Non-steroidal inhibitors of Drosophila melanogaster steroidogenic glutathione S-transferase Noppera-bo

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Introduction

Insect growth regulators (IGRs) can be developed by elucidating the molecular mechanisms of insect-specific biological events. Because insect molting, and metamorphosis are controlled by ecdysteroids, their biosynthetic pathways can serve as targets for IGR development. The glutathione S-transferase Noppera-bo (Nobo), which is conserved in dipteran and lepidopteran species, plays an essential role in ecdysteroid biosynthesis. Our previous study using 17β-estradiol as a molecular probe revealed that Asp113 of Drosophila melanogaster Nobo (DmNobo) is essential for its biological function. However, to develop IGRs with a greater Nobo inhibitory activity than 17β-estradiol, further structural information is warranted. Here, we report five novel non-steroidal DmNobo inhibitors.

Analysis of crystal structures of complexes revealed that DmNobo binds these inhibitors in an Asp113-independent manner. Among amino acid residues at the substrate-recognition site, conformation of conserved Phe39 was dynamically altered upon inhibitor binding. Therefore, these inhibitors can serve as seed compounds for IGR development.

Keywords: insect growth regulator, ecdysone, ecdysteroid, glutathione S-transferase, structure–activity relationship.
the IGR targets that has historically received great attention is ecdysteroid, which plays pivotal roles in regulating the development and physiology of arthropods, including insects.\textsuperscript{2,3} Since ecdysteroids are not present in animals other than arthropods, molecules involved in their biosynthesis, secretion, circulation, and reception may serve as potent targets for IGR development.

Ecdysteroids are biosynthesized from dietary sterols.\textsuperscript{2,3} Each reaction in the biosynthetic pathway of the ecdysteroid 20-hydroxyecdysone (20E) is catalyzed by a specific ecdysteroidogenic enzyme.\textsuperscript{4,6} Several of these enzymes have been identified over the last two decades, including Noppera-bo (Nobo),\textsuperscript{5–7} Neverland,\textsuperscript{8,9} Non-molting glossy/Shroud,\textsuperscript{10} Spook/CYP307A1,\textsuperscript{11,12} Spookier/CYP307A2,\textsuperscript{12} CYP6T3,\textsuperscript{13} Phantom/CYP306A1,\textsuperscript{14,15} Disembodied/CYP302A1,\textsuperscript{16} Shadow/CYP315A1,\textsuperscript{16} and Shade/CYP314A1.\textsuperscript{17} Loss-of-function mutations of genes encoding these enzymes result in developmental lethality. Specifically, in Drosophila melanogaster, complete loss-of-function mutants of nobo, shroud, spook, phantom, disembodied, shade, and shadow, often classified as Halloween mutants, typically exhibit embryonic lethality due to loss of differentiated cuticular structures.\textsuperscript{18} Thus far, the functions of these enzymes have been characterized genetically and some of them have also been analyzed biochemically.\textsuperscript{3,19}

Among the Halloween genes, nobo encodes a member of the epsilon class of cytosolic glutathione S-transferases (GSTs, EC 2.5.1.18; hereafter GSTEs).\textsuperscript{20–22} Typically, glutathione (GSH) binds to the GSH-binding site (G-site) of GST and functions of Nobo.\textsuperscript{22} However, inhibitory effects of compounds other than EST should be identified for IGR development, because EST may act as an endocrine disruptor in vertebrates. Therefore, non-steroidal DmNobo inhibitors identified from a chemical compound library must be analyzed. Here, we report five novel inhibitors and some derivatives of DmNobo as well as the crystal structures of their complexes with DmNobo. These inhibitors do not interact with Asp113 and induce a conformational change of Phe39. Our findings offer a novel strategy for IGR development.

