

# Analysis of clinical *Bordetella pertussis* isolates using whole genome sequences reveals novel genomic regions associated with recent outbreaks in the United States of America

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1 **Abstract**

2 **Background:** Despite high-levels of vaccination, whooping cough, primarily caused by *Bordetella*  
3 pertussis (BP), has persisted and resurged. It remains a major cause of infant death worldwide and  
4 is the most prevalent vaccine-preventable disease in developed countries. To date, most genomic  
5 studies have focused on a small subset of the BP genome, biasing our clinical understanding and  
6 public health awareness.

7  
8 **Methods:** We performed a Genome-Wide Association Study (GWAS) on 76 U.S. BP whole  
9 genomes, including strains from recent outbreaks.

10  
11 **Results:** A GWAS of the 76 BP isolates revealed a sharp increase in genetic variation associated  
12 with the Minnesota 2012 outbreak and identified 52 variants unique to the Minnesota outbreak  
13 and 19 unique to the California and Washington outbreaks. None of the identified variants were  
14 shared between the outbreaks and the vast majority were previously uncharacterized. We further  
15 identified variation associated with pertactin negative strains and acellular vaccination.

16  
17 **Conclusions:** We identified novel genomic regions associated with recent BP outbreaks. Our  
18 results underscore the need for increased whole genome sequencing of BP isolates, which can re-  
19 duce costly misdiagnosis and improve surveillance. The genes containing these variants warrant  
20 further investigation into their possible roles in BP pathogenicity and the ongoing resurgence in  
21 the U.S.

## 22 Introduction

23 The ongoing whooping cough epidemic – caused primarily by *Bordetella pertussis* – is rapidly be-  
24 coming a public health emergency in the United States of America (U.S.) [1]. The recent outbreaks  
25 in 2012 generated the highest number of whooping cough cases observed since the pre-vaccine  
26 era [2], and, in 2014, California had the most cases ever recorded in the state [3].

27 Clinicians rely on genetic-based tests to rapidly and accurately determine whether an individual  
28 is infected with *B. pertussis* and public health officials use genetic results to monitor outbreaks and  
29 the effectiveness of interventions [4, 5]. However, despite advances in next generation sequencing  
30 (NGS) technology and expanded specimen collection, the majority of *B. pertussis* genetic studies  
31 focus on detecting various genetic markers associated with only a handful of known virulence factors  
32 and vaccine components [6, 7]. This approach is unable to identify other genomic regions involved  
33 in *B. pertussis* pathogenicity and/or vaccine resistance potentially responsible for clonal expansion  
34 and persistence in a vaccinated population.

35 To investigate the clinical and public health consequences of using fixed regions of the *B. per-*  
36 *tussis* genome for diagnosis and reporting, we conducted a genome wide association study (GWAS)  
37 on *B. pertussis* genomes isolated during three recent U.S. outbreaks. We detected an increase in  
38 genetic variation in the Minnesota 2012 subpopulation and 31 non-synonymous single nucleotide  
39 polymorphisms (SNPs) not associated with other outbreaks. These findings indicate that it is  
40 possible to differentiate *B. pertussis* subpopulations underlying outbreaks and track the spread of  
41 the bacteria using genomic data. The same methods were used to identify SNPs associated with  
42 pertactin(-) strains and strains isolated during the acellular vaccine period (1998-2012). Import-  
43 tantly, the identified SNPs are not located in genes currently associated with known virulence or  
44 vaccine components and thus represent an opportunity for the design of novel therapeutics.

45 These findings broaden our understanding of the evolutionary forces acting on *B. pertussis*,  
46 the genetic loci undergoing selective pressure and how *B. pertussis* may be evolving in response to  
47 vaccine pressure.

## 48 Methods

### 49 Strains studied

50 We utilized genomic data from 76 *B. pertussis* strains isolated in the U.S. between 1935-2012.  
51 Thirty five strains were collected from various locations in the U.S. between 1935-2005 (referred to  
52 as the historical strains), 19 strains were collected from the 2010 California outbreak [8], 9 strains  
53 were collected from the 2012 Washington outbreak [9], 12 strains were collected from the 2012  
54 Minnesota outbreak [10], and *B. pertussis* strain 18323, isolated in the U.S. in 1946 and for which  
55 a detailed Genbank file is available, was the reference genome used for comparison. Except for the  
56 Minnesota outbreak genome sequences, all genome sequence data was downloaded from the NCBI  
57 Sequence Read Archive. NCBI Sequence Read Archive accession numbers for the genome sequences

58 are listed in eTable 1 in the Supplement. The Minnesota outbreak whole genome sequence data  
59 were obtained from the researchers who isolated and sequenced the Minnesota outbreak strains [10].

