Molecular Evolution of the Two-Component System BvgAS Involved in Virulence Regulation in *Bordetella*

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**Abstract**

The whooping cough agent *Bordetella pertussis* is closely related to *Bordetella bronchiseptica*, which is responsible for chronic respiratory infections in various mammals and is occasionally found in humans, and to *Bordetella parapertussis*, one lineage of which causes mild whooping cough in humans and the other ovine respiratory infections. All three species produce similar sets of virulence factors that are co-regulated by the two-component system BvgAS. We characterized the molecular diversity of BvgAS in *Bordetella* by sequencing the two genes from a large number of diverse isolates. The response regulator BvgA is virtually invariant, indicating strong functional constraints. In contrast, the multi-domain sensor kinase BvgS has evolved into two different types. The pertussis type is found in *B. pertussis* and in a lineage of essentially human-associated *B. bronchiseptica*, while the bronchiseptica type is associated with the majority of *B. bronchiseptica* and both ovine and human *B. parapertussis*. BvgS is monomorphic in *B. pertussis*, suggesting optimal adaptation or a recent population bottleneck. The degree of diversity of the bronchiseptica type BvgS is markedly different between domains, indicating distinct evolutionary pressures. Thus, absolute conservation of the putative solute-binding cavities of the two periplasmic Venus Fly Trap (VFT) domains suggests that common signals are perceived in all three species, while the external surfaces of these domains vary more extensively. Co-evolution of the surfaces of the two VFT domains in each type and domain swapping experiments indicate that signal transduction in the periplasmic region may be type-specific. The two distinct evolutionary solutions for BvgS confirm that *B. pertussis* has emerged from a specific *B. bronchiseptica* lineage. The invariant regions of BvgS point to essential parts for its molecular mechanism, while the variable regions may indicate adaptations to different lifestyles. The repertoire of BvgS sequences will pave the way for functional analyses of this prototypic system.

**Introduction**

*Bordetella pertussis*, the whooping cough agent, is an extremely contagious pathogen that infects the upper respiratory tract of humans and causes an acute infection [1]. The expression of most virulence factors of this Gram-negative bacterium, including adhesins and toxins, is controlled by the BvgAS two-component system (TCS) [2–5]. TCSs regulate major physiological responses in bacteria [6–9]. They are composed of two proteins, a sensor kinase that perceives (a) signal(s) and autophosphorylates a conserved histidine residue, and a response regulator that becomes activated upon phosphorylation by its cognate sensor kinase and often serves as a transcriptional activator [7,10,11]. When phosphorylated, the response regulator BvgA activates the transcription of the virulence-activated genes (vags) [15].

By controlling the intracellular concentration of phosphorylated BvgA, this system mediates a progressive transition between three phenotypic phases, Bvg (avirulent), Bvg (intermediate) and Bvg (virulent) [16–20]. The Bvg phase occurs at 37°C and is necessary for *B. pertussis* to cause respiratory infections in animal models [16,21]. The Bvg phase, in which some adhesins are produced, has been proposed to play a role in transmission and in the initial stages of infection [22]. Under laboratory conditions, negative signals such as nicotine, MgSO4 or low temperature trigger modulation to the Bvg phase [23]. A set of virulence-repressed genes (vrgs) is expressed in that phase [24].

*Bordetella parapertussis* and *Bordetella bronchiseptica* are close relatives of *B. pertussis* [25–28]. Two distinct lineages of *B. parapertussis* cause either a generally milder form of whooping cough in humans or ovine respiratory infections [29–33], and *B. bronchiseptica* causes chronic respiratory infections in various mammals and has also
been occasionally isolated from humans [34,35]. B. pertussis and B. parapertussis are thought to have derived from distinct B. bronchiseptica clones [28,33,36]. The virulence regulons of the three species are similar. They are also controlled by BvgAS and respond to the same negative modulators. B. bronchiseptica can survive outside its hosts in the Bvg− phase, which promotes motility and survival under nutrient-limiting conditions [37–39]. In contrast, B. pertussis and B. parapertussis have no known reservoir other than humans, and the function of their Bvg− phase is unclear. While B. bronchiseptica is fairly responsive to negative modulators, the sensitivity of B. pertussis to modulators appears to vary between isolates, which might suggest that the ability to down-modulate virulence has become dispensable for this species [16].

Because of the diversity of hosts and the different types of infection caused by these three Bordetella species, we investigated here the molecular diversity and evolution of BvgAS among the three Bordetella species. Sequencing of bvgA and bvgS from a number of isolates revealed that BvgA is almost invariant, while BvgS shows marked divergences between phylogenetic groups. Two evolutionary solutions for BvgS clearly appear from the analysis. The VFT2 domain is totally conserved within each of the two major BvgS types, indicating its pivotal role for the function of the protein.

