**RESEARCH PAPER**

**In vivo effects of the pure aryl hydrocarbon receptor antagonist GNF-351 after oral administration are limited to the gastrointestinal tract**

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**BACKGROUND AND PURPOSE**

GNF-351 is a potent aryl hydrocarbon receptor (AHR) antagonist that inhibits dioxin response element-dependent and independent activities. Here, the absorption, metabolism and in vivo AHR antagonist activity of GNF-351 were investigated.

**EXPERIMENTAL APPROACH**

LC-MS metabolomics was used to analyse GNF-351 metabolism in vitro and in vivo. Recombinant drug-metabolizing enzymes were employed to determine the enzymes involved in GNF-351 metabolism. Analysis of target AHR genes was performed to investigate the inhibitory effects of GNF-351 towards AHR activation.

**KEY RESULTS**

Several phase I metabolites were generated after GNF-351 was incubated with microsomes from human or mouse liver and intestine, including two oxidized GNF-351 and one tri-demethylated GNF-351. Poor absorption from the intestine resulted in no detectable levels of GNF-351 in mouse serum (0–6 h) and urine (24 h) and almost all GNF-351 was found in the faeces after 24 h. Analysis of faeces further revealed all the in vitro phase I metabolites. Novel metabolites were detected, including one di-oxidized GNF-351, two oxidized and tri-demethylated GNF-351, one dehydrogenated product of oxidized GNF-351, and one sulfation product of di-oxidized GNF-351. Cytochromes P450 were demonstrated to be the major enzymes involved in metabolism of GNF-351. After oral administration to mice, GNF-351 readily inhibited β-naphthoflavone-induced AHR activation in ileum and colon, but not that in the liver.

**CONCLUSION AND IMPLICATIONS**

While poor absorption and extensive metabolism after oral administration limited the in vivo effects of the pure AHR antagonist GNF-351 in liver, it could be used to inhibit AHR activation in intestine and colon.
Introduction

Aryl hydrocarbon receptor (AHR), a basic-helix-loop-helix Per-Arnt-Sim transcription factor, has a key role in xenobiotic-induced toxicity and carcinogenesis (Barouki et al., 2007; Vondracek et al., 2011; receptor nomenclature follows Alexander et al., 2013). The most high-affinity AHR agonist is the notorious environmental and industrial toxin 2,3,7,8-tetrachlorodibenzo-p-dioxin (TCDD) and TCDD toxicity is due to AHR activation (Fernandez-Salguero et al., 1995; Matsubara et al., 2012). Development of AHR antagonists has become an important topic in recent years because of their potential therapeutic role. For example, the AHR antagonist 2-methyl-2H-pyrazole-3-carboxylic acid prevented TCDD-associated toxicity (Kim et al., 2006). Additionally, AHR antagonists can attenuate Th17 cell development in vivo and subsequent secretion of IL-17 and IL-22 (Veldhoen et al., 2009). GNF-351 (Figure 1A) is a recently developed AHR antagonist with the capacity to inhibit dioxin response element-dependent and -independent activity. Additionally and in contrast to other reported antagonists, such as α-naphthoflavone, GNF-351 does not exhibit partial agonist potential (Smith et al., 2011).

Pharmacokinetic properties are important determinants for the research and development (R&D) of the therapeutic agents, including the absorption and elimination characteristics, and poor pharmacokinetics will limit the development of drugs for clinical applications. For example, the clinical use of rifaximin is limited to gastrointestinal diseases because of its poor absorption (Ma et al., 2007). In contrast, the closely related antibiotic rifampicin has excellent bioavailability and is used to treat systemic infections such as tuberculosis (Li et al., 2013).

Ultra performance liquid chromatography-electrospray ionization-quadrupole time-of-flight mass spectrometry (UPLC®-ESI-QTOFMS)-based metabolomics has become an important tool to determine the metabolism, toxicity and metabolic behaviour of xenobiotics (Johnson and Gonzalez, 2012; Johnson et al., 2012). The present study used metabolomics to elucidate the absorption behaviour and the main metabolic pathway of GNF-351, including identification of metabolites and the major drug-metabolizing enzymes (DMEs) involved in metabolism. The influence of absorption and metabolism of GNF-351 on use of GNF-351 as an efficient AHR antagonist in vivo, was also examined.

