Identification of ketene reactive intermediate of erlotinib possibly responsible for inactivation of P450 enzymes

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Running Title Page

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Abbreviations
BBA, 4-bromobenzylamine; m-CPBA, m-chloroperbenzoic acid; CYP, cytochrome P450; DTT, dithiothreitol; EGFR, epidermal growth factor receptor; EPI, enhanced product ion; ELT, erlotinib; GSH, glutathione; HLMs, human liver microsomes; HPLC-UV, high performance liquid chromatography-ultraviolet; LC-MS/MS, liquid chromatography-tandem mass spectrometry; MBI, mechanism-based inactivation; MRM, multiple reaction monitoring; MS, mass spectrometry; MS/MS, tandem mass spectrometry; NADPH, β-nicotinamide adenine dinucleotide 2′-phosphate reduced tetrasodium salt; NMR, nuclear magnetic resonance; NSCLC, non-small-cell lung cancer; PBS, potassium phosphate buffer; Pd-BaSO₄, palladium on barium sulfate; Pd-C, palladium on activated charcoal; PI, precursor ion.
Abstract

Erlotinib (ELT), a tyrosine kinase inhibitor, is widely used for the treatment of non-small-cell lung cancer in clinic. Unfortunately, severe drug-induced liver injury and other adverse effects occurred during the treatment. Meanwhile, ELT has been reported to be a mechanism-based inactivator of CYPs 3A4 and 3A5. The objectives of this study were to identify ketene intermediate of ELT and investigate the association of the acetylenic bioactivation with the enzyme inactivation caused by ELT. A ketene intermediate was detected in human microsomal incubations of ELT, using 4-bromobenzylamine as a trapping agent. CYPs 3A4 and 3A5 mainly contributed to the bioactivation of ELT. Microsomal incubation study showed that the ketene intermediate covalently modified the enzyme protein at lysine residues and destroyed the structure of heme. The vinyl and ethyl analogs of ELT showed minor enzyme inhibitory effect (less than 20%), while ELT inactivated more than 60% of the enzyme. The present study provided a novel bioactivation pathway of ELT and facilitated the understanding of the mechanisms of ELT-induced mechanism-based enzyme inactivation and liver injury.
Introduction

Erlotinib (ELT, 1, Scheme 1) is a highly potent reversible human epidermal growth factor receptor (EGFR) tyrosine kinase inhibitor (Dai et al., 2005). The proliferation and apoptosis of tumor cells are mediated by the EGFR signaling pathway (Scambia et al., 1995; Veale et al., 1993). Studies have shown that non-small-cell lung cancer (NSCLC) was caused by over-expression of EGFR (Jotte et al., 2015; Massarelli et al., 2013). ELT (Tarceva) was first approved by the Food and Drug Administration in 2004 for the treatment of metastatic NSCLC and advanced pancreatic cancer. Despite the beneficial effects of ELT, this anti-neoplastic drug has been linked to interstitial lung disease and hepatorenal syndrome (Makris et al., 2007; Tammaro et al., 2005). In addition, ELT has been associated with severe hepatic injury. The patients with normal liver function may suffer from hepatic impairment and even hepatic failure when treated with ELT (Saif, 2008; Liu et al., 2005; Huang et al., 2009). These rare but severe cases are often described as idiosyncratic toxicity. Though the mechanisms of ELT induced hepatic injury remain unknown, it has been proposed that covalent modifications of cellular macro-molecules by reactive metabolites might be responsible for the idiosyncratic adverse reactions (Stepan et al., 2011).

In humans, CYPs 3A4 and 3A5 are the critical enzymes in the liver metabolism of ELT while CYPs 1A2 and 2C8 make a minor contribution (Ling et al., 2006). In addition, ELT has been proven to be a mechanism-based inactivator of CYPs 3A4 and 3A5 (Li et al., 2010). Mechanism-based inactivation (MBI) is oftentimes referred to as quasi-irreversibly or irreversibly time-dependent inhibition, and/or suicide
Drug-induced MBI may cause more serious injuries to patients compared to reversible enzyme inhibition. The reason is that the MBI can destroy the structure of enzymes which would have to be regenerated, and loss of enzyme function may raise the possibility of unwanted drug-drug interactions. In theory, MBI arises when drugs are bioactivated by CYPs to generate reactive intermediates that cause irreversible loss of enzyme activity through covalent bonding to apoprotein and/or destroying structure of the heme (Orr et al., 2012; Kalgutkar et al., 2007).

