GlgE is a maltosyltransferase involved in α-glucan biosynthesis in bacteria that has been genetically validated as a target for tuberculosis therapies. Crystals of the Mycobacterium tuberculosis enzyme diffract at low resolution so most structural studies have been with the very similar Streptomyces coelicolor GlgE isoform 1. Although the donor binding site for α-maltose 1-phosphate had been previously structurally defined, the acceptor site had not. Using mutagenesis, kinetics, and protein crystallography of the S. coelicolor enzyme, we have now identified the +1 to +6 subsites of the acceptor/product, which overlap with the known cyclodextrin binding site. The sugar residues in the acceptor subsites +1 to +5 are oriented such that they disfavor the binding of malto-oligosaccharides that bear branches at their 6-positions, consistent with the known acceptor chain specificity of GlgE. A secondary binding site remote from the catalytic center was identified that is distinct from one reported for the M. tuberculosis enzyme. This new site is capable of binding a branched α-glucan and is most likely involved in guiding acceptors toward the donor site because its disruption kinetically compromises the ability of GlgE to extend polymeric substrates. However, disruption of this site, which is conserved in the Streptomyces venezuelae GlgE enzyme, did not affect the growth of S. venezuelae or the structure of the polymeric product. The acceptor subsites +1 to +4 in the S. coelicolor enzyme are well conserved in the M. tuberculosis enzyme so their identification could help inform the design of inhibitors with therapeutic potential.

GlgE is an α-maltose 1-phosphate:(1→4)-α-D-glucan 4-α-D-maltosyltransferase (starch synthase (maltosyl-transferring), EC 2.4.99.16) (1, 2) and a member of the carbohydrate-active enzymes database (CAZy) GH13_3 family (3). It is the third and defining enzyme of the four-step GlgE pathway that is responsible for the biosynthesis of a branched α-glucan (4). This pathway involves the sequential conversion of trehalose into α-maltose (5); α-maltose 1-phosphate (6), an α-1,4-linked α-glucan (1, 2); and finally an α-1,4-linked α-glucan with α-1,6-linked branches (7). The polymer comprises large dendrimers tens of nm in diameter (7). Each polymer molecule possesses a single C chain that bears at least one branch and the sole reducing end (Fig. 1). B chains are branches that themselves bear branches, whereas A chains do not such that A chains are largely at the periphery of the polymer. GlgE-derived α-glucan resembles glycogen except that it comprises shorter linear chains, has a correspondingly higher proportion of α-1,6 linkages, and has relatively fewer A chains. It is the combination of GlgE and the branching enzyme GlgB that defines the structure of the polymer. The key contribution that GlgE makes is to preferentially extend A chains, limiting the opportunity for GlgB to introduce a second branch into B and C chains. However, the structural basis for such specificity was not understood.

The GlgE pathway is widespread among bacteria (8) and has been best characterized in actinomycetes such as streptomycetes (9) and mycobacteria (10). Mycobacterium tuberculosis GlgE has attracted particular attention because of the demonstration that it is a genetically validated drug target (11). Unusually, the killing mechanism involves the hyperaccumulation of α-maltose 1-phosphate to toxic levels rather than the blocking of the production of α-glucan. Toxicity results in pleiotropic effects, and the target of this metabolite remains elusive. Driven by the desire to develop new therapies to tackle tuberculosis (11), some inhibitors of GlgE have been described. For example, substrate and transition state analogues have been reported with IC_{50}/K_i values of >200 μM (12, 13). Other potential inhibitors have been proposed based on theoretical docking calculations (14–16).

A number of crystal structures of GlgE have been solved, which will assist in the development of inhibitors. A high resolution structure of the M. tuberculosis enzyme has proven difficult to obtain (17), so most work has been done with Streptomyces coelicolor GlgE isoform 1 (18–20), which has very similar properties. Additional structures of the Mycobacterium thermoresistibile enzyme have also been reported recently (21), providing an alternative model for the M. tuberculosis enzyme. The enzyme consists of five domains (Fig. 2), four of which are typical of the α-amylase family (22). The catalytic domain A is elaborated by domain B, which forms a lid over the donor site, and inserts 1 and 2. Domain N forms the central core of the protein and much of the interface between the two subunits of...
Acceptor and Secondary Binding Sites of GlgE

the biological dimer. Domain C of the *M. tuberculosis* enzyme is capable of binding a malto-oligosaccharide close to the interface with domain A but remote from the active site (17). There are other examples of *α*-amylase enzymes with a secondary binding site on domain C (23), so this appears to be an evolutionarily conserved feature. Finally, domain S, which is unusual in this family, is able to adopt two conformations that seem to influence the ability of the domain B lid on the neighboring subunit to open and/or close (21). The GlgE dimer forms two distinct clefts on the same face of the structure as the active sites. In addition, there are several surface Ser and Thr residues in the B, S, and N domains of the *M. tuberculosis* enzyme that are subject to phosphorylation by the kinase PknB, leading to negative regulation (24). These residues are distinct from the known glucan binding sites. Therefore, regulation may be mediated through long range electrostatic effects on catalysis and/or on the ability of the donor site lid to open and close.

