## **MINIREVIEW**

### Evolution of Host Adaptation in Salmonella enterica

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#### INTRODUCTION

The question of how bacteria are able to overcome species barriers and adapt to new hosts is central to the understanding of both the origin of infectious diseases and the emergence of new pathogens. The analysis of virulence factors used by different Salmonella serotypes can serve as a powerful model for studying mechanisms of host adaptation because these pathogens are physiologically well characterized and lend themselves to genetic analysis. However, they differ greatly with regard to host range and their degree of host adaptation. Salmonella serotypes are closely related as shown by analysis of orthologous genes. Divergence in the nucleotide sequence of orthologous genes ranges between 3.8 and 4.6% and differences in their deduced amino acid sequences range between 0.7 and 1.3% (108). This close DNA relatedness among Salmonella serotypes is evidence for their clonal origin, and based on the degree of sequence divergence, it can be estimated that a common ancestor of the genus existed about 25 to 40 million years ago. Which factors contributed to the clonal divergence of the genus Salmonella from its common ancestor to give rise to serotypes that differ with regard to their host range?

It has been postulated recently that in the genus *Salmonella* virulence evolved in three phases (11) (Fig. 1). The first phase involved acquisition of *Salmonella* pathogenicity island 1 (SPI 1) by plasmid- or phage-mediated horizontal gene transfer. SPI 1 was likely obtained by a lineage ancestral to all *Salmonella* serotypes, since it is present in all phylogenetic lineages of the genus *Salmonella* but absent from *Escherichia coli* and other related organisms (70, 79, 85). SPI 1 encodes virulence factors that mediate mechanisms used by *Salmonella* serotypes during the intestinal phase of infection, including invasion of intestinal epithelial cells (48, 62, 120), induction of neutrophil recruitment (49, 77), and secretion of intestinal fluid (49). Thus, it is likely that these or related virulence mechanisms may have existed early in the evolution of *Salmonella* serotypes.

Multilocus enzyme electrophoresis and comparative sequence analysis of housekeeping and rRNA genes revealed that the genus *Salmonella* contains two lineages that have diverged considerably from each other during evolution (24, 31, 97). By using genetic distance determined by multilocus enzyme electrophoresis and results of DNA-DNA hybridization studies as criteria, it has been proposed that these lineages represent two distinct species, designated *Salmonella enterica* and *Salmonella bongori* (Fig. 1) (69, 97). The formation of these two species could be considered a second phase in the

evolution of virulence in the genus Salmonella, since it involved not only divergence of their lineages by point mutation but also acquisition of new virulence determinants by horizontal gene transfer. Serotypes belonging to S. enterica possess a second pathogenicity island, designated SPI 2, that is not present in S. bongori serotypes (55, 84). The virulence genes present on SPI 2 have an average G+C content of 41%, much lower than the overall G+C content of the S. enterica genome, which averages 52% (55). The limited phylogenetic distribution and the atypical G+C content of SPI 2 imply that this pathogenicity island was acquired horizontally after *S. enterica* branched from the *S.* bongori lineage (55, 84). A possible mechanism for acquisition of SPI 2 by horizontal transfer is suggested by its insertion into the S. enterica gene encoding tRNA<sup>Val</sup>, a DNA region which may facilitate integration of newly acquired genetic material because tRNA genes serve as an attachment site for bacteriophage (55, 59). Experiments performed with mice revealed that mutations in SPI 1 attenuate S. enterica serotype Typhimurium between 15- and 50-fold after oral injection but have no attenuating phenotype when the intestinal phase of infection is bypassed by intraperitoneal injection (17, 48). In contrast, mutations in SPI 2 attenuate S. enterica serotype Typhimurium more than 10,000-fold even after intraperitoneal inoculation (86, 109). Although the mechanism by which SPI 2 contributes to pathogenicity has not yet been elucidated, these data imply that virulence genes located on SPI 1 and SPI 2 are required at different stages, namely the intestinal and the systemic phases of infection, respectively.

Finally, the lineage of *S. enterica* is postulated to have branched into several distinct phylogenetic groups, which by current nomenclature are considered subspecies. The formation of one of these groups, *S. enterica* subspecies I, involved a dramatic expansion in host range: while *S. bongori* and *S. enterica* subspecies II, IIIa, IIIb, IV, VI, and VII are mainly associated with cold-blooded vertebrates, members of *S. enterica* subspecies I are most frequently isolated from avian and mammalian hosts (2, 93). The host adaptation of *S. enterica* subspecies I to warm-blooded vertebrates characterized a third phase in the evolution of virulence in the genus *Salmonella* and is the focus of this review.

