A NOVEL, SPECIES-SPECIFIC CLASS OF UNCOMPETITIVE INHIBITORS OF γ-GLUTAMYL TRANSPEPTIDASE
Jarrod B. King1, Matthew B. West1, Paul F. Cook2 and Marie H. Hanigan1
From the Department of Cell Biology1, University of Oklahoma Health Sciences Center, Oklahoma City, OK, and the Department of Chemistry and Biochemistry2, University of Oklahoma, Norman, OK
Running Title: Uncompetitive Inhibitor of GGT
Address correspondence to: Marie H. Hanigan, Ph.D., Biomedical Research Center, Room 264, 975 N.E.10th Street, Oklahoma City, OKC 73104, Phone:405-271-3832; FAX:405-271-3813; e-mail: marie-hanigan@ouhsc.edu

Expression of γ-glutamyl transpeptidase (GGT) in tumors contributes to resistance to radiation and chemotherapy. GGT is inhibited by glutamine analogues that compete with the substrate for the γ-glutamyl binding site. However, the glutamine analogues that have been evaluated in clinical trials are too toxic for use in humans. We have used high-throughput screening to evaluate small molecules for their ability to inhibit GGT and have identified a novel class of inhibitors that are not glutamine analogues. These compounds are uncompetitive inhibitors, binding the γ-glutamyl enzyme complex. OU749, the lead compound, has an intrinsic Ki of 17.6 μM. It is a competitive inhibitor of the acceptor glycyl-glycine, which indicates that OU749 occupies the acceptor site while binding to the γ-glutamyl substrate complex. OU749 is more than 150-fold less toxic than the GGT inhibitor acivicin toward dividing cells. Inhibition of GGT by OU749 is species-specific, inhibiting GGT isolated from human kidney with 7- to 10-fold greater potency than GGT isolated from rat or mouse kidney. OU749 does not inhibit GGT from pig cells. Human GGT expressed in mouse fibroblasts is inhibited by OU749 similarly to GGT from human cells, which indicates that the species specificity is determined by differences in the primary structure of the protein rather than species-specific, post-translational modifications. These studies have identified a novel class of inhibitors of GGT, providing the basis for further development of a new group of therapeutics that inhibit GGT by a mechanism distinct from the toxic glutamine analogues.

The mechanism of inherent and acquired resistance of tumors to many forms of treatment involves glutathione. Elevated glutathione levels in tumors have been shown to contribute to resistance to chemotherapy and radiotherapy and prevent the initiation of the apoptotic cascade in tumor cells (1-5). The enzyme γ-glutamyl transpeptidase (GGT, EC 2.3.2.2), which is localized to the cell surface, cleaves the γ-glutamyl bond of extracellular glutathione, enabling the cell to use extracellular glutathione as a source of cysteine for increased synthesis of intracellular glutathione (6). GGT is induced in many human tumors, enhancing their resistance to chemotherapy (7;8). Inhibiting GGT prior to chemotherapy or radiation would sensitize GGT-positive tumors to treatment. However, all known GGT inhibitors are too toxic for use in humans (9;10).

GGT plays an essential role in releasing cysteine from extracellular glutathione. Most cells are unable to take up intact glutathione (6). In GGT knockout mice, the absence of GGT in the renal proximal tubules results in the excretion of glutathione in the urine (11). In these mice, the glutathione in the glomerular filtrate can not be cleaved into its constituent amino acids for reabsorption. GGT knockout mice have a 4500-fold elevation of glutathione in their urine relative to their GGT-wild-type littermates. GGT knockout mice grow slowly and die by ten weeks of age due to a cysteine deficiency.

Inhibiting GGT for as little as 2 hours lowers the intracellular cysteine concentration in GGT-positive tumors (3). Inhibitors of GGT activity could be used prior to the administration of chemotherapy to limit the supply of cysteine to the tumor, thereby blocking the tumor's ability to maintain high levels of intracellular glutathione.

