

1 **A cooperative network of molecular “hot spots” highlights**
2 **the complexity of LH3 collagen glycosyltransferase activities**

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19

20 **ABSTRACT**

21 Hydroxylysine glycosylations are collagen-specific post-translational modifications essential
22 for maturation and homeostasis of fibrillar as well as non-fibrillar collagen molecules. Lysyl
23 hydroxylase 3 (LH3) is the only human enzyme capable of performing two chemically-
24 distinct collagen glycosyltransferase reactions using the same catalytic site: inverting beta-
25 1,O-galactosylation of hydroxylysines and retaining alpha-1,2-glycosylation of galactosyl
26 hydroxylysines.

27 Here, we used structure-based mutagenesis to show that both glycosyltransferase activities are
28 strongly dependent on a broad cooperative network of amino acid side chains which includes
29 the first-shell environment of the glycosyltransferase catalytic site and shares features with
30 both retaining and inverting enzymes. We identified critical “hot spots” leading to selective
31 loss of inverting activity without affecting the retaining reaction. Finally, we present
32 molecular structures of LH3 in complex with UDP-sugar analogs which provide the first
33 structural templates for LH3 glycosyltransferase inhibitor development.

34 Collectively, our data provide a comprehensive overview of the complex network of shapes,
35 charges and interactions that enable LH3 glycosyltransferase activities, expanding the
36 molecular framework for the manipulation of glycosyltransferase functions in biomedical and
37 biotechnological applications.

38

39 **KEYWORDS**

40 Collagen biosynthesis; extracellular matrix; post-translational modifications; collagen
41 glycosylations; glycosyltransferase; lysyl hydroxylase.

42

43 1. INTRODUCTION

44 Collagens are the most abundant proteins in the human body and are highly conserved from
45 sponges to mammals (Luther et al, 2011; Myllyharju & Kivirikko, 2004). The different
46 oligomeric architectures and roles of collagen molecules strongly depend on a variety of post-
47 translational modifications, including proline and lysine hydroxylations, as well highly
48 specific glycosylations of hydroxylated lysines (HyK) (Cummings, 2009; Myllyharju &
49 Kivirikko, 2004). The disaccharide present on HyK contains a highly conserved
50 glucosyl(α -1,2)-galactosyl(β -1,O) glycan moiety, whose identity was discovered in the late
51 sixties (Spiro & Spiro, 1971; Spiro, 1967; Spiro, 1969). Monosaccharidic galactosyl-(β -1,O)
52 HyK have been identified as well, as a result of catabolic reactions carried out by the collagen
53 α -glucosidase, an enzyme highly specific for the disaccharide present on collagenous
54 domains. The role for this enzyme is to localize collagen in the glomerular basement
55 membrane (Hamazaki & Hamazaki, 2016; Sternberg et al, 1982; Sternberg & Spiro, 1980).
56 The spread of glycosylation largely depends on collagen type (Bornstein & Sage, 1980; Spiro,
57 1969; Terajima et al, 2014), on the functional area inside tissues (Moro et al, 2000; Schofield
58 et al, 1971; Toole et al, 1972), on the developmental stage (Rautavuoma et al, 2004; Sipila et
59 al, 2007) and on disease states (Lehmann et al, 1995; Salo et al, 2008; Tenni et al, 1993).
60 Although extensively studied, the precise mechanisms of collagen glycosylation and their
61 biological relevance in collagen homeostasis have remained poorly understood.
62 The identity and exquisite stereochemistry of the Glc(α -1,2)-Gal(β -1,O)-HyK-linked
63 carbohydrate supports the idea that at least two distinct enzyme types are needed to fully
64 incorporate these complex post-translational modifications on collagen molecules (Hennet,
65 2019). The first reaction indeed requires an inverting-type galactosyltransferase (GalT) acting
66 on HyK, whereas the subsequent glucosylation is catalyzed by a retaining-type
67 glucosylgalactosyltransferase (GlcT). Multifunctional lysyl hydroxylase 3 (LH3) was the only

68 enzyme identified as possessing lysyl hydroxylase activity as well as GalT and GlcT activities
69 *in vitro* (Wang et al, 2002a). The glycosyltransferase activities are specific of LH3, as highly
70 homologous LH1 and LH2a/b are not capable of catalyzing these reactions (Heikkinen et al,
71 2000). Conversely, *in vivo* studies have demonstrated that decreased LH3 protein levels
72 and/or pathogenic mutations in LH3 GT domain, exclusively impair the GlcT activity (Ewans
73 et al, 2019; Salo et al, 2008; Savolainen et al, 1981). This occurs secondarily to the LH3
74 p.Asn223Ser, which introduces an additional glycosylation site within the enzyme's GT
75 domain leading to an osteogenesis imperfecta-like phenotype (Salo et al, 2008); and in the
76 recently identified LH3 p.Pro270Leu, which results in a Stickler-like syndrome with vascular
77 complications and variable features typical of Ehlers-Danlos syndrome and Epidermolysis
78 Bullosa (Ewans et al, 2019). Mouse studies have also shown that only the LH3 GlcT activity
79 is indispensable for the biosynthesis of collagen IV and formation of the basement membrane
80 during embryonic development (Rautavuoma et al, 2004; Ruotsalainen et al, 2006), consistent
81 with the presence of additional collagen galactosyltransferases. Two genes encoding for O-
82 galactosyltransferases (*GLT25D1* and *GLT25D2*) were recently identified (Perrin-Tricaud et
83 al, 2011; Schegg et al, 2009). It is of interest that *GLT25D1* and LH3 were proposed to act in
84 concert on collagen molecules (Schegg et al, 2009; Sricholpech et al, 2011; Yamauchi &
85 Sricholpech, 2012). Studies on osteosarcoma cell lines which produce large amounts of
86 fibrillar collagens, showed that the simultaneous deletion of *GLT25D1* and *GLT25D2* resulted
87 in growth arrest due to lack of glycosylation, further indicating that the GalT activity of LH3
88 might not be as essential as its GlcT activity (Baumann & Hennet, 2016).

89 These data support the existence of a specific and highly conserved machinery for collagen O-
90 glycosylation, and sustain the hypothesis that *in vivo* the entire collagen glycosylation
91 machinery may involve distinct proteins and protein complexes for GalT and GlcT reactions.
92 This raises the intriguing question of how this highly conserved process is spatiotemporally

93 regulated at the molecular level. Our current understanding of collagen glycosyltransferases is
94 however restricted to three-dimensional structures of human LH3 in complex with UDP-sugar
95 donor substrates (Scietti et al, 2018) and to few mutagenesis studies focusing on the main
96 hallmarks of Mn^{2+} -dependent glycosyltransferase catalysis (Wang et al, 2002a; Wang et al,
97 2002b).

98 Here, we combine site-directed mutagenesis scanning with biochemistry and structural
99 biology to characterize the glycosyltransferase activities of human LH3. Our data highlight an
100 overall distribution of “hot spots” around the extended glycosyltransferase cavity of LH3,
101 critically involved in both GalT and GlcT functions, and very few amino acid residues
102 capable of selectively abolishing transfer of galactose to HyK without affecting GlcT activity.
103 Finally, we also identify and characterize UDP-sugar substrate analogs acting as inhibitors of
104 LH3 glycosyltransferase activities.

105 Together, our results provide insights into the LH3 glycosyltransferase activities and expand
106 the available structural framework for the development of collagen GalT/GlcT inhibitors.
107 These insights will assist with the manipulation of LH3 protein functions and donor substrate
108 specificity in biomedical applications.

