurements were performed at least three times, and the means and standard errors of the mean are given. Statistical analyses were performed, and significant p values are indicated (***, p < 0.0001, ***, p < 0.001). C and D. Numbers of colony-forming units (CFUs) recovered at the indicated time points from the lungs (C) or the noses (D) of mice infected with BPSM or BP-BvgSΔ65. Prior to inoculation, the bacteria were grown in standard conditions. Three mice were sacrificed at each time point for each bacterial strain. The means and standard errors of the means are shown. Statistical analyses were performed for each data point using the corresponding BPSM data point as a control. Significant p values are indicated (***, p < 0.001; *, p < 0.05). The other data present p-values >0.05.

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we observed in this study that BvgS reporter [12]. However, using the ptx-lacZ closure abolished BvgS kinase activity in both standard and modulated conditions using the ptx-lacZ reporter [12]. In non-modulating conditions, BvgS appears to be in a 'low-kinase' (Bvg<sup>↓</sup>) mode of activity, rather than in the bona fide phosphatase mode. Conversely, introduction of the same S-S bond in Bvg<i>S</i><sub>Δ65</sub>, yielding Bvg<i>S</i><sub>Δ65-SS1</sub>, shifted it to a high-kinase mode of activity, as shown with the ptx-lacZ reporter in standard conditions [20] (Fig 3B). Upon addition of TCEP to the culture, the β-gal activity decreased significantly but not fully, suggesting that S-S bond reduction was most likely incomplete in those conditions.

In the second animal experiment, two Bvg<i>↓</i>-phase strains, BPSM and BP-Bvg<i>S</i><sub>Δ65-SS1</sub>, and two Bvg<i>↑</i>-phase strains, BP-Bvg<i>S</i><sub>Δ65</sub> and BPSM<sub>Δ</sub>65<i>SS1</i>, were thus used to inoculate mice. The bacteria were counted in their lungs and noses over the time course of infection. The colonization profiles of both organs by BP-Bvg<i>S</i><sub>Δ65-SS1</sub> were similar to those of BPSM (Fig 3C and 3D). This indicates that formation of the S-S bond in the VFT1 domains of Bvg<i>S</i><sub>Δ65 SS1</sub> enabled the bacteria to express the virulence factors necessary for colonization and survival in both organs. In contrast, the two Bvg<i>↑</i>-phase strains, BP-Bvg<i>S</i><sub>Δ65new</sub> and BPSM<sub>Δ</sub>65<i>SS1</i>, were rapidly cleared from the lungs but colonized the noses of mice.

In contrast to the first animal experiment, no hemolytic colonies of BP-Bvg<i>S</i><sub>Δ65new</sub> were obtained from either organ at any time point. This supported the hypothesis that the initial BP-Bvg<i>S</i><sub>Δ65</sub> bacterial stock used for the first experiment contained a very small proportion of revertants that were not detected on blood-agar plates prior to inoculation of the mice. In the lungs, BP-Bvg<i>S</i><sub>Δ65new</sub> was rapidly eliminated and was not detectable after day 7. In the noses, this strain did not multiply, and the bacterial load markedly decreased at day 7, followed by a milder decrease thereafter.

In the lungs, BPSM<sub>Δ</sub>65<i>SS1</i> did not multiply and was rapidly cleared, though not quite as quickly as BP-Bvg<i>S</i><sub>Δ65new</sub>. In contrast, BPSM<sub>Δ</sub>65<i>SS1</i> persisted in the noses in a manner similar to BPSM, or even better at day 21. Thus, whereas the recombinant strains that express Bvg<i>S</i> variants displaying intermediate modes of activity were rapidly cleared from the lungs, they survived much longer in the noses. Intriguingly, BPSM<sub>Δ</sub>65<i>SS1</i> persisted in the noses at higher bacterial counts than BP-Bvg<i>S</i><sub>Δ65new</sub>. The observation that the nose colonization profiles of two strains in the Bvg<i>↑</i> phase were not identical indicates that they might display slightly different gene expression patterns.

Expression range of the various recombinant strains. We performed RNA-seq experiments on BPSM<sub>Δ</sub>SS1 and BP-Bvg<i>S</i><sub>Δ65-SS1</sub> to characterize their gene expression profiles (Fig 4A and 4C). An additional strain, BPSM<sub>T733M</sub>, which represents a prototypical Bvg<i>↓</i>-phase-locked strain, was included in the experiments (Fig 4B). The T<sup>733M</sup> mutation, which localizes to the DHp domain of Bvg<i>S</i>, spontaneously occurred in Bordetella bronchiseptica and led to the initial description of the Bvg<i>↑</i> phase [9]. This mutation was introduced in the BPSM chromosome by allelic exchange to serve as the Bvg<i>↑</i> reference strain for RNA-seq.

