Genetics of Lipopolysaccharide Biosynthesis in Enteric Bacteria
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INTRODUCTION

Lipopolysaccharides (LPS) are essential components of bacterial endotoxins which define many of the properties of host-parasite interactions, and they are the immunodominant antigens of most gram-negative bacteria. The early realization that these complex molecules had a unique structure to which the immunological and biological properties could be related led to a tremendous amount of interest and research effort in the 1950s and 1960s. As a result of this effort, by 1970 the LPS species produced by various Salmonella and Escherichia coli serovars had become the best-
studied complex polysaccharides in nature, in terms of both their structure and the biochemistry of their synthesis. Mutants played a central role in this research by providing conditional blocks in common precursors which allowed radioactive sugars and sugar nucleotides to be incorporated into LPS and by introducing defects at specific points in biosynthetic pathways to provide intermediate structures whose biochemical and immunochemical properties could be used to deduce these pathways.

Genetic studies accompanied the biochemical characterization of these mutants, and despite the difficulties imposed by the essential nature of genes involved in the early biosynthetic steps and the deleterious effects of some mutations which affect the later steps, a large number of genes were identified. A comprehensive and well-documented review of the progress in understanding the genetics of LPS synthesis in enteric bacteria was presented by Mäkelä and Stocker in 1984 (93). We have attempted to provide an update to that review, covering the period from 1983 to late 1992; except as required for clarity or emphasis, we have not cited articles appearing prior to 1982.

Three aspects of LPS research have been particularly important during the past decade. The first is the elucidation of the lipid A biosynthetic pathway and the use of conditional mutants and in vitro assays to identify the genes for this pathway. This has led to the discovery of a unique class of glucosamine-based lipids, which in turn has led to a better understanding of the biological effects of endotoxin. The second has been the extensive application of molecular genetics and recombinant DNA technology to the study of the biosynthesis of the oligosaccharide portion of the LPS molecule. This has solidly confirmed earlier work and led to the discovery of new genes whose presence was not anticipated on the basis of traditional genetic and biochemical analysis. Finally, the past decade has been a time in which the tremendous need of microorganisms to generate and maintain cell surface diversity has been recognized and studied. As an immunodominant and physically significant component of the cell surface, LPS has been shown to reflect this need for diversity by its genetic polymorphism and by chemical heterogeneity both at the level of species or serovars and among individual organisms.

In attempting to provide cohesive coverage of these three aspects of LPS research, we have generally limited the scope of our discussion to E. coli K-12, Shigella dysenteriae 1, and various Salmonella serovars, since our intent has been to provide insight into systems which provide the most complete picture rather than to present an overview of all of the enteric organisms which are presently being studied. Sadly, we must note that some interesting and potentially informative systems which were important in the early development of the enteric LPS field have received almost no attention in the past decade. Foremost among these is the phenomenon of the modification of O-antigen structure by temperate phages, which perhaps holds the best hope of allowing the synthesis and translocation of O antigen to be understood at the molecular level. Another neglected topic is the mechanism of form variation. We recommend the earlier review of Mäkelä and Stocker (93) for a stimulating discussion of these systems.

**OVERVIEW**

As is generally true for bacterial complex carbohydrate pathways which have been studied genetically, the genes for the core and O-antigen components of LPS are organized primarily into clusters of contiguous genes. In E. coli K-12 and Salmonella typhimurium these clusters are present in two regions of the chromosome, and in both regions they are near similar clusters of genes involved in capsular polysaccharide biosynthesis (Fig. 1). The rfa gene cluster (5), which encodes LPS core biosynthesis and O-antigen attachment, and the rfe-rff cluster, which includes genes for the biosynthesis of the capsular polysaccharide enterobacterial common antigen (ECA) and some serotypes of O antigen (36, 100), are located in the 81- to 85-min region flanking the origin of chromosomal replication. In E. coli K-12 the rfa cluster consists of 17 genes (74) and the rfe-rff cluster consists of about 11 or 12 genes (36). The 44- to 48-min region contains the rfb cluster, which is involved in O-antigen synthesis and attachment, and the cps cluster, which is involved in synthesis of type 1 capsules such as colanic acid (5). In S. typhimurium the rfb cluster contains 16 genes (65), but this number varies with the sugar composition and complexity of the O antigen. The exact size of the cps cluster is not known, but it is also likely to vary depending on the composition and complexity of the capsule. As seen from the outer circle of genes in Fig. 1, the general areas of the genome which contain these clusters are also rich in individual LPS genes which are not part of the clusters.

The clustering of genes involved in lipid A biosynthesis is not as striking. Several genes acting early in the pathway are located in a large, heterogeneous, complex operon at 4 min, which contains genes related to growth and macromolecular synthesis (166, 167), whereas other genes are scattered around the chromosome.

Figure 2 illustrates some of the general features of three LPS carbohydrate gene clusters for which the complete structure is known. Some shared features deserve comment. First, in all of these clusters only a few gene products have extensive potential transmembrane domains; most of the genes encode soluble proteins or proteins which are thought
to be bound to the inner surface of the cytoplasmic membrane. None of the gene products have signal sequences typical of proteins translocated into the periplasm or outer membrane.

Second, each of the clusters contains some genes encoding products which are strongly conserved across strain and species lines and other genes which are not strongly conserved. In the rfa cluster the strongly conserved group includes most of the genes, since the products of kdtA and rfaBCDFGPQYZ all exhibit identity of 70% or more in amino acid sequence between E. coli K-12 and S. typhimurium LT2 (65, 76, 124). In the rfb clusters there is somewhat more genetic polymorphism, but an example of such a strongly conserved region is the rfbBCDAE block, which encodes synthesis of a nucleotide sugar precursor common to many O antigens (65, 77). Another group of genes can be described as semiconserved, and these include the rfaIJ genes of S. typhimurium and E. coli (124) and some of the Salmonella rfb genes encoding sugar transferases (139). The proteins encoded by these genes are clearly related but show less sequence similarity (on the order of 25 to 50% identity) and can be considered to represent genes which have evolved to encode families of enzymes which catalyze similar reactions but recognize different substrates. Yet another group of genes encode proteins which have conserved structural features such as hydropathy profiles but show very little conservation of primary amino acid sequence across serovar or species lines (less than 25% identity). It is likely that these proteins have evolved for the specific recognition of structural features of the core or O antigen which are variable among different species and serovars, and in cases when their function is known they appear to be involved in late stages of the assembly or translocation of the polysaccharides. This category includes the genes rfaK, rfaL, rfbX, and rfc (21, 75, 77, 139).

In cases when comparable rfa or rfb clusters have been sequenced from different species or serovars, the products of the genes toward the center of the cluster are less strongly conserved whereas the products of the genes at the ends are more strongly conserved (75, 139). If each cluster is considered to be made up of a series of interchangeable modules, this arrangement of conserved regions flanking a nonconserved region may facilitate the exchange of modules within the clusters as a means of generating diversity while preserving basic biosynthetic functions. In several cases, it has been found that the nonconserved regions have a higher A+T content than the conserved regions, which in turn may have a higher A+T content than the flanking non-LPS genes or the chromosme of the organism as a whole (65, 75, 77). This suggests that these clusters may have arisen from a series of ancestral genetic exchanges involving very dissimilar organisms (139).

A simplified schematic overview of representative LPS structures and some of the genes involved in their synthesis is shown in Fig. 3. Only structures and gene functions which are well documented are shown in Fig. 3, and a number of substitutions which are variable or for which the site of attachment is not known have been omitted. A much more detailed picture of a complete S. typhimurium LPS molecule based on a combination of physical measurements and computer reconstruction has recently been presented (71). Readers should use caution and remember that the LPS can be quite heterogeneous and that the structures presented usually represent average or consensus structures rather than distinct molecular species.

FIG. 2. Schematic diagrams, drawn to the same scale, showing the rfa gene cluster of E. coli K-12 and the rfb gene clusters of S. dysenteriae 1 and S. typhimurium LT2 (65, 74, 77). Coding regions are drawn as blocks, with solid blocks indicating proteins which are hydrophobic or have clearly identified membrane-spanning domains. In the S. dysenteriae 1 rfb cluster, two genes lying outside of the cluster are shown to the right. Arrows indicate the direction of transcription. Brackets indicate approximate regions containing genes which are not conserved across species or serovar lines. The scale indicates kilobase coordinates for the Salmonella rfb genes (65).
GENETICS OF LIPID A SYNTHESIS

One of the most important aspects of LPS research in the past decade has been the elucidation in E. coli K-12 of the pathway of biosynthesis of a unique group of phospholipids based on glucosamine rather than glycerol, whose only known function is to serve as precursors of LPS. Biochemical details of the lipid A pathway have been covered extensively in recent reviews (127-131), so this section will be concerned primarily with the genetics of the pathway. A summary of the genes involved is provided in Table 1.

Acylation of UDP-N-Acetylglucosamine

Central role of intermediates. UDP-GlcNAc is a central intermediate in cellulase surface biosynthesis and serves as the common precursor of peptidoglycan and lipid A. It also plays an important role in surface polysaccharide biosynthesis. Addition of GlcNAc to the LPS core plays a key role in the attachment of O antigen in S. typhimurium and perhaps in E. coli (75). The transfer of GlcNAc phosphate from UDP-GlcNAc to the polypropenoid lipid carrier undecaprenol phosphate (referred to here as antigen carrier lipid or ACL) is also involved in initiating the synthesis of rfe-dependent O antigens and ECA (63, 102). Regulation of the synthesis of UDP-GlcNAc is not well studied, but it is noteworthy that glmS, the first gene involved in its synthesis from fructose 6-phosphate, is located in a cluster of genes adjacent to orfC at 84 min (5). R-3-Hydroxymyristoyl-acyl carrier protein (ACP), the other unique early precursor of lipid A, is also a common intermediate since it can be incorporated into both lipid A and glycerol-based phospholipids (132). The tightly interlocked network of common intermediates between the membrane lipid and polysaccharide pathways and the clustering of genes for the biosynthetic pathways together with other elements of macromolecular synthesis make it seem less than serendipitous that the first steps of lipid A biosynthesis were discovered during a search for genes involved in early steps of glycerophospholipid synthesis (164).

lpxA and macromolecular synthesis II operon. The first committed step in lipid A biosynthesis is the transfer of the acyl group from R-3-hydroxymyristoyl-ACP to the 3-OH group of UDP-GlcNAc. This acyltransferase exhibits a remarkable substrate specificity which precisely determines the lipid composition at this substitution site (2). The acyltransferase gene is encoded by the lpxA gene at 4 min on the
TABLE 1. Genes known to be involved in the synthesis of lipid A, listed in the order in which their products are thought to act

| Gene | Function of gene products | Example and reference |
|------|---------------------------|-----------------------|
| lpxA | Transfer of β-hydroxymyristic acid from ACP to 3-hydroxy group of UDP-GlcNac | E. coli (2) |
| lpxC | Gene formerly termed envA; removal of N-acetyl group from UDP-3-hydroxy-myristoyl-GlcNac | E. coli (126) |
| lpxD | Gene formerly termed fir; transfer of β-hydroxymyristic acid from ACP to 2-amino group of decacetylated GlcNac to form UDP-2,3-diacylmyristoyl-glucosamine | E. coli (126) |
| lpxB | Condensation of one molecule of UDP-2,3-diacylmyristoyl-glucosamine with one molecule of lipid X (same compound with UDP removed) to form UMP and lipid A disaccharide with four fatty acids and a single phosphate at the reducing end | E. coli (137) |

*E. coli* K-12 map, as shown by the isolation of a tight conditional lpxA mutant which was temperature sensitive for growth and exhibited temperature-sensitive acyltransferase activity (45). Like conditional mutants in the biosynthesis and attachment of 2-keto-3-deoxyoctulosonic acid KDO (143, 144), this mutant demonstrates the essential nature of the LPS pathway. The lpxA gene (28) is part of a cluster of 11 genes organized into a complex operon which has been termed the macromolecular synthesis II operon (99, 166). In addition to lpxA, this remarkable operon includes polC (dnaE) which encodes the α core subunit of DNA polymerase III (167); the mnh gene encoding RNase H; the lpxB and lpxD genes which also function in the lipid A biosynthetic pathway (35, 99, 126, 167); the cdc gene which encodes CDP-diglyceride synthetase, an essential step in biosynthesis of glycerophospholipids (132); the ompH gene (also referred to as skp or hlpA) which encodes a 16-kDa basic outer membrane protein (80); and an as yet unnamed gene encoding a β-hydroxymyristoyl thioester dehydratase (58, 82, 126). Both of these latter genes are noteworthy. OmpH protein binds both LPS and DNA, and was initially described independently as an LPS-binding protein and as the histone-like protein HNL-1 (1, 82). The main function of OmpH protein remains unknown, although a role has been suggested in protein translocation (165). The myristoyl dehydratase is a 17-kDa protein which shows extensive sequence similarity to the β-hydroxydecanoyl thioester dehydratase encoded by fabA, an important enzyme in unsaturated fatty acid biosynthesis (34). The gene for the myristoyl dehydratase is located between lpxA and lpxD (see below), which encode acyltransferases, and leads to the unique situation in which the genes for three enzymes that independently utilize the same substrate (R-3-hydroxy-myristoyl-ACP) are grouped together and presumably coregulated. The relevance of this regulatory scheme is indicated by the fact that missense mutations in the myristoyl dehydratase gene represented one class of temperature-resistant suppressors of the temperature-sensitive lpxA2 allele (128).

The next steps in the lipid A pathway involve N-deacetylation of the O-acylated UDP-GlcNac and the transfer of a second acyl group from R-3-hydroxy-myristoyl-ACP to the amino group to form UDP-2,3-diacylglicosamine, which is the precursor of the nonreducing end of lipid A. Removal of UDP by either of two different pyrophosphatases results in 2,3-diacylglicosamine-1-phosphate, which is otherwise known as lipid X (128, 129). Historically, it was the accumulation of lipid X in mutants defective in lipid A synthesis which provided one of the first clues leading to elucidation of the lipid A biosynthetic pathway (164).

Very recently, the two genes involved in N-acylation have been identified (126). The N-deacetylase is encoded by the envA gene (5), and it has been suggested that this gene be renamed lpxC. The first mutations in the lpxC (envA) gene were identified on the basis of a defect in cell division which was accompanied by a decrease in plasmid- or chromosomally encoded resistance to drugs such as ampicillin (114). The lpxC (envA) gene is located in a cluster of genes (5) involved in cell division (ftsQAZ) and protein secretion (secA).

The N-acylation is carried out by the product of the fir gene, which has now been renamed lpxD (126). Mutations in lpxD (firA) were first selected on the basis of rifampin sensitivity and independently on the basis of temperature sensitivity and enhanced sensitivity to multiple hydrophobic drugs. These latter mutants were termed ssc in *S. typhimurium* and omsA in *E. coli* and have recently been shown to be allelic to lpxD (firA) (171). The outer membrane phenotype of these mutants is entirely consistent with the identification of LpxD protein as an enzyme involved in lipid A synthesis, rather than a component of the transcriptional machinery as previously suggested (38, 39). LpxD protein shares an unusual region of homology termed an isoleucine patch with LpxA protein and other acyl- and acetyltransferases including the CysE and LacA proteins (39).

