Domain Interactions Control Complex Formation and Polymerase Specificity in the Biosynthesis of the Escherichia coli O9a Antigen

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Sean D. Liston, Bradley R. Clarke, Laura K. Greenfield, Michele R. Richards, Todd L. Lowary, and Chris Whitfield

From the Department of Molecular and Cellular Biology, University of Guelph, Guelph, Ontario N1G 2W1 and the Alberta Glycomics Centre and Department of Chemistry, University of Alberta, Edmonton, Alberta T6G 2G2, Canada

Background: Synthesis of the E. coli O9a polysaccharide requires a multidomain polymerase and a terminator protein. Results: C-terminal deletions in the polymerase compromise fidelity or activity. Conclusion: Complex interactions between polymerase domains and the terminator are essential for accurate synthesis. Significance: These properties determine serotype specificity and may be exploited in glycoengineering to produce glycans with novel structures and applications.

The Escherichia coli O9a O-polysaccharide (O-PS) is a prototype for bacterial glycan synthesis and export by an ATP-binding cassette transporter-dependent pathway. The O9a O-PS possesses a tetrasaccharide repeat unit comprising two α-(1→2)- and two α-(1→3)-linked mannose residues and is extended on a polysisoprenoid lipid carrier by the action of a polymerase (WbdA) containing two glycosyltransferase active sites. The N-terminal domain of WbdA possesses α-(1→2)-mannosyltransferase activity, and we demonstrate in this study that the C-terminal domain is an α-(1→3)-mannosyltransferase. Previous studies established that the size of the O9a polysaccharide is determined by the chain-terminating dual kinase/methyltransferase (WbdD) that is tethered to the membrane and recruits WbdA into an active enzyme complex by protein-protein interactions. Here, we used bacterial two-hybrid analysis to identify a surface-exposed α-helix in the C-terminal mannosyltransferase domain of WbdA as the site of interaction with WbdD. However, the C-terminal domain was unable to interact with WbdD in the absence of its N-terminal partner. Through deletion analysis, we demonstrated that the α-(1→2)-mannosyltransferase activity of the N-terminal domain is regulated by the activity of the C-terminal α-(1→3)-mannosyltransferase. In mutants where the C-terminal catalytic site was deleted but the WbdD-interaction site remained, the N-terminal mannosyltransferase became an unrestricted polymerase, creating a novel polymer comprising only α-(1→2)-linked mannose residues. The WbdD protein therefore orchestrates critical localization and coordination of activities involved in chain extension and termination. Complex domain interactions are needed to position the polymerase components appropriately for assembly into a functional complex located at the cytoplasmic membrane.

Many different macromolecules containing complex carbohydrates (glycoconjugates) are found on the surfaces of living cells. These structures play crucial roles in the interactions between the cell and its environment or in communications and interactions between one cell and another. Glycoconjugates on the surfaces of bacteria are important in survival against factors in the environment or against the protective responses of mammalian or plant hosts. The function of a particular glycoconjugate is dictated by its structure, and the precise order of sugars and linkages in a glycan is dictated by the specificities of glycosyltransferase (GT) enzymes. Although 97 GT families are recognized in the Carbohydrate-Active Enzyme (CAZy) database (1) based on bioinformatics or structural criteria (2), the features dictating acceptor/donor specificity are often unknown and cannot be inferred from either catalytic fold or GT family (3). Most GTs catalyze the formation of a single glycosidic linkage, but increasing numbers of polymerases are being identified. These enzymes sequentially add multiple residues to an acceptor molecule, creating complex glycans of varying lengths without a template. Examples are found in prokaryotes and eukaryotes, and some form highly important products such as cellulose, chitin, hyaluronic acid, chondroitin, and polysaccharide (4, 5). Depending on the enzyme and the complexity of the product, polymerases may contain one or more GT active site(s). Polymerases operate by

* The abbreviations used are: GT, glycosyltransferase; O-PS, O-polysaccharide; GlcNAc, N-acetylglucosamine; Manp, mannose; LB, lysogeny broth; BisTris, 1,5-bis[2-hydroxyethyl]amino)-2-[hydroxymethyl]propane-1,3-diol; TOCSY, total correlation spectroscopy; tROESY, transverse rotating frame Overhauser enhancement spectroscopy; GalpNAC, N-acetylgalactosamine; Glc, glucose; GlcP, gluconic acid; X-Gal, 5-bromo-4-chloro-3-indolyl β-D-galactopyranoside.

† This article contains supplemental Table S1.

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||To whom correspondence should be addressed: Dept. of Molecular and Cellular Biology, Science Complex, University of Guelph, Guelph, Ontario N1G 2W1, Canada. Tel.: 519-824-4120 (ext. 53361); Fax: 519-837-3273; E-mail: cwhitfiel@uoguelph.ca.

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either “processive” or “distributive” mechanisms (6, 7). Processive enzymes catalyze multiple rounds of glucose transfer to a given acceptor while maintaining the growing glycan in a single active site. In contrast, strictly distributive enzymes possess one or more active sites and release the glycan product after each addition. The structural principles that guide efficiency and fidelity of distributive multidomain polymerase enzymes are largely unknown. The polymerase involved in the biosynthesis of the lipopolysaccharide (LPS) O9a O-polysaccharide (O-PS) antigen in Escherichia coli provides a prototype for addressing these questions.

