Chemical bile duct embolization for chemical hepatectomy, long term efficacy and feasibility in rats

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ABSTRACT

Background: Hepatolithiasis is the presence of calculi within the intrahepatic bile ducts. It represents a significant problem in hepatobiliary surgery because of its high recurrence rate and the associated intra-operative and post-operative risks. This study was designed to explore the long-term efficacy of chemical biliary duct embolization (CBDE) to treat recurrent hepatolithiasis and to explain the mechanism of CBDE. Aims and Objectives: To investigate the long term efficacy of chemical bile duct embolization (CBDE) on the targeted hepatic lobe and to explain the mechanism of CBDE to achieve chemical hepatectomy. Materials and Methods: The median biliary duct of rats were embolized with phenol and N-butyl-cyanoacrylate. The short-term (6 weeks) and long-term (12 weeks) effects of chemical bile duct embolization were compared by observing the degree of atrophy, fibrosis, and proliferation of collagen fibers and apoptosis of hepatocytes of the embolized hepatic lobe. The feasibility and effectiveness of chemical hepatectomy were analyzed by histology, Western blot analysis of collagen I fibers and assessment of hydroxyproline content. Results: After 6 weeks of the procedure, destruction of hepatocytes, fibrosis and "self-cut" was seen only in the periphery of the targeted hepatic lobe. Whereas after 12 weeks, complete destruction of hepatocytes, and replacement with proliferative bile ductules and collagen fibers leading to complete fibrosis and "self-cut" phenomenon in the whole targeted hepatic lobe was seen. Collagen I expression in the 6-weeks treatment group and 12-weeks treatment groups were 4 times and 12 times higher than that in the Sham operated (SO) group respectively (P<0.05). In addition, there was increase in hepatic hydroxyproline content (HYP) approximately by sevenfold in 6-weeks treatment group and twentyfold in 12-weeks treatment group after CBDE, when compared to that in the SO group (P<0.05). Conclusion: Chemical bile duct embolization can achieve ideal effect of chemical hepatectomy in the whole targeted lobe.

Key words: Hepatolithiasis, Bile duct, Chemical embolization, Chemical hepatectomy

INTRODUCTION

Hepatolithiasis is commonly encountered in the Asia-Pacific regions, but is considered as a particularly intractable disease because of its recalcitrant nature.1,2 Preventing stone recurrence and biliary re-stenosis after successful removal of intrahepatic stones still remains an unachieved goal in hepatobiliary surgery.3,4 Patients who requires third or fourth biliary surgery for the stone recurrence along with severe biliary stricture, are commonly encountered in East Asia. In fact, approximately two-thirds of patients with hepatolithiasis in Asia require follow-up surgery due to recurrent biliary stricture, bile duct infection, and other refractory factors. The degree of biliary stricture is directly proportional to the frequency of stone recurrence.5 This is further associated with gradual worsening of the
MATERIALS AND METHODS

Experimental program
All animal experiments were approved by the Animal Care and Use Committee of Sichuan University (Chengdu, China). Twenty-six male Sprague-Dawley rats, weighing 220-250 gm provided by Experimental Animal Centre of Sichuan University (Chengdu, China), were randomly divided into three groups depending on their treatment group and time of tissue collection:
(a) Tissue collected 6 weeks after embolization (6-weeks treatment group, n = 10): The median biliary duct was ligated and was injected with 0.15 ml of phenol through a T_{60}G polyethylene catheter (Abbott, Co., Chicago, IL, USA), and aspirated after 15 minutes. Subsequently, 0.15 ml of N-butyl-cyanonacrylate (Tissue adhesive glue, Jingya Co., Beijing, China) was injected quickly with pressure. Rats were sacrificed at the sixth week. Embolized lobe and non-embolized lobe were resected out and preserved for further analysis.
(b) Tissue collected 12 weeks after embolization (12-weeks treatment group, n = 10): The median biliary duct was embolized as above. After retrograde cholangiography, rats were sacrificed at the twelfth week after the model establishment. Embolized lobe and non-embolized lobe were resected out and preserved for further analysis.
(c) Sham operation (SO group, n = 6): The median biliary duct was dissected out only.