**Synthesis and Modification of Lipid A Disaccharide**

**lpxB gene.** The reaction which forms the backbone of lipid A involves the condensation of one molecule of UDP-2,3-diacylglicosamine with one molecule of lipid X to form a β1,6-linked glucosamine disaccharide in which each sugar is substituted with two 3-hydroxyxymyristoyl groups and which carries a single phosphate residue on the 1 position of the reducing sugar. This condensation reaction is catalyzed by the product of the lpxB gene (35). This enzyme is found primarily in the cytosolic fraction and can be assayed in the absence of detergent, although lipid X is found primarily in the membrane and the disaccharide product becomes very hydrophobic (137). The next step in the pathway is addition of a 4'-monophosphate by a specific kinase, generating the rather symmetrical disaccharide with a phosphate at either end which is known as lipid IVα (128). This kinase reaction can be carried out in vitro with labeled ATP, and this provides a very effective method for the generation of radiolabeled precursors which can be used to study subsequent reactions (22).

These reactions lead to the first structure which exhibits some of the characteristic properties which define bacterial endotoxin. Although lipid IVα does not exhibit all of the properties characteristic of endotoxin, it is an agonist of endotoxin and clearly has specific biological properties (49). The ability to use lipid IVα as a precursor and to manipulate its synthesis opens exciting avenues in the development of antimicrobial agents and drugs designed to combat endotoxin shock. Although the structure of lipid A is clearly not the same among all gram-negative bacteria (97, 163), these reactions are probably widely distributed and conserved among many bacterial genera.
TABLE 2. Genes involved in synthesis of the LPS core, listed in the order in which their products are thought to act

| Gene(s) | Probable functions of gene product | Example(s) and reference(s) |
|---------|-----------------------------------|----------------------------|
| kdsA    | Synthesis of KDO-8-phosphate      | E. coli (180), S. typhimurium (144) |
| kdsB    | Synthesis of CMP-KDO from KDO-8-phosphate | E. coli (47) |
| kdsC   | Sequential addition of backbone KDO I and branch KDO II residues to lipid A; possible attachment of branch KDO III to KDO II at later stage† | E. coli (12) |
| rfaE    | Synthesis of ADP-heptose           | S. typhimurium (156) |
| rfaD    | Epimerization of ADP-heptose to L-glycerol-d-manno form | E. coli (29) |
| rfaC    | Addition of Hep I to KDO I of inner core | S. typhimurium (156) |
| rfaF    | Addition of Hep II to Hep I to complete inner-core backbone | S. typhimurium (178) |
| rfaH    | Positive regulator of rfa operon (rfaQGPSBIJYZK) and rfa operon of plasmid F; may act by transcription antitermination; also termed sfbB | E. coli (125) |
| rfaQ    | Addition of partial substituent (perhaps Hep III) to core † | E. coli (74) |
| rfaG    | Addition of first backbone hexose (Glc I) to Hep II | E. coli (118), S. typhimurium (31) |
| rfaP    | Required for transfer of phosphate from ATP to Hep I; may be involved in other core phosphorylation steps † | S. minnesota (106), E. coli (118) |
| tolC    | Required for detergent resistance and LPS cross-linking; may be involved in conversion of core PPEA to phosphate † | E. coli (118) |
| rfaB    | Addition of branch Gal to Glc I | E. coli (124), S. typhimurium (181) |
| rfaI    | Addition of second backbone hexose (Glc II in E. coli K-12 and Gal I in *Salmonella* spp.) to core | E. coli (124), S. typhimurium (24) |
| rfaJ    | Addition of third (terminal) backbone hexose (Glc III in E. coli K-12, Glc II in *Salmonella* spp.) to core | E. coli (124), S. typhimurium (24) |
| rfaY    | Appears to be required for rfaJ action †; exact function unknown | E. coli (124) |
| rfaK    | Addition of GlcNAc to terminal Glc II in *Salmonella* spp.; may add GlcNAc to inner core in E. coli K-12 † | E. coli (74), S. typhimurium (93) |
| rfaS    | Required for synthesis of LOS form of core but not for cores carrying O antigen; may be Rha transferase †; absent in S. typhimurium LT2 | E. coli (74), S. typhimurium (76) |
| rfaZ    | Required for synthesis of LOS form of core but not for cores carrying O antigen; exact function unknown; present in S. typhimurium LT2 but may not function † | E. coli (74), S. typhimurium (92) |
| rfbBCAD | Biosynthetic pathway for TDP-Rha; required for synthesis of LOS form of core | E. coli (78) |
| rfaL    | Required for addition of O antigen to core and in E. coli K-12 for rfa-dependent addition of GlcNAc to terminal Glc III | E. coli (74), S. typhimurium (93) |
| rfe     | GlcNAc-P-ACL transferase required for addition of GlcNAc to terminal Glc III of E. coli K-12 | E. coli (77) |

* Functions which are speculative or not well studied.

GENETICS OF LPS CORE

The arrangement of LPS genes into clusters and operons does not always parallel the biosynthetic pathway. For example, the attachment of the first two residues of KDO to the lipid A precursor IVₐ, which involves a gene in the rfa cluster, takes place before the final acylation of lipid A. Likewise, although the genes in the rfa cluster are primarily involved in synthesis and modification of the LPS core, there are genes in this cluster such as rfaK and rfaL which are intimately involved in the attachment of O antigen to the core. Although the majority of the known genes whose functions are involved exclusively in LPS core synthesis are located in the rfa cluster, there are also genes such as kdsA and rfaE located outside of the rfa cluster which are involved in biosynthesis of sugars unique to the core or which appear to exert a direct effect on core structure. These are also included in the sections below. However, genes involved in biosynthesis and activation of sugar residues which are shared among multiple pathways such as biosynthesis of peptidoglycan or glycogen (44) have not been included. A summary of the genes involved primarily in core synthesis and their proposed functions is given in Table 2.

Synthesis and Attachment of KDO

*kdsA* and *kdsB*. The genes involved in the biosynthesis of KDO are among those which are located outside of the rfa gene cluster. The gene kdsA encodes the KDO-8-phosphate synthetase which combines D-arabinose 5-phosphate and phosphoenolpyruvate to generate KDO-8-phosphate and is part of a short operon located at 27 min (5, 179, 180). Following removal of the phosphate, free KDO is activated to CMP-KDO by the CMP-KDO synthase encoded by the kdsB gene at 85 min (5, 47).

Conditional (temperature-sensitive) kdsA mutants of *S. typhimurium* exhibit a stasis in growth under nonpermissive conditions. Although the cessation in KDO synthesis occurred promptly after a shift to a nonpermissive temperature, the stasis appeared only after a lag of almost a full generation. Surprisingly, this stasis had no deleterious effect and the mutants were able to resume growth after a shift to a permissive temperature (143, 144). These findings have several important implications. First, they provided the most direct evidence that LPS containing KDO is absolutely required for some cell surface assembly or cell growth process. Second, they indicate that continuous synthesis of KDO is not required and that cells can continue to grow until a pool of some cellular component is depleted at a critical level. It is not clear whether the critical component is a pool of LPS intermediates in the cytoplasmic membrane or whether there is simply an excess of LPS in the outer membrane which is sufficient to meet the needs of the cell until it is diluted by growth to a limiting concentration. Finally, in the absence of KDO synthesis the rate of translocation of lipid IVₐ to the outer membrane is low. This is in contrast to heptose-deficient mutants blocked in a later step...
of core assembly, in which translocation of a defective form of LPS to the outer membrane appears to be efficient (113). This suggests that attachment of KDO, or some step in LPS completion which is coupled to or depends upon the attachment of KDO, may be a minimum requirement for efficient translocation of LPS to the outer membrane. It has been possible to demonstrate that there is some transport of precursors which lack KDO to the outer membrane, but this appears to occur at a much lower rate (115). One set of reactions which are known to be coupled to the attachment of KDO are the final steps in acylation of lipid A (23, 129, 172). Unfortunately, nothing is known about the genes involved in terminal acylation, and no mutants blocked in these steps have been described. Thus it has not been possible to establish whether there is a relationship between the terminal acylation of lipid A and efficient translocation of LPS to the outer membrane.

kdA. The kdaA gene is located near the right end of the rfa cluster (26, 154) and encodes a bifunctional enzyme (12) which sequentially transfers two molecules of CMP-KDO to the lipid A precursor lipid IVα. This generates both the backbone KDO residue (KDO I) which is linked α2,6' to the lipid A disaccharide and the first α2,4-linked branch KDO residue (KDO II). The KdaA protein has a hydrophobic sequence capable of serving as a membrane anchor (27), and the isolation of enzymatically active (blue) TpgA fusions at this site in kdaA indicates that this is a transmembrane segment (124). This is consistent with the observations that KdaA is capable of binding hydrophobic compounds and requires a detergent for solubilization (12).

The gene adjacent to kdaA encodes an 18-kDa polypeptide (27). This protein is not required for KDO transferase activity in vitro, and its function remains unknown. Present data indicate that kdaA and the gene for the 18-kDa polypeptide are transcribed as an operon, and that both gene products are essential for viability (126, 147).

It is generally thought that KDO II is substituted by an additional α2,4-linked KDO (KDO III) which is added at a late stage of core completion (142). It should be emphasized that this is very difficult to study because of the labile nature of KDO and its glycosidic linkages, and the actual extent of substitution by KDO III is not known. The gene encoding the hypothetical KDO III transferase has not been identified, and there is the possibility that KdaA is a trifunctional enzyme which also catalyzes the addition of KDO III. This suggestion is based on the properties of a KDO transferase encoded by the gseA gene of Chlamydia trachomatis (11). The KdaA protein shares extensive regions of homology with the GseA KDO transferase, which is clearly trifunctional. In E. coli K-12, the gseA gene can complement a conditional kdaA mutant, and the expression of gseA in E. coli results in the addition of three KDO residues to the LPS core (16). Moreover, E. coli extracts containing GseA protein are capable of transferring at least one additional KDO residue to LPS bearing the two KDO residues added by KdaA (11). When gseA is expressed in E. coli in vivo, the addition of a third KDO residue appears to block the further addition of heptose residues, and it is the presence of this KDO III residue on a molecule lacking heptose which results in the unique Chlamydia epitope which allowed the gseA gene to be cloned in E. coli K-12 (16). The inhibition of heptose attachment explains why in wild-type E. coli K-12 any additional KDO residues would have to be added after the partial completion of the heptose inner-core region. The addition of KDO III is not essential in Salmonella species, since gaiE mutants which lack core sugars distal to the first heptose produce LPS which contains only two KDO residues (84) and these strains grow normally.

Heptose Synthesis and Inner Core

rfaD and rfaE. Although the genetics of the complete pathway (142) has been proposed for the synthesis of 1-glycerol-D-manno-heptose from sedoheptulose-7-phosphate has not been established, the genes for the last two steps in the pathway are known. Activation of heptose occurs before the final epimerization step, and the rfaE gene of S. typhimurium (93) which is located outside of the rfa cluster at 64 min appears to be involved in ADP-heptose synthase activity. This is based on in vitro experiments in which the addition of heptose to KDO₂-lipid IV₅ by an extract from an rfaE mutant could be rescued by the addition of ADP-heptose (156).

The rfaD gene has been cloned and sequenced both in S. typhimurium (150) and in E. coli K-12 (120), and the cloned E. coli K-12 rfaD gene can efficiently complement S. typhimurium mutants (29, 150). The RfaD protein is strongly conserved between the two organisms, but it does not share homology with other Rfa proteins whose sequence is known. rfaD is located near the left end of the rfa cluster, where it is the first of a block of four genes (rfaDFCL) which make up a small operon (147). Mutants lacking rfaD are defective in the epimerization of ADP-D-glycero-D-manno-heptose to ADP-1-glycero-D-manno-heptose. Such mutants accumulate ADP-D-glycero-D-manno-heptose and synthesize defective LPS which lacks LD-heptose but contains a small amount of the incorrect DD form (29). LPS from an rfaD knockout insertion mutant appears heterogeneous on sodium dodecyl sulfate (SDS)-gels with a major band comigrating with heptose-deficient LPS and some minor, more slowly migrating species which are assumed to result from the incorrect incorporation of DD-heptose.

There is a gene encoding a 26-kDa protein adjacent to rfaD at the left end of the rfa cluster (134, 154). The 26-kDa protein product shares a region of homology with bacterial kinases, and insertions into this gene exhibit a detergent-sensitive phenotype, indicating that it is probably involved in LPS synthesis (133). Its exact function is unknown, but it is tempting to speculate that it might be involved in heptose phosphorylation.

rfaC and rfaF. The RfaC and RfaF proteins share regions of sequence homology (150), and both proteins share somewhat less abundant regions of homology with RfaQ and KdaA (26, 27, 119). On the basis of homology, these four proteins could be considered to belong to a family of related proteins, all presumably sugar transferases, which all may act in the lipid A inner-core region. Unlike KdaA protein, RfaQ, RfaF, and RfaC proteins do not have a significant transmembrane domain capable of serving as a membrane anchor. One can argue that a membrane-binding domain is more essential in KdaA protein, since the final steps of acylation of lipid A are not completed until after the addition of KDO (23, 172). The RfaC and RfaF proteins of S. typhimurium share strong homology (more than 70% identity) with the corresponding proteins of E. coli K-12, and a region near the C terminus of RfaC is nearly identical between the two organisms (75). The cloned rfaC gene of E. coli K-12 is capable of very efficiently complementing an rfaC mutation in S. typeimurium (74).

An extract from an rfaC mutant is defective in the addition of heptose to KDO₂-lipid IV₅ in vitro (156), and this supports earlier evidence (93) that rfaC might be the gene.
encoding the heptosyltransferase which adds the first α1,5-linked backbone heptose residue (Hep I) to KDO I. LPS from mutants lacking rfaF exhibits a migration on SDS-gels intermediate between that of completely heptose-deficient LPS from an rfaC mutant and hexose-deficient LPS from an rfaG mutant (117) and is intermediate in heptose content (178). These results, together with the homology between RfaG and RfaE proteins, suggest that RfaG and RfaF are both structural genes for heptosyltransferases and that RfaF is the transferase which adds the second α1,3-linked backbone heptose residue (Hep II) to the inner core (Fig. 3).

Heptose-deficient mutants have a very deleterious deep rough phenotype, and many of those which have been reported appear to be quite leaky in terms of heptose content. Thus it was of interest to determine the null phenotype of heptose-deficient mutants of E. coli K-12, since the possibility existed that a small amount of LPS containing heptose might be necessary for some essential cell function. Neither an Ω insertion in rfaC nor a deletion extending from a restriction site in rfaD to a restriction site in rfaG which removed almost all of the rfa cluster was lethal when crossed onto the chromosome (73). The mutants exhibited a very mucoid deep rough phenotype and produced a truncated LPS which exhibited a single band on SDS-gels and comigrated with LPS from an rfaC mutant of S. typhimurium. To test that the mucoid phenotype was due to the production of colanic acid and to show that mucoidity did not suppress a potentially lethal phenotype resulting from the heptose deficiency, the rfaC:Ω null mutation was transferred into a strain carrying a null cps: : Tn10 mutation. In this background, the rfaC null mutation exhibited a “crunchy” colony phenotype similar to that described for other deep rough mutants (118) and otherwise exhibited a typical deep rough phenotype. This rfaC cps double mutant was not temperature sensitive and grew over a range of 30 to 42°C (73).