**Materials and methods**

1. **Chemical compounds**

A chemical library containing 9,600 compounds was provided by the Drug Discovery Initiative (DDI), The University of Tokyo, Japan. TDP011 (IUPAC name: 4-bromo-2-[(4-(3-methoxyphenyl)-2,2-dimethyl-1H-pyrimidin-6-yl)phenol] and TDP012 (IUPAC name: 1-(4-fluorobenzyl)-1H-thieno[3,2-c][1,1]thiazin-4(3H)-one) were purchased from ChemDiv (Cat# K915-0968 and C200-3720, respectively). TDP044 (IUPAC name: 2,2’-(1,1-ethanediyl)bis(3-hydroxy-5,5-dimethyl-2-cyclohexen-1-one) was purchased from Labotest (Cat# LT00014582). TDP045 (IUPAC name: 5,5-dimethyl-1,3-cyclohexadiene) was purchased from Fluorochem (Cat# 011151). 3,4-DNADCF was prepared as previously described.\textsuperscript{20} TDP013 (IUPAC name: 2-(5-tert-butyl-2-methyl-benzensulfonylamo)-benzoic acid, PubChem CID: 2360625), TDP015 (IUPAC name: 6-(6,7-dihydrothieno[3,2-c]pyridin-5(4H)-yl)-3-pyridinamine, PubChem CID: 1677781) and TDP046 were synthesized as described in Supplementary Information (Schemes S1–S3, Supplementary Methods).

2. **Protein expression, purification, and crystallization**

The DmNobo protein was expressed, purified, and crystallized as previously described.\textsuperscript{22} Briefly, DmNobo was expressed in Escherichia coli BL21 (DE3) harboring the pCold III_DmNobo plasmid and purified by GSH affinity chromatography with Gluthathione Sepharose 4B (Cytiva, Tokyo, Japan) and size-exclusion chromatography with Superdex 200 HiLoad 16/600 (Cytiva). Conditions of the reservoir solution for DmNobo crystallization were optimized as 34% (v/v) PEG400 in 80 mM Bis-Tris (pH 6.4). Crystals of substrate complexes were prepared by soaking them in artificial mother liquor (42.5% [v/v] PPG 400 in 100 mM Bis-Tris [pH 6.4]) containing each of 30 mM compound with or without 10 mM GSH for 2 days.

3. **GST activity inhibition assay**

High-throughput screening for DmNobo inhibitors using a 384-well format (20 µL per well) with 9,600 small molecules from a chemical library (DDI, The University of Tokyo) was conducted as previously described.\textsuperscript{22} Then, in vitro GST activity inhibition assay was performed to determine the 50% inhibitory concentration (IC\textsubscript{50}) value of six chemical compounds (Fig. 2), as described by Koiwai et al.,\textsuperscript{22} with minor modifications. Compound solutions were prepared as follows: TDP011, TDP012,
TDP013, and TDP015 were dissolved to 2.5 mM and TDP045 to 100 mM in DMSO. Dilution series of compounds, ranging from 100 mM to 5.1 µM for TDP016 and from 2.5 mM to 4.9 µM for all other compounds were prepared by two-fold serial dilution with their solvents. Next, 5 µL of each compound solution in the dilution series was mixed with 245 µL of solution A (100 mM sodium phosphate buffer [pH 6.5], 50 ng·mL−1 DmNobo protein), and 100 µL of the mixture was dispensed on a 96-well plate. Then, 100 µL of solution B (2 µM 3,4-DNADCF and 100 mM sodium phosphate buffer [pH 6.5]) was added to each well. In summary, the final reaction system (3,4-DNADCF and 100 mM sodium phosphate buffer [pH 6.5]) was composed of 100 mM sodium phosphate buffer [pH 6.5], 50 ng·mL−1 DmNobo protein, and 100 µL of the mixture was dispensed on a 96-well plate. Then, 100 µL of solution B (2 µM 3,4-DNADCF and 100 mM sodium phosphate buffer [pH 6.5]) was added to each well. In summary, the final reaction system was composed of 100 mM sodium phosphate buffer [pH 6.5], 25 ng·mL−1 DmNobo protein, 1 mM GSH, 0.005% Tween 20, and 1 µM 3,4-DNADCF. The GSH-conjugated product was excited at 485 nm, and the fluorescence intensity at 538 nm was measured every 30 sec for 3 min using Fluoroskan Ascent FL (Thermo Fisher Scientific, Waltham, USA). The IC50 values were calculated as described by Koiwai et al.22 The assay was performed in duplicate.