# LYMPH NODES, A NEW BARRIER ENCOUNTERED IN MAMMALS

Which new barriers did *S. enterica* subspecies I serotypes encounter in birds and mammals? One obstacle may have been the immune system of higher vertebrates, which is more highly developed and shows a higher level of organization than that of cold-blooded vertebrates. For instance, lymphoid aggregates located in the gut lamina propria of cold-blooded vertebrates appear only as solitary units (82) whereas gut-associated lymphoid nodules of mammals and birds are organized in complex

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FIG. 1. Model for the evolution of virulence in the genus Salmonella. The three phases in which virulence evolved in the genus Salmonella since its divergence from the *E. coli* lineage have been proposed previously (11). The phylogenetic tree is not drawn to scale. For explanation see the text.

organs, such as Peyer's patches, tonsils, appendix, or the avian bursa of Fabricius (51, 76). In addition, in birds and mammals B-lymphocyte variants which arise after somatic hypermutation are selected in the germinal centers of lymphoid organs for improving the affinity to their antigen. In contrast, because the lymphoid organs of cold-blooded vertebrates lack germinal centers selection of B cells is impeded, and consequently, antibody affinity does not increase during the immune response (40, 123). Furthermore, the antibody repertoire of lower vertebrates, such as fish, amphibians, and reptiles, is much less diverse than that of mammals (39). B cells selected in germinal centers of higher vertebrates show isotypic switching and are accumulated as memory cells. In cold-blooded vertebrates, on the other hand, immunological memory is poorly developed (39) and repeated immunization with S. enterica induces only the production of immunoglobulin M antibodies in lizards (82), suggesting that isotype switching does not occur during infection with this pathogen.

Finally, the presence of more highly developed peripheral lymphoid filter organs constitutes yet another challenge during infection of warm-blooded hosts. Pathogens that are able to penetrate the intestinal mucosa in fish can spread unchecked to central parts of the body until they are filtered from the blood by phagocytes located in sinusoids of the spleen. During the adaptation to life on land vertebrates developed peripheral lymphoid organs which function as local filtering systems (65). Although less-developed lymph node-like structures have been reported in some reptiles (82), true lymph nodes are present only in mammals and some bird species (41, 42, 51, 123). In mammals, a large number of lymph nodes located at peripheral positions form an highly effective lymph filter system (41, 42). Thus, upon penetration of the intestinal mucosae of mammals, Salmonella serotypes are confronted by an effective barrier to further spread, namely macrophages that line the lymphatic

sinuses of regional lymph nodes. In mammals this host defense mechanism can successfully limit bacterial expansion to the intestine, the gut-associated lymphoid tissue, and the mesenteric lymph node. For instance, humans infected with nontyphoidal *Salmonella* serotypes usually develop an acute gastroenteritis, but in only 1 to 7% of clinical cases do bacteria manage to pass through the mesenteric lymph node and cause bacteremia (19). Therefore, in order to produce systemic disease in warm-blooded hosts, microbial intruders must be able to breach the local defense formed by macrophages of regional lymph nodes.

The capability of S. bongori or S. enterica subspecies II, IIIa, IIIb, IV, VI, and VII serotypes to survive in macrophages of poikilothermic animals has not been studied. Experimental oral exposure of snakes, turtles, or lizards to isolates of S. enterica subspecies I, II, or III resulted in all cases in no overt signs of disease and no colonization of organs other than the intestinal tract (35-37, 111). It has therefore been speculated that Salmonella serotypes evolved in the alimentary tract of reptiles, where they developed from pathogens into commensal organisms (87, 102). In contrast, S. enterica subspecies I serotypes frequently colonize internal organs of warm-blooded animals and the ability to survive and multiply in cells of the reticuloendothelial system correlates with their capability to cause systemic disease in these hosts (8, 45). Since macrophages from distinct homeothermic animal species differ greatly in their abilities to neutralize a particular S. enterica serotype, adaptation to a new host may require adaptation to its mononuclear phagocytes. For instance, the human-adapted S. enterica serotype Typhi is able to survive in vitro in human macrophages but not in murine macrophages, whereas S. enterica serotype Typhimurium, which causes a systemic disease in mice, survives well in vitro in murine macrophages but not in human macrophages (117). Furthermore, these two S. enterica