The gene expression profiles of BPSM<sub>Δ</sub>SS1 and BPSM<sub>T733M</sub> were broadly similar to each other and rather close to that of BP-Bvg<i>S</i><sub>Δ65</sub> (compare Fig 4A and 4B with Fig 1C; S1 Table). Subtle differences were nevertheless found among the Bvg<i>↑</i>-phase strains, notably regarding the levels of expression of specific vags (Fig 1C and Fig 4A–4C, blue ovals; S1 Table). Furthermore, bip<i>A</i> (<i>bp1112</i>), a specific marker of the Bvg<i>↑</i> phase [23], was overexpressed in both BP-Bvg<i>S</i><sub>Δ65</sub> and BPSM<sub>T733M</sub> (purple circles), but not to the same extent in BPSM<sub>Δ</sub>SS1. Altogether, BP-Bvg<i>S</i><sub>Δ65</sub>, BPSM<sub>T733M</sub> and BPSM<sub>Δ</sub>SS1 present transcriptomic patterns characteristic of
the Bvg^ phase, but within a certain range of gene expression levels. The reason why BPSM_SS1 was more efficient at colonizing the mice noses than BP-BvgS_Δ65 is difficult to ascribe to specific differences between their transcriptomes.

BP-BvgS_Δ65-SS1 expressed several vags, including ptx, vag8 (bp2315), tcfa (bp1201), brkA (bp3494) and bfrD (bp856), at significantly lower levels than BPSM, but higher than the three Bvg^_phase strains (S1 Table). The expression of bipA in BP-BvgS_Δ65-SS1 was higher than in BPSM and in the same range as in the Bvg^_phase strains. BP-BvgS_Δ65-SS1 also expressed the genes of the chemotaxis and flagellar operons at higher levels than modulated BPSM (Fig 4C, grey circle). BP-BvgS_Δ65-SS1 is thus closer to the Bvg^+ phase than the other variants but with lower expression levels of several vags than BPSM, although it colonized mice as efficiently as BPSM. Genetic engineering of BvgS can thus generate a range of intermediate phenotypes.

We also performed quantitative real-time polymerase chain reaction (qRT-PCR) experiments to quantify the transcripts of selected genes in the various strains (Fig 5). We included late vags (cya, ptx-S1, tcfa), an early vag (bp1881, i.e. fimB), the intermediate phase marker bipA, and the vrg bp2782. We tested the effect of modulation on gene expression in BP-BvgS_Δ65.

The qRT-PCR data showed that the strains expressing BvgS_Δ65, BvgS_T733M and BvgS_SS1 have similar patterns of expression that represent the Bvg^ phase. This analysis also confirmed that BP-BvgS_Δ65 treated with MgSO_4 and BP-BvgS_Δ65 SS1 are closer to the Bvg^ phase than the three Bvg^_phase strains, although they expressed some vags at lower levels than BPSM. Thus, the BvgS variants analyzed in this work display a range of gene expression patterns between the fully Bvg^+ and Bvg^ phases, which most likely correlate with their respective behaviors in animal colonization.

**Identification of a mutation in hemolytic colonies of BP-BvgS_Δ65**

Finally, we characterized hemolytic colonies of BP-BvgS_Δ65 that appeared in the first animal experiment to test the hypothesis that their bvgS gene harbored a mutation that locks it in the kinase mode. Such ‘constitutive’ mutants were previously reported to spontaneously occur in wt BvgS, with the substitutions mapping in particular in the PAS domain and the linker 1 [28, 29]. We thus PCR-amplified and sequenced the corresponding region of bvgS_Δ65 from a number of hemolytic clones obtained at various time points from the two sites of the mice’s
respiratory tracts. A mutation replacing the CGC codon of the Arg572 residue with a Leu CTC codon was identified in a majority of the clones recovered from the lungs, and in some of those recovered from the noses (Fig 6A). Arg572 is localized in linker X of BvgSΔ65.

To confirm the effect of this mutation on the activity of BvgSΔ65, the R572L substitution was introduced in that variant by site-directed mutagenesis. The recombinant BvgSΔ65-R572L-expressing strain was hemolytic on blood agar plates irrespective of the addition of 50 mM MgSO4 as the modulating agent. It also showed a high level of β-galactosidase (β-gal) activity using the ptx-lacZ reporter fusion that did not respond to modulation (Fig 6B), unlike BP-BvgSΔ65. This confirms that the R572L mutation is sufficient to lock the BvgSΔ65 variant in a kinase state unresponsive to modulation.

