Chemical precision glyco-mutagenesis by glycosyltransferase engineering in living cells

Benjamin Schumann¹£, Stacy A. Malaker¹$, Simon P. Wisnovsky¹$, Marjoke F. Debets¹±, Anthony J. Agbay¹, Daniel Fernandez³, Lauren J. S. Wagner⁴, Liang Lin⁵, Junwon Choi¹#, Douglas M. Fox¹, Jessie Peh¹, Melissa A. Gray¹, Kayvon Pedram¹, Jennifer J. Kohler⁶, Milan Mrksich⁵, Carolyn R. Bertozzi¹,²*¹

¹Department of Chemistry, Stanford University, 380 Roth Way, Stanford, CA94305, USA.
²Howard Hughes Medical Institute, 380 Roth Way, Stanford, CA94305, USA.
³Stanford ChEM-H Macromolecular Structure Knowledge Center, 443 Via Ortega, Stanford, CA.
⁴Department of Chemistry, University of California, Berkeley, CA 94720, USA.
⁵Department of Biomedical Engineering, Northwestern University, 2145 Sheridan Rd, Tech B492, Evanston, IL-60208, USA.
⁶Department of Biochemistry, University of Texas Southwestern Medical Center, Dallas, TX 75390-9038, USA.

£Current address: The Francis Crick Institute, 1 Midland Rd, London NW1 1AT, UK, and Imperial College London, 80 Wood Lane, London W12 0BZ, UK.
±Current address: Lilly Research Laboratories, Eli Lilly and Company, Indianapolis, IN 46285, USA
#Current address: Korea Institute of Science and Technology, Hwarangro 14-gil 5, Seongbuk-gu, Seoul, 02792, Republic of Korea.
$These authors contributed equally.
*Corresponding author: bertozzi@stanford.edu
Abstract: Studying posttranslational modifications classically relies on experimental strategies that oversimplify the complex biosynthetic machineries of living cells. Protein glycosylation contributes to essential biological processes, but correlating glycan structure, underlying protein and disease-relevant biosynthetic regulation is currently elusive. Here, we engineer living cells to tag glycans with editable chemical functionalities while providing information on biosynthesis, physiological context and glycan fine structure. We introduce a non-natural substrate biosynthetic pathway and use engineered glycosyltransferases to incorporate chemically tagged sugars into the cell surface glycome of the living cell. We apply the strategy to a particularly redundant yet disease-relevant human glycosyltransferase family, the polypeptide N-acetylgalactosaminyl transferases. This approach bestows a gain-of-function modification on cells where the products of individual glycosyltransferases can be selectively characterized or manipulated at will.

Main Text: Posttranslational modifications expand the structural diversity of proteins, but are notoriously difficult to study in living systems. Most modifications are refractory to direct genetic manipulation and require reductionist strategies such as in vitro systems or simplified cells. Glycans are the prime example for this: the human glycome is constructed by the combinatorial activity of more than 250 glycosyltransferases (GTs) with both hierarchical and competing activities. On the cell surface, glycans play a central role in modulating signal transduction, cell-cell interactions and biophysical properties of the plasma membrane (1, 2). Understanding and manipulating these processes should thus be possible by adorning specific glycans with editable chemical functionalities. In such a synthetic biology approach, individual GTs could be engineered to accommodate a chemical functionality that is not found in native substrates and not accommodated by other GTs. This “bump-and-hole” tactic has been applied to a range of enzymes by us and others (3-9), but application in the living cell is a substantial technical challenge: the
nucleotide-based substrate analog must be delivered across the plasma membrane and the cell must stably express the correctly localized and folded mutant enzyme. Bump-and-hole engineering is particularly attractive to deconvolve GT families of multiple homologous isoenzymes: the complex interplay of these isoenzymes in the secretory pathway cannot be probed in sufficient detail in in vitro assays.

One of the largest GT families in the human genome is the polypeptide N-acetylgalacosaminyl (GalNAc) transferase family (GalNAc-T-1…T-20). Transferring GalNAc to Ser/Thr side chains, GalNAcTs initiate abundant O-linked glycosylation in the secretory pathway (Fig. 1A) (10, 11). O-GalNAc glycosylation can differ based on cell type or activation stage, and clear disease phenotypes are associated with glycan aberration (12, 13). Unsurprisingly, GalNAcT expression is often associated with tumorigenesis, sometimes with opposite effects on different types of cancer (14, 15). However, the absence of a glycosylation consensus sequence and the variability of glycan elaboration render O-GalNAc glycans challenging to study by mass spectrometry (MS) based glycoproteomics. Thus, the protein substrate specificity of each isoenzyme is poorly studied and largely built on data inferred from synthetic peptide libraries (16). So-called SimpleCells that do not elaborate O-GalNAc glycans make glycoproteins that are easier to enrich and study by MS (17-20). Knock-outs of single GalNAcTs in the SimpleCell background have been profiled by glycosproteomics (18, 21). Similarly, glycopeptides from non-SimpleCells with titratable GalNAcT knock-in have been enzymatically simplified to allow for enrichment and MS (22). These studies have revealed comprehensive glycoproteomics datasets, but suffer from loss of glycan elaboration and laborious genome engineering required for targeted knock-in. Further, the activity of GalNAcT isoenzymes is both redundant and competitive, such that compensation and/or
shift of glycosylation sites occurs upon knock-out (21, 22). A gain-of-function strategy to visualize the products of a particular GT on the living cell is currently elusive.

