# **A Catalytically Essential Motif in External Loop 5 of the Bacterial OligosaccharyItransferase PglB**\*<sup>3</sup>

Received for publication, October 4, 2013, and in revised form, November 5, 2013 Published, JBC Papers in Press, November 25, 2013, DOI 10.1074/jbc.M113.524751

 $\Delta$ Christian Lizak $^{+1}$ , Sabina Gerber $^{+1,2}$ , Daria Zinne $^{\pm}$ , Gaëlle Michaud $^{\mathbb{S}}$ , Mario Schubert $^{+}$ , Fan Chen $^{\P}$ , Monika Bucher $^{+}$ , **Tamis Darbre**§ **, Renato Zenobi**¶ **, Jean-Louis Reymond**§ **, and Kaspar P. Locher**‡3

*From the* ‡ *Institute of Molecular Biology and Biophysics, Department of Biology, ETH Zurich, Schafmattstrasse 20, 8093 Zurich, Switzerland, the* § *Department of Chemistry and Biochemistry, University of Berne, 3012 Berne, Switzerland, and the* ¶ *Department of Chemistry and Applied Biosciences, ETH Zurich, 8093 Zurich, Switzerland*

**Background:** *N*-Linked glycosylation is catalyzed by oligosaccharyltransferase (OST). **Results:** A so far unrecognized sequence motif in the external loop 5 (EL5) of bacterial OST is essential for catalysis. **Conclusion:** EL5 is involved in acceptor substrate binding and in critical interactions with the lipid-linked oligosaccharide. **Significance:** The study defines the dual role of EL5 during catalysis in protein *N*-glycosylation.

**Asparagine-linked glycosylation is a post-translational protein modification that is conserved in all domains of life. The initial transfer of a lipid-linked oligosaccharide (LLO) onto acceptor asparagines is catalyzed by the integral membrane protein oligosaccharyltransferase (OST). The previously reported structure of a single-subunit OST enzyme, the** *Campylobacter lari* **protein PglB, revealed a partially disordered external loop (EL5), whose role in catalysis was unclear. We identified a new and functionally important sequence motif in EL5 containing a** conserved tyrosine residue (Tyr<sup>293</sup>) whose aromatic side chain is **essential for catalysis. A synthetic peptide containing the conserved motif can partially but specifically rescue** *in vitro* **activity of mutated PglB lacking Tyr293. Using site-directed disulfide cross-linking, we show that disengagement of the structurally ordered part of EL5 is an essential step of the glycosylation reaction, probably by allowing sequon binding or glyco-product release. Our findings define two distinct mechanistic roles of EL5 in OST-catalyzed glycosylation. These functions, exerted by the two halves of EL5, are independent, because the loop can be cleaved by specific proteolysis with only slight reduction in activity.**

Glycosyltransferases utilize activated donor sugar substrates for the transfer to various targets, thereby forming glycosidic bonds. The transfer of glycans to the side chain of asparagines, a process termed *N*-linked protein glycosylation, is of major importance for protein folding, cell viability, and organism development (1–3). Protein *N*-glycosylation occurs in all three domains of life  $(4-6)$ , and although there exist a number of variations, the general mechanism is conserved; a lipid-linked



oligosaccharide  $(LLO)^4$  is assembled in a multistep process before being translocated from the cytosolic to the luminal side of the endoplasmic reticulum membrane of eukaryotes or to the periplasmic/external side of the plasma membrane of prokaryotes. The integral membrane enzyme oligosaccharyltransferase (OST) then catalyzes the *en bloc* transfer of the oligosaccharide to asparagine residues located in the consensus sequon  $\text{Asn-Xaa-}\ \text{Ser}/\text{Thr}$  (with  $\text{Xaa} \neq \text{Pro}$ ) of acceptor polypeptides. The OST enzyme in higher eukaryotes is a multiprotein complex with Stt3 as the catalytic subunit (7, 8), but in kinetoplastids and prokaryotes, OST is a single-subunit enzyme that is homologous to Stt3 (9–11). The crystal structure of a fulllength, bacterial Stt3 homolog, the PglB protein of *Campylobacter lari*, revealed the architecture of this class of enzymes (12). PglB contains an N-terminal transmembrane (TM) domain featuring 13 TM segments and a soluble domain facing the periplasm. The TM domain contains two large external loops (EL1 and EL5) that provide non-covalent interactions between the TM and the periplasmic domains. Whereas the role of EL1 appears to be mainly a structural one, EL5 was proposed to be involved in catalytic steps of the process. Recent functional studies have provided a quantitative basis for sequon recognition and binding by PglB; the  $+2$  sequon Ser/Thr of acceptor substrates is recognized by a binding pocket provided by the WWD motif (a diagnostic motif among Stt3 homologs) and a neighboring Ile residue (Ile<sup>572</sup>), suggesting that the  $+2$ Ser/Thr defines the specificity of *N*-linked glycosylation sites (13). The *C. lari* PglB enzyme was also useful for studying the mechanism of activation of the acceptor Asn side chain of sequon (14).

Arguably the least understood feature of the PglB structure (and OST mechanism) is the role of the external loop EL5. Both for substrate binding and Asn activation, the C-terminal half of this loop appears to play an important role by pinning the this work was supported by a grant from NCCR Structural Biology Zurich (to the subset of pray and important fole by printing the \* This work was supported by a grant from NCCR Structural Biology Zurich (to the subsetion ag

K. P. L.) and Swiss National Science Foundation Grants SNF 200020-125020 (to J. L. R.), SNF 200020-124663 (to R. Z.), and SNF 31003A-131075/1 and SNF 31003A-146191/1 (to K. P. L.).

<sup>□</sup>**<sup>S</sup>** This article contains supplemental Tables S1 and S2 and Figs. S1–S6. <sup>1</sup> Both authors contributed equally to this work.

<sup>&</sup>lt;sup>2</sup> Present address: GlycoVaxyn AG, 8952 Schlieren, Switzerland.

<sup>&</sup>lt;sup>3</sup> To whom correspondence should be addressed. Tel.: 41-44-633-3991; Fax: 41-44-633-1182; E-mail: locher@mol.biol.ethz.ch.

<sup>&</sup>lt;sup>4</sup> The abbreviations used are: LLO, lipid-linked oligosaccharide; EL1 and EL5, external loop 1 and 5, respectively; Ni-NTA, nickel-nitrilotriacetic acid; OST, oligosaccharyltransferase; TM, transmembrane; HSQC, heteronuclear single-quantum correlation; HMQC, heteronuclear multiplequantum correlation.

essential residue Glu<sup>319</sup> to the catalytic site and positioning the acceptor Asn correctly for glycosylation (12, 13). The function of the N-terminal half of EL5, which was disordered in the structure of peptide-bound PglB (supplemental Fig. S1, *top state*), remained unclear.

To explore the function of the N-terminal half of EL5 and to investigate conformational rearrangements in the C-terminal half during catalysis, we performed a detailed structure-function analysis of *C. lari* PglB. We used a synthetic, fluorescently labeled peptide that contained a bacterial glycosylation sequon and a previously established *in vitro* assay that not only provided high precision in determining glycosylation turnover but also allowed us to observe very low reaction rates of disfavored PglB mutants and determine changes in sequon binding affinities (13). We thus identified a previously unrecognized, catalytically essential motif, termed Tyr-plug, in the N-terminal half of EL5 that contains a conserved tyrosine residue (Tyr<sup>293</sup>). The motif is essential for catalysis but not for peptide binding.

We had speculated earlier that once the *N*-glycosidic linkage is formed, the glyco-polypeptide will need to dissociate from the enzyme, which will probably require a disengagement of EL5 during the catalytic cycle (supplemental Fig. S1). To investigate this hypothesis, we engineered disulfide cross-links in PglB by introducing cysteines in EL5 and the enzyme core at three distinct locations. The resulting PglB mutants were expressed and purified in large quantities, and sequon binding and glycosylation turnover rates were determined under oxidizing and reducing conditions. The results allowed us to define the dynamics of EL5 during catalysis.

### **EXPERIMENTAL PROCEDURES**

*Construction of Plasmids*—Point mutations in the *pglB* gene of *C. lari* were generated by the QuikChange<sup>TM</sup> method either on a pBAD plasmid (12) for *in vitro* glycosylation studies or on a pMLBAD plasmid (15) for *in vivo* glycosylation assays. For *in vitro* glycosylation, the two endogenous glycosylation sites were removed (N535Q and N556Q) to prevent autoglycosylation of PglB, and all subsequent mutations were based on this construct. We refer to wild type PglB in our *in vivo* and *in vitro* experiments irrespective of the presence of endogenous glycosylation sites because the catalytic site and the peptide binding site are not affected by these mutations. Plasmid DC3 was obtained by digesting plasmid pACYC<sub>pgl<sub>mut</sub> (11) with enzyme</sub> BbvCI and ligation of the insert containing the *pglB* gene into the backbone of plasmid pACYC*pgl*::*pglJ* (16) digested with the same enzyme. Plasmid pCL67 was obtained by digesting a plasmid containing a synthesized GFP loop with a glycosylation site (Invitrogen) with enzymes MfeI and DraI and ligation of the 246-bp insert into plasmid pSN28 (17), where the NheI site was inactivated (by QuikChange<sup>TM</sup>) and which was treated with the same enzymes.

