

Comparative analysis of the genome sequences of *Bordetella pertussis*, *Bordetella parapertussis* and *Bordetella bronchiseptica*

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Bordetella pertussis, *Bordetella parapertussis* and *Bordetella bronchiseptica* are closely related Gram-negative β -proteobacteria that colonize the respiratory tracts of mammals. *B. pertussis* is a strict human pathogen of recent evolutionary origin and is the primary etiologic agent of whooping cough. *B. parapertussis* can also cause whooping cough, and *B. bronchiseptica* causes chronic respiratory infections in a wide range of animals. We sequenced the genomes of *B. bronchiseptica* RB50 (5,338,400 bp; 5,007 predicted genes), *B. parapertussis* 12822 (4,773,551 bp; 4,404 genes) and *B. pertussis* Tohama I (4,086,186 bp; 3,816 genes). Our analysis indicates that *B. parapertussis* and *B. pertussis* are independent derivatives of *B. bronchiseptica*-like ancestors. During the evolution of these two host-restricted species there was large-scale gene loss and inactivation; host adaptation seems to be a consequence of loss, not gain, of function, and differences in virulence may be related to loss of regulatory or control functions.

B. pertussis, *B. parapertussis* and *B. bronchiseptica* colonize the respiratory tracts of mammals. *B. pertussis* is strictly a human pathogen and is the etiologic agent of whooping cough, an acute respiratory disease that is particularly serious in children¹. Despite extensive vaccination programs, whooping cough is still endemic in many countries, causing approximately 285,000 deaths in 2001 (ref. 2). *B. pertussis* strains show little genetic variation, indicating that the species derived from a common ancestor in the recent past, perhaps only a few thousand years ago³. *B. parapertussis* infects both humans and sheep; in human infants it causes whooping cough^{4,5}. Phylogenetic analyses have shown that *B. parapertussis* strains isolated from humans (*B. parapertussis*_{hu}) are distinct from those isolated from sheep (*B. parapertussis*_{ov}; ref. 6). It has been suggested that *B. parapertussis*_{hu} and *B. parapertussis*_{ov} evolved independently from a common ancestor (*B. bronchiseptica*; ref. 6), and there is little or no transmission between the two reservoirs

(sheep and human). Furthermore, *B. parapertussis*_{ov} isolates are genetically diverse, whereas *B. parapertussis*_{hu} isolates are uniform⁶. *B. bronchiseptica* has a broad host range, causing chronic and often asymptomatic respiratory infections in a wide range of animals, but only occasionally in humans⁷.

Many of the virulence factors characterized in the bordetellae are common to all three species. These include adhesins, such as filamentous hemagglutinin (FHA), pertactin, tracheal colonization factor and fimbriae, and toxins, such as adenylate cyclase-hemolysin, dermonecrotic toxin and tracheal cytotoxin. Other virulence factors are expressed by just one of the species, such as pertussis toxin and serum-resistance protein expressed by *B. pertussis* or a type-III secretion system expressed by *B. bronchiseptica* (reviewed in ref. 8).

Expression of many of the proteins implicated in *Bordetella* pathogenesis is regulated in response to environmental stimuli by

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a two-component regulatory system called BvgA/S (reviewed in ref. 8). Environmental stimuli cause the inner membrane sensor histidine kinase BvgS to undergo autophosphorylation before phosphorylating the BvgA protein. Phosphorylated BvgA activates the transcription of a number of virulence-activated genes (vags) by binding to sites in their promoters. One such gene encodes the regulator BvgR that represses the transcription of a number of other genes called virulence-repressed genes (vrgs). Thus, activation of Bvg markedly changes the gene expression profile of the bacterium. Many genes implicated in pathogenesis are upregulated when the Bvg system is active, suggesting that this phase (Bvg-plus) is adopted by bacteria on entering the host during the course of infection. Conversely, the Bvg-minus phase is thought to be expressed when the bacteria are in the environment. A third dis-

tinct phenotypic phase, Bvg-intermediate (Bvgi), is characterized by the lack of expression of the vrgs and by the expression of only a few of the vags. The signals to which Bvg responds *in vivo* are unknown, but *in vitro*, the system is activated by growth at 37 °C and silenced either by growth below 27 °C or by adding sulfate ions or nicotinic acid to the growth medium, regardless of growth temperature.

We sequenced and compared the complete genomes of *B. pertussis* Tohama I, a clinical isolate that has subsequently been used widely for studies in the laboratory; *B. paraptussis* strain 12822, isolated in 1993 from a baby in Germany; and *B. bronchiseptica* strain RB50, isolated from a rabbit. Comparison of the genomes of these bacteria elucidates the genetic background of their differences in disease severity and the factors that determine their host range.