109

110 **RESULTS**

111 *Features and roles of the non-conserved LH3 “glycoloop”*

112 The LH3 N-terminal GT domain shares its fold with Mn^{2+} -dependent GT-A
113 glycosyltransferases, encompassing a UDP-donor substrate binding cavity stretched towards a
114 GalT/GlcT catalytic pocket (Scietti et al, 2018). We firstly inspected the structural
115 organization of the UDP binding cavity to identify signatures for LH3 glycosyltransferase
116 activity by comparing the amino acid sequences for the residues surrounding the UDP-sugar
117 donor substrate with those of GalT/GlcT-inactive LH1 and LH2a/b isoforms. We found that

118 nearly all residues involved in Mn^{2+} and UDP-sugar binding are conserved in orthologs, with
119 the exception of Val80, becoming Lys68 in LH1 and Gly80 in LH2a/b (Fig 1). The presence
120 of a different amino acid side chain surrounding the donor substrate cavity led us to consider
121 whether this could be a discriminating functional feature among the GT domains in LH
122 enzymes. In LH3, Val80 is located in the middle of a flexible “glycoloop” (Gly72-Gly87), not
123 visible in the electron density of the ligand-free LH3 structure and stabilized upon UDP-
124 substrate binding (Scietti et al, 2018). Within this glycoloop, residue Val80 is in close
125 proximity to the ribose ring of the UDP-sugar donor substrates. We hypothesized that
126 introduction of a large, positively charged residue such as Lys in LH1, or alterations due to
127 complete removal of side chain steric hindrance such as Gly in LH2 could lead to inability of
128 binding donor substrates. We therefore generated the LH3 Val80Lys and Val80Gly mutants.
129 Both enzyme variants were found to be folded based on analytical gel filtration and
130 differential scanning fluorimetry (DSF), and showed lysine hydroxylation activity comparable
131 to wild-type LH3 (supplementary Fig S1). Conversely, the mutation resulted in significant
132 reduction of both GalT and GlcT activities when the reaction was carried out in presence of
133 both donor and acceptor (gelatin) substrates (Fig 2, supplementary Table S1). Considering
134 that wild-type LH3 is also capable of activating donor UDP-sugar substrates and release UDP
135 in absence of the acceptor collagen substrate (“uncoupled” activity, as defined in (Scietti et al,
136 2018)), we investigated the impact of the Val80Lys and the Val80Gly mutations also in
137 absence of acceptor substrates. In this case, the experiments yielded minor, but significant
138 reduction in the enzymatic activities between wild-type and mutant LH3 (Fig 2,
139 supplementary Table S1), indicating that the Val80 residue might be involved in the
140 productive positioning of the donor substrate during transfer of the glycan moiety to the
141 acceptor substrate, rather than stabilizing the UDP moiety in the catalytic pocket.

142 To further rationalize the implications of LH3 Val80 in GalT and GlcT activities, we
143 crystallized and solved the 3.0-Å resolution structure of the Val80Lys mutant in complex with
144 Mn^{2+} , and also obtained its 2.3-Å resolution structure in presence of both Mn^{2+} and the UDP-
145 Glc donor substrate (supplementary Table S2). Overall, both structures superimpose almost
146 perfectly with wild-type LH3 for all domains (supplementary Fig S2). The structure of the
147 LH3 Val80Lys mutant bound to Mn^{2+} appeared identical to that of wild-type LH3. In both
148 structures, the glycoloop containing Val/Lys80 could not be modelled in the electron density
149 due to its high flexibility (supplementary Fig S2B). On the other hand, the side chain of the
150 Val80Lys residue could be modelled unambiguously in the experimental electron density of
151 the UDP-donor substrate bound structure (Fig 3A). Despite the increased steric hindrance, the
152 mutated Lys80 residue adopted a conformation compatible with the simultaneous presence of
153 the UDP-Glc in the catalytic cavity. However, similar to what observed for wild-type LH3,
154 the glycan moiety of UDP-Glc was completely flexible and therefore not visible in the
155 electron density (Fig 3A). Collectively, these data are consistent with the alteration introduced
156 by the Val80Lys mutation, impacting partially on the LH3 glycosyltransferase catalytic
157 activities.

158 The glycoloop is a structural feature found exclusively in the GT domains of LH enzymes. It
159 incorporates Trp75, a residue whose aromatic side chain stabilizes the uridine moiety of the
160 donor substrate and, together with residue Tyr114 of the DxxD motif (a distinguishing feature
161 of LH3 GT domain, shared among LH enzymes, (Scietti et al, 2018)) “sandwiches” the donor
162 substrate in an aromatic stacking environment (Fig 1A). Both residues are critical for the LH3
163 GalT and GlcT enzymatic activities (Scietti et al, 2018). The conformation adopted by the
164 LH3 glycoloop in the presence of UDP-donor substrates is however not accompanied by other
165 significant structural changes in surrounding amino acids, with the exception of minor
166 rearrangements of distant residue Trp92 (not conserved in other LH isoenzymes (Fig 1)),

167 whose bulky side chain rearranges pointing towards the aromatic ring of Trp75. Prompted by
168 this observation, we mutated this residue to alanine and found that the presence of this variant
169 did not alter the folding of the enzyme (supplementary Fig S1A-B) nor its LH enzymatic
170 activity (supplementary Fig S1C). Conversely, the mutant showed 40% decrease for both
171 GalT and GlcT activities in presence of donor and acceptor substrate compared to LH3 wild-
172 type (Fig 2, supplementary Table S1); the impact of the Trp92Ala mutation on reactions in
173 absence of acceptor substrate seemed to affect both activities at similar levels, with lower
174 residual GalT (30%) compared to GlcT (40%) (Fig 2, supplementary Table S1). These
175 findings suggest that LH3-specific long-range interactions in the GT domain may contribute
176 to the productive conformations of the glycoloop in donor substrate-bound states.
177 In UDP-sugar bound structures, the glycoloop contacts a poly-Asp sequence (Asp188-
178 Asp191, Fig 1A); this sequence is partially conserved in LH isoforms lacking
179 glycosyltransferase activities (Fig 1B). Mutations of Asp190 and Asp191 were reported to
180 affect the glycosyltransferase enzymatic activities of LH3 (Wang et al, 2002b). Based on LH3
181 crystal structures, such behaviour is expected, since residues Asp190 and Asp191 point
182 towards the GalT/GlcT catalytic cavity. Interestingly, superposition of LH3 molecular
183 structures with GT-A fold glycosyltransferases show that Asp191 caps the N-terminal end of
184 an α -helix in a highly conserved position, with functional relevance in both retaining (Flint et
185 al, 2005; Persson et al, 2001) and inverting enzymes (Charnock & Davies, 1999; Pedersen et
186 al, 2000) (supplementary Fig S3, Table 1). We designed and generated individual alanine
187 mutants for both LH3 Asp190 and Asp191, and found that both variants were compatible with
188 folded and functional LH enzymes (supplementary Fig S1). When tested for GalT and GlcT
189 activity, both these mutants caused severe impairment in activation of donor UDP-sugar (as
190 shown by the strong reduction in uncoupled activity to less than 5% using UDP-Glc as
191 substrate) as well as in transfer of the sugar moiety to the acceptor substrate (Fig 2,

192 supplementary Table S1). However, none of these mutations resulted in a complete inactive
193 LH3 GalT nor GlcT glycosyltransferase. Collectively, these data point towards an extended
194 involvement of the residues of the poly-Asp repeat, and in particular Asp190 and Asp191, in
195 both the positioning and recognition of donor or acceptor substrates.