As shown in Scheme 1, ELT (I) contains a phenyl-acetylene moiety. Metabolism of ELT is mediated by three primary routes in humans: (I) O-demethylation of the methoxy group of ELT, (II) 4-hydroxylation of the aromatic ring of aminophenylacetylene group to p-aminophenol, and (III) hydrolysis to phenyl acetic acid after oxidation of terminal acetylene (Ling et al., 2006). 4-Hydroxylation of the phenyl ring of ELT results in forming para-hydroxyaniline metabolite, which further undergoes oxidation to form a quinoneimine intermediate through loss of two electrons by CYPs. This quinoneimine intermediate has been detected as glutathione (GSH) conjugate by liquid chromatography/tandem mass spectrometry (Li et al., 2010).

Although the bioactivation of ELT by CYPs to a reactive quinoneimine metabolite has been well-defined, the possible bioactivation of this drug containing chemical scaffold terminal alkyne to a ketene intermediate is much less characterized. To our knowledge, the reactive oxirene and ketene intermediates can occur in the oxidation of the acetylene bond by CYPs and are thought to play a part in CYPs
In this study, a novel bioactivation pathway of ELT in the form of an amino-group based 4-bromobenzylamine (BBA) adduct in NADPH-dependent human liver microsomes (HLMs) incubated with ELT was reported. Moreover, the correlation of ketene intermediate formation and enzyme inactivation was investigated.
Materials and Methods

1. Reagents

Erlotinib (ELT, ≥99%) and chloroprotoferrihem was purchased from Dalian Meilun Biotech Co., Ltd. (Dalian, China). 4-Bromobenzylamine (BBA) was purchased from Shanghai Darui Chemical Co., Ltd. (Shanghai, China). m-Chloroperbenzoic acid (m-CPBA), palladium on activated charcoal (10%Pd-C), palladium on barium sulfate (10%Pd-BaSO₄) and quinoline were purchased from Aladdin Reagent Co., Ltd. (Shanghai, China). Pronase E, chymotrypsin, L-lysine, testosterone, 6β-hydroxytestosterone, DL-dithiothreitol (DTT), propranolol and reduced nicotinamide adenine dinucleotide phosphate (NADPH) were purchased from Sigma-Aldrich Co. (St. Louis, MO). Recombinant human P450 enzymes and HLMs were purchased from BD Gentest (Woburn, MA). Formic acid and trifluoroacetic acid were purchased from Fisher Scientific (Springfield, NJ). All organic solvents were obtained from Fisher Scientific (Springfield, NJ). All chemicals were of analytical grade and all solvents were of HPLC grade.

2. Chemical Synthesis of ELT Analogs

The ethyl and vinyl analogs (Scheme 1, 2 and 3) were synthesized by catalytic hydrogenation of ELT, according to the published literature with modification (Ban et al., 2004). To synthesize the ethyl analog (2), ELT (100 mg, 0.26 mmol) was dissolved in 10 mL methanol, and Pd-C (20 mg) was then added as a catalyst. The mixture was stirred in a sealed and hydrogen-purged reactor at room temperature and with pressure of 70 MPa for 4 h. The resulting reaction mixture was condensed
under vacuum after filtration. Silica gel column chromatography with eluate of dichloromethane and methanol (20:1) was used to purify the product. The resulting product was characterized by HPLC-UV, MS/MS and $^1$H NMR. Data and spectrums are shown in supporting information (Supplemental Fig. 1 and 2).