To generate plasmids with EL5 mutations, the restriction sites for enzymes SwaI (S280N) and XhoI (silent) were introduced into the pMLBAD plasmid containing the *pglB* gene by the QuikChange<sup>TM</sup> method. Plasmid EL5 mutant 1 was obtained by cloning of a PCR fragment amplified with oligonucleotides 1 and 2 using a *Wolinella succinogenes* genome sequencing cosmid (18) as template. The PCR product was

digested with enzymes SwaI and XhoI and ligated into the pMLBAD plasmid digested with the same enzymes. The sequence of EL5 mutants 2, 3, 4, 6, and 8 was generated by gene synthesis (Invitrogen), and the pMLBAD plasmids for the corresponding EL5 variants were constructed by subcloning of provided vectors using enzymes SwaI and XhoI. The pBAD construct for EL5 mutant 8 was subcloned from the pMLBAD construct using enzymes NcoI and EcoRI. The mutation S280N, generated for the SwaI restriction site was removed from all final plasmids by the QuikChange<sup>TM</sup> method.

EL5 mutant 7 was obtained by overlap PCR to remove PglB residues 237–280. The upstream PCR fragment was generated with oligonucleotides 3/4, and the downstream fragment was generated with oligonucleotides 5/6. Overlap PCR was performed using the two purified PCR fragments and oligonucleotides 7/8. The overlap PCR fragment was digested with enzymes NcoI and EcoRI for ligation into the pBAD plasmid digested with the same enzymes; the fragment was digested with enzymes MfeI and PstI for ligation into the pMLBAD plasmid digested with enzymes EcoRI and PstI.

Constructs for EL5 mutant 5 were simultaneously created for the pBAD and pMLBAD plasmid using overlap PCR. For pMLBAD, the upstream PCR fragment was generated with oligonucleotides 9/10, and the downstream fragment was generated with oligonucleotides 11/12. For pBAD, the upstream PCR fragment was generated with oligonucleotides 13/10, and the downstream fragment was generated with oligonucleotides 11/14. Overlap PCR was performed using the purified PCR fragments and oligonucleotides 9/12 and 13/14, respectively. The overlap PCR fragments were ligated into zero blunt end TOPO vectors (Invitrogen), and the final expression plasmids were subcloned using enzymes EcoRI/PstI for the pMLBAD and NcoI/EcoRI for the pBAD vector, respectively.

All plasmids were validated by DNA sequencing. A description of the oligonucleotide primers used and the plasmids constructed can be found in supplemental Tables S1 and S2.

*In Vivo Glycosylation Assay*—*In vivo* complementation analysis of PglB mutants was performed as described before (12). Briefly, *Escherichia coli* SCM6 cells were transformed with three separate plasmids carrying 1) the *C. jejuni pglB*<sub>mut</sub> cluster (containing an inactivated *pglB* gene) (11), 2) the glycosylation acceptor protein 3D5 containing a DQNAT acceptor site (19), and 3) *C. lari* PglB, wild type or mutants. Expression and glycosylation of the acceptor protein 3D5 was monitored by SDS-PAGE of periplasmic cell extracts and visualized by mobility shift to increased size in an Immunoblot using an anti-c-Myc antibody (Calbiochem) or by the reactivity of the glycoprotein in an anti-glycan Immunoblot using hR6 antiserum. Immunodetection of PglB expression was performed from whole cell extracts using an anti-HA antiserum (Santa Cruz Biotechnology, Inc.).

*Expression and Purification of PglB Mutants*—Overexpression and purification of PglB mutants was performed as described previously (13). The desalted protein was analyzed by size exclusion chromatography (Superdex 200, GE Healthcare) and, if needed, concentrated up to 26 mg/ml in a 100 kDa molecular mass cut-off Ultra-15 concentrator (Amicon, Millipore). The concentrated sample was reanalyzed by size exclu-



sion chromatography, and protein concentrations were determined by absorbance at 280 nm using diluted samples.

EL5 cross-linking variants were prepared from whole cells. Cells of mutant Pos1 (N321C/L374C) and Pos2 (F311C/F579C) were resuspended in a buffer containing 10 mm Tris-HCl, pH 8.0, 250 mm NaCl, and 10 mm  $\beta$ -mercaptoethanol at a cell mass to buffer ratio of 1:7 ( $w/v$ ). The suspension was supplemented with 1% (w/v) *N-*dodecyl-β-ɒ-maltopyranoside (Anatrace) and sonicated for 5 min on ice. Sonication was repeated after 5 and 15 min, and the protein was solubilized for 1 h at 4 °C. Nickelnitrilotriacetic acid (Ni-NTA) equilibration and washing buffers contained 10 mm  $\beta$ -mercaptoethanol, which was removed in a last washing step prior to elution. The Ni-NTA eluate was supplemented with 10 mm  $MnCl<sub>2</sub>$  and 1 mm  $CuCl<sub>2</sub>$ , incubated for 20 min at 4 °C, and desalted into a buffer containing 10 mm Mes-NaOH, pH 6.5, 100 mm NaCl, 0.5 mm EDTA, 3% (v/v) glycerol, and *N-*dodecyl-β-D-maltopyranoside.

Cells of EL5 cross-linking mutant Pos3 (F308C/K528C) were resuspended as described above, omitting the addition of  $\beta$ -mercaptoethanol during solubilization and purification. The Ni-NTA-bound protein was washed for 20 column volumes, followed by 5 column volumes of imidazole-free wash and a 5-column volume wash containing  $1 \text{ mm } CuCl<sub>2</sub>$ . After a 15-min incubation at  $4^{\circ}$ C, CuCl<sub>2</sub> was washed out, and the protein was eluted and desalted as described above.

For glycosylation turnover and sequon binding measurements of reduced EL5 mutants, cross-linked samples were treated with 80 mm  $\beta$ -mercaptoethanol. Resistance to reduction of the induced disulfides in the presence of 80 mm  $\beta$ -mercaptoethanol in the folded state was demonstrated by inactivation of the reducing agent with 100 mM *N*-ethylmaleimide (Fluka) followed by SDS-PAGE under non-reducing conditions (see "Quantification of EL5 cross-linking efficiencies").

PglB mutant with deletion of TM helices 8 and 9 (mutant 7) was purified from membranes as described above. This mutant was not desalted but was directly concentrated to 3.5 mg/ml after elution from the Ni-NTA column and further analyzed by SDS-PAGE and MALDI-MS to prove the expected molecular weight.

PglB mutant carrying a cleavable EL5 loop (mutant 8) was purified from membranes as described above, and the desalted protein was incubated in a 1:3 molar ratio of PglB/3C protease (prepared in house) for 2 h at 4 °C. The protein was concentrated and repurified by preparative size exclusion chromatography (Superdex 200, GE Healthcare) and reconcentrated for final functional analysis.

*Expression and Purification of Glycoengineered GFP*—Glycoengineered GFP was overexpressed from pCL67 in *E. coli* BL-21 (DE3) Gold cells in 5-liter baffled flasks using Terrific Broth medium supplemented with 1% (w/v) glycerol. Cells were grown to  $A_{600}$  of 2.0 and induced with 0.5 mm isopropyl  $\beta$ -D-1thiogalactopyranoside for 4 h at 37 °C. Cells were harvested by centrifugation, and cell pellets were resuspended in a buffer containing 100 mM Tris-HCl, pH 8.0, and 100 mM NaCl at a cell mass/buffer ratio (w/v) of 1:5. Cells were disrupted by sonication on ice  $(3 \times 3 \text{ min})$ , and GFP was purified from the supernatant by Ni-NTA affinity chromatography and desalted into a buffer containing 100 mm Tris-HCl, pH 8.0, 100 mm NaCl, and 0.5 mM EDTA. The desalted protein was analyzed by size exclusion chromatography (Superdex 200, GE Healthcare), protein concentration was determined by absorbance at 280 nm, and the protein was stored at 4 °C.

