Neisseria meningitidis (Nm) is a leading cause of bacterial meningitis and sepsis. Crucial virulence determinants of pathogenic Nm strains are the polysaccharide capsules that support invasion by hindering complement attack. In Nm W-135 and Nm Y the capsules are built from the repeating units (\(\rightarrow 6\)-\(\alpha\)-\(\text{Gal}-(1\rightarrow 4)-\alpha\)-Neu5Ac-(2\(\rightarrow\)\(\alpha\))\(\rightarrow\))\(\alpha\) and (\(\rightarrow 6\)-\(\alpha\)-\(\text{Gal}-(1\rightarrow 4)-\alpha\)-Neu5Ac-(2\(\rightarrow\)\(\alpha\))\(\rightarrow\))\(\alpha\), respectively. These unusual heteropolymers represent unique examples of a conjugation between sialic acid and hexosyl-sugars in a polymer chain. Moreover, despite the various catalytic strategies needed for sialic acid and hexose transfer, single enzymes (SiaD\(\text{W-135-Y}\)) have been identified to form these heteropolymers. Here we used SiaD\(\text{W-135}\) as a model system to delineate structure-function relationships. In size exclusion chromatography active SiaD\(\text{W-135}\) migrated as a monomer. Fold recognition programs suggested two separate glycosyltransferase domains, both containing a GT-B fold. Based on conserved motifs predicted folds could be classified as a hexosyl- and sialyltransferase. To analyze enzyme properties and interplay of the two identified glycosyltransferase domains, saturation transfer difference NMR and mutational studies were carried out. Simultaneous and independent binding of UDP-Gal and CMP-Sia was seen in the absence of an acceptor as well as when the catalytic cycle was allowed to proceed. Enzyme variants with only one functionality were generated by site-directed mutagenesis and shown to complement each other in trans when combined in an in vitro test system. Together the data strongly suggests that SiaD\(\text{W-135}\) has evolved by fusion of two independent ancestral genes encoding sialyl- and galactosyltransferase activity.

Bacterial meningitis remains a serious threat to global health, accounting for an estimated 170,000 annual deaths worldwide (WHO website). Despite the availability of potent antimicrobial agents, case fatality rates are high (10%) and survivors frequently suffer from sequelae such as neurologic disability or limb loss and deafness (1, 2). Although most cases of meningococcal disease are sporadic, outbreaks are frequently observed and large epidemics occur in the African meningitis belt (2, 3). The predominant serogroups are B and C followed by W-135 and Y (1). Serogroup W-135 meningococci caused an outbreak of disease among pilgrims in 2000 (4) and an epidemic in Burkina Faso in 2002 with over 13,000 cases and more than 1,400 fatalities (5, 6). Furthermore, an increase in serogroup Y disease was observed in the 1990s in North America (7).

Crucial virulence determinants of disease causing Nm species are their extracellular polysaccharide capsules (CPs) that are essential for meningococcal survival in human serum (8). Based on the chemical composition of these polysaccharides at least 12 different serogroups of Nm have been identified, but only six account for virtually all cases of disease. Of these serogroups B, C, Y, and W-135 express CPS structures that contain the acidic nonulosic sialic acid (Sia or Neu5Ac). Although the capsules expressed by serogroups B and C represent sialic acid homopolymers conjugated by \(\alpha(2,8)\)- and \(\alpha(2,9)\)-glicosidic linkages, respectively, the capsules of serogroups W-135 and Y represent heteropolymers built from disaccharide repeating units with sialic acid \(\alpha(2,6)\)-linked to galactose (W-135) or glu-
cose (Y). For example the repeating unit in NmW-135 is \((\rightarrow6)-\alpha-D-Gal-(1 \rightarrow 4)-\alpha-Neu5Ac-(2 \rightarrow 3)\) (9, 10). With sialic acid as an internal sugar conjugated by heteroglycans, the NmW-135 and NmY represent unique structures not found in any other organism so far.

The biosynthesis of these unusual glycoconjugates is achieved by the capsule polymerases, SiaDW-135 and SiaDy (11). The genes encoding these proteins have been cloned (11) and the function of the expressed recombinant proteins was initially characterized \textit{in vitro} and \textit{in vivo} (12). Overall the two proteins share 98% identity and differences are limited to the N-terminal domains. Already after cloning, the remarkable size of the genes had stimulated the hypothesis that they have evolved by fusion domains. Already after cloning, the remarkable size of the genes had stimulated the hypothesis that they have evolved by fusion domains. Already after cloning, the remarkable size of the genes had stimulated the hypothesis that they have evolved by fusion domains.