Host species	Disease	<i>S. enterica</i> subspecies I serotype(s) most frequently encountered	Most susceptible age groups	Typical symptoms or sign(s) of disease	Reference
Humans	Salmonella enteritis Typhoid fever Paratyphoid fever	Typhimurium, Enteritidis Typhi <sup>c</sup> Sendai; Paratyphi A, B, and C <sup>c</sup>	Children (<4 yr) Children and adults Children and adults	Diarrhea, dysentery, fever Septicemia, fever <sup>a</sup> Septicemia, fever <sup>a</sup>	78 78 78
Cattle	Salmonellosis	Typhimurium Dublin	Calves (<8 wk) Calves and adult cattle	Diarrhea, dysentery, septicemia, fever Diarrhea, dysentery, septicemia, abortion, fever	104 99, 104
Poultry	Pullorum disease Fowl typhoid	Pullorum <sup>c,d</sup> Gallinarum <sup>c,d</sup>	Newly hatched birds Growing stock and	Diarrhea, septicemia Diarrhea, comb discoloration, septicemia	27 27
	Avian paratyphoid	Enteritidis, Typhimurium	Newly hatched birds	Diarrhea, septicemia	27
Sheep	Salmonellosis	Abortusovis <sup>c</sup> Typhimurium	Adult sheep Lambs Lambs	Septicemia, abortion, vaginal discharge Diarrhea, dysentery, septicemia Diarrhea, dysentery, septicemia	90 90 90
Pigs	Pig paratyphoid Salmonellosis Chronic paratyphoid	Choleraesuis <sup>c</sup> Typhimurium Typhisuis	Weaned and adult pigs Weaned pigs (<4 mo)	Skin discoloration, septicemia, fever <sup>b</sup> Diarrhea Intermittent diarrhea	106 106 7
Horses	Salmonellosis	Abortusequi <sup>c</sup> Typhimurium	Adult horses Foals Foals	Septicemia, abortion Diarrhea, septicemia Diarrhea, septicemia	110 110 110
Wild rodents	Murine typhoid	Typhimurium, Enteritidis		Septicemia, fever	28

TABLE 1. Diseases caused by Salmonella subspecies I serotypes in humans and higher vertebrates

<sup>a</sup> Diarrhea develops only in about one third of typhoid fever patients and usually several days after the onset of fever.

<sup>b</sup> Diarrhea is not a typical sign of pig paratyphoid but may develop by the third or fourth day of disease.

<sup>c</sup> These serotypes have been most frequently associated with illness in the preantibiotic era but are now rare or have been eradicated in most developed countries. <sup>d</sup> Gallinarum and Pullorum are considered biotypes that belong to the same serotype.

serotypes differ in their mechanism of bacterial internalization and their intracellular trafficking in human and mouse mononuclear phagocytes (3, 60, 61, 91). These data imply that the ability of *S. enterica* serotypes to cause systemic disease is directly related to the capability to withstand an assault by the macrophages of a given host. Thus it appears that mononuclear phagocytes are an important barrier that restricts the host range of *Salmonella* serotypes.

#### COEVOLUTIONARY COURSE TOWARD INCREASED VIRULENCE

Although *S. enterica* subspecies I contains 1,367 different serotypes (93), only one or a few are associated with the majority of cases of illness in a particular avian or mammalian species (Table 1). These serotypes show different degrees of host adaptation. Pathogens that lack host specificity, such as *S. enterica* serotype Typhimurium and *S. enterica* serotype Enteritidis, tend to be more frequently associated with disease in young animals than in adults, suggesting that they are not optimally adapted to cope with a fully mature immune system. Serotypes that are host specific, on the other hand, have acquired the ability to breach defense mechanisms in full-grown animals, as shown by their association, at similar rates, with illness in all age groups (Table 1). Furthermore, host-specific serotypes tend to be more virulent, as illustrated by the fact that they cause higher mortality rates.

