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# How genomics is changing what we know about the evolution and genome of *Bordetella pertussis*

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## Abstract

The evolution of *Bordetella pertussis* from a common ancestor similar to *Bordetella bronchiseptica* has occurred through large-scale gene loss, inactivation and rearrangements, largely driven by the spread of insertion sequence element repeats throughout the genome. *B. pertussis* is widely considered to be monomorphic, and recent evolution of the *B. pertussis* genome appears to, at least in part, be driven by vaccine-based selection. Given the recent global resurgence of whooping cough despite the wide-spread use of vaccination, a more thorough understanding of *B. pertussis* genomics could be highly informative. In this chapter we discuss the evolution of *B. pertussis*, including how vaccination is changing the circulating *B. pertussis* population at the gene-level, and how new sequencing technologies are revealing previously unknown levels of inter- and intra-strain variation at the genome-level.

## Keywords

*Bordetella pertussis*, whooping cough, evolution, DNA sequencing, genomic variation

## Abbreviations

ACV – acellular vaccine  
ACT - adenylate cyclase toxin  
CGH – comparative genomic hybridization  
COG - Clusters of Orthologous Genes  
FHA – Filamentous haemagglutinin  
Fim(2/3) – Fimbrial protein (2/3)  
HGT – horizontal gene transfer  
INDEL – insertions and deletions  
IS – insertion sequence

MLST – multilocus sequence typing  
 MLVA - multilocus variable-number tandem-repeat analysis  
 NGS – next generation sequencing  
 ONT – Oxford Nanopore Technologies  
 PacBio – Pacific Biosciences  
 PERISCOPE - PERTussis Correlates Of Protection Europe (consortium)  
 PFGE – pulsed field gel electrophoresis  
 prn – Pertactin  
 ptx – Pertussis toxin  
 SMRT – single-molecule real-time  
 SNP – single nucleotide polymorphism  
 WCV – whole-cell vaccine  
 WGS – whole genome sequencing

## 1. The ongoing problem of *Bordetella pertussis*

Whooping cough, the infectious respiratory disease caused by the bacterium *Bordetella pertussis*, has been resurgent in many countries for the past two decades. This resurgence comes in spite of a global vaccination programme, with 90% of the target population receiving a single dose of *pertussis*-containing vaccine, and 85% receiving three doses (WHO 2018). In addition, there has been a shift in the epidemiological profile of the disease: whereas once most cases were reported in infants and unvaccinated children, the resurgence is also affecting vaccinated children, adolescents and adults,

**Figure 1** (Strebel et al. 2001; Clark 2014). Data from the Centers for Disease Control, for example, show

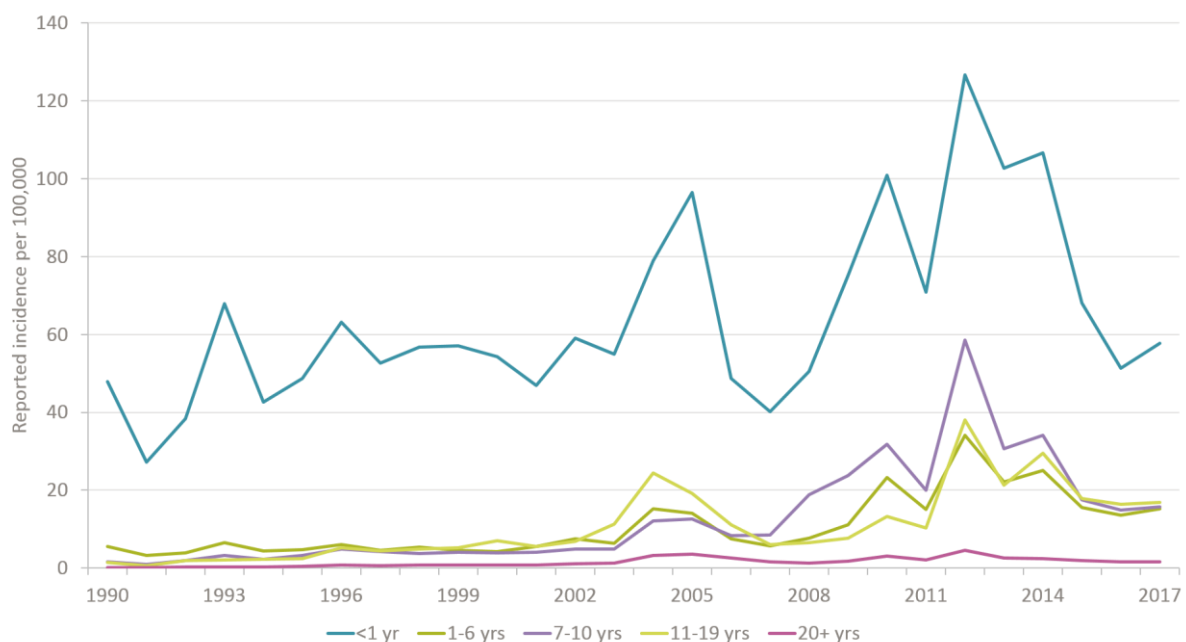


Figure 1: Reported incidence of whooping cough per 100,000 in the USA, 1990-2017, showing increase in all age brackets. Data source: CDC National notifiable diseases surveillance reports (Centers for Disease Control 2019)

that from 1990-1997, the mean incidence of whooping cough per year in 11-19 year-olds was 2.54 per 100,000 people, whereas from 2010-2017, it was 20.43 (Centers for Disease Control 2019).