GGT catalyzes the cleavage of γ-glutamyl compounds and the transfer of the gamma-glutamyl group to an acceptor substrate by a ping-pong kinetic mechanism (12). Glutathione and glutathione-S-conjugates are the most common physiologic substrates of GGT. They serve as the
gamma-glutamyl donor in the initial reaction. In the first reaction the $\gamma$-glutamyl bond of the initial substrate is cleaved, the $\gamma$-glutamyl group becomes covalently bound to the enzyme and the remainder of the substrate is released as the first product. With glutathione as the substrate, cysteinyl-glycine is released and is subsequently cleaved into cysteine and glycine by cell surface dipeptidases. In the second reaction of GGT transpeptidation, the $\gamma$-glutamyl-group is transferred from the $\gamma$-glutamyl-GGT complex to the second substrate (the acceptor). Dipeptides and amino acids have the highest $K_m$ as acceptors. The second substrate with the covalently bound gamma-glutamyl group is released as the second product from the enzyme.

Compounds that inhibit GGT include the glutamine analogues acivicin, 6-diazo-5-oxo-L-norleucine, and azaserine (Fig. 1) (13). Rational design of GGT inhibitors based on studies of the active site has led to the identification of additional $\gamma$-glutamyl analogues. Lherbet and Keillor have designed sulfur derivatives of L-glutamic acid which inhibit GGT (14;15). Han and coworkers have synthesized and tested a series of $\gamma$-(monophenyl)phosphono glutamate analogues which also functioned as inhibitors of GGT (16;17).

Evaluation of several of the glutamine analogues that inhibit GGT has shown them to be toxic (9;10). Acivicin, the most potent inhibitor of GGT that has been tested clinically, is a neurotoxin (18). The neurotoxicity of the glutamine analogues may be due to interference with glutamine in recycling the neurotransmitter glutamate via the glutamate-glutamine cycle. Glutamine is also involved in the synthesis of several nucleotides and complex polysaccharides. Inhibition of these essential synthetic pathways can be toxic to dividing cells. Acivicin, 6-diazo-5-oxo-L-norleucine, and azaserine all cause bone marrow suppression (9). There is no known GGT inhibitor that can be used clinically.

Rather than design glutamine analogues, we have used physiologic conditions to screen libraries of small molecules to identify new inhibitors of GGT. This effort has lead to the identification of a novel class of uncompetitive inhibitors of GGT that are structurally distinct from and less toxic than the glutamine analogues. This new class of compounds occupies the acceptor site, not the $\gamma$-glutamyl site. The lead compound is species-specific, inhibiting human GGT, but with weak to no inhibitory activity towards GGT from mice, rats and pigs.

**Experimental Procedures**

*High throughput screening -* A high throughput method was developed to screen for inhibitors of GGT. The assays were conducted in 96-well plates. The final volume in each well was 100 $\mu$L. The assay buffer contained: 100 mM Na$_2$HPO$_4$ pH 7.4, with 3.2 mM KCl, 1.8 mM KH$_2$PO$_4$, and 27.5 mM NaCl. Each reaction contained 1 mM L-glutamic acid $\gamma$-4-nitroanilide HCl (GpNA) and 40 mM glycyl-glycine (Gly-Gly) (19). GGT is expressed on the cell surface and the assay was initiated by addition of $10^4$ 786-O cells (ATCC CRL-1932, a GGT-positive human renal cell adenocarcinoma line). Formation of the product, p-nitroaniline at 37°C, was monitored continuously at OD$_{405}$ by a Bio-Rad Model 680 Microplate Reader with Microplate Manager 5.2 software (Bio-Rad). One unit of GGT activity was defined as the amount of GGT which released one $\mu$ mole of p-nitroaniline/min at 37°C.

*Screening chemical library:* A 10,000 compound chemical library from the DIVERSet collection (ChemBridge Corporation, San Diego, CA) was screened for inhibitors of GGT. The library was formatted in 96-well plates with 0.25 $\mu$moles test compound per well. Stock solutions were prepared by the addition of 25 $\mu$L dimethyl sulfoxide (DMSO) per well. For each test compound, 5 $\mu$L was added to the assay mixture, resulting in a final concentration of 500 $\mu$M. Glutathione is a competitive inhibitor of the GpNA substrate, and wells containing 400 $\mu$L oxidized glutathione were included on each 96-well plate as a positive control. Compounds which inhibited GGT activity to the same extent or greater than glutathione were scored as positive hits. The positive hits were retested at several concentrations for inhibition of GGT activity. For further testing, individual compounds and their structural analogues were purchased from ChemBridge, Specs (Delft, Netherlands), or Otava Chemicals (Kiev, Ukraine).

