sp²-Iminosugars targeting human lysosomal \( \beta \)-hexosaminidase as pharmacological chaperone candidates for late-onset Tay-Sachs disease

Manuel González-Cuesta\(^a\), Irene Herrera-González\(^a\), M. Isabel García-Moreno\(^a\), Roger A. Ashmus\(^b\), David J. Vocadlo\(^b\), José M. García Fernández\(^de\), Eiji Namba\(^d\), Katsumi Higaki\(^d\) and Carmen Ortiz Melleta\(^a\)  

\(^a\)Department of Organic Chemistry, Faculty of Chemistry, University of Seville, Sevilla, Spain; \(^b\)Department of Chemistry and Department of Molecular Biology and Biochemistry, Simon Fraser University, Burnaby, Canada; \(^d\)Instituto de Investigaciones Químicas (IIQ), Consejo Superior de Investigaciones Científicas (CSIC) - Universidad de Sevilla, Sevilla, Spain; \(^e\)Organization for Research Initiative and Promotion, Tottori University, Yonago, Japan

ABSTRACT

The late-onset form of Tay-Sachs disease displays when the activity levels of human \( \beta \)-hexosaminidase A (HexA) fall below 10% of normal, due to mutations that destabilise the native folded form of the enzyme and impair its trafficking to the lysosome. Competitive inhibitors of HexA can rescue disease-causative mutant HexA, bearing potential as pharmacological chaperones, but often also inhibit the enzyme O-glycosaminidase (GlcNAcase; OGA), a serious drawback for translation into the clinic. We have designed sp²- iminosugar glycomimetics related to GalNAc that feature a neutral piperidine-derived thiourea or a basic piperidine-thiazolidine bicyclic core and behave as selective nanomolar competitive inhibitors of human Hex A at pH 7 with a ten-fold lower inhibitory potency at pH 5, a good indication for pharmacological chaperoning. They increased the levels of lysosomal HexA activity in Tay-Sachs patient fibroblasts having the G269S mutation, the highest prevalent in late-onset Tay-Sachs disease.

1. Introduction

\( N \)-Acetyl-\( \beta \)-hexosaminidase (Hex; EC 3.2.1.52) is a member of the glycosyl hydrolase family 20 (GH20) that catalyses the removal of terminal, non-reducing \( N \)-acetyl-\( \beta \)-D-glucosamine (GlcNAc) or galactosamine (GalNAc) residues from gangliosides, glycoproteins or glycosaminoglycans\(^1\). In humans two Hex isoforms are readily detectable, namely HexA and HexB. The first one is a heterodimer formed by \( \alpha \) and \( \beta \) subunits, encoded respectively by the evolutionary related \( \alpha \)-HEXA and \( \beta \)-HEXB genes, whereas the second one is the \( \beta \) homodimer\(^2\). The thermodynamically less stable \( \alpha \times \beta \) homodimer (HexS) is also formed, but only reaches measurable levels when the \( \beta \) subunit is deficient. Although the \( \alpha \) and \( \beta \) subunits possess independent active sites, dimerisation is a prerequisite for their in vivo biological function. Exclusively the \( \alpha \)-subunit of HexA can hydrolyse the \( \text{G}_{\text{M}}\text{2} \) ganglioside (GM2), an intermediate in the biosynthesis and degradation of higher brain gangliosides, in lysosomes by specifically interacting with the GM2 activator protein (GM2AP) co-factor\(^3\). Disabling mutations in \( \text{HEXA}, \text{HEXB} \) or the gene encoding for GM2AP result in Tay-Sachs disease (TSD; OMIM #272800), Sandhoff disease (SD; OMIM #268800) or the less common AB variant (OMIM #272750), respectively, a subset of lysosomal storage disorders (LSDs) collectively referred to as \( \text{G}_{\text{M}}\text{2} \) gangliosidoses\(^4\). All the three are autosomal recessive conditions associated with phenotypic neurodegeneration and devastating consequences. Currently, there are no effective treatment options for any of these diseases\(^5\). TSD is the best characterised of the \( \text{G}_{\text{M}}\text{2} \) gangliosidoses, with over 100 \( \text{HEXA} \) gene mutations categorised (http://www.hgmd.cf.ac.uk/). The frequency of asymptomatic heterozygotes is 1 in 300 live births in the general population (1 in 30 among Ashkenazy Jews), with a predicted frequency of 1 in 360,000 for homozygotes (1 in 2,900 among Ashkenazy Jews). Different genotypes result in different clinical phenotypes (infantile, juvenile or adult/chronic), with severity generally associating with the level of residual HexA activity permitted by different mutations (<0.5% of normal activity for infantile; 2%–5% for late on-set forms)\(^6\). Based on genotype/phenotype correlations, it has been suggested that 10% of the normal activity could be sufficient to prevent development of clinical symptoms\(^7\). Administration of recombinant HexA (enzyme replacement therapy; ERT) is unlikely to be clinically effective due to the inaccessibility imposed by the blood-brain barrier. Gene therapy using viral vectors has shown some promise in animal models\(^9,10\), but toxicity issues unveiled in non-human primates and low effectivity in clinical trials may seriously thwart translation to hospital settings\(^11,12\). Other approaches, such as the use of gene editing tools or brain permeable inhibitors of glycosphingolipid biosynthesis are under investigation\(^13,14\). Since many of the TSD-causative mutations do not compromise the catalytic site, but target the \( \alpha \) chain of HexA to endoplasmic reticulum-associate degradation (ERAD), the development of pharmacological chaperones (PCs) that can stabilise the native folding of the protein despite its anomalous conformation and restore activity appears attractive\(^15\). Typically, a PC is a small molecule able to bind to the
mutant enzyme at the ER, promote the correct folding and restore trafficking to the Golgi apparatus for maturation and then to the final destination. With few exceptions, most reported PCs developed for LSDs are competitive inhibitors of the target enzyme; they however exert an effector action by dissociating from the corresponding mature enzyme: inhibitor complex in the presence of an excess of substrate in the lysosomes of patient cells.

The prospective hexosaminidase PC candidates reported up to now include carbohydrate-related (glycomimetic) derivatives, among which 2-acetamido-1,2-dideoxyo-norimycin (DNJNAc), 6-acetamido-6-deoxyxastanospermine (NHAC-CAS), O-(2-acetamido-2-deoxy-o-glucopyranosylidene)amino N-phenylcarbamate (PUGNAc) and N-acetylglucosamine-thiazoline (NAG-thiazoline) are representative examples, and noncarbohydrate-based compounds identified after high throughput screening, such as pyrimethamine or the naphtalimide derivative M-31850 (Figure 1). A main problem is that oftentimes those candidates are not hexosaminidase function-specific and also inhibit a similar glycosidase, O-GlcNAcase (OGA), a GH84 enzyme that hydrolyses GlcNAc residues from O-linked glycoproteins. Three different strategies have been envisioned to address this issue: (a) installing substituents on the PC that can provide nonglycone interactions with the enzyme, as it is the case for N-[(di[p-methoxybenzyl]amino)-octyliminomethyl]-N-acetylhexosaminidase (DNJNAc) (DMH-DNJNAc) and (6S)-N-[(octyliminomethylidene)nojirimycin] (N-1-deoxy-6-thiogalactonojirimycin; NOI-NJ) 54 and the naphtalimide derivative M-31850 (Figure 1). Whereas all three approaches have produced selective hexA inhibitors, their application as TSD-associated mutant HexA enhancers remains to be demonstrated. Up to now, pyrimethamine has been the only PC candidate assayed in TSD patients; unfortunately, the results were not satisfactory enough, highlighting the urgent need for developing more efficient and selective PC prototypes.