Morphological and histological analyses
After all rats were sacrificed on 6th week and 12th week respectively, the liver samples were taken and fixed in liquid nitrogen and 10% formaldehyde for further tests. Five micrometer thick formalin-fixed liver tissue slides were prepared for HE staining. The percentage of the fibrosis area was evaluated by Masson staining.

Western blot analysis of collagen I
Hundred micrograms of extracted protein were separated by sodium dodecyl sulfate polyacrylamide gel electrophoresis, and then electrotheretically transferred to polyvinylidenedifluoride membrane and hybridized. The membrane was blocked with 5% non-fat dry milk for 1 h then incubated with collagen I primary antibody (Zymed Co., South San Francisco, California, USA) overnight at 4°C. After rinsing with Tris-buffered saline and Tween 20, the membrane was incubated for 2 h with horseradish peroxidase-conjugated secondary antibody. Immunoreactive bands were visualized with enhanced chemi-luminescence and captured on X-ray film. The results were expressed as the ratio of the expression of target gene to that of β-actin.

Assessment of hydroxyproline content (mg/g liver)
Collagen content was quantitatively determined by hydroxyproline, an amino acid found primarily in collagen and the principal component of the extracellular matrix.
The hydroxyproline content in liver was measured as per the instruction provided by Jamall et al.\textsuperscript{16,17}

**Statistical analysis**

Data analyses were made using SPSS 10.0 software (SPSS Inc., Chicago, and IL, USA). Quantitative data were expressed as mean and standard deviation. Statistical comparisons among means were performed using one-way analysis of variance (ANOVA), and Fisher’s Least Significant Difference analysis was used for within-group comparisons. $P<0.05$ was considered statistically significant.

**RESULTS**

**Histopathological examination**

In the 6-weeks treatment group, the hepatocytes in the peripheral region were almost completely destroyed and replaced with proliferative bile ductules and collagen fibers leaving large islands of residual liver cells in the center portion of the targeted hepatic lobe (Figure 1a). Whereas, in 12-weeks treatment group, the hepatocytes disappeared completely and were replaced by proliferative ductules and collagen fibers leading to complete fibrosis and “self-cut” phenomenon in almost the whole embolized lobe (Figure 1b). Fibrosis levels in both treatment groups were expressed using Masson staining. Blue stained collagen fibers portions were significantly higher in 12-weeks treatment group (Figure 1d) in comparison to that of 6-weeks treatment group (Figure 1c). No liver abscess and alteration in the liver functions were seen during the entire experiment.

**Western blot analysis of collagen I**

Collagen I protein expression in the 6-weeks treatment group was 4 times higher than that in the SO group. Whereas, collagen I expression in the 12-weeks treatment group was 12 times higher than that in the SO group, which was 3 times higher than that in the 6-weeks treatment group (Figure 2) ($P<0.05$).

**Assessment of hepatic hydroxyproline content**

Quantitative determination of hepatic collagen content was performed by assessing the hepatic hydroxyproline (HYP) content after CBDE. There was an approximate sevenfold increase in 6-weeks treatment group and a twentyfold increase in 12-weeks treatment group, when compared to that in the SO group. Additionally, the differences between the 6-weeks treatment group and 12-weeks treatment group were statistically significant ($P<0.05$) (Figure 3).

**DISCUSSION**

Hepatolithiasis is a commonly encountered disease in the Asia-Pacific region, but is still regarded intractable as the long term effects of current available therapeutic approaches are far from satisfactory.\textsuperscript{1,18} Calculi in secondary or tertiary subsidiary branches of bile duct are usually complicated with biliary duct stenosis. Therefore, even after successful stone clearance with choledochoscopy, stone recurrence is inevitable. Biliary stricture causes bile stasis and secondary infections which then stimulates hyperplasia of bile duct thus causing chronic proliferative cholangitis (CPC). Repeated episodes of CPC leads to stone formation, thus forming a vicious cycle of CPC and stone formation.\textsuperscript{2,19} Therefore, treatment of hepatolithiasis must include clearance of stones, correction of biliary stricture and elimination of CPC. Presently, traditional hepatectomy remains the only treatment of choice for hepatolithiasis. Unfortunately, 40% of hepatolithiasis patients have calculi distributed throughout the liver and most patients cannot tolerate multiple lobes/segments resection. Due to these reasons, application of hepatectomy to treat hepatolithiasis is greatly restricted. In previous study, we explored the feasibility and efficacy of filling the diseased bile duct with chemical embolization agent to achieve chemical resection of the diseased bile duct and hepatic segment, and was able to establish the effects of chemical hepatectomy only in the periphery of the targeted hepatic lobe.\textsuperscript{11-15} This study focuses on whether CBDE in the long-term manages to achieve complete fibrosis, “self-cut” and chemical hepatectomy in the whole embolized hepatic lobe.