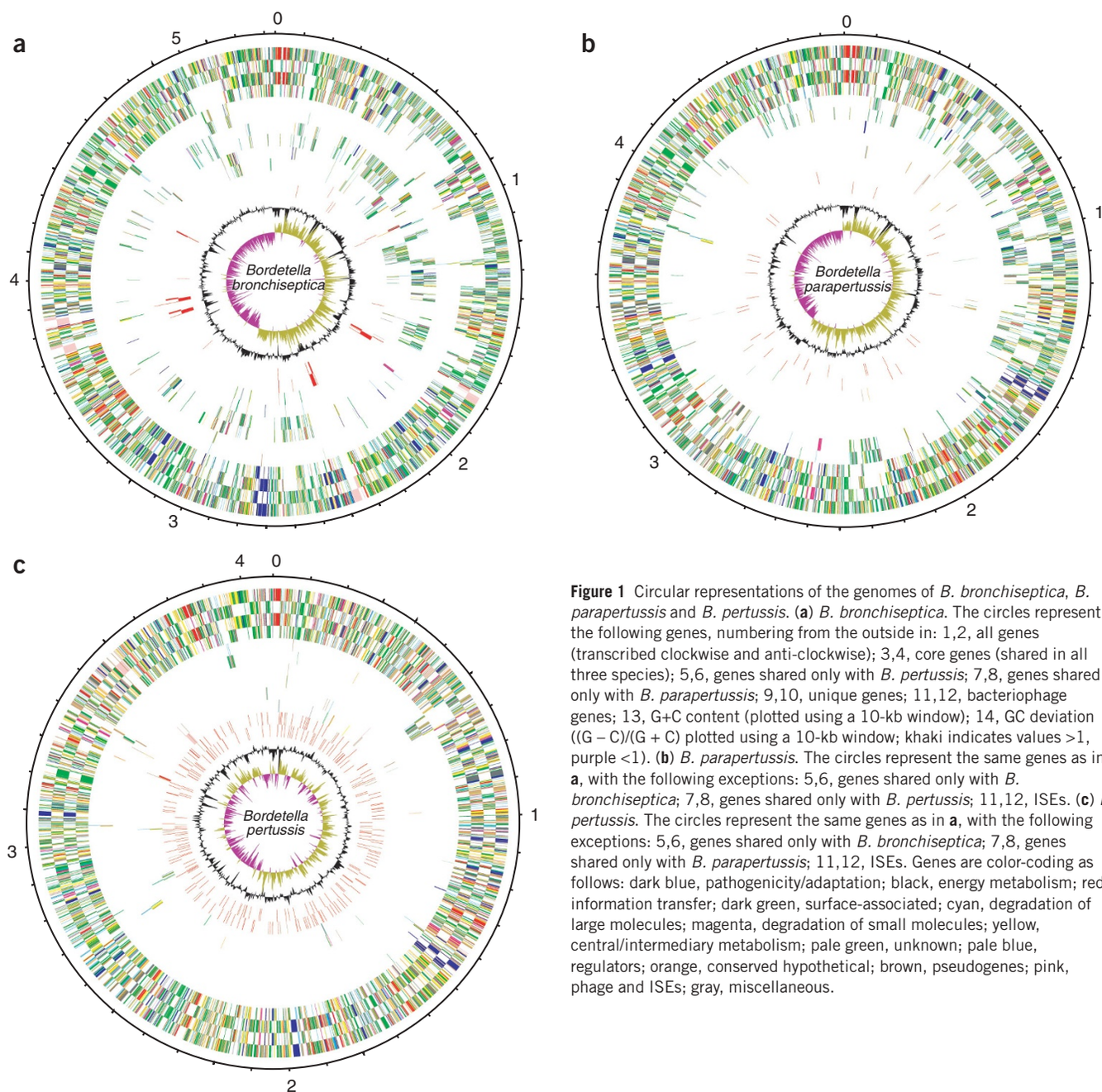


Figure 1 Circular representations of the genomes of *B. bronchiseptica*, *B. paraptussis* and *B. pertussis*. **(a)** *B. bronchiseptica*. The circles represent the following genes, numbering from the outside in: 1,2, all genes (transcribed clockwise and anti-clockwise); 3,4, core genes (shared in all three species); 5,6, genes shared only with *B. pertussis*; 7,8, genes shared only with *B. paraptussis*; 9,10, unique genes; 11,12, bacteriophage genes; 13, G+C content (plotted using a 10-kb window); 14, GC deviation ($(G - C)/(G + C)$ plotted using a 10-kb window; khaki indicates values >1 , purple <1). **(b)** *B. paraptussis*. The circles represent the same genes as in **a**, with the following exceptions: 5,6, genes shared only with *B. bronchiseptica*; 7,8, genes shared only with *B. pertussis*; 11,12, ISEs. **(c)** *B. pertussis*. The circles represent the same genes as in **a**, with the following exceptions: 5,6, genes shared only with *B. bronchiseptica*; 7,8, genes shared only with *B. paraptussis*; 11,12, ISEs. Genes are color-coding as follows: dark blue, pathogenicity/adaptation; black, energy metabolism; red, information transfer; dark green, surface-associated; cyan, degradation of large molecules; magenta, degradation of small molecules; yellow, central/intermediary metabolism; pale green, unknown; pale blue, regulators; orange, conserved hypothetical; brown, pseudogenes; pink, phage and ISEs; gray, miscellaneous.