In the CAZy classification system for glycosyltransferases (cazy.org), SiaDW-135 and SiaDy were assigned to family GT-4, comprising retaining hexosyltransferases, many of bacterial origin (13–15). Allocation to this family was based on primary sequence similarities that the N-terminal domains exhibit with members of the group. A fundamental characteristic in this respect is the motif EX\(\times\)-E, which is highly conserved in retaining hexosyltransferases (16) and present also in the neisserial capsule polymerases SiaDW-135 and SiaDy (12). In contrast, no similarities to CAZy GT-families that contain sialyltransferases or polysialyltransferases were identified. Considering, however, that SiaDW-135 and SiaDy are bifunctional enzymes that not only use donor substrates of significant chemical diversity (this is true with respect to the nucleotide and sugar moieties) but in addition are likely to use different catalytic mechanisms (retaining for the hexose and inverting for the sialic acid transfer) the CAZy assignment must be reassessed.

Accordingly, the present work was focused on bringing new insight to the structure-function relationships of the bifunctional meningococcal capsule polymerases. Using SiaDW-135 as a model system, we have employed fold recognition programs to re-investigate the protein organization. Our refined structural model suggested a series of biochemically and biophysical studies that were carried out and consequently SiaDW-135 was identified as a monomeric protein that comprises two active sites: a galactosyltransferase and a monosialyltransferase. Using natural and artificial acceptors we show that the synthesis of the NmW-135 polymer depends on the alternate activity of these two catalytic domains.

**EXPERIMENTAL PROCEDURES**

**Expression Plasmids and Site-directed Mutagenesis**—The construction of the plasmid pET22b-Strep-NmW-135, to express SiaDW-135 as N-terminal Strep-II and C-terminal His6-tagged protein, was described earlier (12). Expression constructs lacking the N-terminal Strep-II tag were constructed by amplifying the siaDW-135 insert from pHC4 (11). PCR products obtained with the oligonucleotides KS422/KS273 were ligated between the NdeI and XhoI sites of pET22b (Novagen). Mutagenesis of siaDW-135 was performed using the QuiChange site-directed mutagenesis kit (Stratagene) according to the manufacturer’s instructions. The plasmid pET22b-Strep-NmW-135 (12) was used as template in all mutagenesis reactions. Primer sequences are given in Table 1 with the matched base triplets underlined. Mutations and corresponding primer pairs are E307A: KS350/KS351; H934A: KS364/KS365; P935A: KS366/KS367; S972A: KS370/KS371; T973A: KS372/KS373. The mutated siaDW-135 were subsequently subcloned into the BamHI and NotI sites of pET22b-Strep (17) and the sequence identity of all constructs was confirmed by sequencing.

**Expression and Purification of Recombinant SiaDW-135**—Wild-type and mutant variants of SiaDW-135 carrying a C-terminal His\(_6\)-tag (the wild-type protein was produced in two forms with C-terminal His\(_6\)-tag and as double-tagged protein) were expressed in Escherichia coli BL21(DE3) in the presence of 100 \(\mu\)g/ml of carbenicillin. Bacteria were cultivated in autoinducing ZYM-5052 medium (18) and grown for 78 h at 15 °C. Cells were harvested (4000 \(\times\) g; 30 min; 4 °C), re-suspended in binding buffer (50 mM Tris/HCl, pH 8.0, 300 mM NaCl, 40 mg/ml of bestatin, 1 \(\mu\)g/ml of pepstatin, 1 \(\mu\)M PMSF), and disrupted by sonication (BRANSON SONIFIER® W-450D; amplitude 50%; 8 \(\times\) 30 s). Recombinant proteins were bound to HisTrap affinity columns (GE Healthcare) using binding buffer (50 mM Tris/HCl, pH 8.0, 300 mM NaCl, 40 mg/ml of bestatin, 1 \(\mu\)g/ml of pepstatin, 1 mM PMSF), and disruption by sonication (BRANSON SONIFIER® W-450D; amplitude 50%; 8 \(\times\) 30 s) was used to elute the bound proteins. Recombinant proteins were analysed by SDS-PAGE, and purified proteins were used in biochemical assays.
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HCl, pH 8.0, 300 mM NaCl) and eluted in a two-step gradient (50 and 150 mM imidazole in binding buffer). Fractions containing SiaD<sub>W-135</sub> were pooled and applied to a HiLoad 16/60 Superdex 200 pg column (GE Healthcare) for further purification by size exclusion chromatography. Proteins were eluted (50 mM Tris/HCl, pH 8.0, 300 mM NaCl, 2 mM DTT) at a flow rate of 0.5 ml/min and subsequently concentrated to 2 mg/ml using Amicon Ultra centrifugal devices (Millipore; 50 kDa MWCO). The purified proteins were flash-frozen in liquid nitrogen and stored at −80 °C.