**Results**

**Genotyping of bvgA and bvgS in Bordetella**

Full genomic sequences are available thus far for B. pertussis Tohama I, the B. bronchiseptica rabbit isolate RB50, and the human B. parapertussis isolate Bpp12922 [26]. The three genomes are closely related, with orthologous genes displaying little diversity between the three species. However, the bvgS genes markedly differ between them, while in contrast the bvgA genes are highly similar, harbouring small numbers of substitutions all of which are synonymous. To characterize the diversity of bvgA and bvgS, these genes were sequenced from a number of isolates selected from a collection described earlier [29]. In that previous phylogenetic study based on the sequences of housekeeping gene fragments, 4 major complexes of Bordetella were identified [28]. Complex I includes B. bronchiseptica isolates mainly of animal origin and the B. parapertussis ovine isolates (Bppov), complex II all B. pertussis isolates, complex III all B. parapertussis human isolates (Bpphu) and complex IV B. bronchiseptica isolates mainly of human origin. We thus selected isolates to represent these 4 complexes and all the sequence types identified in that study. We attempted to maximize the diversity of hosts, geographic origins and times of isolation.

In the 82 bvgS sequences, 247 distinct single nucleotide polymorphisms (SNPs) on a total of 3,717 bp were found (6.6%). Among these, 147 SNPs are synonymous substitutions (3.8%) and 100 non-synonymous substitutions (2.6%). This apparent diversity masks major differences between the Bordetella complexes (Table 1), as well as between domains of the protein (see below). Thus, bvgS is remarkably conserved in B. pertussis (complex II) with only two non-silent SNPs among all 29 isolates. This is illustrated by a very low diversity index (0.15), which is yet lower than that calculated from the set of housekeeping genes sequenced in [28] (0.63).

In contrast, the genetic diversity indices of both bvgA and the housekeeping genes calculated for complex IV and for complex I are significantly higher than those of B. pertussis (Table 1). Although B. parapertussis is thought to have emerged more recently than B. pertussis [28], bvgS is more diverse in complex III than in complex II, while the contrary is true for the housekeeping genes (Table 1).

bvgA was found to be almost invariant, with 15 SNPs in total (Table 1). Only three SNPs are non-synonymous, but each is found in only one isolate. This indicates an extremely strong pressure on bvgA in Bordetella. Accordingly, genetic diversity in all 4 complexes is lower for bvgA than for housekeeping genes or bvgS. Both bvgA and bvgS are strikingly conserved in B. pertussis (complex II). A phylogenetic tree was constructed based on the bvgS sequences (Fig. 1). This analysis clearly distinguishes two lineages of bvgS. The ‘‘pertussis type’’ (BvgSbp) encompasses bvgS from the 29 B. pertussis isolates as well as from the 12 B. bronchiseptica complex IV isolates. The other type, referred to hereafter as the ‘‘bronchiseptica type’’ (BvgSbs) encompasses bvgS from all 32 isolates that belong to the complex I of B. bronchiseptica and from the 9 isolates of B. parapertussis. Interestingly, the ovine and human B. parapertussis isolates do not form separate clusters but are included in the B. bronchiseptica complex I group, in contrast with previous studies [28,40]. Similarly, the human isolates of B. bronchiseptica complex I are found interspersed among the animal isolates and do not form a separate group.

A comparison of the bvgS-based phylogeny with that based on concatenated fragments from 7 different housekeeping genes described earlier [28] showed that while the overall topology of the two trees are similar, details differ. In addition to the absence of a
Figure 1. Phylogenetic tree based on the DNA sequences of bvgS. The lengths of the branches represent the phylogenetic distances. All human B. pertussis (Bp), B. bronchiseptica (Bb) and B. parapertussis (Bpp) isolates are shown in red. The major complexes identified in an earlier study [28] are boxed in colors, and the numbers in parentheses correspond to the sequence types defined in that study.
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separate complex III encompassing the *B. parapertussis* human isolates in the bvgS-based tree, some of the sequence types (ST) defined based on the housekeeping genes are split between several different branches of the *bvgS*-based tree, e.g. ST6, ST16 and ST19 (Fig. 1). These observations suggests that bvgS has evolved somewhat differentially from the housekeeping genes in the various branches.

Amino acid sequence analyses
To gain insight into the evolution of the functional domains of BvgS, the predicted amino acid sequences were aligned (Fig. 2). The two types of BvgS, BvgSBb and BvgSBp, appear distinctly from this analysis. BvgS is identical among 27 *B. pertussis* isolates, while the two strains widely used in laboratories harbour each one substitution, E705K for Tohama I and I124T for strain 18323,
respectively. BvgS of *B. bronchiseptica* complex IV is most similar to BvgS\textsubscript{Bb}, in the three domains potentially involved in signal perception, while the picture is slightly more blurred in the region corresponding to the phosphotransfer domains. BvgS from *B. parapertussis* resembles BvgS\textsubscript{Bp}, all along the sequence.

Remarkably, no more than two different residues are present at any given sequence position of the protein in all 82 isolates (Supporting information). Thus, in spite of the divergent evolution of BvgS between *B. bronchiseptica* complex I on the one hand, and *B. pertussis* and *B. bronchiseptica* complex IV on the other hand, selective pressure appears to limit the possible substitutions at the variable positions.