## 60 **Whole genome sequencing and phylogenetic analysis**

61 Short read sequences from each of the *B. pertussis* genomes underwent quality checks with FastQC,  
62 had adapters trimmed with Trimmomatic, were mapped to the *B. pertussis* reference genome 18323  
63 with bowtie2 and SNPs called using bcftools. Parsnp [11] was used to further filter the SNPs  
64 to produce 6079 core-genome SNPs, create a whole genome alignment and maximum likelihood  
65 phylogenetic tree of the 76 *B. pertussis* genomes. Given the very high support for the tree and all  
66 internal nodes, we do not discuss the results of a Bayesian phylogenetic analysis as the posterior  
67 distribution of trees was tightly peaked around the maximum likelihood tree.

## 68 **Genome wide association study**

69 In order to identify genomic regions significantly associated with the various outbreaks we conducted  
70 a GWAS on the 6079 core-genome SNPs following established best practices [12]. The 6079 SNPs  
71 from each genome were concatenated and an association between each SNP and the phenotype was  
72 tested by logistic regression implemented in R [12]. Each of the *B. pertussis* groups, e.g. Minnesota,  
73 California, Washington, and historical, were treated as a phenotype and the rest of the genomes used  
74 as controls. Due to their similar number of SNPs/genome and phylogenetic similarity we combined  
75 the California and Washington outbreak strains into a single phenotype while using the remaining  
76 genomes as controls. Strain 18323 was used as the reference genome and its Genbank file used to  
77 annotate the SNPs. Multiple testing was accounted for by applying a Bonferroni correction. The  
78 individual locus effect of a SNP was considered significant if its p-value was smaller than  $\alpha/n$ , with  
79  $\alpha = 0.05$  as the genome-wide false positive rate and  $n$  is the number of SNPs. The genome-wide  
80  $-\log_{10}P$  value value threshold was 5.0.

## 81 **Results**

### 82 ***B. pertussis* genome SNP frequencies in the US from 1935-2012**

83 To investigate possible SNP accumulation trends of *B. pertussis* genomes isolated in the U.S., we  
84 calculated the median number of SNPs/genome for each year we had *B. pertussis* whole genome  
85 sequence data. Figure 1 illustrates that the number of SNPs/genome fluctuated very little each  
86 year around a median of 1940 SNPs/genome until 2012 when there was a 22% increase to 2367  
87 SNPs/genome. Due to the large range of SNPs/genome values in 2012, we separated the strains from  
88 2012 into two groups, those isolated in the Washington outbreak and those isolated in the Minnesota  
89 outbreak. We also grouped the remaining genomes isolated between 1935-2005 by decade.

90 Separating the 2012 strains geographically revealed that the Minnesota strains were the source  
91 of the increase in 2012 SNP counts with a median 2408 SNPs/genome, 24% more SNPs/genome

92 than the 1943 SNPs/genome median of all other periods, including those from the California 2010  
93 (1942 SNPs/genome) and Washington 2011 & 2012 (1947 SNPs/genome) isolates (Figure 2). It is  
94 unlikely the increased SNPs/genome in the Minnesota strains can be attributed to imbalances in  
95 genome coverage that favor finding more SNPs in the Minnesota genomes as the depth of coverage  
96 is 49x for the Minnesota genomes, 90x for the California and Washington genomes and 60x for the  
97 historical genomes. Likewise, the breadth of coverage is 90% for the Minnesota genomes, 88% for  
98 the California and Washington genomes and 88% for the historical genomes.

### 99 **Phylogenetic analysis of *B. pertussis* strains**

100 Greater than 99% of each genome was included in the whole genome alignment that was used to  
101 generate the phylogenetic tree in Figure 3. The genomes cluster roughly in chronological order  
102 with a few exceptions. Twenty four of the 28 California and Washington strains cluster together.  
103 Three of the four California strains are in a clade that contains the 18323 reference genome that  
104 was isolated in 1946. One of the three California strains that clusters with 18323 differs by only  
105 seven SNPs, the other two each by 22 SNPs. This cluster of three California strains with 18323 is  
106 evidence that a subpopulation of *B. pertussis* very similar to strains present 70 years ago are still  
107 in circulation and causing disease.

108 The 12 strains isolated in the Minnesota outbreak form a distinct clade with two distinct  
109 subclusters. One subcluster contains nine genomes that lack a pertactin (*prn1*) signal sequence  
110 and the other cluster contains three genomes in which pertactin (*prn2*) is disrupted by insertion  
111 sequence IS481 or contains a stop codon. The majority of the pertactin(-) strains studied cluster  
112 around the Minnesota clade except for three pertactin(-) strains from the California outbreak found  
113 in the upper half of the tree. This suggests that pertactin(-) strains have arisen independently  
114 multiple times in the U.S., evidence for which has been previously reported [13]. Two of these  
115 pertactin(-) strains cluster with 18323, indicating that *B. pertussis* strains very similar to those  
116 present 70 years ago are still in circulation and have become pertactin(-). The phylogenetic tree  
117 in Figure 3 illustrates the uniqueness of the Minnesota subpopulation and the 18323-associated  
118 cluster of CA strains compared to the rest of the genomes studied.