![Diagram](image_url)

**Fig. 1**: GalNAcT bump-and-hole engineering. A, GalNAcTs initiate O-GalNAc glycosylation. Transfer of GalNAc to a Ser or Thr side chain is followed by downstream glycan elongation. B, the principle of bump-and-hole engineering. Engineered double mutant (DM) GalNAcTs are paired with UDP-GalNAc analogs 1-4 to chemically tag GalNAcT substrates that can be monitored by click chemistry. C, Overview of the steps taken in this study toward GalNAcT bump-and-hole engineering in the living cell. PG = Protecting group.

Here, we equipped living cells with the ability to tag the protein substrates of individual GalNAcTs with chemical, editable functionalities. We made use of the fact that a double mutation (‘‘DM’’,
I253A/L310A double mutant in T-2, fig. 1B) re-programmed GalNAcTs to accept alkyne-containing uridine diphosphate (UDP)-GalNAc analogs such as compounds 1-4 instead of the native substrate UDP-GalNAc (Fig. 1B) (8). We endowed living cells with the capacity to biosynthesize a UDP-GalNAc analog that was complementary to engineered DM-GalNAcTs. Further, we showed that DM-GalNAcTs emulate wild type (WT) GalNAcTs with regard to structure, localization and protein substrate specificity. This approach bestowed a bioorthogonal tag (23-27) on the protein substrates of distinct GalNAcT isoenzymes while conserving the complexity of glycan elaboration in the secretory pathway (Fig. 1C). Precision glycome engineering has widespread applications in biomarker discovery, GT profiling and targeted cell surface engineering.

**Structural basis for GalNAcT engineering**

As bump-and-hole engineering of a GT family in living cells has no precedent, we first set out to understand the structural implications of this approach. All GalNAcTs are type II transmembrane proteins with luminal GT and lectin domains connected through a flexible linker (10, 11, 28). We crystallized the luminal part of DM-T-2 in complex with the native ligands Mn^{2+}, UDP, and the substrate peptide EA2 (PTTDSTTPAPPTTK) at 1.8 Å resolution. Comparison of DM-T-2 and WT-T-2 (PDB 2FFU) revealed complete conservation of both the three-dimensional enzyme architecture and bound ligand structure (Fig. 2A, fig. S1 and Table S1) (29). In DM-T-2, the interdomain linker adopts an extended conformation previously found in the catalytically active WT enzyme (30, 31). The mutant A253 and A310 side chains in the DM enzyme are congruent with the WT side chains I253 and L310, respectively (Fig. 2B). Two glycine residues, G308 and G309, are slightly shifted by 1.2 Å and 1.7 Å (Cα distances), respectively, likely to account for the
changes elsewhere in the active site. DM-T-2 thus retains the native structural properties of the WT-GalNAcT enzyme.

**Fig. 2**: Bump-and-hole engineering conserves folding and substrate binding of GalNAcT-2. A, crystal structure of DM-T-2 at 1.8 Å superposed with WT-T-2 (PDB 2FFU). Bound EA2 substrate peptide is cyan (sticks), Mn$^{2+}$ magenta (sphere) and UDP grey (sticks). Ligands are taken from DM-T-2. For superposition with WT-T-2 ligands, see figure S1A. B, superposition of the UDP-sugar binding site of DM-T-2 and WT-T-2. Electron density is rendered at 1 σ and carved at 1.6 Å. C, depiction of UDP-GalNAc analog 1 in a co-crystal structure with DM-T-2 at 3.1 Å, and UDP-GalNAc in a co-crystal structure with WT-T-2 (PDB 4D0T) (30), as well as WT and mutated gatekeeper residues. D, substrate specificities of DM-T1 and DM-T-2 as determined in an *in vitro* glycosylation assay with detection by SAMDI-MS. For comparison with WT-GalNAcT glycosylation, see figure S2. Data are from one representative out of two independent experiments. A co-crystal structure of DM-T-2, Mn$^{2+}$ and UDP-GalNAc analog 1 at 3.1 Å resolution helped us visualize how the DM-T-2 active site mutations affect enzyme-substrate binding. In comparison with a corresponding WT-T-2/UDP-GalNAc/Mn$^{2+}$/EA2 complex (PDB 4D0T), UDP-sugar
binding is completely conserved (Fig. S1B, C and table S1) (30). DM-T-2 indeed contains a hole that accommodates the alkyne side chain bump in UDP-GalNAc analog 1 (Fig. 2C). The formation of additional van der Waals interactions between enzyme and substrate explain why the $K_M$ of DM-T-2 toward 1 is approximately ten-fold lower than the $K_M$ of WT-T-2 toward UDP-GalNAc (8).