The original vaccination programme, introduced in the 1940s and 1950s, used a whole-cell vaccine (WCV). Initially, cases of the disease appeared to drop significantly. Due to perceived reactogenicity of the WCV (now largely discredited), it was replaced in many countries throughout the 1990s and early 2000s with an acellular vaccine (ACV) (for example: Cherry 1990, 1992, 1996; Cherry et al. 1993; Blumberg et al. 1993; Moore et al. 2004). The acellular vaccine contains one to five *B. pertussis* antigens, including Pertussis toxin (ptx), pertactin (prn), filamentous haemagglutinin (FHA) and the fimbrial proteins Fim2 and Fim3. These days, most developed countries use the ACV, although many developing countries continue to use the WCV. The recent use of the ACV has been strongly implicated in whooping cough's resurgence. However, concerns were raised about waning immunity conveyed by the WCV beginning in the early 1990s and, in many countries, the resurgence does seem to pre-date the switch to the ACV (De Serres et al. 1995; Cherry 1996; de Melker et al. 1997).

Three main potential causes are thought to have contributed to the recent observed increase in whooping cough cases: increased awareness of the disease coupled with improved diagnosis due to a switch from culture-based to PCR- and serology-based techniques; waning immunity, particularly that conveyed by the ACV compared to the WCV; and genetic variations in circulating *B. pertussis* strains, away from the vaccine strains (Sealey et al. 2015; Clark 2014; Ausiello and Cassone 2014). Here we focus on the latter from two different but highly interrelated perspectives: variation at the gene-level, and variation at the genome-level, with particular consideration of how recent developments in genomic research have contributed to our understanding of evolution and variation in a species which traditionally has been described as highly monomorphic (for example: Bart et al. 2010; Mooi 2010).

## 2. The speciation of *Bordetella pertussis*

The study of the bordetellae has focussed largely on the three classical, pathogenic, *Bordetella* species: *B. bronchiseptica*, *B. pertussis* and *B. parapertussis*. However, the *Bordetella* genus contains many additional species which have been isolated from extremely diverse environments, including marine sponges, bioreactors, nitrifying sludge and mural paintings in ancient tombs (Wang et al. 2007; Bianchi et al. 2005; Sun et al. 2019; Tazato et al. 2015). Using previously published 16S sequence data derived from many *Bordetella* species to create a phylogenetic tree, Hamidou Soumana et al. (2017) demonstrated that eight out of the ten clades contained soil-dwelling bordetellae; the permeation of the environmental phenotype throughout the phylogenetic tree hints at a soil-based origin for the *Bordetella* genus.

Key evolutionary milestones within the genus involve species that are capable of both environmental and pathogenic lifestyles, such as the key ancestor to the classical *Bordetella* species, a *B. bronchiseptica*-like bacterium. Multilocus sequence typing (MLST) studies of *B. bronchiseptica* have established two distinct complexes of isolates, complex I and IV. The majority of strains isolated from humans originate from Complex IV (Diavatopoulos et al. 2005; Park et al. 2012). Using the much higher discriminatory power of next generation sequencing (NGS) data, it has been demonstrated that *B. pertussis* and *B. parapertussis* evolved from different complexes, with *B. parapertussis* sharing a more recent ancestor with Complex I and *B. pertussis* with Complex IV (Linz et al. 2016).

Table 1: Characteristics of all classical *Bordetella* closed genomes available on RefSeq, March 2019

Characteristic	<i>B. pertussis</i>	<i>B. parapertussis</i>	<i>B. bronchiseptica</i>
Number of closed genomes	421	18	19
Genome size / Mb <sup>1</sup>	4.11 (4.04 – 4.39)	4.78 (4.77 – 4.90)	5.21 (5.08 – 5.34)
Number of predicted genes <sup>1</sup>	3,979 (3,856 – 4,239)	4,501 (4,490 – 4,574)	4,911 (4,774 – 5,090)
Number of proteins <sup>1</sup>	3,615 (3,425 – 3,866)	4,166 (4,157 – 4,184)	4,804 (4,663 – 4,993)
G+C % <sup>1</sup>	67.7 (67.7 – 67.8)	68.1 (67.8 – 68.1)	68.2 (68.1-68.4)
IS 481 <sup>1</sup>	256 (234 - 273)	-	1 (0-3)
IS 1001 <sup>1</sup>	-	22 (22-28)	1 (0-1)
IS 1002 <sup>1</sup>	8 (5 – 10)	9 (0-9)	0 (0-5)
IS 1663 <sup>1</sup>	17 (16 – 24)	-	2 (0-14)

<sup>1</sup> Figures shown are mean and (range)

Despite their apparently different evolutionary trajectories, *B. pertussis* and *B. parapertussis* cause remarkably similar pathologies in humans. In an example of convergent evolution, the genomes of both species have evolved primarily through genome reduction, mediated through homologous recombination of insertion sequences (IS). As seen in **table 1**, the genome of *B. bronchiseptica* contains very few IS elements; in fact, the reference strain RB50 was originally believed to contain no IS elements at all (Parkhill et al. 2003; Preston et al. 2004). In contrast, *B. parapertussis* genomes contain around 30 IS elements, usually 22 copies of IS 1001 and 9 copies of IS 1002. The *B. pertussis* genome contains the most IS elements: up to ten copies of IS 1002, around 20 copies of IS 1663, and over 200 copies of IS 481. The appearance and expansion of these IS elements is thought to have led to the speciation of *B. pertussis* and, separately, *B. parapertussis*, from their *B. bronchiseptica*-like ancestors (Parkhill et al. 2003; Preston et al. 2004; Diavatopoulos et al. 2005).