In both the 6-weeks and 12 weeks treatment groups, chemical hepatectomy resulted from CBDE exhibited destruction of the biliary duct mucosa and replacement with collagen fibers of the targeted hepatic lobe. Thus, completely eradicating the pathological basis of stone recurrence: chronic proliferative cholangitis (CPC). In the 6-week treatment group, islands of residual hepatocytes were still visible, with signs of
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Chemical hepatectomy only observed in the periphery of the embolized lobe. Whereas in the 12-week treatment group, destruction of hepatocytes, fibrosis and “self-cut” was seen in the whole embolized hepatic lobe. Masson staining revealed collagen fibers were seen in almost the entire embolized liver lobe. This was confirmed quantitatively by western blot analysis of the collagen fiber. The expression level of collagen I at 6 and 12 weeks after CBDE was 4 and 12 times higher than the corresponding levels in the SO group respectively. In addition, the hydroxyproline content in 6-weeks treatment group and 12-weeks treatment group were 7-times and 20-times higher than that in the SO group respectively. The difference of hydroxyproline content in the 6-weeks treatment group and 12-weeks treatment group was statistically significant (P<0.05).

Feasibility of embolization of small bile ducts to treat hepatolithiasis have been confirmed in the past. Patients with multiple biliary calculi with biliary stricture in the inferior branch of the left lateral bile duct were treated with CBDE. Fifteen months later, computed tomography scans demonstrated that the inferior segment of the left lateral lobe had almost completely atrophied and disappeared, with no signs of either hepatic abscess or stone recurrence.

Liver abscess is rarely encountered in totally atrophic lobe. Liver abscess may occur secondary to stone obstruction or incomplete embolization of the first or second subsidiary bile duct. If the bigger bile duct is targeted, occurrence of liver abscesses is likely due to incomplete embolization of the targeted bile duct. Selection of an appropriate embolization agent that can complete embolization hepatic ducts of various sizes is the most important factor for successful chemical hepatectomy via CBDE.

Fu-Yu Li, Jing-Qiu Cheng et al suggested that phenol is not only the most efficient sclerosants of gall bladder mucosa, but is also a powerful antiseptic which might not only assist in ameliorating biliary infection which commonly accompanies hepatolithiasis, but also could commendably destroy the biliary mucosa and induced fibrous conglutination in the bile duct lumen, leading to partial occlusion of the large-size and middle-size biliary ducts, and complete occlusion of the small biliary ducts. However, the biliary system could not be permanently occluded by using phenol or ethanol alone due to the regeneration of biliary mucosa. As an ideal liquid occlusion agent; cyanoacrylate, histoacryl or neoprene has such remarkable liquid-dynamic characteristics that could perfectly fill bile ducts of various sizes, including the acroteric branch. Thus, leading to complete disappearance...
of mucosal epithelium which are entirely replaced by foreign body granuloma and collagen fibers. Additionally, cyanoacrylate occupies the space where stones can occur, thus preventing intrahepatic stone recurrences. All of these suggests that mucosal sclerosant plus tissue adhesive glue results in a more complete and reliable embolization than a mucosal sclerosant alone. In addition, histoacyrl (also called N-butyl cyanoacrylate), a derivative of cyanoacrylate can be dissolved in contrast medium and thus can be visualized under x-ray. This kind of visibility makes CBDE much safer, easier and more accurate in its application.\textsuperscript{5,11,12,15,23,25}

