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1 Activity based probes for functional interrogation of

$_2$ retaining β -glucuronidases

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Abstract

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- Humans express at least two distinct β-glucuronidase enzymes involved in disease: *exo*-acting β-
- 18 glucuronidase (GUSB), whose deficiency gives rise to mucopolysaccharidosis type VII, and endo-
- 19 acting heparanase (HPSE), implicated in inflammation and cancers. The medical importance of these
- 20 enzymes necessitates reliable methods to assay their activities in tissues. Herein, we present a set of
- 21 β-glucuronidase specific activity based probes (ABPs) which allow for rapid and quantitative
- 22 visualization of GUSB and HPSE in biological samples, providing a powerful tool for dissecting their
- 23 activities in normal and disease states. Unexpectedly, we find that the supposedly inactive HPSE
- 24 proenzyme proHPSE is also labeled by our ABPs, leading to surprising insights regarding structural
- 25 relationships between proHPSE, mature HPSE, and their bacterial homologs. Our results
- demonstrate the application of β -glucuronidase ABPs in tracking pathologically relevant enzymes,
- 27 and provide a case study of how ABP driven approaches can lead to discovery of unanticipated
- 28 structural and biochemical functionality.

Introduction

- 30 Retaining β -glucuronidases are enzymes responsible for hydrolytic cleavage of β -linked glucuronides
- 31 from polysaccharide and glycoconjugate molecules, with net retention of anomeric stereochemistry
- 32 at the released glucuronide. Humans express at least two major retaining β -glucuronidases: exo-
- 33 acting GUSB, responsible for cleaving β-linked glucuronides from the non-reducing end of diverse
- 34 glycosaminoglycans (GAGs) in the lysosome, and endo-acting heparanase (HPSE), specifically
- 35 responsible for breakdown of heparan sulfate (HS) in lysosomes and the extracellular matrix (ECM).
- 36 Both enzymes are strongly implicated in disease processes: deficiency of GUSB is the basis of the
- 37 autosomal recessive disease mucopolysaccharidosis type VII (MPSVII), also known as Sly syndrome¹⁻
- 38 ³, whilst HPSE overexpression is linked to a variety of pathologies including inflammation and cancer
- 39 metastasis^{4,5}.
- 40 Although both GUSB and HPSE possess β -glucuronidase activity, these enzymes are dissimilar at the
- 41 sequence level and fall under different families of the Carbohydrate Active enZymes (CAZy)
- 42 classification scheme⁶: GH2 for GUSB and GH79 for HPSE. Structurally, human GUSB is a large
- 43 homotetrameric assembly, with each protomer comprising a $(\beta/\alpha)_8$ barrel domain, a jelly roll domain
- and an Ig constant chain like domain⁷. In contrast, HPSE is a heterodimer comprised of 8 kDa and 50
- 45 kDa subunit chains, which fold to produce a $(\beta/\alpha)_8$ barrel flanked by a smaller β-sandwich domain⁸.
- 46 The mature HPSE heterodimer is formed by proteolytic removal of a 6 kDa linker peptide from a
- 47 single chain proenzyme proHPSE.
- 48 Given the importance of β -glucuronidases in human health and disease, a facile method to visualize
- 49 and quantitate their activity would be of great utility. We have previously reported the
- 50 development of activity based probes (ABPs) based upon the cyclophellitol aziridine scaffold, which
- 51 can be used to specifically detect enzymatic activity for a range of glycosidases⁹⁻¹⁴. These probes
- 52 provide valuable tools to rapidly determine enzyme activities within their native physiological
- 53 contexts
- 54 Herein, we unveil the synthesis of ABPs designed to selectively target and label retaining β-
- 55 glucuronidases. We demonstrate the utility of these probes in quantitating β -glucuronidase activity

- 56 in a range of cell and tissue samples, via both fluorescence and chemical proteomics approaches.
- 57 Unexpectedly, we find that a monosugar β -glucuronidase probe is sufficient to label not only exo-
- 58 acting GUSB, but also endo-acting HPSE, despite binding just one of multiple subsites within the
- 59 HPSE active site cleft. Furthermore, the supposedly inactive proHPSE proenzyme is also labeled by
- 60 ABPs, prompting us to investigate the nature of proHPSE 'inactivation' by its 6 kDa linker, and how
- this structure relates to other GH79 enzymes. Our results demonstrate a wide ranging potential of β -
- 62 glucuronidase ABPs as biological and biomedical tool compounds, and highlight the general power of
- ABPs for driving the discovery of novel biological insights¹⁵.

64 Results

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Glucuronidase specific inhibitor and probe design

- We have previously demonstrated cyclophellitol derived epoxides and aziridines to be powerful
- 67 mechanism based inhibitors for retaining β -glucosidases¹⁶, due to their ability to specifically label the
- 68 enzyme catalytic nucleophile, in a conformation resembling the covalent intermediate of glycoside
- 69 hydrolase reactions. (Fig. 1a, b)¹⁷. Inhibition is typically tolerant to functionalization at the ring
- 70 nitrogen of cyclophellitol aziridines, allowing fluorophore or biotin tagging to create inhibitor probes
- 71 which can label specific glycosidases within complex biological mixtures¹⁸.
- 72 Conceptually, we envisioned that β -glucuronidase specific ABPs could be accessed from
- 73 cyclophellitol by oxidation at the C6 equivalent position, to emulate the carboxylate of glucuronic
- 74 acid (GlcUA). ABPs **1–4** are composed of such a β-glucuronide configured cyclophellitol aziridine,
- bearing a spacer from the aziridine nitrogen terminating in BODIPY-FL, a BODIPY-TMR analog, Cy5, or
- 76 biotin respectively. Alongside these functionalized ABPs we also prepared azide substituted ABP 5
- 77 (the precursor of 1–4), unsubstituted aziridine 6 and cyclophellitol-6-carboxylate 7. (Fig. 1c;
- 78 structures of additional compounds 8–16 used in this study are shown in Supplementary Results,
- 79 **Supplementary Fig. 1**).

GlcUA ABPs target β-glucuronidases in vitro and in situ

- 81 To assess the potency of our β-glucuronidase ABPs in vitro, we first turned to the exo-acting GH79 β-
- 82 glucuronidase AcGH79 from Acidobacterium capsulatum, whose activity is readily followed using the
- 83 fluorogenic substrate 4-methylumbelliferyl-glucuronic acid (4MU-GlcUA)¹⁹. All compounds tested
- 84 were effective inhibitors of AcGH79, with apparent IC50s in the low to sub nM range (Table 1 left
- 85 panel). Core ABP 'warhead' 6 inhibited AcGH79 with apparent IC50 of ~5 nM. This was potentiated
- 86 by further functionalization: apparent IC50 of Cy5 substituted ABP 3 was ~1 nM, whilst 1, 2, 4 and 5
- 87 were all sub-nanomolar inhibitors of AcGH79. Apparent IC50 for epoxide **7** was ~34 nM, consistent
- 88 with lower reactivity of the epoxide moiety compared to aziridines.
- 89 Kinetic parameters for inhibition of AcGH79 were determined using a continuous assay, whereby
- 90 substrate and inhibitor react with enzyme simultaneously (Supplementary Note 1)²⁰, allowing us to
- 91 derive a combined inhibition parameter k_i/K_i for all ABPs tested. k_i/K_i values largely reflected the
- 92 trend seen with IC50s, with the activity of core aziridine 6 potentiated by further functionalization,
- 93 and epoxide 7 substantially less active than aziridines (Table 1 middle panel, Supplementary Fig. 2).