Other genes involved in lipid A or inner-core modification.

Although many of the intermediates in the synthesis of the lipid A-inner-core region of enteric bacteria have been identified within the last decade, the exact structures of lipid A and the inner-core region of complete LPS synthesized by E. coli and S. typhimurium. There are several reasons for this. First, the amphipathic nature of intact LPS makes it insoluble in most useful solvents, and this prevents high-resolution 31P- or 1H-nuclear magnetic resonance analysis and limits analysis by electrospray or fast atom bombardment mass spectrometry. Second, many of the bonds involved in inner-core structures are labile and are broken by mild hydrolysis or deacylation procedures, which are required to convert the molecules to forms more amenable to study. Finally, there is the ever-present problem of heterogeneity, since not all molecules carry the same substituents and the nature of the substituents may change dramatically in response to environmental conditions. Since LPS molecules are relatively stable once synthesized, a culture may carry its entire history imprinted in the structures present in its LPS.

The modifications which are best characterized include a 4-aminoarabinose attached to the 1-phosphate of lipid A (158) and a phosphorylhexosamine (PEA) substituent attached to the 7 position of KDO (61). This PEA substituent is present in deep rough mutants (84) which are thought to contain only two KDO residues, so the PEA is almost certainly attached to KDO II. It is unclear whether additional substituents of PEA or pyrophosphorylhexosamine (PEA) are attached elsewhere in the inner core. At least one phosphate is always present at the 4' position of lipid A (158), but this may be modified further to PPEA (129, 158).

The pmrA gene at 95 min of the S. typhimurium map (151) affects the addition of 4-aminoarabinose to the 1-phosphate of lipid A, with the mutant phenotype being an increase in the percentage of molecules which carry the amino sugar. This mutation was selected on the basis of resistance to polymyxin (93), and it has been suggested that reduction in the negative charge of the lipid A directly reduces its ability to bind polymyxin. The exact nature of pmrA is not known; it may be a regulatory mutation or exert its effect indirectly (93).

It has been observed that the phenotype of deep rough strains carrying a deletion removing genes rfaGSPB1 is not stable, that these strains gradually acquire a reduced hypersensitivity to detergents and novobiocin, and that complete hypersensitivity could be restored by transducing the rfa deletion into a fresh background (118). This suggests that these strains are accumulating suppressor mutations unlinked to the rfa cluster, and analysis of these suppressors might yield new mutations of the pmrA type which affect inner core substituents.

Two interesting new genes which may affect inner-core structure were identified during a screen for genes which are necessary for growth of E. coli at elevated temperature. The htrB gene at 23.4 min encodes a basic polypeptide with a membrane-spanning domain (67, 69). Knockout mutations of htrB exhibit a complex phenotype, including reduced sensitivity to deoxycholate and inability to grow in rich media at temperatures above 32.5°C with lysis and morphological changes occurring above that temperature. The msbB gene at 40.5 min (70) was first identified as a multicopy suppressor of the temperature-sensitive phenotype of htrB. In an htrB+ background, msbB null mutants exhibit an increased resistance to deoxycholate but no growth phenotype. However, an htrB+ msbB double mutant became even more temperature sensitive than htrB alone and showed additional morphological changes. In an rfa+ background htrB did not affect the mobility of LPS on SDS-gels but resulted in a rather dramatic decrease in silver-staining intensity similar to that which is seen with LPS from heptose-deficient mutants. In a strain background carrying a deep rough rfaGSPB1 mutation, the htrB mutation still exhibited a temperature-sensitive phenotype and a decrease in LPS staining intensity (117). In contrast, the msbB mutation caused a significant alteration of LPS gel migration, shifting the bands to a faster migration equivalent to a loss of one or two sugars from the molecules. It did not affect the migration of the truncated LPS in a rfaGSPB1 background but caused a reduction in staining intensity similar to that seen with the htrB mutant (117). The reason for the reduction in staining intensity is not known, but it may reflect changes in the inner-core or lipid A substitution pattern which affect the generation of aldehyde groups by periodate cleavage.

As in the case of the pmrA mutants, the cause of the htrB and msbB effects is not clear and may reflect either direct or secondary effects. Recent studies of htrB indicate that it affects regulation of lipid synthesis (68), and it was suggested that it may be part of a link between growth regulation and the regulation of lipid and LPS synthesis. However, given the lack of knowledge concerning the genetics and the function of inner-core substituents and the regulation of their synthesis, these mutations warrant further investigation.
Genes of Core Completion

rfaG/BIJK family and hexose backbone. The genes rfaG/BIJK are involved in the sequential attachment of glucose and galactose residues to the inner core to provide the structural framework for the outer, or hexose, region of the core. The rfaK gene is involved in the addition of GlcNAc to the terminal core glucose in S. typhimurium and a substituent (possibly also GlcNAc) to a different core site in E. coli K-12. Since this substitution plays a somewhat different role, the functional aspects of rfaK are described in the next section. In E. coli K-12 the protein products of rfaG/BIJK share limited regions of sequence homology and, on a more subjective basis, appear to have a similar structure based on visual comparison of hydropathy plots (119, 124). There is direct evidence only for RfaL and RfaJ that the proteins are sugar transferases, and this evidence is not compelling (124). However, the similarity of these proteins, the lack of evidence for any other genes involved in the transferase functions, and the agreement of the properties predicted for these proteins with the properties of those sugar transferases which have been purified leave doubt that these gene products are the transferases. The predicted proteins are similar in size, and all lack significant hydrophobic regions which could function as membrane-binding domains. They do not have signal sequences, and they yield enzymatically active (blue) fusions on mutagenesis with Tn5acZ but only inactive (white) fusions with TnphoA (124). Their structure and the fact that they act on membrane-bound LPS precursors suggest that they are peripheral membrane proteins attached to the inner surface of the cytoplasmic membrane through their interaction with their LPS substrate and other membrane components.

A galactosyltransferase thought to be the product of either rfaL or rfaB and a glucose transferase thought to be the rfaG product have been extensively purified and characterized from S. typhimurium (142). Both the galactosyl- and glucosyltransferases were shown to form enzymatically active ternary complexes with acceptor LPS and phospholipid. These complexes were specific since the LPS had to have a structure which allowed it to function as an acceptor for the appropriate sugars and both enzymes showed a strong preference for phosphatidylethanolamine (PE) as the phospholipid. In addition, an increase in lateral pressure was observed in an LPS-phospholipid monolayer when the galactosyltransferase was added, indicating that the enzyme was capable of intercalation into the monolayer. This led to a "ducks-on-a-pond" model for sugar transferases (summarized in reference 142), in which the individual transferases were envisioned as floating randomly on the cytoplasmic surface of a fluid phospholipid bilayer and interacting with acceptor LPS in the bilayer by lateral diffusion. Although there is nothing in the structure of the transferases to preclude such a model, more recent genetic studies suggest that in vivo there may also be significant interaction between the transferases (124), and it may be more appropriate to consider the active unit in vivo as a more structured or stoichiometric complex of enzymes acting in sequential fashion on bound or laterally compartmentalized substrates. Such structured complexes could explain why DNA fragments containing one or a few rfa genes are sometimes very difficult to manipulate in multicopy plasmids, a problem which everyone studying this region seems to have encountered.

The rfaG gene of E. coli K-12 can efficiently complement rfaG mutations in S. typhimurium (31, 119). The gene has not yet been sequenced in S. typhimurium, but this result and the similarity of the inner-core structure between the two organisms (60, 142) suggest that their RfAG proteins are likely to exhibit significant homology. There is strong chemical evidence which indicates that in both Salmonella spp. and E. coli K-12, RfaG is involved in the transfer of glucose from UDP-glucose to Hep II (31, 118). Complementation studies of long rfa deletions and the phenotypes of nonpolar mutations in rfaP and rfaQ indicate that RfaG protein does not require the products of either of these flanking genes for its function (118, 124, 125). This clarifies previous results obtained with polar rfaQ::Tn10 mutations which exhibited an RfaG" phenotype (3) and older biochemical studies which suggested that rfaP acted before rfaG (93).

Mutants deleted for rfaG are nonmotile and completely lack flagella and type I pili, and this effect is independent of the presence of rfaP (118). The loss of flagella is also observed with mutants defective for galU, which is necessary for the synthesis of UDP-glucose, and this loss has been interpreted as being due to a defect in transcription of flagellar genes (79). This suggests that the cell has a system which is involved in regulating the expression of flagella (and perhaps pili or other surface components) and which is capable of sensing either the degree of completion of the LPS core or the presence of the protein with its substrate, UDP-glucose. In this context it should be noted that the secretion of TolC protein is also defective in a galU mutant (174). The consequences of the loss of TolC are described in more detail in the next section.

In E. coli K-12 the second backbone hexose is glucose, whereas in Salmonella spp. it is galactose (142). Despite the difference in sugar composition, the following gene assignment is used in both organisms: rfaB is involved in addition of the α1,6-linked branch galactose to glucose I (Glc I), rfaM is involved in addition of the second α1,3-linked backbone hexose (either glucose or galactose), and rfaG is involved in the addition of the terminal α1,2-linked glucose (termed Glc II in Salmonella spp. and Glc III in E. coli K-12). This replaces an earlier terminology in E. coli K-12 in which rfaM and rfaN were used instead of rfaM and rfaJ, respectively (3, 31). The E. coli K-12 gene designations were dropped when it became evident that the arrangement of the genes and the sequence of the proteins they encoded were very similar in both organisms (124, 154).

Since rfaB is involved in the attachment of a branch sugar, the phenotype of rfaB mutants is not obvious in either organism. Since this branch sugar is the only galactose in the E. coli K-12 outer core, it would be predicted that the phenotype of an rfaB mutation would be identical to the phenotype of a null Δgal mutation. This has now been confirmed (124). Somewhat surprisingly, the major LPS band seen in rfaB and Δgal null mutants comigrates with LPS which contains only one hexose and not with LPS from rfaM mutants, which contains two hexoses. This indicates that in E. coli K-12 the RfaB protein cannot efficiently add Glc II in the absence of the branch galactose, regardless of whether the defect is in rfaB or in synthesis of UDP-galactose. Both rfaB and Δgal mutants produce minor LPS bands which migrate as though they have larger cores, but these are absent in mutants in which rfaJ is also absent, and it has been suggested that the absence of the branch galactose on Glc I may cause RfaJ protein to mistake it for Glc II and inefficiently add a terminal glucose (124). This might also explain anomalous phase P1 sensitivity results (124, 148), since this phase is thought to recognize an unsubstituted glucose as one of its receptors.
S. typhimurium rfaB mutants which appear to be completely lacking the branch galactose are able to compete a substantial number of core molecules and even add some O antigen (181). This is consistent with the unique phage resistance phenotype of rfaB mutants (93); these are resistant to phage FO, which requires the GlcNAc substituent attached to the terminal Glc II, but remain somewhat sensitive to smooth-specific phages such as P22. This would suggest that the stringency of RfaI with respect to its requirement for the branch galactose is different between the two organisms. However, an alternative explanation is that the RfaB and RfaI proteins function as a complex and that the S. typhimurium rfaB mutants were missense mutants which produced proteins that lacked transferase activity but were still capable of forming a complex with RfaI (124). The complex would have to include UDP-galactose to explain the similar phenotypes of Δgal and rfaB null mutations in E. coli K-12. To resolve this discrepancy, there is clearly a need to reexamine the biochemical properties of these galactosyl- and glucosyltransferases now that defined mutations are available.

Despite the differences in core structure, DNA fragments carrying rfaIJ of E. coli K-12 expressed from either the physiological rfa promoter or fortuitous plasmid promoters can efficiently complement rfaI mutants of S. typhimurium as measured by completion of a core which allows attachment of O antigen (124). This indicates that the final steps of core completion and O antigen attachment in S. typhimurium are indifferent to whether the second backbone hexose residue is galactose or glucose.

In contrast, only an rfaIJ DNA fragment from E. coli K-12 which also carries the physiological promoter region for these genes can efficiently complement an rfaI mutant of S. typhimurium (124). The most convenient explanation is that S. typhimurium RfaI protein competes effectively with an inefficiently expressed E. coli K-12 RfaI protein and that this results in the synthesis of Salmonella cores containing galactose in the backbone, which cannot be extended by the E. coli K-12 RfaI transferase. This provides the evidence noted above that rfaI and rfaD are the structural genes for the hexosyltransferases. Assuming that this is correct, the change in substrate specificity of the RfaI and RfaJ transferases between E. coli K-12 and S. typhimurium must be due to relatively minor changes in protein structure, given the strong conservation of the sequence of these proteins between the two organisms (24, 124).

In E. coli K-12, the change in electrophoretic mobility of the LPS accompanying the rfaIJ-mediated completion of the core is much greater than can be accounted for by the addition of two hexose residues. This is also the stage in core completion at which heterogeneity becomes evident, as indicated by multiple LPS gel bands (124). This suggests that there are other modifications of the core (perhaps additional KDO, Hep, rhamnose [Rha] or PPEA substituents) whose addition is coupled to the completion of the hexose backbone. The genes involved in these modifications are not known, although it is likely that rfaY (154) is one such gene. The phenotype of defined rfaY mutations is not known, but complementation studies of deletions which include rfaY suggest that rfaY may be necessary for rfaI function (124). It is possible that some mutations in S. typhimurium identified as rfaI are actually in rfaY, since the genes are adjacent in the chromosome and mutations in either gene might have the same phenotype. The sequence of the C-terminal portion of RfaY protein is known in both S. typhimurium and E. coli K-12 (75, 92), and RfaY appears to be strongly conserved between these organisms.

rfaP, tolc and deep rough phenotype. The deep rough phenotype is a complex phenotype exhibited by a subset of rfa mutants. In addition to alterations in the structure of the LPS, the phenotype includes the following: (i) sensitivity to detergents and hydrophobic antibiotics such as actinomycin D and novobiocin; (ii) elevation of the phosphatidylethanolamine (PE) content of the outer membrane to the point where regions of phospholipid bilayer, which are absent in the wild-type outer membrane, can readily be demonstrated; (iii) a dramatic (>90%) reduction in porin proteins, including OmpF, OmpC, PhoE, the Lc porin of lambdoid phages, and LamB, together with an approximately 50% reduction in OmpA protein; (iv) resistance to the LPS-specific phase U3 and the OmpF-specific phase K20; and (v) a decrease in buoyant density of the outer membrane (3, 113, 118). Many (if not all) deep rough mutants also exhibit a mucoid phenotype as a result of the induction of production of a colanic acid capsular polysaccharide. In a deep rough ΔfgaGPSB1 mutant of E. coli K-12 which has been well studied, this induction of capsule synthesis remained temperature and Lon modulated and required the rcsC component of the rcsBC two-component regulatory system (118). Mucoidy has now always been reported as part of this phenotype, and this may reflect the contribution of other factors such as temperature and carbon source to the rather subjective observation of colony mucoidy (118). In E. coli K-12 strain backgrounds carrying null cpx::Trl10 mutations, deep rough mutants invariably exhibit a brittle, “crunchy” colony morphology and a strong tendency to autoagglutinate in liquid culture (118).