196

197 *LH3 shares features with both retaining and inverting glycosyltransferases*

198 After investigating the amino acid residues involved in stabilization of the UDP moiety of
199 donor substrates, we focused on another group of residues within the GT catalytic pocket,
200 opposite to the putative position of the flexible sugar rings of the same substrates (Fig 1A).
201 Many LH3 residues shaping this part of the glycosyltransferase catalytic cavity matched
202 catalytic amino acids found in other GT-A type glycosyltransferases (Fig 1A, supplementary
203 Fig S3) (Ardevol et al, 2016; Lairson et al, 2008). In particular, LH3 Trp145, a residue located
204 in one of the loops of the GT domain uniquely found in LH3, was previously suggested as a
205 possible candidate for modulation of LH3 GalT and GlcT activities. This residue was found to
206 adopt different side chain conformations in substrate-free and substrate-bound structures,
207 affecting the shape and steric hindrance of the enzyme's catalytic cavity (Scietti et al, 2018);
208 interestingly, nearly identical conformational changes were observed when comparing
209 substrate-free and substrate bound LH3 Val80Lys structures (Fig 3B). Mutating the LH3
210 Trp145 residue into alanine strongly reduced both GalT (6% residual) and GlcT (10%
211 residual) enzymatic activities (Fig 2, supplementary Table S1), without affecting other
212 enzyme's properties (supplementary Fig S1). This supports previous hypotheses of a "gating"
213 role for Trp145 in the GT catalytic cavity, assisting the productive positioning of sugar
214 moieties of donor substrates for effective transfer during catalysis. Comparison with
215 molecular structures of other glycosyltransferases (including distant homologs) highlighted
216 that most structurally-related enzymes manage to position aromatic side chains from different

217 structural elements of their fold in their catalytic cavities. Such structural arrangement is
218 reminiscent to that of Trp145 in LH3, but relies on completely different structural features of
219 the glycosyltransferase domain. In particular, similar aromatic residues were found in other
220 glycosyltransferases such as Tyr186 in LgtC from *Neisseria meningitidis*, Trp314 in the N-
221 acetylglucosaminide α -1,3-galactosyl transferase GGTA1, Trp300 in the histo-blood group
222 ABO system transferase, and Trp243 and Phe245 in the two glucoronyltransferases B3GAT3
223 and B3GAT1, respectively (Table 1, supplementary Fig S3). This supports
224 glycosyltransferases being highly versatile enzymes, displaying an impressive structural
225 plasticity to carry out reactions characterized by a very similar mechanism on a large variety
226 of specific donor and acceptor substrates.

227 Our previous structural comparisons of ligand-free and substrate-bound LH3 highlighted the
228 additional possibility of a concerted mechanism involving conformational changes of a non-
229 conserved aromatic residue located on the LH3 surface (Trp148, Fig 1), together with Trp145
230 (Scietti et al, 2018). To investigate such possibility, we mutated Trp148 into alanine. The
231 mutant enzyme was also found to behave like wild-type LH3 in folding and LH activity
232 (supplementary Fig S1). Glycosyltransferase assays showed that this variant had reduced
233 GalT (30% residual) and GlcT (50% residual) activities compared to the wild-type, in
234 presence of both donor and acceptor substrates (Fig 2, supplementary Table S1). Despite less
235 pronounced alterations compared to those observed when mutating Trp145, these data support
236 possible synergistic mechanisms between long-range acceptor substrate recognition on the
237 enzyme's surface and conformational rearrangements in the enzyme's catalytic site.

238 Molecular structures of LH3 in complex with UDP-Glc and Mn^{2+} , showed weak electron
239 density near the UDP pyrophosphate group partially compatible with glycan donor substrates,
240 likely representative of multiple conformations simultaneously trapped in the substrate
241 binding cavity (Scietti et al, 2018). We explored the LH3 catalytic cavity in its proximity,

242 looking for additional amino acids potentially critical for catalysis. In particular, we searched
243 for residues carrying carboxylic or amide side chains, thus capable of acting as candidate
244 catalytic nucleophiles for the formation of a (covalent) glycosyl-enzyme intermediate prior to
245 glycosylation of the acceptor substrate (Gloster, 2014).

246 In retaining-type glycosyltransferases belonging to the GT-6 family, a conserved glutamate
247 has been found to act as a nucleophile (supplementary Fig S3) (Albesa-Jove et al, 2017;
248 Coutinho et al, 2003; Gomez et al, 2012; Lombard et al, 2014; Patenaude et al, 2002). In LH3
249 structures, we noticed that residues Gln192, Asn165 and Glu141 point towards the cavity that
250 accommodates the glycan moiety (Fig 1A). We generated Ala mutations of all these residues,
251 obtaining in all cases folded functional LH enzymes (supplementary Fig S1). When probed
252 for GalT and GlcT activity, we found that both the Asn165Ala and the Gln192Ala mutants
253 were less efficient, but still capable, in activating UDP-donor substrates and performing sugar
254 transfer to acceptor substrates. Conversely, the Glu141Ala mutant was completely deprived of
255 both GalT and GlcT activities (Fig 2, supplementary Table S1). These data suggest Glu141 as
256 essential for catalysis and the surrounding negatively charged pocket composed of Asp190,
257 Asp191, Gln192, Asn165 comprising a broad network of amino acids which may concertedly
258 assist the LH3 glycosyltransferase activity.

259 Proximate to Glu141 in the GalT/GlcT cavity, residue Asn255 is the closest amino acid to the
260 UDP phosphate-sugar bond. Despite being fully conserved in LH isoforms (Fig 1B), this
261 residue is not found in any GT-A-type glycosyltransferases with known structures to date.
262 The side chain of Asn255 consistently points to a direction opposite to the donor substrate in
263 all LH3 structures (Fig 3B, supplementary Fig S2B). We wondered whether the side chain
264 amide group might be involved in catalysis, possibly through recognition of acceptor
265 substrates given the conformation displayed by this side chain. Surprisingly, LH3 Asn255Ala
266 mutants showed that their GalT activity was completely abolished, whereas the GlcT

267 enzymatic activity was reduced to 50% (Fig 2, supplementary Table S1); the protein was
268 properly folded and showed LH activity comparable to LH3 wild-type (supplementary Fig
269 S1). These results identify Asn255 as a possible critical discriminating residue for the two
270 glycosyltransferase activities of LH3, and rule out possible functions for this residue as
271 catalytic nucleophile for retaining-type glycosyltransferase mechanisms, given its major
272 impact restricted to the GalT catalytic activity.

273

274 *Pathogenic LH3 mutations in the LH3 GT domain affect protein folding*

275 Recently a new pathogenic LH3 mutation, Pro270Leu, has been identified and mapped at the
276 interface of the AC and GT domain (Ewans et al, 2019). This residue localizes in a loop
277 which is critical for shaping the GT cavity, although given its position it is unlikely to play
278 direct roles in catalysis. To better understand the impact of this pathogenic mutation on LH3
279 enzymatic activity, we generated a Pro270Leu mutant. In this case, due to very low
280 expression levels, we could not reliably carry out any *in vitro* investigations. Considering the
281 high reproducibility associated to recombinant production of a large variety of LH3 point
282 mutants, this result may indicate that this mutation is likely to severely impact the overall
283 enzyme stability rather than its enzymatic activity, resulting in extremely low protein
284 expression levels *in vitro* and likely also *in vivo*.

285

286 *Molecular structures of LH3 in complex with UDP-sugar analogs provide insights on how* 287 *glycan moieties are processed inside the LH3 catalytic cavity*

288 A frequent limitation associated to molecular characterizations of glycosyltransferases is the
289 high flexibility of the donor substrate glycan moiety within the catalytic cavity. Such
290 limitation becomes even more relevant when the enzyme is capable of processing UDP-sugar
291 molecules in absence of acceptor substrates, such as in the case of LH3. Considering our

292 previous (Scietti et al, 2018) and current co-crystallization results, we wondered whether free
293 UDP, the product of the enzymatic reaction, could remain bound in the LH3 GT domain with
294 the same efficiency as physiological donor substrates even after processing. We therefore
295 compared the binding of free UDP and donor UDP-sugars using DSF and detected a thermal
296 shift of 3.5 °C for free UDP, compared to a 2-2.5 °C shift using UDP-sugar substrates (Fig
297 4A). These results suggested that free UDP may bind to LH3, likely with even higher affinity
298 than UDP-glycan substrates, and that the GalT and GlcT reactions may therefore be affected
299 by product inhibition. To our surprise, the increase in thermal stability did not correlate with
300 efficient trapping of the reaction product in LH3 molecular structures. Independently from the
301 UDP concentration used in co-crystallization and soaking experiments, we never observed
302 any electron density for free UDP, yielding LH3 structures completely identical to ligand-free
303 enzyme (supplementary Fig S4A).