Intriguingly, other hemolytic variants selected for sequencing did not harbor the R572L or any other substitution in the linker X region of BvgSΔ65, and whole-genome sequencing on seven such clones did not identify mutations to account for their hemolytic phenotype. We performed qRT-PCR analyses on a hemolytic variant harboring the R572L substitution in BvgSΔ65 (BP-BvgSΔ65-rev79) and on several other revertants devoid of that substitution. The transcription patterns of the revertant strains showed rather similar trends (Fig 6C). They expressed cyaA at levels similar to, or moderately lower (in BP-BvgSΔ65-rev80) than BPSM, in agreement with their hemolytic phenotypes on blood-agar plates. Similarly, the expression levels of ptx or fimB were similar to or slightly below those of BPSM, whereas those of another vag, tcfa, were lower. The revertants transcribed the Bvg1-phase gene bipA at levels similar to or higher than BPSM. The expression levels of the selected vrg, bp2782, were significantly lower in several of them, but not in BP-BvgSΔ65-rev79 harboring the R572L mutation, than in BPSM.

Discussion
We genetically engineered a chimera, BvgSΔ65, in which the linker X of a BvgS homolog replaces the region encompassing the linker 1, the PAS domain and the linker 2 of BvgS, between the transmembrane and the DHp domains [20]. In standard growth conditions,
recombinant BP-BvgSΔ65 is in the Bvg\(^{-}\) phase, and modulation causes an increase of BvgSΔ65 kinase activity, contrary to its effect on wt BvgS. We thus used the inverted regulation properties of this chimera as a tool to investigate host-pathogen interactions in an animal model of infection. Additional strains harboring other engineered versions of BvgS were also included in mouse colonization experiments. Two distinct groups of strains were thus revealed. BP-BvgSΔ65 and BPSMSS1, which are in the Bvg\(^{-}\) phase, were able to colonize and to survive in the mouse noses but were rapidly cleared from the lungs, while BP-BvgSΔ65 SS1 could colonize and survive in those two compartments, similar to the Bvg\(^{+}\)-phase control strain BPSM. There thus appears to be distinct virulence thresholds for the colonization of the mouse nose, which both Bvg\(^{-}\)- and Bvg\(^{+}\)-phase bacteria can colonize, and the mouse lungs, which only Bvg\(^{+}\)-phase bacteria can. Our transcriptomic analyses showed that those Bvg\(^{-}\)- and Bvg\(^{+}\)-phase strains populate distinct ranges of phenotypes, based on gene expression levels in the various strains. Such recombinant strains are useful tools to finely dissect the interactions of \textit{Bordetella pertussis} with its host.

Fig 6. Characterization of hemolytic variants of BP-BvgSΔ65. A. Numbers of hemolytic colonies selected for sequencing that present the R\(^{572}\)L mutation. The variants were isolated from the lungs (left panel) and noses (right panel) of the mice. The cause of the hemolytic phenotype of the clones that do not carry the above mutation is not identified (ni). B. The \textit{ptx-lacZ} reporter system was used to determine the activities of the BvgSΔ65 R572L compared to BvgSΔ65 in standard conditions (0) or after growth in the presence of 2 mM chloronicotinate (CN 2). The measurements were performed at least three times, and the means and standard errors of the mean are given. Statistically different values relative to BPSM are indicated by \(* * * *\), \(p<0.0001\); ns, \(p>0.05\). C. Quantitative RT-PCR analysis of selected genes in various recombinant strains. The values represent the Log2 values of the mean expression of each gene normalized to \textit{bp3416} using the 2ΔCt method. The error bars represent the Ct standard deviations. BP-BvgSΔ65-rev79 (obtained from a mouse nose at day 14) is a hemolytic clone with BvgSΔ65 carrying the R\(^{572}\)L substitution. BP-BvgSΔ65-rev26 (from nose; day 3), -rev47 (from lung; day 3), -rev54 (from nose; day 7) and -rev80 (from nose; day 14) are other hemolytic revertants without that mutation. The results represent biological triplicates for BP-BvgSΔ65-rev79, biological duplicates for BP-BvgSΔ65-rev80, and single biological sampling for the other strains. In all cases, the measurements were performed in triplicates. Statistical analyses were performed using gene expression in BPSM as a control and indicated as follows: \(* * * *\), \(p<0.0001\); \(* *\), \(p<0.01\); \(*\), \(p<0.05\).