RESULTS

Structure of the genomes

The general properties of the three genomes are shown in **Table 1** and **Figure 1**. Our data indicate that at the genetic level, *B. pertussis* and *B. parapertussis* each derived from a *B. bronchiseptica*-like ancestor. Because strains of *B. pertussis* (and human isolates of *B. parapertussis*) are genetically uniform, it seems possible that each evolved recently from a strain of *B. bronchiseptica*. We therefore estimated the time to a last common ancestor between each pair of genomes based on the pairwise synonymous substitution rates for all ~3,000 genes that are common to the three species. The estimated times to the last common ancestors were 0.27–1.4 million years (My) for *B. bronchiseptica* and *B. parapertussis*, 0.7–3.5 My for *B. bronchiseptica* and *B. pertussis* and 0.8–4.0 My for *B. parapertussis* and *B. pertussis*, suggesting that the genetic uniformity in *B. pertussis* reflects a recent bottleneck rather than recent descent from *B. bronchiseptica*. Alternatively, *B. bronchiseptica* might be sufficiently genetically diverse that some isolates are more similar to the last common ancestor of *B. pertussis* than is strain RB50, whose genome was sequenced here. These estimates have a large range, reflecting the uncertainty over actual substitution rates in bacterial populations, and should therefore be treated with caution.

Consistent with the relative degrees of divergence at the nucleotide level, it is apparent from the gross comparison between the genomes (**Fig. 2**) that *B. bronchiseptica* and *B. parapertussis* are more similar, in terms of overall organization, than *B. bronchiseptica* and *B. pertussis*. The comparison between *B. bronchiseptica* and *B. parapertussis* shows a large degree of collinearity between the genomes, primarily around the replication origin. The region around the terminus has eight rearranged blocks of DNA. Most chromosomal rearrangements

Table 1 General features of the genomes of *B. pertussis*, *B. parapertussis* and *B. bronchiseptica*

	<i>B. pertussis</i>	<i>B. parapertussis</i>	<i>B. bronchiseptica</i>
Size (bp)	4,086,186	4,773,551	5,338,400
G+C content (%)	67.72	68.10	68.07
Coding sequences	3,816	4,404	5,007
Pseudogenes	358 (9.4%)	220 (5.0%)	18 (0.4%)
Coding density (intact genes)	82.9%	86.6%	91.4%
Coding density (all genes)	91.6%	92.2%	92.0%
Average gene size (bp)	978	987	978
rRNA operons	3	3	3
tRNA	51	53	55
IS481	238	0	0
IS1001	0	22	0
IS1002	6	90	0
IS1663	17	0	0

between related bacteria are arranged reciprocally around the origin or terminus of replication, because, it has been suggested, most recombination occurs between, or close to, replication forks⁹ or because other rearrangements may occur but are selectively disadvantageous¹⁰. Most of the rearrangements in *B. parapertussis* (and *B. pertussis*) do not follow this pattern, suggesting that any such selection may be overcome by frequent recombination between large perfect repeats. Indeed, each rearrangement is bordered by identical copies of the insertion sequence elements (ISEs) IS1001 or IS1002 in *B. parapertussis*, suggesting that recombination between these ISEs was the primary cause of the rearrangements.

Compared with *B. parapertussis*, *B. bronchiseptica* has several large discrete regions of unique DNA. Several of these are prophage insertions, but three of them (80 kb around position 750,000, 120 kb around position 1,083,000 and 65 kb around position 4,981,000) clearly are not. Two lines of evidence suggest that these large blocks of DNA are deletions in *B. parapertussis* rather than insertions in *B. bronchiseptica*. First, similar, though not identical, blocks of DNA are absent from *B. pertussis*, but the boundaries of the regions are different in *B. parapertussis* and *B. pertussis*, suggesting that independent deletions of similar regions occurred in the two species. Second, analysis of the *B. bronchiseptica* genome for signals associated with horizontally transferred DNA, such as anomalous G+C content, dinucleotide content or GC-skew¹¹, clearly indicated that such sequences as the prophage were of recent horizontal acquisition but gave no such indication for these three large regions of difference. Similar analysis of the whole *B. bronchiseptica* genome suggested that little of the DNA was probably acquired recently and certainly not enough to account for the differences in size between the three genomes.

Genomic comparison of *B. bronchiseptica* and *B. pertussis* (**Fig. 2**) showed that rearrangements and deletions in *B. pertussis* were similar