We and others have previously shown that replacing the amine-type endocyclic nitrogen atom in iminosugar frameworks by a pseudoamide-type nitrogen (sp²-iminosugars) represents a versatile strategy to achieve highly selective glycosidase ligands for fundamental studies on enzyme mechanisms and for fundamental studies on enzyme mechanisms and for discovery, with applications ranging from cancer and inflammation to antiparasitic agents. sp²-iminosugars are very well adapted to molecular diversity schemes, including modifications in the configurational pattern, the heterocycle framework and the nature of substituents.

The prospective hexosaminidase PC candidates reported up to now include carbohydrate-related (glycomimetic) derivatives, among which 2-acetamido-1,2-dideoxyo-norimycin (DNJNAc), 6-acetamido-6-deoxyxastanospermine (NHAC-CAS), O-(2-acetamido-2-deoxy-o-glucopyranosylidene)amino N-phenylcarbamate (PUGNAc) and N-acetylglucosamine-thiazoline (NAG-thiazoline) are representative examples, and noncarbohydrate-based compounds identified after high throughput screening, such as pyrimethamine or the naphtalimide derivative M-31850 (Figure 1). A main problem is that oftentimes those candidates are not hexosaminidase function-specific and also inhibit a similar glycosidase, O-GlcNAcase (OGA), a GH84 enzyme that hydrolyses GlcNAc residues from O-linked glycoproteins. Three different strategies have been envisioned to address this issue: (a) installing substituents on the PC that can provide nonglycone interactions with the enzyme, as it is the case for N-[(di[p-methoxybenzyl]amino)-octyliminomethyl]-N-acetylhexosaminidase (DNJNAc) (DMH-DNJNAc) and (6S)-N-[(octyliminomethylidene)nojirimycin] (N-1-deoxy-6-thiogalactonojirimycin; NOI-NJ) 54 and the naphtalimide derivative M-31850 (Figure 1). Whereas all three approaches have produced selective hexA inhibitors, their application as TSD-associated mutant HexA enhancers remains to be demonstrated. Up to now, pyrimethamine has been the only PC candidate assayed in TSD patients; unfortunately, the results were not satisfactory enough, highlighting the urgent need for developing more efficient and selective PC prototypes.

We and others have previously shown that replacing the amine-type endocyclic nitrogen atom in iminosugar frameworks by a pseudoamide-type nitrogen (sp²-iminosugars) represents a versatile strategy to achieve highly selective glycosidase ligands for fundamental studies on enzyme mechanisms and for discovery, with applications ranging from cancer and inflammation to antiparasitic agents. sp²-iminosugars are very well adapted to molecular diversity schemes, including modifications in the configurational pattern, the heterocycle framework and the nature of substituents.

Figure 1. Structures of the HexA inhibitors DNJNAc, NHAC-CAS, PUGNAc, NAG-thiazoline, pyrimethamine, M31850, DMH-DNJNAc, LABNAc, Gal-PUGNAc and Gal-NAG-thiazoline, and of the sp²-iminosugars NOI-NJ, pFPhT-DGJ and 6S-NBI-DGJ.

2. Results and discussion

2.1. Design rational and synthesis

We advanced that sp²-iminosugar glycomimetics featuring a substitution profile of stereochemical complementarity to GalNAc would discriminate the lysosomal N-acetyl-β-hexosaminidase from OGA, in view of their respective substrate specificities. The recent implementation of a scalable preparative method for accessing 2-acetamido-1,2-dideoxygalactonojirimycin (DGJNAc), as the corresponding hydrochloride, enabled the direct synthesis of the corresponding thiourea adducts by coupling reactions with butyl, octyl, benzyl or phenyl isothiocyanate. Triethylamine was used as a base to promote deprotonation of the starting iminosugar salt.
The transformations proceeded with total chemoselectivity and good yields, avoiding hydroxyl protection/deprotection steps, in agreement with the categorisation of the thiourea-forming reaction as a click-type conjugation method very suited for molecular diversity oriented schemes (Scheme 1). The resulting hydroxythioureas served at their turn as precursors for the bicyclic derivatives \(5-8\). Intramolecular nucleophilic displacement of the primary hydroxyl group by the thiocarbonyl sulphur atom proceeded spontaneously in methanol at room temperature in the presence of hydrochloric acid, affording the target isothiourea-type sp\(^2\)-iminosugars in yields that ranged from virtually quantitative in the case of \(N'^{\text{alkyl}}\) derivatives \(1-3\) to moderate (41\%) for the \(N'^{\text{phenyl}}\) counterpart \(8\); Scheme 1).

2.2. Glycosidase inhibition properties against commercial enzymes

Compounds \(1-8\) were preliminary assayed for their glycosidase inhibitory properties against a panel of commercial glycosidases covering a variety of configurational and anomeric specificities, including bovine and almond \(\beta\)-glucosidase, yeast \(\alpha\)-glucosidase and isomaltase, \textit{Aspergillus niger} amyloglucosidase, \textit{Escherichia coli} \(\beta\)-galactosidase, green coffee bean \(\alpha\)-galactosidase, \textit{Helix pomatia} \(\beta\)-mannosidase, Jack bean \(\alpha\)-mannosidase and \(\beta\)-hexosaminidases from human placenta, bovine kidney and Jack bean. Only the hexosaminidase activity was significantly affected at concentrations below 1 mM of the sp\(^2\)-iminosugars (Supplementary Table 1). The corresponding Lineweaver-Burk plots were consistent with a competitive inhibition mode in all cases. In the thiourea series, the inhibition constants \((K_i)\) values for the human placenta enzyme were in the low micromolar range (3.0 to 16.0 \(\mu\)M) for the \(N'^{\text{alkyl}}\) substituted members \(1-3\) and down to nM range (0.24 \(\mu\)M) for the \(N'^{\text{phenyl}}\) adduct \(4\). The notably higher inhibition potency of the aromatic thiourea correlates with its higher hydrogen bond-donating character, strongly suggesting the direct implication of the \(N'H\) proton in enzyme-inhibitor complex stabilisation.

The influence of the substituent’s nature in the inhibition properties of the bicyclic iminothiazolidine-type sp\(^2\)-iminosugars \(5-8\) was much less pronounced as compared with the monocyclic thiourea counterparts \(1-4\). The basic character of the isothiourea group \((pK_a 8-9)\) implies that it will be fully protonated at neutral pH in all cases. Electrostatic interactions are then expected to be dominant at this region of the molecule. Indeed, the available structural information on \(\beta\)-N-acetylhexosaminidase reveals the presence of multiple negative charges at the vicinity of the enzyme active site, which is consistent with the much lower \(K_i\) values for the isothioureas as compared with the corresponding neutral thioureas, with the notable exception of the \(N'^{\text{phenyl}}\)thiocarbamoyl DGJNAc derivative \(4\) for the plant hexosaminidase \((K_i 0.065 \, \mu\)M; Supplementary Table 1).