Our analyses revealed significant differences with respect to domain conservation. The first periplasmic domain, VFT1 totals the largest number of variable positions, although it is relatively well conserved within each type. Among the 82 isolates, 37 positions are variable in VFT1 over a total of 255 residues (14.5%). The overwhelming majority of the substitutions between the BvgS\textsubscript{Bb} and the BvgS\textsubscript{Bp} VFT1 are conservative in nature. In very rare instances, more drastic changes were observed, such as a Glu to Lys substitution at position 133 of BvgS\textsubscript{Bp} is invariant for all 41 isolates and a Val to Glu substitution at position 206 for a single complex I isolate (Supporting information). Unlike for B. pertussis, some degree of variation is found within the complex IV VFT1s. Intriguingly, 4 of the substitutions identified in complex IV are not at the positions that differentiate BvgS\textsubscript{Bb} from BvgS\textsubscript{Bp}. They are thus specific of complex IV.

One isolate, B0226, stands out from the *B. bronchiseptica* complex I by a small portion of VFT1. At 7 positions between residues 133 and 166 it shares *B. pertussis* sequences, although it also harbours unique substitutions in that region.

For VFT2 essentially two types of sequence have evolved. VFT2 of BvgS\textsubscript{Bb} is invariant for all 41 *B. pertussis* and *B. bronchiseptica* complex IV isolates, suggesting that this domain is under strong stabilizing selection. VFT2 of BvgS\textsubscript{Bp} also displays a very limited degree of variation. Furthermore, only conservative substitutions are found between the two types except for one at position 404, where Ala in BvgS\textsubscript{Bb} is replaced by Asp in BvgS\textsubscript{Bp}. In total, 19 positions are different between the pertussis type and the *bronchiseptica* type VFT2s over a total of 254 residues (7.5%). There are no intermediates between the two VFT2 types. All these observations argue that VFT2 is pivotal to the function of BvgS and is under strong selective pressure in each lineage.

The PAS domain is well conserved among all isolates, suggesting that it is also under stabilizing pressure. In total only 5 positions are variable, and the few substitutions found in this domain are all conservative. Similar to VFT1 and VFT2, the PAS domain distinguishes *B. bronchiseptica* complex I from *B. pertussis* and *B. bronchiseptica* complex IV. Thus, for all three putative perception domains, *B. bronchiseptica* complex IV clearly sides with *B. pertussis*.

For the His-kinase domain, in contrast, *B. bronchiseptica* complex IV sequences appear to be more similar to BvgS\textsubscript{Bb} than to BvgS\textsubscript{Bp}. Altogether the His-kinase domain is also much more conserved than the two VFT domains, with only 11 variable positions harbouring essentially conservative substitutions over a total of 223 residues (4.9%). This probably points to strong functional constraints on this domain.

The receiver domain is more diverse, with 9 variable positions over a total of 122 residues (7.3%). Intriguingly, the receiver domain of *B. bronchiseptica* complex IV is of the BvgS\textsubscript{Bb} type, unlike the His-kinase domain.

The Hpt domain is well conserved, with only 4 variable positions for a total of 98 residues (4.1%). Intriguingly, at position 1190, the conservative Val to Ala substitution differentiates *B. pertussis* from *B. bronchiseptica* of both complexes I and IV. Altogether, thus, complex IV appears to be intermediate between the BvgS\textsubscript{Bb} and the BvgS\textsubscript{Bp} types for the His-kinase and Hpt domains, in contrast with the VFT1, VFT2, PAS and receiver domains. This suggests that the distinct domains have evolved at different rates.

Distinct functional constraints appear to operate also on the various linkers that join the domains. The transmembrane helix is invariant, except for a unique conservative Gly to Ala change. Similarly, the linkers A and B between the membrane and the PAS domain and between the PAS and His-kinase domains respectively are invariant, indicating that they are under strong selective pressure (Fig. 2). One striking exception to this rule is that a Glu residue is replaced by a Lys residue in Tohama I at position 705 (see below). In marked contrast with the strict conservation of linkers A and B, the linker C between the His-kinase and receiver domains seems to accumulate a disproportionate number of mutations, with 20% positions that are variable. Linker D between the receiver and Hpt domains harbours 2 variable positions over 36 residues.

**Mapping of the substitutions**

In order to map the positions of the substitutions, structural models were constructed for several domains of BvgS based on available X-ray structures of homologous proteins, since the crystal structure of BvgS is not available. VFT1 and VFT2 were modelled on the glutamine-binding protein, a bacterial periplasmic binding protein of an ABC transporter. The overwhelming majority of the substitutions between the two broad phylogenetic groups map to the external surfaces of both VFT1 and VFT2, leaving the potential solute-binding cavities conserved (Fig. 3). In addition, a few substitutions appear to be in the hydrophobic core of VFTs, but they are highly conservative and thus unlikely to affect the structure or stability of the proteins. Thus, the tertiary structures of the two VFT domains and their potential ligand-binding pockets are likely to be totally conserved, while the external surfaces of the proteins are quite different between the two BvgS types. The more drastic substitution between BvgS\textsubscript{Bb} (Asp\textsubscript{104}) and BvgS\textsubscript{Bp} (Ala\textsubscript{104}) is located in an external loop in the second lobe of VFT2. Unlike non-synonymous substitutions, synonymous substitutions are found both in the ligand binding cavities and on the surfaces of the VFTs, indicating differential selective pressure between cavities and surfaces.

