Natural product juglone targets three key enzymes from Helicobacter pylori: inhibition assay with crystal structure characterization

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Abstract

Aim: To investigate the inhibition features of the natural product juglone (5-hydroxy-1,4-naphthoquinone) against the three key enzymes from Helicobacter pylori (cystathionine γ-synthase [HpCGS], malonyl-CoA:acyl carrier protein transacylase [HpFabD], and β-hydroxyacyl-ACP dehydratase [HpFabZ]).

Methods: An enzyme inhibition assay against HpCGS was carried out by using a continuous coupled spectrophotometric assay approach. The inhibition assay of HpFabD was performed based on the α-ketoglutarate dehydrogenase-coupled system, while the inhibition assay for HpFabZ was monitored by detecting the decrease in absorbance at 260 nm with crotonoyl-CoA conversion to β-hydroxybutyryl-CoA. The juglone/FabZ complex crystal was obtained by soaking juglone into the HpFabZ crystal, and the X-ray crystal structure of the complex was analyzed by molecular replacement approach.

Results: Juglone was shown to potently inhibit HpCGS, HpFabD, and HpFabZ with the half maximal inhibitory concentration IC\(_{50}\) values of 7.0±0.7, 20±1, and 30±4 µmol/L, respectively. An inhibition-type study indicated that juglone was a non-competitive inhibitor of HpCGS against O-succinyl-L-homoserine (K\(_i\)=αK\(_i\)=24 µmol/L), an uncompetitive inhibitor of HpFabD against malonyl-CoA (αK\(_i\)=7.4 µmol/L), and a competitive inhibitor of HpFabZ against crotonoyl-CoA (K\(_i\)=6.8 µmol/L). Moreover, the crystal structure of the HpFabZ/juglone complex further revealed the essential binding pattern of juglone against HpFabZ at the atomic level.

Conclusion: HpCGS, HpFabD, and HpFabZ are potential targets of juglone.

Key words
cystathionine γ-synthase; malonyl-CoA:acyl carrier protein transacylase; β-hydroxyacyl-ACP dehydratase; inhibitor type; complex structure

Abbreviation
IC\(_{50}\), the half maximal inhibitory concentration; K\(_i\), the dissociation constant for inhibitor binding. CoA, coenzyme A; ACP, acyl carrier protein transacylase; PDB, Protein Data Bank; HO-HxoDH, D-2-Hydroxyisocaproate dehydrogenase.

Introduction

Helicobacter pylori (H pylori) is a Gram-negative bacterium associated with a number of human diseases, including gastritis, peptic ulceration, and gastric cancer\(^1\). H pylori has been recognized as a pathogenic bacterium that chronically infects approximately 50% of the world’s human population\(^2\). The rapid infection of H pylori has become a severe threat against human health. Usually, the treatment of H pylori infections involves the administration of a combination of antibiotics and other drugs. However, the overuse and misuse of antibacterial agents have resulted in the generation of antibiotic resistant strains. For example, in the UK, the metronidazole resistance was found by 31.7% against H pylori isolates\(^3\). Accordingly, the alarming rise of antibiotic resistance among the key bacterial pathogens has strongly stimulated an urgency to develop novel antibacterial agents acting on new drug targets.

The natural product juglone (5-hydroxy-1,4-naphthoquinone; Figure 1A), from the leaves and unripe hulls of the fruits of Juglans regia, was found to demonstrate cytotoxicities when added in cell cultures\(^4–6\). Such an effect might result from juglone’s involvement in the block of transcription\(^7\), or K\(^+\) channel against cell membrane\(^8\), the inhibition of the peptidyl-prolyl isomerase Pin1\(^7,9\), or the generation of hydrogen peroxide\(^5\). In addition, juglone was found to possess antibacterial properties, including anti-H pylori and
antifungal properties\textsuperscript{[10]}, however, no target information has ever been disclosed.

In the present study, we reported that juglone functions as a multitargeted inhibitor against 3 key enzymes from \textit{H pylori}: cystathionine \(\gamma\)-synthase (HpCGS), malonyl-CoA:acyl carrier protein transacylase (HpFabD), and \(\beta\)-hydroxyacyl-ACP dehydratase (HpFabZ). The analyzed crystal structure of the HpFabZ/juglone complex has further clarified the essential binding feature of juglone against HpFabZ at the atomic level.