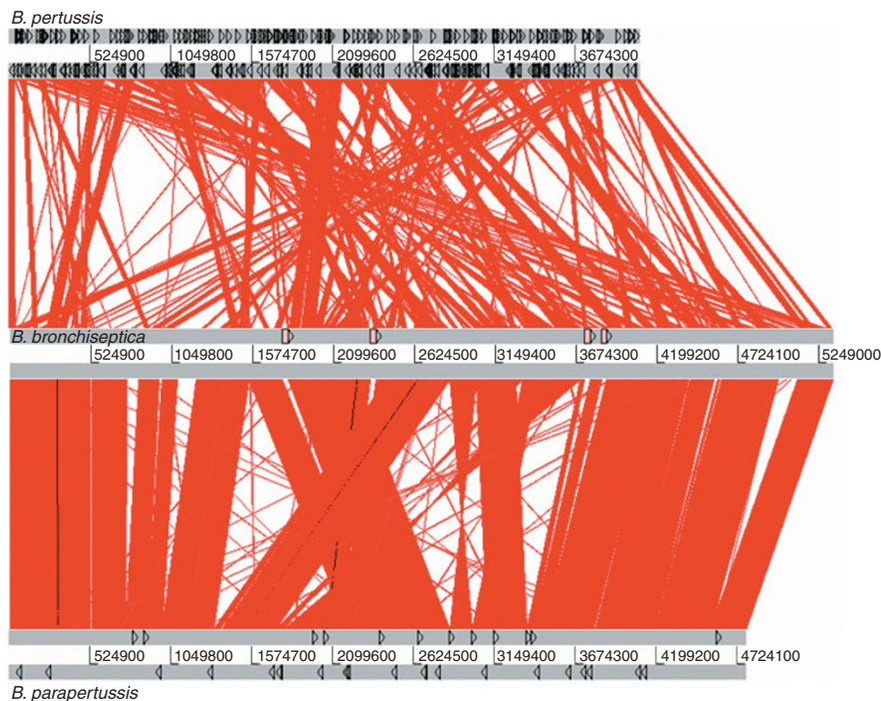


Figure 2 Linear genomic comparison of *B. pertussis*, *B. bronchiseptica* and *B. parapertussis*. The gray bars represent the forward and reverse strands. Top, *B. pertussis*. Black triangles represent ISEs. Center, *B. bronchiseptica*. Pink boxes represent prophage. Bottom, *B. parapertussis*. Black triangles represent ISEs. The red lines between the genomes represent DNA:DNA similarities (BLASTN matches) between the two sequences.

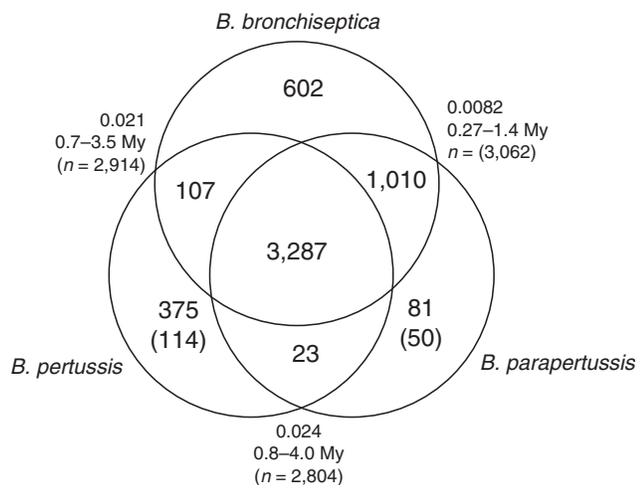


Figure 3 Venn diagram showing gene complements of *B. pertussis*, *B. paraptussis* and *B. bronchiseptica*. Numbers in parentheses indicate numbers of unique genes, excluding those from ISEs. Figures outside the circles indicate the average synonymous substitution rates (number of synonymous substitutions per potential substitution site) for the set of core genes between each pair of organisms and the estimated age of divergence in million years (My) calculated from these rates.

to those in *B. paraptussis* but more extreme, with short blocks of almost perfect conservation broken up by nearly 150 individual rearrangements, 88% of which are bounded by ISEs (primarily IS481). Again, the rearrangements do not follow the usual reciprocal pattern around the origin or terminus of replication. The large degree of rearrangement of overall genome structure is unprecedented for congeneric bacteria.

The *B. pertussis* genome is considerably smaller than that of *B. bronchiseptica*, and, although individual events are difficult to trace, much of this loss is probably due to ISE-mediated deletion events. At some point in its recent evolution, *B. pertussis* seems to have undergone a massive expansion of one family of ISEs (IS481; **Table 1**), and subsequent recombination between these perfect DNA repeats caused a large amount of rearrangement and deletion in the chromosome. Comparison of the genetic maps of several *B. pertussis* strains showed that inversions of large sections of the genome are frequent, presumably owing to such recombination¹². Such ISE expansions have previously been seen in rarely recombining organisms whose effective population size was greatly reduced by an evolutionary bottleneck¹³. In any population, ISEs transpose to novel sites at a certain rate, but most of these novel insertions are lethal or carry a selective disadvantage, and bacteria carrying them are therefore competed out of the population. When the population size is very small, however, the degree of intraspecies competition is reduced, and bacteria carrying such non-lethal mutations are less likely to be competed out, leading to an increase in ISE accumulation in the population.

Gene complements

Comparison of the gene complements of the three species supports the inference from the gross structure that *B. pertussis* and *B. paraptussis* are independent derivatives of a *B. bronchiseptica*-like organism. Apart from ISE transposases, only 114 genes are unique to *B. pertussis* compared with *B. bronchiseptica* and *B. paraptussis*,

and only 50 genes are unique to *B. paraptussis* compared with *B. bronchiseptica* and *B. pertussis* (**Fig. 3**). In fact, most genes that are apparently specific to *B. pertussis* seem to reflect diversity in *B. bronchiseptica* as indicated by microarray analysis (C. Cummings and D. Relman; personal communication).

