Early molecular-recognition events in the synthesis and export of group 2 capsular polysaccharides

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The outer membrane (OM) of almost all Gram-negative bacteria is composed of phospholipids, lipopolysaccharides, proteins and capsular or loosely adherent polysaccharides that together mediate cellular interactions with diverse environments. Most OM components are synthesized intracellularly or at the inner membrane (IM) and thus require an export mechanism. This mini-review focuses on recent progress in understanding how synthesis of one kind of capsular polysaccharide (group 2) is coupled to the export apparatus located in the IM and spanning the periplasmic space, thus providing a transport channel to the cell surface. Although the model system for these investigations is the medically important extraintestinal pathogen Escherichia coli K1 and its polysialic acid capsule, the conclusions are general for other group 2 and group 2-like polysaccharides synthesized by many different bacterial species.

Introduction

Bacterial extracellular or exo-polysaccharides (EPSs) may be homopolysaccharides composed of a single sugar residue or heteropolysaccharides with two or more monosaccharides that may or may not include branching of the main chain. The evolution of these polysaccharides has produced thousands of different somatic or O antigens as part of Gram-negative bacterial lipopolysaccharide (LPS) and hundreds of K (‘kapsel’) or capsular polysaccharides in diverse species. Although most investigators prefer to distinguish polysaccharides by their O antigen, capsular, EPS or enterobacterial common antigen designations, we think that EPS is a suitable term for lumping all polysaccharides synthesized by many different bacterial species. Despite immense structural variation there are just two or three mechanisms for exporting EPSs from their site of synthesis inside the cell to the cell surface in Enterobacteriaceae and others: (i) the group 1/4 or LPS model involves assembly of monosaccharides (building blocks) on a lipid carrier, usually undecaprenol, that is ‘flipped’ or transported across the inner or cytoplasmic membrane and either polymerized at the IM–periplasm junction or exported directly to the outer membrane (OM) if already polymerized in the cytoplasm; and (ii) the group 2/3 capsule model, in which polysaccharides are synthesized inside the cell prior to or during export to the OM without apparent periplasmic intermediates or commingling with components of the LPS pathway. The first model, which is widely variable, and includes the colanic acid or M antigen common to most strains of E. coli and diverse O antigens (Bos et al., 2007), will not be discussed here. [See recent reviews for details of this mechanism, including its ramifications for groups 1 and 4 capsules (Whitfield & Paiment, 2003; Whitfield, 2006).] Both groups 1/4 and group 2 export mechanisms require ATP-binding cassette-like (ABC) transporters to energize polysaccharide export as well as a lipid A flipase for LPS biosynthesis; in at least one case the direct binding of EPS to a specialized domain of the ATPase component of an ABC transporter has been demonstrated (Cuthbertson et al., 2007). By contrast, some polysaccharides, such as cellulose and poly-β-1,6-N-acetyl-d-glucosamine, which are common biofilm matrix molecules in E. coli and other bacteria, do not seem to require ABC transporters, suggesting that the polymerases themselves might direct nascent polysaccharides across the cytoplasmic membrane, representing what may be a distinct translocation mechanism. The reader is directed to a short review by Weigel & DeAngelis (2007) for a discussion of the theoretical problems associated with polysaccharide export in the absence of a mechanism for energizing translocation.

The focus of this mini-review is on the early molecular-recognition events involving heterotypic protein–protein interactions that direct group 2 and probably group 3 capsular polysaccharides to the evolutionarily conserved export apparatus composed of an ABC transporter, periplasmic connector and OM pore, shown on the right in Fig. 1. Whitfield (2006) has summarized the distinctions between groups 1–4 capsules and the reader is directed to that review for further information about capsule classification. The translocation apparatus for groups 2 and 3...
capsular polysaccharides is formally analogous to the
translocators for certain disease and polypeptides that use
the type I protein secretion system shown on the left in
Fig. 1 (Thanabalu et al., 1998); these include substrate-
specific IM ABC transporters, periplasmic connectors, and
an OM exit pore provided by interactions with TolC (Silver
et al., 2001). However, unlike type I secretion systems,
group 2 capsule exporters invariably require additional, or
accessory proteins for polysaccharide translocation (Fig. 1).

For example, a simple BLASTP analysis of the groups 2 and 3
accessory polypeptide, KpsC, yielded 87/100 non- E. coli
‘hits’ with alignment values <1 x 10^-88 in 44 different bacterial species of diverse phylogenetic origins. This result
indicates that capsule biosynthesis is likely to be a property
of many more bacterial species than currently studied,
supporting the overall importance of capsules for medi-
atting interactions with host or environmental surfaces.