The donor site has been structurally characterized using wild-type and mutated forms of GlgE with either maltose (17, 19, 21), α-maltose 1-phosphate (18), or substrate/transition state analogues bound (17, 20). Not only do we have a clear picture of how the donor binds to the enzyme but also the nature of a covalently linked glycosyl-enzyme intermediate produced in the next step in the catalytic cycle (18). Indeed, it has been unequivocally demonstrated that Asp-394 (S. coelicolor numbering) is the nucleophile that attacks the donor to liberate inorganic phosphate by trapping a β-maltosyl-enzyme intermediate. A neighboring Glu-423 acid/base catalytic side chain is well placed to assist by protonating the phosphate leaving group and then deprotonating an incoming acceptor substrate. Loss of activity resulting from mutation of Glu-423 supports this role in catalysis (18). We therefore have a good understanding of the structures of the first two intermediates of what is a retaining double displacement mechanism consistent with other members of the GH13 family of enzymes. Conversely, the acceptor site has not been clearly defined in any studies to date. It is known that a hydrophobic ridge between two clefts is capable of binding cyclodextrins and that the *S. coelicolor* enzyme is inhibited by cyclodextrins by competing with acceptors (19). Therefore, the acceptor site probably overlaps with the cyclodextrin cleft. However, this site is over 10 Å away from the donor site and oriented orthogonally with respect to the donor, making modeling of the entire acceptor site challenging.

Important issues that remain unresolved therefore include how an acceptor binds such that it prefers A chains (Fig. 1) and what role the secondary binding site has. To address these questions, we have used fresh approaches with mutated forms of *S. coelicolor* GlgE isoform 1 to assess the role of specific amino acid side chains in enzyme activity and to obtain new structures with oligosaccharides bound.

Results

Mapping the Acceptor Binding Site—There are two conspicuous clefts that are adjacent to the donor binding site (Fig. 2). The one closest to the donor site is the linear cleft that is formed primarily by domains S and N of the neighboring subunit. To establish whether linear acceptor substrates of GlgE can bind along this linear cleft, we mutated three solvent-exposed amino acid side chains (Ala-58, Gly-60, and Ala-186) along its length to Arg. The substitution with such a large side chain would be expected to compromise the binding of acceptors if the linear cleft were to overlap with the acceptor binding site. By monitoring the production of inorganic phosphate with malachite green, the kinetics of each of the three variants was tested using maltose as the acceptor. In each case, little difference in the kinetics was observed compared with the wild-type enzyme (Table 1). It would therefore seem that short linear acceptors do not bind in the linear cleft. Similarly, when these GlgE variants were assayed with α-glucan polymer isolated from *Streptomyces venezuelae* rather than maltohexaose, their values of *k*<sub>cat</sub>/ *k*<sub>m</sub> were less than 2-fold different from that of the wild-type enzyme (data not shown), suggesting that the linear cleft is not particularly important in the accommodation of the larger acceptor either.
We have previously shown that cyclodextrins bind to a largely hydrophobic ridge between the linear and diagonal clefs over 10 Å from the donor site (Fig. 2) (19). Because cyclodextrins inhibit S. coelicolor GlgE competitively with regard to acceptor substrates, the acceptor binding site might be expected to overlap with this site. Significant interactions between cyclodextrins and GlgE involve the side chains of Trp-448 and Arg-449, so we explored the effect of mutations at these positions. When Trp-448 was substituted by either Ala or Glu, the enzyme lost almost all of its detectable activity (Table 1), consistent with the acceptor binding at this position through van der Waals interactions. When Arg-449 was mutated to Ala, the \( k_{cat}^{app} \) for maltohexaose increased 14-fold with no change in the value of \( k_{cat}^{app} \), showing that binding was somewhat compromised by the loss of some hydrogen bonding interactions. These observations provide additional evidence that the acceptor binding site overlaps with the cyclodextrin binding site.