It has been proposed that vertebrate pathogens which cause high mortality rates are new arrivals that are not well adapted to their hosts, because the death of an animal would destroy their habitat, thereby reducing both their transmissibility and fitness. According to this theory, these pathogens will evolve over time into less-virulent forms, which will reflect better adaptation to their hosts (26). However, in *S. enterica* subspecies I evolution has apparently driven host-pathogen interactions in the opposite direction, since strongly host-adapted serotypes tend to cause higher mortality rates than those with a broad host range. For instance, *S. enterica* serotype Typhi exhibits host specificity for humans in whom it causes typhoid fever, a disease which results in 12 to 32% mortality (78). In contrast, death occurs in less than 0.5% of cases of human illness caused by zoonotic *Salmonella* serotypes, such as *S. enterica* serotype Typhimurium or *S. enterica* serotype Enteritidis (78). What selective forces were responsible for the increase in virulence that accompanied the adaptation of *Salmonella* serotypes to humans?

On the basis of a mathematical model, Anderson and May (5) predicted that if transmissibility in a particular host-pathogen association is linked to a high level of virulence, then the coevolutionary course will be toward high virulence. During coevolution of *Salmonella* serotypes and their human host such a link between transmissibility and high virulence may have been provided by the ability to develop chronic carriage. A chronic carrier has the potential to infect a large number of individuals, thereby increasing transmissibility. Chronic carriage develops more frequently following systemic infection, such as that caused by the host-adapted *S. enterica* serotype Typhi. For instance, 1 to 4% of patients that recover from typhoid fever but only 0.2 to 0.6% of patients with nontyphoidal salmonellosis become chronic carriers (56, 81). Thus, the link between systemic infection and increased transmissibility



FIG. 2. Phylogenetic distribution of virulence genes and virulence of different *S. enterica* subspecies I serotypes in mice and humans. The bacterial isolates shown in this figure have been described previously (23). The left side shows a dendrogram reflecting the phylogenetic relatedness of these strains as reported by Selander and coworkers (23). ET, enzyme type determined by multilocus enzyme electrophoresis (23). The distribution of the *spv* (*Salmonella* plasmid virulence), *lpf* (long polar fimbriae), *sef* (*S. enterica* serotype Entertitidis fimbriae), *agf* (thin aggregative fimbriae), *fim* (type 1 fimbriae), and *pef* (plasmid-encoded fimbriae) operons among these isolates has been determined previously (12, 17). The presence in these strains of sequences related to *viaB* (Vi capsular antigen) is described elsewhere (107). Virulence of these isolates in humans or mice has been reported recently (17, 107). a, although missing in this strain, virulence plasmids are present in most isolates of this serotype.

resulting from chronic carriage may have exerted evolutionary pressure on the development of a high degree of virulence during the adaptation of typhoidal *Salmonella* serotypes (causing typhoid or paratyphoid fever in humans) to the human host.

#### THE VIRULENCE PLASMID IS IMPORTANT FOR SYSTEMIC DISEASE CAUSED BY NONTYPHOIDAL S. ENTERICA SEROTYPES

Although nontyphoidal S. enterica serotypes do not cause enteric fever in humans, some do cause similar systemic infections in warm-blooded animals. In S. enterica subspecies I, the spv (Salmonella plasmid virulence [54]) operon is found in only a few serotypes, including S. enterica serotypes Typhimurium, Enteritidis, Choleraesuis, Gallinarum/Pullorum, Abortusovis, Paratyphi C, and Dublin (17, 75, 94, 95, 103, 122) (Fig. 2). Interestingly, these serotypes are also among those most frequently associated with disease in homeothermic vertebrates (Table 1), suggesting a role for spv during infection of these hosts. The spv operon is required for the systemic phase of disease caused by S. enterica serotype Choleraesuis in pigs (34), S. enterica serotype Gallinarum/Pullorum in chickens (9, 10), S. enterica serotype Dublin in cattle (72, 118), and S. enterica serotypes Typhimurium and Enteritidis in mice (53, 63, 83). Epidemiological evidence provides support for the idea that the *spv* operon is also important for the pathogenesis of extra intestinal infections associated with nontyphoidal Salmonella serotypes in humans (46). Therefore, in a first approximation it appears that the *spv* operon is required for systemic infections caused by nontyphoidal serotypes in warm-blooded animals, including humans (9, 10, 34, 53, 63, 72, 83, 103, 118). Mechanisms for systemic infection that are *spv*-dependent may, however, not be restricted to nontyphoidal serotypes since the *spv* operon is present in *S. enterica* serotype Paratyphi C (17, 94). Typhoidal serotypes which lack the *spv* genes, such as *S. enterica* serotypes Typhi, Paratyphi A, Paratyphi B, and Sendai, produce enteric fever by an *spv*-independent mechanism. The virulence determinants responsible have not yet been identified (94, 100, 122).

