Clinical study on the sensitivity test guided hepato-arterial/portal-vein chemotherapy in patients with unresectable hepatocellular carcinoma

Dongde Wu¹, Shaozhong Wei²*, Chenggang Luo¹, Xinghong Wu³, Yaojun Feng³, Feng Zhang¹, Lei Nie¹, Xiaqin Xia¹

¹Department of Hepatobiliary and Pancreatic Surgery, Cancer Hospital of Wuhan University, Wuhan, Hubei, China.
²Department of Urology Surgery, Cancer Hospital of Wuhan University, Wuhan, Hubei, China.
³Department of Breast Cancer Surgery, Cancer Hospital of Wuhan University, Wuhan, Hubei, China.

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Original Article

Abstract

Purpose: Approximately 20 % of tumors have the opportunity to be resected in patients with hepatocellular carcinoma (HCC) and their prognoses were acceptable. For the unrespectable HCC, however, the outcomes were rather poor because the speciality of the tumor blood supply and the tumor was insensitive to the chemotherapy drug. The objective of this study was to find sensitive drugs for individual patients and determine the safety and antitumor activity of hepatic intra-arterial and portal vein infusion chemotherapy. Methods: A total of 120 patients with the mean patient age of 56 years and with unresectable HCC were randomly divided into experimental group and control group. The experiment group was infused through an intra-arterial and portal vein catheter 3 different drugs were chosen by the results of drug sensitive test, whereas the control group was treated for TACE. The changes in tumor-size and AFP, two-step operation rate, survival rate, and complications were observed in these patients. Results: The tumor size reduced in 28 (47.6%) cases, stabilized in 14 (23.3%) cases, and progressed in 18 (30%) cases in experimental group as measured by CT or MRI after six chemotherapy cycles, whereas the corresponding data was 17 (28.5%) cases, 7 (11.5%) cases and 36 (60%) cases in the control group (P < 0.05). The AFP was declined in 51 cases in experimental group and in 30 cases in control group (P < 0.05). In the experimental group, the median follow-up time was 21 months; the overall survival rates (OS) of 6 months, 12 months, and 18 months were 86%, 72%, and 65%, respectively. In the control group, the median follow-up time was 16 months; the OS rates of 6 months and 12 months were 58% and 40%, respectively. Six patients in the experimental group and 3 patients in the control group had two-step operation. There was no severe incidence of complications in both groups. In the experimental group, 2 (3%) patients had wound infection, 8 cases had the chemotherapy relative diarrhea, and 18 (30%) cases had grade I or II bone marrow suppression. In the control group, the chemotherapy relative diarrheas were 15 (25%) cases and grade I or II bone marrow suppression were 33 (55%) cases. Conclusion: The artery and portal vein pump transfusion chemotherapy guided by drug sensitive test was efficient for HCC treatment. The patients can get longer OS and lower complication incidence.

Keywords: Primary Liver Cancer; Drug Sensitive Test; Regional Chemotherapy

Introduction

Because the liver tumors are not easily found, only less than 20% of the primary liver cancers have the opportunity to be surgically removed. Treatment of hepatic artery chemotherapy embolism (TACE) is one of the therapeutics available to cure the advanced stage unresectable liver cancer; however, the effective rate is about 30% to 40%. Research showed that primary liver cancers have hepatic artery and portal vein dual blood supplies: blood vessel in the center of the tumor is mainly composed of hepatic artery whereas the peripheral part is mainly composed of portal vein. The cancer cells tend to invade the intrahepatic vascular, especially into the portal system, which promote the portal venous embolism and...
have negative effects of conventional TACE. In order to improve the efficiency to cure primary liver cancer, scholars developed the hepatic artery/portal vein pump chemotherapy (HA/PVPC). However, liver cancer is not achesynonymous to chemotherapy. To improve the efficiency of individual patients to HA/PVPC, we used individualized drug to patients based on their sensitive tests (MTT) and perfused the drugs through transartery and portal vein dual channel. We aimed to increase the regional drug concentration and improve the tumor susceptibility to drugs, thus improving the efficiency of liver cancer treatment.