To corroborate our structural interpretation that GalNAcT bump-and-hole engineering does not alter substrate peptide binding, we profiled the substrate specificities of two engineered GalNAcTs, DM-T-1 and DM-T-2 (8, 33-35). A peptide library containing a single Thr residue with randomized neighboring residues was used in in vitro glycosylation experiments and analyzed by self-assembled monolayers for matrix-assisted desorption/ionization mass spectrometry (SAMDI-MS, fig. 2D and fig. S2) (32). We found a striking difference in substrate specificity between DM-T-1 and DM-T-2 that was similar to the differences seen in the corresponding WT-GalNAcTs: T-1 generally preferred hydrophobic amino acids at -1-position and Glu, Ile, Pro, Val or Trp at +1-position of the Thr glycan acceptor, whereas T-2 showed a preference for Pro, Ala, Ser and Thr at -1-position (Fig. S2) (32). These data indicate that bump-and-hole engineering faithfully reports on GalNAcT isoenzyme activity, a crucial prerequisite for a cellular bump-and-hole system.
**Fig. 3:** Engineered GalNAcTs localize to the Golgi compartment and glycosylate protein substrates.  

**A**, expression construct for full-length GalNAcTs under the control of a Dox-inducible promoter. Inverted Tandem Repeats (ITR) are recognized by Sleeping Beauty transposase. WT-T-2 and DM-T-2 were expressed by stably transfected HepG2 cells in a Dox-inducible fashion.  

**B**, fluorescence microscopy of HepG2 cells stably transfected with T-2 constructs, induced with 0.2 µg/mL Dox and subsequently stained.  

**C**, *in vitro* glycosylation of proteins in a membrane fraction by full-length GalNAcTs using UDP-GalNAc analogs. Data are from one representative out of two independent experiments. Experiments were repeated with the membrane fraction of non-
transfected cells and soluble, purified GalNAcTs as an enzyme source (Fig. S3). DIC = Differential interference contrast; TA = Trans-activator.

DM-GalNAcTs glycosylate membrane proteins

We next sought to confirm that DM-GalNAcTs localize to the Golgi compartment and glycosylate proteins in a membrane environment. Full-length WT- and DM-T-1 and -T-2 with a C-terminal VSV-G epitope tag exhibited doxycycline (Dox) dose-dependent expression under the control of a tetracycline-responsive promoter in stably transfected HepG2 cells (Fig. 3A and fig. S3A) (36). All tested GalNAcTs co-localized with the Golgi marker giantin, confirming native localization of engineered GalNAcTs (Fig. 3B and fig. S3B). To assess GalNAcT activity, membrane protein fractions were prepared from WT- and DM-GalNAcT expressing HepG2 cells and used for in vitro glycosylation experiments. After incubation with alkyne-containing UDP-GalNAc analog 1, alkyne-tagged glycoproteins were derivatized with azide-biotin using Cu¹⁻-catalyzed [3+2] “click” cycloaddition, and characterized on a streptavidin blot. Compounds 1, 2 and 3 were specific substrates for DM- but not WT-GalNAcTs. In the presence of these substrates, DM-T-1 and DM-T-2 labeled overlapping and also unique glycoprotein species (Fig. 3C and fig. S3C, D). In contrast, treatment of lysates from WT-T-1 and -2 expressing cells with compound 4 with a shorter alkyne chain led to substantial labeling, consistent with 4 being a substrate for WT-GalNAcTs (8, 26). These results were confirmed using soluble GalNAcTs and a membrane fraction of non-transfected cells (Fig. S3E). Importantly, profiling both enzyme activity and protein specificity of single GalNAcT isoenzymes has been impossible to date even in cell lysates. We next targeted a GalNAcT bump-and-hole platform in the living cell.
**Biosynthesis of a bumped UDP-GalNAc analog**

Among the insightful bump-and-hole studies that have probed various enzyme families, few have been performed in living cells due to the inability to deliver negatively charged substrates across the plasma membrane (5, 6, 9, 37). GalNAc analogs have been fed to cells as membrane-permeable per-acetylated precursors that are deprotected by esterases and converted to UDP-GalNAc analogs via the kinase GALK2 and the pyrophosphorylase AGX1 (Fig. 4A) (26, 27, 38, 39). By delivery of the corresponding protected GalNAc-1-phosphate analog, a GalNAc analog can bypass GALK2, such that only AGX1 is necessary for biosynthesis to the UDP-GalNAc analog. However, in accordance with previous findings (39), bumped UDP-GalNAc analogs 1, 2 and 3 were not biosynthesized from their sugar-1-phosphate precursors in the living cell (Fig. 4A, B).

A lack of AGX1 activity towards synthetic N-acetylglucosamine (GlcNAc) analogs has previously prompted the engineering of AGX1 to recognize the corresponding GlcNAc-1-phosphate analog as a substrate (40). As WT-AGX1 accepts both GlcNAc-1-phosphate and GalNAc-1-phosphate as substrates, we investigated whether AGX1 mutants could biosynthesize UDP-GalNAc analogs 1, 2 and 3 from the corresponding GalNAc-1-phosphate analogs in the living cell. We mutated the gatekeeper residues F381 and F383 to Gly or Ala in FLAG-tagged AGX1 expression constructs (40). GalNAc-1-phosphate analogs were delivered to stably transfected HEK293T cells by virtue of caged precursors 5, 6 and 7. UDP-sugar biosynthesis was then determined by high performance anion exchange chromatography with pulsed amperometric detection (HPAEC-PAD) of cell lysates. Gly and Ala mutants of F383 efficiently biosynthesized bumped UDP-GalNAc analog 1, while neither F381 single mutants nor any F381/F383 double mutants produced 1 despite equal expression levels (Fig. 4A) (40). In contrast to linear alkyne 1, neither methylated alkynes (2 or 3)
were biosynthesized by engineered AGX1^{F383A}, hereby called mut-AGX1 (Fig. 5B and fig. S4B).