Genome reduction was key in the speciation of *B. pertussis*. The significant, IS-mediated, reduction in genome size of the *B. pertussis* genome (around 4.1 Mb) compared to the *B. bronchiseptica* genome (around 5.3 Mb) has led to a streamlined genome, depleted of many metabolic, membrane transport, surface structure synthesis and gene expression regulatory genes when compared to *B. bronchiseptica* genomes (Parkhill et al. 2003). Comparative genomic studies between *B. bronchiseptica* and *B. pertussis* reveal that the latter has around 1200 fewer genes (Parkhill et al. 2003; Linz et al. 2016). In addition, insertions of IS elements into genes which are functional in *B. bronchiseptica* has resulted in the existence of over 350 pseudogenes in *B. pertussis*, compared to only around 20 in *B. bronchiseptica*. This sculpting of the *B. pertussis* genome via IS-mediated homologous recombination has produced a highly specialised pathogenic bacterium which is niche-restricted to the human nasopharynx. Traditionally, *B. pertussis* has been described as a monomorphic species (for example: Mooi 2010; Bart et al. 2010); however, since the advent of whole genome sequencing, genomics has been revealing that the bacterium may be less clonal than previously thought, with the introduction of vaccination and continued homologous recombination between IS elements driving gene- and genome-level variations respectively.

### 3. Vaccination has accelerated *B. pertussis* gene-level evolution

#### 3.1 Changes to circulating alleles

Over the last several decades, allele-typing of selected genes has shown a number of similar trends, characterised largely by the drift of genes away from the vaccine alleles (for example: Mooi et al. 1998; van der Zee et al. 1996; Mooi et al. 2001). One well-reviewed example involves *pertussis toxin*. Prior to the 1990s, the predominant *ptx* promoter allele was *ptxP1*. A new allele, *ptxP3*, was first observed in 1988 (Bart et al. 2010). In *ptxP3*, a SNP in the binding site for *ptx*'s transcriptional regulator, *BvgA*, appears to increase the binding affinity between the promoter and regulator, thus increasing transcription and causing *ptxP3*-carrying strains to produce more *ptx*. The expression of other proteins involved in complement resistance is also altered and, together, these changes increase the transmissibility and severity of the disease caused by *ptxP3*-carrying strains (Mooi et al. 2009; Bart et al. 2010; King et al. 2013; de Gouw et al. 2014). This new allele spread rapidly throughout the 1990s, and *ptxP3* is now present in greater than 90% of recent isolates (Lam et al. 2012; Bart et al. 2010). A thorough screen of 343 strains representing 19 countries and six continents, spanning 90 years of *B. pertussis* isolation, showed that similarly rapid selective sweeps have also occurred in other antigen-related genes, including *ptxA*, *prn*, and *fim3* (Bart et al. 2014a) .

In addition, analysis of 100 strains isolated during a 2012 whooping cough outbreak in the UK, after the introduction of the ACV, showed that the evolution of the antigens included in the ACV is occurring more rapidly than that of the surface proteins not included in the ACV (Sealey et al. 2015). Importantly, this analysis also showed that numerous different strains were circulating during the outbreak, rather than one particularly virulent strain or allelic profile being responsible. The same was also true for strains circulating during outbreaks in the USA, in California in 2010, and in Vermont and Washington in 2012 (Bowden et al. 2014; Bowden et al. 2016). This suggests that the strains circulating during outbreaks tend to be the same as those that circulate during non-outbreak periods, but that some unknown trigger causes an increase in whooping cough cases in regular four-yearly cycles.

Supporting the idea that the recent allelic changes we are seeing in *B. pertussis* are, in part, a response to the introduction of vaccination, Xu et al. (2015) and Du et al. (2016) showed that, in countries where vaccine uptake has been lower or delayed, the rate of change to the allelic profile of circulating strains has also been delayed. In the Philippines, for example, where the WCV is still in use, *prn2* has yet to appear, despite being the allele most frequently seen in ACV-adopting countries (Galit et al. 2015).

### **3.2 The proliferation of antigen-deficient strains**

Another gene-level phenomenon which has been observed recently is the emergence of strains deficient in one or more of the antigens used in the ACV. During the pre-ACV era, antigen-deficient strains were occasionally isolated, albeit at very low frequencies. Individual strains with mutations (deletion, frameshift, and premature stop codons) which resulted in the non-expression of pertactin were, for example, isolated in Europe, North America and Japan in the 1990s (Mastrantonio et al. 1999; Weigand et al. 2018; Miyaji et al. 2013). The landmark study by Bart et al. (2014a), in which all but around 20 strains were isolated prior to 2007, did not identify any strains which were *prn*-deficient (although one, BP310, has subsequently been resequenced by Zomer et al. (2018) and is likely to be deficient). Since the mid-2000s, however, the number of *prn*-deficient strains being isolated globally has increased rapidly. A study of Australian isolates from 2008-2012 showed an increase from 5% *prn*-deficient strains in 2008 to 78% *prn*-deficient strains in 2012 (Lam et al. 2014). Pertactin-deficiency appears to be polyclonal, affecting both dominant *prn* alleles, *prn1* and *prn2*, and arising through several different mechanisms including insertions of IS 481, large deletions, and SNPs, with no single predominant causative mutation (Hegerle et al. 2012; Queenan et al. 2013; Barkoff et al. 2019).

As with changes to allelic profiles, countries with different vaccination strategies appear to be differently affected by the proliferation of antigen-deficiency. A longitudinal study of *prn*-deficiency in European countries between 1998 and 2015, **figure 2**, showed that the number of *prn*-deficient strains is increasing in all screened countries but, the earlier a country introduced the ACV, the higher the percentage of strains currently found to be deficient (Barkoff et al. 2019). The rapid recent



increase in prn-deficiency suggests that, although the deficiency may have always occurred in some strains by chance, it has been strongly selected for by the ACV compared to the WCV. Further supporting the idea of ACV-mediated selection pressure is the sustained decrease of prn-deficient strains circulating in Japan since pertactin was removed from the Japanese ACV in 2012 (Hiramatsu et al. 2017).