304 A report from Kivirikko and colleagues (Kivirikko & Myllyla, 1979) suggested that UDP-
305 glucuronic acid (UDP-GlcA) could act as competitive inhibitor of collagen
306 glycosyltransferases. Based on that, UDP-GlcA was used to isolate LH3 from chicken
307 embryos preparation (Myllyla et al, 1977; Wang et al, 2002a). However, no follow-up
308 biochemical studies could be found in the literature. We used DSF and luminescence-based
309 GalT and GlcT activity assays to investigate whether and how UDP-GlcA could affect LH3
310 enzymatic activity. DSF showed that UDP-GlcA indeed binds weakly to LH3, resulting in a
311 thermal shift of 1-1.5 °C (Fig 4A), highlighting limited stabilization compared to UDP-glycan
312 substrates and free UDP. Enzymatic assays also confirmed the competitive inhibition
313 displayed by this molecule (Fig 4B), with IC₅₀ values in the millimolar range (supplementary
314 Table S1). We also successfully co-crystallized and determined the 2.2-Å resolution crystal
315 structure of wild-type LH3 in complex with Mn²⁺ and UDP-GlcA (supplementary Table 2),
316 and found that the inhibitor could efficiently replace UDP-sugar donor substrates in the

317 substrate cavity (Fig 4C). We observed additional electron density for the glucuronic acid
318 moiety of the inhibitor in the enzyme's catalytic cavity, however this density could not be
319 interpreted with a single inhibitor conformation. Nevertheless, analysis of the experimental
320 electron density for the glucuronic acid moiety unambiguously showed that the inhibitor
321 adopts a "bent" conformation: the glycan moiety is deeply buried in the enzyme's catalytic
322 cavity proximate to residues Lys89, Asp190, Asp191, but distant from the residues found
323 critical for catalysis, including Trp145, Asn255 and Glu141 (Fig 4C), thereby leaving the
324 remaining space in the cavity for accommodating acceptor substrates.

325 Considering the possible conformations adopted by the glucuronic acid moiety based on
326 analysis of the electron density and the close proximity of the glucuronic acid moiety to LH3
327 Val80, we wondered whether the LH3 Val80Lys mutation could interfere with inhibitor
328 binding. We therefore co-crystallized and solved the 2.7-Å resolution structure of LH3
329 Val80Lys mutant in complex with UDP-GlcA (supplementary Table S2), and surprisingly
330 observed partial displacement of the glycoloop, for which we could not observe the typical
331 well defined electron density present in UDP-sugar-bound wild-type LH3 structures (Fig 4D).
332 At the same time, we could not observe improvements in the quality of the electron density
333 for the glucuronic acid moiety, resulting even poorer than what observed in wild-type LH3
334 (Fig 4D). This suggests that the intrinsic flexibility of the sugar-like moiety is not influenced
335 by specific conformations of the glycoloop, but rather by lack of specific protein-ligand
336 interactions that could provide stabilization of the sugar ring in a unique structural
337 arrangement.

338 The lack of a precise conformation for the glucuronic acid moiety observed crystal structures
339 prompted us for a further investigation of another UDP-sugar substrate analog, characterized
340 by lack of the carboxylic moiety of UDP-GlcA: UDP-Xylose (UDP-Xyl). Similar to UDP-
341 GlcA, UDP-Xyl resulted to be a weak inhibitor of LH3 GalT and GlcT activities (Fig 4A-B),

342 with IC₅₀ in the high micromolar range (supplementary Table S1). The 2.0-Å resolution
343 structure of LH3 in complex with Mn²⁺ and UDP-Xyl also showed the inhibitor bound inside
344 the enzyme's catalytic cavity, with weak electron density associated to the sugar moiety
345 suggesting multiple conformations of the xylose moiety attached to UDP, similar to what
346 observed for UDP-GlcA (Fig 4E). Taken together, these results suggest that the LH3 GT
347 binding cavity is capable of hosting a variety of UDP-sugar substrates, and that inhibition
348 likely depends on the reduced flexibility (and therefore increased stabilization) of the ligand
349 within the cavity. In this respect, we could expect that UDP-sugar analogs strongly interacting
350 with side chains proximate to the glycan moieties of UDP-GlcA and UDP-Xyl, may have the
351 potential to become powerful inhibitors of LH3 glycosyltransferase activities.

352

353 **DISCUSSION**

354 Glycosyltransferases are highly versatile, yet very specific enzymes. If carefully inspected,
355 they reveal a series of recurrent features that allow their comparative characterization even in
356 presence of very low sequence/structure conservation. LH3 has been known for long time as a
357 promiscuous enzyme able to exploit both an inverting and a retaining catalytic mechanism *in*
358 *vitro* for the specific transfer of different sugars to at least two different acceptor substrates:
359 the HyK and the GalHyK of collagens (Myllyharju & Kivirikko, 2004). Our *in vitro*
360 investigations highlight multiple areas surrounding the glycosyltransferase catalytic site that
361 can be considered as critical “hot spots” for the LH3 GalT and GlcT activities: the glycoloop,
362 the poly-Asp helix, the acceptor substrate cavity, and the region proximate to the UDP-sugar
363 donor substrate (Fig 1A).

364 LH3 is the only isoform of its family found capable of glycosyltransferase activities, however
365 the strong sequence conservation among human LH isoforms cannot be used to elucidate the
366 key determinants for this additional function. The GT domain has features (such as the DxxD

367 motif, the poly-Asp region, the glycoloop) not found in other glycosyltransferases, yet highly
368 conserved within the LH family (Fig 1B). Computational homology models of homologous
369 LH1 and LH2a/b (Scietti et al, 2019) (supplementary Fig S5) support the possibility of UDP-
370 sugar donor binding and processing. To further investigate LH enzymes' GT domains, we
371 focused on the only non-matching residue present within the whole amino acid sequence
372 directly surrounding the UDP-donor substrate. This residue (Val80 in LH3, corresponding to
373 Lys68 in LH1 and Gly80 in LH2a/b) is located in the middle of the glycoloop, in close
374 proximity to the ribose ring of the UDP-sugar donor substrate(s). Our data were consistent
375 with this residue being important for catalysis, as it assists the positioning of the bound donor
376 substrate and in particular the glycan moiety. However, the results also emphasize how
377 sequence alterations at this site are not sufficient to justify the lack of GalT/GlcT enzymatic
378 activities in homologous LH1/2. We therefore expanded our investigation to the second-shell
379 environment surrounding the donor substrate, and found that non-conserved LH3 Trp92
380 (Leu80 in LH1, Leu92 in LH2a/b) positions its aromatic side chain in a conformation that
381 stabilizes the entire glycoloop to facilitate the enzymatic reactions. Again, mutating this
382 residue did not lead to full loss of the glycosyltransferase activities. This is consistent with the
383 loss of GalT/GlcT functions in LH1 and LH2a/b being associated to a broad set of subtle
384 alterations, possibly involving residues distant from the actual enzyme's catalytic site, likely
385 essential for recognition of collagen acceptor substrates.