It is known that different organisms display distinct spectra of transsulfuration enzymes. Most plants and microbes employ only the forward pathway from cysteine to homocysteine and methionine, and mammals carry out only the reverse transsulfuration, while fungi take up transsulfuration and use only the forward pathway from cysteine to homocysteine or \(L\)-cystathionine. For microorganisms, such a reaction is the first step involved in the transsulfuration pathway that converts \(L\)-cysteine into \(L\)-homocysteine. Since CGS is absent in non-ruminant animals that require a dietary source of \(L\)-homocysteine or \(L\)-methionine\textsuperscript{[12,13]}, it has thus been regarded as an attractive target for antibiotic discovery\textsuperscript{[14]}. However, the \textit{metB} gene seems to be unessential for \textit{H pylori} growth as reported by Salama \textit{et al.}\textsuperscript{[15]}

Fatty acid biosynthesis (FAS) is an essential pathway for the survival of the organism since fatty acid is the major component of cell membranes and possesses important biological functions. In nature, according to the enzymes involved in the pathway, fatty acid biosynthesis is divided into 2 types: type I (FAS I) and type II (FAS II)\textsuperscript{[16-18]}. In the FAS I system, which found in animals, the biosynthesis of fatty acid is catalyzed by a multi-enzyme, which is a single polypeptide with 8 distinct domains. However, in the FAS II system, the reactions are carried out by a series of structurally dissociated enzymes as discovered in most bacteria\textsuperscript{[16,17]}. Due to the large differences between these 2 FAS systems, enzymes like FabD (EC2.3.1.39) and FabZ (EC4.2.1.60) involved in type II fatty acid biosynthesis have been developed as potential targets for the discovery of antibacterial agents\textsuperscript{[17]}. FabD catalyzes the transfer of a malonyl moiety from malonyl-CoA to holo-ACP, forming malonyl-ACP as the elongation substrate for the fatty acid biosynthesis\textsuperscript{[19-22]}, while FabZ is the primary dehydratase that participates in the elongation cycles of saturated and unsaturated fatty acid synthesis\textsuperscript{[23-26]}

It is expected that our current work might help understand the possible antibacterial mechanism for juglone and provide useful structural information for the discovery of an anti-\textit{H pylori} agent by using juglone as a potential multitargeted lead compound.

**Materials and methods**

**Materials** \textit{H pylori} strain SS1 was maintained at our institute. The \textit{Escherichia coli} host strain BL21(DE3) was purchased from Stratagene (Germany). The natural product juglone was from the in-house chemical library established in our laboratory. All other chemicals used were of reagent grade or ultra-pure quality.

**Expression and purification of HpCGS** The HpCGS enzyme was cloned, expressed, and purified according to our recently published work\textsuperscript{[27]}. Briefly, \textit{H pylori} SS1 strain genomic DNA was PCR-amplified from \textit{H pylori} SS1 strain genomic DNA and ligated into a pET28b expression vector (Novagen, Germany) and transformed into BL21(DE3) after restriction digestion. The correct clones were expressed at 37ºC, and the purified HpCGS enzyme was obtained by using buffer C (20 mmol/L Tris-HCl, pH 8.0, 500 mmol/L NaCl, and 120 mmol/L imidazole) as elution buffer.

**Expression and purification of HpFabD** The cloning, expression, and purification of HpFabD were performed according to our published description\textsuperscript{[19]}. The purified HpFabD protein was dialyzed against buffer A (20 mmol/L Tris-HCl, pH 8.0) to remove imidazole.

**Expression and purification of HpFabZ** The cloning, expression, and purification of HpFabZ were based on our recently published work\textsuperscript{[29]}. The purified protein was obtained by dialysis against buffer B (20 mmol/L Tris-HCl, pH 8.0, 500 mmol/L NaCl, and 1 mmol/L EDTA).

**HpCGS inhibition assay** The HpCGS enzyme inhibition assay was carried out according to our recently published study\textsuperscript{[27]}. The \(K_{IC_{50}}\) value of juglone against HpCGS was obtained by fitting the data to a sigmoid dose-response equation using Origin software (OriginLab, Northampton, Massachusetts, USA). Inhibitor type and inhibition constants \(K_i\) and \(K_{IC_{50}}\) were determined by the double-reciprocal (Lineweaver-Burk) plot, and the \(V_{max}\) plot and slopes of the lines from the double-reciprocal plot were used as a function of inhibitor concentrations.
HpFabD inhibition assay  The HpFabD enzyme inhibition assay was performed based on our reported approach\(^\text{[28]}\). To determine the IC\(_{50}\) value of juglone against malonyl-CoA, HpFabD was incubated for 5 min at room temperature with different concentrations of juglone (0–100 µmol/L) in a total volume of 100 µL. To investigate the inhibition mode of juglone against malonyl-CoA, different concentrations of juglone (0, 8, 10 µmol/L) were used, and the reaction was initiated by the addition of malonyl-CoA (5–30 µmol/L). The Ki value was obtained by the Dixon plot.