To synthesize the vinyl analog (3), ELT (100 mg, 0.26 mmol) was dissolved in 10 mL methanol, and Pd-BaSO$_4$ (20 mg) was then added as a catalyst. Aliquots (0.5 mL) of quinoline were added to quench the activity of the catalyst. The mixture was stirred in a sealed and hydrogen-purged reactor at room temperature and with pressure of 70 MPa for 2 h. After reaction, the mixture was subjected to a silica gel chromatography to remove quinoline and the fraction was purified by a semi-preparative HPLC system. The resulting product was characterized by HPLC-UV, MS/MS and $^1$H NMR. Data and spectrums are shown in supporting information (Supplemental Fig. 3 and 4).

3. Microsomal Incubations

ELT (100 μM from a dimethyl sulfoxide stock) and HLMs (1.0 mg protein/mL) were mixed in potassium phosphate buffer (PBS, pH 7.4). BBA (5 mM) (Scheme 1, 4), GSH (20 mM) or L-lysine (40 mM) were then added as a trapping agent, respectively. The total volume of the incubation was 500 μL. The reactions were initiated by adding NADPH (1 mM). Control samples did not contain NADPH. Reactions were terminated by mixing with ice-cold acetonitrile in a volume of 1:1. The resulting mixture was centrifuged at 19,000 g for 10 min at 4 °C to pellet protein. After the supernatants were evaporated to dryness under nitrogen stream, the
The reconstituted solution dissolved in 200 μL of acetonitrile: water (1:9, v/v) was injected into LC-MS/MS for analysis.

The ethyl and vinyl analogs were incubated in the same microsomal system as described above to detect possible BBA and GSH adducts, respectively.

4. Recombinant Human P450 Incubations

ELT was individually incubated with human recombinant P450 enzymes (100 nM), including CYPs 1A2, 2A6, 2B6, 2C9, 2C19, 2D6, 2E1, 3A4, and 3A5. The conditions were the same as the microsomal incubation experiments, except that recombinant P450 enzymes replaced the microsomes. An equal volume of ice-cold acetonitrile which contained propranolol (500 ng/mL) as internal standard was added to quench the reactions. The experiments were performed in triplicate.

5. Chemical Synthesis of ELT-BBA and ELT-Lysine Adducts

Synthesis of ELT-BBA adduct was achieved by initial epoxidation of ELT using m-chloroperbenzoic acid (m-CPBA), based on a reported method about peroxidation studies of acetylenes (McDonald et al., 1964). ELT (4 mg, 10.2 μmol) was dissolved in dichloromethane (10 mL), and m-CPBA (20 mg, 116.3 μmol) and BBA (20 mg, 107.5 μmol) were then added. After stirred for 20 h at room temperature, the reaction mixtures were centrifuged, and the supernatants were condensed under nitrogen stream. The resulting sample was reconstituted with acetonitrile–water (1:9) for LC–MS/MS analysis. Synthesis of ELT-lysine adduct was conducted by the same procedure as described above, except that BBA was replaced by lysine (15 mg,
102.7 μmol).

6. P450 3A Inactivation Studies

To determine whether terminal alkyne of ELT is responsible for P450 time-dependent inactivation, experiments were carried out in HLMs with ELT, ethyl and vinyl analogs as substrates, respectively. Primary incubations (200 μL) included one of the substrates (25 and 50 μM), 0.5 mg/ml HLMs, 1 mM NADPH, 3.2 mM MgCl₂, and 100 mM PBS (pH 7.4). After incubation at 0, 10, 20, and 30 min, the primary mixtures (40 μL) were transferred to the secondary incubation mixtures (120 μL), which included 200 μM testosterone, 1 mM NADPH, 3.2 mM MgCl₂, and 100 mM PBS (pH 7.4). The proportion of dimethylsulfoxide was 0.5% (v/v). An equal volume ice-cold acetonitrile containing propranolol as the internal standard was added to terminate the reaction after 15 min incubation. After centrifugation, the amount of 6β-hydroxytestosterone produced was measured by LC-MS/MS to determine the remaining CYP3A activity.

