Protective effect of *Glechoma hederacea* extract against gallstone formation in rodent models

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Abstract

**Background:** Our current study aimed to evaluate the effect of an *Glechoma hederacea* extract (Hitrechol®) in normal rats and gallstone diseased mice to explore its underlying mechanisms. Normal rats and C57BL/6 mice with/without cholesterol gallstone were used in this study.

**Methods:** To monitor the effect of Hitrechol® on bile secretion, bile flow rates at 15 min interval until 2 h post-dosing in normal rats treated with vehicle and Hitrechol® were compared using multiple t-test with a \( p < 0.05 \) considered as statistically significant different. To further evaluate the effect of Hitrechol® against the development of gallstone in lithogenic diet treated mice, mice were treated with vehicle or Hitrechol® (QD-once daily or TID-three times daily) for 3 weeks followed by comparing the levels of bile composition among the treatment groups. In addition, the anti-oxidative biomarkers in liver and anti-inflammatory biomarkers in serum were detected and compared among all the treatment groups to evaluate the hepato-protective effect of Hitrechol®. The obtained levels of biomarkers and bile composition were compared among different treatment groups using one-way ANOVA tests followed by Tukey’s multiple comparisons with \( p < 0.05 \) considered as statistically significant.

**Results:** Despite no significant impact on the bile flow rate, Hitrechol® TID treatment dramatically decreased size and amount of gallstone crystals and total cholesterol level \( (p < 0.05) \), as well as total bile acid \( (p < 0.05) \) and several types of bile acid \( (p < 0.05) \) levels in gallstone disease model mice. Hitrechol® TID treatment could significantly decrease the frequencies of hepatocyte necrosis and lipid aggregation notably as well as increase the antioxidant enzyme level \( (p < 0.05) \) in the liver.

**Conclusions:** Our findings for the first time demonstrated the beneficial effect of Hitrechol® against gallstone via its litholytic, liver-protective and antioxidant activities.

**Keywords:** *Glechoma hederacea* extract, Hitrechol®, Herbal medicine, Cholesterol gallstone, Lithogenic diet, Bile composition
Background

Gallstone disease is one of the most common digestive diseases, affecting 10–20% of the global adult population [1]. More than 90% of gallstones are cholesterol solid crystals formed in the gallbladder. The factors for causing cholesterol gallstone disease include improved standards of living, chronic high cholesterol diet, and overnutrition. In the normal physiological state, bile is produced in the liver and is secreted into the duodenum to digest food [2]. Bile contains bile salts, phospholipids, cholesterol, proteins, and bilirubin, all in stable equilibrium states [2]. However, under certain pathophysiological conditions, especially the supersaturated bile of cholesterol, the relative excess cholesterol will precipitate as solid crystals, then aggregate, fuse, and eventually form gallstones within a gallbladder-secreted mucin gel that cause disease and complaints [2, 3]. Although laparoscopic cholecystectomy is currently considered the gold standard in treating patients with symptomatic gallstones, new perspectives regarding medical therapy of cholesterol gallstone disease are under discussion. The natural history of the disease and the analysis of the overall costs of therapy are also taking into considerations [4]. Classical oral litholysis like ursodeoxycholic acid (UDCA) has been used for the treatment of cholesterol gallstone disease [4]. Recent studies have raised the possibility that cholesterol-lowering agents that inhibit hepatic cholesterol synthesis or intestinal cholesterol absorption, or drugs acting on specific nuclear receptors involved in cholesterol and bile acid homeostasis for treatment of cholesterol gallstone disease. Other natural plant extraction medicines or components are also reported to have the effect of dissolving gallstone and prevent the occurrence of this disease [5].