*Enzyme isolation -* GGT was isolated from human kidney (National Disease Research Interchange, Philadelphia, PA), male Sprague-Dawley rat kidney and female Balb/c mouse kidney. Tissue
was homogenized in 4 volumes of 25 mM Tris pH 7.5, containing 0.33 M sucrose, 0.2 mM EDTA, 1 μM leupeptin and 1.4 μg/ml aprotinin. A 9000g supernatant was spun at 100,000g for 1 h. The microsomal pellet was homogenized in 25 mM Tris pH 7.35, 0.5% Triton X-100, 1 μM leupeptin, 1.4 μg/ml aprotinin then centrifuged again at 100,000g for 1 h. The supernatant was assayed for GGT activity, aliquoted, and stored at -80°C until further use. All solutions were maintained at 4°C throughout the isolation. The specific activities of GGT were 3.4, 7.4, and 1.5 units/mg protein for human, rat, and mouse preparations, respectively. Prior to use in the GGT assay, the enzyme was diluted in phosphate-buffered saline (PBS) containing 0.025% Triton X-100 and 0.19 mUnits of enzyme were used per assay unless otherwise indicated.

Cloning of human GGT and expression in Pichia pastoris - The soluble domain of human GGT (amino acids 28-569) was amplified by PCR. The forward primer introduced an EcoRI restriction site and a TEV protease-cleavable N-terminal His6 tag, while the reverse primer introduced a NotI restriction site after the stop codon of genomic GGT. The PCR product was digested with EcoRI and Not I and inserted into the corresponding sites of pPICZαA (Invitrogen), generating plasmid pMW-102. This construct was amplified in E. coli DH5α cells. The fidelity of the recombinant ORF was verified by sequencing (Oklahoma Medical Research Foundation’s DNA Sequencing Core Facility, Oklahoma City, OK). The SacI-linearized plasmid, pMW-102, was purified and transformed into wild-type Pichia pastoris strain X-33, selected and induced as recommended by Invitrogen. Secreted, recombinant, His6-tagged, human GGT was collected from the media with a Ni-NTA column. The hexahistidine tag was cleaved with His6-tagged TEV protease. TEV protease and the cleaved histidine tag were separated from human GGT by collecting the flow-through from a second Ni-NTA column purification. The specific activity of the purified enzyme was 397.8 units /mg. Prior to use in the GGT assay, the enzyme was diluted in PBS and 2.8 mUnits of enzyme were used per assay.

Kinetic Studies - The assay buffer contained: 100 mM Na2HPO4 pH 7.4, with 3.2 mM KCl, 1.8 mM KH2PO4, and 27.5 mM NaCl. The concentrations of the substrate GpNA and the acceptor Gly-Gly were varied as indicated. Purified GGT was added to initiate the assay. The reaction was monitored as described for the high throughput screening. Cytotoxicity assays - 786-O (ATCC CRL-1932), a human renal cell adenocarcinoma line, was plated in DMEM supplemented with 5% FBS and penicillin/streptomycin (50 U/mL, 50 μg/mL) at 10⁶ cells per well in 96-well plates. The next day the medium was changed to fresh DMEM containing FBS, antibiotics and the test compound. Equivalent concentrations of DMSO were added to control wells. Cell viability was determined by the MTT assay 72 hr after the addition of the test compound (20).

Cell lines – LLC-PK1 (ATCC CRL 1392), a pig kidney cell line, NRK-52E (ATCC CRL-1571), a normal rat kidney cell line, LLC-MK2 (ATCC CCL-7), a normal kidney monkey cell line, and 786-O cells were grown in complete Dulbecco’s Modified Eagle’s Medium (DMEM) supplemented with 5% FBS (HyClone, Logan, UT), and penicillin/streptomycin (50 U/mL, 50 μg/mL) (Invitrogen, Carlsbad, CA). NIH/3T3 cells transfected with cDNA encoding human GGT were described previously and were grown in DMEM/F12 containing 200 μg/mL of G418 (6). HK-2 (ATCC CRL- 2190), a human cell line derived from immortalized normal proximal tubule cells, were grown in keratinocyte serum-free media (K-SFM) from ATCC (Manassas, VA). The cells were trypsinized off the plates and suspended in PBS for the GGT assay.