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Our initial goal in this study was to determine whether modulation might occur in the host. Closing of the inter-lobe cavity upon ligand binding triggers signaling in other VFT domain-based systems [30, 31]. Several observations have suggested that the VFT1 domains of BvgS might function similarly. Thus, they are open in the available crystal structure, their putative solute-binding cavity is conserved among _B. pertussis_ isolates, and their closing by inter-lobe S-S bond formation down-regulates the kinase activity of wt BvgS [12]. Therefore, we took advantage of the inverted regulation phenotype of BP-BvgSΔ65 to test the hypothesis that VFT1 might bind modulating ligands in the course of infection by _B. pertussis_. As S-S-bond-induced closing of the VFT1 domains of BvgSΔ65 restored kinase activity and a wt-like colonization profile by BP-BvgSΔ65 SS1 in a mouse model of infection (this work), ligand-induced closing of the VFT1 domains of BvgSΔ65 in the respiratory tract of mice might similarly have enhanced colonization by BP-BvgSΔ65. Our animal experiments, however, provided no indication that BP-BvgSΔ65 could colonize mouse lungs in the absence of reversion, arguing that modulation of BvgS activity did not occur in that organ. One caveat to our conclusion is that temporally or spatially restricted _in vivo_ modulation might be missed in animal experiments. Of note, high-level expression of several _vrgs_ in mouse infection were recently reported [32]. Using a fluorescent reporter under the control of the _ptx_ promoter in BP-BvgSΔ65, we also found no evidence that modulation might occur inside macrophages or dendritic cells (our unpublished data), in line with recent proteomic analyses of _B. pertussis_ that showed increased production of some virulence proteins inside human macrophages [33]. Nevertheless, as both Bvg⁺ and Bvg⁻ phase strains can colonize the nose, we cannot exclude the possibility of up-modulation of BvgSΔ65 in the upper respiratory tract of mice. It is also possible that modulating signals are present in specific environments or conditions encountered by the bacteria in the human respiratory tract, but not in that of the mouse.

In the first animal experiment, colonies with hemolytic phenotypes progressively outnumbered non-hemolytic colonies. Transcriptomic analyses of those variants showed increased expression levels of several virulence factors relative to parental BP-BvgSΔ65. The rapid increase of the proportions of those more virulent revertants and their ability to outcompete non-hemolytic clones suggest that they were selected for by innate immune responses in the mice, and that this selective pressure is very strong in the lungs. In the nose, in contrast, non-hemolytic bacteria in the Bvg⁺ phase could survive for up to four weeks, similarly to those in the Bvg⁻ phase. Nevertheless, the occurrence of both BP-BvgSΔ65 and hemolytic revertants in the noses of mice from day 3 _de facto_ resulted in a mixed infection, and the initially low proportions of revertants in the noses increased relative to their Bvg⁺-phase parent over time. This shows that they may have a selective advantage in the upper respiratory tract as well, in agreement with previous reports that the prototypical Bvg⁺-phase-locked strain harboring the T₇₃₃M substitution, was less competitive in both compartments [26]. Nonetheless, as selective pressure is less intense in the nose, it is conceivable that attenuated mutants could persist in that compartment in the absence of more virulent competitors. The report that avirulent _B. pertussis_ clones harboring an IS481 insertion in _bvgAS_ were detected in the nasopharynx of experimentally infected monkeys suggests that even avirulent bacteria might be able to persist for some time in the upper respiratory tract once infection has been established [34].

The transcription profile of BP-BvgSΔ65 SS1 shows that it is not a fully virulent Bvg⁺-phase strain, but an additional intermediate between the _bona fide_ Bvg⁺ and Bvg⁻ phenotypes. Nevertheless, expression of _vrgs_ in that strain appears to be sufficient to establish lung infection, indicating that the virulence phenotype of BP-BvgSΔ65 SS1 is beyond the threshold at which mouse lung colonization can occur. Whether such a moderately virulent strain might be successful in natural human infections is unknown, and it might be interesting to address this question in a baboon model of infection [1]. Its phenotype might be advantageous in the face
of the immune pressure, notably linked to vaccination. The loss of specific virulence factors or lower levels of virulence gene expression have indeed been reported to occur in currently circulating \( B. \) pertussis strains [35–39]. On the contrary, however, immune pressure appears to currently select for strains that express greater levels of pertussis toxin [40]. Thus, distinct virulence factors make different contributions to the success of infection. In this respect, it is worth noting that a live attenuated vaccine candidate, BPZE1, which lacks some \textit{bona fide} virulence factors, efficiently colonizes the respiratory tract of animals and humans without causing disease [41, 42].