The O9a system is a representative of the widespread ATP-binding cassette transporter-dependent assembly pathway (8, 9). The pathway (see Fig. 1A) begins with the transfer of N-acetylglucosamine 1-phosphate to a 55-carbon polyisoprenoid lipid acceptor (undecaprenol phosphate) catalyzed by WecA (10, 11). Undecaprenol diphosphate-GlcNAc is then committed to O9a biosynthesis by the action of WbdC and WbdB, two GDP-mannose (GDP-Manp)-dependent mannosyltransferases that add a trimannose adapter region (12). These enzymes and the resulting structure are conserved in E. coli serotypes O9a, O9, and O8, which produce structurally related O-PSs differing in the type and sequence of linkages in their repeat units (9). A serotype-specific multidomain polymerizing mannosyltransferase (WbdA) extends this adapter to create the precise repeating unit O-PS (12, 13). In E. coli O9a, the peripheral membrane protein WbdD terminates polymerization by adding a methyl phosphate to the non-reducing end of the nascent O9a polymer (14–16). This terminal modification is required for recognition and export of the completed O-PS across the cytoplasmic membrane by its cognate ATP-binding cassette transporter. The recognition event is mediated by the nucleotide-binding domain polypeptide of the transporter, which possesses a serotype-specific carbohydrate-binding module that only binds terminated O-PS chains (17, 18). The WbdD terminator plays an additional pivotal structural role in recruiting WbdA to the membrane (19). The stoichiometry of WbdA:WbdD in active complexes is a critical factor in establishing the chain length distribution of the resulting glycans, and the “variable geometry” model has been developed to explain this process (20). The key element in the model is that complexes with different stoichiometries have different physical sizes that generate a range of glycan chain lengths. Implicit in this model is a structural element that serves as a molecular ruler to establish chain length. This role is fulfilled by an extended α-helical bundle that holds the WbdD trimer together and maintains a critical distance between the catalytic sites of the terminator and its site of interaction with the WbdA polymerase (21). WbdD is therefore a central player in a sophisticated quality control system that dictates the distribution of chain lengths and marks those chains with a terminal export tag. After export, the O-PS is transferred to the lipid A-core portion of LPS, translocated to the cell surface, and inserted into the outer membrane (22, 23).

The WbdA polymerase possesses a modular design comprising two identifiable GT(family)4 domains as defined by the CAZy classification system (1, 2). The two GT4 active sites correlate with the presence of α-(1→2) and α-(1→3) linkages in the O9a tetrasaccharide repeat unit (13) (Fig. 1A). Coexpression of the two domains (with coding sequences cloned in separate plasmids) restores O9a biosynthesis in an E. coli wbdA deletion mutant, but neither domain shows any in vivo activity when expressed alone (13). Surprisingly, when examined in vitro with synthetic oligosaccharide acceptors, the purified N-terminal domain (WbdAN) transfers multiple Manp residues to the synthetic acceptor to create α-(1→2)-linked polypeptide (X-Gal) (40 µg/ml), and isopropyl β-d-thiogalactopyranoside (0.5 mM).

**DNA Methods—**Pwo DNA polymerase (Roche Applied Science) was used to PCR amplify DNA fragments used in cloning. The oligonucleotide primers (Sigma) used for PCR introduced restriction endonuclease sites and epitope tags; their particular features are described in supplemental Table S1. DNA fragments were purified using the PureLink PCR Purification kit (Invitrogen). The PureLink Quick Plasmid Miniprep kit (Invitrogen) was used to purify plasmid DNA. Restriction endonucleases (Invitrogen and New England Biolabs) and T4 DNA ligase (New England Biolabs) were used as described by the manufacturers. PfuUltra high fidelity DNA polymerase (Stratagene) was used for site-directed mutagenesis of plasmids according to the QuikChange method (Stratagene). All DNA constructs were assessed by restriction endonuclease digestion and verified by DNA sequencing. Sequencing reactions were performed by the Genomics Facility at the Advanced Analysis Center (University of Guelph).

**Generation of WbdA Constructs—**Nested C-terminal truncations of WbdA (see Fig. 1B) were generated in pWQ631 using complementary oligonucleotide primers containing base changes that encoded a stop codon and a unique restriction site for screening purposes. PCR products were digested using DpnI and transformed into E. coli Top10. For complementation experiments, each of the plasmids encoding truncated WbdA derivatives also contained sequences encoding the
C-terminal fragment of WbdD (residues 475–708). This fragment has been used in previous work to increase the quantity of O-PS synthesized by WbdA constructs (13) and was used here for the same purpose. For protein expression and purification, DNA fragments encoding WbdA (and its various derivatives) were cloned into pUT18 and pUT18C to create N- and C-terminal fusions of WbdA with the T18 fragment of the adenylate cyclase. To assess protein-protein interactions, DNA fragments were cloned behind the pBAD arabinose-inducible promoter (26). To express WbdA and its C-terminally truncated derivatives, DNA fragments encoding WbdA (and its various derivatives) were cloned into pUT18 and pUT18C to create N- and C-terminal fusions of WbdA with the T18 fragment of the Bordetella pertussis adenylate cyclase.