Glutathione degradation – Incubation mixture contained 1 mM glutathione, 40 mM glycylglycine, and inhibitor in the GGT assay buffer. Reactions were initiated with the addition of 1.9 mU of GGT and incubated at 37°C. The final volume of the reaction was 100 μL. Aliquots of the reaction mixture were removed at the specified time points, immediately acidified with an equal volume of cold 4.31% 5-sulfosalicylic acid, and maintained at 4°C until glutathione concentration was determined by the method of Tietze (21).

Data analysis - Initial velocity data were first analyzed graphically using double-reciprocal plots of initial velocities versus substrate concentration and suitable secondary plots. Data were then fitted using the appropriate equation and the Marquardt-Levenberg algorithm supplied with the EnzFitter program from BIOSOFT, Cambridge, U.K. Kinetic parameters and their corresponding
standard errors were estimated using a simple weighting method.

Data for competitive and uncompetitive inhibition were fitted using eqs 1 and 2.

\[ v = \frac{VA}{K_a(1 + K_i) + A} \]  
\[ v = \frac{VA}{K_s + A(1 + K_{ii})} \]

In eqs. 1 and 2, \( v \) and \( V \) are the measured initial rate and maximum rate, respectively, \( K_a \) is the Michaelis constant for the varied substrate, and \( K_s \) and \( K_{ii} \) are slope and intercept inhibition constants.

The \( LD_{50} \) and 95% confidence intervals of the test compounds in cell lines were calculated with a Prism log (inhibitor) vs normalized response (variable slope) curve fit (Prism, GraphPad Software Inc., San Diego, CA). A two-tailed t-test was used to detect significant differences between OU749 inhibition of human GGT and inhibition of GGT from other species.

RESULTS

Screening chemical library for inhibitors of GGT. A 10,000 compound chemical diversity library from ChemBridge Corporation was screened for inhibitors of GGT activity. Compounds in the DIVERSet library were chosen for maximum diversity of chemical structure. All compounds satisfied several physiochemical properties important for drug discovery: \( MW \leq 500, \text{clogP} \leq 5, tPSA \leq 100, \text{rotatable bonds} \leq 8, \text{hydrogen bond acceptors} \leq 10, \text{and hydrogen bond donors} \leq 5 \) (22).

The initial screen identified 16 hits. When retested, 12 of the compounds were only weak inhibitors and not pursued further. Three of the remaining compounds were derivatives of isoindole-1,3-dione. Additional evaluation of this group of compounds revealed low solubility in aqueous solution and severe toxicity in a cell based assay. These compounds were not pursued further. The last of the initial hits was a novel inhibitor of GGT. The inhibitor, N-[5-(4-methoxybenzyl)-1,3,4-thiadiazol-2-yl]benzenesulphonamide, was designated OU749.

The structure of OU749 reveals that it is not a glutamine analogue (Fig. 2). It inhibited GGT isolated from human kidney in a dose-dependent manner (Fig. 2).

Kinetic analysis of inhibition of GGT by OU749. To determine the mechanism of OU749 inhibition of human kidney GGT, inhibition patterns were obtained varying each of the substrates with the second maintained at a fixed concentration. With GpNA (the first substrate) varied, glygly was maintained at 40 mM (\( K_{glygly} \) is \( 11.4 \pm 1.2 \) mM), while GpNA was maintained at 3 mM (\( K_{GpNA} \) is \( 1.07 \pm 0.04 \) mM) when glygly was varied. Inhibition by OU749 was uncompetitive with respect to GpNA, indicating that it binds to the covalent E-\( \gamma \)-glutamyl complex, the F form of the enzyme (Fig. 3A). The \( K_i \) value obtained was \( 73.8 \pm 2.5 \) \( \mu \)M. Inhibition by OU749 was competitive with respect to glygly indicating that OU749 was occupying the acceptor site (Fig. 3B). A \( K_i \) of \( 17.6 \pm 3.8 \) \( \mu \)M was obtained. The \( K_i \) of 17.6 \( \mu \)M is the intrinsic value (23). The value obtained varying GpNA at a fixed glygly concentration must be the same once corrected for the concentration of fixed reactant. Given, \( appK_i = K_i(1 + [\text{glygly}]/K_{glygly}) \), the calculated value of \( K_i \) is 16.5 \( \mu \)M.