4. Crystal structure determination

Crystals of proper size were picked up with MicroLoop (MiTeGen, New York, USA), flash-frozen in liquid nitrogen, and stored in Uni-Pucks (Molecular Dimensions, Ohio, USA). Diffraction data were collected at beamlines BL-1A, BL-5A, BL-17A, and AR NE-3A of the Photon Factory in KEK (Tsukuba, Japan) and at beamline 5A of the Taiwan Photon Source (Hsinchu, Taiwan). The diffraction datasets collected from the Photon Factory were automatically processed and scaled using XDS,26 POINTLESS,27 and AIMLESS,28 which are integrated in PReMo,29 and those collected from the Taiwan Photon Source were processed and scaled using XDS and AIMLESS. Crystallographic statistics are summarized in supplemental Table S1.

For automated protein crystallography, we developed the software PEINTS (https://github.com/KotaroKoiwai/PEINTS), in which phases of crystal structures can be determined by the molecular replacement (MR) method with MOLREP30 using the crystal structure of the apo form of DmNobo (PDB ID: 6KEN) as a search model. Molecular models were initially refined using REFMAC5.31 Models were built manually using COOT32 and the molecular models were further refined with PHENIX. REFINEM33. Simulated annealing mFo−DFc omit-maps of compounds and GSH were calculated using PHENIX.REFINE. Interactions between DmNobo and GSH or compounds were analyzed using PoseView34 and Discovery Studio Visualizer (Dassault Systèmes BIOVIA, Discovery Studio Modeling Environment, Release 2020, San Diego: Dassault Systèmes, 2019).

The volume of DmNobo binding cavity was calculated using the Channel Finder program in 3V,35 with radii of 4 and 1 Å for the outer and inner probes, respectively. The root-mean-square deviation (RMSD) values from the least-squares fitting of the DmNobo structures were calculated using GESAMT.36 Atom pairs within a 4.0 Å distance were defined as direct contacts. All molecular graphics were prepared using the PyMOL Molecular Graphics System version 1.7.6 (Schrodinger, NY, USA).

The X-ray data and coordinates presented in this study are deposited in the Protein Data Bank (PDB; https://pdbj.org/) under the PDB IDs 7DAX, 7DAY, 7DAZ, 7DB0, 7DB1, 7DB2, 7DB3, and 7DB4.

5. Molecular dynamics simulations

Molecular dynamics (MD) simulations for DmNobo–Apo (PDB ID: 6KEO) and DmNobo–GSH (PDB ID: 6KEN) were performed as previously described.22 The structures were processed to assign bond orders and hydrogenation. The ionization states of GSH at pH 7.0±2.0 were predicted using Epik,37 and H-bond optimization was conducted using PROPKA.38 Energy minimization was performed in Maestro using the OPLS3 force field.39 Preparation for MD simulations was conducted using the Molecular Dynamics System Setup Module of Maestro (Schrodinger, NY, USA). DmNobo–Apo and DmNobo–GSH were subjected to energy minimization and placed in an orthorhombic box with a buffer distance of 10 Å to create a hydration model, and the TIP3P water model40 was used for the hydration model. NaCl (0.15 M) served as the counterion to neutralize the system. The MD simulations were performed using Desmond software, version 2.3 (Schrodinger, NY, USA). The cut-off radii for van der Waals was 9 Å. The cut-off values of the time step, initial temperature, and pressure of the system were set to 2.0 fs, 300 K, and 1.01325 bar, respectively. The sampling interval during the simulation was set to 10 ps. Finally, we performed MD simulations using the NPT ensemble for 100 nsec.