Hitrechol® is an herbal medicine used for gallstone disease since 1970s and was indicated for the treatment of early-stage cholesterol gallstones as well as solitary and multiple cholesterol gallstones. Hitrechol® contains the purified extract of *G. hederacea*, in which saponins, essential oil, and phenolic compounds (such as flavonoids, tannins, caffeic acid, and chlorogenic acid [6]) served as the major components. The chemical structure of the saponin ursolic acid in Hitrechol® is similar to that of UDCA, one of the compositions of bile acids in mice gallbladder reported to be effective in dissolving gallstones by lysis of the cholesterol crystals [7, 8], and in altering the bile secretion [9]. The saponins and the essential oil could significantly decrease total cholesterol level [10, 11]. Some studies have also implied that the active components of *G. hederacea* have other possible pharmacological activities including anti-inflammatory [12], antispasmodic [13], liver-protective [14], choleretic [9], litholytic [15], antioxidant [16], antimicrobial [17] and antitumor [18]. It can reduce the inflammatory process which caused by mechanical irritation attributable to gallstones in the gallbladder wall and help to relax the muscle cells in the bile ducts, facilitating the flow of bile. The plants extract can also exhibit antioxidative effect on damaged liver cells. Although several clinical studies have shown that extracts of *G. hederacea* can decrease gallbladder complaints associated with gallstone [19] and reduce the average stone area, the underlying action mechanisms of Hitrechol® remain unknown.

The current study was proposed aiming to investigate the effect of Hitrechol® against gallstone formation in developed lithogenic disease model mice and to evaluate its effect on bile flow in rats. In addition to total cholesterol and total phospholipid levels in bile, the morphological changes of gallstone crystals and pathological changes in the liver after Hitrechol® treatment were monitored. To further evaluate the liver protective effect of Hitrechol®, the antioxidant and anti-inflammatory biomarkers of catalase activities, and superoxide dismutase, reduced glutathione levels and TNF-alpha were also determined and compared among different treatment groups.

Materials and methods

Quality control and HPLC fingerprint of Hitrechol®

Hitrechol Aihuo Dantong (Hitrechol®) Capsule were sponsored by Pharmazeutische Fabrik Evers GmbH & Co. KG (Batch No. 25003). The major component of Hitrechol® is the extract of *G. hederacea*, which is cultivated in Central Europe, growing in shady places. It is a perennial hairy herb with creeping stem and commonly known as “ground ivy” [17]. Authentication of *G. hederacea* was conducted by expert in botany from Albrecht-von-Haller-Institute for Plant Science, the Georg-August-University Göttingen. Manufacturing of Hitrechol® by Pharmazeutische Fabrik Evers GmbH & Co. KG company, complied with all relevant regulations. Briefly, to prepare the extract, the dry herb was cut into a defined size of 1 cm followed by extraction with analytical pure alcohol (90% v/v) at room temperature (25° C) and subsequent concentration under vacuum. The concentrated extract was then mixed with sunflower oil to form into a homogenized mixture to be encapsulated into a soft gelatin capsule.

Besides using macroscopic and microscopic identity tests, the High-Performance Liquid Chromatography (HPLC) fingerprint indicated the existence of three marker components namely ursolic acid, oleanolic acid and linolenic acid in Hitrechol® (Fig. S1). Moreover, content of ursolic acid, one of the major active marker components, was determined by HPLC with UV detector and listed in the specification of Hitrechol® (Table S1). Characteristics, identity, purity and content assay were listed in Hitrechol® specification of the batch (No.
25003) (Table S1), in which includes appearance, filling variation, disintegration time, TLC fingerprint of active ingredient, purity, microbial limit, and the contents of three active marker components. Test results of the batch comply with all the specifications with sum of ursolic acid and oleanolic acid of 40–150 μg per capsule and linolenic acid of 70–350 μg per capsule.