- 94 Finally, we tested the ability of our probes to inhibit β -glucuronidases in live fibroblast cells. *In situ*
- 95 apparent IC50s were determined for ABPs 2 and 3 to be in the low μM range (~1.7 and ~1.8 μM
- 96 respectively). We were unable to determine in situ apparent IC50s for 1 or 4–7, likely reflecting a
- 97 limited ability of these compounds to permeate the cell membrane (Table 1 right panel).
- 98 Fluorescent labeling of AcGH79 by ABP 1 was readily visualized after running on SDS-PAGE, and
- 99 could be blocked by competition with 2–7, 4MU-GlcUA, or iminosugar 8. Labeling also was abolished
- 100 by SDS denaturation of protein, in line with a mechanism-based mode of action requiring
- 101 catalytically competent enzyme (Supplementary Fig. 3a).
- 102 To dissect the mechanistic mode of action of our probes, we obtained crystal structures of 5 in
- 103 complex with wild type AcGH79, and an inactive AcGH79(E287Q) nucleophile mutant
- 104 (Supplementary Fig. 3b, c). Both complexes showed a single molecule of 5 bound within the active
- site of AcGH79, with no labeling of off-target residues. In wild type AcGH79, reacted **5** was observed
- bound via C1 to the enzyme nucleophile (Glu287) in a 4C_1 conformation, making identical non-
- 107 covalent contacts as previously observed for GlcUA or 2F-GlcUA¹⁹. In the AcGH79(E287Q) mutant, 5
- 108 occupied the same active site position, but was instead found to adopt a 4H_3 conformation, due to
- restricted rotation across the C1-C7 bond imposed by the aziridine. Notably, the ⁴H₃ conformation
- observed for unreacted 5 is the same as that postulated for oxocarbenium-like transition states of
- 111 retaining β -glycosidase substrates during hydrolysis (**Supplementary Fig. 3d**)²¹. The high affinities of
- 112 cyclophellitol derived ABPs for their target enzymes may thus be in part due to their conformational
- 113 mimicry of this transition state²².

ABP profiling reveals GUSB and HPSE as probe targets

- 115 To determine the targets of retaining β -glucuronidase ABPs in complex biological samples, human
- 116 splenic lysates (which we have previously shown to express a range of glycosidases 10-12) were treated
- 117 with one or more ABPs, resolved by SDS-PAGE, and labeled proteins visualized by fluorescent
- scanning (the typical ABP workflow is shown in **Supplementary Fig. 4**).
- 119 Several fluorescent bands were observed in samples treated with Cy5 ABP 3 which were absent in a
- mock (DMSO) control, and which could be competed for by biotin ABP 4 (Fig. 2a). Based on
- 121 literature reports, we tentatively assigned the prominent double bands at ~78–80 kDa as full length
- and C-terminal truncated isoforms of GUSB²³, and the lowest molecular weight band as the ~64 kDa
- isoform of GUSB²⁴; these bands were also identified by an anti-GUSB western blot (**Supplementary**
- 124 Fig. 5a). A band at ~60 kDa did not correspond with any known glucuronidases but could be
- abrogated by pretreatment with β -glucosidase ABP **9**, suggesting this was the lysosomal acid β -
- 126 glucosidase GBA, which is specifically labeled by **9**¹². Correspondingly, this ~60 kDa band was also
- absent in splenic lysates from patients with Gaucher disease, which is characterized by lack of GBA
- 128 activity.

- 129 To unambiguously establish the targets of our ABPs, we carried out a set of chemical proteomics
- 130 experiments using liquid chromatography coupled to tandem mass spectrometry (LC-MS/MS).
- 131 Lysates from normal or Gaucher spleens were incubated with biotin ABP 4, with or without
- 132 pretreatment using 9. Labeled proteins were then pulled down using streptavidin beads, and
- processed using a standard proteomics workflow, utilizing both 'in-gel' and 'on-bead' digest

| 12/ | protocols (Supplemen | tary Fig. 4) T | he 'on head' d | igest protocol was | also applied to human |
|-----|------------------------------|----------------|----------------|--------------------|-----------------------|
| L34 | protocois (Supplemen | Larv rig. 4). | ne on beau u | igest protocol was | also applied to numan |

- fibroblast lysates, in order to assess the efficacy of ABP pulldown in another tissue type.
- 136 Proteomic profiling using the 'in-gel' digest protocol identified GUSB as the predominant splenic
- protein labeled by 4, with particular enrichment in bands corresponding to those previously
- 138 identified by 3 (Fig. 2b). As with fluorescent labeling, we detected GBA at ~60 kDa, which was
- 139 abrogated by pretreatment with 9 and reduced in Gaucher spleen. 'In-gel' proteomic results were
- 140 largely mirrored by those from the 'on bead' protocol, which showed GUSB to be the most abundant
- 141 glycosidase after pulldown from spleen, and also highly enriched after pulldown from fibroblast
- 142 (Supplementary Data Set 1). Covalent modification of the GUSB nucleophile (Glu540) by 4 was
- 143 directly characterized by MS/MS fragmentation of the 13 amino acid peptide containing this residue
- 144 (Supplementary Fig. 5b).
- 145 The high sensitivity of proteomic profiling enabled detection of two glycosidase enzymes in splenic
- 146 lysates not observed by fluorescent labeling with **3**: β-galactosidase GLB1, and endo β-glucuronidase
- 147 HPSE. Both enzymes were substantially less abundant in pull-down fractions compared to GUSB, as
- 148 estimated by their exponentially modified protein abundance index (emPAI) scores (Supplementary
- 149 Fig. 5c). We theorized that GLB1 was likely a weak non-specific target of 4, as it showed the lowest
- emPAI of all detected glycosidases despite robust transcriptional expression reported in spleen
- (Supplementary Fig. 5d)²⁵, and strong histochemical staining of β -galactosidase activity in
- mammalian splenic tissues²⁶. In contrast, HPSE is predicted to be poorly expressed in spleen by
- transcriptomics, yet showed a higher emPAI score than GLB1 after pull-down by 4. These
- observations suggested HPSE was a bona fide target of our ABPs, despite the inability of the
- 155 monosugar probes to make a full complement of interactions within the HPSE endo-acting substrate
- cleft, which normally accommodates at least a trisaccharide²⁷.

β-glucuronidase ABPs label endogenous HPSE and proHPSE

- 158 Following the surprising discovery that splenic HPSE was labeled by 4, we re-examined fluorescent
- 159 labeling of HPSE in cells and tissues expressing higher levels of HPSE. In the first instance, we induced
- 160 HPSE overexpression in HEK293T cells, and probed harvested lysates at set intervals. Two plasmids
- 161 were tested: pGEn1-HPSE, encoding for N-His-Strep-TEV tagged proHPSE, and pGEn2-HPSE, encoding
- 162 for N-His-Avitag-eGFP-TEV tagged proHPSE; HEK293T cells subsequently process these proHPSE
- 163 precursors to mature HPSE.

- 164 Using 3, we tracked increasing expression of a band at ~50 kDa, corresponding to the mass of the
- large HPSE subunit, which contains the nucleophile Glu343 (Fig. 3a)²⁸. Unexpectedly, we also
- 166 detected bands at ~75 kDa for pGEn1-HPSE and ~100 kDa for pGEn2-HPSE, corresponding to the
- 167 masses of proHPSE chains expressed by these plasmids. Western blotting using an anti-HPSE
- 168 antibody confirmed these bands to be proHPSE, indicating that 3 also labeled the supposedly
- inactive proenzyme. Initial comparison of band intensities from ABP labeling using 3 vs. western
- 170 blotting suggested that 3 labeled mature HPSE with greater efficiency than it labeled proHPSE.
- 171 However, the opposite trend was observed when using purified recombinant proHPSE
- 172 (Supplementary Fig. 6a), suggesting that the majority of proHPSE overexpressed by HEK293T cells
- 173 was likely inactive.