*Expression and Purification of HmuT-Q147C*—HmuT-Q147C periplasmic substrate-binding protein was expressed as described previously (20). Cells were resuspended in a buffer containing 50 mM Tris-HCl, pH 8.0, and 150 mM NaCl at a cell mass/buffer ratio ( $w/v$ ) of 1:5, and cells were disrupted by sonication on ice  $(3 \times 3 \text{ min})$ . HmuT-Q147C was purified from the supernatant by Ni-NTA affinity chromatography and desalted into a buffer containing 10 mm Mes-NaOH, pH 6.5, 100 mm NaCl, 0.5 mm EDTA, and 3% (v/v) glycerol. The protein was supplemented with 10 mm tris(2-carboxyethyl)phosphine and stored at  $-20$  °C.

*Quantification of EL5 Cross-linking Efficiencies*—The crosslinking efficiency of PglB mutants Pos2 (F311C/F579C) and Pos3 (F308C/K528C) could be determined from a mobility shift to increased molecular weight after SDS-PAGE in the presence of non-reducing sample buffer. Samples were treated with an excess of *N*-ethylmaleimide (Fluka) prior to the addition of SDS sample buffer. Band intensities of oxidized and reduced species were determined from scanned gels (ImageJ). Mutant Pos1 (N321C/L374C) did not reveal a mobility shift upon cross-linking, and the residual non-cross-linked fraction was quantified by labeling of the free cysteines. The sample was incubated with a 3-fold molar excess of fluorescein 5-maleimide (Thermo Scientific) over cysteines for 10 min at room temperature, and the reaction was quenched with a molar excess of *N*-ethylmaleimide. The protein HmuT-Q147C carrying a single, surface-exposed cysteine (20, 21) was used as an internal standard. HmuT-Q147C labeling was performed after preparative size exclusion chromatography (to remove tris(2-carboxyethyl) phosphine) according to the procedure for PglB labeling. The labeled proteins were mixed in an equimolar ratio and resolved by SDS-PAGE, and fluorescent bands were visualized by using a Typhoon Trio Plus imager (GE Healthcare) with excitation at 488 nm and a 526-nm SP emission filter. Band intensities were determined from fluorescence gel scans (ImageJ). Fluorescence bands were additionally correlated to Immunoblot signals of the same gel. Immunodetection of PglB and HmuT-Q147C was performed using an anti-His<sub>5</sub> antibody (Qiagen).

*Synthesis of DQNAT Acceptor Peptide Labeled with 5-Carboxyfluorescein*—Peptide synthesis was performed as described previously (13).

*Fluorescence Anisotropy Measurements*—Measurements of acceptor peptide binding by fluorescence anisotropy were performed as described previously (13).

*Extraction of LLOs*—Isolation of LLOs containing the *C. jejuni* heptasaccharide was performed as described before (17). LLOs with the truncated disaccharide glycan structure  $(GaINAc-α1,3-diNAcBac-α1-PP-Und)$  were extracted from *E. coli* SCM6 cells carrying plasmid DC3 (containing an inactivated *pglB* gene) by a mixture of chloroform/MeOH of 2:1. Extracts were dried in a rotary evaporator and reconstituted in a buffer containing 10 mm Mes, pH 6.5, 100 mm NaCl, and 1% (w/v) Triton X-100. The concentration of reconstituted LLOs was determined by titrating various amounts of LLO against a



constant amount of fluorescently labeled acceptor peptide in an *in vitro* glycosylation assay (see below).

*In Vitro Glycosylation Assay*—*In vitro* glycosylation assays were performed as described previously (13) using fluorescently labeled acceptor peptide or glycoengineered GFP (10  $\mu$ M each) as acceptor substrate and*C. jejuni* LLO (heptasaccharide) or the disaccharide LLO (35  $\mu$ M each) as donor substrate. For turnover rate determination, a total of 5 samples were taken in 6-min intervals, and the concentration of PglB mutants was adjusted (1 nm to 50  $\mu$ m) so that the reaction was in a linear range. Data were fitted by linear regression to determine turnover rates.

For EL5 peptide titration experiments,  $15 \mu M$  PglB mutant 5 (Tyr-plug mutant) was incubated with various amounts of undecapeptides EL5 (acetyl-VLYQLKFYVFK-NH<sub>2</sub>) and EL5 mut (acetyl-VLYQLKFAVFK-NH<sub>2</sub>) (GenScript), respectively. Five samples were taken at appropriate time intervals (3–10 min) so that the reaction was in a linear range.

For analysis of GFP glycosylation, Triton-PAGE was used. Reactions were stopped by the addition of  $4\times$  native sample buffer (375 mM Tris-HCl, pH 6.8, 250 mM EDTA, 16.5% (w/v) sucrose, 0.1% (w/v) Triton X-100). Samples were diluted 7-fold prior to analysis by Triton-PAGE in minigels  $(8 \times 8 \text{ cm})$  consisting of an 8.5% resolving gel (0.5 M Tris-HCl, pH 8.8, 7% sucrose, 0.1% Triton X-100) and a 5% stacking gel (125 mm Tris-HCl, pH 6.8, 0.1% Triton X-100). Gels were run at 4 °C in a running buffer containing 50 mm Tris and 384 mm glycine. Fluorescent bands for glycosylated and non-glycosylated GFP were visualized by using a Typhoon Trio Plus imager (GE Healthcare) with excitation at 488 nm and a 610-nm emission filter. The amount of formed glyco-GFP was determined analogously to the amount of formed glycopeptides (13).

*Mass Spectrometry*—Mass spectra were acquired using a commercial MALDI-TOF/TOF<sup>TM</sup> mass spectrometer as described previously (22). The measurement was performed in the linear positive ion mode, and ionization was achieved with an Nd:YAG laser (355 nm). The mass spectrum was the average of 1,000 laser shots acquired at random sample positions. The sample (2  $\mu$ M) was directly mixed with the matrix solution (sinapinic acid, 20 mg/ml in water/acetonitrile/trifluoroacetic acid,  $49.95:49.95:0.1$ ) in a 1:1 (v/v) ratio before spotting. A mixture of cytochrome *c* and bovine serum albumin was applied for external calibration. Before data processing, the spectrum was background-subtracted and smoothed using a Savitzky-Golay algorithm (Igor Pro 6.2., Wavemetrics).

*NMR Spectroscopy*—NMR spectra were recorded at 283 K on Avance III 600-, 700-, or 750-MHz Bruker spectrometers equipped with a QCI, TCI, or TXI probe, respectively. The undecapeptide acetyl-VLYQLKFYVFK-NH<sub>2</sub> (GenScript) was dissolved in either  $D_2O$  or 95% H<sub>2</sub>O, 5% D<sub>2</sub>O at a concentration of 4 mM. Standard two-dimensional spectra were recorded for resonance assignment of the peptide: <sup>1</sup>H-<sup>1</sup>H TOCSY (mixing times of 13 and 80 ms),  $^{13}C^{-1}H$  HSQC, long range  $^{13}C^{-1}H$ HSQC, <sup>13</sup>C<sup>-1</sup>H HMQC-COSY, and <sup>1</sup>H-<sup>1</sup>H NOESY spectra (mixing time of 150 ms). All spectra are referenced to 4,4-dimethyl-4-silapentane-1-sulfonic acid, whereas  $^{13}$ C chemical shifts are indirectly referenced using a scaling factor  $\Xi$  of 0.251449530 (23). All spectra were processed with Topspin 3.1

(Bruker) and analyzed by Sparky 3 (T. D. Goddard and D. G. Kneller, University of California, San Francisco). Chemical shift assignments are shown in supplemental Figs. S5 and S6.