**Chemicals and reagents**

Hitrechol Aihuo Dantong (Hitrechol) Capsule were sponsored by Pharmazeutische Fabrik Evers GmbH & Co. KG (Batch No. 25003). Lithogenic diet was purchased from Teklad Diets EVIGO USA (#TD.88051, Cocoa Butter Diet and Purina Mouse Chow). PEG 400 was purchased from Sigma-Aldrich. Assay kit for Cholesterol/Cholesteryl Ester Quantiﬁcation Assay kit (ab65359), Phospholipid Assay Kit (ab234050), Catalase Activity Assay Kit (ab83464), Superoxide Dismutase Activity Assay kit (ab65354), GSH/GSSG Ratio Detection Assay kit (ab138881) and Mouse TNF alpha ELISA Kit (ab208348) were purchased from Abcam, UK. All other reagents were purchased from Sigma-Aldrich.

**Evaluation of the effect of Hitrechol® on bile secretion flow rate in rats**

The experiments were conducted after approval by the Animal Ethics Committee of The Chinese University of Hong Kong (Ref No. 20–007-MIS-4-B) and complied with the guidelines of the Animal Care and Utilization Committee in Hong Kong SAR. Sprague-Dawley rats with 12/12 h light/dark cycles with free access to food and water. The mice were randomly divided into five groups ($n = 6$ / group). Group 1 mice orally received Hitrechol® capsule content (oily form) was dispersed in aqueous solution consisting of 50% PEG400 (1:10 v/v) to serve as the preparation for oral administration.

**Animal treatment**

The animal care and experimental procedures complied with the guidelines of the Animal Care and Utilization Committee in Hong Kong. The experiments were carried out after approval by the Animal Ethics Committee of The Chinese University of Hong Kong (Ref No. 20–007-MIS-4-B). 27 healthy male C57BL/6 mice of body weight about 22 g were kept in a controlled environment with 12/12 h light/dark cycles with free access to food and water. The mice were randomly divided into five groups ($n = 6$ / group). Group 1 mice orally received the same volume of 50% PEG400 in H$_2$O as vehicle control for 3 weeks. Group 3 mice received once daily Hitrechol® at 2.4 mL/kg (assuming the density of Hitrechol® is 1 mg/mL: equal to 2.4 g/kg in mice and 1.2 g/60 kg/day in human). Mice from Groups 2, 4 & 5 were freely access to a lithogenic diet containing approximately 15.8% fat, 1.25% cholesterol, 0.5% sodium cholate, and 7.5% casein for 3 weeks. During the feeding period of the lithogenic diet, mice were orally administrated with Hitrechol® preparation at once daily (QD, Group 4) or three times daily (TID, Group 5) and 50% PEG400 in H$_2$O as vehicle control (Group 2) for 3 weeks. The dosing regimen for Group 1 to Group 5 mice was summarized in Fig. 1.

**Sampling and sample treatment**

At the end of 3-week treatment, mice from Groups 1 to 5 were sacrificed after overnight fasting followed by collecting their liver, gallbladder and bile. The sampling schedule for Group 1 to Group 5 mice was demonstrated in Fig. 1. Briefly, both body weight and liver weight of each mouse was recorded for calculation of Organ Index (Liver weight /Body weight). About 1 g of

**Comparisons of the Relative Bile Flow between control and Hitrechol® treatment groups were conducted using multiple t-test with a $p < 0.05$ considered as statistically significant.**

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the liver from each mouse was minced and then homogenized using ultrasonic probe for 30 s on ice in 2 mL/g phosphate buffer saline for analyses of bile acid composition and antioxidative biomarkers including catalase, superoxide dismutase and reduced glutathione. The collected whole gallbladders were used to detect the bile acid composition, while the collected bile was used to determine the total cholesterol and phospholipid levels. For mice from Groups 1–4, blood was also collected via cardiac puncture followed by obtaining the serum after centrifugation at 8000 g for 3 min for preliminary analysis of antioxidative biomarkers including catalase, superoxide dismutase and reduced glutathione.