**Methods and Materials**

**Patients**
Between October, 2010 and October, 2005, 120 cases of unresectable HCC were randomly divided into experimental group and control group. 60 patients in the experimental group were enrolled to determine the safety and antitumor activity of hepatic intra-arterial and portal vein infusion chemotherapy; another 60 patients were enrolled in the control group. The clinical protocol was reviewed and approved by the Institutional Review Board of the Hubei Cancer Hospital. All study participants provided written informed consent prior to therapy. Eligible study participants were chosen according to the following inclusion criteria:

- Liver function test was Child A-B grade;
- Tumor cannot be resected;
- Eastern Cooperative Oncology Group performance status was 0 or 1;
- Total bilirubin concentration was ≤1.0 mg/dL;
- Serum creatinine concentration was ≤1.5 mg/dL;
- Prothrombin time was <13 seconds; activated partial thromboplastin time was < 30 seconds; and
- Complete blood count was normal. All of the patients before treatment did not use any antitumor drugs.

**Preparation of tumor cells**
Sampling: Specimens of primary liver cancer tissue was obtained from surgical biopsy, placed into RPMI 1640 culture medium which contained streptomycin (100 U/ml) and penicillin streptomycin (100 mg/ml) within 10 minutes after shearing, preserved in 4 °C refrigerator, and processed as described below within 4 hours.

Processing samples: Tissue blocks were placed in a petridish, cut off blood clots and unused tissues, rinsed with hanks solution twice, and then cut into 1 mm³ tissue blocks.

Isolating cells: We crushed the small tissue blocks gently by repeatedly passing through a syringe needle. The tissue homogenate was filtered through 4 layers of gauze followed by centrifugation in low speed centrifuge machine (800 rpm, 5 min) twice.

**Cell culture**
Cells were cultured in RPMI 1640 (Gibco, Invitrogen) supplemented with 10% (v/v) heat inactivated fetal bovine serum (Gibco, Invitrogen) and 1% (v/v) antimicotic antibiotic (Gibco, Invitrogen) at 37°C under a 5% CO₂ atmosphere. The cells adhered to the wall of the culture flasks after 48 hours. After 4 days of incubation, the cell concentration was adjusted to (2 to 5) × 10⁵/ml.

**Cell viability assay (MTT assay)**
100 ul of liver cancer cells were seeded on 96-well plates. For the cells in the experimental group, seven chemotherapy drugs (5-FU/ DDP/ ADM/ MMC/ HCPT/ LOHP/ GEM) were individually added to the cells in triplicate at a concentration ten times higher than that used clinically. No drug was added to the cells in control group. 1640 medium was used as a blank control. Cells were cultured for 48 hours. Then the cells were added 20 ml of MTT (5 mg/ml) and continually cultured for 4 hours before replacing culture medium with dimethyl sulfoxide (100 μL/well) to terminate the reaction. The cells were finally vibrated for 5 minutes and the absorbance (value A) was measured by a micro plate reader at a wavelength of 570 nm.

Drugs inhibition rate = [1 - (average A value in drug treatment wells / average A value in negative control wells)]. A rate value > 50% is defined as hypersensitivity; 30% to 50% as mild sensitive; < 30% as drug resistance.

Sensitive rate (%) = (sensitive cases / total cases of the group) × 100%

**Clinical experiment**
All patients in the experimental group underwent angiographic placement of two catheters for hepato-artery and portal vein. We obtained the hepatic tumor biopsy for pathological diagnosis and then The Bard Access Ports (detailed information showed below) were inserted into the right gastro-omental artery and the right gastro-omental vein to establish regional chemotherapy channels. Seven chemotherapy drugs were used as the experimental drugs (detailed information showed below). We infused each individual in the experimental group by arterial and portal vein pump 2 to 3 drugs which showed highest inhibition to the cancers of the patient in our MTT test above. The drugs were delivered equally in both channels. The regimen was repeated every 3 weeks unless there was a tumor progression, withdrawal of consent, or occurrence of side effects, such as bone marrow suppression, fulminant hepatitis, or other greater than grade 3 toxicities, etc.

The control group patients underwent conventional TACE, chemotherapy regimens were 5 - fluorouracil (1000mg/m²),...
mitomycin (10mg/m²) and cisplatin (100mg/m²), and with iodipin embolization (scheme: MMC+DDP+5-Fu, iodipin embolism).