- 174 HPSE maturation is thought to be mediated by the cysteine protease cathepsin L (CTSL), through
- multiple proteolytic cleavages of the 6 kDa linker peptide²⁹. We tested the effect of CTSL inhibitors
- 176 CAA0225³⁰ (**10**), Z-FY(tBu)-DMK³¹ (**11**) and leupeptin on proHPSE maturation in HEK293T cells by
- incubating cells transfected pGEn1-HPSE with CTSL inhibitors for 2 d and labeling lysates with 3. In
- 178 inhibitor treated cells, we observed modest but dose dependent accumulation of a band ~5 kDa
- 179 below the principal proHPSE band, corresponding to loss of the unstructured N-His-Strep-TEV tag
- followed by blockade of further proteolysis, in line with the role of CTSL in HPSE maturation
- 181 (Supplementary Fig. 6b). However, accumulation of mature HPSE was still detected in the presence
- 182 of all inhibitors tested, suggesting either incomplete CTSL inhibition, or the presence of non-
- cathepsin mediated HPSE maturation pathways also utilized by HEK293T cells.
- 184 We also tracked internalization and processing of proHPSE to mature HPSE in fibroblasts, a process
- 185 which may be utilized by cancer cells to increase their own levels of HPSE: by capturing and
- internalizing extracellular proenzyme³². Within 90 minutes of introducing proHPSE into culture
- 187 medium, internalization and processing of HPSE by fibroblasts was detectable using 3 (Fig. 3b).
- 188 These experiments demonstrate the ability of β-glucuronidase ABPs to detect and track key
- 189 biological processes such as HPSE internalization and maturation.
- 190 Lastly, we reattempted fluorescent labeling of endogenous HPSE in human tissues. As splenic HPSE
- 191 expression was below the fluorescent detection limit, we turned to platelets, which are known to
- 192 contain high levels of mature HPSE³³. Using **3**, we observed labeling of a band at ~50 kDa in platelet
- 193 lysates corresponding to HPSE, in addition to the same GUSB bands previously detected in spleen
- 194 (Fig. 3c). Comparison of fluorescence and western blotting intensities between platelet HPSE and a
- 195 200 fmol recombinant standard suggested fluorescent sensitivity for recombinant HPSE to be in the
- 196 fmol range, somewhat more sensitive than western blotting in our hands. Fluorescent HPSE
- 197 detection in platelet lysates was slightly less sensitive, possibly due to the presence of competing
- 198 protein targets or inactive HPSE in situ. 10 nM of 3 was sufficient to produce a detectable HPSE
- signal in platelets after 30 minutes (**Supplementary Fig. 6c**). Labeling of HPSE, but not GUSB, was
- also improved by the addition of NaCl (**Supplementary Fig. 6d**). Optimum labeling of HPSE by **3** was
- achieved at pH 4.5–5.0, consistent with literature reports of its optimum pH for enzymatic activity³⁴.
- 202 In contrast, optimum pH for labeling GUSB was higher than expected at pH ~5.5–6.0 (Fig. 3d),
- 203 compared to its reported optimum for activity at the lysosomal pH ~4.5³⁵. This unexpected pH of
- 204 GUSB ABP labeling may be due to facile aziridine ring opening occurring independently of a
- 205 protonated acid/base residue³⁶. However, optimum labeling of GUSB at a non-lysosomal pH
- 206 presents its own serendipitous advantages, allowing both GUSB and HPSE to be analyzed either
- jointly or independently of each other through modulation of labeling pH.

Competitive ABP labeling identifies HPSE specific inhibitors

- 209 Because ABPs can detect a complete complement of enzymes in a cellular/environmental sample,
- 210 competitive ABP labeling provides a powerful tool to assess inhibitor efficacy and specificity within a
- 211 single experiment³⁷. We sought to establish whether GlcUA ABPs could be used for the assessment
- 212 of enzyme specific inhibitors, by testing platelet labeling at pH 5.0 (where both GUSB and HPSE
- react) in the presence of a set of known inhibitors.

- 214 In the presence of the monosaccharide-like β -glucuronidase inhibitor siastatin B³⁸, both GUSB and
- 215 HPSE labeling were abrogated in a dose-dependent manner, demonstrating the ability of this

- 216 molecule to outcompete ABP binding in both endo- and exo- acting enzymes. Using quantitated
- band intensities, IC50s for GUSB and HPSE labeling inhibition were measured to be $^{\sim}3.3~\mu M$ and $^{\sim}6.7$
- 218 μM respectively, indicating slightly greater affinity for GUSB by siastatin B (Fig. 4a).
- 219 In contrast, HPSE labeling in platelets was selectively inhibited by competition with heparin (12, IC50
- 220 ~0.17 mg/mL, Fig. 4b), a large polysaccharide which cannot be accommodated by the exo- acting
- 221 active site of GUSB. Selective inhibition was also observed upon competition with HS (13), the
- 222 substrate of HPSE, albeit with slightly lower potency (Fig. 4c, IC50 ~0.50 mg/mL). Negligible
- 223 inhibition was observed for N-Acetyl-O-desulfated heparin (14) (Supplementary Fig. 7a), highlighting
- the importance of sulfation for interactions between heparin/HS and HPSE. A lower degree of
- 225 sulfation in HS vs. heparin may partly account for its slightly weaker abrogation of ABP labeling³⁹.
- 226 We next tested labeling inhibition by GAGs with different linkages and sulfation patterns to heparin
- 227 and HS. Hyaluronic acid (15) and chondroitin sulfate (16) both showed no inhibition of either HPSE
- 228 or GUSB labeling at concentrations sufficient for inhibition by heparin/HS (Supplementary Fig. 7b, c),
- highlighting the critical role of sugar linkage and sulfation in interactions between GAGs and HPSE.
- Taken together, these assays provide proof of principle that GlcUA ABPs are amenable for use in a
- 231 competitive format, to assess inhibition of specific β -glucuronidases within a mixture of related
- 232 activities.

Structural basis of HPSE and proHPSE ABP labeling

- 234 To investigate how efficient labeling of endo-acting HSPE was achieved by a monosugar ABP, we
- 235 obtained crystal structures of both wild type and nucleophile mutant (E343Q) HPSE in complex with
- 236 ABP 5. Complexes of HPSE with 5 were similar to those obtained with AcGH79, showing a single
- 237 molecule of probe occupying the -1 subsite of the HPSE substrate binding cleft (nomenclature
- according to Ref. 40) in reacted 4C_1 (wild type) or unreacted 4H_3 (mutant) conformations
- 239 (Supplementary Fig. 8a). The network of interactions made to the probe was highly similar between
- 240 AcGH79 and HPSE, with the primary difference being a lack of interaction by HPSE to O4 of the
- 241 probe, due to extension of its natural HS substrate towards this position (Supplementary Fig. 8b). A
- 242 —1 subsite C6 carboxylate recognition motif, comprising 3 H-bonds from a tyrosine and two
- 243 consecutive backbone amides, is highly conserved in GH79 β-glucuronidases (Tyr 334, Gln293-
- 244 Gly294 in AcGH79; Tyr391, Gly349-Gly350 in HPSE; Tyr 302, Gly261-Gly262 in the recently
- 245 characterized heparanase from *Burkholderia pseudomallei*)⁴¹. This strong network of H-bonds to C6
- 246 carboxylate likely offsets the absence of only a single H-bond to O4 of the ABP in HPSE compared to
- 247 AcGH79, thus rationalizing robust labeling of HPSE by a monosaccharide probe that only occupies a
- single subsite within its extensive binding cleft. Additional binding affinity may also derive from the
- transition state like 4H_3 conformation adopted by unreacted ABPs.
- 250 We next sought to solve the structure of proHPSE, in order to characterize the basis of its
- 251 'inactivation' by the 6 kDa linker peptide, and to determine how β-glucuronidase ABPs are able to
- 252 circumvent this. Herein, we report the first crystal structure of proHPSE in both apo and ABP
- 253 complexed forms, which together with previously reported HPSE structures completes a structural
- 254 characterization of the HPSE maturation process.
- The proHPSE structure was similar to that of mature HPSE (RMSD: 0.52 Å over $451 \text{ C}\alpha$), with the
- 256 same $(\beta/\alpha)_8$ and β -sheet domains clearly discernible. The 6 kDa linker (110-157) forms a large helical