Figure 1

Chromatography and MS/MS fragmentation of GNF-351. A. The structure of GNF-351; B. Chromatographic separation of GNF-351; C. MS/MS fragmentation pattern of GNF-351.
Methods

Animals
All animal care and experimental procedures complied with protocols approved by the National Cancer Institute Animal Care and Use Committee. The mice were maintained under a standard 12 h light, 12 h dark cycle with water and chow provided ad libitum. All studies involving animals are reported in accordance with the ARRIVE guidelines for reporting experiments involving animals (Kilkenny et al., 2010; McGrath et al., 2010). A total of 22 animals were used in the experiments described here.

In vitro metabolism of GNF-351 in liver microsomes (LMs), intestine microsomes (IMs) and recombinant DMEs
Livers from untreated 6- to 8-week-old male C57BL/6J mice were homogenized to prepare microsomes (MLM) as previously described (Fang et al., 2012). Human liver microsomes (HLM) were purchased from BD Gentest Corp. (Woburn, MA, USA). Human intestinal microsomes (HIM) were purchased from BD Gentest Corp. (Woburn, MA, USA). Mouse intestinal microsomes (MIM) were prepared as described for mouse liver microsomes. The phase I incubation system (200 μL) contained 50 mM Tris-HCl buffer solution (pH = 7.4), 0.5 mg mL⁻¹ HLM, MLM, HIM or MIM, 5 mM MgCl₂, 100 μM GNF-351, and 1 mM freshly prepared NADPH. After 0.5 h incubation at 37°C, the reaction was stopped using 200 μL cold 50% aqueous methanol containing 5 μM chlorpropamide. After centrifuging at 14 000× g for 20 min, a 5 μL aliquot of the supernatant was injected into a Waters UPLC-ESI-QTOFMS system (Waters Corporation, Milford, MA, USA).

UPLC-ESI-QTOFMS
An Acquity C18 BEH UPLC column (Waters Corporation) was employed to separate components in serum, urine, faeces and microsomal incubation samples. The mobile phase consisted of water containing 0.1% formic acid (A) and acetonitrile containing 0.1% formic acid (B). The following gradient condition was used: 100% A for 0.5 min, increased to 100% B over the next 7.5 min and returned to 100% A in the last 2 min. The flow rate of mobile phase was set 0.5 mL min⁻¹. Data were collected in positive ion mode on a Waters Q-Tof Premier mass spectrometer, which was operated in full-scan mode at 50–850 m/z. Nitrogen was used as both cone gas (50 L·h⁻¹) and desolvation gas (600 L·h⁻¹). Source desolvation temperatures were set at 120 and 350°C respectively. The capillary voltage and cone voltage were 3000 and 20 V respectively. The structures of metabolites were elucidated by tandem MS fragmentography with collision energies ranging from 15 to 40 eV.

Multivariate data analysis
MarkerLynx software (Waters Corporation) was used to deconvolute the chromatographic and mass spectrometric data. A multivariate data matrix containing information on sample identity, ion identity (retention time and m/z), and ion abundance was generated through centroiding, deisotoping, filtering, peak recognition and integration. The data matrix was further analysed using SIMCA-P 12.0 software (Umetrics, Kinnelon, NJ, USA). Orthogonal partial least squares data analysis (OPLS-DA) was adopted to analyse the data to identify the major latent variables in the data matrix. Potential metabolites were identified through analysing the ions contributing to the separation of sample groups in the loading scatter plots.