386 Prompted by these observations, we expanded our investigation to the LH3 catalytic cavity
387 expected to host the glycan moieties of the UDP-sugar donor substrates and the acceptor
388 molecules. Residues Asp190 and Asp191 of the characteristic LH3 poly-Asp helix lie in a
389 conserved position compared to other retaining and inverting type GT-A (supplementary Fig
390 S3, Table 1) and our data indicate that both these carboxylate moieties are critical for efficient
391 donor substrate activation and sugar transfer. Although residues matching these positions

392 have been proposed to act as catalytic nucleophiles in retaining type GT-A (Flint et al, 2005;
393 Persson et al, 2001; Wang et al, 2002b), and as catalytic bases in inverting type GT-A
394 (Chamock & Davies, 1999; Pedersen et al, 2000), their distances and the relative orientations
395 with respect to UDP-sugar donor substrates in the LH3 GT domain (Fig 1A) do not support
396 this hypothesis. Nevertheless, these residues play critical roles in recognizing and assisting the
397 proper positioning of donor UDP-sugar substrates, as shown by their proximity to glycan
398 moieties in LH3 co-crystal structures with UDP-sugar analogs (Fig 4C-E), a feature observed
399 also in mutagenesis studies on other GT-A retaining type glycosyltransferases (Lairson et al,
400 2004).

401 On the opposite site of the UDP-binding pocket, LH3 exhibits a non-conserved loop shaping
402 the GT catalytic cavity bearing two aromatic residues, the Trp145 and the Trp148 that seem to
403 act in a concerted way during catalysis, as suggested by comparisons between substrate-free
404 and substrate-bound LH3 molecular structures (Fig 1A). Trp145 is indispensable for both
405 GalT/GlcT activities: its conformational changes seem to respond to the presence and
406 conformational positioning of the donor substrate inside the catalytic cavity. Although located
407 in a loop that is uniquely found in LH3, the Trp145 side chain matches a site frequently
408 occupied by bulky aromatic residues in other GT-A glycosyltransferases that shape a portion
409 of the GT cavity to facilitate donor substrate processing and catalysis. This further highlights
410 the versatility of glycosyltransferases, in which many different structural features have
411 evolved to specifically recognize distinct donor and acceptor substrates, while preserving the
412 ability to carry out the same catalytic reaction. The implication in catalysis of the less
413 conserved Trp148 on the surface of the GT domain is even more intriguing: this residue
414 seems to coordinately respond to the rearrangements of its counterpart Trp145 in the catalytic
415 site, suggesting involvement in recognition of the acceptor substrate prior to its access into the
416 GT cavity. Alternatively, Trp148 may contribute to long-range stabilizing interactions with

417 collagen molecules, while they dock their HyK or GalHyK residues in the acceptor substrate
418 site during the GalT or GlcT reactions, respectively.

419 Catalytic nucleophiles have been clearly identified so far only in the retaining-type
420 glycosyltransferases belonging to the GT-6 family (Coutinho et al, 2003; Lombard et al,
421 2014), such as the α -1,3 galactosyltransferase (GGTA1) where a conserved glutamate is
422 found positioned on the β -face of the donor sugar (Albesa-Jove et al, 2017; Gomez et al,
423 2012; Patenaude et al, 2002) (supplementary Fig S3). Conversely, extensive structural
424 comparisons and mutagenesis experiments have been performed in the O-
425 galactosyltransferase LgtC from *Neisseria meningitidis*, focusing on matching residue,
426 Gln189 (Lairson et al, 2004). However, the role of this residue as catalytic nucleophile was
427 ruled out. This site is occupied by Gln192 in LH3. This residue is next to the poly-Asp helix,
428 distant from the sites occupied by donor substrates and in an arrangement that is not
429 compatible with a direct role in catalysis. However, our mutagenesis data indicate that
430 removal of the Gln192 side chain has a strong impact on LH3 glycosyltransferase activity. In
431 close proximity, we identified two other amino acid residues potentially involved in donor
432 substrate activation or transfer of sugar moieties to the acceptor molecule. Both Asn165 and
433 Glu141 point directly towards the glycan moiety of the donor substrate (Fig 1A). Whilst the
434 Asn165Ala mutation only reduced the glycosyltransferase activity by a factor of two (Fig 2,
435 supplementary Table 1), we found that Glu141 is essential for both GalT and GlcT activities,
436 as the Glu141Ala mutation yields results in a completely inactive LH3 glycosyltransferase
437 (Fig 2, supplementary Table 1). In LH3, Glu141 adopts a conformation corresponding to
438 Asp130 in the O-galactosyltransferase LgtC from *Neisseria meningitidis*, Asp125 in the O-
439 glucosyltransferase YGG1 from rabbit, and Gln247 in the O-glucosyltransferase GGTA1
440 from *Bos taurus* (supplementary Fig 3, Table 1). In LH3, residue Asn255 is the closest amino
441 acid to the UDP phosphate-sugar bond, but in crystal structures its side chain consistently

442 points to a direction opposite to the donor substrate (Fig 1A). When inspecting the molecular
443 structures of other GT-A glycosyltransferases, we noticed that this residue is not conserved
444 (supplementary Fig 3). On the contrary, this residue is fully conserved among human LH
445 isoforms (Fig 1B). Strikingly, the LH3 Asn255Ala mutant showed a complete loss of GalT
446 activity, but partially preserved the GlcT activity. Although the significance of *in vivo* LH3
447 GalT activity is uncertain, such activity is clearly detectable *in vitro* (Scietti et al, 2018; Wang
448 et al, 2002a). The ability of Asn255 to selectively abolish only LH3 GalT activity highlights
449 how LH3 can promiscuously accept and recognize very different acceptor substrates (i.e.,
450 collagen HyK *versus* GalHyK) within the same catalytic site and carry out two
451 glycosyltransferase reactions that rely on different mechanisms (i.e., inverting GalT and
452 retaining GlcT). Presently, LH3 is the only known glycosyltransferase capable of such
453 promiscuity, and our data show that it shares numerous features with both retaining and
454 inverting glycosyltransferases. Collectively, these results suggest the intriguing possibility
455 that LH3 is not just the ancestor of the whole LH family, but may preserve in its sequence
456 features belonging to the evolutionary precursors of both retaining and inverting
457 glycosyltransferases.

458 In addition, the present work provides a set of 3D structures of LH3 in complex with UDP-
459 sugar analogs, which work as mild inhibitors (Fig 4). Despite the high flexibility observed for
460 the glycan moieties of the bound molecules, the new molecular structures presented provide
461 valuable insights for structure-based drug development of inhibitors of LH3 GalT/GlcT
462 enzymatic activities. These molecules may give the spark to innovative therapeutic strategies
463 against pathological conditions characterized by excess collagen glycosylations, such as
464 osteogenesis imperfecta (Raghunath et al, 1994).

465 Together with the mutagenesis scanning of the entire GT catalytic site, our work provides a
466 comprehensive overview of the complex network of shapes, charges and interactions that
467 enable LH3 GalT and GlcT activities.

468

469 **EXPERIMENTAL PROCEDURES**

470 *5.1 Chemicals*

471 All chemicals were purchased from Sigma-Aldrich (Germany) unless otherwise specified.

472

473 *5.2 Site-directed mutagenesis*

474 The LH3 coding sequence (GenBank accession number BC011674.2 - Source Bioscience),
475 devoid of the N-terminal signal peptide was amplified using oligonucleotides containing in-
476 frame 5'-BamHI and 3'-NotI (supplementary Table 3) and cloned in a pCR8 vector, that was
477 used as a template for subsequent mutagenesis experiments. All LH3 mutants were generated
478 using the Phusion Site Directed Mutagenesis Kit (ThermoFisher Scientific). The entire
479 plasmid was amplified using phosphorylated primers. For all mutants the forward primer
480 introduced the mutation of interest (supplementary Table 3). The linear mutagenized plasmid
481 was phosphorylated prior to ligation. All plasmids were checked by Sanger sequencing prior
482 to cloning into the expression vector.