Furthermore, only 23 genes were found in both *B. pertussis* and *B. paraptussis* but not in *B. bronchiseptica* RB50. Thus, these derivative organisms seem to have little net gain of genes. Investigation of these genes (**Supplementary Table 1** online) shows few obvious virulence factors. Conversely, *B. bronchiseptica* has over 600 genes not found in either of the other two species, over 1,000 shared only with *B. paraptussis* and just over 100 shared only with *B. pertussis*. Analysis of the genes that were lost in *B. pertussis* and *B. paraptussis* (**Fig. 4**) indicates that many are involved in membrane transport, small-molecule metabolism, regulation of gene expression and synthesis of surface structures. The three large deletions identified by comparing *B. bronchiseptica* and *B. paraptussis* contain genes that are unique to *B. bronchiseptica* and, therefore, that were probably lost in both *B. pertussis* and *B. paraptussis*. Some of these genes are involved in the transport and metabolism of a wide range of compounds, including amino acids, other aromatic compounds, fatty acids, nitrobenzene/benzoate, polyamines, phosphonates, phthalate, vanillate and mandelate. There is also a cluster of genes encoding a type-IV pilus (BB0776-BB0792), two siderophore receptors and several regulatory proteins.

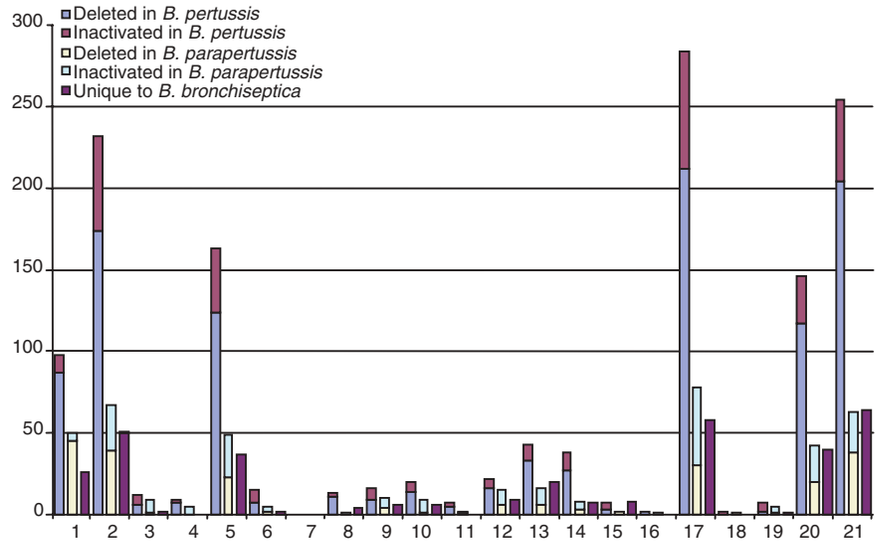
In addition to this substantial gene deletion in *B. pertussis* and *B. paraptussis*, both seem to have lost the function of many other genes through pseudogene formation. In *B. pertussis*, 358 genes have been inactivated by insertion of ISEs, in-frame stop codons or frameshifts; similarly, 200 genes in *B. paraptussis* have been inactivated by these mechanisms. Conversely, *B. bronchiseptica* has only 19 pseudogenes (**Supplementary Table 2** online). Again, analysis of the inactivated genes in *B. pertussis* and *B. paraptussis* indicates that they are predominantly involved in transport, small-molecule metabolism, regulation and surface structures (**Fig. 4**).

Metabolism

B. bronchiseptica can survive in the environment, but *B. pertussis* and *B. paraptussis* are apparently incapable of long-term survival outside the host. Despite this, much of the central and intermediary metabolism is conserved between the three genomes. Consistent with their ability to grow on defined media in culture, the genomes of all three *Bordetella* species seem to encode complete pathways for the biosynthesis of nucleotides, cofactors and amino acids (except cysteine, see below). Generally, *Bordetella* do not use sugars as carbon sources. This is probably because three genes encoding glycolytic functions (glucokinase, phosphofructokinase and fructose-1,6-bisphosphatase) are absent from the *Bordetella* genomes. Genes encoding the gluconeogenesis pathway are present, suggesting that glucose can be synthesized by *Bordetella* spp.

The metabolism of *B. pertussis* has been studied in some detail; it can grow in culture on glutamate and cysteine¹⁴. In laboratory media, cysteine is present at low levels and probably serves as a source of sulfur rather than carbon, suggesting that glutamate is the main carbon source for cultured *B. pertussis*. Given this cysteine auxotrophy, it is not surprising that the genes encoding several of the proteins of the sulfate transport (*cysU* and *cysW*) and cysteine biosynthesis pathway (*cysH* and *cysN*) are pseudogenes in *B. pertussis* and genes related to other key steps (*cysC*, *cysI*, *cysJ* and *cysP*) are absent. Not all the defects found in *B. pertussis* are present in *B. paraptussis* and *B. bronchiseptica*, but the pathway is mutated in

Figure 4 Representation of genes lost in *B. pertussis* and *B. paraptussis*. Blue, genes present in *B. paraptussis* and not *B. pertussis* (genes lost by *B. pertussis*); mauve, pseudogenes in *B. pertussis*; yellow, genes present in *B. pertussis* and not *B. paraptussis* (genes lost by *B. paraptussis*); pale blue, pseudogenes in *B. paraptussis*; purple, genes unique to *B. bronchiseptica* (genes lost by both *B. pertussis* and *B. paraptussis*). Functional classifications are as follows: 1, unknown; 2, conserved hypothetical; 3, cell processes; 4, protection responses; 5, transport/binding proteins; 6, adaptation; 7, cell division; 8, macromolecule degradation; 9, macromolecule synthesis/modification; 10, amino acid biosynthesis; 11, cofactor biosynthesis; 12, central/intermediary metabolism; 13, small-molecule degradation; 14, energy metabolism; 15, fatty acid biosynthesis; 16, ribonucleotide biosynthesis; 17, cell surface; 18, ribosome constituents; 19, pathogenicity; 20, regulation; 21, miscellaneous. Genes from mobile elements (ISEs and bacteriophage) were omitted from the figure.



both. This suggests that an ancestral *Bordetella* could use external sulfate or thiosulfate for cysteine biosynthesis but this ability was since lost. The genes related to synthesis of cysteine from acetyl-coenzyme A, serine and sulfide are present and intact, but the cysteine auxotrophy of *B. pertussis* suggests that this pathway does not function, and an IS481 insertion immediately upstream of the start codon of *cysM* may explain this.