The considerable distance between the cyclodextrin and donor binding sites necessitates several sugar residues to bridge the gap (Fig. 2). To obtain some indication which path the acceptor binding site takes, we mutated two amino acid side chains that flank either side of the direct path between the two sites. We first substituted His-397 to Arg and found that \( k_{cat}^{app} \) was reduced 43-fold (Table 1), strongly suggesting that the orientation of the acceptor is compromised in its ability to attack the glycosyl-enzyme intermediate of this form of the enzyme. The associated \( k_{cat}^{app}/k_{cat}^{on} \) was consequently reduced 33-fold. By contrast, mutation of Glu-527 to Ala had relatively little effect, suggesting that the Glu side chain at this position does not have a significant role in either binding or orienting the acceptor. However, introduction of an Arg at this position did lower the \( k_{cat}^{app}/k_{cat}^{on} \) for the acceptor by about a third, suggesting that the bulkier side chain had a greater reach, leading to a slight distortion of the orientation of the acceptor. Nevertheless, the effect of this substitution was much smaller than the equivalent mutation of His-397.

The acceptor chain length specificity of each of the mutants was evaluated. In each case, the optimum was between five and seven glucose residues in length. This compares with six for the wild-type protein. The variants therefore showed only minor changes in chain length specificity with no obvious trends emerging. Nevertheless, it is clear from the kinetic constants with maltolhexaose that the acceptor binding site overlaps with the cyclodextrin binding site and passes by His-397.

**The Cyclodextrin Site Can Bind a Linear Malto-oligosaccharide**—Attempts to identify the acceptor binding site using protein crystallography have previously been thwarted by the ability of GlgE to disproportionate and ultimately hydrolyze malto-oligosaccharides to form maltose. To prevent GlgE from catalyzing such reactions, a D394A GlgE variant that lacks the nucleophilic side chain (18) was generated. This GlgE variant was co-crystallized (Table 2) with a preparation of maltodextrin, which contains a wide range of both linear and \( \alpha \)-1,6 branched oligosaccharides according to mass spectrometry and NMR spectroscopy. The donor binding site was occupied by maltose as has been observed in many of the published structures. However, a linear malto-oligosaccharide was observed bound to the cyclodextrin binding site for the first time (Fig. 3). Five sugar residues were resolved, four of which occupied very similar positions to those observed with \( \beta \)-cyclodextrin such that they interacted with Trp-448 and Arg-449. The remaining glucose residue was at the non-reducing end and was slightly displaced such that it was more oriented toward the donor binding site. However, this glucose residue was still over 10 Å away from the donor site. Therefore, the complete acceptor site was not defined in this structure.

**A Product-bound Structure**—Co-crystallization and soaks of a different catalytically compromised E423A variant of GlgE, which lacks the acid/base catalytic side chain (18), were carried out with various substrates and/or substrate analogues. In an attempt to obtain a structure of a trapped maltosyl-enzyme intermediate (18) with an acceptor bound in a productive conformation, we soaked a crystal of the E423A variant of GlgE in a 2-methyl-2-deoxy-2-fluoro-\( \alpha \)-maltosyl fluoride. Fortuitously, this yielded a structure where malto-olactose was bound such that it occupied both the donor and cyclodextrin binding sites (Fig. 4), whereas the donor analogue was unexpectedly absent from the crystal. This structure would therefore appear to represent a product-bound species with subsites −2 to +6 occupied. The two sugar residues in the donor site are oriented in a manner similar to those observed with maltose and \( \alpha \)-maltose 1-phosphate (18, 19). The residues occupying subsites +3 to +6 overlap with those seen in cyclodextrins and the maltopentaose described above but are somewhat displaced to allow the product to bind in the appropriate orientation. The maltopentaose interacts with Trp-448 (stacking to the +3 sugar), Arg-449 (3.1-Å H-bonds to O2 and O3 of the +4 sugar), and His-397 (3.2-Å H-bond to O2 of the +1 sugar) but not Glu-527 (>4.7-Å O-O distance) as expected from the kinetics of the variants described above. The conformation of the oligosaccharide was unusually flat and extended (Table 3) compared with amylose or amylopectin (25), which have \( \alpha \)-1,4 torsion angles closer to that of maltose (\( \varphi = 116^\circ \) and \( \psi = -118^\circ \)) (26). This was particularly striking at the linkage where the donor and acceptor glucose residues occupying subsites −1 and +1 connect (\( \varphi = 34^\circ \) and \( \psi = -157^\circ \)), giving the longest distance between the 2- and 3'-hydroxyl oxygens of 4.9 Å (Table 3). This likely reflects the geometry required to allow glycosyl transfer to occur. An essentially identical structure was also obtained by