2.3. Inhibition properties against human lysosomal hex in fibroblast lysates

The \(\beta\)-hexosaminidase from human placenta has been previously used in high throughput screenings to identify inhibitors that could act as pharmacological chaperones against disease-associated hexosaminidase mutants. However, it was recently encountered that the inhibitory activity of glycomimetic derivatives towards lysosomal \(\beta\)-hexosaminidases from different tissues can vary strikingly different, probably because of variances in the proportions of the mature and precursor forms of the enzyme, which may lead to false positives. Evaluation of the inhibition potency towards total Hex and HexA in human fibroblast lysates represents a more reliable test in search for chaperone candidates against...
TSD. The corresponding IC_{50} or K_i values for 1–8 are collected in Table 1. The inhibitory strength followed a similar trend to that encountered for the human placenta enzyme, that is, the bicyclic basic isothioureas 5–8 exhibited much higher inhibition efficiencies than the corresponding neutral thioureas 1–4. Among the later, only the N'-phenyl derivative 4 showed an IC_{50} value in the low micromolar range against total Hex (10.9 μM). The inhibitory potency generally increased when considering HexA, the target glycodelicase in TSD, which was particularly noticeable for the neutral thiourea derivatives. Thus, the N'-butyl and N'-phenyl-thioureas 1 and 4 behaved as potent inhibitors of HexA (K_i 4.7 and 0.91 μM, respectively, at pH 7), in addition to the bicyclic compounds 5–8 (0.41 to 0.99 μM at pH 7).

It is worth remarking that the rescuing capabilities of a PC depend not only on the binding potency to the target misfolded enzyme but also on the dissociation rate of the PC-enzyme complex once trafficking to the lysosome has been restored, so that substrate processing can take over. Interestingly, all the tested compounds exhibited HexA inhibition potencies that were about one order of magnitude higher at neutral (mimicking the endoplasmic reticulum) as compared to acidic pH (mimicking the lysosome environment), which is generally considered a good indication for PC candidates (Table 1). Regarding the HexA/OGA selectivity ratio, except for the N'-benzyliminothiazolidine derivative 7 (IC_{50} 0.66 vs 25 μM), it was over 1000-fold for all compounds (IC_{50} against OGA >100 μM).

The superior HexA inhibitory capabilities of the aromatic thiourea 4 and the thiazolidine derivatives 5–8 strongly suggest the involvement of the NH proton of the former (a better H-bond donor as compared to aliphatic thiourea protons) and of the protonated imonium group of the later in hydrogen bonding/salt bridge interactions stabilising the HexA-inhibitor complex. Similar scenarios have been previously reported for other sp²-iminosugar/glycosidase pairs. Actually, docking experiments support that compound 4 bind to the active site of HexA in a similar manner as NAG-thiazoline in the reported crystal structure. Specifically, the H-bond network involving the NHac and the secondary hydroxyl groups with ASP322, ARG178 and GLU462 is conserved. The C(=S)-N'HPh bond adopts the Z-configuration, with the NH proton close to the carboxylate group of the catalytic residue GLU323, forming an angle compatible with the presence of an H-bond (Figure 3A). A parallel situation is encountered for compound 6, but now the Z-oriented octyl chain must twist to avoid steric clashes and occupy a hydrophobic pocket flanked by TRP392 and TYR421 (Figure 3B). Such twisted conformation becomes less favourable in the aliphatic thiourea series, resulting in a destabilisation of the complex that intensifies with the volume of the NH substituent. Altogether, the data are in agreement with the observed experimental inhibition potency trend: 4–8 > 1 >> 2 and 3.

### 2.4. Chaperoning vs inhibitory properties in normal and late onset TSD fibroblasts

The new PC candidates were further evaluated about the HexA activity effect in wild type (healthy) fibroblasts (Figure 4A) and in TSD fibroblasts from patients hosting the G269S mutation (TSD-GS fibroblasts), the most commonly identified allele in patients with adult-onset TSD and a privileged target for future treatments (Figure 4B). None of the compounds affected significantly normal HexA activity in the healthy cells. Sharply differently, the results showed that the DGJNAc thioureas 1 and 4 and isothiourreas 5–8 behaved as very efficient mutant HexA effectors, being able to elicit activity enhancements in the 200%–300% range at 20 μM in all cases and in the 200%–250% range at only 2 μM concentration in the case of compounds 1, 4, 5, 7 and 8. The later compounds were further evaluated at 200 μM to ascertain whether the inhibitory character compensates the rescuing abilities at high dose. We keep in mind that reaching useful PC

---

**Table 1.** IC_{50} or K_i values for sp²-iminosugars 1–8 human lysosomal hexosaminidases and recombinant human O-GlcNAcase.

| Compound | IC_{50} (μM) | K_i (μM) |
|----------|-------------|----------|
|          | Total Hex b | HexA pH 7/pH 5 | OGA |
| **DGJNAc thioureas** |
| 1        | >100        | 4.7/33.7   | >100 |
| 2        | >100        | >100/100   | >100 |
| 3        | >100        | >100/100   | >100 |
| 4        | 10.9        | 0.91/9.4   | >100 |
| **DGJNAc isothiourreas** |
| 5        | 2.6         | 0.99/9.8   | >100 |
| 6        | 1.3         | 0.58/6.4   | >100 |
| 7        | 2.2         | 0.66/6.9   | 25   |
| 8        | 0.33        | 0.41/3.9   | >100 |

*aIC_{50} and K_i data are the mean of three independent determinations; errors are in the range of 5–10% and are omitted for clarity. Inhibition was competitive in all cases. No inhibition was observed for any compound at 200 μM on lysosomal α-glucosidase, α- and β-galactosidases and α- and β-mannosidases in cell lysates.

*bDetermined in human fibroblast lysates.

---

**Figure 3.** Predicted binding modes of phenylthiourea 4 (A; magenta sticks) and protonated octylminothiazolidine 6 (B; orange sticks) to HexA (coordinates from PDB ID 2GKL; tan sticks) from the molecular docking experiments. Hydrogen atoms from ligands are not shown for clarity. Hydrogen bonds are represented as black dashed lines, and hydrophobic interactions as red dashed lines.
concentrations in the central nervous system generally entails much higher doses in plasma and peripheral organs. The data evidenced a further enhancement of the G269S mutant HexA activity in all cases, reaching 3.4- and 2.7-fold for the thioureas 1 and 4 and up to 4.2-, 3.5- and 4.3-fold for the isothioureas 5, 7 and 8, respectively. The fact that the effector properties of the compounds prevails in a broad range of concentrations supports the notion that they can diffuse out of the lysosome and avoid intralysosomal accumulation, presumably due to their amphiphilic nature. The results highlight the promise of DGJNAc sp²-iminosugars as PCs for late-onset TSD.

To confirm that the mechanism of action involves rescuing and trafficking restoration of the endogenous mutant HexA, immunolabeling and colocalization experiments were further conducted using fibroblast from TSD-GS patients bearing the p.G269S/c.1278insTACT mutation (Figure 5). The left panel shows a series of confocal fluorescence microscopy images after immunostaining of HexA (green), lysosomes (lysosome associated membrane protein 2, red) and the corresponding merge images, in healthy control fibroblast and in TSD-GS fibroblasts. The right panel are the corresponding panels after treatment with 6 at 20 μM. It can be observed that the HexA content in the cells of TD patients is much lower than in the control and that it hardly colocalizes with the lysosomes. After treatment with chaperone 6, the amount of HexA increased significantly and clearly colocalized with this organelle as it happens in the control cells (Figure 5, Supplementary Figure 1). These results complement the in vitro assays and strongly supports that the observed enzyme activity enhancements correspond undoubtedy to the lysosomal HexA. Given that the residual activity of G269S mutant HexA was about 6% that of wild-type HexA, the measured enhancements warrant substrate processing levels well over the 10% of the normal threshold, considered critical for a medically relevant effect, in the 2–200 μM range, with no apparent toxicity. By comparison, pyrimethamine afforded 1.2 and 2.0-fold G269S mutant HexA activity enhancements at 0.5 and 1.0 g/mL in parallel assays, higher concentrations resulting in significant toxicity.