HpFabZ inhibition assay  The enzymatic inhibition assay of the HpFabZ enzyme was monitored by using the reported spectrophotometric method\(^\text{[26,29,30]}\). The IC\(_{50}\) value of the inhibitor was estimated by fitting the inhibition data to a dose-dependent curve using a logistic derivative equation\(^\text{[31]}\). The inhibitor mechanism was determined in the presence of various inhibitor concentrations (0–50 µmol/L). After 2 h incubation, the reaction was started by the addition of crotonoyl-CoA (10–250 µmol/L). The Ki value was obtained from the Dixon plot and secondary plots.

Crystallization and data collection  The crystallization of the HpFabZ/juglone complex was carried out according to our previous work\(^\text{[30]}\). The purified HpFabZ enzyme in 20 mmol/L Tris-HCl (pH 9.0) and 500 mmol/L NaCl was concentrated to ~10 µg/mL. For crystallization, 1 µL of the enzyme mixture was mixed with an equal volume of the reservoir solution containing 2 mol/L sodium formate and 0.1 mol/L sodium acetate trihydrate at pH 3.6–5.6, and benzamidine-HCl was added to a final concentration of 2% (w/v). The mixture was equilibrated against 500 µL of the reservoir solution at 277K by the hanging-drop vapor-diffusion method. Crystals of dimensions 0.5×0.3×0.3 mm\(^3\) were obtained after 7 d. Juglone was added to the original drop to a final concentration of ~20 mmol/L, and the crystals were soaked for 24 h.

Diffraction data were collected at 100K using CuK\(\alpha\) X-ray with a Rigaku R-AXIS IV\(++\) image plate (Rigaku Corp, Tokyo, Japan). Before the crystals were flash-frozen in liquid nitrogen, the drop was dehydrated against 500 µL reservoir solution containing 4 mol/L sodium formate for 24 h. The data were processed using HKL2000\(^\text{[32]}\). The crystallographic statistics are summarized in Table 1. The structure was solved by the molecular replacement approach using the crystal structure of HpFabZ as the search model and refined by the program CNS\(^\text{[33]}\). Electron density interpretation and model building were performed using the computer graphics program “coot”\(^\text{[34]}\).

Results

Juglone is a multitargeted inhibitor against CGS, FabD,

| Structure | Complex |
|-----------|---------|
| Data collection | |
| Space group | P2\(_1\),2\(_1\),2\(_1\) |
| Cell dimensions (Å) | |
| a | 73.9826 |
| b | 100.3689 |
| c | 186.236 |
| α | 90 |
| β | 90 |
| γ | 90 |
| Wavelength | 1.5418 |
| Resolution(Å) \(^1\) | 50.00-2.40 |
| R \(_{free}\) | 0.106 (0.301) |
| R/σ | 13.4 (4.1) |
| Completeness (%) | 99.8 (99.9) |
| Redundancy | 5.7 (5.4) |

| Refinement | |
| N\(_2\) reflections | 55018 (7903) |
| R \(_{free}\)/R \(_{sym}\) \(^3\) | 0.192/0.228 |
| N\(_2\) atoms | 7289 |
| Protein | 63 |
| Benzamidine-HCl | 6 |
| Cl\(^-\) ions | 492 |
| Water molecules | 26 |
| Compound | 20.344 |
| B-factors | 20.71 |
| Protein | 38.2 |
| Benzamidine-HCl | 23.988 |
| Cl\(^-\) ions | 46.33 |
| Water molecules | 0.007 |
| Compound | 1.4 |

\(^1\) Numbers in parentheses represent statistics in highest resolution shell.