Understanding the mechanism of group 2 capsule export is
thus central to how we think about microbes in diseases
resulting from animal and plant infections as well as
bacterial social interactions such as biofilm organization.

In addition to accessory proteins, it is believed that group 2
polysaccharide synthesis requires a membrane-bound
initiator upon which the polysaccharide chain is elongated,
and/or a terminator (Fig. 1), which has been variably
described, sometimes by the same research group, as
phospholipid, phospholipid-linked 3-deoxy-D-manno-
octulosonate (KDO) (Bronner et al., 1993; Finke et al.,
1991), ‘endogenous acceptor’ protein, and undecaprenyl
phosphate (Troy et al., 1975; Weisgerber & Troy, 1990).

The exact nature of the endogenous acceptor and
termination event remain outstanding questions for future
research.

**Building blocks – monosaccharide synthesis,
activation and polymerization or transfer to
acceptors**

Monosaccharides are EPS building blocks synthesized from
simpler precursors by sequentially acting enzymes includ-
ing at least some of and sometimes more than the catalysts
listed in Step 1 of Fig. 2. However, in one prominent case
the building block designated sialic acid is acquired directly
from the host environment, thereby obviating the need for
de novo glucose synthesis (Severi, et al., 2007; Vimr et al.,
2004). Once the glucose units have been synthesized or
obtained from the environment, activation to mono- or
diphosphonucleotide sugars (Step 2), where XTP, XDP and
XMP indicate nucleotide (adenosine, cytosine, thymine or
uridine) tri-, di- and monophosphonucleotides respect-
ively, primes them for the polymerization step. Synthesis of
these ‘high-energy’ nucleotide intermediates is necessary
for polymerization catalysed by the glycosyltransferases
that transfer glycosyl units to appropriate acceptors (Step 3).
In most cases, polysaccharides are synthesized as part of a
glycoconjugate that includes a lipid or protein moiety.

These conjugates of polysaccharides with lipids or proteins
are the molecular forms finally expressed at the cell surface.
Any of the steps involved in synthesis or acquisition of the
building blocks or of activation and polymerization of
these units are potential targets for inhibiting the
expression of capsular polysaccharides. Targeting these
steps for therapeutic or vaccine development offers the
potential to interfere with or block diverse microbial
interactions involved in colonization and disease progress-
ion as well as adherence to environmental surfaces (Vimr
& Steenbergen, 2006).

**Phenotypes of group 2 capsular polysaccharide-export-
deficient mutants as they relate to different export
models**

The genes for group 2 and 3 polysaccharide synthesis are
organized into functional modules, where region 2 for
synthesis is flanked by export genes (regions 1 and 3
comprising the multigenic cluster for biosynthesis of the E.
coli K1 capsular polysialic acid) (Fig. 3A). Arrows in Fig.
3(A) indicate modular transcription directions and shaded
areas regional gene translational coupling, which presum-
ably ensures synthesis of the precise amounts of each
polypeptide required for efficient capsular polysaccharide
biosynthesis. The modular organization of E. coli groups 2
and 3 capsules is observed in other encapsulated bacteria
such as Campylobacter, Haemophilus, Neisseria, Pasteurella
and Actinobacillus spp. (Silver et al., 2001), suggesting a
common origin for the biosynthetic gene clusters in these
and probably other species that synthesize group 2 or
Polysaccharide export

<table>
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<tr>
<th>Step 1:</th>
<th>Precursor(s) → Glycosyltransferase(s)</th>
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<tr>
<td></td>
<td>Dehydrogenase, epimerase, isomerase, kinase, synthase</td>
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<th>Step 2:</th>
<th>XTP + Glycosyltransferase(s) → XDP- or XMP-Glycosyltransferase(s)</th>
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<td>Synthetase</td>
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<th>Step 3:</th>
<th>Activated Glycosyltransferase(s) → Polysaccharide</th>
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<td>Glycosyltransferase(s)</td>
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Fig. 2. Summary of the steps involved in polysaccharide biosynthesis. See text for details.