7. Microsomal Protein Modification and Digestion

ELT (100 μM) was mixed with 1.0 mg/mL HLMs and NADPH (1 mM). The final volume of incubation was 0.5 mL. After 30 min of incubation at 37 °C, the protein samples were denatured by 30 min of heating at 60 °C in a water bath, followed by centrifugation at 19,000 g for 10 min. Ammonium bicarbonate (50 mM, pH 8.0) was used to suspend the pellets, which were then mixed with 5.0 mM DTT. After incubation at 60 °C for 1 h, the resulting mixture was digested with a mixture of
Pronase E (2.5 mg/mL) and chymotrypsin (2.5 mg/mL) in the presence of CaCl$_2$ (5.0 mM) (Wang et al., 2015). After 15 h of incubation for 37 °C, the digestion mixtures were centrifuged at 19,000 g for 10 min and the supernatants were analyzed by LC-MS/MS.

To further investigate lysine-based protein adducts, protein digestion was also conducted after the ethyl or vinyl analogs was incubated with HLMs as described above.

8. Measurement of Native Heme Loss

The determination of loss of the native heme was achieved, according to the reported literature (Lin et al., 2002). ELT (50 μM) or ELT analogs were incubated with 0.5 mg/ml HLMs, 1mM NADPH for 30 min at 37°C. The control samples did not contain NADPH. The final volume of incubation mixtures were 0.5 mL. After incubation, the mixtures were extracted with butanone containing 10% trifluoroacetic acid. The residue was reconstituted in acetonitrile for LC-UV and LC–MS/MS analysis after the extracted solution was evaporated to dryness under nitrogen stream.

9. Analytical Conditions

Metabolites were characterized on a SCIEX 5500 mass spectrometer (SCIEX, Foster City, CA) equipped with an Agilent 1260 system (Agilent Technologies, Santa Clara, CA). A ZORBAX SB-C18 column (5.0 μm, 150 mm × 4.6 mm; Agilent Technologies, Santa Clara, CA) was used to perform HPLC separation. The mobile phase was water and acetonitrile with 0.1% formic acid. The gradient elutions for
analyzing ELT-BBA and ELT-lysine adducts and separating 6β-hydroxytestosterone were shown in Supplemental Table 1 and 2, respectively. All the metabolites were analyzed in a positive mode by multiple reaction monitoring (MRM) scanning via electrospray ionization, based on structurally specific fragmentation: \(m/z\) 595/479 and 597/481 for ELT-BBA adduct, \(m/z\) 556/428 for ELT-lysine adduct, \(m/z\) 305/269 for 6β-hydroxytestosterone and \(m/z\) 261/116 for propranolol as internal standard. Precursor ion (PI) scanning of \(m/z\) 169, 171 were also used for analyzing the ELT-BBA adduct. In addition, SCIEX 4000 Q-Trap (SCIEX, Foster City, CA) equipped with an Agilent 1260 system were used to analyze the MS/MS fragments of metabolites. Multiple reaction monitoring coupled with enhanced product ion (MRM-EPI) mode was employed. The optimized mass spectrometric instrument parameters are shown in supporting information (Supplemental Tab. 3). The Analyst 1.6.2 software was used to analyze all the data.

For the analysis of heme, chromatographic separation was conducted on a C4 column (250 × 4.6 mm, 5 µm, VYDAC, Hesperia, CA) with solvents including 0.1% trifluoroacetic acid in water and 0.05% trifluoroacetic acid in acetonitrile. The gradient was shown in Supplemental Table 4. The eluates were monitored at 400 nm and Q1MS scanning mode for the heme and possible heme adducts formed in the bioactivation of ELT.
Results