\(^2\) \(R_{free}=\Sigma||F_o|-|F_c||/\Sigma|F_o|\)

\(^3\) \(R_{sym}=(\Sigma||F_o|-\Sigma|F_c||)/\Sigma|F_o|\)

and FabZ enzymes  Inhibition against CGS enzyme as evaluated from the enzymatic assay, the natural product juglone was identified to show strong inhibitory activity against HpCGS with an IC\(_{50}\) of 7.0±0.7 µmol/L (Table 2). In a separate control experiment by directly using 2 mmol/L \(\alpha\)-ketobutyrate as the substrate, juglone showed no inhibition activity against D-2-hydroxyisocaproate dehydrogenase (HO-HxoDH) at its concentration, even up to 50 µmol/L, further confirming that juglone is a HpCGS inhibitor. The Lineweaver-Burk plot-based analysis indicated that juglone
prevented the binding of O-succinyl-L-homoserine (L-OSHS) to HpCGS in a non-competitive fashion (Figure 2A), and the inhibition constants \( K_i \) of 24 \( \mu \text{mol/L} \) and \( K_i' \) of 24 \( \mu \text{mol/L} \) (Table 2) were determined from the Dixon plot and secondary plots.

**Inhibition against HpFabD** As indicated in Table 2, juglone could inhibit HpFabD with an IC\(_{50}\) against malonyl-CoA at 20±1 \( \mu \text{mol/L} \) as a non-competitive inhibitor (Figure 2B). The \( \alpha K_i \) value of 7.4 \( \mu \text{mol/L} \) (Table 2) was thereby obtained by the Dixon plot.

**Inhibition against HpFabZ** As indicated in Figure 2C, besides targeting CGS and FabD, juglone was also detected as a competitive inhibitor against HpFabZ by an IC\(_{50}\) of 30±4 \( \mu \text{mol/L} \) (Table 2) with respect to the substrate crotonoyl-CoA. A competitive value of \( K_m \) at different inhibitor concentrations as a function of the inhibitor concentration.

**Crystal structure analysis of the HpFabZ/juglone complex** In order to gain the essential inhibition mechanism at the atomic level for juglone against these 3 enzymes, we tried crystallizing the enzymes in complex with juglone and succeeded in obtaining the crystal of the HpFabZ/juglone complex, which was well determined against 2.4 \( \text{Å} \) level data (PDB code: 3B7J, Figure 3 and Table 1). As shown in the analyzed complex structure, juglone binds to HpFabZ in 2 binding models similar to our previous binding model for HpFabZ binding to the inhibitor\(^{30}\) (Figure 4). In model A, juglone fits into the groove around the entrance (Figure 4A) and locates between the phenol ring of Tyr100 and the pyrrolidine ring of Pro112\(^\prime\), forming a sandwich structure (the prime indicates a residue from the other subunit in the dimer). As shown in Figure 4B, the carbonyl oxygen of juglone forms H-bonds with the nearby water chain, which also forms H-bonds with the back bone carbonyl oxygen atoms of Phe65\(^\prime\), Ile111\(^\prime\), Val 146\(^\prime\), and Phe101, as well as the back bone nitrogen of Val113\(^\prime\). In model B, juglone entered into the middle of the tunnel near the active site of HpFabZ (His58 and Glu72\(^\prime\); Figure 4C) and was stabilized via the hydrophobic interactions between residues Leu21, His23, Ala75, Phe83, Ile98, Val 99, and Phe59\(^\prime\). At this stage, the carbonyl oxygen of juglone

| Enzyme   | Inhibition mode | \( IC_{50}\) (\( \mu \text{mol/L} \)) | \( K_i \) (\( \mu \text{mol/L} \)) | \( K_i' \) (\( \mu \text{mol/L} \)) |
|----------|----------------|---------------------------------|---------------------------------|---------------------------------|
| HpCGS    | Non-competitive | 7.0±0.7                         | 24                              | 24                              |
| HpFabD   | Uncompetitive   | 20±1                            | -                               | 7.4                             |
| HpFabZ   | Competitive     | 30±4                            | 6.8                             | -                               |

**Table 2.** Inhibition data of juglone against CGS, FabD, and FabZ enzymes from \( H. pylori \).
Ala75, the Ne2 of His 23 and His58', and Oε1 and Oε2 of Glu72 (Figure 4D).

Discussion

Recently, the rapid infection of *H pylori* has become a severe threat to human health, and the discovery of new, effective drugs has attracted more and more attention. Many antibacterial agents used for standard pathogens in clinics are subject to high-level resistance resulting from single-step mutations from target enzymes, whereas multitargeted agents might display low potential for rapid endogenous resistance development in pathogenic bacteria[35].

It is reported that juglone could strongly inhibit *H pylori* growth at a low MIC of 1.6 µg/mL[36,37], although no acting target information has ever been disclosed. In the current work, we discovered that juglone functions as a multitargeted inhibitor against 3 key enzymes, HpCGS, HpFabD, and HpFabZ, which provides some clues for the inhibitory mechanism underlying juglone’s anti-*H pylori* activity.