Results

1. Identification of DmNobo inhibitors by high-throughput screening

We performed high-throughput screening for the inhibitors of GSH conjugation activity of DmNobo using the artificial fluorescent substrate 3,4-DNADCF. From 9,600 chemical compounds obtained from DDI,25 we identified six compounds with IC50 values below 10 µM. One of the inhibitors was EST, which has a steroidal backbone similar to ecdysteroids.22,25 The other five compounds did not have a steroidal backbone and were designated TDP011 (IC50 = 6.72±1.48 µM), TDP012 (IC50 = 3.10±0.77 µM), TDP013 (IC50 = 8.94±1.65 µM), TDP015 (IC50 = 3.32±0.81 µM), and TDP044 (IC50 = 0.62±0.52 µM) compared with EST (IC50 = 2.33±0.08 µM) (Figs. 1, 2, Table 1). Since these compounds have various chemical structures, we expected them to bind with DmNobo in a different manner from EST.

2. Crystal structures of DmNobo complexed with the five inhibitors and their derivatives

To examine the structure–activity relationships of the novel inhibitors identified, we determined the crystal structures of their complexes with DmNobo (Supplemental Table S1). The resolutions of these structures were in the range of 1.40 to 1.80 Å, and inhibitor and water molecules could be clearly identified (Supplemental Fig. S1).
TDP044 possesses two dimedone ring structures (Fig. 1). Of note, a DmNobo crystal soaked in a solution containing TDP044 showed clear electron densities corresponding to only one dimedone-ring in either chain A or B (Supplemental Fig. S1 F). Although we have collected data from crystals soaked under various conditions, a clear electron density map for TDP044 was not observed. The reason behind such a change in the crystal remains unclear, possibly because of degradation or oxidation of TDP044 during soaking. Based on this observation, we examined whether dimedone (hereafter TDP045) physically interacted with DmNobo and inhibited its enzymatic activity. We determined the crystal structure of DmNobo complexed with TDP045. Although TDP045 did not exhibit a strong inhibitory activity against DmNobo (Fig. 2), the electron density of TDP045 could be clearly observed (Supplemental Fig. S1 G and H). Therefore, although TDP045 cannot be classified as a DmNobo inhibitor per se, we decided to include TDP045 instead of TDP044 for further crystal structure analysis to understand the possible mode of action of TDP044, a TDP045 derivative.

Although all compounds showed binding at the H-site, they could be categorized into two classes based on the GSH-dependency of their interaction with DmNobo. Crystallographic analysis with and without GSH revealed that the interactions of TDP011, TDP012, and TDP015 with DmNobo were GSH-dependent, while those of TDP013 and TDP045 with DmNobo were GSH-independent (Table 1). For comprehensive comparison, the DmNobo–inhibitor complex structures determined in this study and those reported in previous studies were mutually superposed by least-squares fitting using Cα atoms (Supplemental Table S2). The RMSD values were 0.14–0.48 Å for superpositions of the Cα atoms of chain A and 0.15–0.97 Å for superpositions of the Cα atoms of chains A and B (Supplemental Table S2), indicating that it is reasonable to consider that these crystal structures were essentially the same as one another, and no large-scale conformational changes occurred upon compound and/or GSH binding. Structural comparison, however, revealed several local conformational changes induced by compound binding (see below).

3. Residues interacting with the identified compounds
Interactions between compounds and residues at the H-site were analyzed using PoseView and Discovery Studio (Fig. 3). The analysis revealed that Pro15, Leu38, Phe39, Phe110, Met117, Val121, Leu208, Thr211, Met212, and Val215 were involved in hydrophobic interactions with compounds, while Ser14 and Ser114 formed specific hydrophilic interactions. Leu38 and Met117 interacted with the compounds most frequently. All inhibitors, except TDP045, interacted with three to seven hydrophobic residues at the H-site. Among hydrophobic residues at the H-site, Phe39 largely contributes to hydrophobic interactions with EST.22 However, TDP012 and TDP013 in a GSH-free form did not interact with the side chain of Phe39, suggesting that this side chain is not always necessary for hydrophobic interactions with the compounds.