Group 2-like capsules. Indeed, Silver and his colleagues indicated that homologues of the E. coli K1 ABC transporter (kpsMT), periplasmic connector (kpsE) and OM pore genes from Actinobacillus pleuropneumoniae, Haemophilus influenzae and Neisseria meningitidis complemented the export defect of an E. coli kpsMTDE quadruple mutant, supporting the conclusion that carbohydrate structure has no major effect on recognition of polysaccharides by the respective translocation machineries (Silver et al., 2001). This observation begs the question: how are group 2 and group 2-like polysaccharides recognized by the exporters?

It is useful to think about group 2-polysaccharide export by focusing on three different potential mechanisms. For example, synthesis could be obligatorily coupled to export such that no polymer is made if the translocator is inactive. Whitfield & Paiment (2003) have reviewed export systems for the LPS-like group 1 capsules, where this feedback mechanism appears to apply. By contrast, defects in group 2 and probably group 3 capsule export result in the intracellular accumulation of polysaccharides that are visualized as electron-transparent lacunae in transmission electron micrographs (Cieslewicz & Vimr, 1996, 1997). The intracellular accumulation of these unexported group 2 polysaccharide pools, observed in regions 1 and 3 and neuE mutants, means that EPS translocation is not obligatory for synthesis, suggesting two additional biosynthetic options. (Note that for the purposes of this mini-review we define biosynthesis as the combined processes of polysaccharide synthesis and export.) First, polysaccharides could be synthesized with a common molecular tag directing them to the export apparatus, a mechanism that would be analogous to the C-terminal secretion signals in polysaccharide substrates of type I secretion systems (Thanabalur et al., 1998). Similarly, the polymannans made by E. coli O8 and O9a are synthesized with terminal methyl groups at their non-reducing ends which are recognized by specialized domains of the ATPase components of the ABC transporters, signalling subsequent export (Cuthbertson et al., 2007). The ATPase from each system does not cross-complement, indicating specificity for any particular polysaccharide modification. By contrast, all group 2 and probably group 3 capsule export genes function directly in translocation (as seen in mutants with peripherally distributed lacunae) while others direct the coupling between synthesis and export (as in mutants with large intracellular pools). On the basis of this interpretation, we hypothesized that KpsC interacts directly with the polymerase (NeuS in the K1 system) and guides it to the periplasmic connector, KpsE, defining a series of molecular-recognition events that might effectively couple synthesis to export by positioning the polymerase near the export channel. By contrast, other components of the system, such as KpsT and KpsS, would function passively in the export process. For example, the ATPase component of the ABC transporter might function mainly to decorate the KpsE channel to the OM. In other words, while the polymer presumably passes through the IM component, KpsM, of the exporter, it might not necessarily have to pass through the KpsT ATPase.
**Fig. 3.** Group 2 capsular polysialic acid biosynthesis in *E. coli* K1. (A) Genetic and functional organization of the *kps–neu* gene cluster for polysialic acid biosynthesis adapted from Steenbergen & Vimr (2008); see text for details. (B) Protein–protein interactions between export and synthetic gene products detected by two-hybrid analysis. (C) Model of group 2 capsular polysaccharide biosynthesis. Gene products indicated by black boxes are defined in panel (A) and the text. Open and shaded small circles indicate unacetylated and acetylated sialic acids respectively. The small black circles indicate KDO or unknown moiety while the squiggle-circled P is phosphatidic acid. Triangles indicate CMP; the square is N-acetylmannosamine (ManNAc); Ac, acetyl groups; AcCoA and CoA, acetyl coenzyme A and coenzyme A respectively; Rib-5-P, ribulose 5-phosphate; Ara-5-P, d-arabinose 5-phosphate. Other small molecules have their usual abbreviations. Colours indicate essential or non-essential functions as follows: pink, essential synthetic functions; yellow, non-essential synthetic functions in the K1 system; green, essential exporter functions; blue, essential accessory proteins; purple, atypical OM protein found in some but not all group 2 or group 2-like systems. Specific interactions detected by two-hybrid analysis between NeuS, KpsE, and KpsC are indicated by the overlapping shapes. The specific steps in polysialic acid biosynthesis include the following. Step 1, the first committed step, involves the releasing epimerase, NeuC, producing ManNAc. Step 2: ManNAc interacting with the NeuB–NeuD complex is condensed with phosphoenolpyruvate (PEP) by NeuB while NeuD transfers an acetyl group to most of the nascent sialic acids. Step 3: the N-terminal domain of the synthetase NeuA activates sialic acid for polymerization by coupling it to CMP donated by CTP. Concomitantly with activation the C-terminal synthetase domain (NeuA*) acts as an O-acetyl esterase to remove the acetyl group from most sialic acid substrates. The functional significance of the cyclical acetylation and deacetylation reactions is not presently understood. Step 4: sialic acid residues are polymerized by NeuS while NeuO in most K1 strains reacetylates the growing chain. Step 5: NeuS, NeuC, and NeuE interact such that during or shortly after polymerization the completed polysialic acid is exported through the transmembrane channel to the outer membrane where the phospholipid anchors it to the outer leaflet of the OM.
component. Despite evidence that KpsS can be chemically cross-linked to most other components of the E. coli K5 biosynthetic system (McNulty et al., 2006), the function of this accessory protein appears to be as a bystander rather than a coupler on the basis of the kpsS mutant phenotype (Fig. 4).