A model of the His-kinase domain of BvgS was also constructed. The ATP-binding pocket appeared to be conserved among all isolates, as expected for the activity of the protein (not shown). A drastic substitution located in the linker B that precedes the His-kinase domain distinguishes *B. pertussis* Tohama I from all other *B. pertussis*, *B. bronchiseptica* and *B. parapertussis* isolates. Tohama I harbours a Lys at position 705 instead of a Glu in the 81 others. This linker is predicted to form an $\alpha$ helix immediately preceding the dimerisation helix of the His-kinase domain. Conceivably, this substitution might affect the conformation of this region most likely crucial for signal transduction (see below).

The structure of an Hpt domain in complex with its cognate phosphorylation substrate has been reported [41,42], showing the interface between the two proteins. The BvgS Hpt domain has two interaction partners, the receiver domain and BvgA. The surfaces of Hpt predicted to form interfaces with its partners appear to be conserved as well (not shown). Similarly, neither of the three frequent substitutions between the receiver domains of the two BvgS types map to the predicted interface with Hpt (not shown).
Figure 3. Structural models of VFT1 and VFT2. The models of VFT1 (A) and VFT2 (B) are based on the structure of the glutamine binding protein in an unliganded, open conformation (PDB 1GGG). The elements of secondary structure are shown in ribbon representation in lime green, and the surfaces of the proteins are shown in semi-transparency. The cavity of each of the two domains corresponds to the groove between the two lobes. The positions of the substitutions that distinguish the BvgSBp and BvgSBb types are depicted as red balls. Only the main chain atoms are shown. The rare substitutions have not been represented.

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Regarding BvgA, only 3 unique substitutions are found each in one isolate, namely Glu to Gly, Asp to Gly and Val to Glu at positions 64, 137 and 197, respectively. Sequence alignments of response regulators homologous to BvgA as well as the X-ray structure of one such protein indicate that these three substitutions are at non-conserved positions in the superfamily (not shown). The rarity and the positions of non-synonymous replacements in BvgA strongly indicate that the protein is under stringent selection.

Functional implications of sequence variations

Thus the vast majority of the predicted amino acid differences between pertussis type and bronchiseptica type BvgS are located in the periplasmic domains, essentially at the surface of the two VFT domains that appear to have co-evolved within each lineage. This suggests that the two domains operate together through inter-domain interactions, most likely for proper signal transduction. To test this hypothesis, we determined whether the sequence variations identified between the VFT1 and VFT2 of the two types of BvgS affect its activity or its level of sensitivity to the known negative virulence modulators. BvgS harbouring the BvgSb, periplasmic domain of our laboratory strain B. pertussis BPSM, a Tohama I derivative, was substituted with BvgS harbouring the BvgSb, periplasmic domain from B. bronchiseptica RB50, a typical complex I isolate (Fig. 4A). We also performed single domain exchanges. The activities of chimeric BvgS proteins and their responses to modulation were determined using the reporter gene lacZ placed under the control of the Bvg-regulated promoter of the pertussis toxin operon.

The replacement of both VFT1 and VFT2 of the BPSM BvgS by those of RB50 (BPb1+2) did not affect the level of BvgS activity in the absence of modulation, but it made the recombinant bacteria much more sensitive to nicotinic acid than their parent (Fig. 4B,C). Thus, the BvgSm and BvgSb, periplasmic domains of BvgS influence the response to this negative modulator.

Similarly, the replacement of the BPSM VFT2 by that of RB50 (BPs2), yielding a VFT1b-VFT2b, chimera strongly enhanced the sensitivity of BvgS to both nicotinic acid and MgSO4 (Fig. 4B,C). In contrast, introduction of the RB50 VFT1 into the BPSM BvgS (BPb1) did not modify the response of the recombinant bacteria to either modulator (not shown). Thus, VFT2 is essential in determining sensitivity to modulation.

Because of its drastic character in an otherwise totally invariant linker segment, the substitution identified in the Tohama I BvgS was also investigated. Tohama I has been described as relatively insensitive to modulation compared with other B. pertussis isolates [16], and thus we tested whether Lys705 is the cause for this lack of sensitivity (Fig. 5A). Glu at position 705 in the BPSM BvgS made the recombinant strain more responsive to nicotinate than its parent, with 4 mM nicotinic acid causing the complete loss of BvgS activity (Fig. 5B). The BPb705 substitution also enhanced the responsiveness to MgSO4 (Fig. 5C). Thus, the presence of a Lys at position 705 is responsible for the lack of sensitivity to modulation of Tohama I. This substitution is quite rare in Bordetella, suggesting that the mutation acquired in Tohama I is not particularly advantageous. The other unique substitution identified in Bp18323, 1241T, was also introduced into BPSM (BPb124). The level of β-gal activity and the responsiveness to modulators of BPb124 was identical to that of BPSM, indicating that this is likely a neutral substitution (not shown).