### 119 **Polymorphisms characteristic of three recent U.S. outbreaks of *B. pertussis***

120 Out of 6079 SNPs from the 76 *B. pertussis* strains we identified 52 SNPs in coding regions that  
121 were significantly associated ( $-\log_{10}P$  value  $\geq 5$ ) with the Minnesota outbreak. Of the 52 SNPs,  
122 the 31 non-synonymous or nonsense SNPs characteristic of the Minnesota subpopulation are listed  
123 in Table 1.

124 The Washington outbreak GWAS did not identify any significantly associated SNPs. However,  
125 due to their similar number of SNPs/genome and phylogenetic similarity, we combined the Califor-  
126 nia and Washington outbreak strains into a single phenotype while using the remaining genomes  
127 as controls in a GWAS. From 6079 SNPs we identified 19 SNPs in coding regions that were signifi-  
128 cantly associated ( $-\log_{10}P$  value  $\geq 5$ ) with the combined California/Washington outbreak strains.

129 The 19 SNPs are listed in Table 2. The California/Washington SNPs overlapped with nine SNPs of  
130 the historical strains suggesting that some of the California/Washington strains are more closely re-  
131 lated to the historical strains. This is supported by the way these strains cluster in the phylogenetic  
132 tree.

133 None of the significant Minnesota SNPs overlapped with the significant California/Washington  
134 SNPs or the historical SNPs. However, the *tsr* and *BN118\_0965* genes were found to have SNPs in  
135 both Minnesota and California/Washington populations, though not at the same codon positions.  
136 According to the annotations in the 18323 reference genome Genbank file, *BN118\_0965* encodes a  
137 putative permease component of the branched-chain amino acid transport system and *tsr* encodes  
138 a methyl-accepting chemotaxis protein I.

### 139 **Polymorphisms characteristic of pertactin(-) strains**

140 When all pertactin(-) strains were tested as a phenotype we identified 14 significantly associated  
141 ( $-\log_{10}P$  value  $\geq 5$ ) SNPs in coding regions (Table 3). Twelve of the SNPs were associated with  
142 Minnesota strains and two of the SNPs were associated with the California and Washington strains.  
143 Six of the 12 SNPs were non-synonymous.

### 144 **Polymorphisms characteristic of the pre-, whole cell and acellular vaccine periods**

145 When all strains isolated during the acellular vaccine period (1998-2012, 62 isolates) were tested as a  
146 phenotype we identified a single significantly associated ( $-\log_{10}P$  value  $\geq 5$ ) non-synonymous SNP  
147 in the *BN118\_0189* gene, which encodes for a restriction endonuclease. This SNP is also associated  
148 with the pertactin(-) phenotype, specifically the California/Washington pertactin(-) strains. No  
149 significant SNPs were identified from the pre-vaccine ( $< 1946$ , 3 isolates) and whole cell vaccine  
150 (1946-1997, 10 isolates) phenotypes.

## 151 **Discussion**

152 *Bordetella pertussis* produces multiple toxins in addition to other virulence factors that facilitate  
153 within-host survival by manipulating many aspects of the human immune system [14]. The ma-  
154 jority of comparative genomics studies of *B. pertussis* focus exclusively on these known toxins and  
155 virulence factors. Our study was motivated by the importance of understanding all genes undergo-  
156 ing selective pressure, not just known virulence genes. Our results did detect SNPs in known toxin  
157 and virulence genes, however the putative non-virulence SNPs we describe here are associated with  
158 outbreak isolates at a much higher level of statistical significance and therefore merit reporting.  
159 More than half of the SNPs reside in genes whose protein products are not fully characterized and  
160 whose roles in pathogenesis are unknown. Some are likely housekeeping genes without a role in  
161 pathogenesis, but others, like *BN118\_0462* that is associated with lipopolysaccharide assembly, are  
162 almost certainly involved in pathogenesis.