257 domain which sits directly 'above' the active site cleft, blocking access to the bulky HS substrates of

258 HPSE. The final loop of the linker leading into the 50 kDa subunit (His155-Lys159) is substantially

more disordered than the rest of the protein, as evidenced by higher B-factors for these residues in

the crystallographic model (Supplementary Fig. 9a, b). Mutation studies have established Tyr156 of

- the proHPSE linker to be critical for recognition by CTSL in the first step of HPSE maturation⁴².
- 262 Disorder of the His155-Lys159 loop allows for unencumbered CTSL access to Tyr156 without
- 263 disrupting preexisting secondary structures, consistent with the important role of Tyr156 in HPSE
- 264 maturation.

- 265 Unexpectedly, steric blockage by the linker peptide was found to be incomplete in proHPSE, leaving
- 266 a 'binding pocket' on the protein surface containing exposed catalytic nucleophile and acid/base
- residues, similar to the *exo*-acting active site of AcGH79 (**Fig. 5a**). When compared in a sequence
- 268 alignment, the proHPSE linker corresponds to a loop in AcGH79, which forms the 'rear' face of its
- active site pocket, suggesting that whilst this sequence has expanded in the human enzyme,
- 270 proHPSE still retains some structural characteristics reminiscent of a GH79 exo-glycosidase
- 271 (Supplementary Fig. 8c). ABP 5 was found to bind to this proHPSE 'active-site pocket' in a
- 272 configuration identical to that observed for HPSE. The O4 proximal position, vacant in HPSE, was
- occupied by His155 in proHPSE, contributed by the linker, which blocks off extension towards this
- position by HS substrates (**Fig. 5b, c**). The disordered proHPSE His155-Lys159 loop was slightly
- 275 displaced upon binding 5 (~1.54 Å for Tyr156 Cα), due to steric clashes with the bound ABP
- 276 (Supplementary Fig. 9c).
- 277 As with mature HPSE, proHPSE was inactive against the artificial fluorogenic substrate 4MU-GlcUA,
- 278 indicating it does not possess any additional exo-glucuronidase activity against this substrate which
- 279 is lost upon maturation (Supplementary Fig. 8d). To assess the accessibility of the proHPSE 'pocket'
- 280 compared to mature HPSE, we conducted competitive ABP experiments against recombinant
- proHPSE and HPSE using **3**. As with platelets, siastatin B inhibited ABP labeling of both pro- and
- 282 mature HPSE (Supplementary Fig. 10a), indicating it could efficiently occupy the 'binding pocket' of
- 283 proHPSE as well as HPSE. In contrast, heparin only inhibited labeling of HPSE, albeit with lower
- 284 efficacy than seen in platelets, due to more facile labeling of the recombinant enzyme.
- 285 Unexpectedly, proHPSE labeling was slightly increased at moderate heparin concentrations
- 286 (Supplementary Fig. 10b). Finally, we tested the ability of GlcUA to inhibit labeling of HPSE and
- 287 proHPSE. No substantial inhibition of either proHPSE or HPSE labeling was observed at up to 20 mM
- 288 GlcUA (**Supplementary Fig. 10c**), suggesting that GlcUA cannot occupy the active site of (pro)HPSE
- with sufficient affinity to prevent binding and reactivity of an ABP. Further subsite interactions may
- 290 be required for binding of simple glucuronides to (pro)HPSE.
- 291 It has previously been demonstrated that proHPSE uptake by cells is a HS dependent process, and
- can be disrupted by addition of exogenous heparin⁴³. To investigate possible roles for the proHPSE
- 293 'binding pocket' in proHPSE uptake and maturation, we prelabeled recombinant proHPSE with
- either untagged ABP 6, fluorescent ABPs 1 or 3, or a mock DMSO control, and examined its uptake
- 295 by fibroblasts at 90 or 180 min. In all cases, prelabeled proHPSEs were taken up and processed to
- 296 mature HPSE, as evidenced by western blot and fluorescence of internalized 1 or 3 (Supplementary
- 297 **Fig. 11**). These data indicate that the proHPSE 'pocket' does not participate in the HS interactions
- involved in cellular uptake, and that occupation of the proHPSE 'pocket' does not inhibit HPSE
- 299 maturation.

Discussion

The important role of β-glucuronidases in human biology is highlighted by the pathologies
 associated with aberrant expression of these enzymes. Lack of GUSB activity leads to accumulation
 of glucuronide-containing GAGs within lysosomes in MPSVII (Sly Syndrome). Conversely,

overexpression of HPSE leads to aberrant breakdown of HS in the ECM, causing increased cancer

305 growth and metastasis. Accurate tracking of β -glucuronidase activities is an essential prerequisite

for fully understanding their role in both physiological and disease states.

of previously uncharacterized β -glucuronidase activities in biological samples.

Here we have reported the design and application of novel β -glucuronidase configured ABPs, and demonstrated their broad utility for interrogating activities of these enzymes. We show that ABP profiling is a viable method to assay β -glucuronidase activity in a variety of samples, ranging from recombinant proteins, to complex cell, tissue and organ lysates. Fluorescent labeling provides a facile method for probing β -glucuronidases in tissues with sufficient expression, allowing for tracking of processes such as proenzyme uptake and processing, and how these are affected by biological or pharmacological perturbation. In tissues with lower enzyme abundance, we have demonstrated detection of β -glucuronidases using a proteomic approach, which is also applicable for the discovery

Use of ABPs provides several advantages over more traditional methods to quantitate glycoside hydrolase activities. Compared to techniques such as western blotting, ABPs specifically detect active enzymes, rather than an entire protein complement which may include misfolded or inactive isoforms. Whilst fluorometric or colorimetric assays also provide assessments of enzyme activity, they cannot distinguish between overlapping activities in complex mixtures, which arise from several enzymes or enzyme isoforms active on the same substrate. Indeed, many carbohydrate processing enzymes are processed from precursors into one or more isoforms with differing activities 23,28,44,45.

ABP profiling allows for multiple activities to be visualized and their responses to perturbation or

inhibition to be individually assessed in situ.

Many endo-glycosidases such as HPSE are inactive in traditional activity assays, necessitating the use of expensive specialized substrates and/or cumbersome assay procedures to follow their activities. The discovery that aziridine ABPs label HPSE paves the way for more rapid and practicable methods to assess the activity of this enzyme, and may inspire development of probes to assay other endo-glycosidases. Whilst this current generation of β -glucuronidase ABPs shows some off-target effects against GBA and GLB1, limiting their use in diagnostic applications, further optimization based upon the crystal structures of HPSE (and proHPSE) may lead to improved probes with increased potency and specificity. Optimization efforts will be aided by the use of competitive ABP techniques, which we have demonstrated to be a viable method for assessing selective inhibitors of individual β -glucuronidases.