**TABLE 1**

| GlgE    | \( k_{cat}^{app} \) | \( k_{cat}^{app} \) | \( k_{app}^{cat}/k_{app}^{on} \) |
|---------|-------------------|-------------------|----------------|---|
| Wild type | 9.4 ± 0.5 | 0.49 ± 0.07 | 19000 ± 3000 |
| Linear cleft |
| A58R    | 7.4 ± 0.5 | 0.50 ± 0.09 | 14800 ± 2800 |
| G60R    | 7.2 ± 0.4 | 0.49 ± 0.07 | 15000 ± 2200 |
| A186R   | 9.2 ± 0.5 | 0.60 ± 0.08 | 15300 ± 2800 |
| Acceptor site |
| W448A   | ND* | ND | 94 ± 10 |
| W484E   | ND | ND | 75 ± 10 |
| R449A   | 9.6 ± 0.8 | 6.7 ± 1.0 | 9600 ± 1600 |
| H397R   | 0.220 ± 0.002 | 0.36 ± 0.01 | 580 ± 14 |
| E527A   | 7.2 ± 0.5 | 0.32 ± 0.07 | 22500 ± 2100 |
| E527R   | 6.5 ± 0.3 | 0.52 ± 0.07 | 12500 ± 1800 |
| Secondary site |
| Y646A   | 9.3 ± 0.6 | 0.66 ± 0.11 | 14000 ± 2500 |

* ND, not derived due to lack of enzyme activity.
soaking a crystal with maltoheptaose except that the +6 subsite was not occupied. This showed that it was possible to obtain a product-bound structure in the absence of a donor substrate analogue.

A Newly Identified Secondary Binding Site in S. coelicolor GlgE—The product-bound structures of the E423A variant of GlgE had an additional malto-oligosaccharide bound at a secondary site located on domain C (Fig. 5). Unexpectedly, this is a previously undescribed secondary binding site over 40 Å away from the donor site and over 10 Å away from that recently described for the M. tuberculosis enzyme (17). In each case, five glucose residues were well resolved, clearly showing that they that were wrapped around Tyr-646 of S. coelicolor GlgE. The non-reducing end of this malto-oligosaccharide occupied a pocket on the surface of the enzyme, precluding the ability to accommodate any more glucose residues at this location. By contrast, the reducing end glucose residue was flipped relative to the other residues such that its α-anomeric hydroxyl projected into solvent (Fig. 5). The potential for the malto-oligosaccharide to extend into solvent was supported by the detection of orthogonal electron density suggesting the detection of at least one other glucose residue. However, it was not possible to resolve additional residues unambiguously. The likelihood is that malto-octaose/maltoheptaose was bound to this site but that three/two glucose residues at the reducing end were not well ordered in the crystal. If so, this implies that the secondary binding site is capable of accommodating linear malto-oligosaccharides with a length of greater than five residues.

Similarly, another oligosaccharide was observed in the structure of the D394A GlgE variant obtained with maltodextrin. Again, five glucose residues of an α-1,4-linked malto-oligosaccharide were wrapped around the side chain of Tyr-646 with the reducing end glucose residue being flipped relative to the others (Fig. 6). Remarkably, several additional glucose residues were resolvable within the electron density. In one copy of the GlgE subunit within the crystal, an additional four sugar residues were present that made no direct contact with this subunit but instead interacted with a subunit from a neighboring GlgE homodimer in the crystal lattice. There was an α-1,6 linkage between the reducing end of the proximal pentasaccharide and the second residue from the reducing end of the distal tetrasaccharide. Therefore, the protein-bound pentasaccharide constitutes an A chain emanating from a tetrasaccharide C chain. In the other copy of the GlgE subunit, only the first additional α-1,6-linked distal sugar residue was resolvable.