163 Multiple studies indicate that *B. pertussis* relies on iron uptake systems for growth in vivo [15].  
164 In *B. pertussis*, outer membrane receptors are required for transfer of iron chelates to the periplasm,  
165 followed by transport to the cytoplasm by ATP-binding cassette (ABC) transporters [15]. Bacterial  
166 ABC transporters are an important class of transmembrane transporters known to influence many  
167 cellular processes including antibiotic resistance, nutrient acquisition, adhesion, protein secretion  
168 and have been shown to be important for the virulence of a range of bacterial pathogens [16].  
169 The GWAS of combined California/Washington isolates identified SNPs in both an iron uptake  
170 receptor gene, *bfrI* [17], and ABC transporter genes *BN118\_2154*, *BN118\_0963*, *BN118\_0964* and  
171 *BN118\_0965*. The Minnesota GWAS also identified a gene with putative iron transport func-  
172 tionality, *BN118\_0794*, as well as multiple ABC transporter genes: *BN118\_0138*, *BN118\_2781*,  
173 *BN118\_2428*, and *BN118\_0965*. *BN118\_0964* and *BN118\_0965* are both involved in branch chain  
174 amino acid transport and *BN118\_0965* had SNPs in both California/Washington and Minnesota  
175 populations, though not at the same codon positions. Branch chain amino acid transporters have  
176 been found essential for disease pathogenesis in some bacteria [18].

177 Another gene that contained SNPs detected in both Minnesota and California/Washington pop-  
178 ulations, but at different codon positions, was the *tsr* gene, which encodes a chemotaxis protein that  
179 recognizes the amino acid serine [19]. A total of six SNPs in the Minnesota strains were detected  
180 in the *maeB* gene that encodes NADP-dependent malic enzyme, a protein involved in pyruvate  
181 metabolism and carbon fixation [20]. Multiple SNPs in the *maeB* gene have been previously re-  
182 ported and postulated to be important in *B. pertussis* adaptation [21].

183 In summary, GWAS of the outbreak phenotypes identified several SNPs in genes that encode  
184 iron uptake, transporter, metabolism and chemotaxis proteins associated with in-host survivabil-  
185 ity and virulence that overlapped outbreaks, and other genes that were specific to a particular  
186 outbreak. Compared to 18323, the elevated SNPs/genome of the Minnesota outbreak strains  
187 (2408 SNPs/genome) and the nearly identical CA\_2010\_3- (7 SNPs), CA\_2010\_5- (22 SNPs) and  
188 CA\_2010\_4 (22 SNPs) strains suggest that subpopulations of *B. pertussis* are undergoing differential  
189 selective pressure.

190 Pertactin(-) *B. pertussis* strains have been increasingly reported in the U.S. and several different  
191 genotypes have been attributed to the pertactin(-) phenotype [13]. The pertactin(-) phenotypes  
192 in the strains we studied also included multiple genotypes. The GWAS of the pertactin(-) pheno-  
193 type identified non-synonymous SNPs in the *maeB* and *tsr* genes of the Minnesota strains and in  
194 the *BN118\_2561* and *BN118\_0189* genes of the California/Washington strains. The *BN118\_2561*  
195 and *BN118\_0189* genes encode putative restriction endonucleases. Considering the multiple geno-  
196 types for the pertactin(-) phenotype, and the dissimilarity between *maeB*, *tsr*, *BN118\_2561* and  
197 *BN118\_0189* genes, these are likely confounding SNPs and not correlated or compensatory to the  
198 pertactin(-) phenotype in any way. While these SNPs may be confounding SNPs associated with  
199 the pertactin(-) phenotype, it is possible that since all the pertactin(-) strains were isolated during  
200 the acellular vaccine period that the SNPs are involved with acellular vaccine evasion and adap-  
201 tation. In fact, the SNP in the *BN118\_0189* gene was found to be significantly associated with



202 the acellular vaccine period (1998-2012). No significant SNPs were identified from the pre-vaccine  
203 (<1946, 3 isolates) and whole cell vaccine (1946-1997, 10 isolates) phenotypes. While this may be  
204 due to not having enough samples for each phenotype, it is possible that the whole cell vaccine  
205 successfully blocked *B. pertussis* transmission to such an extent that evolutionary opportunities  
206 were not available for the pathogen to adapt and establish consensus strains able to evade the  
207 vaccine [22]. Analyzing additional samples should strengthen these results.

208 The Minnesota outbreak is alarming for several reasons. The strains isolated had higher  
209 SNPs/genome counts than all other strains, all isolates were pertactin(-), and *B. parapertussis*  
210 was also isolated (although not sequenced) [10]. The same region in Minnesota also experienced  
211 a *B. parapertussis* outbreak in 2014 [23]. This may signal the beginning of a “new normal” for *B.*  
212 *pertussis* and *B. parapertussis* resurgence. There are a handful of *B. parapertussis* genome sequences  
213 publicly available but more are needed for a useful GWAS and comparison to *B. pertussis*.