ABPs also provide powerful tools for characterization of novel enzyme activities, which may escape detection in traditional biochemical experiments. The use of an ABP driven approach in this study lead us to the surprising observation that the HPSE precursor proHPSE is in principle catalytically competent, an entirely unanticipated outcome based on previous studies. We have reported the first structural views of proHPSE, illustrating how its 6 kDa linker restricts access to the active site cleft for HS substrates. This linker does not entirely block access to the catalytic residues of

| 341 | proHPSE, but instead contributes to the formation of an exo-glycosidase like 'binding pocket', which |
|-----|---|
| 342 | can accommodate smaller molecules. It remains to be determined whether this proHPSE 'pocket' is |
| 343 | simply a structural relic from evolutionary expansion of an ancestral GH79 active site loop, or if there |
| 344 | are bona fide endogenous substrates which are hydrolyzed by proHPSE. |
| 345 | In conclusion, we have presented a set of ABPs for functional interrogation of β -glucuronidases in |
| 346 | their native contexts. The application of ABP methodology to carbohydrate processing enzymes |
| 347 | provides a powerful set of tools to study the activity of these key enzymes, and will contribute |

Accession codes

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- 350 Coordinates and structure factors have been deposited in the Protein Data Bank under accession
- 351 codes 5G0Q (AcGH79(wt)-5 complex), 5L77 (AcGH79(E287Q)-5 complex), 5L9Y (HPSE(wt)-5

towards our understanding of fundamental processes in glycobiology.

352 complex), 5L9Z (HPSE(E343Q)-5 complex), 5LA4 (apo proHPSE), 5LA7 (proHPSE-5 complex).

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360 Author contributions

- 361 L.W., J.M.F.G.A., H.S.O. and G.J.D. conceived and designed the experiments. J.J., M.A., W.D. and
- 362 C.v.E. carried out synthesis of probes, with guidance from G.A.v.d.M. and J.D.C.C.. L.W. and Y.J.
- carried out protein expression and structural studies on enzyme-probe complexes. J.J., L.W., W.W.K.
- and C-L.K. carried out gel labeling experiments. J.J. and B.I.F. carried out proteomics experiments. C-
- 365 L.K. and W.W.K. determined IC50 and kinetic parameters for ABPP inhibition. M.v.E. obtained tissue
- 366 samples. L.W., J.J., H.S.O., and G.J.D. wrote the manuscript with input from all authors.

Competing financial interests

368 The authors declare no competing financial interests.

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Methods and Supplementary Information

- 372 Supplementary results containing Supplementary Tables 1 and 2, Supplementary Figures 1–12,
- 373 Supplementary Notes 1 and 2 and Supplementary Data Set 1 are available in the online version of
- this paper.

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- Figure 1 Concept and design of β-glucuronidase targeting ABPs. **a** Generalized schematic of double-displacement mechanism employed by retaining β-glycosidases. **b** Mechanism based inhibition by cyclophellitol derived ABPs. **c** Structures of cyclophellitol, cyclophellitol aziridine, and β-glucuronidase specific ABPs used in this study. Numbering of atomic positions is shown on cyclophellitol.
- 488 Figure 2 ABP labeling of retaining β-glucuronidases in human spleen lysates. a Three isoforms of 489 GUSB are fluorescently labeled by Cy5 ABP 3 in human wild type spleen, along with off target 490 labeling of the β -glucosidase GBA. Labeling of these proteins by 3 can be competed for by biotin ABP 491 4. GBA labeling in wild type spleen is also specifically competed for by 9, and is absent in lysates 492 from Gaucher spleen. b Silver stained SDS-PAGE gel of proteins captured from human wild type 493 spleen by labeling with 4 (with or without competition by 9) followed by streptavidin pulldown. 494 Glycosidase enzymes identified in each gel band by proteomic profiling are listed. Full proteomics 495 datasets for proteins identified by ABP pulldowns are available in Supplementary Data Set 1.
- Figure 3 Human HPSE is readily visualized by fluorescent β-glucuronidase ABPs. a Induced overexpression of HPSE and proHPSE in HEK293T cells can be tracked by ABP 3. Fluorescent labeling by 3 correlates with bands from western blotting using an anti-HPSE antibody. b ABP tracking of uptake and processing of proHPSE to HPSE by fibroblast cells. c Endogenous HPSE in human platelets can be labeled by 3, along with the same GUSB bands as observed in spleen. d pH dependence of HPSE and GUSB labeling in platelet lysates, demonstrating how general or specific enzyme labeling can be achieved by modulating pH.
 - **Figure 4** General and endo-specific inhibition of β-glucuronidases assessed by competitive ABP profiling. **a** Monosugar like β-glucuronidase inhibitor siastatin B can be accommodated *exo* and *endo* acting β-glucuronidase active sites, and competes out ABP **3** labeling of both GUSB and HPSE. **b** Polysaccharide heparin (**12**) only inhibits ABP labeling of HPSE, due to its inability to interact with the *exo* configured active site of GUSB. **c** Selective HPSE inhibition is also achieved by heparan sulfate (**13**). Competitive ABP gels shown are representative of three technical replicates. Plots are mean values \pm s.d. (N=3) for quantitated HPSE and GUSB fluorescent band intensities, normalized to band intensities in the no inhibitor control lane. For all plots, quantitated GUSB fluorescence is a sum of the three assigned bands. n.d.: not determined.
- 512 Figure 5 3-dimensional structure of proHPSE, and its active site interactions with ABP 5. a Ribbon 513 and surface diagram of proHPSE, demonstrating steric blockage of the HPSE binding cleft by the 6 514 kDa linker. An exposed 'pocket' in proHPSE can still interact with small molecules such as 5 515 (highlighted pink for clarity). **b** ABP **5** in complex with proHPSE within its 'binding pocket'. The O4 516 position, where HS substrates would extend in mature HPSE, is blocked by His155, contributed by 517 the linker (colored in green). Density is REFMAC maximum-likelihood/ σ_A weighted 2Fo-Fc contoured 518 to 0.38 electrons/Å³. c Schematic of H-bonding interactions between reacted 5 and proHPSE active 519 site residues. Interactions are identical to those observed for the mature enzyme (Supplementary 520 Fig. 8a, b), except for His155 proximal to O4 of the probe. (nuc.: nucleophile; a/b: acid base).

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Table 1 Apparent IC50 values for *in vitro* and *in situ* inhibition of β -glucuronidase activity by ABPs, and kinetic parameters for inhibition of AcGH79 by ABPs. Data are mean values \pm s.d. (N=3) from three biological replicates.

| Compound | In vitro AcGH79 apparent IC50 (nM) | Kinetic Parameters (AcGH79) k _i /K, (μM ⁻¹ min ⁻¹) | In situ fibroblast apparent IC50 (μM) |
|----------|------------------------------------|---|--|
| 1 | 0.6±0.2 | 25.0±0.7 | >15 |
| 2 | 0.8±0.2 | 18.2±0.9 | 1.7±0.6 |
| 3 | 1.1±0.1 | 14.0±0.8 | 1.8±0.4 |
| 4 | 0.4±0.02 | 5.5±0.2 | >15 |
| 5 | 0.1±0.01 | 18.8±0.7 | >15 |
| 6 | 4.6±0.03 | 3.5±0.2 | >15 |
| 7 | 33.4±3.1 | 0.49±0.05 | >15 |