Fig. 1. Chemical structures of the newly identified DmNobo inhibitors and their derivatives. TDP011, TDP012, TDP013, TDP015, and TDP044 were identified as hit compounds in high-throughput screening. The 6-membered rings of TDP011 are labeled with A, B, and C. Asterisk indicates the GSH-adduct site in crystal structure.
4. **GSH-independent interactions**

TDP013 and TDP045 could bind DmNobo without GSH. Of note, the conformation of these two compounds at the H-site was altered upon GSH binding through interaction or conjugation with GSH (Fig. 4A and 4B).

In the absence of GSH, TDP013 conformation was such that the planes of the two rings were roughly parallel to the α-helices. A large cavity, the G-site, was placed above the bound TDP013, and the unoccupied G-site was filled with water molecules, forming a hydrogen bond network. In the presence of GSH, TDP013 rotated at the H-site, assuming a conformation in which the planes of the rings were roughly perpendicular to the α-helices. This change likely resulted from the clash between the thiol group of GSH and TDP013. The rearrangement of TDP013 induced another conformational change of Phe39 (see below).

In the absence of GSH, TDP045 was located deep inside the H-site. One of the carbonyl groups of TDP045 interacted with Asp113 (2.6 Å), and the other formed a hydrogen bond with Ser114 (2.7 Å). While the IC$_{50}$ value of TDP045 was substantially larger than those of other compounds, its electron density could be clearly observed, suggesting that the binding of TDP045 is stable, at least in the crystal. Upon GSH binding, TDP045 ro-

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**Table 1.** Characters of DmNobo inhibitors

|                  | TDP011 | TDP012 | TDP013 | TDP015 | TDP044 | TDP045 | TDP046 |
|------------------|--------|--------|--------|--------|--------|--------|--------|
| # of bound molecules in the H-site | 1      | 1      | 1      | 1      | 2      | 1$^{(b)}$ | 1      |
| GSH requirement for binding | NO     | YES    | YES    | NO     | YES    | NO     | NO     |
| GS-conjugation | NO     | YES    | YES    | NO     | YES    | NO     | YES$^{(c)}$ |
| Interaction with Asp113 | YES    | NO     | NO     | NO     | YES    | YES$^{(c)}$ | NO     |
| Conformational change of Phe39 | NO     | NO     | YES    | YES    | YES    | NO     | NO     |
| Different conformations of the inhibitor in subunits A and B | NO     | NO     | YES    | YES    | NO     | NO     | NO     |
| IC$_{50}$ (µM)   | 2.33±0.08 | 6.72±1.48 | 3.10±0.77 | 8.94±1.65 | 3.32±0.81 | 0.62±0.52 | >1000 |
|                  |        |        |        |        |        |        | 2.08±0.04 |

$^{(a)}$PDB ID: 6KEP. $^{(b)}$Electron density map for a whole part of TDP044 was not observed clearly. $^{(c)}$TDP045 interacts with Asp113 only when no GSH binds to DmNobo. After conjugation, no interactions with Asp113 were observed.
tated at the H-site such that the C2 atom of TDP045 was conjugated with the thiol group of GSH (Fig. 4C and 4D). The covalent bond between TDP045 and GSH was clearly observed in the electron density map (Supplementary Fig. S1G), which was identical to the electron density map of chemically synthesized GS-dimedone (hereafter TDP046) (Supplementary Fig. S1H), indicating that TDP045 in DmNobo crystals is glutathionylated at the C2 atom.
5. **GSH-dependent interaction**

Under crystallographic conditions in this study, no electron densities of TDP011, TDP012, and TDP015 were observed in the absence of GSH (Supplementary Fig. S2A, B, C). However, electron densities of these inhibitors could be clearly observed in the presence of GSH. In addition, continuous electron densities were observed between these inhibitors and GSH (Supplementary Fig. S1A, B, E). These results suggest that TDP011, TDP012, and TDP015 require GSH for binding with DmNobo at the H-site.