1. Enzyme-Dependent ELT-BBA Adduction

ELT was incubated in HLMs supplemented in BBA as a trapping agent. ELT-BBA adduct (Figure 1 for its structure) eluted at 11.6 min was detected by MRM scanning of $m/z$ 595/479, $m/z$ 597/481 and PI scanning of $m/z$ 169, $m/z$ 171 (Fig. 1E-H). The intensity ratios of these two scan mode (MRM and PI) were around 1:1, respectively. No peaks at the corresponding retention time were observed in the absence of NADPH (Fig. 1A-D) demonstrating that the formation of this adduct was NADPH-dependent. The MS/MS spectrum of the ELT-BBA adduct obtained from MRM-EPI scanning of $m/z$ 595/479 or 597/481 showed the characteristic fragment ions containing bromine at $m/z$ 169/171 (loss of $\text{C}_{22}\text{H}_{28}\text{N}_{4}\text{O}_{5}$), 479/481 (loss of $\text{C}_{6}\text{H}_{14}\text{O}_{2}$), 537/539 (loss of $\text{C}_{3}\text{H}_{5}\text{O}$) and fragment ions containing $N$-phenyl quinazolin-4-amine or oxidized carbonyl moiety without bromine at $m/z$ 266 (loss of $\text{C}_{14}\text{H}_{21}\text{NO}_{3}\text{Br}$) and 294 (loss of $\text{C}_{13}\text{H}_{21}\text{NO}_{2}\text{Br}$) (Fig. 1I). No such adduct was detected after incubation of the ethyl or vinyl analogs mixed with HLMs. This indicated that the adduct was formed by oxidation of terminal alkyne (Fig. 6B and 6C). To further characterize the adduct, ELT was reacted with $m$-CPBA and BBA. After the reaction, a product was detected which showed the same retention time and similar mass spectrum as that of the product formed in HLM incubations with ELT (Fig. 2).

In a separate experiment, the ethyl and vinyl analogs were individually incubated in the HLM incubation system supplemented with GSH as a trapping agent. For the
vinyl analog, two GSH conjugates were detected, including one derived from the epoxide (the representative MS/MS scanning of $m/z$ 719/590,446) (Supplemental Fig. 5A) and the other arising from the quinoneimine metabolite (MS/MS scanning of $m/z$ 717/588,444) (Supplemental Fig. 5B). As expected, only the quinoneimine derived GSH conjugate (MS/MS scanning of $m/z$ 719/590,446) (Supplemental Fig. 6) was detected in microsomal incubations with the ethyl analog.

2. P450 Enzymes Responsible for Bioactivation of ELT

Recombinant human P450 incubations were carried out for determining which P450 enzymes contribute to the oxidation to form the ELT-BBA adduct. CYPs 3A4 and 3A5 were found to be the major enzymes in contribution to the bioactivation of ELT. CYP3A4 displayed the highest catalytic activity, and the adduct formation by CYP3A5 was approximately 80% of that induced by CYP3A4. CYPs 2A6 and 2D6 also slightly catalyzed the ELT-BBA formation. The levels of adduct formation by CYPs 2A6 and 2D6 were about 10% and 13% of that formed by CYP3A4, respectively (Fig. 3).

3. Enzyme Inactivation Effects of ELT and ELT Analogs

Microsomal incubation of ELT at concentrations of 25 and 50 μM caused dramatic drop in CYP3A activity after 30 min by 36% ($P<0.01$) and 63% ($P<0.001$), respectively. Unlike ELT, the ethyl analog induced a slight activity decrease by only 3% and 15% ($P<0.05$), respectively. The vinyl analog showed similar degree in enzyme inactivation (approximately 9% and 20% ($P<0.05$), Fig. 4). The observed
significant loss of enzyme inactivation effect of ELT after saturation of the alkynyl group to vinyl and ethyl groups suggests that the terminal alkyne of ELT was involved in the P450 inactivation.

4. Lysine-based Protein Modification by ELT

ELT was incubated with HLMs. After incubation, the resulting microsomal protein was exhaustively digested by proteinase and then analyzed by monitoring MRM ion pair \(m/z\) 556/428. A lysine-derived adduct was eluted at 7.7 min (Fig. 5D). No such adduct was found in the protein digestion of HLMs in the absence of NADPH (Fig. 5C). The MS/MS characteristic fragment ions of amide moiety of the adduct, such as \(m/z\) 410 and 428 were detected. Product ion \(m/z\) 511 resulting from the loss of carboxyl group of this adduct (Fig. 5G). Fragment ions \(m/z\) 266 and 294 represented the same moieties corresponding to those of the ELT-BBA adduct (Fig. 1I and Fig. 5G). The ethyl or vinyl analog was also incubated with HLMs as described above. No lysine adduct was detected in protein digestion samples (Fig. 6E and 6F). An adduct with the same retention time at 7.7 min was observed in the HLMs incubation in the presence of NADPH and lysine (trapping agent) (Fig. 5B). Synthetic ELT-lysine adduct showed the same chromatographic and MS characteristics (Fig. 5E and 5F) as that of the corresponding adduct in incubation of HLMs as well as microsomal protein digestion.