526 Methods 527 **Chemical probes and inhibitors** 4MU-GlcUA, Leupeptin, Siastatin B, Hyaluronic Acid (15) and Chondroitin Sulfate (16) were obtained 528 from Sigma Aldrich. CTSL inhibitors CAA0225 (10) and Z-FY(tBu)-DMK (11) were obtained from 529 Merck. Heparin (12) and Heparan Sulfate (13) were obtained from Iduron. N-Acetyl-O-desulfated 530 Heparin (14) was obtained from Dextra. Cyclophellitol, cyclophellitol aziridine⁴⁶ and ABP 9¹⁰ were 531 532 synthesized according to described procedures. Syntheses of compounds 1-8 are described in 533 Supplementary Note 2. 534 Tissue and cell samples 535 Gaucher patients were diagnosed on the basis of reduced GBA activity and demonstration of an 536 abnormal genotype. Spleens from a normal subject and a patient suffering from type 1 Gaucher disease were collected after splenectomy and frozen at -80 °C until use. Platelets were collected 537 538 from healthy donors, using EDTA as the anti-coagulant. Platelet rich plasma (PRP) was prepared by centrifugation at 100 g for 20 min at 22 $^{\circ}\text{C}$ to remove red and white blood cells. Platelets were 539 540 isolated from PRP by centrifugation at 220 g for 10 min at 22 °C, and frozen at -80 °C until use. 541 Approval for tissue collection was obtained from the Academisch Medisch Centrum (AMC) and 542 University of York medical ethics committees. Informed consent was obtained from all donors. 543 Primary human fibroblasts (CC-2511) were obtained from Lonza. HEK293T cells (ATCC-CRL-3216) 544 were obtained from the American Type Culture Collection (ATCC). Sf21 and High Five cells for 545 protein production were obtained from Invitrogen. Cells were used as obtained from the supplier 546 without further authentication. All cells used tested negative for mycoplasma contamination. 547 All tissue lysates were prepared in KPI buffer (25 mM potassium phosphate [pH 6.5], supplemented 548 with $1 \times$ cOmplete protease inhibitor cocktail (Roche)). Cells/tissues were homogenized with a silent 549 crusher S equipped with a type 7 F/S head (30,000 rpm, 3 × 7 sec) on ice. Lysate protein 550 concentrations were determined with a Qubit 2.0 Fluorometer (Invitrogen) or Bradford assay using 551 BSA as a standard. Lysates were stored in aliquots at -80 °C until use. 552 Recombinant protein cloning, expression and purification 553 AcGH79 554 The coding sequence of AcGH79 with an N-terminal 6×His tag was cloned into pET28a (Novagen), 555 which was used to transform E. coli BL21-Gold(DE3) (Agilent). Transformants were grown at 37 °C in 556 LB media containing 50 μ g/mL kanamycin to an OD₆₀₀ of 0.8, induced by addition of 1 mM isopropyl 557 β -D-1-thiogalactopyranoside, and protein production carried out out at 25 °C for 12 h. Harvested 558 cells were resuspended in 50 mL AcGH79 HisTrap buffer A (20 mM HEPES [pH 7.0], 200 mM NaCl, 5 559 mM imidazole), lysed by sonication, and lysate clarified by centrifugation at 12000 g. Supernatant

containing AcGH79 was filtered before loading onto a HisTrap 5 mL FF crude column (GE Healthcare) pre-equilibrated with AcGH79 HisTrap buffer A. The loaded HisTrap column was washed with 10

column volumes (CV) of AcGH79 HisTrap buffer A, before eluting with AcGH79 HisTrap buffer B (20

mM HEPES [pH 7.0], 200 mM NaCl, 400 mM imidazole) over a 20 CV linear gradient.

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- 564 Fractions containing AcGH79 were pooled, concentrated using a 30 kDa cutoff Vivaspin concentrator
- 565 (GE Healthcare) and further purified by size exclusion chromatography (SEC) using a Superdex 75
- 566 16/600 column (GE Healthcare) in AcGH79 SEC buffer (20 mM HEPES [pH 7.0], 200 mM NaCl).
- 567 Fractions containing AcGH79 were pooled and concentrated using a 30 kDa Vivaspin concentrator to
- a final concentration of 14.5 mg/mL, and flash frozen for use in further experiments.
- 569 E287Q mutagenesis was carried out using a PCR based method⁴⁷. Mutant protein was purified using
- 570 the same protocol as for wild type protein. Mutagenesis primers are listed in Supplementary Table
- 571 **2**.
- 572 Mature HPSE
- 573 Mature HPSE cloning, expression, purification was carried out as previously described⁸.
- 574 E343Q mutagenesis was carried out using a PCR based method. Mutant protein was purified using
- 575 the same protocol as for wild type protein. Mutagenesis primers are listed in Supplementary Table
- 576 **2**.
- 577 proHPSE
- 578 Insect cells are unable to process proHPSE to mature HPSE, allowing the former to be isolated
- 579 following expression. cDNA encoding for proHPSE, minus the first 35 amino acid codons comprising
- the native signal sequence, was cloned behind a 5' honeybee mellitin signal sequence, 6×His tag,
- and TEV cleavage site, into the pOMNIBac plasmid (Geneva Biotech) using SLIC⁴⁸. pOMNIBac-
- 582 proHPSE was used to generate recombinant bacmid using the Tn7 transposition method in
- 583 DH10EMBacY cells⁴⁹ (Geneva Biotech). Baculovirus preparation and protein expression was carried
- out as previously described for mature HPSE.
- For purification, 3 L of conditioned media was cleared of cells by centrifugation at 400 g for 15 min at
- 586 4 °C, followed by further clearing of debris by centrifugation at 4000 g for 60 min at 4 °C. DTT (1
- 587 mM) and AEBSF (0.1 mM) were added to cleared media, which was loaded onto a HiTrap Sepharose
- 588 SP FF 5 mL column (GE healthcare) pre-equilibrated in IEX buffer A (20 mM HEPES [pH 7.4], 100 mM
- NaCl, 1 mM DTT). The loaded SP FF column was washed with 10 CV of IEX buffer A, and eluted with
- 590 a linear gradient over 30 CV using IEX buffer B (20 mM HEPES [pH 7.4], 1.5 mM NaCl, 1 mM DTT).
- 591 proHPSE containing fractions were pooled and diluted 10 fold into proHPSE HisTrap buffer A (20 mM
- HEPES [pH 7.4], 500 mM NaCl, 20 mM Imidazole, 1 mM DTT), before loading onto a HisTrap 5 mL FF
- 593 crude column pre-equilibrated in proHPSE HisTrap buffer A. The loaded HisTrap column was washed
- with 10 CV HisTrap buffer A, and eluted with a linear gradient over 20 CV using proHPSE HisTrap
- buffer B (20 mM HEPES [pH 7.4], 500 mM NaCl, 1 M Imidazole, 1 mM DTT). proHPSE containing
- fractions were pooled and concentrated to ~2 mL using a 30 kDa cutoff Vivaspin concentrator, and
- 597 treated with 5 μL EndoH (NEB) and 5 μL AcTEV protease (Invitrogen) for >72 h. Digested protein was
- purified by SEC using a Superdex S75 16/600 column in proHPSE SEC buffer (20 mM HEPES [pH 7.4],
- 599 200 mM NaCl, 1 mM DTT). proHPSE containing fractions were concentrated to 10 mg/mL using a 30
- 600 kDa Vivaspin concentrator, exchanged into IEX buffer A via at least 3 rounds of
- dilution/reconcentration, and flash frozen for use in further experiments.