A smaller number of strains deficient in other antigens included in the ACV have also been identified. Bart et al. (2015) and Weigand et al. (2018) identified several geographically independent recent strains which were unable to produce FHA; the same strains were often also prn-deficient. In addition, a handful of strains have been isolated globally which are deficient in both prn and ptx (Bouchez et al. 2009; Williams et al. 2016; Weigand et al. 2018). However, both FHA and Pertussis toxin are thought to play more vital roles in whooping cough disease development than pertactin (Carbonetti 2010; Henderson et al. 2012; Serra et al. 2011). Hence, although occasional strains may develop deficiency in these antigens, the reduced ability of these deficient strains to survive in the host and cause disease

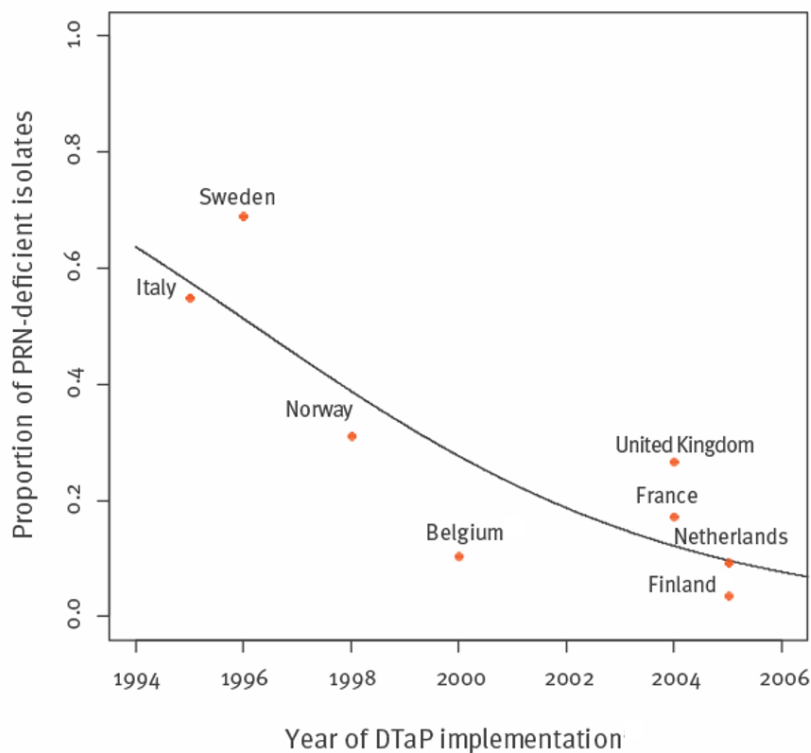


Figure 2: Correlation between the introduction of a primary acellular pertussis vaccine containing pertactin (PRN) in a European country and the proportion of PRN-deficient isolates found in the study, 2012–2015. Reproduced with permission from Barkoff et al. (2019)

and may mean that this kind of antigen deficiency is unlikely to proliferate in the same way as prn-deficiency.

## **4. Recent sequencing advances highlight genome-level variation**

### **4.1 The limitations of short-read sequencing**

Whilst the wide availability of whole genome sequencing throughout the 2000s enabled a variety of high-throughput strain screens, the highly repetitive nature of the *B. pertussis* genome has made the assembly of single-contig *B. pertussis* genome sequences difficult. Genome sequencing requires an organism's genome to be sheared into smaller fragments prior to sequencing; one of the goals of genomics is to fully re-assemble these fragments back into their original order. The goal when sequencing an organism with a single chromosome, like *B. pertussis*, would therefore be to re-assemble the sequenced fragments into a single, closed genome sequence. IS 481, however, together with the smaller number of copies of other repeated regions, such as IS 1002, IS 1663 and the rRNA operon, has confounded attempts to assemble closed *B. pertussis* genomes using short-read sequencing technologies, which sequence fragments shorter than the repeated section. The hundreds of *B. pertussis* genomes assembled using short-read sequencing alone have tended to consist of several hundred fragments, or "contigs": ostensibly one per repeat in the genome. Thus, the majority of high-throughput screens throughout the 2000s were focussed on the gene-level differences between strains already discussed, rather than comparisons on a whole genome-level.

The presence of so many IS elements, however, means that assembly of closed *B. pertussis* genomes could be particularly informative: IS elements are able to move around the genome through homologous recombination, potentially causing genome-level structural changes which may be discernible only through single-contig assemblies (Bentley et al. 2008; Siguier et al. 2014). Despite the discovery that most whooping cough outbreaks tend to be polyclonal in nature, *B. pertussis* remains a relatively clonal species, with a low SNP rate compared to many other bacteria. In other species, in addition to gene-level variations, differences at a whole-genome level are known to contribute to

altered gene expression and phenotypic diversity (Darch et al. 2014; Sousa et al. 1997). IS-mediated rearrangement may affect gene regulation and/or expression in *B. pertussis* by a number of mechanisms, including IS 481's inwards and outwards-facing promoters, as well as changing the distance of genes from the origin of replication (Amman et al. 2018). Limited evidence has already shown that certain genome-level differences can affect phenotype in *B. pertussis* in this way (Brinig et al. 2006).

The speciation of *B. pertussis* from *B. bronchiseptica* via IS element-mediated homologous recombination resulted in a variety of genomic arrangement differences between the two species alongside the reduction in genome size, and it is likely that IS-mediated genomic rearrangement in *B. pertussis* is an ongoing process (Parkhill et al. 2003). Indeed, prior to the advent of long-read sequencing, pulsed-field gel electrophoresis (PFGE) was one of the few methods able to discriminate between highly clonal *B. pertussis* strains: isolate screens using PFGE indicated that strains which seemed otherwise alike could vary significantly in terms of PFGE type (van Gent et al. 2015; Bisgard et al. 2001; Advani et al. 2004; Advani et al. 2013). Although the existence of numerous PFGE types was widely seen, however, it could not be confirmed how different PFGE types arose; they could represent different genomic arrangements, but could also have arisen due to mutagenesis at PFGE restriction sites, for example.