3. Conclusion

Grounding on the current structural knowledge on hexosaminidase-substrate (or inhibitor) complexes, we hypothesised that sp²-iminosugars emulating the substitution profile and stereochemistry of GalNAc would fit in the active site of HexA but not in that of the related enzyme OGA. By equipping the glycomimetic core with suitable appendages, providing additional contacts with the enzyme, the ability to properly folding and rescuing mutant HexA causative of late-onset TSD could be optimised. To put this concept into practice, we conducted the direct conjugation of four different isothiocyanates with DGJNAc. The corresponding thio-urea adducts were thus obtained and subsequently subjected to intramolecular cyclisation to afford bicyclic pyridine-thiazolidine derivatives. Nanomolar competitive inhibitors of HexA at pH 7, with high HexA/OGA selectivity, were identified in both the (neutral) thiourea and (basic) iminothiazolidine series. The inhibitory potency fell down by ten-fold at pH 5, which should facilitate the dissociation of the HexA-glycomimetic complex at the lysosome, where the Hex substrate (namely GM2 ganglioside) accumulates. In agreement with this notion, remarkable HexA activity enhancements were achieved in fibroblasts from TSD patients having the G269S mutation, associated to the late-onset phenotype.

**Figure 4.** Effect in the HexA activity of different concentrations of the thioureas DGJNAc thioureas 1–4 and the isothiourea derivatives 5–8 in wild type (A) and G269S TSD fibroblasts (B) relative to controls in the absence of any compound. Bars represent average values ± SD (n = 3).

**Figure 5.** Effect of compound 6 on the traffic of HexA to lysosomes in WT and in TSD-GS fibroblast. Fibroblasts were treated with 20 μM complexed chaperone. Lysosomal marker (lysosome associated membrane protein 2, LAMP-2). HexA is visualised in green; in the merged images, yellow denotes colocalization in lysosomes. The shown pictures are representative of more than 100 cells in 10 randomly obtained images.
Interestingly, most candidates still enhanced G269S HexA activity in cell assays even at high (200 μM) concentration. Altogether, the results validate the approach of using GalNAc sp²-iminosugar mimics to identify drug-like pharmacological chaperone candidates for the treatment of late-onset TSD.

4. Experimental section

4.1. General methods

Reagents and solvents were purchased from commercial sources and used without further purification. Optical rotations were measured with a JASCO P-2000 polarimeter, using a sodium lamp (λ = 589 nm) at 22 °C in 1 cm or 1 dm tubes. IR spectra were recorded on a JASCO FTIR-410 device. UV spectra were recorded on JASCO V-630 instrument; unit for ε values: mM⁻¹ cm⁻¹. NMR experiments were performed at 300 (75.5) and 500 (125.7) MHz. 1-D TOCSY as well as 2-D COSY and HMQC experiments were carried out to assist signal assignment. In the FABMS spectra, the primary beam consisted of Xe atoms with a maximum energy of 8 keV. The samples were dissolved in m-nitrobenzyl alcohol or thioglycoler as the matrices and the positive ions were separated and accelerated above a potential of 7 keV. NaI was added as a catalyzing agent. For ESI mass spectra, 0.1 pm sample concentrations were used, the mobile phase consisting of 50% ac MeCN at 0.1 ml min⁻¹. Thin-layer chromatography was performed on precoated TLC plates, silica gel 30 F-254, with visualization by UV light and also with 10% H₂SO₄ or 0.2% w/v cerium (IV) sulfate-5% ammonium molybdate in 2 m H₂SO₄ or 0.1% ninhydrin in EtOH. Column chromatography was performed on Chromagel (silice 60 ACC 70-200 μm). All compounds were purified to >95% purity as determined by elemental microanalysis results obtained on a CHNS-TruSpect® Micro elemental analyser (Instituto de Investigaciones Químicas de Sevilla, Spain) from vacuum-dried samples. The analytical results for C, H, N and S were within ±0.5 of the theoretical values. Starting 2-acetamido-1,2-dideoxygalactonojirimycin (DGJNAc) was prepared following a described procedure. 4-Methylumbelliferone (4-MU)-conjugated N-acetyl-β-D-glucosaminide was obtained from Sigma (St. Louis, MO). 4-MU-N-acetyl-β-D-glucosaminide-6-sulfate sodium salt was from Slater and Frith Ltd (Norfolk, UK). LDH cytotoxicity kit was from WAKO (Tokyo, Japan).

4.2. Chemistry

4.2.1. General procedure for the preparation of 2-acetamido-N-(N’-alkyl(aryl)thiocarbamoyl)-1,2-dideoxygalactonojirimycin derivatives (1–4)

To a solution of DGJNAc (102 mg, 0.423 mmol) in pyridine (3.8 ml), (Tokyo, Japan).

4.2.2. General procedure for the preparation of 2-acetamido-N-[2-[2-(1H-imidazol-2-yl)ethyl]phenyl]carbonyl]-1,2-dideoxygalactonojirimycin derivatives (5–8)

The DGJNAc thiourea derivative (1-4, 0.256 mmol) was dissolved in MeOH (80 ml) and a few drops of concentrated HCl were added until pH 1. The solution was stirred at rt until complete disappearance of the starting material (7–24 h). The solvent was removed under reduced pressure and the residue was coevaporated with MeOH several times until neutral pH and purified by column chromatography using the eluent indicated in each case to afford the corresponding bicyclic DGJNAc isothiouries.
2-Acetamido-1,2-dideoxy-5-N,6-S-(N'-butyliminothiobenzylidene)-6-thiogalactonojirimycin (5). Column chromatography, eluent: 70:10:1→30:10:1 CH2Cl2:MeOH:H2O. Yield: 78 mg (quantitative). ESIMS: m/z 538 [M + H]⁺. Anal. Calcd for C15H19N3O3S: C, 51.89; H, 7.78; N, 13.77; S, 10.36.

2-Acetamido-1,2-dideoxy-5-N,6-S-(N'-octyliminothiobenzylidene)-6-thiogalactonojirimycin (6). Column chromatography, eluent: 40:10:1 DCM:MeOH:H2O. Yield: 82.7 mg (96%). ESIMS: m/z 542 [M + H]⁺. Anal. Calcd for C15H23N3O3S: C, 51.99; H, 7.64; N, 13.77; S, 10.36.

2-Acetamido-1,2-dideoxy-5-N,6-S-(N'-benzyliminothiobenzylidene)-6-thiogalactonojirimycin (7). Column chromatography, eluent: 60:10:1→40:1:1 DCM:MeOH:H2O. Yield: 60 mg (quantitative). ESIMS: m/z 546 [M + H]⁺. Anal. Calcd for C15H22N3O3S: C, 51.99; H, 8.17; N, 13.77; S, 10.36.

2-Acetamido-1,2-dideoxy-5-N,6-S-(N'-phenyliminothiobenzylidene)-6-thiogalactonojirimycin (8). Column chromatography, eluent: 80:10:1→40:1:1 DCM:MeOH:H2O. Yield: 27 mg (41%). ESIMS: m/z 544 [M + H]⁺. Anal. Calcd for C15H22N3O3S: C, 51.99; H, 8.17; N, 13.77; S, 10.36.