In the last 10 years, many CGS and FabZ inhibitors have been discovered. However, most of the inhibitors are single-targeted, except some flavonoid derivatives (luteolin and (−)-catechin gallate), which were reported to inhibit *Plasmodium falciparum* by acting as multitargeted inhibitors against FabG, FabZ, and FabF of *Plasmodium falciparum*[38].

In our previous work[27], we reported that the natural products α-lapachone and 9-hydroxy-α-lapachone demonstrated inhibitory activities against HpCGS. Considering the structure similarity between these 2 compounds and juglone, each of them contains a common quinone group (Figure 1). Although the natural products α-lapachone and 9-hydroxy-α-lapachone exhibited no inhibition activities against HpFabD and HpFabZ, as evaluated by the inhibition assay, the quinone group seemed to have some links to the CGS enzyme inhibition.

Previously[30], we reported 2 crystal structures of HpFabZ in complex with 2 small molecular inhibitors, and 2 binding models of an inhibitor against HpFabZ were supposed. In

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**Figure 3.** Stereo view of the omitted electron density map contoured at 1.0σ around juglone. Monomers A/B are yellow/magenta, monomers C/D are blue/orange, and juglone is salmon.

**Figure 4.** Crystal structure of the HpFabZ/juglone complex. (A) binding model A of juglone in the HpFabZ active tunnel. Electrostatic surface of the active tunnel is rendered by a color ramp from red to blue. Critical residues and water molecules that interact with juglone are shown. Dimer A/B is yellow/magenta, and dimer C/D is blue/orange; juglone is salmon. Water and chlorine atoms are shown as red and green spheres. Oxygen and nitrogen are red and blue, respectively. (B) H-bond network around juglone in binding model A. H-bonds are shown as yellow dashes. Residues around juglone are labeled. (C) binding model B of juglone in the active tunnel. (D) H-bond network around juglone in binding model B. Image produced by Pymol[40].

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could form H-bonds with the nearby water chain that also forms H-bonds with the back bone carbonyl oxygens of Ala75, the Ne2 of His 23 and His58', and Oε1 and Oε2 of Glu72 (Figure 4D).
binding model A, the inhibitor is sandwiched between the phenol ring of Tyr100 and the pyrrolidine ring of Pro112', and forms a $\pi-\pi$ interaction. In binding model B, the inhibitor locates into the middle of the tunnel via hydrophobic interactions. Juglone also fits in these 2 models (Figure 3), of which the position in model A is corresponding to the groove around the entrance, and the position in model B is corresponding to the active site of the tunnel near the catalytic residues His58 and Glu72, in agreement with the competitive inhibitory properties of juglone against HpFabZ determined by the inhibition type study. However, in comparison with this previous HpFabZ-inhibitor crystal structure analysis, 2 major differences are found for juglone binding to HpFabZ. First, in the binding model of HpFabZ/juglone, besides the $\pi-\pi$ or hydrophobic interactions, juglone was also stabilized by the H-bonds formed by the water chain between the enzyme and juglone. The other major difference is that in binding model B, juglone is inserted deeper to the tunnel (near Phe83) with a more suitable hydrophobic environment for the binding, different from the previously reported HpFabZ-inhibitor binding case, where the inhibitor formed a sandwich-like conformation with Ile98 and His59' of HpFabZ. It is thus expected that the information based on the HpFabZ/juglone complex would be useful for structure-guided drug design which might provide novel competent HpFabZ inhibitors.

In summary, in the present study, we have discovered that the natural product juglone is a multitargeted inhibitor against 3 key enzymes CGS, FabD, and FabZ from *H. pylori*. The obtained crystal structural data for the HpFabZ/juglone complex has further clarified the essential binding feature of juglone against HpFabZ at the atomic level. Although further experiments, such as a cell labeling assay, are needed to determine whether or not these 3 enzymes are targets responsible for juglone’s anti-*H. pylori* activity, juglone could hopefully serve as a potent lead compound for further inhibitory development as stated, and that the multi-targeted inhibitors might provide low potential for rapid endogenous resistance development.

**Author contribution**

Prof Xu SHEN, Prof Hua-liang JIANG and Prof Li-hong HU designed the research. Yun-hua KONG performed the research, analyzed data and wrote the paper. Liang ZHANG determined the complex structure. Zhen-yi YANG synthesized the compound and Cong HAN contributed the platform for determining MIC values against Hp.

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