The genetics of the deep rough phenotype are now well understood. Mutations resulting in defects in the backbone heptose region of the inner core (rfaCDDEF) and rfaP mutations, which affect the efficiency of the heptose region, exhibit a deep rough phenotype (56, 93, 113). Strongly polar mutations in rfaQ and rfaG also result in a deep rough phenotype (3), but this is probably because they prevent expression of rfaP. Strains with nonpolar rfaQ null mutations are sensitive to phage U3 and resistant to novobiocin (74), and complementation of a ΔfgaGPSB1 mutation with a fragment carrying only rfaP+ restored sensitivity to phage U3 and the ability to grow on EMB agar, as well as substantially increasing resistance to novobiocin (118). These data indicate that the loss of rfaP or the inner-core heptose region on which rfaP acts results in the deep rough phenotype.

The exact function of rfaP and the mechanisms involved in the deep rough phenotype are less clear. Although loss of rfaP may reduce the efficiency of core completion, nonpolar rfaP mutants produce a heterogeneous LPS, indicating that they are still capable of a considerable amount of core completion and, in S. typhimurium, some O-antigen attachment (56, 118). Analysis of LPS from ΔfgaGPSB1 mutants complemented with DNA fragments containing rfaG and rfaP indicate that there are two distinct inner-core functions associated with rfaP+ (118). One of these is the attachment of a phosphoryl substituent (either phosphate or PPEA) to Hep I. The other is the attachment of a branch Hep III to the Hep II residue, a substitution which is present on about half of the molecules. Since all molecules are phosphorylated on Hep I, the phosphorylation has been considered to be the primary function of rfaP, but direct evidence to support this is lacking (93). The observation that the γ phosphate from ATP can be transferred to the Hep I residue of LPS from an RfaP- strain by an Rfa+ extract (106) indicates that a kinase
adds phosphate as the first step in the generation of the phosphoryl substituent, but unfortunately it does not indicate that rfaP encodes the kinase. One problem is that the RfaP protein shows no relationship to known kinases and does not contain a consensus nucleotide-binding site (119). Another is the specificity of the kinase. It is generally thought that another PPEA residue is added to the core late in completion (142), and both the detergent-sensitive phenotype and the phosphate content of the LPS core of rfaP mutants suggest that this additional phosphoryl substituent is also absent (3, 56). It seems unlikely that a single kinase would be involved in phosphorylation at two different sites. An interesting possibility which has not been tested is that the 26-kDa protein encoded by the gene adjacent to rfaDFC heptose cluster is the Hep I kinase and that the partial detergent-sensitive phenotype of null mutations in this gene results from the ability of one or more additional kinases which add phosphate at other sites to suppress the detergent sensitivity. The RfaP protein might be a common subunit which is required for several kinases, or it may have another function entirely to which the kinase activity is coupled.

The steps leading to the generation of PPEA are unclear. Kinetic evidence indicates that PE is the precursor of both the monophosphate and the ethanolamine of the phosphorylethanolamine moiety of PPEA (54), and it has been proposed that the PPEA substituent of LPS arises by transfer of the phosphorylethanolamine moiety from PE to phosphate on Hep I (and probably other sugars) with the resulting generation of diacylglycerol, which is recycled into the phospholipid pool by diacylglycerol kinase (132). However, diacylglycerol kinase in enteric bacteria has not been studied extensively, and the proposed transfer of the phosphorylethanolamine moiety from PE has not been demonstrated in vitro.

Although there are still gaps in our knowledge, enough is known about the function of rfaP to allow speculation on how its loss results in the various effects which make up the deep rough phenotype. A useful approach is to attempt to dissect these effects and analyze them separately, and one way of doing this involves null mutations in the tolC and pss genes.

The tolC gene at 66 min of the E. coli K-12 map encodes a minor outer membrane protein (105). tolC mutants exhibit a complex phenotype including tolerance to colicin E1, reduced expression of the OmpF and Lc porin proteins, a failure to secrete hemolysin (173), resistance to the LPS-specific phage U3 (3), and a sensitivity to detergents and hydrophobic antibiotics which is quantitatively and qualitatively identical to that of deep rough mutants. Unlike deep rough mutants, tolC mutants do not produce truncated or heterogeneous LPS, instead showing a slight sharpening of LPS bands on SDS-gels (152), and there is no alteration of the buoyant density of the outer membrane and no overall defect in expression of outer membrane proteins (3). The inhibition of OmpF protein expression is a regulatory effect and can be suppressed by inactivation of the micF gene (104). Interestingly, the recent study showing a defect in export of the TolC protein by strains with galU mutations (174) suggests that the tolC function may be affected by LPS structure. Although this complicates interpretation of studies of mutants with inner-core defects, it may explain some earlier indications that galU mutants might exhibit a partially deep rough phenotype.

An insertion in tolC has a profound effect on the properties of LPS. The purified TolC- LPS is much more easily dispersed in water than is TolC+ LPS, and a micellar dispersion of TolC- LPS is not flocculated by the addition of divalent cations. Preliminary characterization has been done on LPS from a tolC mutant by using the same strain background (ΔfaGPSBI complemented with a DNA fragment carrying rfaGP) and chromatographic techniques which were used to study the effect of rfaP (152). When the TolC- core oligosaccharide fraction was compared with that of Tol+ LPS by gel filtration, peak II, which includes cores containing Hep I-PPEA, was increased in the TolC- LPS and peak IV, which in Tol+ LPS consists primarily of cores containing Hep I-phosphate, was absent in the TolC- LPS. Although the exact composition of these core fractions has not been determined, these results and the lack of flocculation by divalent cations indicate that Hep I of the TolC- LPS may have only PPEA while Hep I of the TolC+ LPS has both phosphate and PPEA. These results suggest that TolC may be involved in the conversion of PPEA to phosphate after the LPS is exported to the outer membrane. Moreover, these results suggest that the phosphate residue on Hep I plays an important role in binding divalent or polyvalent cations (particularly in the absence of phosphorylated substituents added at later stages of core completion), a function which is critical for the lateral cross-linking of the outer leaflet of the outer membrane to prevent penetration by hydrophobic compounds (113), and that the presence of the bulky ethanolamine shields the phosphates to make them less effective in cross-linking. The role of TolC in generating chelating residues could explain results of studies with E. coli O111 strain 15 which showed that newly synthesized LPS was rapidly converted from an EDTA-nonreleasable form to an EDTA-releasable form after it was exported to the outer membrane (86, 111). The lack of a major effect of tolC mutations on outer membrane proteins or on the buoyant density of the outer membrane indicates that lack of the TolC function does not prevent normal translocation or assembly of major outer membrane proteins or result in a gross perturbation of the outer membrane PE content. An interesting prediction of this hypothesis is that organisms such as Neisseria spp., which appear to be naturally hypersensitive to detergents and hydrophobic compounds, would lack a TolC analog. LPS core oligosaccharides from some strains of Neisseria gonorrhoeae have been shown to lack free phosphate and contain only PPEA and PEA (46, 66, 122).

A null mutation has recently been isolated in pss, the gene encoding phosphatidylserine synthetase, a key enzyme in the pathway for the synthesis of PE (37). In addition to lacking PE, the major phospholipid of E. coli, the pss mutant should lack ethanolamine, since the only known pathway for ethanolamine biosynthesis in E. coli is that involving the conversion of phosphatidylserine to PE (37, 132). Although a very minor alternative pathway for the synthesis of PE cannot be strictly ruled out, the amount of PE synthesized in the mutant is less than 1% of the wild-type level, and this is far below the amount which would be necessary to supply even a single ethanolamine residue on each LPS molecule. Thus this pss mutant can be used to study the role of ethanolamine in LPS synthesis.

The pss mutant appears to have a normal amount of LPS in its outer membrane, but the LPS bands migrate significantly faster (117). This is what would be expected as the result of an increase in net negative charge and a loss of the bulky ethanolamine substituents. The pss mutant strain produced normal amounts of porin and OmpA protein and was not sensitive to detergents or novobiocin. These results indicate that addition of ethanolamine cannot be essential for
the translocation of LPS to the outer membrane and that ethanolamine is not the ligand which is required for assembly of the major outer membrane proteins. The requirement of the pss mutant for a very high level of Mg^{2+} (>100 mM) for growth (37) may be due to the difficulty of translocating LPS to the outer membrane when it lacks the bulky ethanolamine residue as a blocking group on critical phosphate residues. It is interesting in this context that Mg^{2+} can be replaced by nontransportable ions such as Ca^{2+}, indicating that the effect of the divalent cations is at the cell surface (37). Another interesting question with respect to the LPS of the pss mutant is the nature of the phosphorylated substituent present on KDO, since nothing is known about the mechanism of synthesis of PEA.

The results with the rfaP, tolC, and pss mutations suggest the following model to explain the phenotype of deep rough mutations. First, as noted above, the ability of divalent and polyvalent cations to cross-link LPS must play a critical role in resistance to detergents and hydrophobic agents. At minimum, this requirement can be satisfied by a phosphate substituent on Hep I, and this requires both RfaP for the synthesis of heptose-linked PPEA and TolC for the conversion of PPEA to free phosphate after export of the LPS to the outer membrane. Second, some substituent in the heptose region of the core is required for the LPS to serve as a ligand for major outer membrane proteins and to promote their translocation and/or folding in the outer membrane. It is possible that this requirement can be satisfied by either phosphate or PPEA attached to Hep I. Alternatively, it could be the branch Hep III or some other substituent on KDO or lipid A whose attachment depends on RfaP function. In any event, the accumulation of PE in the outer membrane and the accompanying decrease in buoyant density probably reflects a combination of two factors: decreased PE turnover due to the lack transfer of ethanolamine to LPS and a defect in the ability to maintain the phospholipid asymmetry of the outer membrane because of a reduction in LPS cross-linking and a lack of space-filling outer membrane proteins. Finally, cross-linked LPS or LPS containing phosphate as a substituent must be required for the LPS to be a receptor for phage U3. All of these features require the backbone heptose residues, which is why heptose-deficient mutants also exhibit a deep rough phenotype.

Core modification and O-antigen attachment. The R LPS produced by rfaL or his-rfb mutants of S. typhimurium LT2 runs as a rather broad band on SDS-gels even when the Tricine buffer system (83) is used. In contrast, the LPS of the R strain E. coli K-12 often exhibits several sharp bands on Tricine-SDS gels, and these can differ depending on the strain background (78, 125). In genetic stocks in which the his-rfb region is derived from the K-12 strain Y-10 (4), a group which includes such widely used strains as C600, AB1157, AB1133, and MC4100, the LPS runs as three closely spaced bands which for convenience are designated below as bands 1, 2, and 3 in order of increasing gel mobility (78). Strains which are more closely related to wild-type E. coli K-12 produce only bands 1 and 2. As will be described in the next section, the reason for these differences is a defect in the synthesis of TDP-rhamnose (TDP-Rha) in strain Y10 and its derivatives (78). Even though these bands are quite sharp, they are chemically heterogeneous. Their molecular weights are not entirely understood, but their gel migration clearly reflects a pattern of substitutions or alterations of the core due to the action of the rfe gene and the rfa and rfb gene clusters (74, 78, 124). Thus, the ability to resolve changes in these bands which accompany the introduction of defined mutations has provided a very useful tool for studying the roles of rfaK and rfaL and, as described in the next section, for studying the functions of rfaQ, rfaS, and rfaZ.

The phenotype of rfaK mutants has been well studied chemically in S. typhimurium (93). These mutants lack the partial α1,2-linked GlcNAc substituent on the terminal Glc II residue and are resistant to the phage Felix O-1 (FO). There is extensive evidence indicating that GlcNAc is an essential part of the receptor for this phage (88), although the fact that rfaB mutants can be selected by FO resistance is evidence that other core structures may also be necessary (93, 181). S. typhimurium rfaK mutants exhibit a defect in attachment of O antigen. The severity of this defect varies among different rfaK alleles (93), but in the tightest mutants no O antigen is detectable on LPS gels indicating that the requirement is quite stringent (74). In S. typhimurium there is almost no difference in gel migration between LPS from rfaK mutants and LPS from rfaL mutants (74).

The RfaK proteins of E. coli K-12 and S. typhimurium are similar in size and overall structure but share almost no significant regions of sequence homology (75). In fact, the S. typhimurium RfaK protein shares more homology with the RfaK protein of E. coli K-12 than with the RfaK protein of S. typhimurium (75, 119). The gel phenotype of an rfaK mutant is quite different in E. coli K-12 than in S. typhimurium (74). The E. coli K-12 rfaK mutant LPS lacks bands 1 and 2 which are seen in Rfa+ LPS and has only a single band which comigrates with band 3. This suggests that RfaK acts at an earlier stage of core completion in E. coli K-12 than in S. typhimurium. The difference between RfaK of these two organisms is further illustrated by their effect on phase sensitivity. E. coli K-12 is resistant to phase FO but becomes sensitive when it is transformed with a plasmid carrying the S. typhimurium rfaK gene (74). Rfa+ E. coli K-12 is also resistant to phase Br2, a phage which requires an inner-core structure for its receptor (136), but rfaK mutants of E. coli K-12 become sensitive to Br2. Conversely, an rfaK mutant of S. typhimurium is sensitive to Br2 but becomes resistant to Br2 when it is transformed with a plasmid carrying the E. coli K-12 rfaK gene. These results are interpreted as indicating that RfaK is a transferase which adds a sugar substituent to the core in both organisms but adds this substituent to a different site in E. coli K-12 from that in S. typhimurium (74). The ability to modify the phase sensitivity of each by introducing a plasmid carrying the rfaK gene of the other indicates that each organism produces a core capable of being modified by both RfaK enzymes.

The complementation experiments below provide the only evidence suggesting that the E. coli K-12 RfaK protein is also a GlcNAc transferase, since they indicate that the S. typhimurium and E. coli K-12 rfaK genes are functional homologs. The site(s) at which GlcNAc is attached to the K-12 core is not clearly known (74). One study reported that GlcNAc was attached in a β1,6-linkage to Glc III (64), and another reported GlcNAc to be in an α1,7 linkage to a Hep residue and detected only α linkages by nuclear magnetic resonance analysis of core oligosaccharides (60). In E. coli K-12, attachment of GlcNAc to an inner core Hep would be consistent with the phase Br2 sensitivity data (74).

The recent observation that band 1 seen in SDS-Tricine gels of E. coli K-12 LPS is absent in an rfe mutant (77) may eventually resolve the conflict about the site of attachment of GlcNAc. The rfe gene product transfers GlcNAc-phosphate to the dolichol analog ACL, a reaction which appears to initiate the synthesis of ECA and rfe-dependent O antigens
It has been suggested that band 1 results from the transfer of this GlcNAc from ACL to the core, a process which requires rfaL and probably rfbX (77), and this is supported by recent chemical studies of the K-12 core (41a). Thus it is likely that GlcNAc is attached to the K-12 core at two different sites, one attachment requiring only rfaK and the other a more complicated transfer which requires rfaK, rfaL, rfe, and rfbX.