It has been reported that the tricarboxylic acid cycle in the bordetellae is non-functional, based on the observation that in these bacteria, acetyl-coenzyme A and oxaloacetate did not give rise to α -ketoglutarate¹⁴. Genes encoding all the main tricarboxylic acid cycle enzymes are present in all three species, however. Thus, there is no obvious genetic defect that explains these observations. Overall, it seems that the inability of *B. pertussis* and *B. paraptussis* to survive in the environment is probably due to the loss of numerous accessory pathways for the use of alternative nutrient sources, described above, rather than a lack of central metabolic pathways.

Host range and pathogenicity

All bordetellae carry a wide range of determinants involved in host interaction and virulence, many of which have been extensively studied. Examples include the filamentous hemagglutinin FhaB; fimbriae; a large number of autotransporters including the adhesins pertactin, tracheal colonization factor and the serum-resistance protein; numerous toxins including the pertussis toxin, adenylate cyclase-hemolysin, dermonecrotic toxin¹⁵ and tracheal cytotoxin; a type-III secretion system; and lipopolysaccharide and flagella biosynthesis systems (reviewed in refs. 8 and 16). Comparisons of many of these systems in the three species identify differences that might bear on the different host ranges and virulence profiles of these organisms.

FhaB (BP1879) is a large secreted protein involved in attachment to host cells¹⁷ and is a member of a growing family of hemagglutinin/hemolysin proteins present in a number of Gram-negative plant and animal pathogens. An ortholog of *fhaB* is present in each of the three species (BP1879, BB2993 and BPP3027); although each seems to be intact, there are internal variations in sequence and

length of the repetitive internal sequences¹⁸. Two other genes encoding FhaB-like proteins are present in *B. pertussis*, BP2667 and BP2907. BP2667 (*fhaS*) is orthologous to BB2312 and BPP1243, and BP2907 (*fhaL*) to BB1936 and BPP2489. In the former case, there are again differences in the internal sequences between the three *Bordetella* spp., although the *fhaS* orthologs seem to be more similar to each other than are the *fhaL* orthologs. FHA is important in adherence of the bacteria to host cells. It binds directly to ciliary membrane glycosphingolipids that are thought to be host receptors

Table 2 Autotransporters encoded in the genomes of *B. pertussis*, *B. paraptussis* and *B. bronchiseptica*

	<i>B. pertussis</i>	<i>B. paraptussis</i>	<i>B. bronchiseptica</i>
SphB1	BP0216	BPP0417	BB0419
Novel	-	BPP0449	BB0450
Novel	BP0529	BPP0452	BB0452
Novel	-	BPP0735	BB0821
Novel	-	BPP0822	BB0916
Serum-resistance protein	BP3494	<u>BPP0867</u>	<u>BB0961</u>
Pertactin	BP1054	BPP1150	BB1366
BapA/AidB	BP2224	<u>BPP2251</u>	BB1649
VagB	BP2315	BPP2415	BB1864
BapC	BP2738	BPP2591	BB2033
Phg	BP1767	BPP1998	BB2246
Novel	<u>BP1793</u>	BPP2022	BB2270
SphB3 (serine protease)	BP1110	BPP2053	BB2301
Novel	<u>BP2627</u>	BPP1256	BB2324
SphB2	BP1660	<u>BPP2745</u>	BB2741
Novel	BP1344	<u>BPP2678</u>	BB2830
Novel	<u>BP1610</u>	BPP2975	BB2941
Novel	-	BPP1618	BB3110
Novel	-	BPP1617	BB3111
TcfA	BP1201	-	BB3291
BapB	BP1200	BPP1815	BB3292

Pseudogenes are underlined.

Table 3 TonB-dependent ferric complex receptors encoded in the genomes of *B. pertussis*, *B. paraptussis* and *B. bronchiseptica*

	<i>B. pertussis</i>	<i>B. paraptussis</i>	<i>B. bronchiseptica</i>
BfeA	BP2901	BPP2495	BB1942
BfrA	–	–	BB4761
BfrB	BP2016	BPP2396	BB1846
BfrC	BP3663	BPP0079	BB0078
BfrD	BP0856	BPP3376	BB3826
BfrE	BP0857	<u>BPP3375</u>	BB3825
BfrF	BP0736	BPP3688	BB4122
BfrG	BP2922	BPP1078	BB1294
BfrH	BP1138	BPP3206	BB3658
BfrI	BP1962	BPP2334	BB1785
BfrZ	–	–	BB4744
BhuR	BP0347	BPP4185	BB4655
FauA	BP2463	BPP3450	BB3900
Novel	BP0456	<u>BPP4370</u>	BB4956
Novel	–	BPP2457	BB1905
Novel	<u>BP3101</u>	<u>BPP0746</u>	BB0832

Pseudogenes are underlined.

for *Bordetella* adherence¹⁹. Differential binding of the three *Bordetella* species to ciliated cells from different hosts suggests that host specificity may be partly dependent on this receptor-ligand interaction²⁰. The identification of additional FHA-like genes that vary between *Bordetella* species is of interest to the study of the host-pathogen interaction and host specificity.