Activity Assays—Radioactive incorporation assays were performed as described (12, 19). Briefly, purified recombinant proteins (7.5 µg) were assayed in reaction buffer (20 mM Tris/HCl, pH 8.0, 10 mM MgCl<sub>2</sub>, 1 mM DTT) in the presence of 1 mM radiocarbon labeled CMP-[<sup>14</sup>C]Neu5Ac (0.13 mCi/mmol, GE Healthcare) and 2 mM UDP-Gal (Sigma). Additionally, 2 mM of an artificial oligosaccharide acceptor (i.e. sialylactose, Sigma; Neu5Ac, and the dimer and trimmer of α(2,8)-Neu5Ac, Nacalai Tesque) or 0.4 mg/ml of hydrolyzed W-135 CPS were included in a total volume of 25 µl. Samples were incubated at room temperature and enzymatic activity was determined at appropriate time intervals by mixing 5-µl aliquots of the reaction solution with 5 µl of chilled ethanol (96%). Samples were spotted on Whatman 3MM CHR paper and the chromatographically immobile <sup>14</sup>C-labeled reaction products were quantified by scintillation counting following descending paper chromatography as described (20).

To visualize the size distribution of the synthesized W-135 polysaccharides, purified SiaD<sub>W-135</sub> (5–15 µg) was assayed in reaction buffer (20 mM Tris/HCl, pH 8.0, 10 mM MgCl<sub>2</sub>, 1 mM DTT) in the presence of 1 mM CMP-Neu5Ac (GERBU), 2 mM UDP-Gal, and hydrolyzed W-135 CPS (0.16 µg/µl) as the oligosaccharide acceptor structure in a total volume of 37.5 µl. Samples were incubated at room temperature and reactions were stopped at appropriate time intervals by addition of 1 mM sucrose. Synthesized products were separated by PAGE (25%) and subsequently detected using a combined Alcian blue/silver staining procedure as described (20).

For the in vivo testing of mutants strain Nm-WUE171 was used and the plasmids were generated on the basis of pIP2 as previously described (12). Briefly, the HpaI/SphI fragment of wild-type siaD<sub>W-135</sub> comprising position 19–2943 was replaced by the respective fragments of the pET22b-Strep-Nm-W-135 mutants. Correct integration of the mutated DNA into the meningococcal chromosome was confirmed by PCR and subsequent DNA sequencing. A siaD knock-out mutant designated WUE2661, used as a negative control in this study, resulted from the transformation of strain WUE171 with pHCl4siaD::TnMax5 (21). Phenotypic analysis of capsule expression was performed as described recently with mAb1509 specific to serogroup W-135 and PorA antibody P1.10 (12).

STD NMR Experiments—All NMR experiments were performed on a Bruker 600 Ultrashield<sup>TM</sup> at 280 K, equipped with a standard triple resonance CryoProbe, in deuterated 20 mM Tris buffer, supplemented with 20 mM MgCl<sub>2</sub> (D<sub>2</sub>O), pH 8.4. 200 µl of sample in Shigemi tubes (Shigemi Co.) were used for all acquisitions. STD spectra were acquired using 150 µg of SiaD<sub>W-135</sub> and saturation time of 2 s at an on-resonance frequency of −1 ppm and off-resonance of 33 ppm. A WATERGATE sequence was used to suppress the residual HDO signal. The on- and off-resonance spectra free induction decays were stored and processed separately and the subtraction of the on-resonance and off-resonance spectrum resulted in the STD NMR spectrum. All STD NMR effects were calculated according to the formula 

\[
A_{\text{STD}} = \frac{I_0 \times I_{\text{sat}}}{I_0 - I_{\text{STD}}} = I_{\text{STD}}/I_0
\]

and the strongest STD NMR signal was set to 100%.

RESULTS

Fold Recognition Tools Predict Two GT-B-like Glycosyltransferase Domains in SiaD<sub>W-135</sub>—The homology of the N-terminal domain with hexosyltransferases of CAZy family GT-4, led to the grouping of SiaD<sub>W-135</sub> into this GT class (11, 12). Considering, however, that SiaD<sub>W-135</sub> catalyzes the formation of a heteropolymer (Fig. 1D) by use of two significantly different donor substrates (UDP-Gal and CMP-Sia) and two different catalytic mechanisms (galactosyltransferases are retaining and sialyltransferases inverting enzymes), the organization of SiaD<sub>W-135</sub> was reinvestigated with the help of the structure prediction tool Phyre (22). Two GT-B folds were identified, the first spanning N-terminal residues 1–399 and the second residues 763–1037 in the C-terminal domain (Fig. 1, A–C). Although primary sequence identities to the respective template structures were low (≤13% and ≤10%, respectively), high confidence matches (100% each) were reported and indicated an accurate prediction of the overall fold (22). Closer inspection of the predicted models revealed in addition that highly conserved amino acid clusters known to be part of the active sites in either hexosyl- (EX,E motif (16)) or sialyltransferases (HP- and S-motif (23–25)) localize along the deep active site crevices that separate the two β/α/β Rossmann-like domains in GT-B glycosyltransferases (Fig. 1, B and C). Subsequent experiments therefore addressed the question if SiaD<sub>W-135</sub> represents a chimera with two enzymatic functions.