The Bard Access Ports was purchased from West Amelia Earhart Drive, Salt Lake City, Utah, USA. The cytotoxic drugs included fluorouracil (5 FU, Tianjin Jinyao acid co., LTD), cisplatin (DDP, Yunnan Gejiu Supertrack Bio-Pharmaceutical co., LTD), Adriamycin (ADM, Zhejiang Hisun pharmaceutical co., LTD), mitomycin (MMC, Jiangsu Henrui Medicine co., LTD), Hydroxy camptothecin (HCPT, Hubei Huangshi Feiyun co., LTD), oxaliplatin (L-OHP, Jiangsu Henrui Medicine co., LTD) and gemcitabine (GEM, Hubei Halsky pharmacy co., LTD)

Curative effect evaluation
Curative effect evaluation includes changes in serum tumor marker AFP, tumor size and number, liver function, complications, and overall survival (OS).

Follow-up
Local therapeutic efficacy was evaluated by contrast-enhanced dynamic CT scanning after 2 courses of chemotherapy or in the case of clinical suspicion for recurrence. Clinical tumor recurrence and response was assessed according to the Response Evaluation Criteria in Solid Tumors (RECIST version 1.0)^2

Toxicity Assessment
Toxic effects were assessed during hospitalization of each patient and at the patient baseline according to the National Cancer Institute Common Toxicity Criteria guidelines, version 2.0. CBCs. Liver function status, complications include chemotherapy relative diarrhea and I to II bone marrow stifled was inspected.

Statistical methods
Clinical responses between two groups were compared by chi-square test. Kaplan-Meier analysis was applied to assess OS, and the significance of differences in survival curves was determined by the log-rank test. OS was defined as the period from the date of chemotherapy to the date of death. P < 0.05 was considered significant for all analyses. The SPSS software package, version 10.0 (SPSS Inc., Chicago, IL, USA) was used for statistical analysis.

Results

Cell culture and drug sensitivity test
Primary tumor cells prepared from the specimens of primary liver tumors obtained by biopsy turned into rapid logarithmic growth phase after cultured for 48 to 72 hours (Figure 1). After 4 days of incubation, chemotherapy drugs were added at the top of the rapid logarithmic phase in the MTT assay. As shown in Figures 2 and 3, different drugs showed different inhibitions to the growth of cancer cells from different patients. In addition, the cells were disrupted and chromatin-mustered when the chemotherapy drugs were added for 24 hours (Figure 4).

| Table 1: Drug sensitivity test of experimental group |
| --- |
| Drug | Tumor suppression rate (%) | Drug-resistant rate (%) |
| E-ADM | 36.7±4.5 | 28.8 |
| HCPT | 33.8±9.4 | 31.5 |
| MMC | 29.4±6.6 | 33.2 |
| 5-FU | 38.5±7.1 | 25.4 |
| LOHP | 40.9±6.3 | 21.6 |
| DDP | 50.2±8.8 | 18.3 |
| GEM | 32.3±3.9 | 27.1 |

FIG. 1: Growth curve of liver cancer cells cultured in RPMI 1640 (Gibco, Invitrogen) supplemented with 10% (v/v) heat inactivated fetal bovine serum and 1% (v/v) antimotic antibiotic at 37°C under a 5% CO2 atmosphere.

FIG. 2: Inhibition of liver cancer cell growth by chemotherapy drugs added at the top of the logarithmic growth phase.
Clinical observation

120 cases of unresectable HCC were randomly divided into experimental group and control group, the clinical pathological features of patients in both groups were not significant different (Table 2). The mean patient age at the time of treatment was 56 (range, 47 - 65) years. Bard Access Ports were inserted into the right gastro-omental artery and the right gastro-omental vein to establish regional chemotherapy channels for all patients (Figure 5). Individual patients in the experimental group were infused by arterial and portal vein pump 3 drugs which showed highest inhibition to the cancers of the patient in our MTT test above, whereas the control group patients underwent TACE. In experimental group, the tumor size reduced in 28 (47.6%) cases as measured by CT or MRI after six chemotherapy cycles (Figure 6), stabilized in 14 (23.3%) cases, and progressed in 18 (30%) cases. The corresponding data was 17 (28.5%), 7(11.5%) and 36 (60%) cases in the control group (P < 0.05). 6 patients in the experimental group and 3 in control group had the opportunity of two-step operation (P < 0.05) (Table 3). There was no severe incidence of complications in both groups. In the experimental group, 2 (3%) cases of patients had wound infection, 8 cases had the chemotherapy relative diarrhea and 18 (30%) cases had I° to II° bone marrow stifled. In the control group, The chemotherapy relative diarrhea was seen in 15 (25%) cases and I° to II° bone marrow stifled was seen in 33 (55%) cases (Table 4). For prognosis, in the experimental group, the median follow-up time was 21 months, OS rates of 6 months, 12 months, 18 months were 86%, 72%, and 65% respectively. In the control group, the median follow-up time was 16 months, OS rates of 6 months and 12 months were 58% and 40%, respectively (Figures 7).
**TABLE 2**: Patient characteristics: patient and tumor features of the experimental group compared with control group