We thus concluded that bumped GalNAc-1-P(SATE)_{2} precursor 5 can be used in conjunction with mut-AGX1 to deliver UDP-GalNAc analog 1 to the living cell and establish a GalNAcT bump-and-hole system.

**Fig 4:** Substrate delivery to the cytosol of living cells. *a*, schematic of substrate delivery. Non-permissive steps are indicated by crossed arrows. The epimerase GALE interconverts UDP-GlcNAc and UDP-GalNAc. *b*, HPAEC-PAD traces of extracts from HEK293T cells stably expressing WT-AGX1 or mut-AGX1 and fed with the indicated compounds. Dashed boxes indicate retention times of standards in separate reference runs (Fig. S4B). The product of potential epimerization of 1 by GALE, compound 8, is marked with an arrowhead. Data is of one experiment and was repeated for compound 5 in HEK293T cells transiently transfected with AGX1 constructs, as well as stably transfected K-562 cells (Fig. S4B and fig. S5). Insert: epimerization to 8 is
suppressed in GALE-deficient K-562 cells expressing mut-AGX1 and fed with 5, but not cells carrying a control single guide (sg)RNA. A reference trace of compound 1 is shown. Data are of one representative out of two independent experiments.

In HPAEC-PAD chromatograms of cells biosynthesizing UDP-GalNAc analog 1, we consistently found a satellite peak that eluted after 1 (arrowhead in figure 4B). The epimerase GALE maintains a cytosolic equilibrium between UDP-GalNAc and UDP-GlcNAc and has been shown to also accept azide-containing analogs (38, 41). We found that GALE epimerizes UDP-GalNAc analog 1 to the corresponding UDP-GlcNAc analog 8 (Fig. 4A): first, HPAEC-mass spectrometry confirmed that the satellite peak was caused by an isomer of UDP-GalNAc analog 1 with 328 m/z [M-2H]²⁻. Second, CRISPR-mediated GALE-KO in the K-562 background abrogated the satellite peak (Fig. 4B and fig. S5). In contrast, epimerization was still observed in GALE-containing cells. Alkyne-containing UDP-GlcNAc analog 8 might result in labeling of GlcNAc-containing glycans on the cell surface, such as N-linked glycans. We thus concluded that epimerization of 1 to 8 must be accounted for in glycoproteomics experiments, for example by enzymatic abrogation of cell surface N-glycans during sample processing.

A glycosyltransferase bump-and-hole-system in living cells

With a GalNAcT bump-and-hole system and a method for cellular substrate delivery of 1 in hand, we set out to probe isoenzyme-dependent glycosylation in the living cell. We cloned FLAG-tagged WT- or mut-AGX1 under a constitutive promoter into an expression vector containing VSV-G-tagged WT or DM versions of GalNAcT-1 or T-2 under the control of a Dox-inducible promoter (Fig. 5A). This setup allowed us to systematically assess any potential background protein labeling when UDP-GalNAc analog 1 could not be biosynthesized, or when 1 was biosynthesized but DM-
GalNAcTs were absent. To preclude any labeling due to epimerization of 1 to UDP-GlcNAc analog 8, we first used GALE-KO K-562 cells for labeling. Cells were supplemented with GalNAc to compensate for the loss of UDP-GalNAc biosynthesis (Fig. S5A). GalNAcT expression was induced (Fig. S6A), cells were fed with caged GalNAc-l-phosphate analog 5, and the cell surface was reacted with clickable Alexa488-picoly azide in the presence of a non-membrane-permissive Cu(I) complex (3, 42). After gating for positive VSV-G signal, flow cytometry showed a more than 15-fold fluorescence increase when both DM-T-2 and substrate 1 were present over controls lacking either (Fig. 5B and fig. S6B). In GALE-containing cells, a more than 2-fold higher signal was still measured over control cells when a functional bump-and-hole system was present (Fig. S6C).