In Salmonella spp. and in E. coli K-12, GlcNAc is considered to be a partial substituent (93, 142). In E. coli K-12, however, the RfaK protein appears to play a very significant role in core completion since the LPS species which accumulates in an rfaK mutant is smaller than the bulk of the LPS molecules with completed cores. Since the GlcNAc content of the completed core is sometimes quite low, this raises the question of whether GlcNAc is added only to a subset of cores or, alternatively, is added to all of the cores and then removed from some of them during subsequent steps of core completion and preparation for O-antigen attachment. The initial studies with Salmonella spp., which were done to establish the linkage of O antigen to the core, involved small oligosaccharides purified after partial hydrolysis and do not address the question of the complete structure of the core to which the O antigen is attached (110). A study of phage F0 inactivation by LPS purified from various Salmonella strains indicated that LPS from S strains was more than an order of magnitude less effective at neutralization of the phage than was LPS from an R strain, and this was paralleled by the phage sensitivity of the bacteria from which the LPS was isolated (88). This difference was interpreted as being due to steric hindrance by the long O polysaccharide chains present in LPS from the S strains, but, alternatively, this difference could be due to the absence of GlcNAc from the LPS molecules which carry O antigen. Such a trimming coupled to maturation would be consistent with the oligosaccharide biosynthetic pathways found in eukaryotic organisms (43) and could explain the stringent requirement of O antigen attachment for a “partial” core substituent.

In S. typhimurium, rfaL mutants make no S LPS but produce complete O-antigen side chains attached to ACL and R LPS which has the chemical and serological properties expected of complete cores. On this basis, rfaL has been proposed to encode a component of the O-antigen ligase which is involved in the transfer of O antigen from ACL to an LPS acceptor (93). Recent data support this conclusion, although the role of rfaL appears somewhat broader than originally envisioned. In E. coli K-12 the LPS from rfaL mutants lacks band 1 and rfaL strains carrying the plasmid-encoded rfp gene from S. dysenteriae 1 fail to produce the Rfp-modified top band which replaces band 1 (74). Band 1 and the Rfp-modified band are now thought to represent transfer of one- and two-sugar partial O units from ACL to LPS (77). In some strains, type 1 capsular polysaccharides are attached to the LPS core, and this also requires rfaL (91).

A better description of the function of rfaL is that it is required for the transfer of a variety of polysaccharides from ACL to the LPS core, and on this basis it can be predicted that rfaL might also be necessary for synthesizing the LPS-linked form of ECA, which has been found in some rough strains (102). The RfaL protein has a structure predicted to be an integral membrane protein with 10 or more transmembrane segments (75, 92), and it has been proposed that it may serve as an organizer in the membrane which binds LPS and interacts with other proteins involved in late steps of core completion and O-antigen attachment (74).

Such a function does not preclude enzymatic activity as well, possible activities being the trimming of GlcNAc or the substituent added by RfaQ protein.

Although the RfaL proteins of S. typhimurium and E. coli K-12 have a nearly identical structure predicted on the basis of hydropathy, these proteins resemble the RfaK proteins in that they share almost no regions of significant sequence homology (75). Plasmids carrying either rfaL alone or rfaK plus rfaL from E. coli K-12 did not complement an S. typhimurium rfaL mutant, even though the E. coli K-12 rfaC gene which was present on the same plasmids efficiently complemented a S. typhimurium rfaC mutation (74). More complex cross-complementation was observed with rfaK. A plasmid carrying E. coli K-12 rfaK was capable of partially complementing the production of an O-antigen ladder by an rfaK mutant of S. typhimurium, but only when the plasmid also contained rfaL from E. coli K-12 (74). Taken together, these results suggest that RfaL has two distinct strain-specific functions: one requires an interaction with a specific modification of the core by RfaK, and for this rfaK and rfaL must be compatible. The other function requires a specific interaction between RfaL and another component which is probably also a core protein. Interestingly, when an rfaK mutant of S. typhimurium is complemented by a plasmid carrying both rfaK and rfaL from E. coli K-12, all of the bands in the resulting O-antigen ladder migrate more rapidly than the bands produced by an Rfa’ strain of S. typhimurium (74). The basis for this phenomenon is unknown.

rfaQSZ and core heterogeneity. The genes rfaQ, rfaS, and rfaZ were not detected in early mutant studies of Salmonella spp. and were found only when the rfa clusters of S. typhimurium LT2 and E. coli K-12 were sequenced (92, 119, 124, 154). The ability to manipulate core heterogeneity by introducing plasmids carrying genes for the synthesis of S. dysenteriae 1 O antigen has been very useful in studying the function of these rfa genes in E. coli K-12 (74).

rfaQ is the first gene in the 10-gene rfa operon which determines the outer core. The 5’ end of the gene has been sequenced in both E. coli K-12 and S. typhimurium, and the protein products are strongly conserved (more than 70% identity), indicating that the gene is similar in each organism (76). In E. coli K-12 the gene has been inactivated by nonpolar Tn10 insertions and by an in-frame deletion removing most of the coding sequence. These mutants are sensitive to phage C21, indicating that there has been a change in the core, but they are not sensitive to novobiocin or deoxycholate and produce normal amounts of the major outer membrane proteins. These mutations have a subtle but very interesting gel phenotype, which suggests that RfaQ is a transferase that adds a sugar or other substituent to the core (74). In a Y10-derived background such as strain AB1133, the LPS of an rfaQ mutant still exhibits three bands on gels, but band 3 exhibits a small but significant shift to a faster migration. In addition, band 1 is consistently less intense in an rfaQ mutant, as is the Rfp-modified band seen when an rfp’ plasmid is introduced into the strain. When a larger plasmid carrying the S. dysenteriae 1 rfp region is introduced into an rfaQ mutant, only a barely detectable amount of O antigen ladder is produced. These results suggest that the transfer of partial O units, O monomers, or polymeric O antigen from ACL to the core is inefficient in rfaQ mutants. However, unlike the shift observed for band 3, in an rfaQ mutant the migration of the small residual amounts of band 1, Rfp-modified band, and O-ladder bands is the same as in the rfa’ parent. The most convenient explanation for these observations is that RfaQ adds a
which dysenteriae which although a other mid, for the basis of its attaching required from and of the LOS band which first defined the LOS band as a unique form of R LPS and suggested that band 3 represented an earlier stage in its synthesis (74).

Additional evidence that the LOS band is the result of a unique biosynthetic pathway was provided by the properties of rfaS and rfaZ mutants (74). In Escherichia coli K-12 strains derived from Y10 which are unable to synthesize TDP-Rha, rfaS and rfaZ mutations have little or no phenotype. However, when rfaS and rfaZ mutations were introduced into strains with plasmids carrying the S. dysenteriae rfb genes that complemented the TDP-Rha defect, they exhibited a very substantial phenotype (74). The LOS band disappeared, and there was an accumulation of material in one or more bands which migrated faster than LOS. However, the intensity and migration of the O-antigen ladder were unaffected by the rfaS and rfaZ mutations, indicating that although these genes are required to produce the LOS band, they are not necessary for the synthesis of the LPS cores to which O antigen is attached.

E. coli K-12 strains such as W1485, W3110, and 679, which are closely related to the K-12 wild type (4), have a partial rfb cluster which contains all of the genes necessary for synthesis of TDP-Rha (77, 78, 112). These strains do not require rfb genes from other organisms to produce LOS, and the shift in their LPS migration following the introduction of an rfaQ mutation indicated that about half of band 2, the most abundant LPS band in these strains, was LOS. When these strains were transduced to rfaS or a deletion removing the rfb region was introduced, they lost the LOS band and acquired band 3, and their LPS gel phenotype became essentially identical to that of Y10-derived strains such as AB1133 (78). These results suggest that the addition of Rha to the core is an essential step in the conversion of band 3 to LOS and that rfaS encodes a function which is necessary for the addition of Rha to the core. Since rfaS mutants have a slight phenotype in Rha− strains (124, 125), it is unlikely that rfaS is the structural gene for a Rha transferase. Recent structural studies on E. coli K-12 LPS have shown that Rha is linked α1,5 to one of the inner-core KDO residues, probably KDO II (59, 60).

It is interesting to speculate that LOS may contain a terminal Hep IV residue attached to Glc III in addition to Rha in the inner core, since Hep IV was detected in strain W3100, which is Rha+ and is closely related to wild-type K-12 (62). In contrast, β-linked GlcNac was found attached to Glc III of strain OT99, a Rha− strain which is derived from Y10 (41a). The definition of LOS in E. coli K-12 is that it is a form of R LPS which is produced as an alternative to S LPS and is not a substrate for the attachment of O antigen (74). The molecular species which contain β-linked GlcNac attached to Glc III can be considered to be an intermediate stage in the synthesis of S LPS; as will be described in a later section, this GlcNac becomes the first sugar at the reducing end of the O antigen produced by E. coli K-12. S. dysenteriae 1 hybrid strains (41a, 77), and it is likely to be the first sugar at the reducing end of other rfe-dependent O antigens which are produced by E. coli. Since the Hep IV and β-linked GlcNac substitutions are both attached to the 6 position of Glc III (41a, 62), they must be mutually exclusive. Thus, the presence of Hep IV would be consistent with the definition of LOS stated above. However, we must emphasize that for the moment this is only speculation, and it would be a formidable task to confirm this speculation by proving chemically that Rha and Hep IV reside on the same LPS molecule. There are two reasons for this. First, as noted above, the LPS of Rha+ strains is heterogeneous, and probably no more than 50% of the LPS is of the LOS form (78). Thus, bulk analysis of the LPS cannot provide proof. Second, the bond between the branch KDO on which the Rha substituent is located and the remainder of the core which would contain the Hep IV is cleaved by the mildest conditions of hydrolysis which can be used to remove lipid A (60). Until an alternative technology such as enzymatic cleavage of LPS or sophisticated mass spectrometry capable of analyzing intact LPS molecules becomes available, there is no effective way to characterize LPS molecules which might contain both substituents.

S. typhimurium LT2 does not appear to produce an LOS band. This may be due in part to the fact that S. typhimurium LT2 lacks an rfaS gene, although the rfaP and rfaB genes which flank rfaS in E. coli K-12 are strongly conserved between E. coli K-12 and S. typhimurium LT2 (76). The region between rfaP and rfaB of S. typhimurium LT2 is what has been termed a gray hole (36), a region which does not encode a functional protein but appears to contain remnants of an ancestral gene. Like the rfaS gene of E. coli K-12, this gray-hole region in S. typhimurium LT2 between rfaP and rfaB has a remarkably high A+T content (76). The core bands at the bottom of an S. typhimurium LT2 O ladder are like those seen in an rfaZ mutant of E. coli K-12, and this suggests that rfaZ may also be nonfunctional in this organism.
Many questions remain about the LOS band besides its chemical structure. It is not known how widely a similar form of LPS is distributed among other enteric strains. A similar band has been seen in LPS of S isolates of E. coli (76) and in poultry strains of S. enteritidis (121). By analogy to other genes in the rfa and rfb clusters, the high A+T content of rfaS indicates that the sequence of the RfaS protein may not be conserved across strain or species lines. As this region of the rfa cluster is sequenced in other enteric organisms, it will be interesting to learn whether rfaS is conserved and whether the ability to produce a LOS has any effect on the ability of the organisms to survive or to colonize various host environments. It will also be interesting to learn whether organisms which produce a LOS have the ability to shift back and forth between the production of S LPS and LOS.

**Regulation of Core Synthesis**

Arrangement of transcriptional units. The first study of rfa transcriptional units was done by examining the complementation of mini-Tn10 insertions into rfaQ, rfaG, and rfaP by cloned fragments of various lengths. These fragments contained the promoter region between rfaQ and kdaA, the rfaQ gene, and one to six additional downstream genes. Regardless of the site of the mutation, the size and complexity of the LPS increased with the size of the complementing DNA fragments. This provided formal proof that the genes were arranged in an operon transcribed from a promoter downstream of rfaQ (3). There was evidence of expression of genes downstream of the Tn10 insertions, which was interpreted as indicating the presence of secondary promoters within the operon, but subsequent studies suggest that these effects may have been due to incomplete polarity of the insertions or promoter activity provided by Tn10. A more extensive study has been done by using polar ω cassettes to interrupt transcriptional units and Tn10 insertions to monitor gene expression (147). This study identified three operons: a pair of divergently transcribed operons at the right (kdaA) end and a single operon at the left (rfaD) end. The divergently transcribed operons at the right end were a short operon consisting of kdaA and the 18K gene and a long operon extending from rfaQ at least through rfaZ. The operon at the left end includes rfaDFCL.

There was no evidence for secondary promoters in the nine-gene span from rfaQ through rfaZ. However, the last reporter fusion in that series was located near the junction of rfaK and rfaZ, and it is not clear that it monitored rfaK expression. There is evidence in another study indicating a weak secondary promoter for rfaK: when a deletion of genes rfaGPSBI marked with a polar omega cassette is complemented with a longer fragment containing the downstream promoter region and the genes from rfaQ through rfaJ, the largest LPS species seen on a gel were larger than those produced by an rfaK mutant (124). It has not been possible to test for this promoter directly since no reporter fusions which are known to be located within rfaK have been isolated.

Recent primer extension studies have provided direct evidence that divergent transcripts originate in the untranslated region between rfaQ and kdaA (26). Two leftward and two rightward transcript start sites were identified. These sites do not overlap but are close enough to share common regulatory elements. As anticipated for highly regulated promoters, the regions upstream of these start sites do not contain strong promoter consensus sequences. Within the untranslated region there are four blocks ranging from 15 to 30 bp in which the sequence is nearly identical between E. coli K-12 and S. typhimurium LT2 (76), suggesting that this region contains important regulatory information.

Less is known about the untranslated region between the 26K gene and rfaD. However, there is one very remarkable aspect to this region. The rfaD gene exhibits heat shock regulation, and one laboratory cloned this gene while screening for novel heat shock-regulated genes (134). This is the first, and to our knowledge the only, case in which a gene in a carbohydrate biosynthetic pathway exhibits heat shock regulation. Of the many genes involved in the synthesis of core sugar nucleotides, rfaD is the only one included in the rfa cluster. Together with the unique heat shock regulation, this suggests that rfaD may play an important role in the regulation of core synthesis.

**Role of rfaH (sfrB).** The rfaH (sfrB) gene lies at 87 min, near ubiqu and fadA, and encodes an 18-kDa basic protein which shows no similarity to other regulatory proteins that have been studied. Despite several attempts, the protein has not been purified or visualized on gels, suggesting that it may be unstable. Insertion and amber mutations in rfaH have been reported (6, 32, 174), indicating that it is not essential, but mutants with a strong phenotype are sickly and readily accumulate suppressors. The phenotype is not consistent among various alleles, with the most severely affected alleles exhibiting both poor growth and a partial deep rough phenotype.