214 In a time when whole genome sequencing is becoming rapidly cheaper, the current *B. pertussis*  
215 surveillance, diagnostic and molecular typing techniques are found severely lacking. The CDC  
216 does not publish any PCR assay protocols or standards for pertussis detection, multilocus sequence  
217 typing (MLST) and multilocus variable number tandem repeat analysis (MLVA) assays only address  
218 a tiny fraction of the genome and pulsed-field gel electrophoresis (PFGE) has been found to be  
219 more discerning than MLST and MLVA combined [9]. In fact, PFGE typing suggests that pertussis  
220 isolates are evolving more rapidly on a genomic scale than in the few genes and repeat regions  
221 targeted by MLVA and MLST [9]. The GWAS we performed is precisely one way to uncover the  
222 strain diversity residing in the rest of the pertussis genome missed by MLST and MLVA assays  
223 and make more definitive conclusions about correlation. As such, it has become obvious to us  
224 that whole genome sequencing technology provides the means with which to replace the lack of  
225 a unified pertussis typing scheme and an opportunity to create a universal format for recording  
226 and reporting the molecular epidemiology of *B. pertussis* isolates that allows for more efficient  
227 comparisons between epidemics and countries.

228 To our knowledge, the findings reported here comprise the first GWAS, SNP count analysis and  
229 phylogenetic tree construction using whole genome *B. pertussis* sequences from strains isolated in  
230 the U.S. We are confident that additional GWAS will produce significant insights into *B. pertussis*  
231 resurgence and play an important role in surveillance diagnostics and vaccine design when applied  
232 to whole genome sequences of clinical samples in the future.

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320 **Figures**

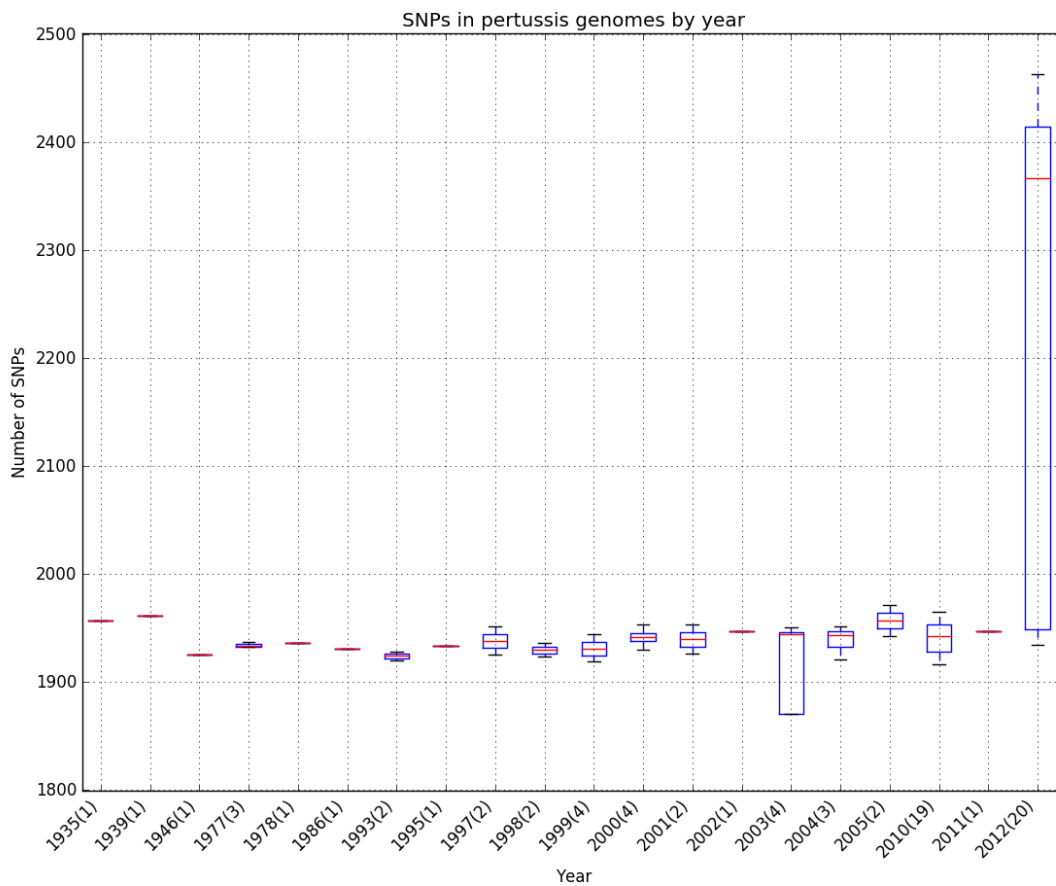


Figure 1: **Median SNPs/genome in all *B. pertussis* strains** Boxplots show median SNPs/genome studied grouped by year. Numbers in parentheses indicate number of genomes for the given year.