|                | Experimental group | Control group | P-value |
|----------------|--------------------|---------------|---------|
| n              | 60                 | 60            |         |
| **Age (Y)**    |                    |               |         |
| Median (range) | 56 (47-65)         | 55 (46-64)    |         |
| Male : Female  | 53:7               | 11:1          |         |
| Cirrhosis      | 56 (93.3%)         | 54 (90%)      |         |
| **Primary tumor pathologic stage** |                    |               |         |
| II             | 21                 | 27            |         |
| III            | 39                 | 33            |         |
| **Tumor size (cm)** |                |               |         |
| Median (range) | 9 (8 - 13.7)       | 9.3 (8.7 - 13.3) |         |
| **Tumor number** |                  |               |         |
| 1              | 23                 | 37            | > 0.05  |
| ≥2             | 21                 | 39            |         |
| **Hepatitides** |                  |               |         |
| Hepatitis B    | 57 (95%)           | 58 (96.7%)    |         |
| Hepatitis C    | 1 (1.7%)           | 1 (1.7%)      |         |
| HepatitisB/C   | 0 (0%)             | 1 (1.7%)      |         |
| Other          | 2 (3.3%)           | 0 (0%)        |         |
| AFP (Log10ng/ml) | 4.33 ± 2.52       | 4.18 ± 2.66   |         |
| **Child-pugh** |                    |               |         |
| A              | 38 (63.3%)         | 40 (66.7%)    |         |
| B              | 22 (36.7%)         | 20 (33.3%)    |         |
| C              | 0 (0%)             | 0 (0%)        |         |
| **Neoplasm**   |                    |               |         |
| Single-tumor   | 6 (10%)            | 4 (6.7%)      |         |
| Multi-tumor    | 54 (90%)           | 56 (93.3%)    |         |

**FIG. 5**: Bard Access Ports were inserted into the right gastro-omental artery and the right gastro-omental vein to establish regional chemotherapy channels. A. Hepatic artery catheter was inserted into the blood vessel to liver tumor, the liver segment appeared blue after methylene blue staining; B. Operation of insertion of chemotherapy catheter; C. Artery images showing that the chemotherapy catheter was in the proper hepatica artery; D. Vein images showing that the chemotherapy catheter was in the portal vein.
FIG. 6: The tumor size reduced in experimental group as measured by CT or MRI after six chemotherapy cycles. A. MRI showed that the tumor was located in segments I and VII; B. The tumor size reduced and the blood supply decreased obviously after 6 cycles of chemotherapy; C. Tumor located in segment VII and the cells growing into inferior vena cava (IVC) via right hepatic vein; D. The tumor size reduced and the embolus in IVC and right hepatic vein disappeared after 6 cycles of chemotherapy.

### TABLE 3: Clinical effect comparisons between experimental and control groups

| Group       | n  | CT/MRI | AFP | Two-step | P-value |
|-------------|----|--------|-----|----------|---------|
|             |    | Reduce | stabilize | progress | decrease | rise/ stabilize | operation (n) |          |
| experimental group | 60 | 28     | 14    | 18       | 51      | 9              | 6            | < 0.05    |
| control group     | 60 | 17     | 7     | 36       | 30      | 30             | 3            |           |

### TABLE 4: Complications comparison between experimental and control groups

| Group       | n  | wound infection | abdominal pain | hepatic failure | myelosuppression | P-value |
|-------------|----|----------------|----------------|-----------------|------------------|---------|
| experimental group | 60 | 2              | 8              | 9               | 18               | < 0.05  |
| control group     | 60 | 0              | 25             | 41              | 33               |         |
FIG. 7: OS (Overall survival, months) comparison between experimental group and control group

Discussion

China is a country in which the morbidity of HCC is the highest in the world, and the latest survey data showed that the mortality rate of PHC is the highest in Hubei province. Only less than 20% of the patients with PHC had the opportunity to be respected. Hepatic trans catheter arterial chemoembolization (TACE) is currently the important therapy of the middle and late stage of PHC patients with a 30%~40% effective rate. Research has showed that blood supply to primary liver cancer is through both hepatic artery and portal vein: the tumor center is mainly composed of hepatic artery, and the peripheral part is mainly composed of portal vein. The surrounding part is the most active part of the cancer. HCC cells invade intrahepatic vascular extremely common, especially the portal vein, taking shape of portal vein embolus and giving rise to intrahepatic metastasis. This is the reason why conventional TACE cannot achieve satisfactory results.