Thus, closed *B. pertussis* genome sequences may validate and further reveal genome-level differences between strains which otherwise appear to have highly similar or identical DNA content. The recent availability of long-read sequencing techniques, which can produce sequencing reads longer than 1,000 bp, has therefore revolutionised our ability to discover and investigate genome-level variations in *B. pertussis*.

#### **4.2 Long-read sequencing shows extensive inter-strain genome rearrangement**

The first study to take advantage of long reads utilised Pacific Biosciences (PacBio) sequencing to produce closed, fully annotated, genomes for two *B. pertussis* strains: BP1917 and BP1920 (Bart et al.

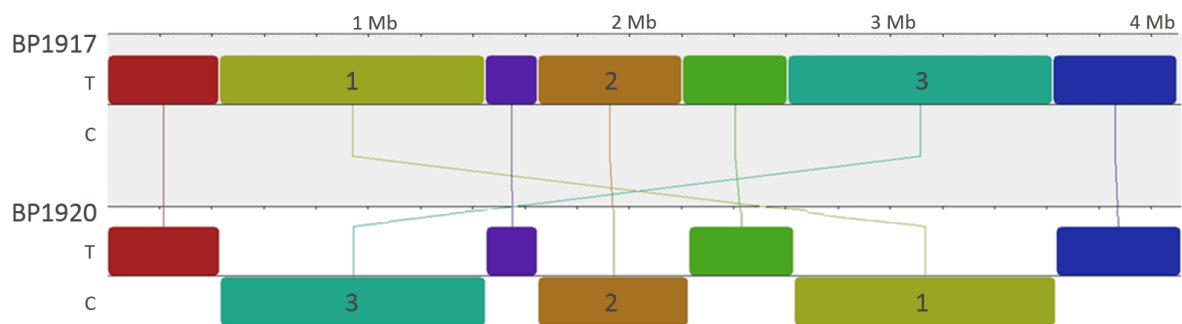


Figure 3: Different arrangements of the *B. pertussis* genome seen in BP1917 and BP1920. Three large sections of the genome (1, 2, and 3) are inverted in BP1920, as shown by their appearance on the complementary strand when the two whole genome sequences are aligned using progressiveMauve (Darling et al. 2010). The Genbank accession numbers for the genome sequences produced by Bart et al. (2014b) for BP1917 and BP1920 are CP009751 and CP009752, respectively.

2014b). The arrangement of these two strains differed significantly, with three large inversions and a variety of deletion and/or insertion events between the pair, **figure 3**. Having proven the ability of long reads to close the genomes of BP1917 and BP1920, Bart et al. (2015) next sequenced 11 *B. pertussis* strains which represented the pandemic *ptxP3* lineage, again using PacBio sequencing to produce 10 kb-long reads. This cohort, which also included several strains deficient in *prn* and/or *FHA*, were characteristically similar in terms of SNPs but again showed significant differences in genome arrangement. As is common for a developing technology, the cost of PacBio sequencing has rapidly decreased. Thus, higher-throughput strain screens have become increasingly feasible. **Figure 4** shows the dramatic increase in closed genome sequences for the classical *Bordetella* species available from the NCBI's RefSeq database since 2014. Bowden et al. (2016) conducted the first whooping cough outbreak screen using long-read sequencing alongside short-read sequencing in hybrid, sequencing 31 strains which had circulated during US whooping cough outbreaks in 2010 and 2012. The hybrid approach has been shown to improve the accuracy of assemblies produced using long-read sequencing, which still have an intrinsically higher error rate than short-read-only assemblies, particularly in homopolymeric tracts (Au et al. 2012; Koren et al. 2012). In the 31 genomes studied, 21 different arrangement profiles were observed; most consisted of inversions around the origin of replication. Bowden et al. also validated the arrangements using whole genome optical mapping and found that, in all cases, the boundaries between rearranged sections were composed of a repeated element: an insertion sequence, or the rRNA operon. The vast majority of the boundaries were IS 481