\( B. \) pertussis has conserved its capability to shift between the \( \text{Bvg}^{-} \) phase and the \( \text{Bvg}^{+} \) or \( \text{Bvg}^{i} \) phases in the laboratory. However, the relevance of the latter phases in the host and the conditions that trigger phenotypic modulation in its current lifestyle remain unclear. For \( B. \) pertussis, the shift to the \( \text{Bvg}^{i} \) phase might be a remnant of an ancestor able to survive outside mammalian hosts, as the \( \text{Bvg}^{i} \) phase of \textit{Bordetella bronchiseptica} allows its growth and dissemination in amoeba as an environmental reservoir [43]. The occurrence and the role of the \( \text{Bvg}^{i} \) phase in the infection remain unclear as well. The presence of antibodies directed against \( \text{Bvg}^{i} \)-phase-specific antigens in the sera of convalescent children was reported [44] but contradicted by subsequent work [26]. An \textit{in vivo} shift to the \( \text{Bvg}^{i} \) phase in the nasopharynx might be favored by the somewhat lower temperature there than in the rest of the respiratory tract, but a recent study revealed that virulence factor expression in \( B. \) pertussis persists at suboptimal temperatures [45]. Several studies including this one have shown that \( \text{Bvg}^{i} \)-phase locked \( B. \) pertussis strains are proficient for infection of the mouse nose. However, \( \text{Bvg}^{-} \)-phase locked \( B. \) bronchiseptica remains proficient for infection and transmission in a pig model of disease, casting doubt on the idea that the \( \text{Bvg}^{i} \) phase contributes to transmission [27]. It is possible that the bacteria transiently shift to the \( \text{Bvg}^{i} \) phase before or after transmission, but the \( \text{Bvg}^{i} \) phase passes undetected by the immune system.

It was proposed that virulence genes expressed by \( B. \) pertussis could be classified in four classes, and that \textit{vrgs} were expressed mainly in absence of phosphorylated \( \text{BvgA} \), that is, in the \( \text{Bvg}^{i} \) phase [4]. Nevertheless, we showed in this study that expression of several \textit{vrgs} was at similarly high levels in the \( \text{Bvg}^{i} \)-phase strains as in the modulated, \( \text{Bvg}^{-} \)-phase wt strain. In contrast, \textit{vrgs} were not overexpressed in the \( \text{Bvg}^{i} \)-phase-locked strain described earlier [23]. Whether these differences stem from the different genetic backgrounds of the strains used in the two studies remains to be determined. It is also possible that the \textit{vrgs} form distinct subgroups that are regulated in slightly different ways, as shown for \textit{vags} [24].

The majority of hemolytic revertants of \( BPSM-\text{BvgS}_{\text{Δ65}} \) isolated from the mice had a specific mutation in linker \( X \) that results in the substitution of an Arg residue with a Leu. In the subset of \( \text{BvgS} \) homologs devoid of a PAS domain, the linker \( X \) harbors two antagonistic coiled-coil registers, and regulation of activity implies interconversion between the two marginally stable coiled coils defined by those registers [20]. In \( \text{BvgS}_{\text{Δ65}} \), Arg572 appears to be involved in repulsive interactions between the two helices in the kinase register of the coiled coil, but not in the phosphatase register [20]. Therefore, the removal of unfavorable interactions by the Arg to Leu replacement is likely to stabilize the two-helix coiled coil in the kinase register, which might lock \( \text{BvgS}_{\text{Δ65}} \) in the kinase state of activity.

Finally, some of the hemolytic revertants from \( BPSM-\text{BvgS}_{\text{Δ65}} \) isolated from the noses of mice at early time points, did not harbor the \( R^{572} \) point mutation to account for their hemolytic phenotype. While whole-genome sequencing did not enable us to identify any other specific mutation, restoration of a hemolytic phenotype may have arisen from genomic reshuffling of insertion sequences or epigenetic modifications. The observation that the hemolytic phenotype appears to be maintained upon sub-culturing suggests that the \( \text{BvgS}_{\text{Δ65}} \) variant might generate bistability, in particular in the mouse nose. Of note, bistable phenotypes have
been reported in the lower respiratory tract of mice infected with specific mutants of the broad-host range pathogen *Bordetella bronchiseptica* [46].

**Supporting information**

**S1 Table. Raw data of the RNA seq experiments.** The RPKM for each open reading frame are provided for the 6 conditions. The corrected p-values and the fold changes (FC) are also provided. Fold changes > 2 or >-2 (in log2) are considered to be significant. (XLSX)

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**References**

1. Pinto MV, Merkel TJ. Pertussis disease and transmission and host responses: insights from the baboon model of pertussis. J Infect. 2017;74 Suppl 1:S114–S9. https://doi.org/10.1016/S0163-4453(17)30201-3 PMID: 28646950.

2. Locht C, Antoine R, Jacob-Dubuisson F. *Bordetella pertussis*, molecular pathogenesis under multiple aspects. Curr Opin Microbiol. 2001; 4(1):82–9. PMID: 11173039

3. Stibitz S. The *bvg* regulon. In: Locht C, editor. Bordetella Molecular Biology Norfolk, U.K.: Horizon Bioscience; 2007. p. 47–68.
4. Jones AM, Boucher PE, Williams CL, Stibitz S, Cotter PA. Role of BvgA phosphorylation and DNA binding affinity in control of Bvg-mediated phenotypic phase transition in *Bordetella pertussis*. Mol Microbiol. 2005; 58(3):700–13. https://doi.org/10.1111/j.1365-2958.2005.04875.x PMID: 16238621.