4.4. Commercial enzyme inhibition assays

Inhibition constant (Kᵢ) values were determined by spectrophotometrically measuring the residual hydrolytic activities of the glycosidases against the respective o- (for β-galactosidase from E. coli) or p-nitrophenyl α- or β-D-glycopyranoside (for other glycosidases) in the presence of the iminosugars. Each assay was performed in phosphate buffer or phosphate-citrate buffer (for α- or β-mannosidase, amyloglucosidase or β-N-acetylc glucosaminidase) at the optimal pH for the enzymes. The reactions were initiated by addition of the enzyme to a solution of the substrate in the absence or presence of various inhibitor concentrations. The mixture was incubated for 10–30 min at 37°C or 55°C (for amyloglucosidase) and the reaction was quenched by addition of 1 m Na2CO3. Reaction times were appropriate to obtain 10%–20% conversion of the substrate in order to achieve linear rates. The absorbance of the resulting mixture was determined at 405 nm. Approximate values of Kᵢ were determined using a fixed concentration of substrate (around the Kᵡ value for the different glycosidases) and various inhibitor concentrations. Full Kᵢ determinations and the enzyme inhibition mode were determined from the slope of Lineweaver–Burk plots and double reciprocal analysis.

4.5. Profiling of the inhibitory selectivity towards lysosomal glycosidases by in vitro enzyme assay using human fibroblast lysates

For determination of lysosomal enzyme activities in cell lysates, cells were scraped into ice-cold H2O (10⁶ cells mL⁻¹) and lysed by sonication. Insoluble materials were removed by centrifugation at 15,000 rpm for 5 min and protein concentrations were determined with Protein Assay Rapid Kit (WAKO, Tokyo, Japan). 10 μL of the lysates in 0.1% Triton X-100 in distilled water were incubated at 37°C with 20 μL of the substrate solution in 0.1 M citrate buffer, pH 4.5, in absence or presence of increasing concentrations of the chaperones. The substrates were 4-methylumbelliferone (4-MU)-
conjugated β-D-glucopyranoside (for GCase), α-D-glucopyranoside (for α-glucosidase), α-D-galactopyranoside (for α-Galase), β-D-galactopyranoside (for β-galactosidase), N-acetyl-β-D-glucosaminide (for total β-hexosaminidase), N-acetyl-β-D-glucosaminide 6-sulfate sodium salt (for HexA) and α-N-acetyl-D-galactosaminide for α-N-acetylgalactosaminidase. HexA activities in cell lysates were also determined in 0.1 M citrate buffer at pH 5 or pH 7, supplemented with sodium taurocholate (0.8% w/v). The reactions were terminated by adding 0.2 mL of 0.2 M glycine sodium hydroxide buffer (pH 10.7). The liberated 4-methylumbelliferone was measured in the black-well plate with a Perkin Elmer Luminescence Spectrometer (excitation wavelength: 340 nm; emission: 460 nm). One unit of enzyme activity was defined as nmol of 4-methylumbelliferone released per hour and normalised for the amount of protein contained in the lysates.

4.6. Cloning, expression and purification of hOGA and inhibition assay

The gene of the longest isoform of hOGA (hOGA-L; Uniprot Accession number: O60502) was synthesised in a codon-optimised form for recombinant expression in Escherichia coli. The successful expression construct encoded the N-terminal region of hOGA, comprising amino acids 11–396, and the C-terminal region, comprising amino acids 535–715. The N-terminal construct was cloned in the vector pACYC-Duet (Millipore) using the sequence and ligation-independent cloning method in frame with an N-terminal His6 tag. The C-terminal construct was cloned into the vector pETYSBLIC3C27 with an N-terminal His6 tag followed by a 3 C-protease cleavage site using the same method. The nucleotide sequences of all made constructs were confirmed by sequencing. Both vectors were simultaneously transformed into E. coli BI21(DE3)-Gold (Agilent) for subsequent protein expression. Cells were grown in 2L TB-medium to an OD600 of ~1.0 and protein synthesis was then induced by adding IPTG to a final concentration of 1.0 mM. Protein expression was carried out at 16 °C with an induction time of 20h. The cells were harvested by centrifugation at 4,500 × g for 20 min, flash frozen and stored at −20 °C until required. For purification of the hOGA complex, cells were resuspended in 50 mM HEPES pH 7.0, 750 mM NaCl, 20 mM imidazole, and 0.5 mM DTT (resuspension buffer). Cells were lysed using a French Press at 25 Kps. The lysate was cleared by centrifugation at 50,000 g for 1 h, and the supernatant was passed through a 10 mL HisTrap FF column (GE Healthcare) pre-equilibrated with resuspension buffer. The bound hOGA was purified by gradient elution over 10 column volumes using 0% to 50% of elution buffer (50 mM HEPES pH 7.0, 750 mM NaCl, 500 mM imidazole and 0.5 mM DTT). hOGA-containing fractions were combined, concentrated by ultrafiltration using Vivaspin columns (Sartorius) with a molecular weight cut-off (MWCO) of 30 kDa, and applied to a Superdex S200 column (GE Healthcare) pre-equilibrated with size-exclusion buffer (10 mM HEPES pH 7.0, 250 mM NaCl, 1 mM DTT). Fractions corresponding to the dimeric form of hOGA were combined and concentrated to 20 mg/mL by ultrafiltration with a Vivaspin (MWCO: 30 kDa) column, flash frozen using liquid nitrogen, and stored at −80 °C until required. Western blotting, conducted as previously reported, showed an intense band corresponding to hOGA and low intensity bands arising from truncated forms of the enzyme (Supplementary Figure 11).

Inhibition potencies for compounds 1–8 against hOGA enzyme were determined by measuring the change in fluorescent signal corresponding to the rate of hydrolytic activity against the artificial substrate, resorufin N-acetyl-β-D-glucosaminide. hOGA activity assays were performed in a buffer of 20 mM HEPES, 5 mM EDTA, 150 mM KCl, pH 7.1 and 0.2 mM [hOGA]. Km = 25 μM was measured with these conditions. Inhibition assays were run at 0.2 mM [hOGA] in the same buffer, in the presence or absence of various concentrations of inhibitors and at a fixed substrate concentration of 25 μM and 1% DMSO. First, inhibitor was serially diluted to the desired range of concentrations in 2% DMSO buffer. Next, 50 μL of inhibitor solutions at various concentrations in 2% DMSO Buffer was added to 50 μL of 0.8 nM enzyme and allowed to incubate at 25 °C for 5 min. 100 μL of 50 μM and 1% DMSO substrate was then added and reaction mixture was immediately mixed and aliquoted in 45 μL triplicates to a CORNING 384 well black plate. Fluorescence signal was measured continuously for 20 min at 37 °C in a BioTek Neo 2 Plate reader set at excitation and emission wavelengths of 572 and 610 nm, respectively. Max reaction rates for all inhibitor concentrations were calculated within Gen5 BioTek reader software. % Activity was subsequently calculated for each inhibitor concentration against the max reaction rate of uninhibited hOGA. Except for compound 7 (IC50 25 mM), none of the compounds displayed significant hOGA inhibition at concentrations up to 100 μM.