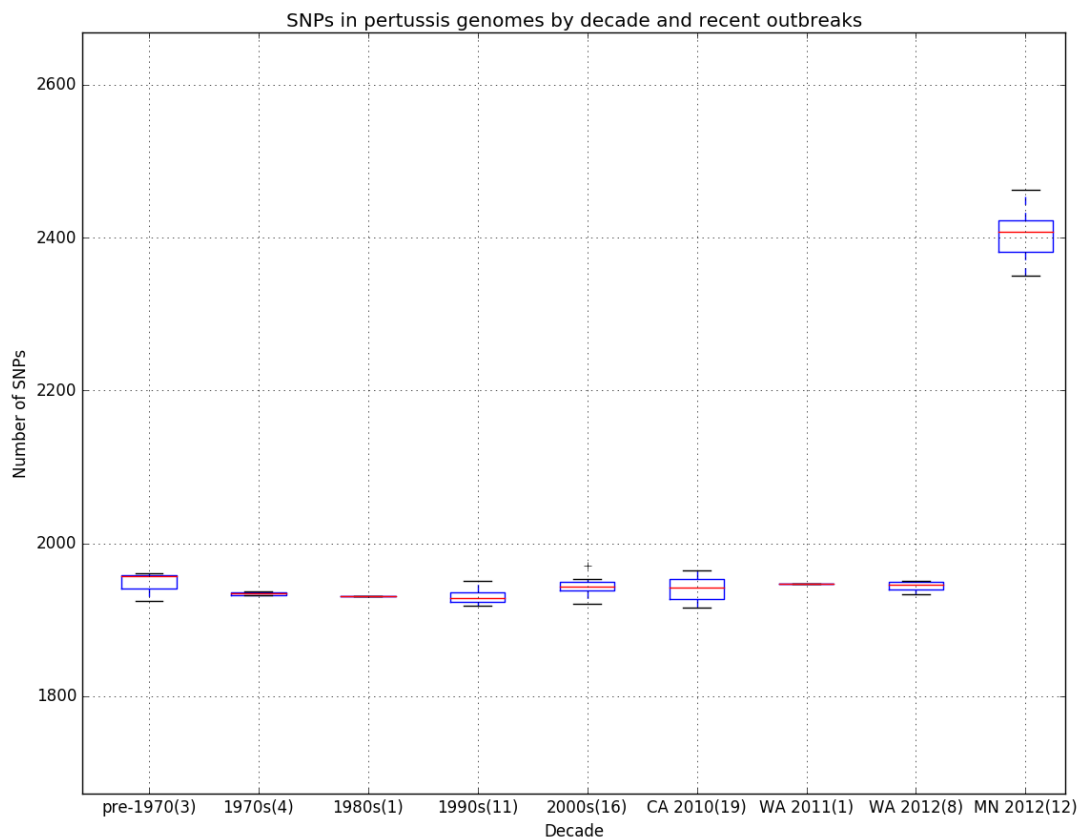


Figure 2: **Median SNPs/genome in all *B. pertussis* strains** Boxplots show median SNPs/genome studied grouped by decade and by outbreak. Numbers in parentheses indicate number of genomes for the given decade/outbreak.