## **RESULTS**

*Cross-Linking EL5 to PglB Core near the Peptide Binding Site Abolishes Catalysis*—We first mutated the three cysteines in the sequence of *C. lari* PglB by mutagenesis (C17A, C30A, and C350S) and tested the *in vitro* activity and the sequon binding affinity of the resulting Cys-less enzyme (Table 1 and Fig. 1*C*). Cys-less PglB bound the bacterial glycosylation sequon DQNAT with an unchanged affinity compared with wild type enzyme ( $K_d$  of 1.18  $\pm$  0.02  $\mu$ M compared with 1.02  $\pm$  0.06  $\mu$ M for wild type PglB (13)) but showed a 2.2-fold reduction in glycosylation rate (0.69  $\pm$  0.02 peptides/s compared with 1.50  $\pm$ 0.04 peptides/s for wild type PglB (13)). We then designed covalent cross-links to restrict the motion of EL5 (Gly<sup>282</sup>–Gln<sup>330</sup>) by engineering double cysteine mutants at three distinct locations of PglB. The first cross-link we engineered was the double mutant N321C/L374C, attaching EL5 to TM helix 12, close to the peptide binding region (Fig. 1*A*, *Pos1*). Oxidative disulfide bond formation at Pos1 was determined to be 96–98% efficient by calibrated fluorescein labeling of unreacted cysteines, and the disulfide bond could be completely reduced upon the addition of 80 mm  $\beta$ -mercaptoethanol (Table 1 and supplemental Fig. S2*A*). This cross-link strongly affected peptide binding because we observed a 45-fold reduction in affinity for the oxidized compared with the reduced enzyme (Table 1 and Fig. 1*B*). Because the cross-linking efficiency was not 100%, our data suggested that the cross-linked species is unable to bind the peptide (or binds it with an unmeasurably low affinity) and that the remaining, non-cross-linked PglB (which has a  $K_d$  of 2.6  $\pm$ 0.2  $\mu$ <sub>M</sub>) accounts for the determined  $K_d$  of 117  $\pm$  11  $\mu$ <sub>M</sub> in the oxidized sample. This interpretation is consistent with the observed 40-fold increase in peptide glycosylation rate upon reduction of the disulfide cross-link at Pos1 (Table 1).We could exclude nonspecific effects by  $\beta$ -mercaptoethanol in the reduced sample because no significant effects on peptide binding or glycosylation were observed for the Cys-less mutant (Table 1 and Fig. 1*C*). In addition, the presence of 80 mm  $\beta$ -mercaptoethanol in the reaction did not affect PglB stability (supplemental Fig. S2*B*).

To evaluate the effect of substrate peptide size on the activity of our cross-linked PglB variants, we investigated the *in vitro* turnover of a GFP construct that contained the bacterial glycosylation sequon DQNAT flanked by flexible GGS sequences in a surface-exposed loop (Fig. 2*A*). Engineered GFP versions have previously been used both as a markers to monitor the *N*-glycosylation status of eukaryotic cell lines and as *in vitro* substrates of *C. jejuni* PglB (17, 24). Our *in vitro* glycosylation of GFP resulted in a mobility shift to increased molecular weight in native Triton-PAGE, and the educt/product ratio of the reaction could be directly quantified from fluorescence band intensities (Fig. 2*B*). We found Cys-less PglB to have a turnover rate of 6.7  $\pm$  0.4 GFP molecules/h (Table 1). The EL5 cross-link at Pos1 resulted in a virtually unchanged activity in the presence of reducing agents (7.5  $\pm$  0.6 GFP molecules/h) but a 40-fold decrease upon disulfide cross-linking (Table 1). Considering



## TABLE 1

#### **Acceptor peptide binding and glycosylation turnover rates of PglB variants containing EL5 disulfide cross-links**

Conditions specified as oxidized refer to copper-induced disulfide cross-linking, and those specified as reduced indicate the presence of 80 mm  $\beta$ -mercaptoethanol. Dissociation constants  $(K_d)$  were derived from curves shown in Fig. 1, *B* and *C*, and turnover rates were derived from three independent experiments. The indicated errors represent the S.E. of the fit for each turnover rate and dissociation constant determination.



<sup>a</sup> Values for relative affinity and relative turnover refer to the ratio between the reduced and the oxidized condition of the same mutant. In the case of mutants for Pos2 and Pos3 cross-linking, they refer to the ratio b

 $\frac{b}{c}$  Cross-linking efficiencies were determined by labeling of free cysteines with fluorescein 5-maleimide and subsequent quantification of fluorescence band intensities from scanned gels (supplemental Fig. S2A).

<sup>c</sup> Cross-linking resulted in a size shift after SDS-PAGE, and cross-linking efficiencies were determined from band intensities of Coomassie-stained gels (supplemental Fig. S2, *C* and *D*).



FIGURE 1.**Disulfide cross-linking of EL5 to core PglB.** *A*, schematic representation of *C. lari* PglB (PDB code 3RCE) with the TM domain in *blue*, the periplasmic domain in *green*, and bound acceptor peptide in *yellow*. EL5 connecting TM9 and TM10 is *colored red*, with the structurally disordered segment (25 residues) depicted by a *red dashed line*. Residues selected for cross-linking are shown in a *ball* and *stick representation* and *colored cyan*. The pairwise mutation to cysteines at three distinct positions is illustrated by *orange dashed circles*, and corresponding residues are indicated. *B* and *C*, peptide binding of different  $c$ ross-linking mutants quantified by fluorescence anisotropy. Purified PglB was titrated into a solution containing 1  $\mu$ M fluorescently labeled peptide variant (containing the sequon DQNAT) and 10 mm MnCl<sub>2</sub>. Data points reflect the mean of 20 measurements of the same sample. *Error bars*, S.D. Curves labeled "oxidized" indicate samples that were cross-linked with CuCl<sub>2</sub> during the purification process, whereas "reduced" indicates samples that contained 80 mm  $\beta$ -mercaptoethanol.





FIGURE 2. *In vitro* **glycosylation of glycoengineered GFP.** *A*, schematic representation of glycoengineered GFP (PDB code 1EMA) where a bacterial gly-cosylation site was introduced after Lys156 (*red loop*). The introduced site (DQNAT sequon) flanked by a GGGGS spacer is indicated *above* the image. *B*, determination of *in vitro* glycosylation turnover rate of glycoengineered GFP by Cys-less PglB. Glycosylated GFP (*g1*) is separated from non-glycosylated GFP (*g0*) after Triton-PAGE. Bands were visualized by a fluorescence gel scan at 488-nm excitation and 610-nm emission. The concentration of substrate GFP was 10  $\mu$ m, and PglB concentration was 1.3  $\mu$ m. The amount of glyco-GFP formed over time was determined from band intensities of the fluorescence gel scan, where the sum of the signals for glycosylated and non-glycosylated GFP for each lane was defined as 100%.

that our engineered GFP protein probably has a lower affinity for PglB than the chemically synthesized, fluorescently labeled substrate peptide, the  $\sim$ 40-fold increase in activity both in peptide and GFP glycosylation strongly suggested that the crosslinked PglB species (Pos1) is indeed completely inactive and that the remaining, non-cross-linked species  $(\sim 2.5\%)$  accounts for the observed basal activity in the oxidized sample. Therefore, we concluded that cross-linking EL5 at Pos1 completely inactivated the enzyme.

*Cross-Linking EL5 Distant from the Peptide Binding Site Decreases Glycosylation Rate*—We next investigated two crosslinks that attached EL5 to the periplasmic domain, more remote from the sequon binding site than Pos1 (Fig. 1*A*). Disulfide bond formation efficiencies at Pos2 and Pos3 were determined to be 94 and 95%, respectively, but both cross-links could only be completely reduced under denaturing conditions upon the addition of SDS (supplemental Fig. S2, *C* and *D*). We therefore compared the activities of PglB cross-linked at Pos2 and Pos3 with those of Cys-less PglB.We found that cross-linking at Pos2 or Pos3 reduced the peptide affinities 15- and 22-fold, respectively. In analogy to the cross-link at Pos1, the observed binding curves could be interpreted as the cross-linked species being unable to bind peptide and the remaining, small fraction of non-cross-linked PglB being fully active. However, the determined glycosylation rates suggest a distinct conclusion; the cross-link at Pos2 showed a turnover rate of 0.38  $\pm$  0.01 peptides/s, which is only 1.8-fold slower than that of Cys-less PglB, indicating that even the cross-linked species must be catalytically active and therefore able to bind peptide substrate. This observation is even more pronounced in the cross-link at Pos3, because the turnover rate for the substrate peptide was only 1.3-fold lower compared with Cys-less PglB. We also determined GFP substrate turnover rates for cross-links at Pos2 and Pos3. In both cases we found a more pronounced reduction in GFP than in peptide glycosylation (7.8-fold lower for Pos2 and

6.7-fold lower for Pos3; Table 1). We concluded that crosslinking EL5 at some distance from the peptide binding site has a strong effect both on sequon binding and on glycosylation of protein substrates but only a moderate effect on glycosylation of small peptide substrates.

*A Short Motif in EL5, the Tyr-plug, Is Essential for PglB Function*—In the *C. lari* PglB structure with bound acceptor peptide (12), the first 24 residues of EL5 (Leu<sup>283</sup>-Ala<sup>306</sup>) were disordered and not visible in the electron density (supplemental Fig. S1). Considering their location, it is tempting to speculate that these residues might interact with the glycan of bound LLO. A sequence alignment revealed that EL5 from *W. succinogenes* has the lowest similarity among different bacterial PglB homologs (Fig. 3*B*), and because the *W. succinogenes* glycan exhibits structural differences compared with that of *C. lari* (25–27), we replaced the EL5 sequence of *C. lari* PglB with the corresponding sequence of *W. succinogenes* (mutant 1). When analyzing the resulting *in vivo* activity, we found only a marginal reduction in acceptor substrate glycosylation compared with wild type *C. lari* PglB, suggesting that the EL5 sequence does not directly account for glycan specificity (Fig. 3*C*). We next created several EL5 versions where we inserted, replaced, and deleted residues in the disordered region (Fig. 3, *A* and *B*). An insertion of 8 amino acids at the beginning or end of the disordered region (mutants 6 and 2, respectively) did not impair PglB activity. Unexpectedly, neither did a deletion of transmembrane helices 8 and 9 (mutant 7), which alters the location of the first residue of EL5 (Fig. 3, *A–C*). It is notable that despite a reduction in mass by 5.2 kDa (determined by analyzing the purified PglB variant by MALDI-MS (22), mutant 7 and WT PglB have similar electrophoretic mobilities in SDS-PAGE (supplemental Fig. S3).