4.7. Cell culture and chaperone treatment

Human skin fibroblasts from normal subject were cultured as described. Human fibroblasts from TSD patient with p.G269S/HEXA mutation were obtained from Coriell Cell Repositories (Camden, NJ). For measurement of chaperone effects, cells were cultured in the medium with or without compounds (1–8) for 4 days and then the cell lysates were subjected for the lysosomal enzyme assay as described above. The enzyme activities were normalised with protein concentrations measured by protein assay rapid kit (Wako, Tokyo, Japan).

4.8. Immunofluorescence microscopy

Immunofluorescence microscopy was performed using standard methods as previously described.40 Coverslips were analysed using a fluorescence microscope (Leica DMRE, Leica Microsystems GmbH, Wetzlar, Germany). Deconvolution studies and three-dimensional projections were performed using a DeltaVision system (Applied Precision, Issaquah, WA) with an Olympus IX-71 microscope. The deconvolved images were derived from optical sections taken at 30 nm intervals using a 60× PLAPON objective with a 1.42 numerical aperture. More than 100 cells in 10 randomly obtained images were evaluated in each experiment to confirm reproducibility.

4.9. Statistical analysis

All results are expressed as mean±SD of three independent experiments, each conducted in triplicate. The measurements were statistically analysed using the Student’s t test for comparing 2 groups. The level of significance was set at p < 0.05.

Author contributions

The manuscript was written through the contributions of all authors. All authors have given approval to the final version of the manuscript.
Disclosure statement

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

Funding

This work was financially supported by MCIN/AEI/10.13039/501100011033 and “ERDF A way of making Europe” [Grants numbers PID2019-105858RRI-100 and RTI2018-097609-B-C21] and the Junta de Andalucía [Grant number P20_00166]. The authors are grateful to the Canadian Institutes of Health Research [MOP-123341], the Natural Sciences and Engineering Research Council of Canada [RGPIN-06466] and the JSPS KAKENHI Grant 17K10051 for support of this research. DJV thanks the Canada Research Chairs Program for support as a Tier I Canada Research Chair in Chemical Biology. The CITIUS (University of Seville) is also acknowledged for technical support. M.G.-C. and I.H-G. are supported by FPI and FPU fellowships, respectively [Grant numbers BES-2017–079676 and FPU17/03147, funded by MCIN/AEI/10.13039/501100011033 and by “ESF Investing in your future”]. RAA is supported by a trainee award from the Michael Smith Foundation Health Research (MSFHR).

ORCID

José M. García Fernández http://orcid.org/0000-0003-2323-6887
Carmen Ortiz Mellet http://orcid.org/0000-0002-7676-7721

References

1. Wendeler M, Sandhoff K. Hexosaminidase assays. Glycoconjug J 2009;26:1364–52.
2. Triggs-Raine B, Mahuran DJ, Gravel RA. Naturally occurring mutations in GM¿ gangliosidosis: a compendium. Adv Genet 2001;44:199–224.
3. Mahuran DJ. The GM¿ activator protein, its roles as a co-factor in GM¿ hydrolisis and as a general glycolipid transport protein. Biochim Biophys Acta 1998;1393:1–18.
4. Gravel RA, Kaback MM, Proia RL, et al. The GM¿ gangliosidoses, In: Scrivé CR, Beaudet AL, Sly WS, Valle D. The metabolic and molecular bases of inherited disease, In: Scriver CR, Beaudet AL, Sly D WS, Valle, eds. The metabolic and molecular bases of inherited disease. New York: McGraw-Hill; 2001:3827–3876.
5. Bley AE, Giannikopoulos OA, Hayden D, et al. Natural history of infantile GM¿ gangliosidosis. Pediatrics 2011;128: e1233–1241.
6. Neudorfer O, Pastores GM, Zeng BJ, et al. Late-onset Tay-Sachs disease: phenotypic characterization and genotypic correlations in 21 affected patients. Genet Med 2005;7:119–23.
7. Sheth J, Mistri M, Datar C, et al. Expanding the spectrum of HEXA mutations in Indian patients with Tay-Sachs disease. Mol Genet Metab Rep 2014;1:425–30.
8. Leinekugel P, Michel S, Conzelmann E, Sandhoff K. Quantitative correlation between the residual activity of beta-hexosaminidase A and arylsulfatase A and the severity of the resulting lysosomal storage disease. Hum Genet 1992;88:513–23.
9. Karumuthil-Melethil S, Kalburgi SN, Thompson P, et al. Novel vector design and hexosaminidase variant enabling self-complementary adeno-associated virus for the treatment of Tay-Sachs disease. Hum Gene Ther 2016;27:509–21.
10. Cachón-González MB, Wang SZ, Lynch A, et al. Effective gene therapy in an authentic model of Tay-Sachs-related diseases. Proc Natl Acad Sci USA 2006;103:10373–8.
11. Golebiowski D, van der Bom IMJ, Kwon CS, et al. Direct intracranial injection of AAVrh8 encoding monkey N-acetyltetrahexosaminidase causes neurotoxicity in the primate brain. Hum Gene Ther 2017;28:510–22.
12. Choudhury SR, Hudry E, Maguire CA, et al. Viral vectors for therapy of neurologic diseases. Neuropharmacology 2017;120:63–80.
13. Leal AF, Benincore-Flořez E, Solano-Galarza D, et al. GM¿ gangliosidoses: clinical features, pathophysiological aspects, and current therapies. Int J Mol Sci 2020;21:6213.
14. Cox TM. Lysosomal diseases and neuropsychiatry: opportunities to rebalance the mind. Front Mol Biosci 2020;7:177.
15. Dersh D, Iwamoto Y, Argon Y. Tay-Sachs disease mutations in HEXA target the x chain of hexosaminidase A to endoplasmic reticulum-associated degradation. Mol Biol Cell 2016;27:3813–27.
16. Iacono R, Minopoli N, Ferrara MC, et al. Carnitine is a pharmacological allosteric chaperone of the human lysosomal x-glucosidase. J Enzyme Inhib Med Chem 2021;36:2068–79.
17. Tran ML, Génisson Y, Ballereau S, Dehoux C. Second-generation pharmacological chaperones: beyond inhibitors. Molecules 2020;25:3145.
18. Porto C, Ferrara MC, Meli M, et al. Pharmacological enhancement of x-glucosidase by the allosteric chaperone N-acetyl-cysteine. Mol Ther 2012;20:2201–011.
19. Sánchez-Fernández EM, García Fernández JM, Ortiz Mellet C. Glycomimetic-based pharmacological chaperones for lysosomal storage disorders: lessons from Gaucher, GM1-gangliosidoses and Fabry diseases. Chem Commun 2016;52:5497–515.
20. Parenti G, Andria G, Valenzano KJ. Pharmacological chaperone therapy: preclinical development, clinical translation, and prospects for the treatment of lysosomal storage disoders. Mol Ther 2015;23:1138–48.
21. Platt FM. Emptying the stores: lysosomal diseases and therapeuetic strategies. Nat Rev Drug Discovery 2018;17:133–50.
22. Tropak MB, Reid SP, Guiral M, et al. Pharmacological enhancement of beta-hexosaminidase activity in fibroblasts from adult Tay-Sachs and Sandhoff patients. J Biol Chem 2004;279:13478–87.
23. Li H, Marcelo F, Bello C, et al. Design and synthesis of acetamido tri- and tetra-hydroxyazepanes: potent and selective beta-N-acetylgalactosaminidase inhibitors. Bioorg Med Chem 2009;17:5589–604.
24. Blériot Y, Tran AT, Prencipe G, et al. Synthesis of 1,2-trans-2-acetamido-2-deoxyhomoinosugars. Org Lett 2014;16:5516–9.
25. Steiner AJ, Schitter G, Stütz AE, et al. 2-Acetamino-1,2-dideoxyo-norimycin-l-lysine hybrids as hexosaminidase inhibitors. Tetrahedron Asymmetry 2009;20:832–5.
26. Maegawa GHB, Tropak M, Buttnar J, et al. Pyrimethamine as a potential pharmacological chaperone for late-onset forms of GM2 Gangliosidosis. J Biol Chem 2007;282:9150–61.
27. Tropak MB, Blanchard J, Withers SG, et al. High-throughput screening for human lysosomal beta-N-Acetyl hexosaminidase inhibitors acting as pharmacological chaperones. Chem Biol 2007;14:153–64.
28. Guo P, Chen Q, Liu T, et al. Development of unsymmetrical dyads as potent noncarbohydrate-based inhibitors against human β-N-Acetyl-d-hexosaminidase. ACS Med Chem Lett 2013;4:527–31.