Expression of WbdA and Its C-Terminal Truncated Derivatives—5-ml cultures of E. coli CWG1105 (ΔwbdA) transformed with pWQ631-based plasmids were grown in LB medium at 37 °C until the A600 reached 0.3. Growth was continued at 20 °C until the A600 reached 0.5 after which expression was induced overnight at 20 °C with L-arabinose (0.01%, w/v) and incubation was continued for 6 h. Culture volumes containing equivalent amounts of cells (as determined by optical density) were subjected to centrifugation at 5000 × g for 10 min, and the pellets were resuspended in SDS-PAGE loading buffer (27).

Purification of Decahistidine-tagged WbdA Derivatives—This protocol was adapted from methods described previously (13). In brief, protein was purified from exponential phase 500-ml cultures of E. coli BL21 transformed with plasmids pWQ492, pWQ769, and pWQ590 encoding WbdA-His10, WbdA E317A-His10, and WbdA E758A, respectively. These plasmids are all pBAD derivatives (26), and gene expression was induced overnight at 20 °C with L-arabinose as described above. Cells were collected by centrifugation at 100,000 × g for 60 min to remove cell membranes. Purification was performed using an AKTA Explorer system (GE Healthcare) and a 5-ml HiTrap chelating HP column charged with Ni2+ (GE Healthcare). Individual lysates containing His10-tagged WbdA derivatives were loaded at 4 ml/min, and the column was washed sequentially with 5 column volumes of buffer A containing 0, 50, 75, 125, and 250 mM imidazole. Fractions containing the His10-tagged WbdA derivatives were pooled, exchanged into Buffer B (20 mM BisTris, 250 mM NaCl, 5% (v/v) glycerol, pH 7.0) supplemented with Complete Mini EDTA-free protease inhibitor tablets (Roche Applied Science). Cells were lysed using an EmulsiFlex homogenizer (Avestin), and the lysate was cleared by centrifugation at 5000 × g for 10 min. The cell-free lysate was treated with 5 μl of Benzonase nuclease (Novagen) for 15 min at room temperature and then centrifuged at 100,000 × g for 60 min to remove cell membranes. Purification was performed using an AKTA Explorer system (GE Healthcare) and a 5-ml HiTrap chelating HP column charged with Ni2+ (GE Healthcare). Individual lysates containing His10-tagged WbdA derivatives were loaded at 4 ml/min, and the column was washed sequentially with 5 column volumes of buffer A containing 0, 50, 75, 125, and 250 mM imidazole. Fractions containing the His10-tagged WbdA derivatives were pooled, exchanged into Buffer B (20 mM BisTris, 50 mM NaCl, pH 7.0), concentrated using a Vivaspin column (50,000 molecular weight cutoff; Sartorius Biolab Products), and stored for subsequent use in in vitro enzyme