In addition, bile from Group 2 and Group 5 were immediately examined under a polarized microscope for cholesterol crystals morphology observation. Also, about 0.2 g of the liver from each mouse in Group 2 and Group 5 was immediately fixed in 4% pre-cooled paraformaldehyde for further histology treatment and observation. The median lobe of each mice livers from the above treatment groups were isolated and divided into two parts for Hematoxylin and Eosin (H&E) staining and Oil Red O staining, respectively. The first part of the median lobe was fixed in 4% paraformaldehyde at 4 °C until with further dehydration and embedding in paraffin wax. After being sectioned at 5 μm thickness and deparaffinized with xylene and graded alcohols, the mounted tissue sections were then stained with H&E for visualizing with 200 times magnification by a microscope (Eclipse Ti-E, Nikon, Japan) equipped with a digital video camera (DS-Qi2, Nikon, Japan) controlled by the NIS-Elements Imaging software program. Inflammation, necrosis, steatosis, cholestasis, and hemorrhage were major aspects during pathological observation [20]. The second part of the median lobe was stored at −80 °C until the frozen section (6 μm) was obtained. Red Oil O staining was conducted to dye the lipids in liver section to red color. Two hundred times magnification was utilized for observation using the same microscope and imaging system mentioned for Red Oil O staining observation [20].

**Evaluation of anti-lithogenic effect of Hitrechol®**

**Analyses of bile composition** In order to evaluate the effect of Hitrechol® on the composition of bile, levels of the total cholesterol, total phospholipids (0.2–0.5 μL) collected from Groups 1–5 were measured using Cholesterol/Cholesteryl Ester Quantitation Assay kit (ab65359) and Phospholipid Assay Kit (ab234050) according to protocols provided by the manufacturer (Abcam, USA) as described before [21, 22].

**Analyses of bile acid composition in gallbladder and liver** To evaluate the effect of Hitrechol® on altering the bile acid composition, levels of the 12 bile acids in gallbladder and liver of mice from Groups 1–5 were measured and compared. Briefly, about 1 ml of 70%
Acetonitrile was added to the collected whole gallbladder organ followed by centrifugation at 13,000 g for 10 min before LC/MS/MS analyses of the bile acids levels. In addition, two volumes of acetonitrile were added to the prepared liver homogenate of mice from Groups 1–5 followed by centrifugation at 13,000 g for 10 min and vacuum drying of the supernatant. The obtained residue was then reconstituted in 70% acetonitrile followed by LC/MS/MS analyses of the bile acid levels in liver.

A previously reported LC/MS/MS method was adopted with modifications for simultaneous determination of 12 bile acids for the mentioned samples [23]. Briefly, the LC/MS/MS system consisted of an Agilent 6430 triple quadrupole mass spectrometer with an electrospray ionization source (ESI), Agilent 1290 pump and auto-sampler (Agilent Technologies Inc., USA). Chromatographic separation of the four analytes were achieved on a Welch Materials Ultimate HPLC XB-C18 (2.1 × 100 mm, 3 μm) analytical column. The mobile phase of water containing 7.5 mM ammonium formate adjusted with NaOH to pH = 7.0 (A) and methanol (B) were used with a gradient elution from 40% solvent B to 90% solvent B within 22 min. The flow rate was set at 0.3 ml/min. The injection volume of each sample was 20 μl. All bile acids were detected under the negative ionization mode with the following mass spectrometer source settings: ion spray voltage of −4000 V; source temperature of 600 °C, collision gas pressure set at high; Q1/Q3 resolution set at high and interface heater turned on. The multiple reaction monitoring (MRM) transitions for each analyte as well as their respective optimum parameters, such as precursor ion, product ion, fragment voltage (Frag) and collision energy (CE) were listed in Table 1.

Data analyses
All data reported in this study were expressed as mean value with standard deviation except for Organ Index, total cholesterol, total phospholipid and total bile acid levels, where mean value with standard error was adopted. The obtained levels of biomarkers and bile composition were compared among different treatment groups using one-way ANOVA tests followed by Tukey’s multiple comparisons with \( p < 0.05 \) considered as statistically significant.