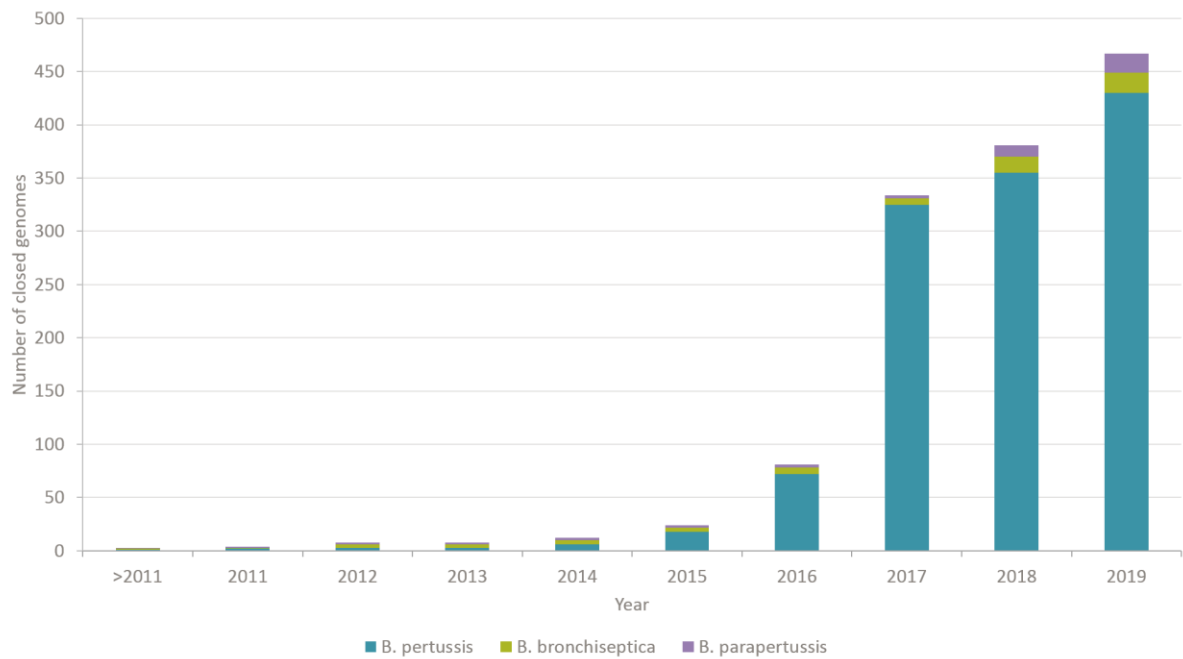


Figure 4: Increase in numbers of closed classical Bordetellae genome sequences (available on RefSeq) since the commercial introduction of long-read sequencing technologies

(89%), whilst the rest were composed of an rRNA operon, IS *1002* or a combination of IS *1002* and IS *481* together.

The most thorough investigation of *B. pertussis* genomic rearrangement to date also used a hybrid assembly strategy, combining PacBio long reads with Illumina short reads to close the genomes of 257 strains, dating from 1939 to 2014 (Weigand et al. 2017). When clustered based on their arrangement profiles, most isolates clustered according to allelic profile; for example, most *ptxP1* strains shared similar arrangements with other *ptxP1* strains. This clustering indicates that most structures are relatively stable, as supported by a clinical isolate which showed the same structure before and after 11 serial passages. Furthermore, these findings suggest that lineages are conserved not just in terms of SNPs, but also in genomic arrangement. Interestingly, Weigand et al. note that, on average, only half of their predicted IS *481* target sites are occupied in any given genome, suggesting a potential for further IS-mediated structural changes in future generations, assuming these sites are not non-permissive.

## **5. How else might *B. pertussis* generate diversity through genome-level variation?**

The primary metric commonly used to assess diversity of bacterial species is SNPs. However, *B. pertussis* is a textbook example of a clonal bacterial pathogen: variation, when judged by SNPs alone, is extremely limited, even taking into account the accelerated mutation of *B. pertussis* genes since the introduction of vaccination. Bart et al. (2010) estimated the mutation rate between *B. pertussis* isolates to be 1 SNP per 8,675 bases, compared to, for example, 1 SNP per 3,000 bases in *Mycobacterium tuberculosis*, and 1 SNP per 6,700 bases in *Escherichia coli* O157:H7 (Fleischmann et al. 2002; Gutacker et al. 2002; Zhang et al. 2006). Diversity within a species is vital for its survival, in order to drive adaptation; this is particularly true for pathogens, which are under pressure from the immune system (Mooi 2010). Therefore, a prominent question in *B. pertussis* genomics is: despite limited SNP diversity, how does *B. pertussis* generate diversity? The large numbers of closed genomes assembled using long-read sequencing have proven that rearrangements are a rich source of genome-level diversity, but can genomics also reveal other types of genome-level variation?

### **5.1 Harnessing deletion as a driver of diversity**

King et al. (2010) analysed the size of *B. pertussis* genomes of strains isolated over a 60-year period, demonstrating that genome streamlining has been an ongoing process, with recently isolated strains having smaller genomes and higher numbers of pseudogenes. Thus, *B. pertussis* is described as a species which is still undergoing genome reduction. Like genomic rearrangement, reduction is driven primarily by homologous recombination between insertion sequences. The large numbers of homologous IS elements in *B. pertussis* therefore produce a fertile mutational landscape capable of the generation of diversity.

Many bacterial species also generate diversity through the gain of genes, often by horizontal gene transfer (HGT), resulting in fluid gene content of the population, enabling the population to effectively respond to evolutionary bottlenecks that may arise over long or short timescales. In *B. pertussis*,

however, HGT appears to occur very rarely (Linz et al. 2016). Gene content of a species or genus is often analysed using a “pangenome” approach, which consists of analysing which genes are consistently present (the core genome) in the population, and which genes are variably present (the accessory genome) (reviewed in Medini et al. 2005; Rouli et al. 2015). A number of studies have undertaken this analysis in *B. pertussis* using either comparative genomic hybridization (CGH) (for example: Zhang et al. 2006; Caro et al. 2006; Heikkinen et al. 2007; King et al. 2008) or NGS (for example: Park et al. 2012; Ding et al. 2017). These have shown that, despite extremely limited HGT and otherwise high levels of clonality between strains, *B. pertussis* maintains a moderate accessory genome, largely through gene loss rather than gene gain. For example, the most comprehensive pangenome study to date, using CGH on 171 *B. pertussis* strains, revealed that 15% of the genes present in the population appeared variably in the 171 strains studied. (King et al. 2010). By using a set of probes which included *B. bronchiseptica* and *B. parapertussis*, King et al. were able to avoid biasing their analysis towards genes that were present only in the *B. pertussis* reference strain, Tohama I, which has been shown to lack over 45 kb of the accessory genome of the population (Caro et al. 2008; Bouchez et al. 2008).