### TABLE 1

| Strain or plasmid | Genotype or property | Source or Ref. |
|-------------------|----------------------|----------------|
| **E. coli**       |                      |                |
| Top10             | F′ c MrA, Δ(mrr-ladRMS-mccBC), φ80, lacZAM15, ΔlacX74, deoR, recA1, araD139, Δ ara-leu)7697, gall, galk, rpsL mutations, endA1 | Invitrogen      |
| BL21             | B′ dcmomp7 lacZΔM15 tetR (maltB–p12) lacY1, Δ(V, lacI, lacZΔM15, tetR (maltB–p12)) | Novagen         |
| BH1101           | lacZΔM15, recA1, endA1, glvV44A(S), Na⁺ | 54             |
| CGW1105          | 909KΔ; trp hsd rpsL, rpsL S, mcr A, ΔwbdA; Sm³; Tc² | 12             |
| **Plasmid**      |                      |                |
| pBAD24          | Plasmid vector with L-arabinose-inducible promoter; Ap⁰ | 26             |
| pKT25          | Bacterial two-hybrid vector with sequence encoding the T25 fragment of B. pertussis adenylate cyclase upstream of a multiple cloning site; Km⁰ | 54             |
| pUT18         | Bacterial two-hybrid vector with sequence encoding the T18 fragment of B. pertussis adenylate cyclase downstream of a multiple cloning site; Ap⁰ | 54             |
| pUT18C        | Bacterial two-hybrid vector with sequence encoding the T18 fragment of B. pertussis adenylate cyclase downstream of a multiple cloning site; Ap⁰ | 54             |
| pKT25-Zip     | pKT25 derivative encoding the CyaA T25 fragment fused to the leucine zipper domain of the yeast Gcn4 protein; Km⁰ | 54             |
| pUT18-Zip      | pUT18 derivative encoding the CyaA T18 fragment fused to the leucine zipper domain of the yeast Gcn4 protein; Ap⁰ | 54             |
| pWQ284     | pBAD24 derivative containing a chloramphenicol resistance cassette; Cm⁰ | 18             |
| pWQ486     | pKT25 derivative encoding T25-WbdD, Km⁰ | 19             |
| pWQ487     | pUT18C derivative encoding T18-WbdA, Ap⁰ | 19             |
| pWQ492     | pBAD24 derivative encoding WbdA-His10, Ap⁰ | 19             |
| pWQ589     | pBAD24 derivative encoding a kanamycin resistance cassette; Km⁰ | 13             |
| pWQ590     | pBAD24 derivative encoding His10-WbdA, Ap⁰ | 13             |
| pWQ591     | pWQ284 derivative encoding His10-WbdA, Ap⁰ | 13             |
| pWQ592     | pWQ631 derivative encoding WbdA Δ727–840, Km⁰ | 13             |
| pWQ630     | pWQ631 derivative encoding WbdA Δ727–840, Km⁰ | 13             |
| pWQ631     | pWQ589 derivative encoding WbdA Δ727–840, WbdA, Km⁰ | 13             |
| pWQ765     | pWQ591 derivative encoding WbdA Δ727–840, WbdA (frameshift mutation after amino acid 635, now encodes S⁵⁶⁵RPRQRTYWHAG–); Km⁰ | This study |
| pWQ766     | pWQ592 derivative encoding WbdA Δ427–840, Ap⁰ | This study |
| pWQ769     | pWQ592 derivative encoding WbdA Δ427–840, H11002, Ap⁰ | This study |
| pWQ770     | pWQ631 derivative encoding WbdA Δ475–708, WbdA Δ1–435, Km⁰ | This study |
| pWQ771     | pWQ631 derivative encoding WbdA Δ475–708, WbdA Δ1–435, Km⁰ | This study |
| pWQ772     | pWQ631 derivative encoding WbdA Δ475–708, WbdA Δ1–435, Km⁰ | This study |
| pWQ773     | pWQ631 derivative encoding WbdA Δ475–708, WbdA Δ1–435, Km⁰ | This study |
| pWQ774     | pWQ631 derivative encoding WbdA Δ475–708, WbdA Δ1–435, Km⁰ | This study |
| pWQ775     | pUT18C derivative encoding WbdA Δ475–708, WbdA Δ1–435, Ap⁰ | This study |
| pWQ776     | pUT18C derivative encoding WbdA Δ475–708, WbdA Δ1–435, Ap⁰ | This study |
| pWQ777     | pUT18C derivative encoding WbdA Δ475–708, WbdA Δ1–435, Ap⁰ | This study |
| pWQ778     | pUT18C derivative encoding WbdA Δ475–708, WbdA Δ1–435, Ap⁰ | This study |
| pWQ779     | pUT18C derivative encoding WbdA Δ475–708, WbdA Δ1–435, Ap⁰ | This study |
| pWQ780     | pUT18C derivative encoding WbdA Δ475–708, WbdA Δ1–435, Ap⁰ | This study |
| pWQ781     | pUT18C derivative encoding WbdA Δ475–708, WbdA Δ1–435, Ap⁰ | This study |
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Reactions. The concentrations of WbdA-His$_{10}$ (30 mg/ml), WbdA$_{E317A}$-His$_{10}$ (4.5 mg/ml), and His$_{10}$-WbdA$_{N}$ (5 mg/ml) were estimated from A$_{280}$ values, and the theoretical extinction coefficients were predicted by ProtParam (28).

Size Exclusion Chromatography—Purified WbdA-His$_{10}$ in Buffer A containing 250 mM imidazole was dialyzed with Buffer C (50 mM BisTris, 150 mM NaCl, pH 7.0). WbdA-His$_{10}$ (0.2 mg at 2 mg/ml) was eluted through a Superose 6 HR 10/30 size exclusion column (GE Healthcare) with Buffer C at 0.5 ml/min. Molecular mass was estimated from a standard curve derived from the standards β-amylase (200 kDa), alcohol dehydrogenase (150 kDa), bovine serum albumin (66 kDa), ovalbumin (44 kDa), and carbonic anhydrase (29 kDa).

**Bacterial Two-hybrid Interactions**—E. coli BTH101 was transformed with the appropriate two-hybrid plasmid combinations. The transformed cell suspensions were serially diluted in LB and plated on LB agar containing X-Gal, antibiotics, isopropyl β-D-thiogalactopyranoside, and 3-aminobenzoic acid (if required). Plates were incubated at 30 °C for 48 h before two-hybrid interactions were assessed visually for the presence of blue colonies. For quantitative hybrid interactions were assessed visually for the presence of antigen, whereas acceptor B (H9251) represents the repeat unit of the O$_9$a mannose residue as expected.