In order to better visualize the scope of protein labeling by our bump-and-hole system, we reacted cell surfaces with a clickable version of the infrared dye CF680 to profile labeled cell surface glycoproteins by in-gel fluorescence. We compared protein labeling patterns of GALE-containing K-562 cells stably expressing T-1 or T-2 constructs. Here, N-linked glycans were removed by PNGase F treatment prior to analysis to reduce background fluorescence (Fig. S7A). A profound band pattern was observed when functional bump-and-hole pairs (5, mut-AGX1 and DM-GalNAcTs) were present (Fig. 5C and fig. S7B). Mut-AGX1 was required for labeling, confirming that our experiments probe enzymatic glycosylation rather than non-specific protein modification (43). Fluorescence was of similar intensity as in cells treated with well-characterized alkyne-containing N-acetylneuraminic acid precursor Ac4ManNAI k (44). Furthermore, the presence of DM-GalNAcT protein was essential, as omission of Dox induction prevented fluorescent labeling (Fig. S7B). Importantly, DM-T-1 and DM-T-2 produced slightly different band patterns, especially between 25 and 37 kDa. The most intense 110 kDa band migrated at slightly higher molecular
weight when glycosylated by DM-T-1 instead of DM-T-2, indicating that T-1 potentially labeled more sites of this particular protein. Digestion with the mucin-selective protease StcE completely removed this band in T-2 labeled samples (Fig. S7C), confirming labeling of mucin-type proteins that are rich in O-GalNAc glycosylation (45). Discrete band patterns were obtained from in-gel fluorescence experiments when GALE-KO cells were used and a functional bump-and-hole pair was present. In contrast, pulsing the same cells with GlcNAc analog 9, a precursor of UDP-GlcNAc analog 8, led to a diffuse background whenever mut-AGX1 was present, but independent of the GalNAcT construct used (Fig. S7D). To profile the identity of labeled cell surface glycoproteins, we synthesized clickable, acid-cleavable biotin-picolyl azide 10 as an enrichment handle of cell surface glycoproteins (Fig. 5D). Following lysis and PNGase F treatment, O-glycoproteins were enriched and digested on-bead with trypsin to profile the corresponding unmodified peptides by mass spectrometry. The myelogenous K-562 cell line expresses mucins and mucin-like proteins that are rich in O-GalNAc glycans such as CD36, CD43, CD45, Glycophorins A and C, and MUC18 (https://www.proteinatlas.org/). We found all these proteins enriched (log prob > 5 for DM and < 5 for WT samples, or at least 40 units higher for DM than controls) from lysates of cells carrying functional T-1 or T-2 bump-and-hole systems (5, mut-AGX1, DM-GalNAcT) over control cells lacking any component (Fig. S8). It is noteworthy that we found mainly cell surface glycoproteins with abundant O-glycosylation in these experiments, probably because DM-GalNAcTs were in constant competition for protein substrates with the corresponding endogenous WT-GalNAcTs in these cells.
**Fig. 5:** Selective bioorthogonal labeling of the living cell surface with bump-and-hole engineered GalNAcTs. A, GalNAcT and AGX1 co-expression construct and workflow of cell surface labeling. Red star depicts a fluorophore. B, labeling analysis of K-562 GALE-KO cells by flow cytometry of MB488-picoly azide labeled and intracellular VSV-G-stained cells. Data are individual values from three independent experiments, means ± SEM of MB488 median fluorescence intensity of VSV-G positive cells. Statistical analysis was performed by two-tailed ratio paired t-test. C, labeling analysis by in-gel fluorescence of PNGase F-treated lysates from metabolically labeled K-562 cells. Data is representative of three independent experiments. D, schematic of glycoprotein enrichment and on-bead digest. The bifunctional molecule 10 bears an acid-labile diphenyldisiloxane moiety. E, exemplary mass spectrometry data: mass spectrum (HCD) of a fully elaborated glycopeptide from SERPIN5A (site Thr39), and further examples from T-2-specific sites from STC2 (Thr28) and APOE (S308). FA = formic acid; MFI = mean fluorescence intensity.
The use of GalNAcT-KO SimpleCells has enabled the mapping of glycosylation sites that are introduced by several GalNAcTs including T-1 and T-2, but by design is not suited to profile the diversity of glycan structures beyond the initiating GalNAc. Our bump-and-hole approach can contribute this information if the bumped GalNAc analog is recognized and extended by downstream GTs. To probe this, we used bump-and-hole pairs to profile the glycoproteome of T-1-KO or T-2-KO HepG2 cells (21). Cells were transfected with GalNAcT-1 or T-2 and Dox-induced before labeling with GalNAc-1-phosphate analog 5. As glycoproteins are abundant in the HepG2 secretome (21), we clicked acid-labile enrichment handle 10 onto proteins in conditioned cell culture supernatants. Following on-bead tryptic digest to release non-glycosylated peptides, glycopeptides were liberated by acidic cleavage of the enrichment handle, and analyzed by mass spectrometry. We used the presence of a 491.2238 m/z ion in higher-energy collisional dissociation (HCD) as an indicator for a bumped glycan to trigger peptide sequencing by electron-transfer dissociation (ETD, fig. 5E and fig. S9). Following an automated search for glycopeptides containing the GalNAc analog alone or as a part of longer glycans, we obtained raw glycopeptide hits which we manually validated. Seventy-seven spectra were found of DM-T-2-modified glycopeptides that corresponded to 37 peptides with 27 glycosylation sites and the presence of up to four different glycan compositions per glycopeptide, the structures of which were inferred based on biosynthetic considerations (Data S1) (18). In contrast, four glycopeptides were found as modified by DM-T-1. Control cells fed with DMSO or containing WT GalNAcTs and fed with GalNAc-1-phosphate analog 5 showed no hits at all, or one (WT-T-2) and two (WT-T-1) glycopeptides, respectively. Several known glycosylation sites in the SimpleCell glycoproteomics dataset by Schjoldager et al. were confirmed herein, and new sites were revealed (Data S1) (19). For instance, we found hitherto unannotated T-2 glycosylation sites in the proteins ITIL2, AMBP,
Complement C3 and C4, GPC3 and STC2, and Laminin subunit gamma-1, among others (Fig. 5E and Data S1). T-2 is involved in lipid homeostasis and was found to glycosylate certain apolipoproteins, including ApoE at Thr307 and Ser308, in HepG2 cells (17, 21). Both sites were confirmed herein and, additionally, found to be elaborated up to the trisaccharide Neu5Acα-Galβ-GalNAc-Ser/Thr or tetrasaccharide Neu5Acα-Galβ-(Neu5Acα)GalNAc-Ser/Thr structures (Data S1) (18). Our data resolved ambiguity about O-glycosylation of apolipoprotein A-1 (ApoA1) in the peptide sequence 220-ATEHLSTLSEK-230 (21): previous data showed that upon T-2-KO, glycosylation of Thr221, Thr226 and Ser228 is slightly increased, but the reason for this was elusive (21). Our gain-of-function approach revealed that DM-T-2 glycosylated Ser225 while DM-T-1 glycosylated Thr221, indicating that T-2-KO cells lose Ser225 glycosylation that is compensated by increased glycosylation at the other sites. We attribute the smaller number of annotated glycopeptides in DM-T-1 cells versus DM-T-2 to the lower expression of T-1 versus T-2 GalNAcTs at 0.5 µg/mL Dox and/or the fact that T-1 glycosylates extracellular matrix proteins which may not be sufficiently soluble in conditioned media (Fig. 3B and fig. S3A). We repeated the experiment using DM-GalNAcTs and treatment with caged GalNAc-1-phosphate analog only, and found similar results (Data S2). Such detailed analysis is facilitated by the gain-of-function chemical tagging strategy that emulates native GalNAcT activity as a result of our bump-and-hole approach.