5. Nakamura MM, Liew SY, Cummings CA, Brinig MM, Dieterich C, Relman DA. Growth phase- and nutrient limitation-associated transcript abundance regulation in *Bordetella pertussis*. Infect Immun. 2006; 74(10):5537–48. https://doi.org/10.1128/IAI.00781-06 PMID: 16988229.

6. Deora R, Bootsma HJ, Miller JF, Cotter PA. Diversity in the *Bordetella* virulence regulon: transcriptional control of a Bvg-intermediate phase gene. Mol Microbiol. 2001; 40(3):669–83. PMID: 11359572.

7. Scarlato V, Arico B, Prugnola A, Rappuoli R. Sequential activation and environmental regulation of virulence genes in *Bordetella pertussis*. EMBO J. 1991; 10(12):3971–5. PMID: 1718746.

8. Veal-Carr WL, Stibitz S. Demonstration of differential virulence gene promoter activation in vivo in *Bordetella pertussis* using RIVET. Mol Microbiol. 2005; 55(3):788–98. https://doi.org/10.1111/j.1365-2958.2004.04418.x PMID: 15661004.

9. Cotter PA, Miller JF. A mutation in the *Bordetella bronchiseptica* bvgS gene results in reduced virulence and increased resistance to starvation, and identifies a new class of Bvg-regulated antigens. Mol Microbiol. 1997; 24(4):671–85. PMID: 9194696.

10. Deora R. Differential regulation of the *Bordetella* blpA gene: distinct roles for different BvgA binding sites. J Bacteriol. 2002; 184(24):6942–51. https://doi.org/10.1128/JB.184.24.6942-6951.2002 PMID: 12446644.

11. Uhl MA, Miller JF. Central role of the BvgS receiver as a phosphorylated intermediate in a complex two-component phosphorylase. J Biol Chem. 1996; 271(52):33176–80. PMID: 8969172.

12. Dupre E, Herrou J, Lensink MF, Wintjens R, Vagin A, Lebedev A, et al. Virulence Regulation with Venus Flytrap Domains: Structure and Function of the Periplasmic Moiety of the Sensor-Kinase BvgS. PLoS Pathog. 2015; 11(3):e1004700. PMCID: 4352136. https://doi.org/10.1371/journal.ppat.1004700 PMID: 25738876.

13. Dupre E, Lesne E, Guerin J, Lensink MF, Verger A, de Ruyck J, et al. Signal Transduction by BvgS Sensor-Kinase: Binding of Modulator Nicotinate Affects Conformation and Dynamics of Entire Periplasmic Moiety. J Biol Chem. 2015; 290:23307–319. https://doi.org/10.1074/jbc.M115.655720 PMID: 26203186.

14. Jacob-Dubuisson F, Wintjens R, Herrou J, Dupré E, Antoine R. BvgS of pathogenic Bordetellae: a paradigm for sensor kinase with Venus flytrap perception domains. In: Gross R, Beier D, editors. Two-component system in bacteria. Norfolk, UK: Caister Academic Press; 2012. p. 57–83.

15. Herrou J, Bompard C, Wintjens R, Dupre E, Willery E, Villeret V, et al. Periplasmic domain of the sensor-kinase BvgS reveals a new paradigm for the Venus flytrap mechanism. Proc Natl Acad Sci U S A. 2010; 107(40):17351–5. PMCID: 2951421. https://doi.org/10.1073/pnas.1006267107 PMID: 20855615.

16. Quiocho FA, Ledvina PS. Atomic structure and specificity of bacterial periplasmic receptors for active transport and chemotaxis: variation of common themes. Mol Microbiol. 1996; 20(1):17–25. PMID: 8861200.

17. Oh BH, Panditi J, Kang CH, Nikaido K, Gokcen S, Ames GF, et al. Three-dimenisonal structures of the periplasmic lysine/arginine/ornithine-binding protein with and without a ligand. J Biol Chem. 1993; 268(15):11348–55. PMID: 8496186.

18. Lesne E, Krammer EM, Dupre E, Locht C, Lensink MF, Antoine R, et al. Balance between Coiled-Coil Stability and Dynamics Regulates Activity of BvgS Sensor Kinase in Bordetella. MBio. 2016; 7(2): e02089. PMCID: 4810494. https://doi.org/10.1128/mBio.02089-15 PMID: 26933056.