In vivo investigation of GNF-351 inhibition of β-naphthoflavone (BNF)-induced AHR activation
Twenty-one male C57BL/6J mice were divided into three groups: control group (n = 7), BNF group (n = 7), and BNF + GNF-351 group (n = 7). For administration of BNF, mice were given BNF, 5 mg·kg⁻¹, dissolved in corn oil. The
corresponding vehicle was used as control. For the BNF + GNF-351 group, GNF-351 (5 mg·kg$^{-1}$) dissolved in coin oil was given by oral gavage 5 min before the dose of BNF. The mice were killed 12 h later and the liver, duodenum, jejunum, ileum and colon were taken for mRNA preparation.

**Real-time quantitative PCR measurements of mRNA**

Real-time quantitative PCR (qPCR) was performed to investigate the expression of Cyp1a1 mRNA. RNA was isolated from

### Table 1

List of detected metabolites in *in vitro* incubation system or mouse faeces

| Symbol | RT (min) | Observed m/z | Formula | Mass error (ppm) | Identity |
|--------|----------|--------------|---------|------------------|----------|
| I      | 5.38     | 412.225      | C$_{20}$H$_{21}$N$_5$[H$^+$] | 0.0     | GNF-351         |
| II     | 5.38     | 269.154      | C$_{13}$H$_{12}$N$_3$[H$^+$] | 9.3     | GNF-351 fragment|
| III    | 5.38     | 144.082      | C$_{10}$H$_7$N$_2$[H$^+$]  | 4.9     | GNF-351 fragment|
| IV     | 5.38     | 227.105      | C$_{11}$H$_{13}$N$_2$[H$^+$] | 2.2    | GNF-351 fragment|
| V      | 5.38     | 370.179      | C$_{20}$H$_{21}$N$_5$[H$^+$] | 2.7    | GNF-351 fragment|
| VI     | 5.15     | 428.219      | C$_{20}$H$_{21}$N$_5$O[H$^+$] | -2.1   | Oxidized GNF-351|
| VII    | 4.89     | 428.220      | C$_{20}$H$_{21}$N$_5$O$_2$[H$^+$] | -2.1  | Oxidized GNF-351|
| VIII   | 4.78     | 370.180      | C$_{13}$H$_{12}$N$_3$[H$^+$] | 2.7    | Tri-demethylated GNF-351|
| G1     | 4.86     | 588.259      | C$_{20}$H$_{21}$N$_5$O$_3$[H$^+$] | 3.2    | GNF-351 glucuronide|
| G2     | 4.12     | 588.253      | C$_{20}$H$_{21}$N$_5$O$_3$[H$^+$] | -7.0   | GNF-351 glucuronide|
| F1     | 4.60     | 444.214      | C$_{20}$H$_{21}$N$_5$O$_3$[H$^+$] | 2.7    | Dioxidized GNF-351|
| F2     | 4.19     | 386.174      | C$_{20}$H$_{21}$N$_5$O$_3$[H$^+$] | 2.8    | Oxidized and tri-demethylated GNF-351|
| F3     | 3.75     | 386.172      | C$_{20}$H$_{21}$N$_5$O$_3$[H$^+$] | -2.3   | Oxidized and tri-demethylated GNF-351|
| F4     | 3.09     | 402.169      | C$_{20}$H$_{21}$N$_5$O$_3$[H$^+$] | 3.0    | Dioxidized and tri-demethylated GNF-351|
| F5     | 4.06     | 426.204      | C$_{20}$H$_{21}$N$_5$O$_3$[H$^+$] | -0.5   | Dehydrogenated product of oxidized GNF-351|
| F6     | 4.20     | 524.174      | C$_{20}$H$_{21}$N$_5$SO$_3$[H$^+$] | 4.6    | Sulfation product of dioxidized GNF-351|