**Discussion**

The cellular glycoproteome is subject to disease-related alterations. Yet, we still lack methodology to selectively visualize, modify or sequence either a certain glycan subtype or the product of a certain GT. Bump-and-hole engineering has been artfully employed to obtain insight into the protein substrates of single kinases (4, 46), ADP-ribosyltransferases (5, 6), and methyltransferases
(37, 47, 48), among others (9). Only few of these systems have been established in the living cell, as cellular delivery of bumped substrates is rarely straightforward, especially when substrates are nucleotide-based. Herein, we established the first GT bump-and-hole system in the living cell to profile GalNAcTs, the largest GT family in the human genome. The fascinating biology of GalNAcTs, with multiple isoenzymes and a profound cross-talk between glycosylation and proteolysis (34, 49), prompted us to develop an isoenzyme-specific chemical reporter. This approach allowed us to expand our knowledge on T-1 and T-2 glycosylation sites by adding information on the nature of the mature glycans after extension by other GTs.

The era of quantitative biology demands great specificity in tools to probe cellular processes or expressed antigens. Biological tools, including antibodies, lectins, engineered binding proteins, or silenced hydrolases have been employed to profile or isolate cellular glycans (45, 50). While powerful probes, these proteins suffer from various drawbacks, including the dependence of binding on the structural context or the requirement to simplify glycans before performing binding studies. Additionally, all conventional strategies for glycan pull-down or visualization rely on post-glycosylation derivatization or binding. As a consequence, information on GT isoenzyme specificity is already lost or – in the context of knock-out cell lines – may have been obscured by partial redundancy. This “specificity space” is covered by our bump-and-hole approach that enables GT- and glycosite-specific labeling. As GalNAcT engineering retains both three-dimensional structure and peptide glycosylation preferences, this strategy probes glycosylation in an unbiased fashion without context dependence. Thereby, we obtained for the first time information on both site and structure of glycans initiated by a single GT isoenzyme in a single experiment.
Taken together, we now have a strategy in hand to specifically modify GT isoenzyme-dependent glycosylation sites on the living cell.

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**Supplementary Materials:**

Materials and Methods

Figures S1-S9

Table S1

Data S1 and S2
**Fig. S1.** A, overlay of ligands bound in DM-T-2/EA2/UDP/Mn^{2+} (ligands cyan) and a published crystal structure of WT-T-2/EA2/UDP/Mn^{2+} (ligands olive, PDB 2FFU), with electron density (map rendered at 1 σ and carved at 1.6 Å) taken from the DM-T-2 co-crystal structure. Mn^{2+} ions are colored in magenta and overlay completely. B, crystal structure of DM-T-2/1/Mn^{2+} in the hexameric unit cell. UDP-GalNAc analog 1 is rendered in sphere representation. C, superposition of the active sites of WT-T-2 (PDB 4D0T, orange) with UDP-GalNAc (light brown) and DM-T-2 (blue) with UDP-GalNAc analog 1 (grey). Gatekeeper residues are rendered in stick representation. Mn^{2+} ions are magenta and overlay completely.
**Fig. S2.** Substrate specificities of DM-T1 and DM-T-2 and comparison with WT enzymes as determined in an *in vitro* glycosylation assay with detection by SAMDI-MS. WT data corresponds with reported substrate specificities, and two replicates are shown for DM enzymes with the full library or a focused sub-library.
Fig. S3. A, WT-T-1 and DM-T-1 are expressed by stably transfected HepG2 cells in a Dox-inducible fashion. B, fluorescence microscopy of HepG2 cells stably transfected with T-1 constructs, induced with 2 µg/mL Dox and subsequently stained. C, streptavidin blot from figure 3C depicted in two different intensities. D, Western blot to assess expression of GalNAcTs in lysates used in C. E, in vitro glycosylation was repeated with a membrane fraction from untransfected HepG2 cells, using soluble GalNAcTs to perform glycosylations (8). Reactions were performed with or without a two-fold excess of UDP-GalNAc over UDP-sugars 1, 2, 3 and 4.
**Fig. S4.** Biosynthesis of UDP-GalNAc analog 1 by mut-AGX1. 