There is a lack of knowledge about the phenotypic impact of gene deletions in *B. pertussis*, however. As the cost of sequencing has plummeted, the frequency and ease with which genomes, and their constituent mutations, are published has far outpaced the publishing of their phenotypic impact. To cope with the deluge of data, ontology schemes strive to categorize genes into functional groups and estimate their function based on sequence homology. A variety of nucleotide polymorphisms in *B. pertussis*, such as those in *ptxP3*, have had their phenotypic impacts analysed, with some providing clear fitness advantages in the mouse model (Mooi et al. 2009). In contrast, many key gene deletions have yet to be experimentally characterized in *B. pertussis*. Using the ontology scheme Clusters of Orthologous Genes (COG), King et al. (2010) showed that, as expected, housekeeping genes were underrepresented in the deleted genes, whilst genes of unknown function were overrepresented by

25%. There therefore remains genetic “dark matter”, genes with unknown function which are overrepresented in gene deletions.

Whilst *B. pertussis* is undisputedly undergoing genome reduction, evolution acts on phenotypes rather than genotypes. Therefore, it is likely that the *B. pertussis* genome is undergoing streamlining as an effect of certain phenotypes being selected against. It has been theorised that the transcriptional and translational cost of superfluous genes far outstrips the mere cost of DNA replication of such regions (Adler et al. 2014). *B. pertussis* maintains many seemingly functionless pseudogenes, despite the vast majority being shown to be transcriptionally inactive *in vitro* and in the mouse model (Bart et al. 2010; King et al. 2008; de Gouw et al. 2014). This supports the idea that the deletion of some genes provides a greater fitness benefit than the deletion of some others. Nonetheless, there is also evidence that the DNA content of the species is under selection, as pseudogenes have been shown to be mildly enriched in gene deletions, suggesting that streamlining of the DNA is also favoured to some extent (Kuo and Ochman 2010). Thus, the process of *B. pertussis* genome streamlining is likely to be a balance between entirely passive and entirely directed.

There have been five deleted regions, totalling over 50 genes, that have been deleted in all recently isolated clinical strains in comparison to the reference strain Tohama I (King et al. 2010; Heikkinen et al. 2007; Bouchez et al. 2008). In addition to clinical strains, Bart et al (2014a) also investigated two strains that were used to make WCVs. In one of the vaccine strains, the five deleted regions were present; if these regions impact cell surface antigens, the immunity conveyed by the WCV could therefore also be affected. This highlights the clinical importance of understanding the continual evolution of *B. pertussis*, which is in part driven by genome reduction.

## **5.2 Harnessing duplication as a driver of diversity**

Homologous recombination between IS elements has not only caused rearrangements and deletions in the *B. pertussis* genome, but also duplications ranging from single genes to large, multi-gene, regions. The general paradigm under which duplications occur is that a gene is duplicated, thus freeing



the second copy from the purifying selection of the original copy of the gene, potentially allowing it over time to evolve a new function. However, the second copy of the gene may also maintain the same function of the first gene. These types of events are “canonical” duplications that are well documented in the bacterial kingdom (for example: Ohta 1989; Lynch 2002; Magadum et al. 2013). Taking the genes from Tohama I and clustering them based on 90% nucleotide homology using the tool CDHIT (available as a web server: [http://weizhong-lab.ucsd.edu/cdhit\\_suite/cgi-bin/index.cgi?cmd=cd-hit-est](http://weizhong-lab.ucsd.edu/cdhit_suite/cgi-bin/index.cgi?cmd=cd-hit-est)), it can be seen that Tohama I maintains two copies of nine separate genes ranging from 97% to 100% homology (excluding IS elements and rRNA genes) The maintenance of these duplications provides further evidence that it is not the genome size of the bacterium itself that is the primary target of streamlining but certain phenotypic traits which are coded in the DNA.

Before the cost of NGS rapidly decreased in the late 2000s and early 2010s, duplications were primarily inferred by increased spot intensity in CGH or disturbances to southern blotting or PFGE patterns. Using these techniques, a number of multi-gene duplications were serendipitously discovered in the *B. pertussis* population. Using the power of long-read sequencing technologies, a number of studies had revealed further duplications, bringing the total to 13 serendipitously discovered mutations (Dalet et al. 2004; Caro et al. 2006; Heikkinen et al. 2007; Weigand et al. 2016; Dienstbier et al. 2018; Ring et al. 2018; Weigand et al. 2018). A recent study by Abrahams et al. (In preparation) systematically analysed the *B. pertussis* population in search of large multi-gene duplications. Previously published short-read sequencing data were utilised and read depth abnormalities were used to predict duplications. In the 473 strains analysed, over 400 duplications were found.

Abrahams et al. (In preparation) presents a deep description of duplications in *B. pertussis*. In addition to the quantity of duplications, over 90% of duplications were found at 11 “hotspot” loci but with varying gene contents, similar to a situation described previously in *M. tuberculosis* (Weiner et al. 2012). Interestingly, when the CNVs at each hotspot loci were mapped to a phylogenetic tree based on core genome SNPs, they appeared not to be vertically inherited and instead appeared to occur

spontaneously many times at similar loci with subtly different gene content in each mutation, suggesting the existence of a potential phenotypic driver at those loci.

Large multi-gene duplications are known to be unstable in the bacterial kingdom, and this has also been demonstrated in *B. pertussis*. For instance, Dalet et al. (2004) noticed that subculturing a single isolate produced both high haemolytic and average haemolytic single colonies. Further analysis showed that colonies with high haemolysis had a duplication of the locus encoding adenylate cyclase, a key virulence factor. Dalet et al. further demonstrated that subculturing a strain with a duplication produced colonies with a single copy of the locus, thus indicating a mixed population, ostensibly caused by an unstable locus. This early study used PFGE and southern blotting to screen colonies for copy number of the locus. These “pre-genomics” tools provide high quality data, but largely answer very specific research hypotheses, in contrast to sequencing experiments, which shed light on a vast range of research questions. Abrahams et al. used ultra-long nanopore sequencing reads (over 3,000 reads longer than 50 kb) to confirm the presence of between 1 and 5 copies of a single locus within an otherwise clonal population. This tentatively supports the findings of Dalet et al., showing that in a single sample there exists a variety of genetic configurations of a single locus. This study demonstrates the potential of long-read sequencing to not only confirm long-predicted genomic structural variations in *B. pertussis*, but also to play a key role in the discovery and further investigation of a variety of entirely unpredicted genomic phenomena. The next steps in understanding these new genomic phenomena in *B. pertussis* should aim to elucidate the existence and extent of any phenotypic effects stemming from large duplicated regions, as well as any contribution they make to whooping cough virulence.