**RESULTS**

**In Vitro Activity of the C-terminal GT4 Domain**—Previously, we identified the N-terminal GT4 domain (WbdA$^N$) as the source of α-(1→2)-mannosyltransferase activity (12). However, no in vitro mannosyltransferase activity was detected with the C-terminal (putative) GT4 domain (WbdA$^C$) in either the presence or absence of its N-terminal partner. The observation that a full-length protein bearing a mutation in the EX$_2$E motif of the putative C-terminal GT4 active site (WbdA$^{E758A}$; Fig. 1B) could synthesize α-(1→2)-linked polymannose led us to test the prediction that the C-terminal GT4 domain possessed α-(1→3)-mannosyltransferase activity. We therefore investigated the activity of the corresponding mutant (WbdA$_{E317A}$), inactivated in the N-terminal EX$_2$E motif, using synthetic oligosaccharide acceptors. This construct showed activity with both acceptors (data not shown) as was found with wild type WbdA, but only acceptor B was examined in depth because its structure (α-Man$_{(1→3)}$-α-Man$_{(1→3)}$-β-GlcpNAC) represents the conserved reducing terminal tri saccharide of the O$_9$a antigens. Standard reactions were performed for 30 min in 250-μl reaction volumes with 10 μM enzyme, 0.5 mM acceptor, and 5 mM GDP-Man$_p$ in Buffer D (50 mM HEPES, 20 mM MgCl$_2$, 1 mM dithiothreitol, pH 7.5). Product generation was initially verified by thin layer chromatography using AL SIL G TLC plates (Whatman) developed in ethyl acetate:water:1-butanol:acetic acid (5:4:4:2.5). Fluorescent reaction products were detected with a hand-held UV lamp. Reaction mixtures were diluted in 1 ml of water and loaded onto a C$_{18}$ Sep-Pak cartridge (Waters). The cartridge was washed extensively with water, and the products were eluted in 2 ml of 60% (v/v) aqueous acetonitrile and concentrated using a SpeedVac concentrator. MALDI-TOF mass spectra of the reaction products were obtained on a Bruker ultraflxXtreme MALDI-TOF/TOF in negative ion mode. All NMR spectra were acquired in D$_2$O at 27 °C on an Agilent VNMR 700-MHz spectrometer equipped with a cryoprobe. The spectra were referenced to an external standard of acetone (2.22 ppm for 1'H and 31.07 ppm for 13'C at 27 °C).

**Protein and LPS PAGE**—For routine monitoring of proteins, cells were solubilized in SDS-PAGE loading buffer (27). LPS samples were prepared by digestion of these whole-cell lysates with proteinase K (30). Both protein and LPS samples were heated at 100 °C for 10 min prior to analysis by SDS-PAGE in Tris-glycine buffer (27). Silver staining was used to visualize LPS (31), and Simply Blue (Invitrogen) was used to visualize protein. For Western immunoblotting, protein and LPS samples were transferred to nitrocellulose membranes (Protran, PerkinElmer Life Sciences). LPS immunoblots were probed with O9a-specific rabbit antiserum (14) and alkaline phosphatase-conjugated goat anti-rabbit secondary antibody (Cedarlane Laboratories) and developed using nitroblue tetrazolium and 5-bromo-4-chloro-3-indolyl phosphate (Roche Applied Science) substrates. Protein immunoblots were probed with WbdA-specific rabbit antiserum (19) or mouse pentahistidine-specific monoclonal antibody (Qiagen), and the secondary anti bodies were either horseradish peroxidase-conjugated goat anti-rabbit (Sigma) or horseradish peroxidase-conjugated goat anti-mouse (Jackson ImmunoResearch Laboratories). Protein immunoblots were developed using Luminata Classic (Millipore) chemiluminescent substrate.

Bioinformatics Analyses—Molecular modeling of WbdA$^C$ was performed using the Phyre$^2$ server in the intensive mode (32). Conserved domains were identified by the NCBI conserved domain database (13, 33).
indicative of H3 of the ring with an anomeric signal at 5.24 ppm. No signals at 5.08 ppm, and the signal at 5.08 ppm showed a correlation to H3 of the ring with an anomeric signal at 5.12 ppm, the signal at 5.14 ppm showed a correlation to H3 of the ring.

In the tROESY spectrum, additional NMR spectra, including COSY, TOCSY, and tROESY spectra (data not shown). Specifically, in the tROESY spectrum, the signal at 5.14 ppm showed a correlation to H3 of the ring with an anomeric signal at 5.24 ppm. No signals indicative of α-(1→2) linkages were observed. These results were consistent with the values predicted by the CASPER NMR analysis program (34–36) (accessed October 28, 2014) (Table 2). We then performed a reaction with the same acceptor incubated with WbdAE317A and WbdAN. When analyzed in isolation, WbdAN extends a synthetic acceptor to generate larger products are possible (Fig. 2C).

A Region within the C-terminal Part of WbdA Is Required for in Vivo Activity of the N-terminal GT4 Domain—With the activities of the two GT4 domains now assigned, we turned our attention to the molecular basis for the lack of detectable in vivo activity of WbdAN despite its ability to restore authentic O9a production in cells co-transformed with plasmids expressing WbdAC (13). Furthermore, when examined in vitro, WbdAN showed robust poly-α-(1→2)-mannosyltransferase activity, and the same activity was replicated in vivo in cells expressing WbdA<sup>E758A</sup>, which possesses a mutation in the catalytically essential EX-E motif in the C-terminal GT4 active site (13). As reported previously, lysates of cells expressing WbdA<sup>E758A</sup> lack the typical O9a LPS banding pattern (reflecting size increments of a tetrasaccharide O-PS repeat unit), and LPS molecules with the aberrant α-(1→2)-linked O-PS no longer reacted with anti-O9a antibodies in Western immunoblots (Fig. 3, A and B). The same LPS phenotype was evident in a strain expressing a wbdA construct where a stop codon was inadvertently introduced, creating a protein (designated WbdA*) that possessed a frame-shift mutation at residue 635 and terminated prematurely at residue 649 (Fig. 3, A and B). The collective results suggest that in vivo poly-α-(1→2)-mannosyltransferase activity of the N-terminal GT4 domain is dependent on a region of the C-terminal domain and that this is missing in WbdAN.