**A.** HPAEC-PAD traces of extracts from HEK293T cells transiently expressing AGX1 constructs, and fed with GalNAc-1-phosphate analog 5 or DMSO. All traces are normalized to the retention time of ADP-Glc as an external standard. Data is of a single experiment.

**B.** Full traces of the data displayed in Fig. 4B. Asterisk indicates an artefact from solvent filling.
Fig. S5. GALE-KO cells are deficient in epimerization of UDP-GalNAc and UDP-GlcNAc, as well as chemically modified analogs. A, HPAEC-PAD traces of GALE-KO or control sgRNA-transduced K-562 cells stably transfected with the indicated FLAG-tagged AGX1 constructs and fed with different concentrations of GalNAc or GlcNAc. Expression of AGX1 (FLAG) and GALE were analyzed by Western blot. Samples were re-blotted with a higher concentration of GALE antibody to assure absence of GALE in KO cells. Data are from one experiment. 

B, cells were fed with DMSO or compound 5, and UDP-sugar production was measured by HPAEC-PAD. GALE-KO contain elevated levels of UDP-Gal as cells are supplemented with galactose to maintain viability and UDP-Gal cannot be epimerized to UDP-Glc. Expression levels of AGX1 (FLAG) and GALE are analyzed by Western blot. Data are of one representative out of two independent experiments. 

C, GALE-KO or control sgRNA-transduced K-562 cells stably transfected with the indicated AGX1 constructs were fed with the indicated compounds, and UDP-sugar production was measured by HPAEC-PAD. Data are from one experiment.
**Fig. S6.** A, gating scheme for flow cytometry experiments in Fig. 5. B, primary flow cytometry data of the experiment in figure 5C. Two technical replicates are shown. C, primary flow cytometry data of “wild type” K-562 cells after induction with 0.5 µg/mL Dox and gating on VSV-G-positive cells. Two technical replicates are shown for cells treated with compound 5. At least 500 gated cells were used for analysis per sample in B and C. D, statistical analysis of the experiment in C. Data are individual values from three independent experiments, means ± SEM of MB488 median fluorescence intensity of VSV-G positive cells. Statistical analysis was performed by two-tailed ratio paired t-test.
Fig. S7. Labeling analysis of K-562 cells by in-gel fluorescence. A, cells expressing AGX1 and GalNAcT-2 constructs were labeled as in figure 5C and analyzed by in-gel fluorescence. Treatment of lysates with PNGase F significantly shifts certain background bands to lower molecular weight. Arrow indicates PNGase F band in Coomassie stain. B, full gel of the experiment depicted in figure 5C. The sialic acid precursor Ac4ManNAk was used as a positive control, and the effect of omitting Dox was investigated. C, treatment of lysates prepared as in A with the glycoprotease StcE. Arrow indicates StcE band in Coomassie stain. D, dissecting GlcNAc vs. GalNAc labeling by using probes 5 and 9. GALE-KO K-562 cells expressing AGX1 and GalNAcT-2 constructs were treated with Dox or left untreated, and fed with compounds 5 or 9. Labeling was performed as in figure 5C. Wild type K-562 cells prepared as in A were used to compare labeling patterns. Data are from one representative out of three independent experiments.
Fig. S8. GalNAcT bump-and-hole pairs selectively label highly O-glycosylated proteins. Heat map represents log prob of detected proteins after enrichment of cell surface proteins of K-562 cells expressing mut-AGX1 and indicated GalNAcT constructs and fed with the indicated compounds.
Fig. S9. Exemplary mass spectra from glycopeptides after modification by DM-T-2. HCD (mainly glycan fragmentation) and ETD (mainly peptide fragmentation) spectra are shown, and ions are annotated. A, glycosylation at Thr28 of STC2 as a newly-identified T-2-specific modification (63). B, confirmation of Ser308 as a T-2-specific modification of ApoE. C, extension of chemically tagged GalNAc by elaborating glycosyltransferases on T39 of SERPIN5A. Legend depicts tentative structural assignments of oxonium ions. Loss of 161 m/z in HCD and 194 m/z peak in ETD likely depicts fragment masses of the triazole-based linker.
### Table S1.
Crystallographic data statistics.