A similar analysis of OU749 inhibition, carried out with a highly purified preparation of human GGT expressed in yeast yielded similar results (Fig. 3C, D). The \( appK_i \) of OU749 as a competitive inhibitor of glygly was 25 \( \pm 2 \) \( \mu \)M, while that obtained varying GpNA was 58.2 \( \pm 1.6 \) \( \mu \)M. A fit of the data displayed in Fig. 3A-D, to the equation for noncompetitive inhibition (\( v = VA/[K_a(1 + I/K_i) + A(1 + I/K_{ii})] \)) does give a finite, but very high \( K_i \) (>200 \( \mu \)M) binding to the free form of the enzyme (E form). This is not surprising since the site for glygly, and thus OU749 must be present in the E form. However, the much weaker binding, > 10-fold, is consistent with synergy of binding of OU749 in the presence of the \( \gamma \)-glutamyl moiety.

Structure activity studies. A series of structural analogues of OU749 were evaluated as inhibitors of GGT. The \( appK_i \) for each compound, obtained by varying GpNA, was determined experimentally with GGT isolated from human kidney. Compounds that had no inhibitory activity aided in defining elements that were essential for inhibition (Fig. 4). The data revealed that inhibition of GGT activity required the benzene ring attached to the
sulfonamide group (Fig. 4, compound 1). Addition of a spacer carbon atom between the sulfur and the benzene ring eliminated inhibitory activity (Fig. 4, compound 2). Removing the carbon spacer atom between the thiadiazole ring and the end ring (Fig 4, compound 3) or substitution of the thiadiazole ring with more complex structures eliminated inhibitory activity (Fig 4, compounds 4 and 5). A core structure containing all the essential elements is shown in Fig 5. The $K_i$ for the unsubstituted core structure is $99.7 \pm 3.1 \mu M$. Inhibition is enhanced by the substitution of either the X or the Y position with chlorine (Fig 5, compound 2, 4). Substitution of both the X and the Y position with chlorine provides the most potent inhibitor with an alpha $K_i$ of $28.7 \pm 1.0 \mu M$ (Fig 5, compound 1). Addition of a methoxy group at the Y position of the core compound yields our lead compound OU749 which is the third most potent inhibitor of the analogues tested (Fig 5 compound 3). Substitution of the X position with either a methoxy compound or a nitroso group weakened the inhibition (Fig 5, compounds 5, 7-9). Addition of an acetamide group at the X position eliminates inhibition (Fig 5, compound 10).

Toxicity of OU749. The glutamine analogues that inhibit GGT activity are toxic to dividing cells. We evaluated the toxicity of OU749 and several of its structural analogues towards cells in log growth using 786-O cells, a human renal tumor cell line. Cells were grown in the presence of the test compounds for 3 days. As shown in Table 1, the glutamine analogue, acivicin, had an LD$_{50}$ of $0.81 \mu M$. All four of the compounds tested (Fig. 5, compounds #1, 2, 3 and 6) were at least an order of magnitude less toxic than acivicin. OU749 (compound #3) was more than 150-fold less toxic than acivicin. These data emphasize the reduced toxicity of GGT inhibitors that are not glutamine analogues.

Although our structure activity analysis revealed two compounds that were more potent inhibitors of GGT than OU749 (Fig. 5, compounds 1 and 2), these compounds were 17- to 20-fold more toxic than OU749 making them less promising as candidates for further development for clinical use. Therefore OU749 continued as our lead compound in further characterization studies of inhibition by this class of compounds.

Species specificity of GGT inhibition by OU749. OU749 inhibits GGT isolated from human kidney in a dose-dependent manner. However, OU749 is 7-fold less potent as an inhibitor of GGT isolated from rat kidney and 10-fold less potent inhibiting GGT from mouse kidney (6A). The species specificity of GGT inhibition by OU749 was further evaluated with cells lines from five different species (Fig 6B). OU749 showed dose-dependent inhibition of GGT in the two human kidney cell lines, 786-O a renal tumor cell line and HK-2, an immortalized renal proximal tubule cell line. Inhibition of GGT in the human cell lines was of similar potency to the inhibition of GGT isolated from human kidney. GGT in the rat kidney cell line NRK-52E was only weakly inhibited by OU749, equivalent to the weak inhibition of GGT isolated from rat kidney. GGT in the pig kidney cell line LLC-PK1 was not inhibited by OU749. GGT in the monkey kidney cell line LLC-MK2 was only weakly inhibited by OU749 similar to the inhibition of GGT in rat kidney cells.