**Figure 2**

Identification of phase I metabolites in MLMs and phase II metabolites in HLMs. A. Scores plot of a PCA model from MLM phase I incubation mixture without and with GNF-351. B. OPLS loading S-plot of MLM phase I incubation mixture; The p(corr)[1] values represent the interclass difference, and the p[1] values represent the relevant abundance of ions. C. Scores plot of a PCA model from HLM phase II incubation mixture without and with GNF-351. D. OPLS loading S-plot of HLM phase II incubation mixture; The p(corr)[1] values represent the interclass difference, and the p[1] values represent the relevant abundance of ions. Representative ions (I, II, III, IV, VI and G1) were given.
cell cultures or animal tissues using TRIzol reagent (Invitrogen, Carlsbad, CA, USA), and cDNA generated from 1 μg RNA with a SuperScript II™ Reverse Transcriptase Kit (Life Technologies, Grand Island, NY, USA). qPCR was carried out using SYBR green PCR master mix and ABI Prism 7900HT Sequence Detection System (Applied Biosystems, Foster City, CA, USA).

Data analysis
The experimental data are given as mean ± SEM. Statistical analysis was carried out using GraphPad Prism 5.0 (GraphPad Software, Inc., La Jolla, CA, USA). Comparisons between two groups were performed using a 2-tailed unpaired Student’s t-test or Mann–Whitney U-tests. Means of more than two groups were compared using one-way ANOVA.

Materials
Alamethicin, NADPH, UDPGA as uridine 5′-diphosphoglucuronic acid trisodium salt were purchased from Sigma-Aldrich (St. Louis, MO). GNF-351 was synthesized as previously described (Smith et al., 2011). All other reagents were of the highest grade commercially available.

Results
In vitro metabolism profile of GNF-351 in liver and intestine
The parent compound, GNF-351, analysed using UPLC-ESI-QTOFMS, eluted at a retention time of 5.5 min (Figure 1B), with an m/z of 412.225, in agreement with a match for C₂₅H₂₅N₇ with a mass error of 0.0 ppm (Table 1). The main MS/MS fragmentation ions of GNF-351 contained 370, 281, 269, 239, 227 and 144, with the fragmentation pattern shown in Figure 1C. In vitro metabolism of GNF-351 was assessed using a microsomal incubation system and the data deconvoluted and analysed by principal components analysis (PCA). This model revealed two clusters corresponding to in vitro MLM phase I incubation samples in the presence and absence of GNF-351 (Figure 2A). The OPLS-DA loading plot showed the major ions contributing to the separation containing the parent compounds, the fragments of parent compounds and metabolites (Figure 2B). Detailed information on the I–VIII ions are shown in Table 1, and the trend plot of these ions (I–VIII) displayed in Supporting Information Fig. S1. Separation was also detected between MLM (or HLM)
incubation samples and the samples without LMs and without NADPH (Supporting Information Fig. S2). For the glucuronidation incubation system, the PCA model could separate the HLM incubation samples with and without substrates (Figure 2C). Besides the contribution of ions derived from the parent compounds and fragments towards this separation, two glucuronides were also observed to contribute to the separation (Figure 2D and Table 1). The representative chromatograms are displayed in Supporting Information Fig. S3. These two glucuronides were not detected in the MLM incubation system.

The chemical structures of the metabolites were elucidated on the basis of MS/MS fragmentation patterns (Figure 3). The oxidized metabolites VI and VII were identified at retention times of 5.15 and 4.89 min, and gave a match for C_{24}H_{25}N_{7}O with the mass error of −2.1 ppm. The same MS/MS fragments were produced for these two metabolites, containing ions of 410, 386, 368, 269, 227 and 160 Da. Compared with the fragments of the parent compound, the fragment ion 160 was 16 Da higher than one fragment ion of parent compound 144, indicating the addition of an oxygen atom in the upper fraction of GNF-351 as indicated (Figure 3). Ion VIII eluted at 4.78 min, and displayed a protonated molecule at m/z 370.180, which gave a match for C_{21}H_{19}O_{7} with the mass error of 2.7 ppm. The main fragments of metabolite VIII contained 239, 227, 210 and 144 ions. The major fragment of parent compound 269 was not found in the fragments of this metabolite, indicating that metabolite VIII was tri-demethylated GNF-351 and the tri-demethylation reaction occurred in the lower fraction of GNF-351, as shown in Figure 3. The GNF-351 glucuronides (G1 and G2) were eluted at 4.86 and 4.12 min, respectively, and yielded a match for C_{30}H_{33}N_{7}O_{6} with the mass error of 3.2 and −7.0 for G1 and G2. The ion 412 was 176 Da lower than the molecular ion 588, indicating the loss of one molecule of glucuronic acid, further demonstrating the formation of glucuronides. All the detected phase I metabolites were found in the products of the HIM and MIM incubations.