| Parameters                              | T-2 I253A/L310A with UDP, Mn\(^{2+}\) and EA2 (PDB 6E7I) | T-2 I253A/L310A with Mn\(^{2+}\), UDP-GalNAc analogue 1 (PDB 6NQT) |
|-----------------------------------------|----------------------------------------------------------|---------------------------------------------------------------------|
| a, b (Å)                                | 69.31                                                    | 116.58, 120.13                                                      |
| c (Å)                                   | 169.78                                                   | 247.39                                                             |
| α, β, γ (°)                              | 90. 90, 120                                             | 90, 90, 90                                                         |
| Resolution range (Å)                    | 56.7-1.80                                                | 39.0-3.05                                                          |
| Space group                             | P6₁ (1 mol/ASU)                                         | P 2₁ 2₁ 2₁ (6 mol/ASU)                                             |
| Wavelength (Å) Synchronous source       | 0.9774/ALS BL5:0.1                                      | 0.9753/SSRL BL7-1                                                 |
| Number of measured/unique reflections   | 230,556/30,854                                          | 286,630/64,645                                                     |
| \(R_{merge}\) (%)                       | 10.7 (47.1)                                             | 13.5 (77.1)                                                       |
| Completeness (%) and Multiplicity       | 93.6 (93.7)/5.8 (5.6)                                   | 97.1 (98.3)/9.0 (3.0)                                             |
| Mean I(σ)                               | 9.5 (0.7)                                                | 7.7 (1.6)                                                         |
| Mean I half-set correlation coefficient | 0.994 (0.674)                                           | 0.989 (0.645)                                                     |
| Refined statistics                      |                                                         |                                                                    |
| Reflections used, total/test set        | 37,096/1,998                                            | 61,083/3,345                                                      |
| Crystallographic \(R_{merge}\)/\(R_{merge}\) | 16.7/21.6                                               | 23.1/28.6                                                          |
| r.m.s.d. bond lengths (Å)               | 0.013                                                   | 0.016                                                              |
| r.m.s.d. bond angles (°)                | 1.61                                                    | 2.06                                                               |
| Number of protein atoms/total atoms     | 3,899/4,314                                             | 22,695/23,138                                                     |
| B-factor statistics \(\AA^2\)           |                                                         |                                                                    |
| Overall Wilson B-factor                  | 16.5/11.6                                               | 62.9/63.8                                                         |
| Protein, average by chain ID            | 17.0 (chain A)                                          | 54.0 (A), 63.7 (B), 62.7 (C), 62.8 (D), 74.1                     |
|                                          |                                                         | (E), 82.1 (F)                                                      |
| Ligand, average by chain ID             |                                  Peptide (62 atoms): 17.6                               | Cmpd. 1 (43 atoms): 50.5 (A) 64.9 (B), 57.1                      |
|                                          |                                  UDP (25 atoms): 10.9                                   | (C), 59.6 (D), 72.7 (E), 77.7 (F)                                 |
|                                          |                                  Manganese (1 atom): 10.1                                | Manganese (1 atom): 49.3 (A) 30.6 (B), 48.7                       |
|                                          |                                  22.0 (327 waters)                                      | (C), 50.8 (D), 47.7 (E), 73.2 (F)                                 |
| Solvent/other atoms                     |                                                         | 34.2 (189 waters)                                                 |
| Ramachandran Statistics Protein geometry |                                                         |                                                                    |
| Most favored and additional allowed (%) | 99.3 (434 of 437 non-Proline non-                  | 96.4 (2,475 of 2,567 non-Proline non-                          |
|                                          | Glycine residues)                                   | Glycine residues)                                                 |
| Generously allowed (%)                   | 0.2 (1 residue)                                        | 2.2 (56 residues)                                                 |
| Outliers (%)                            | 0.5 (2 residues)                                       | 1.4 (36 residues)                                                 |

Figures in parenthesis relate to the outer shell.

\(^*\) \(R_{merge} = \frac{\Sigma_{hkl} \Sigma_{j=1}^N |I_{hkl} - I_{hkl}(j)|}{\Sigma_{hkl} \Sigma_{j=1}^N N I_{hkl}(j)}\), where \(N\) is the redundancy of the data. In parentheses, outermost shell statistics at these limiting values: 1.85 – 1.80 Å in GalNac T2 with EA2 and UDP and 3.21 - 3.05 Å in GalNac T2 UDP-GalNAc analog 1.

\(^{†}\) \(R_{factor} = \frac{\Sigma_{hkl} ||F_{obs}|-|F_{calc}||}{\Sigma_{hkl} |F_{obs}|}\), where the \(F_{obs}\) and \(F_{calc}\) are the observed and calculated structure factor amplitudes of reflection \(hkl\).

\(^{‡}\) \(R_{free} = \) is equal to \(R_{factor}\) for a randomly selected 5.0 % subset of the total reflections that were held aside throughout refinement for cross-validation.

\(^{§}\) According to Procheck.
Data S1. Glycoproteomics data of secretome from HepG2-T1−/− and HepG2-T2−/− cells stably transfected with combinations of AGX1 and T-1 or T-2, respectively, and treated with compound 5 or DMSO.
Data S2. Glycoproteomics data of secretome from HepG2-T1\textsuperscript{−/−} and HepG2-T2\textsuperscript{−/−} cells stably transfected with mut-AGX1 and DM-T-1 or -T-2, respectively, and treated with compound 5.