The Purified Recombinant SiaD<sub>W-135</sub> Is Monomeric and an Active Bifunctional Polymerase—To obtain a suitable enzyme source for the planned biochemical and biophysical studies, SiaD<sub>W-135</sub> carrying a C-terminal His<sub>6</sub> tag was purified in two steps (IMAC and SEC) (Fig. 2A). The protein eluted as a monomer with an apparent molecular mass of 120 kDa (calculated molecular mass 121.137 kDa) from the calibrated SEC column (Fig. 2B). Activity of the purified enzyme was confirmed in a radiocarbon incorporation assay using CPS isolated from Nm-W-135 (for reference see CPS in Fig. 3, A and B) as acceptor. As already described for the partially purified SiaD<sub>W-135</sub> (12) significant incorporation was only detected when CMP-[<sup>14</sup>C]Neu5Ac was combined with the second donor sugar UDP-Gal (data not shown).

To unequivocally demonstrate that the recombinant SiaD<sub>W-135</sub> catalyzes the synthesis of W-135 polymers under the in vitro conditions, the isolated CPS (CPS in Fig. 3, A and B) was partially hydrolyzed to yield oligomers (CPS(h) in Fig. 3A) and the oligomers were used to prime the polymerase reaction in the presence of “cold” substrates. Long chain products obtained in this reaction were displayed in high percentage PAGE (Fig. 3A) and, following a time course experiment, in 10% SDS-
PAGE (Fig. 3B). To visualize polymerization products and enzyme the 10% SDS-PAGE was simultaneously stained with Alcian blue and silver. Oligomeric acceptors were not retained in the low percentage gel, but long chain polymers were detectable after 10 min. Moreover, this analysis demonstrated that the protein band corresponding to purified SiaD\textsubscript{W-135} remained unaltered during the reaction.

**Mutagenesis Studies Indicate Two Distinct Active Sites in SiaD\textsubscript{W-135}**—Site-specific mutations were generated to interrogate the functional nature of the identified GT-B like domains in SiaD\textsubscript{W-135}. Targets in this study were the highly conserved EX\textsubscript{7}E motif that represents a characteristic of the nucleotide recognition domain in retaining GTs (16) and the recently identified bacterial “sialyl motifs” HP and SS/T (23–25) presumed to be involved in substrate binding (26) (see Fig. 1). To inactivate these motifs Glu-307 (the first residue in the EX\textsubscript{7}E motif), His-934 (HP motif) as well as Ser-972 and Thr-973 (SS/T motif) were individually exchanged to alanine.
Obtained mutants were tested in vivo and, with CPS(h) as acceptor, also in vitro (Fig. 4). For the in vivo testing capsule expression of a NmW-135 wild-type strain and the respective SiaDW-135 mutants generated via homologous recombination were quantified in a well established whole cell ELISA (12, 27). In Fig. 4A results are shown as relative values with the starting strain (NmW-135171-wt) set to 100%. The capsule-deficient strain WUE 2661-∆CPS served as a negative control.