## **6. What is the future for *B. pertussis* genomic research?**

Changes to the allelic profile of circulating *B. pertussis* strains have been recorded for many decades, in the pre-genomics era and beyond. The wide-spread availability of whole genome sequencing since the early 2000s, though, has enabled the screening of larger numbers of strains isolated over the last

hundred years, thus allowing us to understand, longitudinally, the extent to which *B. pertussis* has been evolving on the gene-level. As seen above, there is evidence that many of these gene-level changes, in terms of both allelic profile and antigenic deficiency, have been influenced by the introduction of first the WCV and then the ACV. Since the 2010s, long-read sequencing has allowed us to investigate *B. pertussis* on a new level, that of the whole genome. The existence of a wide variety of inter-strain genomic rearrangements is now well-established, and more recent evidence has begun to show that other types of genome-level differences, such as large tandem duplications, also exist. However, the contribution of these observed gene- and genome-level variations to observed phenotypes is yet to be fully understood. In addition to informing our understanding of the continued evolution of the *B. pertussis* genome, a more thorough understanding of *B. pertussis* genomics could also contribute significantly to the future of whooping cough prevention strategies (Cherry 2019).

Long-read sequencing will have a major part to play in any future investigation of *B. pertussis* genomic variations. Until recently, high-throughput long-read sequencing was restricted to larger laboratories which could afford a PacBio sequencing system, thus all the early long-read studies described here took place at national health laboratories: the CDC in the US, and the Centre for Infectious Diseases Control in the Netherlands. Oxford Nanopore Technologies (ONT) sequencing may provide a more accessible alternative for smaller laboratories and, indeed, two studies from late 2018 have shown the feasibility of assembling multiple closed *B. pertussis* genomes using ONT sequencing in hybrid with Illumina sequencing (Ring et al. 2018; Bouchez et al. 2018). In addition, ultra-long ONT sequencing has recently revealed yet further *B. pertussis* structural complexity, in the form of highly mixed populations (Abrahams et al. In preparation). Thus, it is likely that future studies of *B. pertussis* genome structure will utilise both PacBio and ONT sequencing. For investigating the most complicated structural features, such as very long tandem duplications, there will likely be a preference for ONT sequencing, as there is theoretically no upper limit to the length of sequencing read which could be produced by nanopore sequencing (Schmid et al. 2018). It is also likely that any studies utilising long reads to investigate *B. pertussis* will use them in hybrid with a more accurate short-read technology,

although improvements to both long-read technologies mean that highly accurate long-read-only assemblies are on the horizon, which could enable both base-level and genome-level interrogations using a single technology (Wenger et al. 2019; Oxford Nanopore Technologies 2018).

Alongside sequencing, there will still remain a place for other genomics tools, such as PFGE and optical mapping. The most recent survey of *B. pertussis* genomic diversity in the US, by Weigand et al. (2019), demonstrates the potential for such a wholistic approach. Using a combination of short-read sequencing, long-read sequencing, multilocus variable-number tandem-repeat analysis (MLVA), PFGE and optical mapping, Weigand et al. were able in a single study to characterise the gene- and genome-level profiles, including allelic-profile, antigen deficiency, genome arrangement and the existence of several large tandem duplications, in 170 strains isolated between 2000 and 2013. Such detailed analyses will likely provide a springboard for future studies, for both the continued surveillance of *B. pertussis* evolution, and the investigation of any correlation between genotypic, genomic and phenotypic differences.

## **7. Concluding remarks**

In summary, genomics has shown, and continues to show, that *B. pertussis* is not necessarily the entirely monomorphic species it is traditionally believed to be. Although the allelic profile of *B. pertussis* changed in response to the introduction of the WCV, and more rapidly since the switch to the ACV, diversity at the gene-level remains very limited when compared to many other bacteria. However, the wide availability of WGS, and particularly the more recent long-read sequencing technologies, have revealed dynamic and substantial genome-level variations, both between and within strains. Future work may utilise a wholistic approach, focussing on the further elucidation and phenotypic characterisation of both gene- and genome-level phenomena together, ultimately informing our understanding of how diversity is generated in species with limited base-level inter-strain variation and, perhaps, the role this has played in the resurgence of whooping cough.

## Figure Legends

**Figure 1:** Reported incidence of whooping cough per 100,000 in the USA, 1990-2017, showing increase in all age brackets. Data source: CDC National notifiable diseases surveillance reports Centers for Disease Control (2019)

**Figure 2:** Correlation between the introduction of a primary acellular pertussis vaccine containing pertactin (PRN) in a European country and the proportion of PRN-deficient isolates found in the study, 2012–2015. Reproduced with permission from Barkoff et al. (2019)

**Figure 3:** Different arrangements of the *B. pertussis* genome seen in BP1917 and BP1920. Three large sections of the genome (1, 2, and 3) are inverted in BP1920, as shown by their appearance on the complementary strand when the two whole genome sequences are aligned using progressiveMauve (Darling et al. 2010). The Genbank accession numbers for BP1917 and BP1920 are CP009751 and CP009752, respectively.

**Figure 4:** Increase in numbers of closed classical *Bordetellae* genome sequences (available on RefSeq) since the commercial introduction of long-read sequencing technologies

**Table 1:** Characteristics of all classical *Bordetella* closed genomes available on RefSeq, March 2019

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