The Secondary Binding Site Kinetically Assists the Extension of Polymers but Is Not Essential—To establish the role of the secondary binding site of S. coelicolor GlgE, we generated a Y646A variant that would disrupt the binding of oligosaccharides to this site. The kinetic constants of this variant were very similar to those of the wild-type enzyme when using maltohexaose as the acceptor substrate (Table 1). Perhaps this is to be expected because the secondary binding site is quite remote from the donor and acceptor sites, and initial rates are measured in the enzyme assay when oligosaccharides have not yet reached significant lengths.
A linear malto-oligosaccharide can bind to the cyclodextrin binding site. Co-crystallization of a D394A variant of GlgE with maltodextrin yielded a structure with maltose bound to the donor site and maltopentaose bound to the cyclodextrin binding site as indicated by the omit electron density map (1.95-Å resolution; contoured at −3.0 φ). The residues of the maltopentaose superpose well with those of cyclodextrins in complex with GlgE (Protein Data Bank accession codes 3ZST, 3ZT6, and 3ZT7) (19). There is less correspondence with the acceptor subsites seen in the product-bound structures described here (Fig. 4); for this reason, the subsite labels are shown in parentheses (see also Fig. 9). Interactions between the maltopentaose and GlgE involved (from left to right) Trp-448, Arg-449, Tyr-445, Pro-428, Arg-427, and Thr-426 (backbone CO) from one subunit and Gly-84 and Pro-83 from the neighboring subunit. The two subunits are distinguished by coloring, oxygen atoms are shown in red, carbon atoms are shown in green for the malto-oligosaccharide, and the amino acids mutated in this study are shown in magenta. The orientation of GlgE is identical to that of Fig. 2.

FIGURE 3. A linear malto-oligosaccharide can bind to the cyclodextrin binding site. Co-crystallization of a D394A variant of GlgE with maltodextrin resulted in only the former being detected in the structure. Malto-octaose was bound to both the donor and cyclodextrin binding sites. Each subsite resulted in only the former being detected in the structure. Malto-octaose superpose well with those of cyclodextrins in complex with GlgE (Protein Data Bank accession codes 3ZST, 3ZT6, and 3ZT7) (19). There is less correspondence with the acceptor subsites seen in the product-bound structures described here (Fig. 4); for this reason, the subsite labels are shown in parentheses (see also Fig. 9). Interactions between the maltopentaose and GlgE involved (from left to right) Trp-448, Arg-449, Tyr-445, Pro-428, Arg-427, and Thr-426 (backbone CO) from one subunit and Gly-84 and Pro-83 from the neighboring subunit. The two subunits are distinguished by coloring, oxygen atoms are shown in red, carbon atoms are shown in green for the malto-oligosaccharide, and the amino acids mutated in this study are shown in magenta. The orientation of GlgE is identical to that of Fig. 2.

FIGURE 4. A product-bound structure. Soaking a crystal of the E423A variant of GlgE with malt-octaose and 2-deoxy-2-fluoro-α-maltosyl fluoride resulted in only the former being detected in the structure. Malto-octaose was bound to both the donor and cyclodextrin binding sites. Each subsite from −2 to +6 is highlighted along with the amino acids mutated in this study. A, malt-octaose was clearly resolved as indicated by the omit electron density map (2.5-Å resolution; contoured at −3.0 φ). It was bound to the subsites −2 and −1 of the donor site (18, 19) and +1 to +6 of what would be the acceptor site. The locations of the catalytic nucleophile (Asp-394) and acid/base amino acid (Glu-423 mutated to Ala in this case) are indicated. B, when the protein is depicted with a molecular surface, it is apparent that the C6 primary hydroxyl groups of the glucose residues project toward the protein surface in subsites +1 to +5. By contrast, it is feasible that an α-1,6-linked malto-oligosaccharide could emanate from the +6 subsite and be accommodated by the linear cleft as indicated by the yellow arrow. In each panel, the two subsites are distinguished by gold and slate coloring, oxygen atoms are shown in red, and carbon atoms are shown in green for the malto-oligosaccharide and in magenta for amino acids mutated in this study. The orientation of GlgE is identical to that of Fig. 2.

To test whether the secondary binding site becomes important when GlgE extends a polymer, the assays were repeated using α-glucan. In this case, it was clear that disruption of the secondary binding site had an impact on activity. The value kcat/Km increased from 3.5 ± 0.2 to 0.85 ± 0.07 s−1, whereas that of kapp went up from 0.33 ± 0.05 to 2.5 ± 0.4 mg ml−1. Thus, the value of kcat/km decreased 31-fold.

The impact of the secondary binding site on the linear chain lengths of a branched polymer was then assessed. The S. coelicolor GlgE Y646A variant was mixed with the S. coelicolor branching enzyme GlgB isoform 1 and exposed to amylose allowed the constituent linear chain length to be determined using mass spectrometry. It was apparent that the GlgE variant, in combination with GlgB, generated linear chain lengths (mean and chain length distributions of 8.3 ± 2.8) indistinguishable from those of the wild-type enzyme (7.9 ± 2.8) in back-to-back experiments. Therefore, the secondary binding
site does not seem to have a role in defining the structure of the polymer in vitro.