As described above, the genome of *B. bronchiseptica* contains a type-IV pilus biosynthesis operon that is absent in the other two species. Other fimbrial systems are also encoded by the genomes. The previously described chaperone-usher fimbrial system (*fimA-fimD*) is present in all three, although the fimbrial gene *fimA*, which has been inactivated by a deletion in *B. pertussis* (BP1880; ref. 21) and by a frameshift in *B. paraptussis* (BPP3026), is apparently intact in *B. bronchiseptica* (BB2992). Other isolated fimbrial genes exist: the serotype 3 subunit (*fim3*; ref. 22) is intact in all three species; *fimN*²³ is present in *B. bronchiseptica* and *B. paraptussis* (although with a variable C terminus) but deleted from *B. pertussis*; *fimX*²⁴ is intact in *B. pertussis* and *B. bronchiseptica* but frameshifted in *B. paraptussis*; the serotype 2 fimbrial subunit (*fim2*; ref. 25) is present in all three species but with a variable C terminus in *B. pertussis*; and there is a novel subunit upstream of *fimN* present in *B. bronchiseptica* and *B. paraptussis* but deleted from *B. pertussis*.

Variation is also seen in the complement of autotransporters. These are members of a large family of exported proteins that encode an integral outer-membrane pore, enabling them to cross the outer membrane²⁶. The bordetellae encode 21 of these proteins, a few (serum-resistance protein BrkA, pertactin, SphB1, TcfA and Vag8) with documented functions in host interaction and virulence^{26,27}. Each of the three species contains a different complement of autotransporters (Table 2). *B. bronchiseptica* includes what might be considered the complete set of genes, albeit with one represented by a pseudogene. *B. paraptussis* and *B. pertussis* each have fewer autotransporter genes and more autotransporter pseudogenes than *B. bronchiseptica*. As with the variation in sequence of the FHA and fimbrial proteins, this variable complement of autotransporter proteins may have a direct effect on the ability of each species to interact with, and cause disease in, different hosts.

Iron acquisition is of great importance to mammalian pathogens, as iron is present in limiting conditions in the host, and any iron that is present is sequestered by heme and other iron-chelating compounds. Many pathogens produce iron-chelating compounds called siderophores to acquire iron, and the three bordetellae each contain an operon encoding the production, export and uptake of the siderophore alcaligin²⁸ (BP2456-BP2463, BB3893-BB3900 and BPP3443-BPP3450). Iron from siderophores and host iron-binding complexes is internalized through TonB-dependent outer-membrane ferric complex receptors; the three *Bordetella* species encode up to 16 of these proteins (Table 3), suggesting that they scavenge host iron-binding complexes and xenosiderophores in addition to using their own. As with the autotransporters, *B. bronchiseptica* encodes a complete set of these receptors, with *B. paraptussis* and *B. pertussis* having fewer genes and more pseudogenes. The genes encoding several of these receptors have adjacent cognate regulators (AraC, AsnC family and MarR family transcriptional regulators, as well as a two-component system that consists of a sensor protein and an extracytoplasmic function sigma factor). Like the TonB-dependent receptors, all these regulators are intact in *B. bronchiseptica*. In contrast, a few of them are pseudogenes in *B. pertussis* and *B. paraptussis* even when their cognate TonB-dependent receptor genes are intact. Again, it is possible that these different iron-uptake system gene complements affect the range of environments in which the bacteria can survive.

Three structures of great importance for virulence and host interaction in Gram-negative pathogens are type-III secretion systems, O-antigens and flagella; each of these show species specific variations among the three *Bordetella* species. Type-III secretion systems are specific mechanisms for the export of virulence factors into host cells²⁹. The three bordetellae encode a novel type-III secretion system for which the secreted effectors have not yet been identified^{30,31}. *B. bronchiseptica* encodes the full, intact operon (BB1609-BB1638), which is expressed. The operon in *B. paraptussis* (BPP2212-BPP2241) contains two identifiable pseudogenes in regulatory and structural proteins (BPP2241 and BP2262) and is not expressed. Curiously, the operon in *B. pertussis* does not seem to be expressed^{30,31} but contains no clearly identifiable pseudogenes. It is possible that its inactivity is dependant on *cis*- or *trans*-acting regulatory mutations.