As this difference could not be explained by reduced protein expression, stability, or purity (see Fig. 4B), it is likely that integration of mutant proteins into the capsule synthesis complex in vivo creates conditions that favor functionality (12). Importantly, however, if the purified mutants were brought together in one reaction mixture activity was restored (>70% of the wild-type). Because each mutant still contains one functional domain (sialyltransferase in the case of mutant E307A and galactosyltransferase in the case of S972A), this complementation in trans strongly supported the hypothesis that SiaDW-135 represents a chimeric enzyme.
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Sia\textsubscript{D\textsubscript{W-135}} Elongates \(\alpha(2,8)\)-Linked Sialic Acid Trimers—Because defined oligosaccharides consisting of \((-\rightarrow 6)-\alpha-D-Gal-(1\rightarrow 4)-\alpha-D-Neu5Ac-(2\rightarrow)\) were not readily available, further analysis of the catalytic functions of Sia\textsubscript{D\textsubscript{W-135}} was preceded by the search for a low molecular weight acceptor molecule. The compounds tested are indicated in Fig. 5A. Neither the free monosaccharides (Sia and Gal) nor lactose were extended by the enzyme, whereas a slight but reproducible activity was seen with DP2 (Neu5Ac-\(\alpha(2\rightarrow 8)\)-Neu5Ac) and 2,3-sialyllactose (Neu5Ac-\(\alpha(2\rightarrow 3)\)-D-Gal-\(\beta(1\rightarrow 4)\)-D-Glc). However, if DP3 (trimer of \(\alpha(2,8)\)-linked Neu5Ac) was used as acceptor, the activity escalated. To better understand this unexpected finding, the binding modes of DP1, DP2, and DP3 were investigated by STD NMR. This method records saturation transfer by spin diffusion from the target protein to binding ligands. Ligand protons making close contacts to the protein surface receive higher saturation resulting in high STD NMR signal intensities. Substrate binding profiles can thus be determined based on the relative saturation of ligand epitopes (28).

In Fig. 6 the NMR spectra of the compounds are shown. The bottom panels (Fig. 6, a, g, and m) display the \(^1\)H NMR spectra of the soluble compounds. The respective STD NMR spectra obtained for ligands in complex with the recombinant Sia\textsubscript{D\textsubscript{W-135}} are shown in Fig. 6, b, h, and n. From the combined spectra it is immediately clear that all three ligands (DP1, DP2, and DP3) bind to Sia\textsubscript{D\textsubscript{W-135}}. A closer inspection of the data, however, revealed that the \(\alpha\)-anomer of the free sialic acid was bound with much higher affinity than the \(\beta\)-anomer. This effect is easiest seen for the H3eq protons. Although in solution \(\sim 5\%\) of DP1 attains \(\alpha\)- (at 2.45 ppm) and \(\sim 95\%\) \(\beta\)-configuration (at 1.95 ppm), signals of comparable intensities were observed if the protein was present (Fig. 6a). A similar effect could be seen for the singlet of the N-acetamido group at 1.76 (\(\beta\)-anomer) and 1.73 ppm (\(\alpha\)-anomer) (Fig. 6, e and j). The STD NMR data thus strongly suggest that free Neu5Ac (DP1) is bound to the nucleotide sugar binding pocket.

The spectra obtained with DP2 and DP3 revealed that the nonreducing sugar received more saturation than the middle (Sia2 in DP3) and reducing sugar (Sia1). This effect again was reflected by both the H3eq and the N-acetamido group protons (see Fig. 6, i–l for DP2 and o and p for DP3).

Although the data collected so far do not suffice to draw a detailed picture of the acceptor binding site in Sia\textsubscript{D\textsubscript{W-135}}, the observation that DP3 is an efficient acceptor and fully accommodated in the binding pocket supports the notion that only DP3 has the capacity to make contacts to all sites involved in binding and presenting the acceptor to the catalytic center.

Moreover, with this defined acceptor at hand we re-evaluated the hypothesis of consecutive polymer building by the two GT-B folds. Therefore, the wild-type Sia\textsubscript{D\textsubscript{W-135}} as well as mutants E307A and S972A were analyzed in vitro with DP3 as acceptor. With UDP-\(^{14}\)C\textsubscript{Gal} and cold CMP-Neu5Ac long

\(^{14}\)C\textsubscript{Gal} onto DP3 (lane 2) by S972A (comprises an active galactosyltransferase domain) was observed, whereas all other reactions (lanes 3–5) remained negative. These data clearly demonstrate that DP3 is selectively recognized by the galactosyltransferase domain and that formation of long CPS requires the iterative activity between galactosyl- and sialyltransferase domain.