Although all members of S. enterica subspecies I that are able to produce lethal infection in mice possess the *spv* operon, its introduction into S. enterica serotype Typhi does not confer mouse virulence to this host-restricted pathogen (103). Furthermore, the spv operon is present in most isolates of S. enterica serotype Gallinarum/Pullorum, which does not cause disease in mice. These data suggest that host-restricted S. enterica serotypes lack additional virulence factors that are encoded on the chromosome of serotypes with a broad host range. Furthermore, in addition to its presence in several serotypes of S. *enterica* subspecies I, the spv operon has also been detected in some isolates of S. bongori and S. enterica subspecies IIIa and IV (98), and thus the expansion in host range observed for subspecies I cannot be explained merely by the presence of these virulence genes. Therefore, the ability to cause systemic disease in a warm-blooded host is apparently a complex phenotype that cannot be attributed to acquisition of a single virulence determinant. It has been shown recently by subtractive hybridization analysis that 20% of the genome of S. enterica serotype Typhimurium is not present in S. enterica serotype Typhi and vice versa (67). By comparison with the similarly sized E. coli genome, the genome of S. enterica serotype Typhimurium can be estimated to contain approximately 4,400 genes. Serotype-specific DNA may thus encode as many as 880 genes, some of which may contribute to determining the



FIG. 3. Linear genetic maps of virulence plasmids of *S. enterica* subspecies I serotypes. Plasmids were linearized at their conserved *par* (partitioning) region (29, 113, 114). Regions of homology shared among plasmids are shown as closed bars, which are positioned according to their locations on the *S. enterica* serotype Typhimurium virulence plasmid. The distribution among virulence plasmids of *spv* (*Salmonella* plasmid virulence) (6, 66, 122), *repC* (replicon C) (113), *pef* (plasmid encoded fimbriae) (12, 47), *repB* (replicon B) (113), and the *tra* (conjugative plasmid transfer) region (66, 101) have been reported recently. Areas of the plasmid of *S. enterica* serotype Typhimurium have been described previously and are shown as open bars (68, 80). The presence on the *S. enterica* serotype Dublin plasmid of sequences with homology to *feaH* and *feaI*, two genes involved in biosynthesis of K88 fimbriae in *E. coli*, have been reported serotype Start and serotype Start and serotype Typhimurium plasmid are reported elsewhere (25, 80, 101). Dashed lines indicate the position of two DNA regions, namely the *pef* and *spv* operons, which are shown in more detail above the *S. enterica* serotype Typhimurium plasmid (1, 47, 52). The positions of genes are indicated by arrows.

host range. A combination of some of these genes, which are likely to be distributed throughout the chromosome, might determine the ability to infect a particular host. This idea may explain why attempts to extend the host range of host-restricted *S. enterica* serotypes by transferring small segments of the genome from a broad-host-range serotype have not succeeded.

Although the distribution of the spv operon has not been determined for all phylogenetic lineages of the genus Salmo*nella*, its localization on large serotype-specific plasmids in S. enterica subspecies I suggests that this operon was obtained by horizontal gene transfer. Additional regions of virulence plasmids, such as the origin of transfer and the tra genes, hint at conjugation as a possible mechanism by which the spv operon was horizontally transmitted (25, 89, 101). Evidence for the presence of the spv operon on a predecessor plasmid that gave rise to the virulence plasmids found among extant S. enterica serotypes comes from analysis of their patterns of homology. For instance, the virulence plasmids of S. enterica serotypes Enteritidis and Choleraesuis could be seen as variants of the virulence plasmid of S. enterica serotype Typhimurium that were generated by deletion events which occurred during their divergence from a common predecessor (Fig. 3) (25, 80, 101, 120). Similarly, a 25-kb DNA region containing the spv operon is conserved between virulence plasmids of S. enterica serotypes Dublin and Typhimurium, suggesting a common ancestry (18, 68). But where did the virulence plasmids of Salmonella serotypes originate?