GGT is heavily glycosylated (24). To determine whether the sensitivity of GGT to inhibition by OU749 was determined by the primary structure or by post-translational modifications such as glycosylation, we tested the sensitivity of human GGT expressed in mouse NIH/3T3 fibroblasts to inhibition by OU749 (Fig. 6B). The data revealed that human GGT expressed in mouse cells was inhibited by OU749 to the same extent as human GGT expressed in other human cells. Therefore the sensitivity of GGT to inhibition by OU749 was determined by the peptide sequence rather than species-specific post-translational modifications. Data in Figure 2 further confirm that the primary structure rather than post-translational modifications determine the degree of inhibition of GGT by OU749. GGT isolated from human kidney (Fig 2 A,B) is inhibited by OU749 to the same extent as human GGT expressed in yeast (Fig 2, C,D).

OU749 inhibits the cleavage of glutathione by GGT. The standard assay for GGT activity monitors the release of p-nitroaniline from the first substrate, γ-glutamyl-p-nitroanilide. To confirm that OU749 inhibits the cleavage of glutathione, the primary physiologic substrate of GGT, we monitored the breakdown of glutathione by human, kidney GGT in the presence of the inhibitor. OU749 inhibited the cleavage of glutathione in a dose-dependent manner (Fig 7A).
OU749 also inhibited the cleavage of oxidized glutathione (GSSG) by human GGT (data not shown). To determine whether the species specificity of inhibition of GGT by OU749 was also relevant to the physiologic substrate, OU749 was evaluated for its ability to inhibit the degradation of glutathione by rat GGT. Rat GGT cleaved glutathione in a time dependent fashion as had been observed for human GGT (Fig. 7B). OU749 was unable to inhibit degradation of glutathione by GGT from rat kidney (Fig. 7B). These data corroborate the data regarding the species specificity of OU749 obtained using the synthetic substrate, GpNA.

**DISCUSSION**

We have identified a novel class of GGT inhibitors that are not glutamine analogues. Kinetic studies of the lead compound OU749 revealed that the mechanism of inhibition was uncompetitive relative to the $\gamma$-glutamyl substrate, indicating that the inhibitor bound the enzyme-substrate complex. In contrast, to competitive inhibitors, which lose potency as substrate concentration builds, uncompetitive inhibitors become more potent as the substrate concentration rises in an inhibited open system. Data from the GGT knockout mice show that in the absence of GGT activity, glutathione levels decrease in tissues but the concentration of glutathione in the serum rises more than 6-fold, likely due to the inability of cells to cleave glutathione and recover the amino acids (11). In addition, the glutathione concentration in the urine increases more than 2400-fold as the glutathione transits the proximal tubules intact in the absence of GGT. Westley and Westley have argued that uncompetitive inhibitors are superior to competitive inhibitors for instituting change in open systems such as those found in vivo where substrate concentrations rise with enzyme inhibition (25). Many uncompetitive inhibitors function by locking the ES complex in a state after initial product release but before conversion of the enzyme back to the native form. In GGT, the enzyme-substrate complex consists of the glutamyl group covalently bound to the enzyme following release of the remainder of the $\gamma$-glutamyl substrate. GGT from *E. coli* has been crystallized (26). Analysis of the crystal structure of *E. coli* GGT and the gamma-glutamyl-enzyme intermediate revealed that the N-terminal threonine of the small subunit is the catalytic nucleophile in the enzymatic reaction (27). This threonine is conserved in human, rat, mouse and pig. Site directed mutagenesis of human GGT has identified four amino acids that are essential to GGT activity (one in the large subunit, Arg 107, and three in the small subunit, Asp-423, Ser-451 and Ser-452) (28-30). All four of these amino acids are identical in human, rat, mouse and pig.