FIGURE 5. A, the trimer of \(\alpha(2,8)\)-linked sialic acid residues (DP3) is efficiently elongated by Sia\textsubscript{D\textsubscript{W-135}}. As defined W-135 CPS oligomers were not readily available, the compounds as listed were analyzed for their capacity to prime the Sia\textsubscript{D\textsubscript{W-135}} reaction in the radioactive incorporation assay. In this experiment DP3 was found to be efficiently elongated by Sia\textsubscript{D\textsubscript{W-135}}, whereas DP2 and 2,3-sialyllactose (Sia-Lac) were only weak acceptors. The monosaccharides Gal and Sia as well as the disaccharide lactose (Lac) were not used by the enzyme. Specific activities were calculated from three independent experiments. B, asking if DP3 is an acceptor for both glycosyltransferase domains present in Sia\textsubscript{D\textsubscript{W-135}}, the transfer reactions catalyzed by the single mutants E307A and S972A were analyzed in comparison to the wild-type enzyme. Both nucleotide sugars were present in these reactions, with only one radioactive labeled as indicated. Long polymer chains were synthesized in reaction mixtures with wild-type Sia\textsubscript{D\textsubscript{W-135}} (lane 1) and both single mutants (lane 6). In contrast, a single radioactive spot, representing the transfer of one
radioactively labeled polymers (CPS) were synthesized in the reactions with wild-type enzyme (lane 1) and if the two single mutants E307A and S972A were present in one reaction mixture (lane 6). A single transfer of \([^{14}C]\text{Gal}\) onto the acceptor DP3 (lane 2) was catalyzed by mutant S972A, harboring an intact galactosyltransferase domain, whereas mutant E307A, harboring a defective galactosyltransferase, did not produce a radioactive product (lane 4). Importantly, with CMP-\([^{14}C]\text{Neu5Ac}\) and cold UDP-Gal none of the mutants produced a radioactive signal (lanes 3 and 5), thus clearly evidencing that Gal transfer to DP3 is prerequisite to generate the acceptor recognized by the sialyltransferase domain. The data provided in this paragraph prove the iterative activity of the two GT-B folds in SiaDW-135.

**Independent Acceptor and Nucleotide-Sugar Binding to SiaDW-135**—To further analyze the concept of two glycosyltransferase domains in one polypeptide chain the binding of the nucleotide sugars (UDP-Gal and CMP-Neu5Ac) to the protein was investigated. \(^1\text{H}\) NMR spectra of UDP-Gal and CMP-Neu5Ac (Fig. 7, a and c) and STD NMR spectra of UDP-Gal and CMP-Neu5Ac in the presence of SiaDW-135 (Fig. 7, b and d) were acquired and demonstrated in line with earlier data (29) the relevance of H1 Rib in the interactions of sugar nucleotides with proteins. H1 Rib as well as H5 in the nucleotide base received high saturation in both UDP-Gal and CMP-Neu5Ac. Of note, H5 Ura and H1 Rib STD NMR signals were overlapping in UDP-Gal and could not be separated. Confirming the specificity of SiaDW-135 for galactose H4 Gal received 57% saturation and, remarkably, also the anomeric proton in Gal received considerable saturation (62%). In the case of CMP-Neu5Ac, the N-acetamido group (1.82 ppm) was found to be strongly involved in the recognition event by the enzyme. Presence of equimolar concentrations of UDP-Gal did not change the STD NMR signals obtained for CMP-Neu5Ac (Fig. 8), thus excluding co-location of the nucleotide-sugar binding sites. This fact is particularly highlighted by the methyl protons of the N-acetamido group of CMP-Neu5Ac (Fig. 8c).

Next we analyzed if addition of the acceptor DP3 alters nucleotide sugar binding profiles. As previous experiments (see Fig. 5B) had shown that DP3 is not elongated by the sialyltransferase domain, data were recorded in the absence of UDP-Gal to prevent the start of the catalytic reaction. In Fig. 9 three STD...
To determine epitopes in the donor sugars CMP-Neu5Ac (a and b) and UDP-Gal (c and d) that make contact to the protein, an STD NMR study was performed. In line with previous data this experiment demonstrated the importance of H1 Rib for nucleotide sugar binding. This position together with H5 in the nucleotide base received the highest saturation in both UDP-Gal and CMP-Neu5Ac. In accordance with the specificity of SiaDW-135 for galactose H4 Gal received 57% saturation and, remarkably, also the opposite site of the pyranose ring, the anomic proton H1 Gal, received considerable saturation (62%). For experimental details, see Fig. 6.

To find out if binding of one nucleotide sugar impacts binding of the second, the STD NMR spectrum of CMP-Neu5Ac in complex with SiaDW-135 was recorded in the absence (a) and presence (b) of an equimolar concentration of UDP-Gal. No change in the position or intensity of any signal was seen, thus indicating independent binding of the nucleotide sugars. To highlight this fact, an overlay is shown for the intense signal produced by the tightly bound N-acetamido group (c). For experimental details, see Fig. 6.
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FIGURE 9. STD NMR spectra of SiaD<sub>W-135</sub> in complex with CMP-Neu5Ac and the acceptor DP3. To evaluate if binding of the acceptor DP3 influences the binding of CMP-Neu5Ac, three STD NMR spectra were overlaid. In black and green the spectra obtained for SiaD<sub>W-135</sub> in complex with CMP-Neu5Ac and DP3, respectively, are shown. The spectrum obtained when both substrates were present in equimolar concentrations is shown in red. The absolute STD NMR intensities revealed only a minor (~15%) reduction in the saturation transfer to the H3eq proton of CMP-Neu5Ac when DP3 was present. Similarly, a reduction of ~20% was seen for the H3eq protons of DP3 under these conditions. The faintness of the differences observed in the presence of the second substrate argued for independent binding sites. For experimental details, see Fig. 6.