The first rfaH mutants were observed in S. typhimurium as a subclass of phage F0-resistant mutants (93). The inability of rfaH mutants to transfer labeled Gal from UDP-Gal to Gal-deficient LPS cores indicated that rfaH might encode a Gal transferase, which led to some confusion in the S. typhimurium LPS literature. In E. coli K-12, mutants designated sfrB were selected on the basis of lack of expression of the tra genes of plasmid F. These mutants were sensitive to phage C21 and were sometimes resistant to phage U3, which indicated an LPS defect (93). The confusion was resolved when it became apparent that rfaH and sfrB were allelic (93) and that RfaH protein was a positive regulator which regulated the expression of several enzymes encoded by the rfa cluster as well as the tra operon of F (19, 33, 125, 140).

In E. coli, SfrB was found to prevent the premature termination of tra operon transcripts at an internal terminator site, which led to the suggestion that it acts as a transcriptional antiterminator analogous to the n or Q proteins of λ (13). Despite considerable effort in several laboratories, this hypothesis remains unproven; however, there is now evidence from studies of the long rfa operon which provide support for an antitermination mechanism. Suppressors of rfaH map to rho, suggesting that RfaH acts at Rho-dependent termination sites (42). This is consistent with the observations that the ω cassettes which have Rho-independent terminators are completely polar when inserted into the rfa long operon, whereas Tnlac insertions, which should sometimes exhibit Rho-dependent polarity as a result of polypeptide chain termination, appear to be entirely nonpolar (125, 147). Reporter fusions inserted throughout the rfa long operon are regulated by rfaH (19, 125), and there is evidence for multiple termination sites (19). The locations of Tnlac fusions in the long rfa operon of E. coli K-12 have been determined by sequence analysis, and RfaH-dependent fusions have been obtained in all of the genes of the operon except rfaY and rfaK. The extent to which these fusions were regulated increased with the distance from the promoter, with a fusion in rfaQ showing only a 2- to 3-fold
increase in rfa + over rfa whereas a fusion in rfaZ showed a 10- to 15-fold increase (123). Experiments have been done to determine the similarity of the rfaH system to the antitermination systems of λ by examining the effect of various constructions on the rfaH dependence of a Tnlac insertion in rfaB (125). A pair of overlapping deletions extending from a HindIII site near the middle of rfaG to a ClaI site very near the 5' end of the rfaQ coding region had no effect on rfaH dependence (123). However, replacement of the region upstream from this ClaI site with a lac promoter resulted in expression which was dependent on a lac inducer but exhibited no rfaH dependence over a wide range of inducer concentrations (123). This suggests a model in which the action of RfaH is similar to that of the Q protein of λ, which modifies RNA polymerase while it transcribes a site located very near the promoter. Unfortunately, such a site for RfaH action has not yet been identified.

All of the evidence cited above is consistent with a model in which RfaH protein acts to prevent the termination of transcription at multiple Rho-dependent sites throughout the long rfa operon. Unlike RfaH protein, there is evidence that the enzymes of LPS synthesis may be quite stable (149). Since LPS is also quite stable and there is no way to store excess LPS or its precursors, there is a need to prevent expression of the genes for these enzymes at inappropriate times in the growth cycle. One can envision RfaH as acting as a kind of deadman's throttle, allowing synthesis of the LPS enzymes only when it is being actively synthesized itself. It will be interesting to know more about the factors which regulate the synthesis of RfaH protein.

Whether RfaH protein acts directly on operons other than the tra operon of F and the long rfa operon remains an open question. It has been reported that RfaH decreases termination in the rpl-rpoB intercistronic region (135) and, more recently, that RfaH was required for expression of a plasmid-encoded hemolysin (6). There is now evidence that the effect on hemolysin expression may be indirect and mediated by the effect of rfaH mutations on LPS core genes. The outer membrane form of ToIC protein is required for hemolysin secretion, and rfaH or galU mutations which result in a truncated LPS prevent the export of the ToIC protein to the outer membrane (174). This is somewhat similar to the effect of galU or rfaG mutations on the expression of flagella noted earlier. Considering the multiplicity of signaling mechanisms in bacteria, it would not be unreasonable for cells to monitor the synthesis of such an important component as LPS and to use this parameter to adjust the rate of synthesis of other cell surface components. Cells could accomplish this by sensing the completed LPS molecules in the cytoplasmic membrane, by sensing the rate of export or assembly of an outer membrane protein such as a porin which requires LPS as a ligand to promote correct folding, or by sensing the levels of key enzymes such as RfaG or RfaB together with their sugar nucleotide substrates. In proposing new targets for the action of RfaH, it is necessary to show that other mutations such as galU which affect LPS structure do not mimic its effect.

Effect of temperature. In enteric bacteria, temperature is a parameter which has a strong effect on the size distribution and abundance of various molecular species of LPS. This is most evident in E. coli K-12 strains carrying plasmid-encoded O-antigen genes. On the basis of staining intensity on gels, the median size of the long O-antigen polymers decreases by about 2 to 3 O units as temperature is increased from 30 to 42°C, whereas the number of total molecules which carry O units also tends to decrease (73). In K-12 strains without O-antigen plasmids the distribution of R LPS species also appears to be affected by temperature (125). In strains derived from strain Y10, which produce three major R bands, bands 1 and 3 decrease and band 2 increases as the temperature is increased. Some of this may be due to differential regulation of genes within the rfa long operon, perhaps at the level of translation, since the expression of in-frame Tnlac fusions to rfaA and rfaP were found to be affected by temperature (125).

GENETICS OF O-ANTIGEN SYNTHESIS

A summary of the genes involved in O-antigen synthesis and attachment is given in Table 3. Enteric bacteria of genera such as Salmonella, Escherichia, Shigella, and Klebsiella can be divided into classes based on the gene which is involved in attachment of the first sugar of the O unit to ACL. At the present time, only two such groups are known, the rfbP-dependent class and the rfe-dependent class (63, 77, 93, 141), although there will almost certainly be additional classes as more organisms are studied in detail. The rfbP-dependent class includes O groups A, B, C2, D, and E1 of Salmonella species. The prototype for the rfbP-dependent class is the group B organism S. typhimurium. This is the organism with which most of the initial biochemical and genetic studies of O-antigen synthesis were done (93, 142), as well as the first organism in which the rfb gene cluster and the rfe gene were sequenced (30, 65).

The O-specific side chains of these rfbP-dependent strains consist of a repeating unit of three, four, five, or six sugars which are often branched, and there are 25 to 40 of these repeating units in the longest O-specific polymers. In the members of this class, synthesis of a monomeric O unit is initiated by the rfbP-dependent transfer of Gal-phosphate from UDP-Gal to ACL-phosphate to form Gal-pyrophosphate-ACL. The remainder of the O unit is then built up by the sequential transfer of single sugars from their appropriate sugar nucleotides. Monomers are then combined into ACL-linked polymers by repeated transfer of a growing chain of O units onto newly synthesized monomeric O units attached to ACL, with the result that in a growing chain the O unit attached to ACL at the reducing end of the O polymer is the most recently synthesized (63, 111, 142). The rfbP-dependent strains do not require the rfe gene for synthesis of O antigen (64, 93). However, all of these strains require rfe for the synthesis of ECA, a polymer with a trisaccharide repeating unit (90), which, like O antigen, is assembled on ACL (102).

The rfe-dependent class of strains includes S. dysenteriae 1, Salmonella groups C1 and L, E. coli serotypes O8 and O9, and Klebsiella serotype O1. The O antigens of these strains are much more heterogeneous in structure. In S. dysenteriae 1, the O unit contains four sugars and its synthesis appears to be initiated by the rfe-dependent transfer of GlcNAc-phosphate to ACL, which is followed by the sequential addition of the remaining sugars. The rfb cluster of S. dysenteriae 1 does not contain an rfbP gene, but in other respects it resembles the rfb clusters of the rfbP-dependent Salmonella serovars (77). The Salmonella group C1 O antigen has a backbone containing GlcNAc and four mannose residues, with the penultimate mannose substituted by glucose. The rfb cluster of Salmonella group C1 resembles that of S. dysenteriae 1 in that it does not contain an rfbP gene or analog (82, 83).

The O antigens of the rfe-dependent E. coli and Klebsiella
TABLE 3. Genes or blocks of genes involved in the synthesis of O units and their assembly into O antigens attached to the LPS core

| Gene(s) | Probable function of gene products | Example(s) and reference(s) |
|---------|-----------------------------------|-----------------------------|
| rfbBCAD | Synthesis of TDP-rhamnose from glucose-1-phosphate | Salmonella (65) and Shigella (77) spp. |
| rfbKM | Synthesis of GDP-mannose from mannose-6-phosphate | Salmonella spp. (65) |
| rfbFGHI* | Synthesis of CDP-4-keto-3,6-dideoxyglucose (common precursor of dideoxy sugars) from CDP-glucose | Salmonella spp. (139) |
| rfbJ | Synthesis of CDP-abequose from common precursor in Salmonella group B | Salmonella spp. (139) |
| rfbS | Synthesis of CDP-paratose from common precursor in Salmonella groups A and D | Salmonella spp. (139) |
| rfbE | Synthesis of CDP-tyvelose from CDP-paratose in Salmonella group D | Salmonella spp. (139) |
| rfbP | Initiates synthesis of O unit by addition of galactose-phosphate to ACL to form galactose-pyrophosphate-ACL; may be bifunctional and encode RfbT function involved in transfer of O unit to LPS core | Salmonella spp. (65) |
| rfe | Initiates synthesis of O unit by addition of GlcNAc-phosphate to ACL to form GlcNAc-pyrophosphate-ACL | Shigella spp. (77) |
| rfbN | Synthesis of O unit; rhamnosyl transferase in groups A, B, D, and E of Salmonella spp. | Salmonella spp. (139) |
| rfbQ | Synthesis of O unit; rhamnosyl transferase I in S. dysenteriae 1 | Shigella spp. (77) |
| rfbR | Synthesis of O unit; rhamnosyl transferase II in S. dysenteriae 1 | Shigella spp. (77) |
| rfbU | Synthesis of O unit; mannosyl transferase in groups A, B, and D of Salmonella spp. | Salmonella spp. (139) |
| rfbV | Synthesis of O unit; abequosyl transferase in groups B and D of Salmonella spp. | Salmonella spp. (139) |
| rfp | Synthesis of O unit; plasmid-encoded galactosyltransferase in S. dysenteriae 1 | Shigella spp. (74) |
| rfbX | Assembly or transfer of O units; exact function unclear | Salmonella (139) and Shigella (77) spp. |
| rfc | Polymerization of O units on ACL by repeated transfer of growing chain to single O unit on ACL; may play role in ligation of polymeric O antigen to core | Salmonella (21) and Shigella (77) spp. |
| rfaL | Required for ligation of polymeric O antigen, single O units, or partial O units to core | E. coli (74) and Salmonella (92) spp. |
| rol (cll) | Regulation of chain length of polymeric O antigen | E. coli (9) and Salmonella (8) spp. |

* Brackets indicate genes which have not yet been located.

The serotypes listed above have a simpler structure. These consist of neutral homopolymers of Man or Gal, in some cases with modifications such as alternating anomic linkages or simple branch substratutions. The exact mode of synthesis of the homopolymeric O antigens is unclear, and some studies have suggested a continuous polymerization on the nonreducing end rather than the discontinuous buildup from smaller repeating units seen with the rfe-independent polymers (63). A single sugar residue which is different from the sugar residues making up the bulk of the polymer is found linking the reduced end of the polymer to ACL. There is some controversy over the identity of this initiating sugar. It was originally described as glucose (63), but studies on rfe showing that it is a GlcNAc-phosphate transferase (101) and recent analytical studies on an rfe-dependent E. coli O antigen (141) indicate that the initiating sugar will invariably be GlcNAc. In support of continuous polymerization, no mutants have been found in the rfe-dependent systems which attach a single O unit to LPS comparable to the “semirough” LPS which is produced by rfc mutants in the rfe-independent systems. However, an argument against continuous polymerization of the O antigen is that rfe-dependent LPS also migrates as a discontinuous ladder on gels.

Despite the very substantial differences in the structure and mechanism of biosynthesis, the synthesis of all of these types of O antigens requires an rfb gene cluster located in the 40-min region between his and the capsular polysaccharide (cps) genes. This section is concerned primarily with the structure of these rfb gene clusters and with the rfc and rfe genes which work with them.

**O-Antigen Genes of rfbP-Dependent Salmonella Serovars**

The Organization of rfb in S. typhimurium. As shown in Fig. 2, the rfb cluster of S. typhimurium consists of 16 contiguous genes which are transcribed in the counterclockwise (cps-to-his) direction (65, 139). Although there are accessory genes in regions flanking both ends of this cluster which may have a role in O-antigen synthesis, a fragment extending from the noncoding region upstream of rfbB through the rfbP gene is sufficient to confer O-antigen production on a Hiss-cps strain (65). The cluster is flanked at one end by cps genes and galF (Orf2.8 in Fig. 2) and at the other by gnd, rol, and the his operon, so that the order is cps-galF-rfb-gnd-rol-his (8, 18, 65).

The O unit of group B strains consists of a four- or five-sugar repeat (63). The backbone consists of Man, Rha, and Gal (Fig. 3) and is identical to that of groups A, D, and E1; in addition, the Man is substituted by the dideoxyhexose abequose (Abe) and the Gal is partially substituted by Glc. The Abe residue is also acetylated. The rfb cluster contains genes encoding the biosynthesis of GDP-Man, TDP-Rha and CDP-Abe. With the exception of TDP-Rha, these nucleotide sugars are not involved in synthesis of the LPS core or other essential cellular structures. Although carbohydrate gene clusters do not invariably include genes for the synthesis of the unique sugar nucleotides which they require, inclusion of such genes is a common strategy in O-antigen and capsular polysaccharide clusters. The cluster also includes four sugar transferase genes and the rfbX gene which is found in all rfb clusters and encodes an integral membrane protein involved in O-antigen assembly. The cluster is saturated; with the
exception of rfbX, all of the genes can be assigned functions as sugar transferases or in sugar nucleotide biosynthesis.

Sugar biosynthetic genes of the S. typhimurium rfb cluster. The genes for the biosynthesis of the sugar nucleotides are organized into three blocks within the rfb cluster. There are four genes involved in the synthesis of TDP-Rha from Glc-1-phosphate (93), and these are the four genes rfbBCAD at the 3′ end of the cluster (65, 139, 146). In earlier studies it was difficult to distinguish the last two enzymes in the TDP-Rha pathway, the epimerase encoded by rfbC and the dehydrogenase encoded by rfbD, and defects in either enzyme were termed rfbD. An assay for these two activities is now available (146), and the genes have been assigned by homology of their products to analogous enzymes encoded by the strM and strL genes of Streptomyces griseus (139). Immediately adjacent to the TDP-Rha biosynthetic genes are five genes involved in the synthesis of CDP-Abe from Glc-1-phosphate (65, 139). Three of these have been assigned as rfbFGJ, and the remaining genes, rfbH and rfbI, are thought to be the two unassigned open reading frames (87.6 and Orf10.4 present in this block. The two genes involved in the synthesis of GDP-Man from Man-6-phosphate are rfbMK near the other end of the cluster. All of these proteins have a structure consistent with soluble proteins (65).