5. Native Heme Loss during Enzyme Inactivation

The native heme of standard solution eluted at approximately 11.8 min and was
monitored by UV at 400 nm (Fig. 7A) and showed the MH$^+$ ion at \( m/z \) 616 (Fig. 7B). There was an about 70% decrease ($P<0.001$) in native heme of the ELT inactivated sample (Fig. 7C and 7D; Fig. 8). No marked decrease was observed in native heme of both ethyl (~5% decrease) and vinyl analog (~25% decrease) incubated samples (Fig. 7E-H; Fig. 8).
Discussion

ELT, an EGFR inhibitor used in the treatment of advanced NSCLC, was reported to cause severe liver injury during its clinical use (Saif, 2008; Liu et al., 2005; Huang et al., 2009). ELT can also inhibit CYP3A in a time-dependent manner. Formation of reactive metabolites have been implicated in the mechanism-based inactivation of enzymes and idiosyncratic toxicity. ELT was reportedly biotransformed to a quioneimine metabolite while ketene intermediates were not detected using N-acetyl-lysine as a trapping agent (Li et al., 2010). Despite this, we speculated that the terminal alkyne group of ELT would be oxidized to the corresponding ketene intermediate which may play a role in CYP3A inactivation. Ketenes are chemically reactive and unstable, trapping techniques are in need for the metabolite identification. Ketene intermediates reportedly modified N-terminal α-amino groups of amino acid residues and ε-amino group of lysine residues of apoproteins (Chan et al., 2012; Banzon et al., 1995). Thus, we tested several primary amines as trapping agents, and BBA was found to be the one which is the most effective. Additionally, BBA contained an element bromine, and we took the advantage of its 1:1 natural isotope pattern. This made it easier to monitor targeting analytes by mass spectrometry (Wang et al., 2014), by which we succeeded in detecting ELT-BBA adduct in MRM mode by monitoring of ion pairs m/z 595/479, 597/481, and m/z 169, 171 (Fig. 1). The observed adduct clearly indicates the production of ELT-derived ketene intermediate. Moreover, one of the three primary metabolism routes of ELT was oxidation of terminal acetylene followed by hydrolysis to phenyl acetic acid which
was proved to be the stable metabolite of the ketene intermediate (Ling et al., 2006). Formation of the ELT-BBA adduct was found to require the NADPH in the HLM incubation indicating that P450 enzymes were involved in bioactivation of ELT.

To better understand the metabolic activation of ELT, the ELT-BBA adduct was chemically synthesized by reacting ELT with \textit{m}-CPBA and BBA. \textit{m}-CPBA is an oxidant agent widely used in epoxidation reactions. By analogy to the epoxidation of alkenes, the first product of an acetylene would be an oxirene and then the reactive ketene was generated by rearrangement of the oxirene (McDonald et al., 1964). Unfortunately, we failed to obtain enough amount of the ELT-BBA adduct for NMR characterization, due to the poor yield of the chemical synthesis.