The mechanism underlying chemical hepatectomy via CBDE may involve the following aspects. The embolized bile duct obviously distends and compresses the adjacent blood vessels, thus reducing the blood supply to the embolized hepatic lobe.\textsuperscript{15,26,28} Reduction in blood supply to the embolized lobe and deposition of large amount of intrahepatic bile salts due to biliary obstruction activates hepatocellular apoptosis via upregulation of Fas and Bax, which then activates capsase-3 and capsase-9. Fas- triggered apoptosis activates hepatic stellate cells (HSC), which secretes TGF-\(\beta\)\textsubscript{1} and TIMP\textsubscript{1} and consequently synthesize large amount of collagen fibers. Additionally, TGF-\(\beta\)\textsubscript{1} and TIMP\textsubscript{1}, can accelerate hepatocellular apoptosis and inhibit hepatocellular regeneration. Thus, a perpetual cycle of hepatocytes death and proliferation of bile ductules and collagen may develop. It is important to note that CBDE not only permanently embolize, destroy the biliary tree and reduce the blood supply of target liver segment, but also compresses the microvasculature of hepatocytes and replaces hepatocytes by large number of proliferated intrahepatic bile ductules and collagen fibers. With the acceleration of hepatic fibrosis, the blood supply of embolized lobe reduces. Liver cells can thus be induced ongoing injury and apoptosis. Therefore, hepatic fibrosis process caused by CBDE may progress continuously until complete hepatic fibrosis and “self-cut” chemical hepatectomy is achieved.\textsuperscript{12,27,29-31}

CBDE for chemical hepatectomy may provide an innovative and less invasive therapeutic approach in treating hepatolithiasis and preventing postoperative recurrences of intrahepatic stones. Chemical hepatectomy of the targeted liver segment is more selective than the traditional hepatectomy, as it can be implicated accurately for subsegmental hepatectomy and preserve significant functional hepatocytes. However, just like chemical cholecystectomy, CBDE is feasible but not recommended routinely. We must emphasize that cholangioscopic lithotomy or open hepatectomy are still the best treatments of choice for routine hepatolithiasis.\textsuperscript{7,23,28} However, most of the late-stage hepatolithiasis patients cannot tolerate multiple hepatic lobe or segment resections, since intrahepatic calculi can cause many pathological lesions to the liver such as secondary biliary cirrhosis, multiple calculi, recurrent cholangitis, repeated operation, etc. In addition, recurrent hepatolithiasis with cholangitis in these late-stage hepatolithiasis patient can lead to development of apical epithelial hyperplasia, which may progress to cholangiocarcinoma. Thus, early eradication of this kind of chronic proliferative inflammation can prevent occurrence of concomitant intrahepatic cholangiocarcinoma. Therefore, application of CBDE in these conditions not only avoids trauma and risks of conventional hepatectomy, but also expands the operative indications for those late-stage hepatolithiasis patients who are at a high risk for hepatectomy.\textsuperscript{5} Furthermore, if patients with multiple calculi combined with a biliary stricture are only treated by endoscopic cholelithotomy, recurrence of the intrahepatic stone is foreseeable. Thus, application of CBDE after cholecdochoscopic lithotomy would certainly reduce re-operations for recurrent hepatolithiasis. However, the targeted biliary duct in these patients should be limited to the third subsidiary bile ducts, which are easy to occlude completely without subsequent development of cholangitis or liver abscess. If the first or the second subsidiary bile duct is chosen as the target biliary duct, one has to be more careful, as there is a greater risk of suppurative cholangitis or liver abscess due to incomplete embolization.\textsuperscript{3,5,8}

In our study, we were able to conclude that CBDE after 12 weeks was able to express fibrosis, proliferation of bile ductules and “self-cut” of the whole embolized lobe thus achieving the ideal effect of chemical hepatectomy. We believe CBDE might be able to partially replace conventional hepatectomy as a more feasible, effective and safe approach to treat recurrent hepatolithiasis. However, relative literatures suggesting the result are very few and are still in its preliminary stage. Thus, further research and analysis about its clinical significance, indications, contraindications, precautionary measures and long-term effectiveness is definitely required prior to its clinical application.

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**Conflict of interest**

All authors declare that they have no conflict of interest.

**Ethical approval**

All animal experiments were approved by the Animal Care and Use Committee of Sichuan University (Chengdu, China).
REFERENCES


Authors Contribution:

AS and YZ - Contributed to the data acquisition and analysis and drafted the manuscript. WJM, HF and QY - Contributed in histology, western blot analysis and data acquisition; WZ, YQZ and CNS - Were involved in the revision of the manuscript; LFY and HM - Contributed to the study design and revision of the manuscript.

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