29. Liu T, Guo P, Zhou Y, et al. A crystal structure-guided rational design switching non-carbohydrate inhibitors’ specificity between two β-GlcNAcase homologs. Sci Rep 2014;4:6188.

30. Colussi DJ, Jacobson MA. Patient-derived phenotypic high-throughput assay to identify small molecules restoring lysosomal function in Tay-Sachs disease. SLAS Discov 2019;24:295–303.

31. Hart GW, Housley MP, Slawson C. Cycling of O-linked beta-N-acetylglucosamine on nucleocytoplasmic proteins. Nature 2007;446:1017–22.

32. Roth C, Chan S, Offen WA, et al. Structural and functional insight into human O-GlcNAcase. Nat Chem Biol 2017;13:610–4.

33. Wang L, Bharti R, Kumar R, et al. Small molecule therapeutics for tauopathy in Alzheimer’s disease: walking on the path of most resistance. Eur J Med Chem 2021;209:112915.

34. Ho CW, Popat SD, Liu TD, et al. Development of GlcNAc-inspired iminocyclitiols as potent and selective N-acetyl-beta-hexosaminidase inhibitors. ACS Chem Biol 2010;5:489–97.

35. Rountree JSS, Butters TD, Wormald MR, et al. Design, synthesis, and biological evaluation of enantiomeric beta-N-acetyl-hexosaminidase inhibitors LABNAC and DABNAC as potential agents against Tay-Sachs and Sandhoff disease. ChemMedChem 2009;4:378–92.

36. Glawar AFG, Martinez RF, Ayers BJ, et al. Structural essentials for β-N-acetylhexosaminidase inhibition by amides of prolines, piperocic and azetidine carboxylic acids. Org Biomol Chem 2016;14:10371–85.

37. Stubbs KA, Macauley MS, Vocadlo DJ. A selective inhibitor Gal-PUGNAc of human lysosomal beta-hexosaminidases modulates levels of the ganglioside GM2 in neuroblastoma cells. Angew Chem Int Ed Engl 2009;48:1300–3.

38. Amorelli B, Yang C, Rempel B, et al. N-Acetylhexosaminidase inhibitory properties of C1-homologated GlcNAc- and GalNAc-thiazolines. Bioorg Med Chem Lett 2008;18:2944–7.

39. Osher E, Fattal-Valevski A, Sagie L, et al. Effect of cyclic, low dose pyrimethamine treatment in patients with late onset Tay Sachs: an open label, extended pilot study. Orphanet J Rare Dis 2015;10:45.

40. Ríquez-Cuadro R, García Fernández JM, Nierengarten JF, Ortiz Mellet C. Fullereone-sp2-iminosugar balls as multimodal ligands for lectins and glycosidases: a mechanistic hypothesis for the inhibitory multivalent effect. Chemistry 2013;19:16791–803.

41. Abellán Flos M, García Moreno MI, Ortiz Mellet C, et al. Potent glycosidase inhibition with heterovalent fullerene: unveiling the binding modes triggering multivalent inhibition. Chemistry 2016;22:11450–60.

42. García- Moreno MI, Ortega-Caballero F, Ríquez-Cuadro R, et al. The impact of heteromultivalency in lectin recognition and glycosidase inhibition: an integrated mechanistic study. Chemistry 2017;23:6295–304.

43. Ortiz Mellet C, Nierengarten JF, García Fernández JM. Multivalency as an action principle in multimodal lectin recognition and glycosidase inhibition: a paradigm shift driven by carbon-based glyconanomaterials. J Mater Chem B 2017;5:6546–56.

44. Gueder N, Allan G, Telliez MS, et al. sp2-iminosugar ω-glucosidase inhibitor 1-C-octyl-2-oxa-3-oxocastanospermine specifically affected breast cancer cell migration through Stim1, β1-integrin, and FAK signaling pathways. J Cell Physiol 2017;232:3631–40.

45. Sánchez-Fernández EM, Gonçalves-Pereira R, Ríquez-Cuadro R, et al. Influence of the configurational pattern of sp2-iminosugar pseudo N-, S-, O- and C-glycosides on their glycoside inhibitory and antitumor properties. Carbohydr Res 2016;429:113–22.

46. Allan G, Ouadid-Ahidouch G, Sánchez-Fernández EM, et al. New castanospermine glycoside analogues inhibit breast cancer cell proliferation and induce apoptosis without affecting normal cells. PLoS One 2013;8:e76411.

47. Schaeffer E, Sánchez-Fernández EM, Gonçalves-Pereira R, et al. sp2-iminosugar glycolipids as inhibitors of lipopolysaccharide-mediated human dendritic cell activation in vitro and of acute inflammation in mice in vivo. Eur J Med Chem 2019;169:111–20.

48. Sánchez-Fernández EM, García- Moreno MI, Arroba AI, et al. Synthesis of polyfluoroalkyl sp2-iminosugar glycolipids and evaluation of their immunomodulatory properties towards anti-tumor, anti-leishmanial and anti-inflammatory therapies. Eur J Med Chem 2019;182:111604.

49. Herrera-González I, Sánchez-Fernández EM, Sau A, et al. Stereoselective synthesis of iminosugar 2-deoxy(thio)glycosides from bicyclic iminoglycal carbamates promoted by cerium(IV) ammonium nitrate and cooperative Brensted acid-type organocatalyst. J Org Chem 2020;85:503–5047.

50. Sevšek A, Celen M, Erjavec B, et al. Bicyclic isourea derivatives from 1-deoxynojirimycin are potent inhibitors of β-glucocerebrosidase. Org Biomol Chem 2016;14:8670–3.

51. Mena-Barragán T, García- Moreno MI, Nanba E, et al. Inhibitor versus chaperone behaviour of D-fagomine, DAB and LAB sp2-iminosugar conjugates against glycosidases: a structure-activity relationship study in Gaucher fibroblasts. Eur J Med Chem 2016;121:880–91.

52. Sánchez-Fernández EM, Álvarez E, Ortiz Mellet C, García Fernández JM. Synthesis of multibranchd australian derivatives from reducing castanospermine analogues through the Amadori rearrangement of gem-diamine intermediates: selective Inhibitors of β-Glucosidase. J Org Chem 2014;79:11722–8.