Since both VFT2 and the linker preceding the HK domain determine the sensitivity to modulators, we tested whether their effects are cumulative. Therefore, a new chimeric strain, BPb2+705 was constructed that harbors both the VFT2b domain and Glu at position 705 (Fig. 5A). BPb2+705 was hypersensitive to modulators, with as low as 0.25 mM nicotinic acid (Fig. 5B) or 5 mM MgSO4 (Fig. 5C) being sufficient for complete modulation.

Hypermodulator B. pertussis in an animal model of infection

No such hypersensitive variant has been characterized in natural isolates so far [16]. Thus, it was interesting to determine
whether this phenotype might affect \textit{B. pertussis}'s ability to colonize the respiratory tract in an animal model of infection. BPSM or BPBb2+E705 were instilled intranasally to two groups of mice, and the numbers of bacteria recovered from the lungs were counted after various periods of time. No significant difference was observed between the two strains (not shown). To determine whether BPBb2+E705 might be disadvantaged in a mixed infection, mice were also inoculated intranasally with equal numbers of the two bacterial strains together. The two strains colonized the lungs of mice in the same manner (Fig. 6). Thus, in this model of experimental infection, \textit{B. pertussis} hypersensitive to modulation does not appear to be defective for colonization.

\section*{Discussion}

In this work, we investigated the molecular evolution of the BvgAS two-component system, which is central to the regulation of \textit{Bordetella} virulence. BvgA is remarkably conserved. In contrast, the sequences of BvgS differ markedly between species, and in particular can be separated into two lineages of \textit{Bordetella}. Interestingly, one of the two lineages includes all \textit{B. pertussis} isolates and a subset of \textit{B. bronchiseptica} isolates previously identified as forming a distinct \textit{B. bronchiseptica} complex (called complex IV), mainly isolated from humans \cite{28}. Our results confirm the hypothesis that complex IV strains are an evolutionary intermediate between animal and human adapted \textit{Bordetella} from which \textit{B. pertussis} evolved \cite{28}. The degree of divergence of BvgS varies significantly between lineages, as shown by the genetic diversity indices. In \textit{B. pertussis}, BvgS displays an extremely low level of diversity, indicating that it has reached equilibrium in this highly successful human colonizer or has undergone a recent selective sweep. With a broad range of hosts, complex IV may be using a different evolutionary strategy, and its BvgS may still be in a process of diversification.

Our analyses do not provide evidence of an absolute correlation between the type of \textit{bvgS} allele found in a given \textit{Bordetella} lineage and the hosts of its members. Thus, two bona fide human pathogens, \textit{B. pertussis} and \textit{B. parapertussis}, harbour BvgS of the BvgSBp type and of the BvgSBb type, respectively, although the latter type otherwise corresponds to \textit{B. bronchiseptica} complex I, with isolates of essentially animal origin. Similarly, the ovine and human isolates of \textit{B. parapertussis} have the same BvgS type although they infect distinct hosts. It is interesting that BvgS is monomorphic in \textit{B. pertussis} but not in \textit{B. parapertussis}, as \textit{B. pertussis} is assumed to be older \cite{28}. This may reflect that \textit{B. parapertussis} is not yet optimally adapted to its host, unlike \textit{B. pertussis} \cite{33}.

Remarkably, the various domains of BvgS display widely different degrees of diversity, which shows that they are under different functional constraints, and/or parts of the protein are subject to diversifying selection in some lineages. The periplasmic domains of BvgS appear to have evolved most extensively. However, their evolution has been restricted to external surfaces, strongly arguing that the binding paradigm for Venus Fly Trap
domains applies to BvgS. Purifying selective pressure on the 
cavities argues that they bind specific ligands. The prevalent model 
proposes that BvgS is in a permanently activated state in the 
absence of ligand, unlike classical sensors of TCS. If this is correct, 
then the conservation of the cavities is important essentially for the 
perception of negative signals. However, we have obtained preliminary data that specific substitutions in the VFT1 cavities 
markedly decrease the activity of BvgS (our unpublished data). 
This supports the idea that the cavities of the VFT domains also 
perceive positive signals, at odds with the generally accepted model. 