Evidence for directed coupling of group 2 capsular polysaccharide synthesis to the translocation apparatus

Previous studies of group 2 polysaccharide biosynthesis have used immunological, chemical cross-linking and radiation target analysis methods to confirm that accessory proteins and the kpsMTED exporter channel were likely to interact heterotypically with synthetic components such as the polymerases (Andreischeva & Vann, 2006; McNulty et al., 2006; Vionnet et al., 2006). However, these studies provided no coherent model for thinking about the early biosynthetic events coupling polysaccharide synthesis to the exporter, or even if there was coupling between the processes. In an attempt to distinguish between the post-synthetic and directed coupling models discussed above, we used expression of recombinant polysialic acid depolymerase to determine whether cytoplasmic synthesis of the polysaccharide was protected from degradation. In essence, this in vivo protection assay assessed the relative intimacy of the relation between synthesis and export. The depolymerase, which functions by an endo-hydrolytic mechanism, would clip nascent chains prior to export, thus reducing or eliminating surface capsule expression. However, if the molecular associations involved in synthesis and export are intimate, the polysaccharide would be protected and the capsule expressed regardless of depolymerase. The results indicated that capsular polysialic acid synthesis was protected during polymerization and export, supporting the directed coupling model. This conclusion was supported by a series of control experiments showing that any cytoplasmic intermediates would have been susceptible to the depolymerase in vivo, establishing the in vivo protection assay as a novel method for investigating group 2 capsule biosynthesis (Steenbergen & Vimr, 2008).

Two-hybrid analysis of homo- and heterotypic interactions in group 2 capsule biosynthesis

Most two-hybrid systems for detecting protein or peptide interactions rely on activation of various transcriptional regulators to provide the reporter activity. By contrast, reconstituting the N- and C-terminal domains of the Bordetella pertussis adenylate cyclase (Cya) through interacting homo- or heterotypic fusions results in cyclic AMP

Fig. 4. Phenotypes of export-deficient acapsular mutants. Thin sections of representative bacteria with the indicated defects in export gene were examined by transmission electron microscopy as previously described by Cieslewicz & Vimr (1996, 1997). Phenotypes and their interpretations are described in the text. (A) Wild-type, where the capsule is not visible due to the fixation procedure, (B) kpsF mutant, (C) kpsT mutant, (D) kpsS mutant, (E) kpsC mutant, and (F) kpsE mutant.
synthesis and activation of *E. coli* catabolite-activator-protein-dependent genes such as those for maltose and β-galactosidase utilization (Karimova *et al.*, 1998). Thus, as long as the reporter strain lacks its own *cya*, activation is solely dependent on reconstituting the *B. pertussis* fragments through proximity mediated by the interacting fusion partners. The salient feature of this system for analysing bacterial multi-protein membrane assemblies is that the signalling cascade is spatially separated from the transcriptional reporter. For example, Landant and colleagues have provided evidence that the *E. coli* cell division apparatus involves multiple low-affinity interactions analogous to formation of the eukaryotic synapse (Karimova *et al.*, 2005). Using this two-hybrid system, we analysed a variety of protein–protein interactions using plasmids expressing combinations of kps–kps, kps–neu and neu–neu fusions (Steenbergen & Vimr, 2008). The conclusions shown in Fig. 3(B) indicate that the polymerase functions as a monomer, consistent with radiation target analysis (Vionnet *et al.*, 2006), but has affinity for both the IM–OM KpsE connector and the KpsC adaptor, which also interacts with itself. Note that the adaptor function of KpsC is used in the sense that it alters (adapts) the polymerase in some as yet unknown way so that polymerization is coupled to the exporter. In the absence of the adaptor, any group 2 polymer synthesized cannot be exported. If these interactions are interpreted correctly, they imply that positioning of group 2 polysaccharide synthetic functions relative to the export channel occurs through heterotypic interactions, possibly explaining the promiscuous nature of group 2-like polysaccharide export systems (Silver *et al.*, 2001). This hypothesis implies that the polymerases in different group 2 systems include conserved regions mediating the various interactions. It may be possible to identify these regions with the current two-hybrid system through mutation or by making peptide fusions, but ultimately crystal structures may be necessary if the interacting domains are strictly dependent on tertiary folding.