*Figure 1. Biosynthesis of the O9a antigenic polysaccharide in E. coli and a schematic representation of the WbdA constructs used in this study.* In A, the structure of the undecaprenol diphosphate (Und-PP)-linked biosynthetic intermediate is shown. Sugars are represented using the nomenclature established by the Consortium for Functional Glycomics. The enzymes responsible for the formation of each part of the glycan are identified. The polymerase, WbdA, is solely responsible for the extension of the repeat unit region of the glycan, extending a trisaccharide adapter. B describes the WbdA active site mutants, domains, and truncated proteins whose function was examined. Domain predictions were identified using the NCBI Conserved Domain Database (33). This allowed the separation of WbdAN and WbdAC, which retain activity and synthesize authentic O9a polysaccharide when coexpressed in E. coli ΔwbdA (13). From MS data. This compound generated major anomic peaks at 5.24, 5.14, 5.12, and 5.08 ppm. These signals could be assigned to Manp with α-(1→3) linkages through two-dimensional NMR spectra, including COSY, TOCSY, and tROESY spectra (data not shown). Specifically in the tROESY spectrum, the signal at 5.14 ppm showed a correlation to H3 of the ring with an anomeric signal at 5.12 ppm, the signal at 5.12 ppm showed a correlation to H3 of the ring with an anomeric signal at 5.08 ppm, and the signal at 5.08 ppm showed a correlation to H3 of the ring with an anomeric signal at 5.24 ppm. No signals indicative of α-(1→2) linkages were observed. These results were consistent with the values predicted by the CASPER NMR analysis program (34–36) (accessed October 28, 2014) (Table 2). We then performed a reaction with the same acceptor incubated with WbdA<sup>E317A</sup> and WbdAN. When analyzed in isolation, WbdAN extends a synthetic acceptor to generate α-(1→2)-linked polymannose (13), but in combination, these two constructs reconstituted authentic O9a biosynthesis. The 1H NMR spectrum of these products revealed anomeric signals at 4.99, 5.08, 5.25, and 5.33 ppm (Fig. 2D). These values are consistent with the predictions from the CASPER program for the O9a repeat unit (Table 2), the published values for the authentic O9a glycan (37), and the in vitro products obtained with the same acceptor and full-length WbdA (12). The MS spectrum revealed a series of products, each differing by a single mannose residue (m/z 162). The largest products of significant quantity showed 24 added residues, equivalent to six (tetrasaccharide) repeat units of the authentic O9a glycan, but additional peaks with intensities near the background level suggest that larger products are possible (Fig. 2C).
To determine the minimal portion of \( \text{WbdA} \) required for \textit{in vivo} poly-\( \alpha-(1 \rightarrow 3)\)-mannosyltransferase activity, a series of C-terminal truncations of \( \text{WbdA} \) was constructed guided by \( \text{WbdA}^{*} \) as a starting point and the location of the N-terminal GT4 domain as an end point (Fig. 1B). \textit{E. coli} \( \Delta \text{wbdA} \) cells expressing the \( \text{WbdA}^{1–635} \) construct (equivalent to \( \text{WbdA}^{*} \)) produced the \( \text{O}-(1 \rightarrow 3)\)-polymannose glycan as expected. The shortest functional construct was \( \text{WbdA}^{1–568} \), whereas \( \text{WbdA}^{1–547} \) and proteins with more extensive C-terminal deletions were all unable to restore any O-PS synthesis. All of the constructs generated \( \text{WbdA} \) proteins (Fig. 3, C and D), indicating that inactivity did not simply reflect selective proteolytic degradation of some truncated versions of \( \text{WbdA} \). The expression level of the truncated products was slightly less than those of the full-length proteins based on Western immunoblotting. This does not necessarily correlate with protein amount due to loss of epitopes (through truncation) recognized by the rabbit anti-\( \text{WbdA} \) antibodies, but it is consistent with the absence of visible bands in the stained SDS-polyacrylamide gel for any construct shorter than the full-length protein. A reduction in the amount of \( \text{WbdA} \) may explain the slight downward shift in the size distribution of the \( \text{O}-(1 \rightarrow 3)\)-polymannose glycans they generate (Fig. 3A) because alterations in stoichiometry of \( \text{WbdA:WbdD} \) are known to effect chain length (14, 20).