The external surfaces of the VFT domains have evolved 
considerably, arguing that they undergo a lesser selective pressure 
or are even diversifying. However, because there are no 
intermediate types between the BvgS\textsubscript{Bb} and BvgS\textsubscript{Bp} types, the 
evolutionary pressure must have selected for functional combina-
tions of the two VFT surfaces. In other words, the surfaces may 
have co-evolved in each of the two major lineages because 
interactions between the two domains must be fine-tuned for 
optimal function. The combination of VFT\textsubscript{1Bp} and VFT\textsubscript{2Bp} in 
particular appears to significantly increase the sensitivity of BvgS 
to modulators. We thus propose that the sensitivity to modulators 
depend on the efficiency of signal transduction between the two 
VFT domains and between VFT2 and the cytoplasmic portion 
through the membrane segment. The membrane-proximal 
position of VFT2 most likely makes it especially important in this 
respect. Similarly, the two types of BvgS harbour distinct sets of 
cytoplasmic domains, although the situation is more blurred in 
the case of complex IV. This suggests that there may be small 
functional differences in the phosphorylation cascade between the 
two types. 

The region of the protein with the lowest degree of diversity 
comprises the transmembrane segment, the following cytoplas-
mic linker and the PAS and His-kinase domains, emphasizing the 
functional constraints on that region. The transmembrane 
segment must transmit signals perceived by the periplasmic 
domain to the cytoplasmic domain in a purely mechanical 
manner, such as a rotational or a piston-like motion. This signal 
must then be conveyed mechanically by the linker to the PAS 
domain, and then from the PAS domain to the His-kinase domain. 
PAS is thus central to transduction between periplasmic and His-
kinase domains. In agreement with this model, several mutations 
that make BvgS insensitive to negative modulating signals map to 
the linker and PAS domains [43,44]. Following phosphorylation of 
the His-kinase, signal transmission to BvgA is enzymatic, and thus 
functional constraints are exerted on the interfaces between 
domains, for interactions in trans and/or in cis, rather than on 
linkers [45]. In agreement with this idea, the linkers between the 
domains involved in the phosphorylation cascade have diverged 
more rapidly, most likely because they do not undergo strong 
selective pressures. 

Only 3 rare substitutions were identified in BvgA. BvgA forms 
many interactions that are all important for its function. It forms 
homodimers, interacts with the Hpt domain for phosphotransfer 
and with its target DNA sequences and the RNA polymerase. 
This most likely explains why BvgA undergoes such a strong stabilizing 
pressure in pathogenic Bordetella. 
The importance of responsiveness to modulating agents for the 
function of BvgS \textit{in vivo}, and the relevance of nicotinate and 
MgSO\textsubscript{4} are unclear. The Bvg phase is most likely more 
important for \textit{B. bronchiseptica} to survive outside the host and cause 
chronic infections [46,47] than it is for the lifestyle of \textit{B. pertussis}. Nevertheless, the fact that both types of BvgS have retained similar 
sensitivities to negative modulators argues that this phenotype is 
nevertheless relevant for both species. Thus, no natural \textit{B. pertussis} 
isolates appear to be “locked” in the Bvg\textsuperscript{+} phase, even though such 
strains can arise by simple point mutations in the laboratory 
[43,44,48]. Therefore, the ability to down-modulate virulence 
factor expression must confer some advantage upon the 
bacterium, which is supported by the fact that Tohama I, with 
its low sensitivity to modulators, is the exception among \textit{B. pertussis}. 
Conversely, we wondered whether \textit{B. pertussis} with a BvgS variant 
hypermutable to modulation would be impaired for colonisation. 
Surprisingly, this strain appeared to be functional in the mouse 
model of infection. The observation that no naturally occurring 
\textit{Bordetella} isolate has been described to harbour such a hyper-
responsive BvgS nevertheless suggests that this phenotype might be 
detrimental. The animal model used in this study does not 
reproduce all the features of a human infection, and in particular 
bacterial transmission. Better models may be needed to detect 
suble regulation defects. 

Materials and Methods 

Ethics statement 

All animal experiments were performed at the animal facility of the 
Institut Pasteur de Lille (number A59-35-064, Lille, France) 
according to the rules of the European Community Council 
guidelines (86/609/EEC) for laboratory animal experimentation. 
The animal protocol was approved by the local institutional review 
board (Comité d’Ethique en Expérimentation Animale Nord-Pas-
De-Calais, CEEA 03/2009). 