MBI is often referred to as “suicide inactivation”. The cytochrome P450s might often be targets of the reactive intermediates generated in the active site of enzyme (Hollenberg et al., 2008). In previous study, ELT has proved to be a mechanism-based inactivator of CYPs 3A4 and 3A5. CYPs 3A4 and 1A1 were reported to catalyze the formation of the quioneimine metabolite of ELT, but ELT failed to show inhibitory effect on CYP1A1 (Li et al., 2010). The present study demonstrated that both CYPs 3A4 and 3A5 catalyzed the formation of the ketene intermediate of ELT. It is important to define the role of the two reactive metabolites in the enzyme inactivation. To this end, we replaced the alkyne of ELT with ethyl and vinyl groups and evaluated their inhibitory effects on the enzymes. Both compounds only revealed very weak enzyme inactivation. Structurally, the vinyl group might also be oxidized to epoxide possibly participating in protein modification,
due to its constrained bond angle (Pearce et al., 2002). To address this possibility, the vinyl analog was incubated in HLMs supplemented with GSH as a trapping agent. Two GSH conjugates were detected by LC-MS/MS after incubation (Supplemental Fig. 5A and B). One resulted from the corresponding epoxide metabolite, and the other arose from the quinoneimine intermediate (Li et al., 2010). Additionally, similar microsomal incubations were carried out to determine the metabolic activation of the ethyl analog. Only one GSH conjugate was detected (Supplemental Fig. 6), and the observed molecular ion of the conjugate matched the molecular weight of the GSH conjugate derived from the corresponding quinoneimine. The vinyl and ethyl analogs showed little enzyme inhibitory effect (less than 20%), while ELT inactivated more than 60% of the enzyme in a time-dependent manner. This indicates that the formation of ketene, together with quinoneimine intermediate, contributed to CYP3A inactivation, and the ketene intermediate played the major role.

Previous studies on the drug induced MBI have led to the suggestion that the different mechanisms of inactivation via covalent binding to apoprotein and/or destroying the native heme group (Ortiz de Montellano et al., 1985; Lin et al., 2010; Blobaum, 2006). Therefore, we analyzed the products of microsomal protein digestion samples pretreated with ELT and measured the relative amount of the native heme loss to better understand the association between bioactivation of ELT and enzyme inactivation. The observed decrease in the amount of the heme after microsomal incubations suggests that the reactive intermediate of ELT would modify the heme, at least partially responsible for the loss of the enzyme activity. Apart
from the measurement of the native heme loss, we also attempted to check for the possible heme adducts, using HPLC-UV and MS method. No stable heme adduct was detected. We propose that unknown heme fragment or unstable heme adducts were formed in the inactivation reaction so that they cannot be detected. This kind of heme destruction has also been reported by other mechanism-based inactivation studies (Amunugama et al., 2012; Lin et al., 2013). Taken together, our findings suggest that ELT causes inactivation through covalent binding of the bioactivated ELT to the apoprotein of CYPs and the native heme destruction.

Previous clinical studies showed that ELT induced liver injury, which might also be associated with the metabolic activation process. According to the tissue distribution studies (Li et al., 2010), liver was most likely to be toxic target organ, due to accumulated ELT. However, the definitive relationship between the reactive metabolites (ketene and quioneimine) of ELT and liver toxicity remains unknown, though the ketene intermediate played a major role in the enzyme inactivation.

In conclusion, a reactive ketene intermediate was detected in incubations of ELT with HLMs. CYPs 3A4 and 3A5 mainly contributed to the acetylenic bioactivation of ELT. The MBI of the enzymes by ELT was caused by formation of the ketene intermediate. These data are helpful to justify possible drug-drug interactions and to facilitate the understanding of the mechanisms of hepatotoxic action of ELT reported in clinic.
Authorship Contributions:

Participated in research design: Zheng

Conducted experiments: Zhao, Li, and Yang

Performed data analysis: Zhao, Chen, and Peng.

Wrote or contributed to the writing of the manuscript: Zhao and Zheng.
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Figure legends

**Figure 1.** Characterization of ELT-BBA adduct generated from HLM incubations. Extracted ion (m/z 595/479 and m/z 597/481) chromatograms obtained from LC-MS/MS analysis of HLM incubations containing ELT and BBA in the absence (A and B) or presence (E and F) of NADPH. Extracted ion chromatograms of PI scans of m/z 169 and m/z 171 obtained from LC-MS/MS analysis in the absence (C and D) or presence (G and H) of NADPH. (I) MS/MS spectrum of ELT-BBA adduct generated in microsomal incubations obtained from LC-MS/MS.