483

484 *5.3 Recombinant protein expression and purification*

485 Wild-type and mutant LH3 coding sequences were cloned into the pUPE.106.08 expression
486 vector (U-protein Express BV, The Netherlands) in frame with a 6xHis-tag followed by a
487 Tobacco Etch Virus (TEV) protease cleavage site. Suspension growing HEK293F cells (Life
488 Technologies, UK) were transfected at a confluence of 10^6 cells ml^{-1} , using 1 μg of plasmid
489 DNA and 3 μg of linear polyethyleneimine (PEI; Polysciences, Germany). Cells were
490 harvested 6 days after transfection by centrifuging the medium for 15 minutes at 1000 x g.
491 The clarified medium was filtered using a 0.2 mm syringe filter and the pH was adjusted to
492 8.0 prior to affinity purification as previously described (Scietti et al, 2018). All proteins were
493 isolated from the medium exploiting the affinity of the 6xHis tag for the HisTrap Excel (GE

494 Healthcare, USA) affinity column. The purified protein was further polished using a Superdex
495 200 10/300 GL (GE Healthcare) equilibrated in 25 mM HEPES/NaOH, 200 mM NaCl, pH
496 8.0, to obtain a homogenous protein sample; peak fractions containing the protein of interest
497 were pooled and concentrated to 1 mg mL⁻¹.

498

499 *5.4 Crystallization, data collection, structure determination and refinement*

500 Wild-type and mutant LH3 co-crystallization experiments were performed using the hanging-
501 drop vapor-diffusion method protocols as described in (Scietti et al, 2018), by mixing 0.5 µL
502 of enzyme concentrated at 3.5 mg mL⁻¹ with 0.5 µL of reservoir solutions composed of 600
503 mM sodium formate, 12% poly-glutamic-acid (PGA-LM, Molecular Dimensions), 100 mM
504 HEPES/NaOH, pH 7.8, 500 µM FeCl₂, 500 µM MnCl₂, supplemented with 1 mM of the
505 appropriate UDP-sugar analogs (UDP, UDP-glucose, UDP-glucuronic acid, UDP-xylose).
506 Crystals were cryo-protected with the mother liquor supplemented with 20% ethylene glycol,
507 harvested using MicroMounts Loops (Mitegen), flash-cooled and stored in liquid nitrogen
508 prior to data acquisition. X-ray diffraction data were collected at various beamlines of the
509 European Synchrotron Radiation Facility, Grenoble, France and at the Swiss Light Source,
510 Villigen, Switzerland. Data were indexed and integrated using *XDS* (Kabsch, 2010) and
511 scaled using *Aimless* (Evans & Murshudov, 2013). Data collection statistics are summarized
512 in Suppl. Table 2. The data showed strong anisotropy and therefore underwent anisotropic
513 cut-off using *STARANISO* (Tickle et al, 2018) prior to structure determination and
514 refinement. The structures were solved by molecular replacement using the structure of wild-
515 type LH3 in complex with Fe²⁺, 2-OG and Mn²⁺ (PDB ID: 6FXM) (Scietti et al, 2018) as
516 search model using *PHASER* (McCoy et al, 2007). The structure was refined with successive
517 steps of manual building in *COOT* (Emsley et al, 2010) and automated refinement with

518 *phenix.refine* (Adams et al, 2010). Model validation was performed with *MolProbity* (Chen et
519 al, 2010). Refinement statistics for the final models are reported in supplementary Table 2.

520

521 *5.5 Evaluation of LH3 GalT/GlcT enzymatic activity*

522 LH, GalT and GlcT activities were tested using luminescence-based enzymatic assays with a
523 GloMax Discovery (Promega, USA) as described in *Scietti et al.* (Scietti et al, 2018). Minor
524 modifications have been done for GalT/GlcT competitive inhibition assays, where 1 μ L of a
525 mixture of 250 μ M $MnCl_2$, 500 μ M UDP-galactose (GalT assay) or UDP-glucose (GlcT
526 assay) and increasing concentrations of either UDP-GlcA or UDP-Xyl were initially added to
527 the enzyme and gelatin substrate to start the reactions. All experiments were performed in
528 triplicates. Control experiments were performed in the same conditions by selectively
529 removing LH3. Data were analyzed and plotted using the GraphPad Prism 7 (Graphpad
530 Software, USA).

531

532 *5.6 Differential Scanning Fluorimetry (DSF)*

533 DSF assays were performed on LH3 wild-type and mutants using a Tycho NT.6 instrument
534 (NanoTemper Technologies GmbH, Germany). LH3 samples at a concentration of 1 mg/mL
535 in a buffer composed of 25 mM HEPES, 500 mM NaCl, pH 8. Binding assays were
536 performed by incubating LH3 with 50 μ M $MnCl_2$ and 5 mM free UDP or UDP-sugar donor
537 substrates or their analogs. Data were analyzed and plotted using the GraphPad Prism 7
538 (Graphpad Software, USA).

539

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543 useful discussions and M. Miao for support in crystallization experiments.

544

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554 of the report and in the decision to submit this article for publication.

555

556 **AUTHOR CONTRIBUTIONS**

557 AC generated and purified LH3 mutants with support from SF, carried out biochemical
558 investigations and crystallized LH3 mutants; LS purified and crystallized wild-type LH3 in
559 complex with UDP-sugar analogs; FDG performed LH3 enzymatic activity assays; LS, FDG
560 and FF carried out structural refinement and analyses; TR was part of the study group
561 focusing on LH3 pathogenic variants. FF designed the study with help from AC, FDG and
562 LS. AC and FF wrote the paper, with contributions from all authors.

563

564 **CONFLICT OF INTEREST STATEMENT**

565 The authors declare no competing interests.

566

567

568 **Table 1: List of glycosyltransferase enzymes used for comparisons with human LH3.**

569 The list includes the indication of the catalytic bases and nucleophile residues as proposed in

570 the original papers describing the various glycosyltransferases, with the corresponding residue

571 number in human LH3 based on structural superpositions.

Protein Name	Type	PDB ID	catalytic base residue	nucleophile acceptor residue	corresp. LH3 residue	reference paper
LgtC - GALACTOSYL TRANSFERASE LGTC (<i>N. meningitidis</i>)	retaining	1GA8				Persson et al., 2001 10.1038/84168
GYG1 - Glycogenin (<i>O. cuniculus</i>)	retaining	1LL2		Asp163	Asp191	Gibbons et al., 2002 10.1016/S0022-2836(02)00305-4
mgs - Mannosylglycerate synthase (<i>R. marinus</i>)	retaining	2BO8				Flint et al., 2005 10.1038/nsmb950
GALNT10 - Polypeptide N-acetylgalactosaminyl transferase 10 (<i>H. sapiens</i>)	retaining	2D7R		Gln346	Asp190	Kubota et al., 2006 10.1016/j.jmb.2006.03.061
GGTA1 - N-Acetyllactosaminide α -1,3-galactosyl transferase (R365K) (<i>B. taurus</i>)	retaining	5NRB		Glu317	Gln192	Albesa-Jove et al., 2017 10.1002/anie.201707922
ABO - Histo-blood group ABO system transferase (<i>H. sapiens</i>)	retaining	1LZI		Glu303	Gln192	Patenaude et al., 2002 10.1038/nsb832
spsA - PROTEIN (SPORE COAT POLYSACCHARIDE BIOSYNTHESIS PROTEIN SPSA)	inverting	1QGQ	Asp191		Asp191	Charnock et al., 1999 10.1128/JB.183.1.77-85.2001
MGAT1 - N-acetylglucosaminyl transferase I (<i>O. cuniculus</i>)	inverting	1FOA	Asp 291		Asp191	Unligil et al., 2000 10.1093/emboj/19.20.5269
Mfng - Manic Fringe glycosyltransferase (<i>M. musculus</i>)	inverting	2JOB	Asp 232		Asp191	Jinek et al., 2006 10.1038/nsmb1144
B3GAT3 - GLUCURONYLTRANSFERASE I	inverting	1FGG	Glu281		Asp190	Pedersen et al., 2000 10.1074/jbc.M007399200
B3GAT1 – Galactosylgalactosylxylosyl protein 3-beta-glucuronosyltransferase 1		1V84				Kakuda et al., 2004 10.1074/jbc.M400622200