The galF gene, which was mapped within or near the rfb cluster, was originally thought to be involved in the modification of UDP-Gal pyrophosphorylase, the product of the galU gene (65, 93, 108). The evidence for this was the presence of an altered form of the enzyme, which was eliminated by deletions which removed rfb genes. Recently, a comparison of GenBank protein sequences revealed that Orf2.8, encoded by a gene adjacent to rfbB (65), shared substantial regions of homology with GalU protein (138), suggesting that Orf2.8 is an alternate form of UDP-Gal pyrophosphorylase. Although it has not been tested directly, it seems very likely that the gene for Orf2.8 is galF and that its function is to provide a more efficient way of supplying the large amount of UDP-Gal required for synthesis of both a Gal-containing O antigen and a Gal-containing capsular polysaccharide. It is likely that it is the last gene in the cps cluster and not an rfb gene, since it is separated from the rfb cluster by a nontranscribed region which may contain a promoter for the rfb cluster. E. coli K-12 may also contain a galF gene, which could explain the somewhat leaky LPS phenotype of galU mutants (152).

Sugar transferases of the S. typhimurium rfb cluster. Three of the four rfb genes thought to encode sugar transferases have been assigned on the basis of the transfer of sugars from sugar nucleotides to ACL or ACL-linked oligosaccharide intermediates (18, 139). The location of rfbP, the gene for the transfer of galactose phosphate to ACL, at the 3′ end of the rfb cluster was anticipated on the basis of the properties of deletions extending into rfb from his (111). rfbP encodes the first function in the assembly of the O unit which commits ACL to the LPS pathway, an arrangement which was fortuitous for the early genetic study of the rfb cluster (93, 111). Deletions extending into rfb from the his side were easily selected, and these did not have the deleterious phenotype which might otherwise have resulted from accumulation of an ACL-bound intermediate or from the seques-

termination of ACL. The protein encoded by rfbP is relatively hydrophobic, with at least five apparent transmembrane domains (65), and both its structure and its function are different from those the rfb transferases which add the distal sugars. The hydropathy profile of the amino-terminal 250 residues of RfbP protein is very similar to that of the Rfc protein, which performs an analogous function by adding GlcNAc phosphate to ACL (78).

rfbN, rfbU, and rfbV determine the Rha, Man, and Abe transferases, respectively, and are arranged together in a block adjacent to rfbX. These transferase genes and rfbP are arranged in the gene cluster in the reverse order of the sequence in which the transferases act, an arrangement which appears to be followed in other rfb clusters which have been studied (139). The distal sugar transferases resemble the putative rfa hexosyltransferases in that their predicted structure generally does not contain significant hydrophobic segments which could function as membrane anchors or transmembrane domains. Thus it is likely that these proteins function as peripheral membrane proteins on the cytoplasmic face of the cytoplasmic membrane. None of these proteins show significant homology to proteins encoded by the rfa locus, which is not surprising since they recognize different sugar nucleotides and a different lipid acceptor. Two other functions involved in synthesis of group B O antigen, the O acetylation of Abe characteristic of group B strains and the glycosylation of the O antigen, are encoded by genes which lie outside of the rfb cluster, by the oafA gene which lies near but not within rfb and by the oafR-oaeE cluster near purE at 11 min, respectively (93).

Polimerization and role of the rfc gene. The polymerization of O units is one of the first aspects of O-antigen synthesis to be investigated both biochemically and genetically. There are two reasons for this. First, the formation of a glycosidic bond between adjacent O units results in a new structure which can be recognized both as a new antigenic factor and by loss or acquisition of a phage receptor. It was recognized early on that temperate phages have the ability to manipulate the polymerization system, and it was shown that such phages encode both an inhibitor of the host O polymerase and a new polymerase which forms a different linkage. This new linkage can differ from that of the host in the carbons which are involved as well as in the anomeric configuration (93). These phage-encoded polymerase functions are expressed in the lysogenic state, and, like phage-encoded porins, they represent a mechanism for preventing the nonproductive neutralization of phages by nearby uninduced lysogens or debris from the lysing cell when a lysogen is induced.

The second reason has to do with the rather distinctive semirough (SR) phenotype of mutants in which polymerization is defective. Tight mutants of this type produce LPS in which a large proportion of the molecules carry a single O unit while the remainder have R cores which carry no O antigen (93). The proportion of LPS with R cores which lack O antigen is about the same in S and SR strains, indicating that ligation of a single O unit to the core and translocation of the resulting LPS to the cell surface can be very efficient. In S. typhimurium, mutations resulting in SR LPS mapped to a locus outside of rfb between 18 and 34 min (93). This locus was presumed to encode the O polymerase and was termed rfc. More was learned about this phenomenon by examining the behavior of hybrids between various Salmonella serovars (93). When the rfb cluster of group B was transferred into group C1 or C2 strains, only SR LPS of the group B serotype was observed. In a reciprocal experiment, when the rfb cluster of group C1 was transferred into a group B strain, S LPS was produced. Subsequently, it was demonstrated genetically that the Rfc function of Salmonella group E1 mapped to the rfb locus (93).

These experiments suggested that in all three of these
serovars, C1, C2, and E1, the rfc gene might be located within the rfb cluster whereas in other groups such as group B it might be located elsewhere on the chromosome. This has now been partially confirmed by the cloning and sequencing of the rfc gene in group B and the identification of an rfc gene within the rfb cluster of a strain of group C2 (20, 21, 30). Both the group B and the group C2 Rfc proteins are very hydrophobic proteins with 10 or 11 potential transmembrane domains, indicating that they are integral membrane proteins. In the group C2 strain, the rfc gene is located in a nonconserved region near the middle of the rfb cluster adjacent to two genes which encode Man transferases (21). A gene encoding an open reading frame which may be an Rfc protein has also been found in the rfb cluster of group E1, although its location between the rfbD and gnd genes is different from in group C2 (175). However, the situation remains unclear for group C1, since there is no evidence for a gene encoding a similar protein in its rfb cluster (82, 83) and there have been no reports of mutants exhibiting an SR phenotype isolated from group C1 or from other strains producing simple or homopolymeric rfc-dependent O antigens similar to those of group C1. Thus the possibility remains that the O antigens of these organisms are polymerized by a different mechanism, which does not require an Rfc-like protein.

The polymorphism of rfc is not limited to Salmonella serovars. As shown in Fig. 2, an rfc gene is located adjacent to rfbX in the rfb cluster of S. dysenteriae 1, an organism which produces an rfc-dependent O antigen. The S. dysen- teriae 1 rfc gene was identified on the basis of mutational studies as well as on the structural similarity of its product to the Salmonella Rfc proteins. Finding that the polymerase genes are only occasionally a part of the rfb cluster is not surprising, given observations with O hybrid strains that the polymerase has a relaxed specificity with respect to minor differences in the structure of the O units, such as the nature of the dideoxy sugar substituent, and the fact that several temperate phages carry polymerase genes which under some conditions can substitute for host genes (93).

Modification of O polysaccharide. The commonest forms of O-polysaccharide modification involve O acetylation or glucosylation of O-antigen sugar residues or a change (either positional or anomeric) in the linkage between the O units. These changes result in antigenic differences which were detected primarily by serological methods, although they may also result in resistance to O-specific phages. All three of these modifications can be the result of lysogeny by temperate phages (93), and the gene from the temperate phage Sf6 which results in the acetylation of Rha in the O antigen of S. flexneri has recently been studied in detail (25, 155, 170). In group B Salmonella strains, glucosylation involves the chromosomal locus oafA at 43 min and acetyl- lation involves the oafC and oafR genes located in the 12- to 13-min region. There has been no recent work on the genetics of these phenomena in Salmonella strains, and readers are directed to an earlier review (93) for more detailed information.

The rfb cluster of Salmonella group C2 contains a gene termed orfJ4.9 (21), which encodes a protein whose deduced amino acid sequence shows strong similarity to the products of the lacA and nodL genes, both of which are known to be acetyltransferases (138). As yet, no function has been as- cribed to the OrfJ4.9 protein and no comparable gene has been found in the other Salmonella rfb clusters which have been sequenced. A gene located in or near the partial rfb region of E. coli K-12 results in the acetylation of the RhaII residues of O antigens which are produced when the cloned rfb clusters of S. flexneri 2a and 3a are expressed in E. coli K-12 (182). The true function of this gene is not clear, since the E. coli K-12 rfb cluster is not sufficient for the production of its own O antigen.

Polymorphism among rfbD-dependent Salmonella serovars. The rfb clusters of five rfbD-dependent Salmonella sero- groups have been sequenced. Groups B, A, D, and E1 all share a basic Man-Rha-Gal backbone structure, whereas group C2 has a Rha-Man-Man-Gal backbone. Additionally, the groups differ in the nature of the dideoxyhexose substit- uent: none in E1, Abe in B and C2, paratose in A, and tyrvulose in D. The biosynthesis of the dideoxyhexoses requires the products of four genes (rfbFGHI) to convert Glu-1-phosphate to the common precursor CDP-4-keto-3,6- dideoxy-D-Glc (I) and three additional genes to synthesize the final products. The rfbD gene product converts I to CDP-Abe, the product of rfbS converts I to CDP-paratose, and rfbE product converts CDP-paratose to CDP-tyrvulose (93, 139).

Group E1 has the smallest rfb cluster of the five sero- groups, since it lacks all of the dideoxyhexose genes and has only three sugar transferase genes. It contains two additional genes of unknown function, one of which may be an rfc analog (139, 175). The other groups share a conserved region adjacent to rfbBCAD, which includes the four shared dideoxyhexose genes rfbFGHI (Fig. 2) and either rfbJ (groups B and C2) or rfbE (groups A and D). The presence of rfbE in group D was not anticipated since its O antigen contains paratose instead of tyrvulose, but this was explained by the presence of an early frameshift mutation in this gene. Otherwise the rfb clusters of groups A and D are identical, suggesting that group D has acquired this mutation recently (89). The group C2 cluster is the largest, since it includes an additional Man transferase, the rfc gene and the putative acetylase gene noted in the previous section, and at least one additional gene of unknown function to bring the total number of genes to 19 (21). The structures of all of these clusters are quite close to what would be predicted on the basis of previous genetic and biochemical studies (93), and there are few additional genes.

As noted above, a polysaccharide gene cluster will often contain biosynthetic genes for sugar nucleotides which it requires. This can lead to duplication of function, an exam- ple being the duplication of genes for mannose biosynthesis between the rfb and cps gene clusters in some Salmonella serovars (93, 157). A consequence of this duplication is that one set of sugar biosynthetic genes might become lost, resulting in strains which have a similar function at different genetic loci. An example of this is seen in genes for the dTDP-Rha biosynthetic pathway. In S. typhimurium rfbA and rfbB encode the enzymes for the first two steps in the Rha biosynthetic pathway. These are also the first two steps in the synthesis of the furcosamine derivative 4-acetamido-4,6-dideoxy-D-Gal, which is part of the trisaccharide repeating unit of ECA. The O antigen of Salmonella group C1 LPS does not contain Rha, and the group C1 rfb cluster does not have a block of Rha biosynthetic genes corresponding to rfbBCAD of S. typhimurium (83). There is evidence that in group C1 a gene analogous to rfbA (and presumably an analog of rfbB) is located in the rfe-rff cluster instead of rfb (87). It is not known whether the group C1 genes are related to the conserved rfbA and rfbB genes found in other Salmo- nella serovars and in S. dysenteriae 1 (77, 139) or whether the rfe-rff cluster of group C1 contains analogs of rfbCD, the last two genes in the dTDP-Rha pathway.
O-Antigen Genes of Shigella dysenteriae 1

Genetic structure of the rfb cluster. As shown in Fig. 3, the O antigen of S. dysenteriae 1 has a repeating unit consisting of Rha-Rha-Gal-GlcNAc (40). Expression of this O antigen in E. coli K-12 requires the E. coli K-12 rfe gene (77) and the S. dysenteriae 1 rfp gene (74, 159). In S. dysenteriae 1 the rfp gene is carried on a 9-kb multicopy plasmid which is invariably found in virulent strains and is required for O-antigen production (51, 176, 177). In addition, expression of the O antigen in an E. coli K-12 strain deleted for the his-cps region requires a cloned fragment containing eight contiguous genes from the chromosomal rfb cluster of S. dysenteriae 1 (77, 160, 161). The 8.9-kb rfp fragment shown in Fig. 2 is one of the smallest rfb clusters found to date which is capable of directing O-antigen expression in an rfp-deleted host. The left end of the cluster contains the rfbBCAD genes, which are necessary for TDP-Rha synthesis. These show substantial homology to their counterparts in Salmonella strains (65). The next two genes are rfx and rfc, and their protein products show structural similarity but only a low level of sequence homology to their Salmonella counterparts. The two genes designated as rfbRQ have been identified as encoding the Rha II and Rha I transferases, respectively (77, 160). The fragment encodes an additional open reading frame of unknown function, designated as Or1, and part of another open reading frame which appears from restriction site analysis to lie between rfbRQ and gna. Neither Or1 nor the partial product of this additional open reading frame share significant structural homology with other known O-antigen gene products.

Roles of rfe and rfp. As noted above, the rfb cluster encodes only the sugar transferases which add the two distal sugars of the O unit. The two transferases for the proximal sugars are thought to be encoded by rfe and the plasmid gene rfp (77).

Introduction of rfp into E. coli K-12 resulted in an increase in the galactose content of the LPS core, and low-resolution SDS-gel analysis indicated that the migration of part of the LPS was reduced (153, 159). On this basis, it was suggested that rfp encoded a galactosyltransferase (159), and this has now been confirmed by the demonstration that overproduction of rfp leads to increased expression of Gal transferase activity (17). The Tricine-SDS gel system allowed the rfp-induced change in gel migration of the LPS to be examined more precisely. This showed that rfp reduced the migration of band 1 by an amount consistent with the addition of one sugar whereas the migration of bands 2 and 3 showed no change (74).

The size of this Rfp-modified band is two sugars too large to serve as a precursor for the addition of O units (74). This, together with the properties of an rfe::Tn10 mutation, indicated that the Rfp-modified band is not a precursor but instead is an LPS molecule to which a two-sugar partial O unit has been transferred. The rfe mutation eliminated both band 1 and the Rfp-modified band, as well as eliminating the ability of E. coli K-12 to express the S. dysenteriae 1 O antigen. This indicated that the rfp-modified band involved the addition of both GlcNAC and Gal to the core (17, 77). Moreover, when derivatives of the S. dysenteriae 1 rfb cluster with mutations in the gene encoding rhnamosyl II transferase or in rfc are introduced together with rfp, they result in bands which are one and two sugars larger, respectively, than the rfp-modified band (77, 160). These intermediates are also seen in a leaky rfb mutant with an insertion in the 5′ untranslated region, which presumably contains the rfb promoter (65). Thus all four of the transferase activities can be accounted for. Earlier work envisioned the buildup of the O unit as involving the transfer of sugars directly to the LPS core (160), but the involvement of rfe and the structural and functional similarity of Rfe to RfbP indicate that both partial and complete O units are first assembled by ACL and then transferred to the LPS core. Judging by gel band intensity, the efficiency of transfer of the partial O units appears to increase significantly as each sugar is added to the O unit (77). This indicates a preference for complete O units and may be one of the reasons why the transfer of partial O units was not detected in earlier studies.