TDP011 structure has three rings (A to C). The electron density map of the DmNobo–TDP011 complex suggested that conjugation with GSH opened the B-ring, resulting in a two-ring structure (Fig. 5A and Supplementary Fig. S3). The TDP011 part of the conjugated structure formed no direct or indirect hydrogen bond interactions with the protein. In the TDP012 complex, the TDP012 part of the conjugated structure formed two hydrogen bonds with Ser14 and Ser114 (Fig. 5C and 5D).

The interaction of TDP015 with DmNobo was rather unique. Two TDP015 molecules bound a single H-site of DmNobo (Fig. 5E). The electron density map suggested that the amino group in the aminopyridine moiety of one TDP015 molecule formed a covalent bond with the Sγ atom of GSH. The conjugated TDP015 molecule showed a kinked conformation, and the other non-conjugated TDP015 molecule showed a stretched conformation. Only one hydrogen bond was formed between the TDP015 molecules and DmNobo (Fig. 3E).

6. **Comparisons of compound conformations between subunits**

DmNobo is a dimeric protein, and each subunit harbors an H-site for compound binding. There are minor structural and/or physicochemical differences between subunits A and B. GS–TDP011 (hereafter, a compound in the crystal structure conjugated with GSH is indicated by the addition of GS), and GS–TDP013 showed conformational differences between subunits A and B. Simulated annealing-omit maps of the compounds were rather weak (Supplementary Fig. S1). These observations are consistent with the IC₅₀ values of these compounds (Fig. 2, Table 1), which were larger than those of the other inhibitors. Structural and/or physicochemical differences between the two H-sites may affect TDP011 and TDP013 conformation.
It has been reported that DmNobo interacts with EST via the hydrogen bond with Asp113 and the hydrophobic interactions with other amino acids of H-sites. While the newly identified compounds also formed hydrophobic interactions with DmNobo (Fig. 3), the mode of these hydrophobic interactions differed between EST and the newly identified compounds. Structural comparisons revealed that atoms of the compounds were frequently located in the space that the EST atoms did not occupy. Three small spaces were noted around EST (Fig. 6). First, space A was located below EST, that is between EST and surface A that was composed of Met117, Val121, Thr211, Met212, and Val215. Second, space B was located at the side of EST, that is between EST and surface B that was lined by Arg13, Ser14, Pro15, Leu38, Phe39, and Leu208. Finally, space C was located on the other side of EST, that is between EST surface C that was lined by Phe110, Asp113, Ser114, and Thr169. Atoms of the compounds

Fig. 5. TDP011, TDP012, and TDP015 can bind DmNobo with GSH. Positions of TDP011, TDP012, TDP015, and GSH in the DmNobo pocket are illustrated. The carbon structure of GSH is shown in yellow in A, C, and E. Rainbow colors in B, D, and F represent temperature factors (B-factors, Å²) of the inhibitor and protein atoms, respectively. (A and B) TDP011 (A, carbon in blue). (C and D) TDP012 (C, carbon in red). (E and F) TDP015 (E, carbon in orange).
utilized these spaces (or the inner surface of the H-site) (Fig. 6). Since most of the compounds have a non-planar structure, they need to use parts of spaces A, B, and C to be accommodated at the H-site. In some cases, space B was expanded by the conformational change of Phe39 (see below).

8. Conformational changes of residues at H-site
Structural comparison between the apo and compound-bound forms revealed conformational changes of residues at the H-site. DmNobo shows a conformational change of the side chain of Asp113 upon EST binding, forming a hydrogen bond with Asp113. Meanwhile, most of the compounds identified in this study did not form hydrogen bonds with Asp113. Therefore, no conformational changes were observed in Asp113 of DmNobo upon binding with any compound, except TDP045 (Fig. 7A).