Sequencing of the circulating infectious \textit{Bordetella} 
isolates 

The \textit{B. pertussis}, \textit{B. bronchiseptica} and \textit{B. parapertussis} isolates 
were selected from the collection described in [28]. \textit{bvgS} was sequenced 
for 28 isolates of \textit{B. pertussis}, 8 of \textit{B. parapertussis} (5 \textit{Bpp}\textsubscript{Bb} and 3 
\textit{Bpp}\textsubscript{Bp}), 31 of \textit{B. bronchiseptica} from complex I and 12 of \textit{B. 
bronchiseptica} from complex IV [28] (Table S2). \textit{bvgA} was sequenced 
for 27 isolates of \textit{B. pertussis}, 5 \textit{B. parapertussis} isolates (5 \textit{Bpp}\textsubscript{Bb} 
and 3 \textit{Bpp}\textsubscript{Bp}), 34 \textit{B. bronchiseptica} isolates from complex I and 13 from 
complex IV (Table S3). The strains were cultivated on BG blood 
agar medium during 16 hours for \textit{B. bronchiseptica} and 48 hours for 
\textit{B. pertussis} and \textit{B. parapertussis}. Chromosomal DNA was extracted 
using the Illustra\textsuperscript{TM} bacterial genomic Prep Mini Spin Kit (GE 
Healthcare) according to the manufacturer’s instructions. For 
PCR amplification, several partially overlapping PCRs were 
performed by using several pairs of primers, \textit{bvgA}'-Up and 
\textit{bvgA}'-Lo, \textit{VFT1}'-Up and \textit{VFT1}'-Lo, \textit{VFT2}'-Up and \textit{VFT2}'-Lo, 
\textit{PAS/HisKin-Up} and \textit{PAS/HisKin-Lo}, \textit{R-UP} and \textit{R-Lo} and \textit{Hpt-
Up} and \textit{Hpt-Lo} (Table S1). The following conditions for the mixes 
were used: HotStar Taq DNA polymerase in the presence of Q 
buffer and MgCl\textsubscript{2} (QIagen), 30 cycles of 1 min at 95°C, 1 min at 
57°C and 1 min at 72°C. The PCR products were purified by 
using the PCR Purification Kit or the Gel Extraction Kit (Qiagen), 
depending on the degree of purity of the amplicons. The DNA 
fragments were sequenced by Genoscreen using an ABI 377 
sequencer (Lille, France). The sequences were reassembled and 
compared by using the DNAstar software. A number of targets 
were amplified and sequenced twice to determine the level of error 
introduced by the PCR and sequencing steps. In particular, all 
fragments with a unique sequence type were checked in 
this manner. No discrepancy was obtained in any case. 

Sequence data analysis 

The genetic diversity for each complex was calculated using the 
Shannon-Wiener index of diversity using the following formula in
which \( p_i \) is the frequency of the \( i \)th type:

\[
\text{Shannon-Wiener index} = - \sum p_i \ln p_i
\]

Single nucleotide polymorphisms (SNP) and their synonymous or non-synonymous characteristics were evaluated by using the DnaSP software. Neighbour-joining trees were constructed by using the www.phylogeny.fr software.

Models of the BvgS domains

The PDB codes of the X-ray structures used to build models for the various domains of BvgS were 1GGG (VFTs), 2C2A (His-kinase), 2AYX (receiver domain) and 2AOB (Hpt domain), and 3C3W for BvgA. The Modeller software was used on the following portions of BvgS: residues 33–287 for VFT1, 288–541 for VFT2, 726–946 for the His-kinase domain, 974–1095 for the receiver domain and 1133–1228 for the Hpt domain.

Construction of chimeric B. pertussis strains

B. pertussis BPSM is a Tohama I-derivative that is resistant to streptomycin [49]. The \( bvgS \) deletion strain BPSM\textsubscript{ΔbvgS} was constructed as follows. The 5’ and 3’ extremities of the \( bvgS \) locus were amplified by PCR using the BPSM chromosome as template and the oligonucleotides \( bvgA\text{-Up} \) and \( bvgA\text{-Lo} \), and BvgS\text{-Up} and BvgS\text{-Lo} as primers, respectively (Table S1). The amplicons were inserted directly into pCR\textsuperscript{®} II-TOPO (Invitrogen) and sequenced. They were then successively introduced as EcoRI\textsuperscript{−}KpnI and XbaI\textsuperscript{−}HindIII fragments into pUC19, yielding pUC19\textsubscript{ΔbvgS}. The resulting 1.0 kb EcoRI\textsuperscript{−}HindIII fragment was introduced into the EcoRI\textsuperscript{−}HindIII sites of pSORTP1, a mobilizable plasmid used for conjugation [50]. Conjugation was performed on BG-blood agar plates containing 10 mg/ml MgCl\textsubscript{2} for 6–7 hours, and co-integrates were selected on BG-blood agar plates containing 10 \( \mu \)g/ml gentamycin and 30 \( \mu \)g/ml nalidixic acid to prevent growth of the \( E. coli \) donor. Allelic exchange was selected by two successive steps as described [50]. After 4 to 5 days growth on selective media, isolated non-haemolytic streptomycin-resistant colonies, characteristic of the Bvg\textsuperscript{−} phase were analysed by PCR to confirm the deletion.