The genes *finO*, *traY*, and *repA* are present on the *E. coli* F plasmid and the virulence plasmids of *S. enterica* serotypes Enteritidis and Typhimurium (101). Comparison of the nucleotide sequences determined for the *finO*, *traY*, and *repA* genes of several natural isolates of *E. coli* and *S. enterica* revealed similar, average, pairwise differences within and between species (21). These data are indicative of one or more recent horizontal transfer events between *E. coli* and *S. enterica* and suggest that during formation of *S. enterica* subspecies I the *spv* 

operon was obtained from a F-like plasmid pool (21) shared with ancestral *E. coli* isolates and possibly other organisms. As proposed for ancestral *E. coli* strains (87), some of the organisms that contributed to this F-like plasmid pool may have been associated with higher vertebrates long before host adaptations to mammals or birds developed in *S. enterica* subspecies I. For example, the presence on *S. enterica* virulence plasmids of *tlpA*, a gene encoding a regulatory protein that is sensitive to changes from 28°C to physiological temperatures of 37 to 42°C, implies that the donor was already able to sense temperature differences encountered upon entry into warmblooded vertebrates (58). The identity of this ancestral virulence plasmid donor organism, if extant, remains to be discovered.

#### ADHESION TO THE MUCOSAL SURFACE AS A MECHANISM FOR HOST ADAPTATION

Salmonella serotypes initiate infection by attaching to the intestinal mucosa of the host. Recent evidence suggests that the recognition of intestinal surfaces by adhesins may contribute to the host adaptation of Salmonella serotypes. A nonfimbrial adhesin which may play a role in host adaptation is encoded by *invH*, a gene which is present within SPI 1 in S. enterica (22). The idea that invH is involved in adherence rests on the finding that mutational inactivation reduces both attachment to and invasion of cultured epithelial cells by S. enterica serotypes (4). In contrast, mutations in invA and invE, two genes that are part of the type III secretory apparatus encoded on SPI 1, render S. enterica serotypes noninvasive but have no effect on adhesion to cultured epithelial cells (48, 50). The observation that SPI 1-mediated adhesion and invasion are independent events which can be genetically separated suggests that entry of S. enterica into epithelial cells is a twostep process and that InvH may function as an adhesin which mediates an initial attachment step required for invasion (4, 20, 44).

A comparison of the effect on virulence of a mutation in invH with those in genes encoding the invasion-associated type III export apparatus illustrates the importance of adherence during interaction with different hosts. S. enterica serotype Dublin secretes SopB, a virulence factor which is required for eliciting inflammation and fluid secretion in calves, and its translocation into host cells can be abolished by a polar mutation in sipB, a gene located on SPI 1 (49, 121). Similarly, mutations in invA, invB, invC, and sipC render S. enterica serotypes Enteritidis and Typhimurium avirulent in newly hatched chicks (96, 115). Thus, the type III exporter encoded by SPI 1 is apparently required for S. enterica virulence in both chickens and cattle. However, a mutation in the S. enterica serotype Typhimurium invH gene has no effect on oral virulence or colonization of chicks but significantly reduces the severity of enteritis in calves, suggesting a role of the encoded adhesin in adaptation to bovine but not avian hosts (96, 119, 120).

In chicks, the role of InvH in mediating an initial attachment step required for invasion is likely fulfilled by an alternate adhesin. The various fimbrial operons present in S. enterica are possible candidates for encoding such alternate attachment factors involved in host recognition (15, 32). Like binding mediated by InvH, attachment through fimbrial adhesins has been shown to affect entry of S. enterica serotype Typhimurium into some epithelial cell lines (4, 14, 43, 57, 64, 112). In a murine intestinal organ culture model, adhesins encoded by the pef (plasmid-encoded fimbriae) and lpf (long polar fimbriae) operons mediate tissue tropism of S. enterica serotype Typhimurium to villous small intestine and Peyer's patches, respectively (13, 16). These data suggest that adhesins help to select which epithelial surfaces are colonized by S. enterica in a given animal, a mechanism which is likely to be of importance for determining host range. Analysis of bacterial binding to the mucosa may, in some cases, be complicated by the large number of apparently redundant adhesins. For instance, in the mouse the finding that multiple fimbrial operons are required for full virulence of S. enterica serotype Typhimurium suggests that alternative attachment factors can compensate for the loss of a single adhesin (116).