**Figure 2.** Characterization of ELT-BBA adduct generated from chemical synthesis. Extracted ion chromatogram of MRM scans of m/z 597/481 obtained from microsomal incubations (A) and chemical synthesis (C). MS/MS spectrum of ELT-BBA adduct generated in microsomal incubations (B) and chemical synthesis (D) obtained from LC-MS/MS.

**Figure 3.** Formation of ELT-BBA adduct in individual recombinant P450 enzyme incubations containing NADPH, ELT and BBA. Data represent the mean ± S.D. (n = 3).

**Figure 4.** Time-dependent inactivation of CYP3A by ELT and ELT analogs. HLMs (0.5 mg/ml) was incubated with the following substrates: vehicle (▪, black), ELT (◆, red), vinyl analogs (●, blue) and ethyl analogs (▲, green), at concentrations of 25 (A) and 50 (B) μM in the presence of NADPH at 37 °C for 0, 10, 20 and 30 min. The residual enzymatic activity at each time point was determined through calculating peak area ratio of the analytes compared with the internal standard, and the enzymatic
activities at 0 min were normalized to 100% at each concentration. Data represent the mean ± S.D. (n = 3). *P<0.05, **P<0.01, ***P<0.001 represent student's t-test P values compared with vehicle groups.

**Figure 5.** Characterization of ELT-lysine adduct. Extracted ion chromatogram of MRM scans of m/z 556/428 obtained from LC-MS/MS analysis of HLM incubations containing ELT and lysine in the absence (A) or presence (B) of NADPH. Extracted ion chromatograms obtained from the analysis of the proteolytic digestion of human microsomal protein exposed to ELT in the absence (C) or presence (D) of NADPH. Extracted ion chromatogram of MRM scans of m/z 556/428 obtained from chemical synthesis (E). MS/MS spectrum of ELT-lysine adduct generated in chemical synthesis (F) and proteolytic digestion (G) obtained from LC-MS/MS.

**Figure 6.** Formation of BBA and lysine based adduct with ELT and ELT analogs.

Extracted ion chromatograms of PI scans of m/z 171 obtained from LC-MS/MS analysis of HLM incubations containing ELT (A), ethyl analog (B) and vinyl analog (C). Extracted ion chromatograms of MRM scans of m/z 556/428 obtained from the analysis of the proteolytic digestion of human microsomal protein exposed to ELT (D), ethyl analog (E) and vinyl analog (F).

**Figure 7.** HPLC profile of the native heme analyses. HPLC chromatogram (A) at 400 nm and Q1MS spectrum (B) of the standard heme. HPLC chromatograms of HLM incubations containing ELT (C), vinyl analog (E) and ethyl analog (G) in the presence of NADPH. HPLC chromatograms of HLM incubations containing ELT (D), vinyl analog (F) and ethyl analog (H) in the absence of NADPH.
Figure 8. Native heme loss of ELT, vinyl analogs and ethyl analogs in the absence and presence of NADPH. Relative contents (%) of heme loss was determined through calculating peak area of the analytes compared with the vehicle groups. The values of vehicle groups were normalized to 100%. Data represent the mean±S.D. (n=3). ***P<0.001 represent student's t-test P value.
Scheme legends

Scheme 1. Structures of ELT (1), ethyl analog (2), vinyl analog (3) and BBA (4).

Scheme 2. Proposed mechanism of ELT metabolic activation.
Figure 2

[Image of a figure with four panels labeled A, B, C, and D. Each panel shows a graph with intensity values and time or mass-to-charge (m/z) values, labeled with peaks at specific masses and intensities.]
Figure 3
Figure 4

A

B

Log % Activity Remaining

Vehicle
Ethyl analogs
Vinyl analogs
Erlotinib

Time (min)
Figure 5
Figure 6

A

B

C

D

E

F
Figure 7
Figure 8

![Figure 8](image-url)
Scheme 1

1

2

3

4

Br

NH₂
Scheme 2

Erlotinib (ELT) → P450 oxidase → Quinoneimine → GSH

Oxirene + Ketene → ELT-BBA adduct

Heme → Native heme loss → Mechanism-based inactivation → Heme → NH₃-Protein → Hepatotoxicity