Kinetic studies further revealed that while binding the enzyme-$\gamma$-glutamyl complex, OU749 occupies the acceptor site. The acceptor site is not well defined. To date, the crystal structures of GGT isolated from only *E. coli* and *H. Pylori* have been published (27,31,32). No information on mutational analysis that alters the acceptor site has been published. Our finding that inhibition by OU749 is species specific with high affinity for human, lower affinity for monkey, rat and mouse and no inhibition of pig GGT may aid in the delineation of the acceptor site. The sequence of GGT from human, rat, mouse, and pig have been published. Alignment of these sequences is included in the supplementary data. However, no pattern is apparent in the sequence alignment that would provide insight into amino acids or regions of the polypeptide sequence that are critical to inhibitor binding. Alignment of the GGT peptide sequences does reveal that one of the seven potential N-glycosylations sites in human GGT (Asn 266) is not present in rat, mouse or pig. Modeling the mammalian sequences based on the crystal structure of the bacterial enzyme does not highlight any of the amino acids as important in inhibitor binding. There are other reports of species specific inhibitors of GGT, although the effect of sulfhydral reagents on GGT activity appears to have some species specificity (33).

Structural alterations of OU749 increased the inhibitory activity of the compound but not without accompanying increases in toxicity. The in vitro toxicity profile of OU749 is favorable. In a dividing cell model, OU749 is 150-fold less toxic than acivicin which was abandoned after phase I clinical trials due to toxicity.

In addition to its role in cancer therapy, GGT also plays a critical role in drug metabolism due to the ubiquitous presence of glutathione. GGT is an essential enzyme in the formation of mercapturic acids in the kidney and initiates the activation of
halogenated alkenes and other drugs to potent kidney toxins through this pathway (34;35). Cisplatin has been shown to be bioactivated to a renal toxin through the mercapturic acid pathway (36;37). Inhibition of GGT during cisplatin-based chemotherapy would not only sensitize the tumors to the therapy, it would also block the kidney toxicity of cisplatin. Additional clinical conditions for which a GGT inhibitor may have therapeutic benefit include cardiovascular disease and asthma as nitric oxide is transported in the blood as a glutathione conjugate and requires GGT activity for its release (38;39). Finally, GGT is one of two enzymes that metabolize leukotriene C4 to leukotriene D4, a mediator of inflammation common to many diseases (40).

Previous clinical studies have attempted to overcome drug resistance in tumors by inhibiting glutathione synthesis with buthionine sulfoximine (41), an inhibitor of the rate-limiting enzyme in the synthesis of glutathione. However, there was no depletion of cysteine in the body with this protocol. Cysteine’s sulfur, the active nucleophilic group of the glutathione molecule, binds and inactivates reactive, electrophilic compounds. Free cysteine will also react with, and inactivate chemotherapy drugs (42). Inhibition of GGT reduces both intracellular glutathione and depletes cysteine levels, increasing the sensitivity of the tumor to the drug. Studies in mice have shown that inhibiting GGT for as little as 2hr selectively lowers the intracellular cysteine concentration in GGT-positive tumors (3). Mena and colleagues used acivicin to deplete tumor GSH in combination with aggressive therapy and achieved complete cure of metastatic melanoma to the liver in 90% of test animals (4). Development of less toxic GGT inhibitors, such as OU749, hold great promise for enhanced cancer therapy.

REFERENCES


**FOOTNOTES**

We gratefully acknowledge the assistance of Terri McHugh, Department of Cell Biology, OUHSC for assistance with the enzyme assays, and Kostyantyn Bobyk in the Department of Chemistry and Biochemistry, University of Oklahoma for assistance with enzyme kinetics calculations and preparing the graphs for Figure 3. This work was supported by grants HR03-044 from the Oklahoma Center for the Advancement of Science and Technology and RO1CA57530 from the National Cancer Institute, NIH to M.H.H. and the Grayce B. Kerr endowment to the University of Oklahoma to support the research of P.F.C.
FIGURE LEGENDS

Fig. 1. Glutathione and glutamine analogues that inhibit GGT. Shown are the structures of glutathione (1) and the GGT inhibitors azaserine (2) and acivicin (3).

Fig. 2. Inhibition of GGT by OU749. Structure of OU749 (left panel) and substrate velocity curves (right panel) for the inhibition of human kidney GGT by 0 (■), 15.2 μM (▲), 31.3 μM (▼), 62.5 μM (◇), 125 μM (○) OU749 in the presence of 40 mM glycylglycine. Data shown are the average of triplicate values ± S.D. (for many points the error bars are smaller than the symbol).