The repertoire of adhesins expressed by a pathogen is thought to determine which structures are recognized and bound on the surface of intestinal epithelial cells. S. enterica serotype Typhi utilizes the cystic fibrosis transmembrane conductance regulator as a receptor for internalization by intestinal epithelial cells. In contrast, S. enterica serotype Typhimurium appears to use a different epithelial cell receptor for mucosal translocation, because mutations in the cystic fibrosis transmembrane conductance regulator do not reduce the invasiveness of this pathogen for epithelial cell lines or the intestinal wall of transgenic mice (92). This use of different receptors during infection implies the involvement of distinct adhesins which may contribute to differences in host range observed for S. enterica serotypes Typhi and Typhimurium. New combinations of adhesion determinants, generated through horizontal gene transfer and deletion events, may thus have contributed to shifts in host range during evolution of S. enterica serotypes (12).

#### THE HOST SPECIFICITY GAMBIT

The process of adapting to a host may involve not only acquisition of virulence determinants, such as adhesins or the virulence plasmid, but also loss of gene function. The *S. enterica* biotypes Gallinarum and Pullorum are both members of the same *S. enterica* subspecies I serotype (antigen formula

1,9,12:-:-) and show host specificity for poultry and aquatic birds. Multilocus enzyme electrophoresis and comparative sequence analysis revealed that S. enterica biotypes Gallinarum and Pullorum are closely related (71). It has been speculated that their lineage evolved from an S. enterica serotype Enteritidis-like ancestor that had a broad range of hosts, including birds. Since divergence from this ancestor, the ability both to mediate mannose-sensitive hemagglutination (MSHA) and to express flagella (and hence motility) was lost in the S. enterica serotype Gallinarum/Pullorum lineage as a result of point mutation (33, 71, 73, 88). Interestingly, strains of S. enterica serotype Enteritidis and S. enterica serotype Typhimurium that are isolated from cases of avian disease are also frequently nonmotile and lack MSHA (30, 38), suggesting that the niche occupied in birds may select against type 1 fimbriation and flagellation. Although it is arguably an advantage during infection of avian hosts, the simultaneous loss of motility and MSHA results in 100-fold-reduced virulence of S. enterica serotype Typhimurium in mice (74). It is tempting to speculate that the selection for point mutations in type 1 fimbrial and flagellar biosynthesis genes that occurred during the adaptation to avian hosts may account in part for the loss of virulence in mice in the S. enterica serotype Gallinarum/Pullorum lineage (Fig. 2). Thus, in S. enterica subspecies I more-complete adaptation to a particular vertebrate host appears to have evolved, in some cases, at the expense of virulence traits that were important for infection of a wider spectrum of animals, resulting in the loss of virulence for some species. The result of this adaptation was a narrowing of the host range and the development of host specificity.

However, the development of host specificity in *S. enterica* biotype Gallinarum cannot be explained by these point mutations alone. Other important changes which may have reduced virulence in mice have also been found. For instance, *S. enterica* serotype Gallinarum/Pullorum appears to be incapable of entering the follicle-associated epithelium of murine Peyer's patches, is internalized by murine macrophages by a mechanism different from that of *S. enterica* serotype Typhimurium, and is unable to survive and multiply in cells of the mouse reticuloendothelial system in vivo and in vitro (3, 8, 91). Thus, loss of virulence is a complex phenotype which involved multiple changes that accumulated in the lineage of *S. enterica* serotype Gallinarum/Pullorum during its evolution toward host specificity.

#### PERSPECTIVE

Despite the rapid progress in identifying some of the key elements of *S. enterica* virulence by studying the pathogenesis of murine typhoid, our knowledge about genes and mechanisms involved in the expression of host specificity is still very limited. Studies on the distribution of pathogenicity islands, fimbrial operons, and capsular biosynthesis genes among S. enterica serotypes suggest that during evolution, new combinations of virulence determinants arose through multiple horizontal transfer events (Fig. 2), a process which may have driven the development of host adaptation. In addition, deletion events and sequence divergence by point mutation were likely among the events which contributed to changes in the host ranges of S. enterica serotypes. Thus, adaptation to an animal species is a complex phenotype that doubtless involves a large number of gene products. It is not clear which genetic changes account for the adaptation to a particular mammalian or avian species, and it is likely that many of the responsible virulence mechanisms have not been identified to date. For instance, which virulence factors allowed S. enterica subspecies I to

breach the defense formed by regional lymph nodes and to spread to the systemic sites of infection? It is also not known which virulence factors are important during infections that remain localized in the intestine and mesenteric lymph node, such as the gastroenteritis caused by most *S. enterica* subspecies I serotypes in humans. Future research into the genetic basis of host adaptation will provide answers to these and other open questions regarding the pathogenesis of *S. enterica*.

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