19. Lesne E, Dupre E, Locht C, Antoine R, Jacob-Dubuisson F. Conformational changes of inter-domain linker mediate mechanical signal transmission in sensor-kinase BvgS. J Bacteriol. 2017; 199(18): e00114–17. https://doi.org/10.1128/JB.00114-17 PMID: 28507245.

20. Lesne E, Dupré E, Lensink MF, Locht C, Antoine R, Jacob-Dubuisson F. Coiled-coil antagonism regulates activity of Venus flytrap-domain-containing sensor-kinases of the BvgS family. mBio. 2018; 9(1): e02052–17. https://doi.org/10.1128/mBio.02052-17 PMID: 29487240.

21. McClure R, Balasubramanian D, Sun Y, Bobrovskyy M, Sumby P, Genco CA, et al. Computational analysis of bacterial RNA-Seq data. Nucleic Acids Res. 2013; 41(14):e140. https://doi.org/10.1093/nar/gkt444 PMID: 23716638.

22. Herrou J, Debrée AS, Willery E, Renaud-Mengen G, Locht C, Mooi F, et al. Molecular evolution of the two-component system BvgAS involved in virulence regulation in *Bordetella*. PLoS One. 2009; 4(9): e6996. https://doi.org/10.1371/journal.pone.0006996 PMID: 19750014.

23. Cummings CA, Bootsma HJ, Relman DA, Miller JF. Species- and strain-specific control of a complex, flexible regulon by *Bordetella* BvgAS. J Bacteriol. 2006; 188(5):1775–85. https://doi.org/10.1128/JB.188.5.1775-1785.2006 PMID: 16484188.
24. Coutte L, Huot L, Antoine R, Slupek S, Merkel TJ, Chen Q, et al. The multifaceted RisA regulon of *Bordetella pertussis*. Sci Rep. 2016; 6:32774. PMCID: 5020355. https://doi.org/10.1038/srep32774 PMID: 27620673

25. Moon K, Bonocora RP, Kim DD, Chen Q, Wade JT, Stibitz S, et al. The BvgAS Regulon of *Bordetella pertussis*. MBio. 2017; 8(5):e01526–17. PMCID: 5635692. https://doi.org/10.1128/mBio.01526-17 PMID: 29018122

26. Vergara-Irigaray N, Chavarrri-Martinez A, Rodriguez-Cuesta J, Miller JF, Cotter PA, Martinez de Tejada L, Huot L, Antoine R, Slupek S, Merkel TJ, Chen Q, et al. The multifaceted RisA regulon of *Bordetella pertussis* during experimental respiratory infection. Infect Immun. 2005; 73(2):748–60. https://doi.org/10.1128/IAI.73.2.748-760.2005 PMID: 15664913.

27. Nicholson TL, Brockmeier SL, Loving CL, Register KB, Kehrli ME Jr., Stibitz SE, et al. Phenotypic modulation of the virulent Bvg phase is not required for pathogenesis and transmission of *Bordetella bronchiseptica* in swine. Infect Immun. 2012; 80(3):1025–36. PMCID: 3294661. https://doi.org/10.1128/IAI.06016-11 PMID: 22158743

28. Miller JF, Johnson SA, Black WJ, Beattie DT, Mekalanos JJ, Falkow S. Constitutive sensory transduction mutations in the *Bordetella pertussis* bvgS gene. J Bacteriol. 1992; 174(3):970–9. PMID: 1732230.

29. Manetti R, Arico B, Rappuoli R, Scarlato V. Mutations in the linker region of BvgS abolish response to environmental signals for the regulation of the virulence factors in *Bordetella pertussis*. Gene. 1994; 150(1):123–7. PMID: 7959037.

30. Tam R, Saier MH Jr. Structural, functional, and evolutionary relationships among extracellular solute-binding receptors of bacteria. Microbiol Rev. 1993; 57(2):320–46. PMID: 8336670.

31. Felder CB, Graul RC, Lee AY, Merkle HP, Sadee W. The Venus flytrap of periplasmic binding proteins: an ancient protein module present in multiple drug receptors. AAPS PharmSci. 1999; 1(2):E2. https://doi.org/10.2153/apsp010202 PMID: 11741199.

32. van Beek LF, de Gouw D, Eleveld MJ, Bootsma HJ, de Jonge MI, Mooi FR, et al. Adaptation of *Bordetella pertussis* to the Respiratory Tract. J Infect Dis. 2018. 217(12):1987–96. https://doi.org/10.1093/infdis/ify125 PMID: 29528444

33. Lamberti Y, Cafiero JH, Surman K, Valdez H, Holubova J, Večerek B, et al. Proteome analysis of *Bordetella pertussis* isolated from human macrophages. J Proteomics. 2016; 136:55–67. https://doi.org/10.1016/j.jprot.2016.02.002 PMID: 26873878

34. Karataev GI, Sinyashina LN, Medkova AY, Semin EG, Shevtsova ZV, Matua AZ, et al. Insertional inactivation of virulence operon in population of persistent *Bordetella pertussis* bacteria. Russian Journal of Genetics. 2016; 52(4):370–7.