Results
Effect of Hitrechol® on bile secretion flow rate in rats
As shown in Fig. 2, compared with the control group, slight elevation of bile flow in Hitrechol® treated group was observed at 105 min and 120 min post dosing. However, there is no statistically significant difference between the two groups after a multiple t-test conducted for each time point.

Effect of Hitrechol® against the development of gallstone in normal and cholesterol gallstone model mice
Comparison of liver organ index from different treatment groups
As demonstrated in Fig. 3 (a), liver index of mice in all lithogenic diet groups were significantly increased compared to that of mice in normal diet groups. Among the three lithogenic groups (Groups 2, 4 & 5), it was found that Hitrechol® TID treated group (Group 5) showed a trend of decrease in liver index compared to the vehicle treatment group (Group 2), however there was no statistically significant difference.

| Bile acids            | Precursor Ion | Product Ion | Frag (V) | CE (V) | Polarity |
|-----------------------|---------------|-------------|----------|--------|----------|
| Glycin-conjugated CA  | 464           | 74          | 100      | 70     | Negative |
| Glycin-conjugated CDCA/UDCA/DCA | 448       | 74          | 220      | 70     | Negative |
| Taurin-conjugated CA  | 514           | 80          | 320      | 90     | Negative |
| Taurin-conjugated CDCA/UDCA/DCA | 498       | 80          | 125      | 80     | Negative |
| Unconjugated CA       | 407           | 407         | 300      | 5      | Negative |
| Unconjugated CDCA/UDCA/DCA | 391       | 391         | 280      | 5      | Negative |

CA Cholic acid; CDCA Chenodeoxycholic acid; UDCA Ursodeoxycholic acid; DCA Deoxycholic acid
Effect of Hitrechol® on Total cholesterol and phospholipid levels in bile

As shown in Fig. 3 (b), the total cholesterol levels in vehicle (Group 2) and Hitrechol® (QD) treated lithogenic diet groups (Group 4) were significantly elevated in comparison to that from normal diet groups (Groups 1 and 3). Although there was no significant difference in total cholesterol level between vehicle treated lithogenic diet group (Group 2) and Hitrechol® (QD) treated lithogenic diet group (Group 4), Hitrechol® TID treatment (Group 5) significantly decreased total cholesterol level in comparison to both vehicle (Group 2) and Hitrechol® (QD) (Group 4) treated lithogenic diet groups.

Figure 3 (c) demonstrated the comparison of total phospholipid among different treatment groups. It was noticed that the total phospholipid levels in vehicle and Hitrechol® (QD) treated lithogenic diet group (Group 4) were significantly elevated in comparison to that from normal diet groups (Groups 1 and 3). A trend of decrease in total phospholipid level was noticed in Hitrechol® TID (Group 5) in comparison to vehicle treated lithogenic diet group (Group 2) and Hitrechol® (QD) treated Lithogenic diet group (Group 4), however, there was no statistically significant difference.

Gallstone morphological and liver pathological observations after TID Hitrechol® treatment

Since the Hitrechol® TID treatment demonstrated significant improvements in antioxidative biomarkers and bile composition, we further observed their morphological and pathological changes. As shown Fig. 4 (a), more gallstone crystals were noticed in vehicle treated lithogenic diet mice than that of Hitrechol® TID treatment under 4 times magnification microscopy. Also, the morphology of crystals in vehicle treated lithogenic diet mice was found more rigid and larger under 40 times magnification microscopy. In addition, in Fig. 4 (b), despite observed severe hepatocyte necrosis in vehicle treated lithogenic diet group, minimal necrosis was found in TID Hitrechol® treated lithogenic diet group. Moreover, as demonstrated in Fig. 4 (c), oil droplets were in dark red color, prominent and widespread in vehicle treated lithogenic diet liver tissue. However, oil droplets in Hitrechol® TID treated lithogenic diet liver tissue were observed in lighter red color, smaller size compared with vehicle treated lithogenic diet one which indicated that Hitrechol® TID treatment could notably decrease cell necrosis chances and lipid aggregation in the liver.