**TABLE 2**

Comparison of the chemical shifts for the anomeric protons of the products generated in \textit{in vitro} reactions with those predicted by the CASPER database for the structures shown

|       | A   | B   | C   | D   | E   |
|-------|-----|-----|-----|-----|-----|
| 1, Experimental | 5.14 | 5.12 | 5.08 | 5.24 | --  |
| 1, CASPER    | 5.11 | 5.07 | 5.07 | 5.20 | 4.74 |
| 2, Experimental | 4.99 | 5.25 | 5.33 | 5.08 | --  |
| 2, CASPER    | 5.02 | 5.29 | 5.33 | 5.09 | --  |

*The HOD peak at 4.76 ppm is too large to see whether the anomeric signal for \( \beta\)-D-GlcNAc is present.*

To determine the minimal portion of \( \text{WbdA} \) required for \textit{in vivo} poly-\( \alpha-(1 \rightarrow 2)\)-mannosyltransferase activity, a series of C-terminal truncations of \( \text{WbdA} \) was constructed guided by \( \text{WbdA}^{*} \) as a starting point and the location of the N-terminal GT4 domain as an end point (Fig. 1B). \textit{E. coli} \( \Delta \text{wbdA} \) cells expressing the \( \text{WbdA}^{1–635} \) construct (equivalent to \( \text{WbdA}^{*} \)) produced the \( \alpha-(1 \rightarrow 2)\)-polymannose glycan as expected. The shortest functional construct was \( \text{WbdA}^{1–568} \), whereas \( \text{WbdA}^{1–547} \) and proteins with more extensive C-terminal deletions were all unable to restore any O-PS synthesis. All of the constructs generated \( \text{WbdA} \) proteins (Fig. 3, C and D), indicating that inactivity did not simply reflect selective proteolytic degradation of some truncated versions of \( \text{WbdA} \). The expression level of the truncated products was slightly less than those of the full-length proteins based on Western immunoblotting. This does not necessarily correlate with protein amount due to loss of epitopes (through truncation) recognized by the rabbit anti-\( \text{WbdA} \) antibodies, but it is consistent with the absence of visible bands in the stained SDS-polyacrylamide gel for any construct shorter than the full-length protein. A reduction in the amount of \( \text{WbdA} \) may explain the slight downward shift in the size distribution of the \( \alpha-(1 \rightarrow 2)\)-polymannose glycans they generate (Fig. 3A) because alterations in stoichiometry of \( \text{WbdA:WbdD} \) are known to effect chain length (14, 20).
In Vivo Poly-α-(1→2)-mannosyltransferase Activity Correlates with the Ability of WbdD to Recruit the Truncated WbdA Derivatives—Previously, we demonstrated that WbdA activity was dependent on protein-protein interactions with WbdD, which recruits the polymerase to a functional complex on the cytoplasmic membrane (19). The C-terminal region of WbdD involved in this interaction was localized to a predicted 30-residue C-terminal helix, but the region of WbdA involved was not investigated. The in vivo activities for the truncated WbdA proteins could be explained by an essential interacting region in the C-terminal part of WbdA that is deleted in some constructs, and bacterial two-hybrid results proved this hypothesis to be correct. Interactions were measured between the T25-WbdD fusion protein and WbdA variants fused to an N-terminal T18 fragment. This was based on the high level of β-galactosidase activity seen with the full-length proteins (Table 3), consistent with data published previously (19). With the full-length proteins, these positions of the CyaA fragments had no effect on WbdA or WbdD activity (19). Positive interactions were detected between T25-WbdD and T18-WbdA<sup>1-635</sup> and between T25-WbdD and T18-WbdA<sup>1-568</sup>, whereas none of the constructs unable to synthesize the poly-α-(1→2)-mannan demonstrated any significant interaction with WbdD.

Interactions between WbdD and the C-terminal WbdA Domain Require the Participation of the N-terminal Domain—The two-hybrid data reinforce the need for a productive association between WbdA and WbdD for polymerase activity. Because co-expression of WbdAN and WbdAC restored O9a biosynthesis in an E. coli ΔwbdA mutant (13), an interaction between WbdAN and WbdD was anticipated. Unexpectedly, this combination showed no interaction (Table 3), so two-hy-
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WbdA is a two-domain polymerizing GT responsible for the synthesis of a glycan containing a tetrasaccharide repeat unit. The N-terminal GT4 catalytic site has α-(1→2)-mannosyltransferase activity, and we now can assign α-(1→3)-mannosyltransferase activity to the C-terminal GT4 site. The in vitro product profiles obtained using WbdA at different substrate concentrations are consistent with WbdA being a distributive enzyme, releasing product after each mannosyl transfer reaction (12). Reproducible production of a tetrasaccharide in which two α-(1→2)-linked Manp residues are always followed by two α-(1→3)-linked Manp residues represents an interesting challenge for such enzymes. Enzymes exist where a single GT site sequentially adds a defined number of residues to an acceptor. For example, PglH is a UDP-Gal pNac-dependent GT involved in N-glycosylation of proteins in Campylobacter jejuni (38), and it is proposed that its ability to reliably add three residues is precisely controlled by the relative binding affinities of the enzyme for the growing acceptor (39). An involvement of the active sites of WbdA in its fidelity is suggested by analysis of the WbdA protein from serotype O9a and its homolog from serotype O9 with the latter O-PS containing a pentasaccharide repeat unit (three α-(1→2)-linked Manp residues followed by two α-(1→3)-linked Manp residues). A mutation (C80R) in the

**DISCUSSION**

WbdA Exists as a Monomer—Although WbdA and WbdD clearly interact and WbdD is a trimer (16), the possibility existed of further homotypic interactions involving WbdA. In two-hybrid analyses, no interactions could be detected regardless of the locations of the CyaA fragment reporters (data not shown). To confirm this, WbdA-His10 was purified and examined by size exclusion chromatography. The majority of the protein was eluted as a monomer with only a (variable) trace of aggregate evident in the profile (Fig. 5).