NMR spectra are superimposed: 1) SiaD<sub>W-135</sub> in complex with CMP-Neu5Ac (black), 2) SiaD<sub>W-135</sub> in complex with DP3 (green), and 3) SiaD<sub>W-135</sub> in complex with equimolar amounts of DP3 and CMP-Neu5Ac (red). Although the simultaneous presence of acceptor (DP3) and nucleotide-sugar (CMP-Neu5Ac) induced a slight reduction in the absolute STD NMR intensities (~15% for the H3eq proton of CMP-Neu5Ac and ~20% for all H3eq protons in DP3) the STD NMR signal intensities remained predominately unchanged and suggested that DP3 binds to a distinct domain.

The above result was confirmed when the catalytic reaction of SiaD<sub>W-135</sub> was allowed to proceed in the presence of equimolar concentrations of UDP-Gal, CMP-Neu5Ac, and DP3 (Fig. 10a). After 2.5 h acquisition time, CMP-Neu5Ac was almost completely consumed and a product peak representing the W-135 polymer became visible by 1H NMR (Fig. 10c). The STD NMR spectrum recorded for the active enzyme in the presence of all substrates clearly revealed independent binding sites for all three substrates (Fig. 10b).

DISCUSSION

In this study we show that bifunctionality of SiaD<sub>W-135</sub> is the result of the presence of two catalytic domains in one polypeptide chain. Based on the conserved structural motifs catalytic functions could be predicted for the two GT-B folds identified by bioinformatics tools. Rational mutations confirmed the predictions and proved galactosyl- and sialyltransferase activity for the N and C terminally located subdomains, respectively. Moreover, by combining point mutated proteins, in which one of the putative catalytic functions was disturbed, 70% of wild-type activity was restored. In size exclusion chromatography the active recombinant enzyme migrated with an apparent molecular mass of 120 kDa, indicating that the monomer is the functional unit. In STD NMR studies, the two GT-B folds were shown to bind their sugar-nucleotide substrates (CMP-Neu5Ac and UDP-Gal) independently and simultaneously. With DP3, a trimer of α(2,8)-linked sialic acids, a suitable minimal primer for the polymerase reaction has been identified and used in in vitro activity assays as well as in STD NMR studies to demonstrate that chain elongation needs the successive activity of the hexosyl- and sialytransferase domains.

Glycosyltransfeases (GTs) catalyze the synthesis of complex products (e.g. polymers) in the absence of templates. Product control is an enzyme-inherent feature installed by the specificity of the GTs for their substrates (nucleotide-sugar and acceptor) and the type of catalyzed glycosidic linkage. In 1968 this observation led to the formulation of the paradigm of “one enzyme-one linkage” (30). Although the identification of GTs that in the absence of a second catalytic center conjugate one sugar with two types of glycosidic bonds (31–33) challenged the paradigm, SiaD<sub>W-135</sub> and SiaD<sub>Y</sub> remain unique because these enzymes conjugate sugars of very different chemical nature (the disparity is even higher in the GT substrates CMP-Neu5Ac and UDP-Gal/Glc) and use different catalytic mechanisms to form the glycosidic bonds (inversion of the anomeric center in the case of sialic acid and retention in the case of the hexoses). Not surprising thus that these large enzymes were suggested to represent fusion products of ancestral genes (11, 12).

The N-terminal GT-B fold in SiaD<sub>W-135</sub> and SiaD<sub>Y</sub> exhibits sequence similarity with retaining hexosyltransferases of CAZY family GT-4. Common to members of this group is the EX,E motif, a part of the so called nucleotide recognition domain 1α (16, 34). Mutation of the first conserved glutamic acid residue in the EX,E motif has been demonstrated to inactivate retaining GTs (35–38). In line with these earlier data we show here that the position (Glu-307) is essential also in SiaD<sub>W-135</sub>. The replacement E307A abolished activity in vitro and in vivo. Of importance, position four within the EX,E motif (aa 310; proline in NmA<sub>W-135</sub> and glycine in Nm<sub>Y</sub>) was previously shown to determine UDP-sugar specificity in these bifunctional enzymes (12). The STD NMR data obtained in the current study show that H4 Gal receives high saturation (Fig. 7). The specificity for galactose in SiaD<sub>W-135</sub> can thus be explained by a tight contact formed between the C-4 hydroxyl group of Gal and proline 310 in the sugar nucleotide binding pocket.