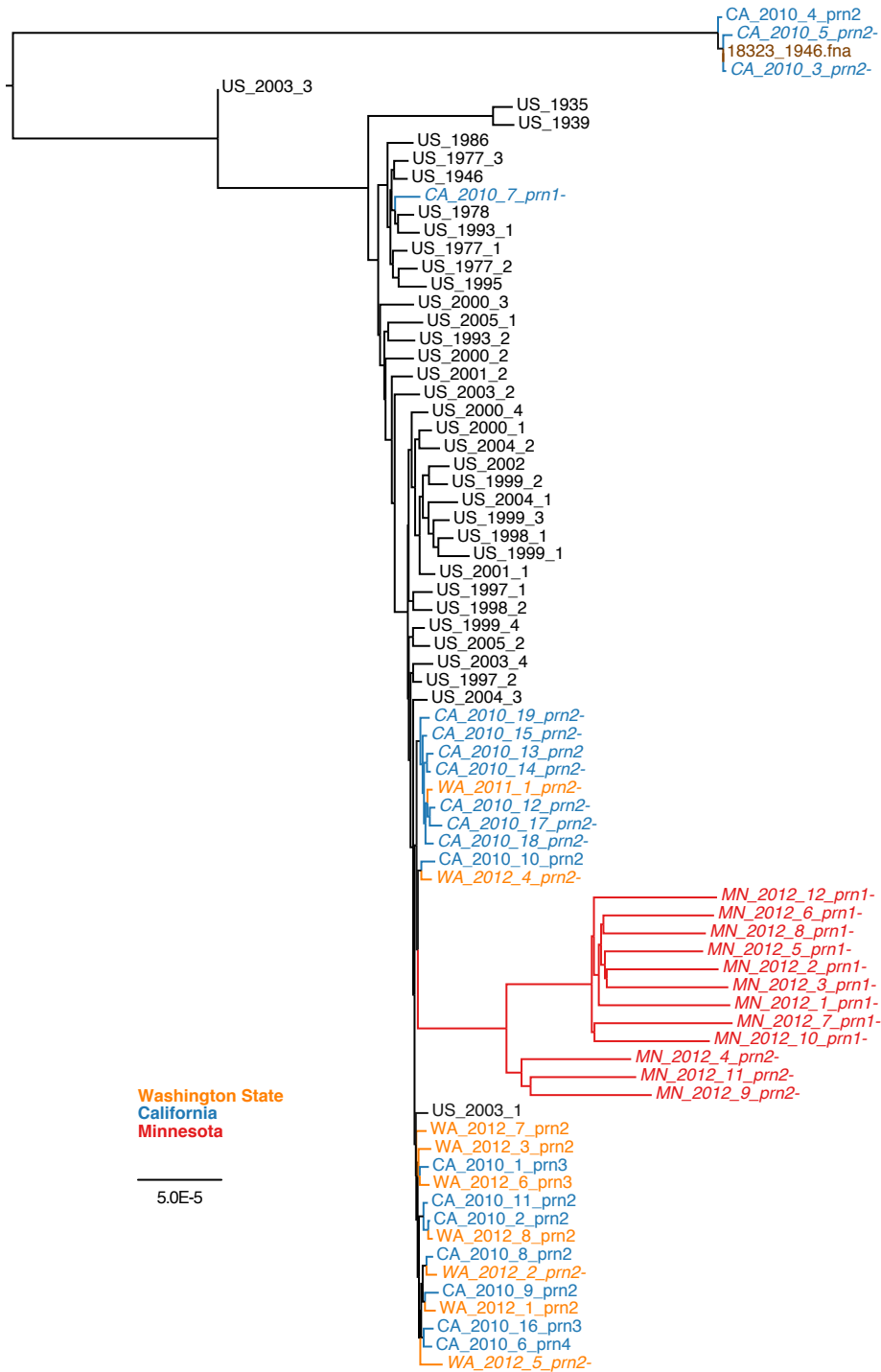


Figure 3: **Phylogenetic tree of 76 *B. pertussis* genomes based on whole genome alignment** Genomes are color-coded: black indicates historical genomes; red, Minnesota outbreak; orange, Washington State outbreak; blue, California outbreak; and brown, reference genome. The Minnesota, California, and Washington outbreak genomes are labeled with their state, year, sample number, and pertactin allele. If the pertactin gene was disrupted, there is a - sign after the pertactin allele and the label is italicized. The unit of scale is substitutions/site.

Table 1: **Non-synonymous and Nonsense SNPs significantly associated with the Minnesota outbreak strains.** Gene and protein annotations are from the 18323 reference genome Genbank file.

Gene	Protein	SNP Type	Codon position	Reference amino acid	Non-reference amino acid
tsr	methyl-accepting chemotaxis protein I	Non-synonymous	498	Ala	Val
putA	bifunctional proline oxidoreductase/transcriptional repressor	Non-synonymous	1222	Ala	Val
nuoD	respiratory-chain NADH dehydrogenase, 49 kDa subunit	Non-synonymous	313	Ile	Ser
mutL	DNA mismatch repair protein	Non-synonymous	356	Ala	Val
maeB	NADP-dependent malic enzyme	Non-synonymous	628	Gly	Ser
maeB	NADP-dependent malic enzyme	Non-synonymous	628	Gly	Asp
maeB	NADP-dependent malic enzyme	Non-synonymous	625	Gln	Arg
maeB	NADP-dependent malic enzyme	Non-synonymous	643	Ser	Thr
lnt	putative apolipoprotein N-acyltransferase	Non-synonymous	15	Ile	Thr
hmp1	ribonuclease E	Non-synonymous	653	Thr	Ala
fliK	flagellar hook-length control protein	Non-synonymous	338	Ala	Thr
dadA	D-amino acid dehydrogenase small subunit	Non-synonymous	258	Arg	His
ctaD	cytochrome c oxidase polypeptide I	Non-synonymous	239	Thr	Ala
carB	carbamoyl-phosphate synthase large chain	Non-synonymous	796	Ser	Gly
argD	putative acetylornithine aminotransferase	Non-synonymous	111	Gly	Ala
BN118_3679	putative inner membrane protein	Non-synonymous	51	Leu	Pro
BN118_3610	TolA protein / Proline-rich inner membrane protein	Non-synonymous	237	Ala	Pro
BN118_3610	TolA protein / Proline-rich inner membrane protein	Non-synonymous	237	Ala	Glu
BN118_3090	conserved hypothetical protein	Non-synonymous	65	Val	Leu
BN118_2919	probable enoyl-CoA hydratase/isomerase	Non-synonymous	6	Ser	Leu
BN118_2793	conserved hypothetical protein	Non-synonymous	264	Ala	Thr
BN118_2781	conserved hypothetical protein	Non-synonymous	176	Leu	Phe
BN118_2428	ABC transport protein, inner membrane component	Non-synonymous	110	Arg	His
BN118_2296	putative membrane protein	Non-synonymous	75	Pro	Gln
BN118_2252	putative outer membrane protein	Non-synonymous	481	Pro	Ser
BN118_1681	probable LysR-family transcriptional regulator	Non-synonymous	160	Asp	Glu
BN118_0965	putative permease component of branched-chain amino acid transport system	Non-synonymous	454	Gly	Asp
BN118_0794	putative periplasmic protein	Non-synonymous	199	Gly	Asp
BN118_0462	putative exported protein	Non-synonymous	154	Ala	Val
BN118_0138	putative exported protein	Non-synonymous	69	Ala	Glu
BN118_3321	conserved hypothetical protein	Nonsense	163	Gln	STO