As described above, one carbonyl group of TDP045 formed a hydrogen bond with Asp113 (2.6 Å) and the other interacted with Ser114 (2.7 Å). The $\chi_1$ angle of Asp113 was rotated by $\sim 25^\circ$, resulting in the formation of a hydrogen bond. In the DmNobo–TDP045–GSH complex, Asp113 did not show a conformational change. GSH conjugation of TDP045 changed its orientation at the H-site, losing the interaction with Asp113.

While the newly identified compounds, except TDP045, induced conformational changes of Asp113, some led to conformational changes of Phe39 (Fig. 7A). The bromophenol group of TDP011, the fluorobenzyl group of TDP012, the tert-butyl group of TDP013, and the thienopyridine group of the kinked TDP015...
molecule clashed with the side chain of Phe39; therefore, binding of TDP011, TDP012, and TDP015 rotated the side chain of Phe39, exposing Thr211 and Val215 on the inner surface of H-site. The thienopyridine group of the kinked TDP015 molecule interacted with Val215, and the main chain atoms around Val215 showed significant shifts from their original position (Fig. 7B).

9. **Phe39 flexibility calculated by MD simulation**

Structural analysis showed that the inhibitors TDP011, TDP012, TDP013, and TDP015 induced a conformational change of Phe39 from a closed to an open form. However, the crystal structure of the DmNobo–EST–GSH complex did not show a conformational change of Phe39. Our previous MD simulations of the DmNobo–EST–GSH complex suggested that Phe39 is stable and remained in a closed form. Here, we performed MD simulations for DmNobo in apo and GSH-complexed forms (Fig. 8) and demonstrated that Asp113 was stable at the EST-bound form during simulation; however, the open form of Phe39 was often simulated (Fig. 8C and 8D). These results suggest that Phe39 is mobile without EST and can easily undergo a conformational change.

**Discussion**

In this study, we identified five small non-steroidal compounds that inhibit the GSH-conjugating activity of DmNobo. The crystal structures of DmNobo complexed with these inhibitors and their derivative suggested that the H-site of DmNobo harbors hydrophobic targets for compounds other than the previously identified ones using EST. These compounds and their complex structures with DmNobo will be beneficial for developing novel IGRs targeting Nobo and new drugs against other GSTs.

Our previous study showed that Asp113 is essential for the binding of DmNobo with EST, indicating that this residue can serve as a target for IGR development. Among the five compounds identified in this study, only one compound—TDP045—interacted with Asp113 for binding. Notably, however, the carbon skeletons with functional groups, which are required for interaction with DmNobo, largely differ between EST and TDP045. EST contains an aromatic ring with a hydroxyl group, while TDP045 contains a non-aromatic ring with carbonyl and/or hydroxyl groups. Since these functional groups of TDP045 form a hydrogen bond with Asp113, DmNobo is unlikely to require an aromatic ring for forming a complex through this residue. Considering that Asp113 of DmNobo is essential for ecdysteroid biosynthesis, an unknown endogenous substrate of DmNobo containing non-aromatic A-rings likely serves as an intermediate in ecdysteroid biosynthesis. Notably, a recent study reported that DmNobo shows isomerase activity against Δ5-androstene-3,17-dione, in which a ketone group is located on a cyclohexane group in the A-ring.

Crystal structures of DmNobo with the identified inhibitors suggested that hydrophobic residues, particularly Leu38, Phe39, Leu208, and Met117, at the H-site mainly contribute to the binding of these inhibitors (Fig. 3). Among these hydrophobic residues, Phe39 is unique. It undergoes a conformational change upon inhibitor binding (TDP011, TDP012, and TDP015), such that Thr211 and Val215 on the inner surface of the H-site are exposed. A buried hydrophobic surface composed of Thr211 and Val215 can, therefore, serve as a target for DmNobo inhibitors. Despite the observed conformational changes of Phe39, we do not think that similar conformational changes occur in vivo to bind an unidentified substrate of Nobo. Since Phe39 is highly conserved among Nobo proteins, it is reasonable to consider...
that the substrate of Nobo interacts with Phe39 in the closed form. Conversely, Thr211 and Val215 are not conserved, suggesting that the unidentified substrate is unlikely to interact with Thr211 and Val215.