Sequence alignments further suggested that four aromatic residues in EL5 are conserved in bacterial homologs of PglB, whereas the middle section of EL5 varies in length and sequence (Fig. 3*B*). Replacement of the non-conserved region of EL5 by a flexible  $(GGS)_{3}$  sequence (mutant 3) did not affect the *in vivo* activity of PglB, and deletion of this region (mutant 4) only slightly reduced catalysis. However, we found that when a stretch of 8 amino acids, including the four aromatic residues, was replaced with a 3C cleavage site, PglB activity was completely abolished (mutant 5; Fig. 3, *A–C*). We termed this short stretch the "Tyr-plug" because of the contained tyrosines. When we analyzed the Tyr-plug mutant (mutant 5) in greater detail by previously established *in vitro* assays, we unexpectedly found that peptide substrate binding was only slightly affected (1.6-fold weaker binding than WT PglB), whereas glycosylation turnover was reduced 14,000-fold (Table 2 and Fig. 4*A*).

*Tyr<sup>293</sup> Has a Key Role in the Function of the Tyr-plug*—In mutant 5, all four aromatic residues of the Tyr-plug  $(Tyr^{288})$ , Phe<sup>292</sup>, Tyr<sup>293</sup>, and Phe<sup>295</sup>) were replaced. These residues are generally conserved among bacterial PglB homologs (Fig. 3*B*), and because aromatic residues are frequently involved in sugar interactions via  $\pi$ -stacking (28–30), we wished to clarify if the aromatic side chains were responsible for the observed drop in PglB reactivity. We thus created various alanine double mutations and tested their activities*in vivo*. We found that all double mutants that contained the Y293A mutation showed impaired



FIGURE 3. **Functional analysis of the disordered part of EL5.** *A*, surface representation of *C. lari* PglB (PDB code 3RCE) with the TM domain in *blue*, the periplasmic domain in *green*, bound acceptor peptide in *yellow*, and EL5 in *red*. The disordered part of EL5 is indicated by a *red dashed line*. *Cyan lines* indicate regions of replacements, *black lines* indicate sequence deletions, and *brackets* indicate insertions. The *white dashed line* visualizes the removal of TM helices 8 and 9. *Numbers* correspond to the sequences shown in *B* and describe mutations introduced into EL5 from the C to the N terminus, successively away from the active site. *B,* sequence alignment of EL5 in bacterial PgIB homologs. The sequence corresponding to the disordered part of EL5 is indicated by a *red dashed line.*<br>Conservation of the catalytically important residue Glu<sup></sup> highlighted in *magenta*. Other conserved residues in the disordered region are shown in *gray*. For mutant 1, EL5 from *C. lari* PglB (Gly282–Gln330) was replaced by EL5 from *W. succinogenes* EL5 (Ala283–Ile329). *Lines*, *brackets*, and *numbers* are as in *A*. *C–E*, *in vivo* glycosylation assay in *E. coli* to investigate the activity of different EL5 mutants. Shown are immunoblots, detecting acceptor protein 3D5 (*top*), bacterial *N*-glycans (*middle*), and PglB (*bottom*). PglB mutants are indicated *above t*he *lanes.* Glycosylation results in a mobility shift from the non-glycosylated (g0) to the glycosylated form of the acceptor protein (g1).<br>Functional PgIB is partially autoglycosylated at Asn<sup>535</sup> and A triplicate, and representative samples are shown. *C*, *numbers above* the *lanes* correspond to the *numbers* shown in *A* and *B*.

## TABLE 2

**Peptide binding affinities and turnover rates for different EL5 mutants** Dissociation constants  $(K_d)$  were derived from curves shown in Fig. 4. The errors represent the S.E. of the fit. The relative turnover refers to the activity of an individual mutant compared withWT PglB. Activities were determined using fluorescently labeled acceptor peptide, and individual turnover rates were derived from three independent experiments.



function (Fig. 3*D*). A subsequent analysis of single mutants confirmed that Y293A leads to almost complete loss of activity, whereas the other mutations showed very little to no effect (Fig.

3*D*). Although tyrosine showed the strongest activity, the aromatic residues tryptophan, phenylalanine, and even histidine were tolerated at position 293 (Fig. 3*E*).

Because an aromatic side chain at position 293 was essential for PglB function, we initially assumed that Tyr<sup>293</sup> forms a stacking interaction with one of the distal GalNAc residues of the LLO glycan, particularly because Gal-configured units represent ideal stacking partners for aromatic residues (30). To study the effect of a truncated glycan structure, we took advantage of an engineered version of the *C. jejuni pgl* gene cluster, yielding a LLO donor with a disaccharide structure (GalNAc-  $\alpha$ 1,3-diNAcBac- $\alpha$ 1-PP-Und) (16). We used extracts of this short LLO variant along with WT *C. jejuni* heptasaccharide





FIGURE 4. Peptide binding of different EL5 mutants quantified by fluorescence anisotropy. Purified PglB was titrated into a solution containing 1  $\mu$ M fluorescently labeled peptide variant (containing the sequon DQNAT) and 10 mm MnCl<sub>2</sub>. Data points reflect the mean of 20 measurements of the same sample. *Error bars*, S.D. *A*, binding curve of PglB with replacement of the essential EL5 Tyr-plug sequence element (mutant 5, *violet curve*) compared with WT PglB (*black curve*, as presented in Ref. 13). *B*, binding curve of PglB with a cleavable EL5 (mutant 8 as presented in Fig. 6). The *black curve* represents PglB with an intact EL5 (uncleaved), whereas the *blue curve* results from PglB with cleaved EL5.

### TABLE 3

#### **Relative glycosylation turnover rates of different LLO donor substrates and PglB Tyr293 mutants**

The relative turnover (-fold reduction) refers to the activity of an individual reaction compared with the reaction of WT PglB and LLO with the heptasaccharide structure. Activities were determined using fluorescently labeled acceptor peptide, and individual turnover rates were derived from three independent experiments. *Red hexagon,* di-*N*-acetylbacillosamine (diNAcBac); *yellow square, N*-acetylgalactosamine; *blue circle,* glucose.



LLO at equal concentrations in an *in vitro* glycosylation assay and determined glycosylation turnover rates for different Tyr<sup>293</sup> mutants. Using full-length *C. jejuni* LLO, we found reductions in turnover of 44-fold and 7,000-fold when  $\text{Tyr}^{293}$ was mutated to Phe and Ala, respectively (Table 3), confirming our *in vivo* findings that Tyr<sup>293</sup> predominantly accounts for the effect of the Tyr-plug mutant (mutant 5) and that replacing Tyr<sup>293</sup> by other aromatic amino acids affects PglB activity much less than removing the aromatic ring. Using the disaccharide LLO, we found a 2-fold reduction for WT PglB, which we interpret to be a result of slightly different biophysical properties of the truncated LLO. To our surprise, the turnover rates of mutants Y293F and Y293A were 41-fold and 8,000-fold reduced with the disaccharide LLO (Table 3). This excluded an interaction of Tyr<sup>293</sup> with one of the distal GalNAc residues of the LLO.