**In vitro screening of phase I and phase II enzymes involved in the metabolism of GNF-351**

Among the tested phase I enzymes, CYP1A1/2 and CYP3A4/5 were identified as the major enzymes involved in the formation of metabolite VI (Figure 4). For the formation of metabolite VII, CYP1A1, CYP1A2, CYP2D6, CYP3A4 and CYP3A5 exhibited higher catalytic activities than the other tested phase I enzymes. CYP1A1 and CYP1A2 played a major role in the formation of metabolite VIII. UGT1A4 was demonstrated to be the major UGT form involved in the glucuronidation of GNF-351. UGT1A3 also exerted catalytic activity towards the formation of glucuronide-1.

**Figure 4**

*In vitro recombinant enzymes screening of phase I DMEs and UGTs involved in the formation of metabolites VI, VII, VIII, G1 and G2. For screening of phase I DMEs, the incubation system contained 50 mM Tris-HCl buffer solution (pH = 7.4), 2 pmol CYP isoforms or 5 mg FMO isoforms, 2 mM MgCl_{2}, 100 μM noscapine and 1 mM freshly prepared NADPH. For identification of UGT isoforms, the incubation system (200 μL) contains 50 mM Tris-HCl buffer solution (pH = 7.4), 5 μg of each UGT form, 2 mM MgCl_{2}, 100 μM noscapine, 1 mM freshly prepared UDPGA. The relative abundance (peak area ratio of metabolites/internal standard) is given.*
Low absorption of GNF-351 and metabolite profiles of GNF-351 in faeces

Serum was taken at 0, 0.5, 2.0, 4.0 and 6 h after oral gavage of 5 mg·kg\(^{-1}\) GNF-351. GNF-351 was not detected at any of these time points, indicating poor absorption (Figure 5). GNF-351 was also not found in 24 h urine, further indicating poor absorption of GNF-351. Given that GNF-351 can only be detected in faeces (24 h), a metabolomics study was

Figure 5
Detection of GNF-351 in serum (0–6 h), urine (24 h) and faeces (24 h). The peak of GNF-351 should appear at a retention time of 5.38 min, and the corresponding molecular ion ([M + H]\(^+\) = 412.225) was employed to extract the peak.

Figure 6
Identification of metabolites in 24 h faeces from mice dosed with 5 mg·kg\(^{-1}\) GNF-351. A. Scores plot of a PCA model from mouse faecal samples without and with GNF-351. B. OPLS loading S-plot of mouse faecal samples; The p(corr)[1] values represent the interclass difference, and the p[1] values represent the relevant abundance of ions. Representative ions (I, VI, VII, F1 and F5) were given.
performed to search for metabolites of GNF-351. Separation was observed between the control group and GNF-351-treated group using the PCA model (Figure 6A). From the OPLS-DA loading plot (Figure 6B), new metabolites (F1–F6), in addition to the metabolites detected using the in vitro incubation system, were found. The trend plot of F1–F6 is given in Supporting Information Fig. S4.