For the second GT-B fold, shown in this study to comprise an α(2,6)-monosialyltransferase, no similarity to members in any of the CAZY families could be identified. Nevertheless, the used fold recognition tool properly located the previously identified conserved motifs HP and SS/T (23–25) along the active site cleft that separates the two Rossmann subdomains (see Fig. 1C). The HP motif is broadly conserved in bacterial sialytransferases and was shown to be involved in CMP binding and/or enzyme catalysis (39). The more C terminally located S-motif contains a highly conserved serine and a second less conserved serine or threonine residue. In CAZY family GT-80 this motif was shown to coordinate the CMP-phosphate group in the intermediate state carbennium ion (24, 26, 40). In accordance with this essential functionality, point mutation of the crucial Ser-972 in SiaD<sub>W-135</sub> fully inactivated the enzyme.

Still searching for homologues of the SiaD<sub>W-135/Y</sub> sialytransferase domain we performed an extended BLAST analysis and...
identified weak homologies to several bacterial open reading frames encoding proteins of unknown function. All identified open reading frames contain the HP and S motifs and, as schematically shown in Fig. 11, occur in the company of two genes encoding enzymes that locate immediately upstream of the sialyltransferase activity in the sialylation pathway. The accompanying enzymes are the sialic acid synthase (41) and the CMP-sialic acid synthetase (42). Because functionally linked genes are often organized in operons in bacteria, the identified ORFs are likely to encode sialyltransferases. Surprisingly, the multifunctional sialyltransferase \( PmST1 \) from \( Pasteurella multocida \) was not identified in this initial screen although the enzyme has been demonstrated to attain a GT-B fold (43).

Besides identification of bifunctionality in single domain GTs the presence of two independent binding sites has also been shown. Examples are the hyaluronan synthase from \( P. multocida \) (44) and the \( E. coli \) K4 chondroitin polymerase (K4CP) (45). However, in marked difference to SiaD\(_{W-135/Y}\) these polymerases are hexosyltransferases only and belong to the GT-A structural superfamily.

Binding analyses carried out by STD NMR provided conclusive evidence that the nucleotide sugar binding pockets in SiaD\(_{W-135}\) are independent from each other (Fig. 8). In both domains the nucleotide moieties (CMP and UDP) receive highest saturation, whereas the sugar moieties bind weaker. This binding mode has been previously described and was suggested to facilitate the transfer process (29, 46, 47). A particular feature in CMP-Neu5Ac binding is the tight contact that the \( N \)-acetamido group of sialic acid makes to the protein. This tight contact may be necessary to prevent the negatively charged acetamido group from interfering with the transfer process.

Unexpectedly we found DP3, but neither DP2 nor 2,3-sialyllactose, an efficient acceptor to start the polymerization reaction in SiaD\(_{W-135}\). Although additional experimental work is needed to understand the primer quality of DP3, the current
STD NMR data demonstrate that DP3 like DP2 is entirely bound to the acceptor site of the enzyme. Because DP2 is not an efficient acceptor, we speculate that only DP3 has the necessary geometry (e.g., length and topology of functional groups) to make contacts to all sites involved in the binding and presentation of the acceptor to the catalytic center.

The nonreducing end sialic acid of DP3 (the sugar where transfer occurs) received the highest saturation when bound to SiaDW-135. In contrast, if DP3 was complexed with the recombinant NmB capsule polymerase (a sialyltransferase produces a homopolymer of \( \beta \)-linked sialic acids), the saturation was found to be lowest for the nonreducing end sugar (47). Understanding the molecular reasons for these differences needs additional work, but it should be mentioned at this point that DP3 in the case of SiaDW-135 is the acceptor of the galactosyltransferase domain, which may have different stereochemical requirements compared with sialyltransferases.

Not clear at this moment is if the two GT-domains in SiaDW-135 comprise individual acceptor binding sites or share a common one. Although the STD NMR experiments carried out in the presence of two (CMP-Neu5Ac and DP3) or even three (CMP-Neu5Ac, UDP-Gal and DP3) substrates demonstrate an independent binding of acceptor and donor substrates, the way the growing polymer is bound to the enzyme remains an open question. Initial trials made to separately express the hexosyl- and sialyltransferase domains failed to produce active proteins. Combined with the knowledge that full-length proteins with only one functional domain were able to complement each other in trans (Fig. 4C) this latter finding suggests that the large linker domain that connects the GT-B fold in SiaDW-135 (see Fig. 1A) may have a function in forming a proper acceptor binding pocket. Our current work focuses at answering this question. With the current study we identified the bifunctional SiaDW-135 as a chimeric enzyme comprising two independent catalytic domains that work in succession to produce the highly unusual W-135 capsular polymer.

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