Effect of Hitrechol® on bile acid compositions in gallbladder of lithogenic treated groups

As shown in Fig. 5 (a), among the three lithogenic diet groups, significant decrease in the amount of GCA, TCA and GCDC/GDCA/GUDCA in the whole gallbladder were found after TID but not QD Hitrechol® treatment. For the total bile acid level in the three lithogenic diet groups (Groups 2, 4 & 5), as demonstrated in Fig. 5 (b), significant reduction in total bile acid level was found in Hitrechol® (TID) group in comparison to that in vehicle treated group. Although Hitrechol® QD treatment group showed a trend of decrease compared to vehicle treated group, no statistical difference was found.
Anti-oxidative and anti-inflammatory effects of Hitrechol®

Since our preliminary evaluation found no significant difference in the serum level of CAT, SOD and GSH among the treatment groups of Groups 1, 2, 3 & 5, our subsequent comparison focused on their levels in the liver. As indicated in Fig. 6 (a), all the lithogenic diet groups (Groups 2, 4 & 5) had significantly decreased Catalase (CAT) activities level when compared with normal diet groups (Groups 1 and 3). In addition, it was also noticed that both QD and TID Hitrechol® (Groups 4 and 5) treatments significantly elevated the liver CAT activities level. As indicated in Fig. 6 (b), all the lithogenic diet groups (Groups 2, 4 & 5) exhibited significantly decrease in Superoxide dismutase (SOD) inhibition rates when compared with that from normal diet groups (Groups 1 and 3). It was also noticed that significant increase in SOD level in TID but not QD Hitrechol® treated group in comparison to vehicle treated lithogenic diet group. In Fig. 6 (c), both vehicle-treated lithogenic diet group (Group 2) and QD Hitrechol® treated normal diet group (Group 4) were found decrease in Reduced glutathione (GSH) levels compared to vehicle treated normal diet group (Group 1). In comparison to vehicle treated lithogenic group (Group 2), significant increase in Reduced GSH after Hitrechol® (QD) and Hitrechol® (TID) treatments, but no significant difference was found between QD and TID Hitrechol® treatment.

Discussions

Gallstone formation is caused by the precipitation of bile components formed within the gallbladder with cholesterol precipitations as the most common one. Although Hitrechol® is used for the treatment of cholesterol gallstone disease since 1970s, the underlying mechanisms of actions of Hitrechol® for cholesterol gallstone treatment are still not clear. In this study, we investigated the effect of Hitrechol® on the bile flow in normal rats and compositions of bile alteration after Hitrechol® treatment in gallbladder in normal and cholesterol gallstone diseased mice. The morphology changes of gallstone and pathology changes in liver tissues cholesterol gallstone diseased mice provided more insights for the actions of Hitrechol® and specific antioxidative and anti-inflammatory biomarkers were measured to understand the hepatoprotective effect of Hitrechol®.

Based on the previous report on bile secretion increase by ursolic acid in normal rats [9], it was expected that Hitrechol® with ursolic acid as a major component might also be able to increase the bile secretion in liver via enhancing the bile flow. However, in the current experiment, despite a trend of slight elevation of bile flow for each time interval in Hitrechol® treated group observed at 105 min and 120 min postdosing, no statistical
significance was found in comparison to that from the control group in normal rats. Such discrepancy may be due to the different treatment doses of ursolic acid. In the above-mentioned study [9], the lowest dose of ursolic acid was 7.5 mg/kg, whereas a much lower dose of ursolic acid (120 mg/kg Hitrecol® containing about 0.096 mg ursolic acid) was given to the rats in our current study.