Fig. 3. Kinetic Analysis of GGT inhibition by OU749. Double-reciprocal plots of the initial velocities of human kidney GGT (A, B) or human GGT transfected into and isolated from Pichia pastoris (C, D) in the presence of 40 mM glycylglycine (A, C) or 3 mM GpNA (B, D) with 0 (■), 15.2 μM (▲), 31.3 μM (△), 62.5 μM (◇), 125 μM (○) OU749. Data shown are average of triplicate values ± S.D. (for many points the error bars are smaller than the symbol).

Fig. 4. Structural analogues of OU749 that do not inhibit GGT activity.

Fig. 5. Inhibition of GGT by structural analogues of OU749.

Fig. 6. Species-specific inhibition of GGT by OU749. A. Inhibition of GGT from human kidney (●), rat kidney (△) and mouse kidney (□) by OU749. Inhibition of human GGT was significantly more potent than inhibition of rat or mouse GGT (p = 0.03). B. Inhibition of GGT in 786-O human renal adenocarcinoma cells (●), HK-2 normal human kidney derived cells (▲), and NIH/3T3 mouse fibroblast transfected with human GGT cDNA (●), LLC-MK2 monkey kidney cells (○), NRK-52E rat kidney cells (△) and LLC-PK1 pig kidney (▼) cells. There was no significant difference in inhibition among the cell lines expressing human GGT, but OU749 inhibition of GGT in human cell lines was significantly more potent than inhibition of GGT in monkey, rat or pig cell lines (p ≤ 0.04).

Fig. 7. Species-Specific Inhibition of glutathione degradation by OU749. Cleavage of glutathione by human GGT (A) or rat GGT (B) in the presence of 0 (■), 62.5 μM (◇), 125 μM (○), 250 μM (★) or 500 μM (□) OU749. Reactions contained 1.9 mU GGT, 40 mM glycylglycine and 1 mM glutathione. OU749 inhibited glutathione breakdown by human GGT but did not inhibit glutathione breakdown by rat GGT.
Table 1. Toxicity of GGT Inhibitors towards Dividing 786-O Cells

<table>
<thead>
<tr>
<th>Compound</th>
<th>LD_{50} (μM)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Acivicin</td>
<td>0.81</td>
</tr>
<tr>
<td>Fig 5 Compound #1</td>
<td>7.6</td>
</tr>
<tr>
<td>Fig 5 Compound #2</td>
<td>6.5</td>
</tr>
<tr>
<td>Fig 5 Cmpd #3 (OU749)</td>
<td>128</td>
</tr>
<tr>
<td>Fig 5 Compound #6</td>
<td>71</td>
</tr>
</tbody>
</table>
Figure 1
Figure 4

(1) (2)

(3)

(4) (5)
### Core Structure

![Core Structure Diagram]

<table>
<thead>
<tr>
<th>Compound #</th>
<th>Substitution</th>
<th>Inhibition AppK (μM)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>X=Cl, Y=Cl</td>
<td>28.7 ± 1.0</td>
</tr>
<tr>
<td>2</td>
<td>Y=Cl</td>
<td>43.3 ± 1.6</td>
</tr>
<tr>
<td>3 (OU749)</td>
<td>Y=OCH₃</td>
<td>73.8 ± 2.5</td>
</tr>
<tr>
<td>4</td>
<td>X=Cl</td>
<td>75.9 ± 1.7</td>
</tr>
<tr>
<td>5</td>
<td>X=NO₂, Y=Cl</td>
<td>74.3 ± 6.9</td>
</tr>
<tr>
<td>6</td>
<td>None</td>
<td>99.7 ± 3.1</td>
</tr>
<tr>
<td>7</td>
<td>X=NO₂, Y=OCH₃</td>
<td>114 ± 8.4</td>
</tr>
<tr>
<td>8</td>
<td>X=OCH₃, Y=OCH₃</td>
<td>226 ± 12.7</td>
</tr>
<tr>
<td>9</td>
<td>X=OCH₃, Y=OCH₃, Z=OCH₃</td>
<td>1,115 ± 100</td>
</tr>
<tr>
<td>10</td>
<td>X=NHOCH₃, Y=OCH₃</td>
<td>No inhibition</td>
</tr>
</tbody>
</table>
Figure 6

A.  

![Graph A](image)

B.  

![Graph B](image)