**FIGURE 4.** Active synthesis of α-(1→2)-polymannose precludes restoration of O9a with an active WbdAC α-(1→3)-mannosyltransferase domain in trans. E. coli CWG1105 cells were co-transformed with compatible plasmids encoding the identified WbdA C-terminal deletions and WbdAC (A and B). C and D show the results of complementation experiments where plasmids encoding derivatives with EX3 catalytic site mutations were co-transformed with plasmids expressing the individual domains. The LPS products in whole-cell lysates were revealed by silver-stained SDS-PAGE (A and C) and Western immunoblotting with antibodies specific for the O9a antigen (B and D).
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FIGURE 5. Size exclusion chromatography of purified WbdA-His10. WbdA-His10 was purified (inset) and eluted through a Superose 6 HR 10/30 column. The predominant peak corresponded with an estimated molecular mass of 120 kDa consistent with monomeric WbdA-His10 (theoretical molecular mass, 96.1 kDa). A minor amount of the protein eluted before the 200-kDa standard. The molecular masses of the protein standards are as follows: β-amylase, 200 kDa; alcohol dehydrogenase, 150 kDa; bovine serum albumin, 66 kDa; ovalbumin, 44 kDa. mAU, milli-absorbance units.

FIGURE 6. Structural model of WbdAC highlighting the region of interaction with WbdD. Residues 440–832 of WbdA were modeled with high confidence using Phyre2 in the intensive mode. The region of interaction identified by deletion analysis (highlighted in red) corresponds to a surface-exposed α-helix located between Val548 and Ala556. GDP-Manp (colored blue) was modeled into the active site through structural alignment with mycobacterial PimA (Protein Data Bank code 2GEJ), a GDP-Manp-dependent GT4 mannosyltransferase that is involved in the biosynthesis of phosphoinositol mannosides (49), that was crystallized in the presence of GDP-Manp. Glutamate residues of the WbdA EXE motif are colored green.

formation of the binding pocket of the C-terminal GlcPA-transferase, creating an unusual orientation of the substrate in the catalytic site. The significance of this feature is also unknown. Clearly, KfoC and WbdA both demonstrate complex domain interactions essential for function, but additional crystal structures are needed to determine the mechanistic consequences and establish whether any generic architectural principles are involved.

Organization of WbdA into a functional active biosynthesis complex requires its interaction with WbdD; there is no known terminator component for KfoC (19). This process presumably engages WbdA at the membrane with the natural undecaprenol diphosphate-linked oligosaccharide acceptor and creates the proper structural context for chain termination (14–16, 20). Previously, the interaction domain in WbdD was localized to a 30-residue α-helix at the C terminus (19). In analysis of the C-terminal deletions of WbdA, interaction was still possible with WbdA1–568 but was lost in WbdA1–547. This region can be modeled using other GT4 enzymes as a scaffold, and the structure of mycobacterial PimA, a GDP-Manp-dependent GT4 mannosyltransferase that is involved in the biosynthesis of phosphoinositol mannosides (49) can be used to predict substrate binding locations (Fig. 6). The region of interaction identified by deletion analysis extends from residues Val548–Ile568 (highlighted in red) and encompasses a surface-exposed α-helix located between Val548 and Ala556 that represents an excellent candidate for interaction. Interestingly, the interactions between WbdA and WbdD are complex. Although the WbdA C domain possesses the interaction site and shows correct catalytic activity when expressed with its WbdAN partner, interactions between WbdAC and WbdD were dependent on the presence of WbdAN. As expected, WbdAN could only interact with WbdD when WbdAC provided the interaction site. Productive interactions were compromised when inactivated GT domains were present. The results indicate that the separated GT domains are able to recreate the correct configuration for authentic O9a polymerase activity in vivo. However, the same is not true in vitro as the same combination can only generate the α-(1→2)-polymannose glycan (13). In this situation, WbdAC is not appropriately positioned to participate in polymerization.
and properly constrain the activity of WbdAN presumably because WbdD is absent. These results presumably highlight the critical scaffold and orientation role played by WbdD in vitro and will only be fully appreciated if the structure of the WbdA-WbdD heterocomplex can be solved. Although in vitro O9a synthesis was restored by combining WbdAN and WbdA$^{E317A}$, the same combination did not work in vivo. The environment in vitro (high enzyme and substrate concentrations and a soluble acceptor) is radically different from the in vivo situation, making it impossible to directly relate the in vivo and in vitro data. However, the results do reinforce the need to consider both in vivo and in vitro situations when assigning functions to individual GT active sites in multidomain polymerases.

GTs have applications in emerging technologies, including glycoengineering in bacteria (50), and bacterial GTs that use sugar nucleotide donors are important in these efforts because of the vast range of unique specificities as well as their potential to build human glycan mimics. Polymerases are being investigated for various roles in enzymatic (and chemoenzymatic) syntheses (e.g. (4, 51–53) where the goal is to produce “glyco-products” with defined chemical and physical characteristics. Understanding the structure and function of multidomain polymerases will be essential for their successful application in glycoengineering efforts.

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