| 603 | Overexpression of HPSE in HEK293T cells |
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| 604 605 606 607 608 609 | pGEn1-HPSE and pGEn2-HPSE plasmids were obtained from the DNASU repository 50 . HEK293T cells were grown in DMEM media supplemented with 10% newborn calf serum (NBCS; Sigma) and 1% penicillin/streptomycin (Sigma). 20 μg DNA was transfected into HEK293T cells at $^{\sim}80\%$ confluence, using linear PEI at a ratio of 3:1 (PEI:DNA). At relevant timepoints cells were washed with PBS, harvested into KPI buffer using a cell scraper, and pelleted by centrifugation at 200 g for 5 min at 4 $^{\circ}$ C. Cell pellets were frozen at -80 $^{\circ}$ C prior to use. |
| 610 611 612 | For CTSL inhibition experiments, transfections were carried out as above, except media was exchanged 7 h post transfection for DMEM supplemented with CTSL inhibitor or vehicle only control (0.05% v/v EtOH). |
| 613 | proHPSE fibroblast uptake experiment |
| 614 615 616 617 618 619 | Primary human fibroblasts were grown in DMEM/F12 media supplemented with 10% NBCS and 1% penicillin/streptomycin. Cells at 80% confluence were washed with PBS, before exchanging into DMEM/F12 media supplemented with proHPSE to 10 μ g/mL final concentration. At relevant timepoints, cells were washed with ice cold PBS twice, harvested into KPI buffer using a cell scraper, and pelleted by centrifugation at 200 g for 5 min at 4 °C. Cell pellets were stored at -80 °C prior to use. |
| 620 621 622 623 624 625 | For uptake experiments with prelabeled proHPSEs, prelabeling was carried out in McIlvaine citrate/phosphate buffer [pH 5.0], 300 mM NaCl in 200 μ L volume, using 50 μ M proHPSE and 200 μ M ABP. Reactions were incubated for 1 h for 37 °C, then excess ABP removed by desalting using a 40 kDa MWCO Zeba spin column (Thermo). Extent of prelabeling by 1 and 3 was quantified by comparison of protein and fluorophore UV/Vis absorption values. Extent of prelabeling by 6 was estimated by testing residual reactivity to 3 (Supplementary Fig. 11b). |
| 626 | Enzyme activity and inhibition assays |
| 627 628 629 630 631 632 633 | Recombinant AcGH79 enzyme activity was assayed using 1.67 ng protein in 150 mM McIlvaine buffe [pH 5.0]. To determine apparent IC50 values, 25 μ L AcGH79 was preincubated with a range of inhibitor dilutions for 30 min at 37 °C, followed by addition of 100 μ L 4MU-GlcUA solution to give final concentrations of 260 pM AcGH79 and 2.5 mM 4MUGlcUA. Reactions were carried out for 30 min at 37 °C, quenched with 200 μ L of 1 M NaOH-glycine [pH 10.3], and 4-MU fluorescence measured using a LS55 Fluorometer (Perkin Elmer) at λ_{ex} 366 nm and λ_{em} 445 nm. Apparent IC50 values were determined in Prism (GraphPad) using a one phase decay function. |
| 634 635 636 637 638 639 640 | Kinetic parameters for inhibition of AcGH79 were determined using a continuous method (Supplementary Note 1 ; also Ref. 20). AcGH79 was added to pre-warmed mixtures of 4MU-GlcUA and ABP, to give final concentrations of 260 pM AcGH79 and 2.5 mM 4MU-GlcUA in a final reaction volume of 125 μ L. Reactions were incubated at 37°C. At set timepoints, aliquots of reaction mixture were transferred to 96-well microplates (Greiner), quenched with 1 M NaOH-glycine [pH 10.3], and 4MU fluorescence measured immediately using a LS-55 Fluorometer. The apparent rate of inactivation (k_{obs}) was calculated for each ABP concentration by fitting with the exponential function |
| 641 | [4MU]=A*(1-e^($-k_{obs}$ *t)). The resulting plot of k_{obs} vs. [ABP] was fitted using a linear function, which |

- gives the combined apparent inhibition parameter ki/K'_{l} as the gradient. k_{i}/K_{l} was derived from k_{i}/K'_{l}
- by correcting for the presence of competing 4MU-GlcUA substrate, using the relationship
- $K'_{i}=K_{i}(1+[S]/K_{M})$, where [S] = 2.5 mM and K_{M} = 18.2 μM. All fittings were carried out using Prism.
- 645 In situ fibroblast IC50s were determined by incubating human fibroblast cells with a range of
- 646 inhibitor dilutions for 2 hours, followed by 3 × washing with PBS and harvesting into KPI buffer
- 647 supplemented with 0.1% Triton X-100. Harvested cells were pelleted by centrifugation at 200 g for 5
- 648 min at 4 °C, and pellets stored at –80 °C prior to use. Enzymatic reactions and apparent IC50
- 649 calculations were performed as described for the in vitro IC50 determination experiments, but with
- 650 5 μg of total lysate protein per reaction.

Fluorescent labeling

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- 652 Initial abeling reactions were carried out in McIlvaine buffer [pH 5.0], except for pH range
- 653 experiments, which were carried out in McIlvaine buffer at the stated pHs. Typically 200 fmol
- 654 recombinant protein was used for labeling AcGH79, HPSE and proHPSE (20 nM in 10 μL final reaction
- 655 volume). 20 μg total protein was used for labeling cell/tissue lysates, except for HEK293T
- 656 overexpression experiments, where 10 μg total protein was used. Unless otherwise specified,
- 657 labeling reactions were carried out by incubation with 100 nM fluorescent ABP in a reaction volume
- 658 of 10 μL for 1 h at 37 °C. Gels were scanned for ABP-emitted fluorescence using a Bio-Rad ChemiDoc
- 659 MP imager with the settings: Cy2 (λ_{EX} 470 nm, bandpass 30 nm; λ_{EM} 530 nm, bandpass 28 nm) for 1,
- 660 Cy3 (λ_{EX} 530 nm, bandpass 28 nm; λ_{EM} 605 nm, bandpass 50 nm) for **2**, and Cy5 (λ_{EX} 625 nm,
- bandpass 30 nm; λ_{EM} 695 nm, bandpass 55 nm) for **3**.
- 662 For labeling rate experiments with pro- and mature HPSE, reactions were carried out as above,
- 663 except 2 pmol recombinant protein was incubated with an equimolar amount of 3 at 37 °C. At
- 664 specified timepoints, aliquots were removed from the reaction and denatured by boiling in Laemmli
- 665 buffer. Denatured samples were stored on ice until all timepoints were collected, and run together
- 666 on SDS-PAGE.
- 667 For competition experiments, optimized labeling reactions were carried out in McIlvaine buffer [pH
- 668 5.0], 300 mM NaCl. Protein samples were preincubated with inhibitor for 60 min at 37 °C prior to
- 669 addition of 100 nM 3 for labeling. Platelet lysates were labeled at 37 °C for 1 h, recombinant
- 670 proteins were labeled at 37 °C for 30 min. Following labeling, samples were denatured by boiling
- 671 with Laemmli buffer for 5 min, and resolved by SDS-PAGE. Gels used for quantitation were scanned
- using a laser based Bio-Rad FX molecular imager, using the λ_{EX} 635 nm external laser and 690BP
- 673 emission filter. Images were analyzed using Quantity One (Bio-Rad). Full-length images of all
- fluorescent gels used in this study can be found in **Supplementary Fig. 12**.

Chemical proteomics

- 3 mg total protein from human wild type spleen, Gaucher spleen lysate, or human fibroblast lysate
- was incubated with either 10 μ M **4**, 10 μ M **9** for 30 min followed by 10 μ M **4**, or a vehicle only
- 678 control (0.1% DMSO). All labeling reactions were carried out for 30 min at 37 $^{\circ}$ C in 500 μ L McIlvaine
- 679 buffer [pH 5.0], before denaturation by addition of 125 μL 10% SDS and boiling for 5 min. Samples
- 680 were prepared for pull-down with streptavidin coupled DynaBeads (Invitrogen) as described
- previously⁵¹. Following pull-down the samples were divided: 1/3 for in-gel digest and 2/3 for on-

bead digest. In-gel digest samples were eluted by boiling beads at 100 °C in 30 μL Laemmli buffer.