Based on the indication of Hitrechol®, gallstone mice model was adopted for the current study for the evaluation of its impact on the gallstone formation. The successful development of the gallstone diseased mice model was verified by the gallstone formation shown in Fig. 4 (a) after mice were sacrificed in lithogenic diet treated groups. Obvious differences between mice treated with normal diet and treated with lithogenic diet were noticed in the biomarkers such as total cholesterol level and total phospholipid level [23]. In the normal diet groups, Hitrechol® treatment had minimal impact on the bile compositions such as the total cholesterol level, total phospholipid level and the total bile acid level with no significant changes in liver biomarkers including CAT,
SOD and reduced-GSH levels, indicating the safe use of Hitrechol® with no severe side effects. Therefore, we focused on the subsequent comparisons on bile composition changes and biomarker changes in liver among the three lithogenic diseased groups. It was noted that the organ index for liver was all significantly...
increased in the three lithogenic groups, further suggesting the existence of liver injury. Moreover, in comparisons to the vehicle treated lithogenic group, the anti-lithogenic and liver protection effects of Hitrechol® both QD and TID treated groups could be significantly noticed in bile compositions and relevant biomarkers. For Hitrechol® QD treatment, it could potentially exhibit liver protection effect by increasing the CAT activity and reduced-GSH content in liver. Although no statistically significant decrease in total cholesterol and no significant increase in phospholipid levels after Hitrechol® QD treatment were noted, the gallstones were found in general to be smaller in size comparing to that in the vehicle treated group. After increasing the dosing frequency of Hitrechol® to TID, more obvious protective effect against the gallstone formation was noted. In addition, as demonstrated in Fig. 4 (b) and Fig. 4 (c) of liver histology samples, Hitrechol® TID treatment could notably decrease cell necrosis chances and lipid aggregation in liver caused by high fat and high cholesterol lithogenic diet, which may be contributed by oleanolic acid (70–350 μg/0.2 g capsule, 0.035–0.175% in Hitrechol®), a hepatoprotective drug for over 20 years in China [6, 24]. Besides the liver histology observations, we also noticed that the anti-oxidative effect in liver indicated by changes in CAT, SOD and Reduced GSH levels were further improved in Hitrechol® TID treatment group compared to Hitrechol® QD and vehicle treatment groups due to the liver protection effects of Hitrechol®. The anti-lithogenic effect of Hitrechol® could be reflected by the significant reduction of total cholesterol levels and slightly decreased phospholipid levels in the liver in Hitrechol® TID treatment group. As we mentioned previously [2], in normal physiological condition, bile salts, phospholipids, cholesterol, proteins, and bilirubin were all in stable equilibrium states. The decreased cholesterol level together with the increased phospholipid-to-bile acid ratio after Hitrechol® treatment is considered to lower the chances of cholesterol precipitation and subsequent gallstone formation. A previous study reported that higher ratio between phospholipid and bile acid could help to increase the solubility of cholesterol [25]. The mean value of phospholipid-to-bile acid ratio in vehicle treated groups was 4.35, which was increased to 4.91 after Hitrechol® QD treatment and significantly increased to 6.32 after Hitrechol® TID treatment. Such increase ratio between phospholipid-to-bile acid after Hitrechol® treatment could help to increase the solubility of cholesterol in bile and prevent the formation of cholesterol crystals. In addition, the decreased cholesterol level might also be contributed by linoleic acid (40–150 μg/0.2 g capsule, 0.02–0.075% in Hitrechol®), which was an essential fatty acid belonging to the omega-3 fatty acids group. It was reported that linoleic acid could lower the total cholesterol, low-density lipoprotein (LDL) cholesterol and high-density lipoprotein (HDL) cholesterol levels significantly [26, 27]. Normally, bile secretion was promoted to dissolve or lysis the gallstone when taking a high