F1 eluted at 4.6 min and exhibited a protonated molecule at 444.214 Da. The matched molecular formula was C24H25N7O2 (mass error = −1.8 ppm), and the MS/MS fragmentation indicated that this metabolite was di-oxidized GNF-351 in which the di-oxidation reaction occurred in the upper fraction of GNF-351 as indicated (Figure 7). Metabolites F2 (Rt = 4.19 min, m/z = 386.174) and F3 (Rt = 3.75 min, m/z =
386.172) gave a match for C₂₁H₁₉N₇O with a mass error of 2.8 and −2.3 ppm respectively. According to the MS/MS fragments and proposed fragmentation mechanism, F₂ and F₃ were likely to be derived from oxidation and tri-demethylation of GNF-351. Metabolite F₄ exhibited a molecular ion at m/z 402.169, with a good match for the molecular formula C₂₁H₁₉N₇O₂. Metabolite F₄ was further identified to be di-oxidized and tri-demethylated GNF-351 based on the proposed MS/MS fragmentation mechanism. Metabolite F₅ eluted at 4.06 min with the protonated molecule to be 426.204, which was 2 Da less than oxidized GNF-351, thus indicating the formation of a dehydrogenated product of oxidized GNF-351 (Figure 7). The protonated molecule of F₆ (m/z = 524.174) was 80 Da higher than metabolite F₁, indicating sulfation of F₁. The fragment of F₆ 256 was 80 Da higher than the fragment of F₁ 176, further supporting the occurrence of sulfation in the upper fraction of F₁. The proposed structure of F₆ is given in Figure 7.

In vivo inhibition by GNF-351 of BNF-induced AHR activation

According to an earlier report (Smith et al., 2011), GNF-351 exhibited the strongest inhibition potential after a 12 h treatment, and did not show inhibition after 24 h. Thus, the treatment time in our experiments was chosen to be 12 h. BNF (5 mg·kg⁻¹) did not induce the expression of Cyp1a1 mRNA in duodenum (A), jejunum (B), liver (C), ileum (D), and colon (E). GNF-351 (5 mg·kg⁻¹, p.o.) was given 5 min before the administration of BNF (5 mg·kg⁻¹, p.o.). The data are means ± SEM (n = 7 for each group). **P < 0.01; ***P < 0.001; N.S., not significant. 18S RNA was used as internal standard.

Discussion

There is a drastic increase in the cost for R&D of a new drug to advance to the clinic, and non-optimal drug ADME (absorption, distribution, metabolism and excretion)
properties are important reasons for the high attrition rate of promising candidates. During the early evaluation of drugs, preclinical studies can result in the possible failure of candidates with poor ADME properties (Pellegatti, 2012). In the present study, metabolomics was used to elucidate the metabolic pathway with use of in vitro phase I and phase II microsome incubation systems. Several phase I metabolites were identified in human and mouse microsomes, including two oxidized GNF-351 (VI and VII) and one tri-demethylated GNF-351 (VIII). For the glucuronidation of GNF-351, there was a species-related difference between results from human and mouse microsomes. From the MS/MS fragmentation patterns, detailed structural information could not be obtained for the glucuronides (G1 and G2). Given that the structure of GNF-351 contains many nitrogen atoms, the glucuronidation sites might occur in these nitrogen atoms as previously reported (Kaivosaari et al., 2011). UGT1A4 was the major UGT form involved in glucuronidation of GNF-351, further supporting the occurrence of the N-glucuronidation reaction. N-glucuronidation of drugs has often been reported to exhibit species difference. Therefore, species scaling from animals to humans for the glucuronidation of GNF-351 should be carefully considered.

Drugs must fulfil some important properties to achieve adequate delivery after oral administration, and intestinal absorption is a key determinant. The present study revealed that GNF-351 was not detected in serum for up to 6 h, indicating that orally administered GNF-351 was not effectively absorbed from the gut either due to poor desolution of the compound, low rates of uptake (transport) and/or high efflux

Figure 9
Summary of the present study. Although GNF-351 is an antagonist in vitro, its poor absorption and metabolic profile limits its in vivo effects. Red colour indicates the CYPs that are significantly affected by AHR activation.
from the gut. Given that the GNF-351 was found in faeces but not in urine at 24 h, the metabolomics analysis was performed on faeces. Besides the phase I metabolites detected in the in vitro system, more phase I metabolites were detected, including di-oxidized GNF-351 (F1), oxidized and trimethylated GNF-351(F2 and F3), di-oxidized and trimethylated GNF-351(F4), and a dehydrogenated product of oxidized GNF-351 (F5). Additionally, a new phase II metabolite, a sulfation product of dioxidized GNF-351 (F6) was detected.