*S. coelicolor* possesses two copies of the glgE gene, making the genetic manipulation of the GlgE pathway in this organism less tractable than in the closely related *S. venezuelae*, which has only one copy (8, 9). The GlgE homologues from *S. coelicolor* (isoform 1) and *S. venezuelae* share 79% amino acid sequence identity together with very high conservation around the secondary binding site including the key Tyr residue (Fig. 8). It would therefore be expected that the *S. venezuelae* enzyme also possesses the secondary binding site and that the Tyr to Ala mutation would disrupt the function of this site. To determine whether the secondary binding site is required for the function of GlgE in vivo, a glgE-null mutant of *S. venezuelae* (9) was complemented with an *S. venezuelae* glgE gene coding for the equivalent Tyr to Ala GlgE variant (Y665A). Without complementation with an intact glgE gene, the mutant strain is known not to produce α-glucan and to exhibit a delayed growth phenotype due to the accumulation of α-maltose 1-phosphate. However, the strain complemented with the GlgE variant grew normally and produced α-glucan in a manner similar to the wild-type strain (Fig. 7). This suggests that the secondary binding site is not essential in vivo in the conditions tested despite a potential loss of activity.

α-Glucan isolated from the complemented strain resembled that isolated from the wild type according to electron microscopy (Fig. 7). The mean chain length of the polymer from the complemented mutant strain (7.9 ± 2.0) was similar to that isolated from the wild type (7). In addition, the degree of digestion by β-amylose (19%) was also comparable with that of the wild-type polymer (21.0 ± 4.7%). Therefore, the secondary binding site does not seem to be important in defining the gross or fine structure of the polymer.

**Discussion**

**Acceptor Binding Site**—The key aim of this study was to identify the acceptor binding site. By using mutagenesis to prevent GlgE from slowly hydrolyzing malto-oligosaccharides, it was possible to obtain a structure with malto-octaoase bound to the active site (Figs. 4, 5, and 9). The occupancy of the donor subsites –2 and –1 together with subsites +1 to +6 indicated that this represents a product-bound structure that reveals the locations of the first six acceptor subsites. Although the accep- tor and cycloextrin binding sites overlapped, the registration of the glucose residues differed. This helps explain the unexpectedly orthogonal orientation of cycloextrins (Fig. 2) and unproductively bound malto-pentaose (Fig. 3) relative to the donor binding site.

We have previously shown that GlgE preferentially extends malto-oligosaccharides with a length of at least four and an optimum of six residues (19). This fits well with the observation that the +3 to +6 acceptor subsites have a particularly high propensity to bind oligosaccharides including α-cycloextrin, β-cycloextrin (19), and unproductively bound malto-pentaose...
ultimately has a diameter of tens of nm. However, GlgE variants with a disrupted linear cleft did not seem to be kinetically compromised when acting on α-glucan polymer, leaving any potential role for the linear cleft unclear.

Consistent with these observations, we have previously shown that GlgE preferentially extends A chains, resulting in most B chains remaining short and bearing only one branch (7). The acceptor binding site of GlgE contrasts with a much more open one that accommodates only two glucose residues in a classical glycogen synthase (27). This allows B and C chains in classical glycogen to be more readily extended and made available for additional branching by GlgB (28).

The only GlgE variant to have a significantly different $k_{cat}$ was the one with the amino acid substitution adjacent to the +1 subsite at His-397 (Table 1). This is consistent with the orientation of the attacking nucleophilic sugar at this subsite needing to be optimal. The unusual conformation of the linkage between the −1 and +1 glucose residues is likely to reflect the appropriate orientation. Interestingly, the +1 pyranose ring is orthogonal to that of the −1 ring such that their respective hydroxyl groups are maximally separated. Perhaps the avoidance of hydrogen bonding interactions between these glucose residues is important in achieving the optimal orientation.

Having already obtained the structure of GlgE with a trapped β-maltosyl intermediate (18), we attempted to obtain such a structure with an acceptor also bound. Despite repeated attempts, we were unable to obtain a ternary Michaelis complex with both a maltosyl-enzyme intermediate and the +1 subsite occupied. It therefore remains to be seen whether such a structure would reveal a $^{3}$H$_{2}$-type conformation of the −1 sugar ring predicted to promote nucleophilic displacement of the catalytic Asp-394 residue. Distortion could potentially be mediated through the H-bonding interaction between the O6 hydroxyl of the −1 sugar ring with the hydroxyl group of Tyr-357 of the B domain, which forms a lid over the donor binding site. Structural changes like this could provide a mechanism through which GlgE could gate maltosyl transfer to an oligosaccharide rather than to water, analogous to gating in other glucosyltransferases (29).