![Fig. 6](image_url) Comparisons of Catalase (CAT) activities (a), Superoxide dismutase (SOD) inhibition rates (b) and Reduced glutathione (GSH) levels (c) in liver from different treatment groups for evaluating the anti-oxidative effect of hitrechol®,. $n = 4$ to 6 for each treatment group. Data presented in Mean ± SD, $^{\wedge}p < 0.05$ compared to VEH + ND group, $^{#}p < 0.05$ compared to VEH + LGD group analyzed by one-way ANOVA followed by Tukey’s multiple comparisons test. VEH: Vehicle; ND: Normal diet; LGD: Lithogenic diet; H (QD): Hitrechol® once daily; H (TID): Hitrechol® three times daily. VEH + ND: Group 1; VEH + LGD: Group 2; H (QD) + ND: Group 3; H (QD) + LGD: Group 4; H (TID) + LGD: Group 5

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cholesterol diet. As expected, the total bile acid level in all the three lithogenic diet treated groups was noticed significantly increased compared to that of normal diet treated groups. In this study, after Hitrechol® TID treatment, the frequency of occurrence was significantly lowered with altered morphology of gallstone. Also, the levels of GCA, TCA and GCDCA/GDCA/GUDCA and total bile acid in gallbladder significantly decreased when compared to that from vehicle-treated group. It was worth noting that the UDCA (ursodeoxycholic acid) amount in bile acid composition showed a trend of increase after Hitrechol® TID treatment, which could be due to the presence of ursolic acid in Hitrechol®. The chemical structure of ursolic acid (40–150 μg/0.2 g capsule, 0.02–0.075% in Hitrechol®) is similar to that of ursodeoxycholic acid (UDCA), one of the endogenous bile acids that had been reported to be effective in dissolving gallstones via lysis of the cholesterol crystals [8].

Conclusions
Hitrechol® dose-dependently demonstrated the anti-lithogenic effect and exhibited obvious liver protection effect through elevating the antioxidant biomarkers in cholesterol gallstone disease mice. Among the tested dosing regimens, Hitrechol® TID could effectively decrease the occurrence of gallstone formation and enhance the liver protection effect via altering the bile composition, elevating the antioxidative biomarkers and inhibiting IFN-gamma release.

Abbreviations
CA: Cholic acid; CDCA: Chenodeoxycholic acid; DCA: Deoxycholic acid; GSH: Reduced glutathione; H: Hitrechol®; LGD: Lithogenic diet; LC/MS/MS: Liquid chromatography with tandem mass spectrometry; N: Normal diet; OD: Once daily; SOD: Superoxide dismutase; TID: Three times daily; UDCA: Ursodeoxycholic acid

Supplementary Information
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Additional file 1: Fig. S1. HPLC Chromatogram of the standard mixture of the major components in Hitrechol (a) and Hitrechol® capsules (b).
Table S1: Certificate of Analysis of Hitrechol® (Batch No. 25003).

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Not applicable.

Authors’ contributions
MX conducted mouse experiments, data collection, analysis and interpretation of the data and drafting the manuscript; MBY conception and design of the work and conducted rat experiments; XYJ helped to analysis the histology samples; YNA, DL and YFL did the animal dosing and sample treatment; ZZ designed the experiment and provided critical revision of the manuscript. The author(s) read and approved the final manuscript.

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Availability of data and materials
All data generated during this study are included in this article. The fingerprint profile of the major components in Hitrechol® and its certificate of analyses are provided as supplementary information Fig. S1 and Table S1, respectively.

Declarations

Ethics approval and consent to participate
The experiments were carried out after approval by the Animal Ethics Committee of The Chinese University of Hong Kong (Ref No. 20-007-MIS-4-B) which in compliance with both the guidelines of the Animal Care and Utilization Committee in Hong Kong SAR and the guidelines 2.0 of the ARRIVE (28). Manufacturing of the Hitrechol® complied with all relevant regulations.

Consent for publication
Not applicable.

Competing interests
There are no conflicts of interest among the authors.

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