Phase I DMEs involved in the metabolism of GNF-351 were identified to decipher the complete metabolic pathway. Among the tested phase I enzymes, CYP1A1, CYP1A2, CYP3A4 and CYP3A5 were demonstrated to be the main enzymes involved in GNF-351 metabolism. Notably, CYP1A1 and CYP1A2 can be significantly induced via AHR activation (Kohle and Bock, 2007), which accelerates the metabolism of GNF-351, further weakening the AHR antagonist activity of GNF-351. Additionally, CYP3A4 and CYP3A5 are involved in the metabolism of many clinical drugs, and the activity can be easily influenced by compounds, such as the inhibitor of CYP3A4 ketoconazole (Fuchs et al., 2013) and the inducers of CYP3A4 and CYP3A5, rifampicin and rifaximin (Ma et al., 2007). Therefore, the risk of drug–drug interactions should be considered in the R&D process of GNF-351. Individual differences in these enzymes also could result in different rates of metabolism of GNF-351. For example, women might exhibit higher metabolic activity than men towards GNF-351 because of higher CYP3A4 activity (Scandlyn et al., 2008). All these factors would complicate the metabolic response of different individuals towards GNF-351 in different exposure environments. It should be also noted that CYP3A4 is highly expressed in human intestines (Granvil et al., 2003), and the present enzyme phenotyping results obtained from recombinant human enzyme sources indicated even more metabolism in human intestines, and thus less antagonistic activity in humans than in mice could be predicted.

As is expected, the in vivo efficiency of GNF-351 as an antagonist was limited by poor absorption and complex metabolic factors. Earlier studies have shown that GNF-351 inhibited AHR activation in murine hepatoma-derived reporter cells H1L1.1c2 with an IC50 value of 116 nM. For a humanized PXR transgenic mouse weighing 25 g, its blood volume is 2 mL (8% of body weight). If the oral bioavailability of GNF-351 was 1%, the plasma concentration of GNF-351 was calculated to reach approximately 1.5 μM after oral gavage of 5 mg·kg⁻¹ GNF-351. The in vivo results indicated that this dose of GNF-351 gave orally did not exhibit antagonism of BNF-induced AHR activation in liver, indicating that low absorption and/or rapid metabolism GNF-351 had prevented an adequate concentration of GNF-351 in the liver to bind and inhibit AHR. The antagonism by GNF-351 of intestinal AHR was also investigated. BNF did not significantly activate AHR in the duodenum and jejunum, but exhibited significant activation of AHR in the ileum. While oral GNF-351 did not show measurable inhibition towards BNF-induced AHR activation in liver, it did inhibit BNF-induced AHR activation in ileum and colon.

In conclusion, the absorption properties and detailed metabolic pathway of GNF-351 were evaluated in the present study (Figure 9). The results suggested that structural modifications to GNF-351 are required in order to improve its oral absorption and metabolic behaviour. For example, a pro-drug structure, linking GNF-351 to bile acids, might increase oral absorption significantly (Rais et al., 2010). Taken together, the information in the present study will be beneficial for further development of GNF-351 as a structure-based antagonist of the AHR in liver.

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Conflicts of Interest

None.

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Supporting information

Additional Supporting Information may be found in the online version of this article at the publisher’s web-site:

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Figure S1 Trend plot of the ions I-VIII. MLM incubated with (+S) and without (−S) GNF-351.

Figure S2 PCA scores scatter plot for HLMs incubation system (dot), MLMs incubation system (triangle), the incubation without NADPH (diamond), incubation without microsomes (circle), and incubation without GNF-351 (inverted triangle).

Figure S3 Comparison of GNF-351 glucuronidation between HLMs and MLMs.

Figure S4 Trend plot of the ions F1–F6.