53. García- Moreno MI, Ortiz Mellet C, García Fernández JM. Synthesis of calystegine B2, B3, and B4 analogues: mapping the glycosidase inhibitory activity relationships in the 1-deoxy-6-oxacalystegine series. Eur J Org Chem 2004;2004:1803–19.

54. Tiscornia G, Lorenzo Vivas E, Matalonga E, et al. Neuronalpathic Gaucher’s disease: induced pluripotent stem cells for disease modelling and testing chaperone activity of small compounds. Hum Mol Genet 2013;22:633–45.

55. de la Mata M, Cotán D, Oropesa-Avila M, et al. Pharmacological chaperones and coenzyme Q10 treatment improves mutant β-glucocerebrosidase activity and mitochondrial function in neuronalpathic forms of Gaucher disease. Sci Rep 2015;5:10903.

56. García- Moreno MI, de la Mata M, Sánchez Fernández EM, et al. Fluorinated chaperone-β-cycloextrin formulations for β-Glucocerebrosidase activity enhancement in neuronalpathic Gaucher disease. J Med Chem 2017;60:1829–42.
57. Mena-Barragán T, Narita A, Matías D, et al. pH-Responsive pharmacological chaperones for rescuing mutant glycosidases. Angew Chem Int Ed Engl 2015;54:11696–700.
58. Yu Y, Mena-Barragán T, Higaki K, et al. Molecular basis of 1-deoxygalactonojirimycin arylthiourea binding to human α-galactosidase A: pharmacological chaperoning efficacy on Fabry disease mutants. ACS Chem Biol 2014;9:1460–9.
59. Suzuki H, Ohto U, Higaki K, et al. Structural basis of pharmacological chaperoning for human β-galactosidase. J Biol Chem 2014;289:14560–8.
60. Takai T, Higaki K, Aguilar-Moncayo M, et al. A bicyclic 1-deoxygalactonojirimycin derivative as a novel pharmacological chaperone for GM1 gangliosidosid. Mol Ther 2013;21:526–32.
61. Ríquez-Cuadro R, Matsumoto R, Ortega-Caballero F, et al. Pharmacological chaperones for the treatment of α-mannosidosis. J Med Chem 2019;62:5832–43.
62. de la Fuente A, Ríquez-Cuadro R, Verdauguer X, et al. Efficient stereoselective synthesis of 2-acetamido-1,2-dideoxy-yallonojirimycin (DNJNAc) and sp(2)-iminosugar conjugates: novel hexosaminidase inhibitors with discrimination capabilities between the mature and precursor forms of the enzyme. Eur J Med Chem 2016;121:926–38.
63. de la Fuente A, Mena-Barragán T, Farrar-Tobar RA, et al. Stereoselective synthesis of 2-acetamido-1,2-dideoxy-yallonojirimycin (DNJNAc) and ureido-DNJNAc derivatives as new hexosaminidase inhibitors. Org Biomol Chem 2015;13:6500–10.
64. Gławar AFG, Best D, Ayers BJ, et al. Scalable syntheses of both enantiomers of DNJNAc and DGNJAc from glucuronolactone: the effect of N-alkylation on hexosaminidase inhibition. Chem Eur J 2012;18:9341–59.
65. Grabosch C, Kleinert M, Lindhorst TK. Glyco-SAMs by "dual click": thiourea-bridged glyco-OEG azides for cycloaddition on surfaces. Synthesis 2010; 2010:828–36.
66. Carbajo-Gordillo AI, Jiménez Blanco JL, Benito JM, et al. Click synthesis of size- and shape-tunable star polymers with functional macrocyclic cores for synergistic DNA complexation and delivery. Biomacromolecules 2020;21:5173–88.
67. Abbas SY, Al-Harbi RAK, Sh El-Sharief MAM. SAM. Synthesis and anticancer activity of thiourea derivatives bearing a benzodioxole moiety with EGFR inhibitory activity, apoptosis assay and molecular docking study. Eur J Med Chem 2020;198:112363.
68. Doğan SD, Gündüz MG, Doğan H, et al. Design and synthesis of thiourea-based derivatives as Mycobacterium tuberculosis growth and enoyl acyl carrier protein reductase (InhA) inhibitors. Eur J Med Chem 2020;198:112363.
69. He ZX, Huo JL, Gong YP, An Q, et al. Design, synthesis and biological evaluation of novel thiosemicarbazone-indole derivatives targeting prostate cancer cells. Eur J Med Chem 2021;210:112970.
70. Nurieva EV, Alexeev AA, Zefirova ON. Cyclic isothiourea in drug design. Chem Heterocycl Comp 2021;57:889–99.
71. Mena-Barragán T, García-Moreno MI, Sevšek A, et al. Probing the Inhibitor versus chaperone properties of sp2-iminosugars towards human β-glucocerebrosidase: a picomolar chaperone for Gaucher disease. Molecules 2018;23:927.
72. Maier T, Strater N, Schuette CG, et al. The X-ray crystal structure of human beta-hexosaminidase B provides new insights into Sandhoff disease. J Mol Biol 2003;328:669–81.
73. Lemieux MJ, Mark BL, Cherney MM, et al. Crystallographic structure of human beta-hexosaminidase A: interpretation of Tay-Sachs mutations and loss of GM2 ganglioside hydrolysis. J Mol Biol 2006;359:913–29.
74. Bateman KS, Cherney MM, Mahuran DJ, Tropak M, et al. Crystal structure of β-hexosaminidase B in complex with pyrimethamine, a potential pharmacological chaperone. J Med Chem 2011;54:1421–9.
75. Lieberman RL, D’aquino JA, Ringe D, Petsko DA. The effects of pH and iminosugar pharmacological chaperones on lysosomal glycosidase structure and stability. Biochemistry 2009;48:4816–27.
76. Nödling AR, Jakab G, Schreiner PR, Hilt G. 31P NMR spectroscopically quantified hydrogen-bonding strength of thioureas and their catalytic activity in Diels–Alder reactions. Eur J Org Chem 2014; 2014:6394–8.
77. Nickisch R, Gabrielsen SM, Meier MAR. Novel access to known and unknown thiourea catalyst via a multicomponent-reaction approach. ChemistrySelect 2020;5:11915–20.
78. Aguilar M, Gloster TM, García-Moreno MI, et al. Molecular basis for beta-glucosidase inhibition by ring-modified calystegine analogues. ChemBioChem 2008;9:2612–8.
79. Aguilar-Moncayo M, Gloster TM, Turkenburg JP M, García-Moreno MI, et al. Glycosidase inhibition by ring-modified castanospermine analogues: tackling enzyme selectivity by inhibitor tailoring. Org Biomol Chem 2009;7:2738–47.
80. Brumshtein B, Aguilar-Moncayo M, Juan M, Benito JM, et al. Cyclodextrin-mediated crystallization of acid β-glucosidase in complex with amphiphilic bicyclic nojirimycin analogues. Org Biomol Chem 2011;9:4160–7.
81. González-Cuesta M, Siddhu P, Ashmus RA, et al. Bicyclic picomolar OGA inhibitors enable chemoproteomic mapping of its endogenous post-translational modifications. J Am Chem Soc 2022;144:832–44.
82. Trost O, Olson AJ. Autodock Vina: improving the speed and accuracy of docking with a new scoring function, efficient optimization, and multithreading. J Comput Chem 2010;31:455–61.
83. Sastry GM, Adzhigirey M, Day T, et al. Protein and ligand preparation: parameters, protocols, and influence on virtual screening enrichments. J Comput Aided Mol Des 2013;27:221–34.