Finally, product release would presumably require the domain B lid to open. Indeed, the elevated $B$ factors of this lid in the product-bound structure are consistent with this likelihood. Tantalizingly, a structure of $M$. thermoresistible GlgE with maltose bound shows a different conformational state of the $S$ domain together with a loss of electron density of the adjacent $B$ domain of the neighboring subunit, suggesting disorder in the lid and the existence of an open conformation (21).

**Secondary Binding Site**—As discussed above, the linear and diagonal clefts could accommodate elements of the polymer to allow non-reducing ends to gain access to the acceptor site. Their apparent inability to specifically recognize and bind malto-oligosaccharides may simply reflect the need for these spaces to be available for a wide variety of different α-glucan substructures. By contrast, we have identified a new secondary binding site on domain C. This site appears capable of binding either linear or branched oligosaccharides (Figs. 5, 6, and 9). Incidentally, there are only a few other protein structures with branched oligosaccharides bound thus far reported (30, 31).
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A likely role for this secondary site is to guide the polymer toward the active site (32). This possibility is supported by the activity of GlgE with a polymeric acceptor being compromised when the secondary site is disrupted. Surprisingly, whether linear or branched, the reducing end of an oligosaccharide projects between the donor sites of the two subunits of GlgE rather than the expected non-reducing end of an acceptor, so how an acceptor polymer is guided to the donor site remains unclear. There has been no indication that the enzyme is processive or subject to allosteric regulation by carbohydrates, so the secondary binding site is less likely to be involved in such phenomena. Another possible function of the secondary site is to co-locate the enzyme and the polymer within the cell. However, this site is not essential in Streptomyces because the bacterium was still able to grow and to generate α-glucan with the same fine structure when the site was disrupted.

A somewhat similar but distinct secondary binding site has also been observed on domain C of M. tuberculosis GlgE (17). In both cases, five glucose residues wrap around an aromatic amino acid side chain (S. coelicolor Tyr-646 versus M. tuberculosis Phe-631). Surprisingly, neither of the two secondary sites is spatially conserved, being over 10 Å apart in the superposed structures. This implies evolutionary pressure for this function to exist even if there is no structural conservation. Although the oligosaccharide appears to be bound to the M. tuberculosis enzyme in the reverse orientation, the resolution of this structure is only 3.9 Å. Therefore, it seems feasible that it is in fact in the same orientation as that seen in the S. coelicolor enzyme, a possibility acknowledged by the authors of the other study (17).

Significance for Therapies against Tuberculosis—GlgE is an interesting enzyme not least because it has been genetically validated as a potential target in therapies for tuberculosis (1). GlgE from S. coelicolor shares many properties with the enzyme from M. tuberculosis and other mycobacteria (1, 17–19, 21). Although a structure of the M. tuberculosis enzyme has been reported (17), the S. coelicolor enzyme crystallizes more readily to give higher resolution structures. To mimic the M. tuberculosis enzyme better, a V279S amino acid substitution has been made in the S. coelicolor donor site (20). Having defined the acceptor site in the S. coelicolor enzyme, it is now possible to explore whether it is conserved in the M. tuberculosis enzyme. Each of the key amino acid residues of subsites +1 to +4 within domain A are conserved (e.g. His-397, Trp-448, Arg-449, Tyr-445, and Pro-428; S. coelicolor numbering), whereas those of subsites +5 and +6 in domain N of the neighboring subunit are quite different (Gly-84 and Pro-83 are spatially substituted by Gln and Glu, respectively). It is likely that these differences lie behind the lack of inhibition of the M. tuberculosis enzyme by cyclodextrins (19). Although the acceptor site is much more open than that of the donor site, it is nevertheless feasible to develop inhibitors that bind to the acceptor subsites, and our new findings allow this to be done in a more rational way.

Experimental Procedures

Protein Production and Enzyme Assays—Recombinant Histagged S. coelicolor GlgE isoform 1 was produced in Escherichia coli and purified using nickel affinity and size exclusion chromatographies as described previously (1, 18, 19). Variants were generated using a QuikChange Lightening site-directed mutagenesis kit (Agilent Technologies) as described previously (18). GlgE activity was determined in triplicate by monitoring the production of inorganic phosphate using malachite green as described previously (1, 18, 19). Maltohexaose concentrations up to 2 mM and α-glucan up to 2.5 mg ml⁻¹ were used, ranges in which data conformed to the Michaelis-Menten equation. The α-glucan was isolated from S. venezuelae as described previously (7). The values of $k_{cat}$ and $K_m$ are apparent because the enzyme conforms to a ping-pong (also known as substituted enzyme) kinetic mechanism (1). Errors reflect S.E. of the fit. Synthetic α-glucan was generated using GlgE and GlgB with α-maltose 1-phosphate and maltohexaose as described previously (7). Linear lengths of constituent chains were determined by treating polymers with isomylase and analyzing the products using mass spectrometry as described previously (7).