35. Bouchez V, Hegerle N, Strati F, Njamkep E, Guiso N. New Data on Vaccine Antigen Deficient *Bordetella pertussis* Isolates. Vaccines (Basel). 2015; 3(3):751–70. PMCID: 4586476.

36. Lam C, Octavia S, Ricafort L, Sintchenko V, Gilbert GL, Wood N, et al. Rapid increase in pertactin-deficient *Bordetella pertussis* isolates, Australia. Emerg Infect Dis. 2014; 20(4):626–33. PMID: 3966384. https://doi.org/10.3201/eid2004.131478 PMID: 24655754

37. Martin SW, Pawloski L, Williams M, Weening K, DeBolt C, Qin X, et al. Pertactin-negative *Bordetella pertussis* strains: evidence for a possible selective advantage. Clin Infect Dis. 2015; 60(2):223–7. https://doi.org/10.1093/cid/ciu788 PMID: 25301209.

38. Otsuka N, Han HJ, Toyoizumi-Ajisaka H, Nakamura Y, Arakawa Y, Shibayama K, et al. Prevalence and genetic characterization of pertactin-deficient *Bordetella pertussis* in Japan. PLoS ONE. 2012; 7(2):e31985. 3279416. https://doi.org/10.1371/journal.pone.0031985 PMID: 22348138

39. Pawloski LC, Queenan AM, Cassiday PK, Lynch AS, Harrison MJ, Shang W, et al. Prevalence and molecular characterization of pertactin-deficient *Bordetella pertussis* in the United States. Clin Vaccine Immunol. 2014; 21(2):119–25. PMCID: 3910938. https://doi.org/10.1128/CVI.00717-13 PMID: 24256623

40. de Gouw D, Hermans PW, Bootsma HJ, Zomer A, Heuvelman K, Diavatopoulos DA, et al. Differentially expressed genes in *Bordetella pertussis* strains belonging to a lineage which recently spread globally. PLoS ONE. 2014; 9(1):e84523. PMCID: 3885589. https://doi.org/10.1371/journal.pone.0084523 PMID: 24416242

41. Thorstensson R, Trolfors B, Al-Tawil N, Jahmzat M, Bergström J, Ljungman M, et al. A phase I clinical study of a live attenuated *Bordetella pertussis* vaccine BPZE1: a single centre, double-blind, placebo-controlled, dose-escalating study of BPZE1 given intranasally to healthy adult male volunteers. PLoS One. 2014; 9(1):e83449. https://doi.org/10.1371/journal.pone.0083449 PMID: 24421886

42. Locht C, Papin JF, Lecher S, Debre AS, Thalen M, Solovay K, et al. Live Attenuated Pertussis Vaccine BPZE1 Protects Baboons Against *Bordetella pertussis* Disease and Infection. J Infect Dis. 2017; 216 (1):117–24. https://doi.org/10.1093/infdis/jix254 PMID: 28535276
43. Taylor-Mulneix DL, Bendor L, Linz B, Rivera I, Ryman VE, Dewan KK, et al. *Bordetella bronchiseptica* exploits the complex life cycle of *Dictyostelium discoideum* as an amplifying transmission vector. PLoS Biol. 2017; 15(4):e2000420. PMCID: 5389573. https://doi.org/10.1371/journal.pbio.2000420 PMID: 28403138

44. Martinez de Tejada G, Cotter PA, Heininger U, Camilli A, Akerley BJ, Mekalanos JJ, et al. Neither the Bvg- phase nor the vrg6 locus of *Bordetella pertussis* is required for respiratory infection in mice. Infect Immun. 1998; 66:2762–8. PMID: 9596745

45. Seydlova G, Beranova J, Bibova I, Dienstbier A, Drzmisek J, M, et al. The extent of the temperature-induced membrane remodeling in two closely related *Bordetella* species reflects their adaptation to diverse environmental niches. J Biol Chem. 2017; 292(19):8048–58. PMCID: 28348085. https://doi.org/10.1074/jbc.M117.781559 PMID: 28348085

46. Bone MA, Wilk AJ, Perault AI, Marlatt SA, Scheller EV, Anthouard R, et al. *Bordetella* PirSR regulatory system controls BvgAS activity and virulence in the lower respiratory tract. Proc Natl Acad Sci U S A. 2017; 114(8):E1519–E27. PMCID: 5338435. https://doi.org/10.1073/pnas.1609565114 PMID: 28167784