Eluted proteins were separated by SDS-PAGE, and visualized by silver staining using the SilverQuest kit (Invitrogen). Bands were excised by scalpel and treated with gel digestion buffer (10 mM NH₄HCO₃, 5% ACN, 1mM CaCl₂, 10 ng/μL trypsin) at 37 °C overnight. The resulting trypsin-digested peptides were desalted using stage tips, followed by evaporation of ACN and resuspension into 70 μL sample solution (95:3:0.1 H₂O:ACN:TFA) for LC-MS analysis. Samples were analyzed with a 2h gradient of 5–25% ACN on a nano-LC, hyphenated to an LTQ-Orbitrap. Peptides were identified via

the Mascot protein search engine.

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On bead digest samples were treated with trypsin digestion buffer (100 mM Tris [pH 7.8], 100 mM NaCl, 1 mM CaCl₂, 2% ACN and 10 ng/ μ L trypsin) at 37 °C overnight with shaking. Trypsin digested peptides were desalted using stage tips, and analyzed as above. For identification of GUSB active-site peptides, trypsin digested beads were further treated with endoproteinase Glu-C digestion buffer (100 ng/ μ L Glu-C in PBS) at 37 °C overnight with shaking. Digested peptides were desalted and analyzed as above.

Peptide identification data were stringently filtered at a false discovery rate (FDR) of 1% and an MS/MS assignment Mascot score >40. Full proteomics data are available in Supplementary Data Set 1. The data presented in Supplementary Data Set 1 show: the ranking of the proteins by the protein score, the UniProt accession number, trivial name of the protein, protein score, protein mass predicted from the RNA sequence, the amino acid coverage of the protein achieved by MS/MS sequencing, the query number of the peptide in the LC/MS run, the experimentally determined m/z of the peptide, the measured molecular weight, the charge state z, the predicted molecular weight, the delta accuracy between predicted and experimentally determined mass, the MS/MS assignment Mascot score, the expectancy value, peptide sequence and emPAI value (which gives an approximate relative estimation of peptide abundance). Full proteomics data are available in Supplementary Data Set 1.

Western blotting

Proteins resolved by SDS-PAGE were transferred to a PVDF membrane using a Trans-Blot Turbo 708 709 system (BioRad). Membranes were blocked in 5% BSA for 1 h at rt, then incubated with anti-HPSE 710 (AbCam ab59787) or anti-GUSB (AbCam ab103112) at 1:1000 dilution in 5% BSA at 4 °C overnight. 711 Membranes were washed 3 × with TBST (50 mM Tris [pH 7.5], 150 mM NaCl, 0.05% Tween 20), then 712 incubated with HRP conjugated goat-anti-rabbit (Sigma A0545) at 1:5000 dilution in 5% BSA for 1 h 713 at room temperature. Membranes were washed again 3 × with TBST, blots visualized using 714 Amersham prime ECL reagent (GE Healthcare), and recorded using a Bio-Rad ChemiDoc XRS. Full-715 length images of all blots used in this study can be found in **Supplementary Fig. 12**.

Protein crystallization

717 AcGH79

AcGH79 was tested against a range of commercial crystallization screens. Well diffracting crystals of wild type AcGH79 were obtained by the sitting drop vapor diffusion method at 20 °C using 0.8–1.2 M 0.5:9.5 NaH₂PO₄:K₂HPO₄ (v/v) at a protein:well ratio of 700:500 nL. Crystals of AcGH79 E287Q

721 mutant were obtained using 1.2–1.5 M 1.0:9.0 NaH₂PO₄:K₂HPO₄ (v/v) at a protein:well ratio of

Comment [L1]: This seemed like a reasonable place to move this statement (from the Data Availability Statement below). However, if you think there is a better place for it in the Figure legend or in the running online methods, please move it.

Comment [12]: Moved to slightly earlier in paragraph, just before the description of the columns in the data set.

Suppl data set 1 is also called out in relevant parts of the main text (e.g. L142) and the caption for Fig. 2.

Comment [L3]: This seemed like a reasonable place to move this statement (from the Data Availability Statement below). However, if you think there is a better place for it in the Figure legend or in the running online methods, please move it.

Comment [14]: Removed the word 'gels', as this is the section about western blotting.

A similar statement about gels has been added to the previous page (L672), in the 'fluorescent labeling' subsection of the methods.

- 722 500:500 nL. Crystals typically appeared after 1 day. Crystals were cryoprotected using 2 M lithium
- 723 sulfate prior to flash freezing in liquid N₂ for data collection.
- 724 Mature HPSE
- 725 Mature HPSE was crystallized and data collected as previously described⁸.
- 726 proHPSE

- 727 proHPSE at 10 mg/mL was tested against a range of commercial crystallization screens. Thin plate
- 728 crystals were found in the JCSG screen, which were used to microseed subsequent rounds of crystal
- 329 screening⁵². Well diffracting single crystals were obtained by the sitting drop vapor diffusion method
- 730 at 20 °C using 100 mM succinate [pH 7.0], 17% PEG3350 and 1:250 diluted seed stock at a
- 731 protein:seed:well ratio of 700:100:400 nL. Crystals typically appeared after 1 day and grew to
- 732 maximum size within 10 days. proHPSE crystals were cryoprotected using mother liquor solution
- 733 supplemented with 25% ethylene glycol prior to flash freezing in liquid N₂ for data collection.

Xray data collection and structure solution

- 735 Xray diffraction data were collected at 100 K at beamlines i02 (5LA4, 5LA7), i03 (5G0Q, 5L77, 5L9Y)
- 736 and i04 (5L9Z) of the Diamond Light Source UK. Reflections were autoprocessed with the xia2
- 737 pipeline⁵³ of the CCP4 software suite, or manually processed using XDS⁵⁴ and Aimless⁵⁵. Apo
- 738 proHPSE was solved by molecular replacement with the mature HPSE model (5E8M) using MolRep⁵⁶,
- 739 followed by alternating rounds of manual model building and refinement using Coot and REFMAC5
- 740 respectively^{57,58}.
- 741 For all complexes with ABP 5, crystals were soaked in their respective mother liquors supplemented
- 742 with 2–5 mM **5** for 1–3 h at 20 °C. Soaked crystals were cryoprotected using their respective
- cryoprotectant solutions prior to flash freezing in liquid N_2 for data collection.
- 744 Complexes were solved by molecular replacement with their respective apo structures, followed by
- rounds of manual model building and refinement using Coot and REFMAC5. Ligand coordinates were
- 746 built using jLigand⁵⁹. Active site diagrams were generated using ccp4mg⁶⁰. Ribbon and protein
- 747 surface diagrams were generated using PyMOL.

748 Accession codes

- 749 Coordinates and structure factors have been deposited in the Protein Data Bank under accession
- 750 codes 5G0Q (AcGH79(wt)-5 complex), 5L77 (AcGH79(E287Q)-5 complex), 5L9Y (HPSE(wt)-5
- 751 complex), 5L9Z (HPSE(E343Q)-5 complex), 5LA4 (*apo* proHPSE), 5LA7 (proHPSE-5 complex).

752 Data availability

- 753 All other data generated or analyzed during this study are included in this published article (and its
- 754 Supplementary Information files), or are available from the corresponding authors on reasonable
- 755 request.

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Comment [15]: This is a verbatim repeat of L350-352. I originally added this information here as part of the data availability statement.

Is it still necessary?

Methods references

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