Recombinant BPSM strains containing chimeric \( bvgS \) genes were constructed as follows. Successive portions of the \( bvgS \) locus of BPSM were amplified by PCR in order to introduce restriction sites using silent mutations at specific sites corresponding to junctions between structural domains. The sequences of the oligonucleotides used as primers are given in Table S1. In addition, an EcoRI restriction site naturally occurring in the region of the gene coding for the PAS/His-kinase portion of \( bvgS \) was removed by overlapping PCR using the pairs of primers ΔEcoRI- Up and PAS/HisKin-Lo, and ΔEcoRI-Lo and PAS/HisKin-Up. The various amplicons were sequenced and introduced successively into pUC19. The resulting “mosaic” \( bvgS \) locus includes the 5’ and 3’ extremities of the operon remaining in BPSM\textsubscript{ΔbvgS} and necessary for allelic replacement in that strain. The mosaic \( bvgS \) locus allowed us to replace selectively the genetic cassettes encoding each of the two VFT-like domains and the PAS/His-kinase domains of the BPSM \( bvgS \) by the corresponding cassettes of B. pertussis 18323 (to introduce Ghu705) or B. bronchiseptica RB50. This procedure generated a number of \( bvgS \) variants that were then excised from pUC19 by restriction with EcoRI and HindIII and introduced into the EcoRI and HindIII sites of pSORTP1. Each \( bvgS \) variant was then introduced into BPSM\textsubscript{ΔbvgS} by allelic exchange, using conjugation as described above. Our criterion for the selection of recombinant clones was the restoration of hemolysis, which can be easily detected on BG-blood agar.

Constructions of \( ptx\text{-}lacZ \) transcriptional fusions and measurement of \( \beta \)-galactosidase activity

A recombinant pFUS plasmid, harbouring \( lac\text{-}z \) in transcriptional fusion with the sequence of the first gene of the pertussis toxin operon [51], was introduced into the different strains by conjugation, and the integrants were selected on BG blood agar containing 100 \( \mu \)g/ml streptomycin and 10 \( \mu \)g/ml gentamycin. The recombinant strains were grown in modified Stainer-Scholte medium (SS) [52] containing the relevant antibiotics. After overnight growth at 37°C under rotating agitation, the bacterial suspension was used to initiate cultures in 10 ml of SS medium containing increasing concentrations of nicotinate or MgSO\textsubscript{4}. The inoculation volume was adapted to compensate for slower growth in the presence of high concentrations of nicotinate. The bacteria were grown until the cultures reached an \( OD_{600} \) of 1.5. They were harvested by centrifugation and broken by using a Hybaid Ribolysing apparatus (35 s at speed 6 in tubes containing 0.1 mm silica spheres as the lysing matrix). \( \beta \)-galactosidase activities were measured as described [51]. The experiments were performed in triplicate.

Murine respiratory tract infections

Experimental infections were performed at biosafety level 2 facilities with 8 week-old female BALB/c mice purchased from Charles River Laboratories.

Two groups of mice slightly sedated by intraperitoneal pentobarbital injection were inoculated with 1.6\( \times \)10\textsuperscript{5} bacteria from BPSM and BP\textsubscript{Bb2\textsuperscript{E705}}, suspensions, by depositing 20 \( \mu \)l droplets into the nostrils. Groups of four animals per strain were sacrificed at days 0, 7, 14, 21 and 28 post-inoculation, and lungs were removed. Lung colonization was quantified by homogenizing the entire lungs in 2 to 5 ml phosphate-buffered saline (PBS), plating 100 \( \mu \)l aliquots of serial dilutions of the lung suspensions onto BG blood agar with 100 \( \mu \)g/ml of streptomycin, and counting the colonies after 4 days of incubation at 37°C.

Mixed infection essays were performed with BPSM carrying a gentamycin resistance marker at the 3’ end of the \( hct/\beta\text{RTF} \) operon [53] to distinguish between this strain (called hereafter BPSMG\textsuperscript{WT} and BP\textsubscript{Bb2\textsuperscript{E705}}). In this case, the inoculum consisted of 1.6\( \times \)10\textsuperscript{5} CFU of each bacterial strain suspended in a final volume of 20 \( \mu \)l. Five animals per time point were sacrificed at days 0, 3, 7, 14, 21 and 28 post-inoculation and lung colonization was quantified as described above. Serial dilutions of each suspension were plated onto BG blood agar with 100 \( \mu \)g/ml of streptomycin or with 10 \( \mu \)g/ml of gentamycin. Colonies grown on gentamycin (corresponding to BPSMG\textsuperscript{WT}) or BP\textsuperscript{Bb2\textsuperscript{E705}} were subtracted from the number of colonies present on the streptomycin-containing medium (corresponding to both BPSMG\textsuperscript{WT} and BP\textsuperscript{Bb2\textsuperscript{E705}}) to determine the number of BP\textsuperscript{Bb2\textsuperscript{E705}} bacteria present in the mice lungs.

Supporting Information

Table S1 Oligonucleotides used in this study

Found at: doi:10.1371/journal.pone.0006996.s001 (0.07 MB PDF)

Table S2 bvgS sequences

Found at: doi:10.1371/journal.pone.0006996.s002 (0.35 MB RTF)

Table S3 bvgA sequences
Author Contributions
Conceived and designed the experiments: JH FJD RA. Performed the experiments: JH ASD EW. Analyzed the data: JH FJD RA. Contributed reagents/materials/analysis tools: FRM. Wrote the paper: JH GRM CL FRM FJD RA.

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References