This interpretation of the roles of rfe and rfp and of the transfer of partial O units to the core has been strengthened by a very recent study (41A) which examined the structure of core oligosaccharides produced by various E. coli K-12-S. dysenteriae 1 hybrid strains. The results on the K-12 background strain confirmed the earlier finding of GlcNAC linked to the core in a β configuration (64) and showed that in hybrids containing only the rfp gene this GlcNAC was substituted with Gal. In hybrids which had both the rfp gene and derivatives of the S. dysenteriae 1 rfb cluster which supplied one or both Rha transferases, this Gal was additionally substituted with one or two Rha residues.

Role of rfxB. The rfxB genes which have been studied in Salmonella and Shigella strains encode large, hydrophobic proteins with multiple membrane-spanning domains (65, 77, 89, 139). Mutations which inactive rfxB have been difficult to study. They are deleterious in strains in which all of the other genes involved in synthesizing an O unit are functional, and thus they are unstable on plasmids and are difficult to isolate and manipulate (77). In addition, E. coli K-12 has one or more chromosomal genes which encode rfxB analogs, and this has further complicated the analysis of rfxB mutations carried on plasmids. A plasmid which carried the S. dysenteriae rfb cluster with a nonpolar rfxB::Tmalac insertion mutation resulted in a significant amount of O antigen when it was introduced into an E. coli K-12 strain background with an intact his-rfb region, indicating that E. coli K-12 had an rfxB analog which was able to effectively complement the mutation in the plasmid. However, when this rfxB::Tmalac plasmid was tested in a strain background deleted for the his-rfb region, some SR LPS was made but there was no detectable O ladder. The his-rfb deletion also affected the production of band 1 in strains without rfb plasmids and reduced the amount of Rfp-modified band (77, 78). Unfortunately, the possibility that the slightly leaky phenotype of the strain deleted for his-rfb is due to partial complementation by an rfxB analog is part of another carbohydrate cluster (for example, rfe-rff) cannot be ruled out.

The observation in the his-rfb-deleted background that the inactivation of rfxB in the plasmid led to a complete loss of S LPS but only a partial reduction of SR LPS suggests that the primary role of rfxB is not in the assembly of the monomeric O unit on ACL. Likewise, the observation that these mutants are much less efficient in synthesizing SR LPS than are rfe mutants indicates that the primary role of rfxB is unlikely to be in polymerization of O antigen but suggests that its role may be in the ligation of O units to the LPS or some other part of the process (for example, the flip-flop of O units on ACL to the periplasmic face of the cytoplasmic membrane) which may be necessary for the efficient transfer of O antigen to LPS.
O-Antigen Genes of E. coli

Genetic analysis of rfb region of E. coli K-12. As noted above, there is genetic evidence that the his region of wild-type E. coli K-12 contains an intact rfbBCAD cluster (4, 77, 78), a functional analog of rfbX (77), a rol gene (78), and a gene encoding an O-antigen acetylase (182). Analysis of sequences from the GenBank data base indicates that the insertion element IS51 (168) lies 5' to gnd (103) and is probably in the distal end of the rfb cluster. There are no significant open reading frames between the IS5 and gnd, so it is not clear how the insertion is related to the lack of rfb function.

rfb clusters from other E. coli serovars. Functionally complete rfb gene clusters have been cloned from a number of different serovars, including O2 (109), O4 (52, 53), O7 (95, 169), O9 (72, 162), O101 (57), and O111 (7). Most of these have not yet been sequenced or analyzed genetically to the point at which significant comparisons to the rfb clusters in Salmonella spp. can be made.

An interesting rfb cluster which has been partially analyzed is that of the E. coli O7:K1 strain VW187 (94). The distal end of this very large cluster contains the two genes rfbM and rfbK involved in GDP-mannose synthesis, which were identified by the similarity of their products to those of the corresponding Salmonella genes. These genes are located immediately adjacent to gnd, unlike the situation in the Salmonella rfb clusters, where rfbM and rfbK are separated from gnd by the rfbP gene. This suggests that E. coli O7:K1, like S. dysenteriae 1, uses a gene other than rfbP to initiate the synthesis of O units. Since the O7 O antigen contains GlcNAc, it is possible that this organism also used the rfe gene for this function.
cally when core synthesis was blocked (107), and it has been shown with doubly conditional mutants that ACL-linked polymer accumulated during inhibition of core synthesis can be chased into LPS (98). Nevertheless, in both systems some caution must be used in interpreting the significance of the presence of material destined for the outer membrane in the periplasmic compartment, since this may represent a default pathway rather than an indication of true intermediates.

The models in which periplasmic O antigen is a direct precursor of LPS on the surface of the cell require the LPS to flip-flop across a membrane twice, once to flip the core-lipid A complex across the cytoplasmic membrane from the cytoplasmic face to the periplasmic face prior to attachment of O antigen and a second time to flip the complete LPS molecule across the outer membrane from the periplasmic face to the external face (129). Although it is not too difficult to imagine the flip-flop of four or five sugars attached to ACL, it is somewhat more difficult to imagine a flip-flop of LPS. At minimum, the lipid A portion of LPS has six acyl chains attached to a rigid, planar disaccharide head group, and this makes it much more likely to form stable bilayer structures than the single, long, flexible acyl chain of ACL. In addition, LPS has a number of widely spaced charged groups, a bulky core of at least 10 sugars, and, in the case of the complete molecule which must be flipped across the outer membrane, a long extended O-polysaccharide chain of perhaps 100 sugars.

A model which should be added to those that have been proposed is one in which the lipid A portion of the LPS molecule remains on one side of the bilayer and part of the core and/or part of the O chain extend through a protein channel into the periplasmic compartment. The core has the size and flexibility to lie flat on the surface of the membrane or to be folded back into the bilayer, and the outer core plus a single O unit are long enough to extend across the bilayer through a protein channel (71). Protein channels which allow polysaccharide chains to cross a membrane bilayer are not uncommon in bacteria, a good example being the LamB protein, which is involved in maltodextrin transport. If translocation to the outer membrane involves a shuttle vesicle or Beyer’s bridge mechanism, one requirement of this model is that channel proteins which coat the polysaccharide as it passes through the membrane would have to accompany the LPS until translocation to the outer membrane was complete, and these proteins would have to be recycled to the cytoplasmic membrane at the end of the cycle.

A useful approach to these models would be to determine the topology of the active sites of enzymes involved in O-antigen assembly and attachment to the LPS core. Unfortunately, this will be difficult with some of the proteins which are likely to be involved, since the RfaL, RfxX, Rfc, RfbP, Rfe, and Rol proteins are all large, hydrophobic proteins with multiple membrane-spanning domains. An alternative approach might be to study the location and properties of the proteins involved in the addition of acetyl (182) or glucosyl (55, 93) modifications to the core, since there is evidence which indicates that these substituents are added after the O unit has been synthesized and at least partially polymerized. This approach would be particularly interesting in view of a recent study of the glucosylation which determines O-antigen factor 12 in *S. typhimurium*, since an electrophoretic comparison of the O-antigen ladders of glucosylated and nonglucosylated LPS suggested that glucosylation was present only in O units distal to the sixth O unit in the polymer (55). Since new units are added to the proximal end of the polymer, this finding is very strongly suggestive of some form of protection or compartmentalization of the O antigen during one stage of its synthesis.

**Attachment of O antigen to the LPS core.** On the basis of the early studies which demonstrated that O units were synthesized and polymerized on ACL and genetic studies of the polymerase, it was implied that the polymerization reaction and the transfer (more commonly termed ligation) were distinct processes involving different gene products (93). The predicted phenotype of mutants lacking O ligase is the presence of polymeric O antigen bound to ACL but no O antigen attached to LPS. This is the phenotype exhibited by *rfaL* mutants and by a few mutants with mutations at the *rfa* locus which were designated as *rft* mutants (93). No gene corresponding to *rft* has been found among the *rfb* clusters which have been sequenced, although a direct role for *rftX* in ligation cannot yet be ruled out. The *Salmonella rft* mutants have been found to have a defect in *rfbP*, suggesting that the large protein encoded by this gene may be bifunctional (138).

This raises the interesting question of how strains which lack an *rfbP* gene, such as *Salmonella* group C1 (82), *S. dysenteriae* 1 (77), and probably *E. coli* O7:K1 (94), can transfer O antigen from ACL to LPS. The answer may lie in a hydrophobic protein, designated as o349, encoded by a gene which lies adjacent to *rfe* (36, 101). Joining of the reading frames of *rfe* and half of the gene encoding o349 resulted in a larger protein with a hydrophathy profile which was remarkably similar to that of RfbP protein, suggesting that *rfbP* arose by a similar fusion of ancestral genes (77). More work is clearly needed to sort out the genetics of the ligation reaction.

In addition to considering the role of various gene products in the ligation reaction, it is useful to examine two different models for the ligation pathway. We term these the single-step and multistep models, and a highly speculative comparison of the roles of the various gene products in each of these models is given in Table 4. The single-step model is the traditional model for ligation developed more than two decades ago and involves the *rfe*-dependent polymerization of O antigen on ACL followed by a single-step ligation reaction in which completed O polymers are transferred from ACL to the terminal core sugar of the acceptor LPS. In this model, ligation of polymeric O antigen does not directly involve *rfe*. The multistep model begins with the ligation of a single O unit to the terminal core sugar of the acceptor LPS to form SR LPS. Polymers are then transferred to this single O unit in an *rfe*-dependent reaction which is analogous to the polymerization reaction. Both models require a ligase to transfer a sugar at the reducing terminus of an O unit from ACL to the LPS core and a polymerase to generate ACL-linked polymers containing multiple O units, so the models are equivalent in complexity. Both models begin with the synthesis of a pool of ACL-linked O units, and the structure of the sugar transferases indicates that these are assembled on the cytoplasmic face of the cytoplasmic membrane. Beyond this step, neither model imposes any obvious constraints on whether reaction take place on the cytoplasmic or periplasmic face of the membrane or within protein channels embedded in the membrane.

The multistep model has some advantages in explaining the large amount of SR LPS observed in smooth strains and *rfe* mutants, as well as the apparently efficient transfer of partial O units in rough strains. The presence of an additional three- to five-sugar extension on the core provided by the ligation of a single O unit to the core prior to the attachment
TABLE 4. Comparison of gene functions predicted by the single-step and multistep models for O-antigen ligation

| Gene or phenotype | Function predicted by: |
|-------------------|------------------------|
| rfaL              | Component of ligase which adds polymerized O antigen to core |
| RfbT*             | Component of ligase which adds polymerized O antigen to core |
| rfc               | Polymerization of O units by repeated transfer of growing chain to single O units on ACL |
| rol (cld)         | Regulates kinetics of polymerization by Rfc versus ligation to core by RfaL and/or RfbT |
|                   | Regulates kinetics of polymerization by Rfc versus ligation to core by RfaL |

* This phenotype is used to indicate the RfbT function proposed for the bifunctional rfbP gene product of group B Salmonella spp. (138) or the product of a gene from the rfe-eff cluster in E. coli K-12-S. dysenteriae 1 hybrids (77).

of polymeric O antigen might be advantageous in the early stages of a model in which periplasmic O antigen is tethered across the cytoplasmic membrane to lipid A on the cytoplasmic face. None of these arguments are compelling in favor of the multistep model. A much more important reason for considering both models is that they differ in their requirements in a reconstituted in vitro system. In the multistep model, SR LPS would be required as the O-antigen acceptor and the reaction would require Rfc protein, whereas in the single-step model, R LPS would be used as the acceptor and Rfc protein would not necessarily be required.

In contrast to the parts of the LPS biosynthetic pathway, reconstitution systems have not provided a useful way of studying the ligation reaction. This is likely to change now that most of the genes involved LPS assembly have been cloned and sequenced, since in vitro genetics provides a means of constructing strains tailored for reconstitution studies which have precisely defined enzymatic lesions or produce LPS acceptors in which assembly has been terminated at specific stages. In addition, knowledge of the structure of the gene products can identify which functions are likely to require intrinsic transmembrane proteins, which must be supplied as membrane fragments or detergent extracts, rather than the more peripheral proteins, which can be supplied by soluble extracts and which are more amenable to synthesis in large amounts by expression systems.

CONCLUDING REMARKS

The past decade has seen the development of a nearly complete catalog of LPS genes. This has both exposed our ignorance and provided a powerful tool for future studies. In its early years, LPS research was dominated by in vitro reconstitution studies in which mutants, cell extracts, and labeled precursors were combined to work out complicated biosynthetic pathways. This tradition has continued more recently with the nearly complete elucidation of the lipid A biosynthetic pathway. Now the availability of sequence data on LPS genes and the ease with which defined mutations can be constructed and crossed onto the chromosome allow these reconstitution experiments to be repeated with different goals and at a much higher level of resolution. These are probably the most important experiments to be undertaken, since they offer a way of studying questions such as the organization of multienzyme complexes in the membrane and the mechanism of the final steps of assembly and translocation of LPS, which have previously been intractable.

This period has also been a time when there has been a more acute recognition of the tremendous flexibility and adaptability of the gram-negative cell surface, and this has changed the way we view LPS. The complex oligosaccharides which decorate glycoproteins in eukaryotic organisms have been likened in function to the bar codes used to mark items in a supermarket (41), and it is useful to extend this analogy to LPS and other complex bacterial polysaccharides. These molecules provide a genetically inexpensive way to allow cells to specifically recognize each other, to be recognized by hosts, to colonize favorable host environments, and to evade or confuse predators. Originally, LPS pathways were viewed as linear pathways designed to produce a single major product, and the heterogeneity resulting from the presence of partial branch substituents could be explained on the basis of the inefficiency of some of the biosynthetic enzymes. New evidence indicates that LPS pathways are branched so that multiple products can be produced, and the addition or removal of partial sugar substituents such as Rha, Hep, or GlcNAc may be used at branch points to target molecules for different pathways. This shows much more similarity to the kinds of mechanisms which are used to target glycoforms in eukaryotic systems (43).

An important aspect of this flexibility is preliminary but growing evidence suggesting that repeating polymers such as O antigens, ECA, and colanic acids, which are assembled on undecaprenol (ACL) carriers, may be interchangeably transferred to the LPS core and to smaller acceptors such as simple phospholipids. This can allow some strains to express these polymers either as LPS, which is a firmly bound, integral component of the outer membrane, or as microcapsular polysaccharides, which are much easily solubilized and dispersed in water as micellar suspensions. Thus the same polymer can be found in both a bound and a diffusible form, in some ways analogous to human blood group substances. The biological importance of this strategy is underlined by a recent discovery in the Rhizobium-legume colonization system. Bacterial nodulation-specific (nod) genes which are responsible for the species or cultivar specificity of nodulation were found to encode enzymes for the site-specific sulfation of a diffusible complex polysaccharide (145). This complex polysaccharide was termed a LOS, but it is attached to a phospholipid anchor rather than lipid A and has a structure more similar to that of ECA.
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