*A Synthetic Peptide Containing the Tyr-plug Confers Functionality*—Our *in vivo* glycosylation studies suggested that the exact location of the Tyr-plug (including  $Tyr^{293}$ ) within EL5 is not important for catalysis because insertions before and after the motif were tolerated. This finding strongly suggested that the Tyr-plug had a defined three-dimensional conformation during catalysis.We therefore performed solution NMR analyses of a synthetic undecapeptide containing the Tyr-plug sequence (peptide EL5-11, VLYQLKFYVFK). The assignment of a corresponding two-dimensional NOESY spectrum of peptide EL5-11 did not reveal any long range NOEs, indicating the absence of a secondary structure (supplemental Fig. S4). The lack of a specific conformation of peptide EL5-11 was further confirmed by the assignment of a <sup>15</sup>N-<sup>1</sup>H HSQC spectrum showing a small dispersion of obtained signals (supplemental Fig. S5). Nevertheless, the synthetic EL5-11 peptide did retain certain properties needed to support catalysis, because it could, to some extent, rescue the impaired function of a Tyr-plug mutant *in vitro*. When we determined glycosylation rates of the Tyr-plug mutant, we found that supplementing the reaction with peptide EL5-11 increased the glycosylation activity up to 3.6-fold at 100  $\mu$ M peptide concentration. The activity decreased at higher peptide concentrations, which may be caused by limited peptide solubility or by nonspecific interactions with the detergent micelle (Fig. 5*B*). Considering the 14,000-fold reduction in activity for the Tyr-plug mutant (Table 2), the rescuing effect of the EL5-11 peptide might appear marginal. However, it is a highly specific effect depending on the presence of  $\text{Tyr}^{293}$ , because a control peptide containing the Y293A mutation did not affect the activity of the Tyr-plug mutant (Fig. 5*B*). A limited local concentration of peptide EL5-11 caused by diffusion and by competition against the existing, mutated EL5 (Fig. 5*A*) might explain this small effect.

*The Two Functions of EL5 Are Independent*—To investigate if the two functions of EL5 in catalysis are independent, we introduced a proteolytic cleavage site into EL5. The eight amino acid insertions in mutants 2 and 6 (and the sequence replacement of mutant 5) all represent 3C protease cleavage





FIGURE 5. **Complementation of impaired EL5 function by a synthetic EL5 peptide.** *A*, *surface representation* of PglB, as shown in Fig. 3*A*, with replacement of the essential Tyr-plug indicated in *cyan*. *B*, the turnover rate of the Tyr-plug mutant (mutant 5) was determined at various concentrations of synthetic EL5 peptide. Data points reflect single turnover rate determinations. *Error bars*, S.E. of each fit. The sequence of used EL5 peptides is shown *above* the diagram.





FIGURE 6. **Design and analysis of PglB with a cleavable EL5.** *A*, amino acid sequence of cleavable EL5 (mutant 8). The non-conserved region of EL5 (boldface type) was replaced by a 3C proteolytic cleavage site (red) flanked by a (GGS)<sub>2</sub> element (blue) on both sides. The *underlined residues* are still ordered in the PglB structure. *B*, *in vivo* glycosylation assay in *E. coli* to investigate the activity of PglB with a cleavable EL5. Shown are immunoblots detecting acceptor protein 3D5 (*top*), bacterial *N*-glycans (*middle*), and PglB (*bottom*). PglB mutants are indicated *above* the *lanes*. Glycosylation results in a mobility shift from the non-glycosylated (*g0*) to the glycosylated form of the acceptor protein (*g1*). Functional PglB is partially autoglycosylated at Asn<sup>535</sup> and Asn556, resulting in two additional bands (*g1* and *g2*). *C*, SDS-PAGE of uncleaved and cleaved PglB. Fragments resulting from proteolytic cleavage are indicated with *arrows*. *D*, size exclusion chromatograms of purified and concentrated PglB EL5 cleave samples after peptide binding (fluorescence anisotropy) measurements. *Black profile*, PglB with intact EL5; *red profile*, sample with cleaved EL5. Note that the superimposed chromatograms are not normalized for the amount of protein loaded onto the gel filtration column. The analysis is used exclusively to assess monodispersity and integrity of this PglB mutant.

sites, but neither construct was found to be efficiently cleaved by 3C protease, possibly due to steric effects. We therefore designed a mutant where 10 non-conserved EL5 residues were replaced by a 3C cleavage site flanked by (GGS)<sub>2</sub> spacer sequences on both sides (mutant 8; Fig. 6A). This mutant was active *in vivo*, could be cleaved efficiently with 3C protease *in vitro*, and showed good monodispersity before and after protease treatment (Fig. 6, *B–D*). Upon cleavage, peptide binding was reduced 2.6-fold relative to the uncut variant (Table 2 and Fig. 4*B*). In addition, cleavage only resulted in a marginal effect on glycosylation turnover, because the cut version revealed a 1.3-fold lower turnover rate than the uncut protein (Table 2). We conclude that the catalytically essential Tyr-plug in EL5 does not have to be

covalently connected to the structurally ordered, peptidebinding segment of EL5 to fulfill its function.

## **DISCUSSION**

The long periplasmic loop EL5 was a remarkable yet insufficiently understood feature of the PglB structure. The C-terminal half of EL5 is involved in generating a tight binding site for acceptor peptide, suggesting that in a peptide-free state, PglB might adopt a distinct conformation where EL5 was disengaged from the core enzyme, as depicted in a three-step reaction cycle (supplemental Fig. S1).We tested this hypothesis using cysteine cross-linking experiments to restrict the flexibility of EL5. We indeed found that a cross-link at the peptide binding site (Pos1) abolished sequon binding and catalysis. In this PglB variant,





FIGURE 7. **Proposed function of the Tyr-plug in PglB.** *A*, *surface representation* of PglB *colored* as in Fig. 3*A*. Binding of LLO donor substrate results in an interaction between the reducing end sugar and the Tyr-plug (*thick red line*). *Red hexagon*, diNAcBac; *yellow square*, *N*-acetylgalactosamine; *blue circle*, glucose; *circled P*, phosphate group; *gray line*, undecaprenyl (*Und*). *B*, mechanistic hypothesis of Tyr-plug function. *Yellow dashed line* indicates the sequon backbone.<br>Hydrogen bonds between PgIB residues Asp<sup>56</sup> and Glu<sup>31</sup> attack on the anomeric carbon (*orange circle*). The C-2 acetamido group of the reducing end sugar can be accommodated in a cavity of conserved residues in the active site of PglB (*blue dashed line*). The Tyr-plug is proposed to interact with the reducing end sugar and with the PglB surface. This is expected to restrict the rotation of the *O*-glycosidic bond and the first O–P bond (*red arrows*), increasing catalytic efficiency. *R*, oligosaccharyl; *Und*, undecaprenyl.

EL5 may be unable to achieve an open conformation that is required for peptide binding, or the sequon binding site might have been distorted by the cross-link. The other two cross-links (positions 2 and 3), which are not overlapping with the peptide binding site, showed a reduced but clearly measurable activity. Our data allow the following conclusions. 1) Sequon binding can occur even when EL5 is only partially disengaged from the enzyme. 2) Sequon binding is not the rate-limiting step during catalysis *in vitro* because the strong reduction of peptide binding affinity (22-fold for Pos3) only leads to a slight reduction (1.3-fold for Pos3) in turnover. This interpretation is in agreement with previous observations that a replacement of the  $+2$ Thr in glycosylation sequons by Ser, which reduced peptide binding 4-fold, only lowered turnover 1.2-fold (13). 3) The larger the peptide substrate, the stronger the effect of EL5 disengagement (for a comparison of peptide *versus* GFP glycosylation, see Table 1). 4) Partial disengagement of EL5 is sufficient for the release of glyco-products from the enzyme. Notably, product dissociation can only occur if either the polypeptide or the sugar moiety passes through the opening provided by EL5. Our observation that glycosylated GFP can be released even in the presence of a disulfide cross-link at Pos3 suggests that the *N*-glycan moiety of glyco-products is pulled through the provided opening, because GFP would be unable to fit.

X-ray structures of the archaeal Stt3 homolog AglB from *Archeoglobus fulgidus* in two crystal forms have recently been reported (31). Neither AglB structure contained bound acceptor peptide or LLO substrate. The structures revealed a transmembrane fold that is similar to that of our *C. lari* PglB structure. The entire loop EL5 is disordered in the AglB structure with bound zinc and sulfate ions, in line with our finding that EL5 is disengaged from the enzyme core in the apo-state of PglB. In the other AglB structure, the entire loop EL5 is ordered, including the N-terminal half, which is located at a distance of  $\sim$ 15–20 Å from the active site of the enzyme, suggesting that a considerable conformational change would be required for interaction with the reducing end sugar. In addition to large scale conformational rearrangements of EL5, conformational fluctuations within the extracellular domain of the archaeal Stt3 homolog from *Pyrococcus furiosus* (AglB) have been reported to be important for function (32), suggesting that the extracellular/periplasmic domain probably adjusts its conformation upon substrate binding and EL5 engagement.