Crystallography—Crystals of S. coelicolor GlgE isoform 1 were obtained by vapor diffusion from 15% (w/v) polyethylene glycol 3350, 0.2 M sodium citrate, and 15% (v/v) ethylene glycol as described previously (18, 19). For the E423A variant, complexes were obtained by soaking the crystals in crystallization solution containing 5 mM ligand for 5 min prior to mounting. In the case of the malto-octaose product-bound structure, both malto-octaose and 2-deoxy-2-fluoro-α-maltosyl fluoride were present in the soaking solution, but only the former was visible in the resultant complex. For the D394A variant, the ligand-bound structure was obtained by co-crystallization with 5 mM maltodextrin. Crystals were flash cooled in LithoLoops (Molecular Dimensions) by plunging into liquid nitrogen and stored in Unipuck cassettes (MiTeGen) prior to transport to the synchrotron. The crystals were subsequently transferred robotically to the goniostat on station I04-1 or I24 at the Diamond Light Source (Oxfordshire, UK) and maintained at 100 K with a Cryojet cryocooler (Oxford Instruments). Diffraction data were recorded using a either a Pilatus 4 M detector (Dectris) and processed using the xia2 expert system (33). X-ray data collection statistics are summarized in Table 2. The data sets for the E423A variant complexes were isomorphous with that previously collected for the same variant complexed with maltose (Protein Data Bank accession code 4CN6) (18). Therefore, the latter was used as a starting point for modeling these structures, which were completed through several iterations of refinement in REFMAC5 (34) and manual adjustment in Coot (35). In the latter stages, translation libration screw (TLS) refinement was used with a total of eight TLS domains, which were defined using the TLS motion determination server (36). By contrast, the data set derived from co-crystallization of the D394A vari-

2 The abbreviation used is: TLS, translation libration screw.
ant with maltodextrin was not isomorphous with the above data sets (or any other data set previously obtained for this protein) despite belonging to the same space group and having similar cell parameters. This structure was solved by molecular replacement with Phaser (37) using one subunit taken from the wild-type structure of GlgE isoform 1 co-crystallized with α-cyclodextrin (Protein Data Bank accession code 3ZST) (19) as the starting template. This gave two molecules per asymmetric unit forming the familiar homodimer but having distinctly different crystal contacts presumably dictated by the binding of branched malto-oligosaccharides to the secondary surface binding site. This starting model was refined as for the two complexes described above. Refinement statistics are summarized in Table 2. Simulated annealing omit electron density maps were calculated for the various ligands as follows. The ligands were deleted from the coordinates of the final models, and these were used as inputs to simulated annealing refinement with PHENIX (38) from a starting temperature of 5000 K after applying small random shifts to the models (“shake” term set to 0.3). The resultant $\Delta F_{\text{obs}} - \Delta F_{\text{calc}}$ difference electron density maps are shown in Figs. 3, 4A, and 6. All structural figures were prepared using CCP4mg (39).

Microbiology—The glgE-null mutant of S. venezuelae ATCC10712 was generated using the Redirect PCR targeting method (40) where the coding region of glgE was replaced with an apramycin resistance cassette as described previously (9). The mutant was complemented with glgE cloned into the plasmid pMS82 (41) as described previously (9) except that glgE was first mutated using the QuickChange kit to allow substitution of Tyr-665 with Ala (S. venezuelae numbering). Strains were grown for 2 days on maltose/yeast extract/malt extract/John Innes Centre borehole water 2 solid medium (42) at 28 °C as described previously (9). Colonies were sectioned, stained for α-glucan using periodic acid-thiocarbazohydrazide-silver proteinate, and imaged using transmission electron microscopy as described previously (9). α-Glucan was isolated from the complemented strain as described previously and analyzed as described above (7).

Author Contributions—S. B. coordinated the study and wrote the paper. K. S., C. E. M. S., and D. M. L. edited the manuscript. K. S., M. T., A. G., and A. M. R. carried out the kinetic experiments summarized in Table 1. K. S., C. E. M. S., and D. M. L. carried out the crystallography. J. E. B. carried out the electron microscopy. K. S. and F. M. carried out the microbiology. All authors approved the final version of the manuscript.

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