**Model of group 2 capsular polysialic acid biosynthesis in *E. coli* K1 and summary of outstanding questions concerning the dynamics of capsule export**

Although the details of building block assembly in the K1 system have been worked out for some time (Vimr *et al.*, 2004), since 2004 a series of previously unidentified acetyl transferases and O-acetyl esterases have been discovered that modify the polysialic acid chain or its precursors before or during export (Steenbergen *et al.*, 2006). However, acetylation modifies structure without affecting export. These modifications of the polysaccharide or its precursors suggest that the synthetic components of group 2 capsule biosynthesis may themselves be associated with the IM, and indeed the bifunctional NeuD protein has been shown to interact physically with NeuB and to exist in an IM-bound configuration (Anunziato *et al.*, 1995; Daines & Silver, 2000). Thus, all of the synthetic and export functions required for capsule biosynthesis might be interacting in a super-complex that we have designated the sialosome (Steenbergen & Vimr, 2008). It remains to be determined if the spatial organization of any synthetic proteins operating prior to the polymerase, NeuS, has a direct impact on export. Fig. 3(C) shows the synthetic steps involved in building-block synthesis, assembly and modification of the K1 capsule, and the homo- and heterotypic interactions between NeuS, KpsC and KpsE described above connecting synthesis to the export channel.

Are there any other gene products besides those shown in Fig. 3(C) that affect group 2 capsular polysaccharide export? McNulty *et al.* (2006) and Silver *et al.* (2001) provided evidence that RhsA and TolC, respectively, affected capsule synthesis or export. In particular, RhsA was suggested to be a carbohydrate-binding protein that could be chemically cross-linked to other components of the export channel and might connect capsule synthesis to export. However, it is difficult to rule out pleiotropic effects on membrane architecture as the explanation for RhsA or TolC phenotypes, especially since polypeptides such as GlgB and SucA involved in entirely separate pathways were also found cross-linked to the *E. coli* K5 export apparatus (McNulty *et al.*, 2006). Despite these outstanding questions and experimental ambiguities it would be useful to probe the export channel by cross-linking nascent capsular polysaccharide to protein components of the channel. It might be possible to approach this question by using a synchronized biosynthetic system making radiolabelled polysaccharide followed by cross-linking the carboxyl groups of nascent polysialic acid to adjacent amino groups in the channel. Such a synchronized system has been developed (Vimr, 1992) and already used to determine the extracellular structure of polysialic acid by NMR (Azurmendi *et al.*, 2007). Unfortunately, even if three-dimensional structures of group 2 export components were to become available, as in the group 1 system (Whitfield & Naismith, 2008), it is unlikely that static images of the components would tell us much new about the outstanding questions raised above regarding the dynamics of polysaccharide biosynthesis. We think that the two-hybrid system described above, and continued genetic manipulation of capsule genes coupled with simple biochemical and cell physiological approaches, will allow researchers to accurately infer the dynamics of group 2 capsule export.

While a complete *in vitro* reconstitution of the export process appears impossible at this time, our current understanding of the minimum set of components required for translocation suggests that progress might be made in the near future. In conclusion, there have been three fundamental questions in group 2 capsule biosynthesis. First, how is polysaccharide synthesis initiated/terminated? Second, is the coupling of synthesis to export directed or does it involve a potential cytoplasmic intermediate? Third, how is polysaccharide recognized by the export apparatus? While an answer to the first question remains mysterious, the second clearly involves directed
coupling mediated by heterotypic protein–protein interactions (Steenbergen & Vimr, 2008). Finally, given the immense structural polysaccharide diversity, export seems to require a signal that would presumably reside in the linkage to the terminal phospholipid moiety or that is the only common feature in different systems. However, our results provide an alternative explanation: the juxtapositioning of the polymerase through heterotypic protein–protein interactions might be sufficient to couple synthesis to export without the need for a specific export signal.

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References