In contrast to the C-terminal half, any role of the N-terminal half of EL5 in catalysis was completely unknown, and the corresponding segment was disordered in the x-ray structure of sequon-bound *C. lari* PglB (12). Through a combination of systematic mutagenesis and *in vivo* activity analysis, we now identified a previously unrecognized motif that proved essential for catalytic activity. Unexpectedly, mutations in the newly found Tyr-plug had stronger effects on PglB activity than many of the previously described mutations at the active site, even of catalytically essential residues (13, 14). However, replacing the Tyrplug sequence did not affect substrate peptide binding. This suggested a role in the interaction with the LLO molecule, the glycan donor of the reaction. Within the Tyr-plug, Tyr<sup>293</sup> has a critical effect, but it was dependent on the immediate sequence context. The observation that Tyr<sup>293</sup> could be replaced by other aromatic residues supported the notion that residue 293 might be involved in an interaction with a sugar moiety of the LLO substrate, because aromatic residues are frequently involved in  $\pi$ -stacking with carbohydrates (28–30). However, our experi-

ments excluded a direct interaction of  $\text{Tyr}^{293}$  with one of the distal sugar residues of the *C. jejuni* LLO, which agrees both with the observation that *C. jejuni* PglB can transfer diverse carbohydrate structures*in vivo* (33, 34) and with *in vitro* studies of eukaryotic OST that identified the dolichol-linked monosaccharide Dol-PP-GlcNAc as the minimal glycosylation unit (35).

On the one hand, the exact position of the Tyr-plug within EL5 seemed to be irrelevant, because insertions, replacements, and even deletions in its vicinity were tolerated. On the other hand, the essential function of  $Tyr^{293}$  could not be taken over by any of the other aromatic residues within the Tyr-plug (Tyr<sup>288</sup>, Phe292, or Phe295). The inescapable conclusion was that the Tyr-plug adopts a defined three-dimensional conformation, at least during one of the rate-limiting steps in catalysis. We could indeed rescue (albeit to a limited extent) the effect of a Tyr-plug replacement in PglB by adding a synthetic peptide containing the Tyr-plug motif. However, this peptide did not adopt a defined structure on its own, as we could show by solution NMR analysis. This suggests that the Tyr-plug only adopts a defined conformation in the context of the enzyme and in the presence of bound LLO (Fig. 7*A*).

What might the essential catalytic role of the Tyr-plug be? Fig. 7 illustrates our mechanistic hypothesis. Once LLO substrate is bound to PglB, its pyrophosphate moiety is bound by interactions with the conserved residue  $Arg<sup>375</sup>$  and the divalent metal cation, both of which are essential for catalysis (Fig. 7*B*) (12, 13). Although the pyrophosphate is now fixed, the reducing end sugar is not, because a rotation around the *O*-glycosidic bond and around the O–P bond of the pyrophosphate moiety would be allowed (*red arrows* in Fig. 7*B*), both of which would prevent proper placement of the glycan for a nucleophilic attack by the activated carboxamide group of the Asn side chain of the sequon. We therefore propose that the function of the Tyr-plug is to engage in contacts both with the active site of PglB and with the reducing end sugar of LLO, forcing the reducing end sugar into a reactive conformation and thus increasing the rate of the transfer reaction. The aromatic side chain of  $Tyr^{293}$  is essential and might directly stack against the reducing end sugar diNAcBac (Fig. 7B). In addition, Tyr<sup>293</sup> might interact with the C-4 acetamido group of diNAcBac, because methyl groups of acetamido moieties are often recognized by aromatic residues in different lectins (30, 36–39). A tentative modeling of the LLO with the diNAcBac properly aligned for a nucleophilic attack places the C2-acetamido group into a PglB cavity that contains residues conserved in all Stt3 homologs. This cavity might also be involved in restricting sugar rotation, in particular rotation around the anomeric bond (Fig. 7*B*). The C2-acetamido group of the reducing end sugar is indeed essential both for bacterial and eukaryotic LLO substrates (34, 40).

Does the Tyr-plug also exist in archaeal and eukaryotic Stt3? Although the chemical reaction and the basic mechanism are conserved among OST homologs, it is currently unclear if eukaryotic and archaeal enzymes use a similar Tyr-plug to restrict the motion of the reducing end sugar during the catalytic step. Sequence alignments would support this hypothesis, because Stt3 homologs also contain aromatic residues (including Tyr and Phe) in EL5 at a similar distance from the catalytically essential Glu<sup>319</sup> residue, which is also conserved. However, it will be very challenging to study these effects*in vivo* or*in vitro*. Most eukaryotic OSTs contain seven additional subunits (41) that might contribute to binding of the LLO and the correct positioning of the reducing end sugar moiety. Additionally, the glycan structure transferred by eukaryotic OST (GlcNAc<sub>2</sub>Man<sub>9</sub>Glc<sub>3</sub>) is different and much bigger than its bacterial counterpart, which could imply that steric effects restricting sugar motions play an additional role. Furthermore, the reducing end sugar is a GlcNAc in eukaryotic LLOs and even a hexose in some archaeal LLOs (4, 31, 42, 43), suggesting mechanistic diversity.

In summary, our study shows that EL5 of PglB has two distinct functions.Whereas the N-terminal half of EL5 is supposed to form a critical interaction with the LLO during catalysis, the C-terminal part contributes to peptide binding and contributes the conserved active site residue Glu<sup>319</sup> that binds both the divalent metal and the acceptor Asn of the sequon. Intriguingly, the two functions of EL5 require no covalent link between the N- and C-terminal halves of the loop, as shown by our ability to specifically cleave EL5 with only minor effects on activity. Although the importance of the Tyr-plug for catalysis is now established, insight into the mechanistic details will require structural evidence, in particular a LLO-bound co-crystal structure of OST.

*Acknowledgments—We thank E. Weber-Ban and R. Glockshuber for access to the fluorometer and M. Aebi for helpful discussions. We are grateful to the D-BIOL NMR Platform for access to NMR spectrometers. We thank A. Geerlof for providing the protocol for 3C protease expression and purification and A. Zeltina for providing HmuT-Q147C cells.*

#### **REFERENCES**

- 1. Helenius, A., and Aebi, M. (2004) Roles of *N*-linked glycans in the endoplasmic reticulum. *Annu. Rev. Biochem.* **73,** 1019–1049
- 2. Kornfeld, R., and Kornfeld, S. (1985) Assembly of asparagine-linked oligosaccharides. *Annu. Rev. Biochem.* **54,** 631–664
- 3. Varki, A. (1993) Biological roles of oligosaccharides. All of the theories are correct. *Glycobiology* **3,** 97–130
- 4. Eichler, J. (2013) Extreme sweetness. Protein glycosylation in archaea. *Nat. Rev. Microbiol.* **11,** 151–156
- 5. Nothaft, H., and Szymanski, C. M. (2010) Protein glycosylation in bacteria. Sweeter than ever. *Nat. Rev. Microbiol.* **8,** 765–778
- 6. Schwarz, F., and Aebi, M. (2011) Mechanisms and principles of *N*-linked protein glycosylation. *Curr. Opin. Struct. Biol.* **21,** 576–582
- 7. Nilsson, I., Kelleher, D. J., Miao, Y., Shao, Y., Kreibich, G., Gilmore, R., von Heijne, G., and Johnson, A. E. (2003) Photocross-linking of nascent chains to the STT3 subunit of the oligosaccharyltransferase complex. *J. Cell Biol.* **161,** 715–725
- 8. Yan, Q., and Lennarz, W. J. (2002) Studies on the function of oligosaccharyl transferase subunits. Stt3p is directly involved in the glycosylation process. *J. Biol. Chem.* **277,** 47692–47700
- 9. Hese, K., Otto, C., Routier, F. H., and Lehle, L. (2009) The yeast oligosaccharyltransferase complex can be replaced by STT3 from *Leishmania major*. *Glycobiology* **19,** 160–171
- 10. Nasab, F. P., Schulz, B. L., Gamarro, F., Parodi, A. J., and Aebi, M. (2008) All in one. *Leishmania major* STT3 proteins substitute for the whole oligosaccharyltransferase complex in *Saccharomyces cerevisiae*. *Mol. Biol. Cell* **19,** 3758–3768
- 11. Wacker, M., Linton, D., Hitchen, P. G., Nita-Lazar, M., Haslam, S. M., North, S. J., Panico, M., Morris, H. R., Dell, A., Wren, B. W., and Aebi, M. (2002) *N*-Linked glycosylation in *Campylobacter jejuni* and its functional