572

573

574 **FIGURE LEGENDS**

575 **Figure 1: Features of the LH3 glycosyltransferase (GT) domain.** (A) Cartoon
576 representation of the LH3 GT domain (PDB ID: 6FXR) showing the key residues shaping the
577 catalytic site as sticks. The PolyAsp motif (brown) and the Glycoloop (cyan) involved in
578 binding of UDP-sugar donor substrates are shown. The residues implicated in the catalytic
579 activity and investigated in this works are colored, while the residues depicted in grey have
580 already been shown to be essential in Mn^{2+} (purple sphere) and UDP (black sticks)
581 coordination. (B) Sequence alignment of human LH1, LH2 and LH3 GT domains,
582 highlighting similarities and differences in the amino acid residues within the active site.
583 Residues shown in Fig 1A are indicated with a triangle. Colored boxes indicate the PolyAsp
584 motif (brown) and the Glycoloop (cyan).

585

586 **Figure 2: Evaluation of the effect of LH3 GT domain mutations in the GT site on**
587 **glycosyltransferase activities.** Evaluation of the GalT activity (A) and GlcT activity (B) of
588 LH3 mutants compared to the wild-type. Each graph shows the enzymatic activity detected in
589 absence (i.e., “uncoupled”) or in presence of gelatin, used as acceptor substrate. The plotted
590 data are baseline-corrected, where the baseline was the background control. Error bars
591 represent standard deviations from average of triplicate independent experiments.

592

593 **Figure 3. Structural characterization of the LH3 Val80Lys mutant.** (A) Crystal structure
594 of the LH3 Val80Lys mutant in complex with UDP-glucose and Mn^{2+} . Electron density is
595 visible for the mutated lysine and the UDP donor substrate (green mesh, $2F_o-F_c$ omit electron
596 density map, contoured at 1.3σ). Catalytic residues shaping the enzyme cavity are shown as
597 sticks, Mn^{2+} is shown as purple sphere. Consistent with what observed in the crystal structure
598 of wild-type LH3, the glucose moiety of the donor substrate is not visible in the experimental

599 electron density. (B) Superposition of wild-type and Val80Lys LH3 crystal structures in
600 substrate-free (cyan and yellow, respectively) with UDP-glucose bound (marine and orange,
601 respectively) states. Notably, the conformations adopted by the side chain of Trp145 upon
602 ligand binding are consistent in the wild-type and in the mutant enzyme. As the glycoloop is
603 flexible in substrate-free structures, the side chains of Val/Lys80 are visible only in the in
604 UDP-sugar bound structures.

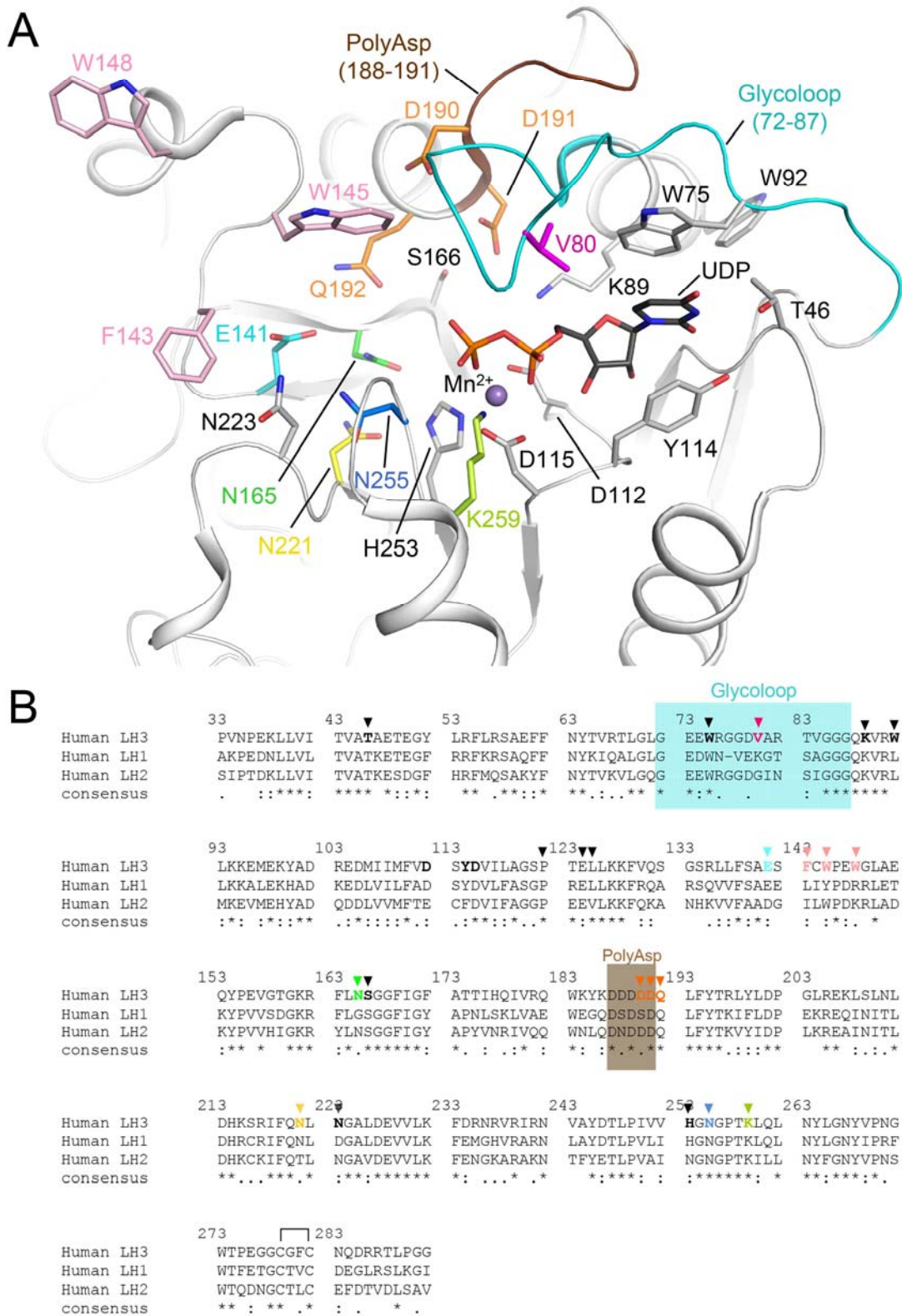
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606 **Figure 4. Characterization of UDP-sugar analogs.** (A) Thermal stability of LH3 wild-type
607 (solid green) using differential scanning fluorimetry (DSF) in presence of various Mn^{2+} and
608 several UDP-sugars. A prominent stabilization effect is achieved in presence of the biological
609 donor substrates UDP-galactose (solid blue), UDP-Glucose (solid purple) and free UDP (solid
610 black). A milder stabilization effect is also obtained with UDP-xylose (red dash) and UDP-
611 glucuronic acid (green dash). (B) Evaluation of GalT and GlcT enzymatic activities of LH3 in
612 the presence of increasing concentrations of UDP-GlcA or UDP-Xyl. (C) Crystal structure of
613 LH3 wild-type in complex with Mn^{2+} and UDP-glucuronic acid shows clear electron density
614 for UDP ($2F_o-F_c$ omit electron density maps, green mesh, contour level 1.2 σ). The
615 glucuronic acid (shown in yellow) can be modelled even if with partial electron density. (D)
616 Crystal structure of the LH3 Val80Lys mutant in complex with Mn^{2+} and UDP-glucuronic
617 acid. Whereas the UDP backbone can be modelled in the electron density (black sticks) ($2F_o-$
618 F_c omit electron density maps, green mesh, contour level 1.2 σ), in this case no electron
619 density is present for the glucuronic acid (shown in yellow). In addition, the portion of the
620 glycoloop containing the mutated lysine is flexible from residue 79 to 83 (shown as cyan
621 spheres). (E) Crystal structure of LH3 wild-type in complex with Mn^{2+} and UDP-xylose.
622 Similar to UDP-GlcA, UDP shows clear electron density ($2F_o-F_c$ omit electron density maps,

623 green mesh, contour level 1.2σ), whereas partial density is shown for the xylose moiety

624 (shown in pink).