TDP015 has a unique binding mode. Two TDP015 molecules bind at the H-site. Of these two, the kinked TDP015 molecule interacts with the exposed Thr211 and Val215 as well as with Phe39, while the other TDP015 molecule forms hydrophobic interactions with residues lining the H-site and π–π interactions with the kinked molecule. Despite these interactions, however, the IC₅₀ value of TDP015 was comparable to that of the other compounds. This may be due to entropy loss caused by the binding of two molecules and lack of interaction with Asp113. Considering the higher temperature factors for the inhibitor atoms (Fig. 5), hydrophobic interactions may not be sufficient for stable binding of these inhibitors. Conformational differences be-

Fig. 8. Molecular dynamics simulations of DmNobo in apo and GSH complex forms. Molecular dynamics simulation of DmNobo–Apo and DmNobo–GSH was performed for 100 nsec at 300 K with water molecules. (A) Distance between the Oδ atom of Asp113 and the O3 atom of a dummy EST molecule. The dummy EST molecule (dumEST) was placed in the models at 0 nsec by superposition of Ca atoms of chain A of DmNobo–EST–GSH. The coordinate of the O3 atom of the dumEST was extracted, and the distance between the Oδ atom of Asp113 at each frame and the O3 atom of the dumEST was calculated. The distances in DmNobo–Apo (black) and DmNobo–EST–GSH (blue) were plotted. (B) Snapshots of Asp113 at the indicated frames. The Ca atoms of chain A of the crystal structure of DmNobo–Apo (gray) were superimposed with those of an MD model of DmNobo–Apo (MD: Apo) at 0 nsec. Ca atoms of chain A of DmNobo–EST–GSH (MD: EST–GSH) were superposed with those of DmNobo–Apo. The absolute values of the χ₁ torsion angle of Phe39 of chains A (black line) and B (gray line) of DmNobo–Apo (C) and DmNobo–GSH (D) were plotted. Representative MD models of DmNobo–GSH with open- and closed-form Phe39 are shown, whose absolute values of χ₁ torsion angle are ∼75° at 0.0 nsec and ∼175° at 5.0 nsec.
between subunits A and B of TDP013, GS-TDP013, and TDP011 (Fig. 4) also support this notion. These weak interactions could be compensated by conjugation with GSH. To improve the affinity of TDP015 to DmNobo, (1) a linkage between the two TDP015 molecules may be required to avoid entropy loss and (2) a hydrophilic group may be added to facilitate interaction with Asp113.

In this study, we found that compounds, TDP011, TDP012, TDP015, and TDP045 were glutathionylated in the presence of GSH. The glutathionylation should not be enzymatically reacted because it needs sulfenylation of the S atom, and may be due to oxidation during crystal preparation. However, these structures can be informative for further development of inhibitors with higher activity. Indeed, from the crystal structure of DmNobo–TDP045–GSH, we could design a novel inhibitor against the G-site of DmNobo, GS-dimedone (TDP046 in this study). By application of the conjugated products, we expect that we can also develop novel inhibitors with higher specificity and inhibitory activity.

In conclusion, we identified five novel Nobo inhibitors, which might be useful for developing new IGRs targeting ecdysteroid biosynthesis. These inhibitors interact with the H-site of DmNobo mainly through hydrophobic interactions. Structural analysis revealed strategies to increase their affinities to DmNobo.

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Electronic supplementary materials

The online version of this article contains supplementary materials (Supplemental Schemes S1–S3, Figs. S1–S3, Tables S1, S2), which are available at http://www.jsagest.jst.go.jp/browse/jpestics/.

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