transfer into *E. coli*. *Science* **298,** 1790–1793

- 12. Lizak, C., Gerber, S., Numao, S., Aebi, M., and Locher, K. P. (2011) X-ray structure of a bacterial oligosaccharyltransferase. *Nature* **474,** 350–355
- 13. Gerber, S., Lizak, C., Michaud, G., Bucher, M., Darbre, T., Aebi, M., Reymond, J. L., and Locher, K. P. (2013) Mechanism of bacterial oligosaccharyltransferase. *In vitro* quantification of sequon binding and catalysis. *J. Biol. Chem.* **288,** 8849–8861
- 14. Lizak, C., Gerber, S., Michaud, G., Schubert, M., Fan, Y. Y., Bucher, M., Darbre, T., Aebi, M., Reymond, J. L., and Locher, K. P. (2013) Unexpected reactivity and mechanism of carboxamide activation in bacterial *N*-linked protein glycosylation. *Nat. Commun.* **4,** 2627
- 15. Lefebre, M. D., and Valvano, M. A. (2002) Construction and evaluation of plasmid vectors optimized for constitutive and regulated gene expression in *Burkholderia cepacia* complex isolates. *Appl. Environ. Microbiol.* **68,** 5956–5964
- 16. Linton, D., Dorrell, N., Hitchen, P. G., Amber, S., Karlyshev, A. V., Morris, H. R., Dell, A., Valvano, M. A., Aebi, M., and Wren, B. W. (2005) Functional analysis of the *Campylobacter jejuni N*-linked protein glycosylation pathway. *Mol. Microbiol.* **55,** 1695–1703
- 17. Kowarik, M., Numao, S., Feldman, M. F., Schulz, B. L., Callewaert, N., Kiermaier, E., Catrein, I., and Aebi, M. (2006) *N*-Linked glycosylation of folded proteins by the bacterial oligosaccharyltransferase. *Science* **314,** 1148–1150
- 18. Baar, C., Eppinger, M., Raddatz, G., Simon, J., Lanz, C., Klimmek, O., Nandakumar, R., Gross, R., Rosinus, A., Keller, H., Jagtap, P., Linke, B., Meyer, F., Lederer, H., and Schuster, S. C. (2003) Complete genome sequence and analysis of *Wolinella succinogenes. Proc. Natl. Acad. Sci. U.S.A.* **100,** 11690–11695
- 19. Lizak, C., Fan, Y. Y., Weber, T. C., and Aebi, M. (2011) *N*-Linked glycosylation of antibody fragments in *Escherichia coli*. *Bioconjug. Chem.* **22,** 488–496
- 20. Mattle, D., Zeltina, A., Woo, J. S., Goetz, B. A., and Locher, K. P. (2010) Two stacked heme molecules in the binding pocket of the periplasmic heme-binding protein HmuT from *Yersinia pestis*. *J. Mol. Biol.* **404,** 220–231
- 21. Woo, J. S., Zeltina, A., Goetz, B. A., and Locher, K. P. (2012) X-ray structure of the *Yersinia pestis* heme transporter HmuUV. *Nat. Struct. Mol. Biol.* **19,** 1310–1315
- 22. Chen, F., Gerber, S., Heuser, K., Korkhov, V. M., Lizak, C., Mireku, S., Locher, K. P., and Zenobi, R. (2013) High-mass matrix-assisted laser desorption ionization-mass spectrometry of integral membrane proteins and their complexes. *Anal. Chem.* **85,** 3483–3488
- 23. Markley, J. L., Bax, A., Arata, Y., Hilbers, C. W., Kaptein, R., Sykes, B. D., Wright, P. E., and Wuthrich, K. (1998) Recommendations for the presentation of NMR structures of proteins and nucleic acids (IUPAC Recommendations 1998). *Pure Appl. Chem.* **70,** 117–142
- 24. Losfeld, M. E., Soncin, F., Ng, B. G., Singec, I., and Freeze, H. H. (2012) A sensitive green fluorescent protein biomarker of *N*-glycosylation site occupancy. *FASEB J.* **26,** 4210–4217
- 25. Dell, A., Galadari, A., Sastre, F., and Hitchen, P. (2010) Similarities and differences in the glycosylation mechanisms in prokaryotes and eukaryotes. *Int. J. Microbiol.* **2010,** 148178
- 26. Nothaft, H., and Szymanski, C. M. (2013) Bacterial protein *N*-glycosylation. New perspectives and applications. *J. Biol. Chem.* **288,** 6912–6920
- 27. Jervis, A. J., Butler, J. A., Lawson, A. J., Langdon, R., Wren, B. W., and Linton, D. (2012) Characterization of the structurally diverse *N*-linked

glycans of *Campylobacter* species. *J Bacteriol.* **194,** 2355–2362

- 28. Morales, J. C., Reina, J. J., Díaz, I., Aviñó, A., Nieto, P. M., and Eritja, R. (2008) Experimental measurement of carbohydrate-aromatic stacking in water by using a dangling-ended DNA model system. *Chemistry* **14,** 7828–7835
- 29. Oldham, M. L., Khare, D., Quiocho, F. A., Davidson, A. L., and Chen, J. (2007) Crystal structure of a catalytic intermediate of the maltose transporter. *Nature* **450,** 515–521
- 30. Weis, W. I., and Drickamer, K. (1996) Structural basis of lectin-carbohydrate recognition. *Annu. Rev. Biochem.* **65,** 441–473
- 31. Matsumoto, S., Shimada, A., Nyirenda, J., Igura, M., Kawano, Y., and Kohda, D. (2013) Crystal structures of an archaeal oligosaccharyltransferase provide insights into the catalytic cycle of *N*-linked protein glycosylation. *Proc. Natl. Acad. Sci. U.S.A.* **110,** 17868–17873
- 32. Nyirenda, J., Matsumoto, S., Saitoh, T., Maita, N., Noda, N. N., Inagaki, F., and Kohda, D. (2013) Crystallographic and NMR evidence for flexibility in oligosaccharyltransferases and its catalytic significance. *Structure* **21,** 32–41
- 33. Feldman, M. F., Wacker, M., Hernandez, M., Hitchen, P. G., Marolda, C. L., Kowarik, M., Morris, H. R., Dell, A., Valvano, M. A., and Aebi, M. (2005) Engineering *N*-linked protein glycosylation with diverse O antigen lipopolysaccharide structures in *Escherichia coli*. *Proc. Natl. Acad. Sci. U.S.A.* **102,** 3016–3021
- 34. Wacker, M., Feldman, M. F., Callewaert, N., Kowarik, M., Clarke, B. R., Pohl, N. L., Hernandez, M., Vines, E. D., Valvano, M. A., Whitfield, C., and Aebi, M. (2006) Substrate specificity of bacterial oligosaccharyltransferase suggests a common transfer mechanism for the bacterial and eukaryotic systems. *Proc. Natl. Acad. Sci. U.S.A.* **103,** 7088–7093
- 35. Tai, V. W., and Imperiali, B. (2001) Substrate specificity of the glycosyl donor for oligosaccharyl transferase. *J. Org. Chem.* **66,** 6217–6228
- 36. Weis, W., Brown, J. H., Cusack, S., Paulson, J. C., Skehel, J. J., and Wiley, D. C. (1988) Structure of the influenza virus haemagglutinin complexed with its receptor, sialic acid. *Nature* **333,** 426–431
- 37. Wright, C. S. (1984) Structural comparison of the two distinct sugar binding sites in wheat germ agglutinin isolectin II. *J. Mol. Biol.* **178,** 91–104
- 38. Wright, C. S. (1990) 2.2 Å resolution structure analysis of two refined *N*-acetylneuraminyl-lactose–wheat germ agglutinin isolectin complexes. *J. Mol. Biol.* **215,** 635–651
- 39. Schubert, M., Bleuler-Martinez, S., Butschi, A., Wälti, M. A., Egloff, P., Stutz, K., Yan, S., Collot, M., Mallet, J. M., Wilson, I. B., Hengartner, M. O., Aebi, M., Allain, F. H., and Künzler, M. (2012) Plasticity of the  $\beta$ -trefoil protein fold in the recognition and control of invertebrate predators and parasites by a fungal defence system. *PLoS Pathog.* **8,** e1002706
- 40. Tai, V. W., O'Reilly, M. K., and Imperiali, B. (2001) Substrate specificity of *N*-Acetylglucosaminyl(diphosphodolichol) *N*-acetylglucosaminyl transferase, a key enzyme in the dolichol pathway. *Bioorg. Med. Chem.* **9,** 1133–1140
- 41. Kelleher, D. J., and Gilmore, R. (2006) An evolving view of the eukaryotic oligosaccharyltransferase. *Glycobiology* **16,** 47R–62R
- 42. Calo, D., Kaminski, L., and Eichler, J. (2010) Protein glycosylation in Archaea. Sweet and extreme. *Glycobiology* **20,** 1065–1076
- 43. Abu-Qarn, M., Yurist-Doutsch, S., Giordano, A., Trauner, A., Morris, H. R., Hitchen, P., Medalia, O., Dell, A., and Eichler, J. (2007) *Haloferax volcanii* AglB and AglD are involved in *N*-glycosylation of the S-layer glycoprotein and proper assembly of the surface layer. *J. Mol. Biol.* **374,** 1224–1236