O-antigens are the membrane-distal domains of some lipopolysaccharides and are involved in many bacterial species in evasion of host innate and acquired immunity mechanisms. *B. bronchiseptica* and *B. paraptussis* have been reported to express an identical O-antigen³², but variable modifications to the terminal sugar residue of the O-antigen polymer were recently discovered³³. These modifications involve, for example, formylation of the amino groups of the terminal sugar residue. The precise pattern of modification differs between different strains of *B. bronchiseptica* and correlates with different reactivities of the O-antigens to monoclonal antibodies. Within the O-antigen biosynthesis locus of *B. bronchiseptica* RB50 reported here is a region that differs from the partial locus sequenced from strain CN7635E and reported previously³⁴. This alternative region of the locus contains genes that are predicted to code for formyl transferases, and these are probably responsible for the variable O-antigen modifications between strains. The 'modification region' from *B. paraptussis* sequenced here is identical to the partial locus of *B. bronchiseptica* CN7635E previously reported, and these two strains share an identical reactivity profile to monoclonal antibodies against O-antigen, which is different from that of *B. bronchiseptica* RB50 (A.P., unpublished observations). Thus, this genetic

and the increased virulence of *B. pertussis* over *B. parapertussis* do not seem to be due to recent acquisition of host interaction factors or virulence determinants. There is little evidence overall of recent large-scale acquisition of genes by any of the species. Conversely, it seems that many of the individual traits of the organisms could be due to independent mechanisms of large-scale gene inactivation and loss in the two derivative species, *B. parapertussis* and *B. pertussis*. The limited host range of these species compared to that of a *B. bronchiseptica*-like putative ancestor can be easily explained by the documented loss of host interaction mechanisms. In addition, the increased virulence for humans of *B. pertussis* could also be due to an overexpression or constitutive expression of virulence traits that are subject to tighter temporal or environmental control in the other species or to the removal of structures or systems that might hinder virulence or survival in the single host. It may be that *B. pertussis* followed this evolutionary path owing to the opportunities for increased transmission rate provided by the increase in size and density of the populations of its specific host, *Homo sapiens*. This would support the hypothesis that, when transmission rates are high, the ability to survive in the environment and keep damage to the host to a minimum are no longer selectively advantageous⁴².

METHODS

DNA preparation and sequencing. *B. pertussis* Tohama I was a gift from S. Stibitz (Food and Drug Administration, Bethesda, Maryland). We grew bacteria on Bordet Gengou agar (Difco) supplemented with 15% defibrinated horse blood (TCS Biologicals) at 37 °C for 3 d. We isolated DNA by the agarose plug method⁴³. We obtained the initial genome assembly from 87,500 paired-end sequences (giving 8.9-fold coverage) derived from three pUC18 genomic shotgun libraries (with insert sizes of 1.0–4.0 kb) using dye terminator chemistry on ABI377 automated sequencers. We used 2,560 paired-end sequences from a pBACe3.6 library with insert sizes of 10–20 kb (a clone coverage of 4.7-fold) as a scaffold. We generated another 41,700 sequencing reads during finishing and verified the final assembly by comparison with the published restriction map⁴⁴.

B. bronchiseptica strain RB50 was isolated from a rabbit and was a gift from J. F. Miller (University of California Los Angeles). We obtained the initial genome assembly from 99,000 paired-end sequences (giving 7.95-fold coverage) derived from two pUC18 genomic shotgun libraries (with insert sizes of 1.0–2.0 kb) using dye terminator chemistry on ABI377 and ABI 3700 automated sequencers. We used 4,480 paired-end sequences from two pBACe3.6 libraries with insert sizes of 15–23 kb and 23–48 kb (a total clone coverage of 12.7-fold) as a scaffold. We generated another 22,660 sequencing reads during finishing.

B. parapertussis strain 12822 was a gift from U. Heininger (der Friedrich-Alexander-Universität Erlangen-Nürnberg, Germany). We obtained the initial genome assembly from 83,850 paired-end sequences (giving 8.6-fold coverage) derived from a pUC18 genomic shotgun library (with insert sizes of 2.0–4.0 kb) using dye terminator chemistry on ABI 3700 automated sequencers. We used 3,020 paired-end sequences from a pBACe3.6 library with insert sizes of 23–48 kb (a clone coverage of 11.0-fold) as a scaffold. We generated another 5,450 sequencing reads during finishing.

Annotation and analysis. We assembled, finished and annotated the sequences as described previously⁴⁵, using Artemis⁴⁶ to collate data and facilitate annotation. We compared the DNA and encoded protein sequences of the three species using the Artemis Comparison Tool (K. Rutherford, unpublished data). We identified orthologous gene sets by reciprocal best-match FASTA comparisons. Pseudogenes had one or more mutations that would prevent complete translation and were identified by direct comparison between the three genomes. We then checked each of the inactivating mutations against the original sequencing data. We calculated synonymous substitution frequency (Ds) values from orthologous gene pairs from the core gene set according to the method of Nei and Gojobori⁴⁷ and estimated ages of divergence (Age = Ds/mutation rate) from the mutation rates calculated by Whittam⁴⁸ or Guttman and Dykhuizen⁴⁹.

URLs. The Artemis Comparison Tool is available from <http://www.sanger.ac.uk/Software/ACT/>. The full annotation of each of the sequences and further information are available from http://www.sanger.ac.uk/Projects/B_pertussis/.

EMBL accession numbers. *B. pertussis*, BX470248; *B. parapertussis*, BX470249; *B. bronchiseptica*, BX470250.

Note: Supplementary information is available on the Nature Genetics website.

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COMPETING INTERESTS STATEMENT

The authors declare that they have no competing financial interests.

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