Table 2: **SNPs significantly associated with the California and Washington outbreaks.** Gene and protein annotations are from the 18323 reference genome Genbank file.

Gene	Protein	SNP Type	Codon position	Reference amino acid	Non-reference amino acid
tsr	methyl-accepting chemotaxis protein I	Synonymous	184	Ala	Ala
bfrI	putative ferrisiderophore receptor	Non-synonymous	219	Arg	Gln
BN118_2634	threonine synthase	Non-synonymous	337	Glu	Gly
BN118_2561	hypothetical protein	Non-synonymous	135	Val	Gly
BN118_2262	putative phospholipase	Non-synonymous	120	Ala	Gly
BN118_2256	putative transferase	Non-synonymous	362	Ala	Gly
BN118_2154	putative ABC-transport protein, ATP-binding component	Non-synonymous	70	His	Pro
BN118_1839	putative oxidoreductase	Synonymous	43	Arg	Arg
BN118_1806	probable probable aldehyde dehydrogenase	Non-synonymous	88	Ala	Gly
BN118_0965	putative permease component of branched-chain amino acid transport system	Non-synonymous	411	Ala	Gly
BN118_0965	putative permease component of branched-chain amino acid transport system	Non-synonymous	574	Met	Val
BN118_0965	putative permease component of branched-chain amino acid transport system	Synonymous	75	Leu	Leu
BN118_0964	putative ATP-binding component of branched-chain amino acid ABC transporter	Non-synonymous	257	Val	Ala
BN118_0964	putative ATP-binding component of branched-chain amino acid ABC transporter	Synonymous	114	Gly	Gly
BN118_0963	putative ATP-binding component of ABC transporter	Non-synonymous	152	Arg	Cys
BN118_0961	probable oxidoreductase	Non-synonymous	142	Arg	His
BN118_0960	putative monooxygenase	Non-synonymous	41	Arg	Leu
BN118_0659	conserved integral membrane protein	Synonymous	500	Gly	Gly
BN118_0189	conserved hypothetical protein	Non-synonymous	152	Arg	Gly



Table 3: **SNPs significantly associated with pertactin(-) strains.** Gene and protein annotations are from the 18323 reference genome Genbank file.

Gene	Protein	SNP Type	Codon position	Reference amino acid	Non-reference amino acid
BN118_0189	conserved hypothetical protein	Non-synonymous	152	Arg	Gly
BN118_2561	hypothetical protein, nicking endonuclease	Non-synonymous	135	Val	Gly
ctaD	cytochrome c oxidase polypeptide I	Synonymous	420	Lys	Lys
maeB	NADP-dependent malic enzyme	Non-synonymous	628	Gly	Asp
maeB	NADP-dependent malic enzyme	Non-synonymous	628	Gly	Ser
maeB	NADP-dependent malic enzyme	Non-synonymous	625	Gln	Arg
maeB	NADP-dependent malic enzyme	Synonymous	640	Asn	Asn
tsr	methyl-accepting chemotaxis protein I	Non-synonymous	498	Ala	Val
tsr	methyl-accepting chemotaxis protein I	Synonymous	510	Ala	Ala
tsr	methyl-accepting chemotaxis protein I	Synonymous	509	Ala	Ala
wbpO	capsular polysaccharide biosynthesis protein	Synonymous	19	Gly	Gly
wbpO	capsular polysaccharide biosynthesis protein	Synonymous	20	Tyr	Tyr
wbpO	capsular polysaccharide biosynthesis protein	Synonymous	18	Leu	Leu
wbpO	capsular polysaccharide biosynthesis protein	Synonymous	21	Val	Val