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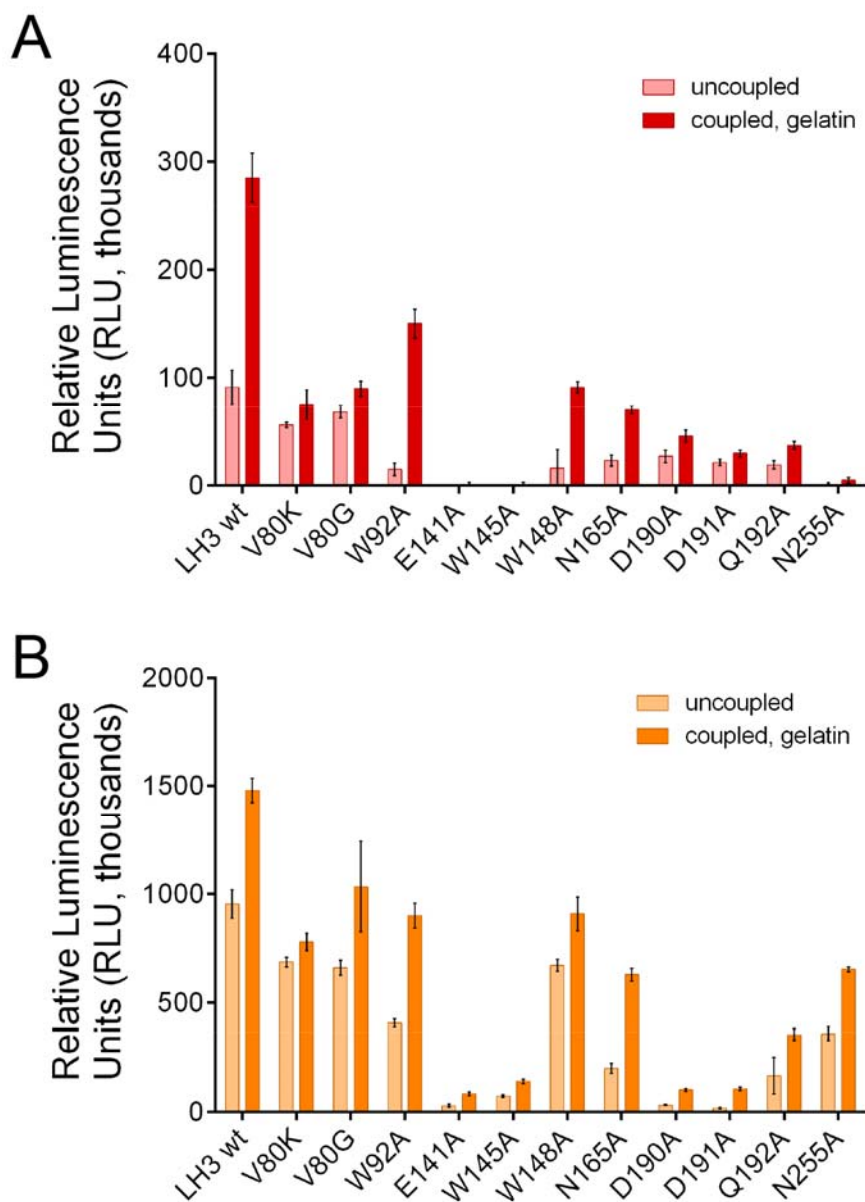


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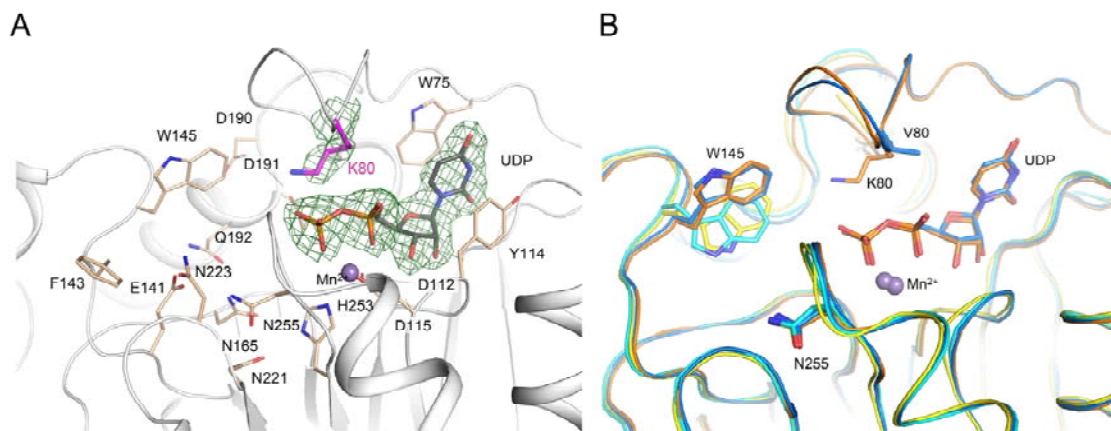
Figure 1



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Figure 2



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Figure 3

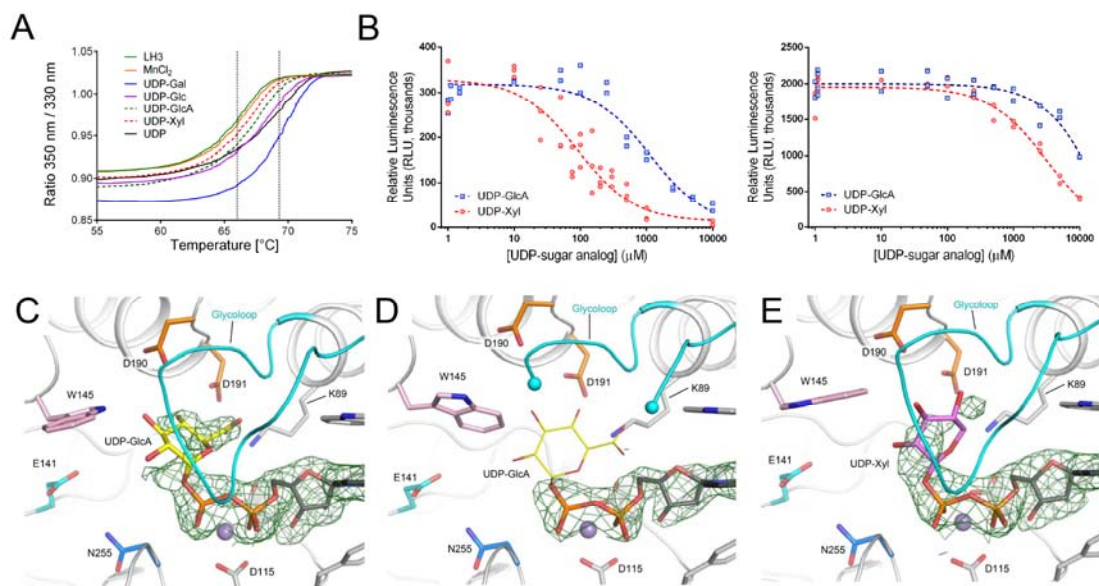


Figure 4

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635

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