The efficiency of cancer chemotherapy is decided by two factors. The first is drug concentration in the tumor, and the second is the drug effectiveness. Thus, it is important to improve the drug efficiency. In the early 1950's, Bierman first reported hepatic artery perfusion chemotherapy for primary liver cancer. Storer reported chemotherapy of hepatic neoplasms via umbilical portal vein in 1966 and other scholars adopt the hepatic artery or portal vein catheter chemotherapy for the sake of increasing the medicine concentration in tumor, and achieved good efficiency. Here, we used drug sensitivity test to select individualized drugs and perfused the sensitive drugs to individual patients by hepatic artery and portal vein channels, giving consideration to improve the regional drug concentration and tumor susceptibility to drugs and achieve high effectiveness of treatment of PHC.

Surgical resection is the preferred treatment for liver cancer, but because of the low removal rate, the 5-year survival rate of entire population of liver cancer is less than 5%. Molecular biology research proved that portal venous endothelial cells have specific adhesion molecules which can bind the various components in the blood, including cancer cells, to promote the transfer and spread of tumor cells in the portal vein. Clinical studies also demonstrated that hepatic artery embolism chemotherapy and portal vein regional chemotherapy of hepatocellular carcinoma after surgical removal can significantly decrease the rate of recurrence and metastasis, and regional chemotherapy through hepatic artery is safe and effective. Reports for colorectal cancer liver metastasis in patients with arterial perfusion 5-Fu can obtain curative effect. With hepatic arterial infusion of fluorouracil plus systemic irinotecan for unresectable hepatic metastases from colorectal cancer, the patients have a response rate of 72% and the median survival time is 49.8 months. Case reports also give us encouraging results. For patient with gastric cancer and breast cancer which is metastasized to liver, regional infusion chemotherapy gains curative effect. In neuroendocrine carcinoma liver metastasis, regional infusion chemotherapy is superior to systemic chemotherapy also. Primary hepatic carcinoma with III/IV segment portal venous tumor emboli resort to hepatic artery perfusion of cisplatin treatment after resection reached relapse-free survival more than 1 year. Regional infusion chemotherapy has better prospective than systemic chemotherapy of liver cancer, but there have been no correspond-
ing treatment standards.17 Solid tumors metastatic to the liver had received good prognosis with oxaliplatin, fluorouracil and leucovorin infused from hepatic artery.18

For unresectable hepatocellular carcinoma, with continually infusion of low dose fluorouracil and cisplatin by hepatic artery, the liver tumor remission rate was 39%, and the median survival was 15.9 months.19 Tanaka’s research showed that hepatic arterial infusion chemotherapy is a safe, effective method to liver cancer patients.20 The curative effect is better and side-effect is lower with the chemotherapy catheter infusion.21

The sensitivity of tumor cells to chemotherapy drugs can be obtained by drug susceptibility test.22 Naitoh et al reported after multicenter clinical trial that culture-drug sensitivity test predicts the outcome of patients undergoing chemotherapy for advanced gastric cancer and can be used as the method to evaluate chemosensitivity.23 Taylor found that the survival rate of the group of malignant tumor patients treated with sensitive drugs was obviously higher than that of the drug resistant group and simple surgical treatment group. This research suggested that the drug sensitivity test in vitro guided individualized chemotherapy is effective.24

Another question debated is whether the result of the drug susceptibility reaction in vitro is accordance with that in vivo. In vitro results and the in vivo effectiveness may not match exactly25, but recent studies have reported that 85.7% consistency of susceptibility test in vitro and clinical observation.26 Kabeshima and Taylor’s research showed that the patients’ long-term survival rate was obviously higher in drug-sensitive group than that of drug resistant group and simple surgical group, this indicates that drug sensitivity test in vitro can play an important role in directing the individualized chemotherapy and improving the curative effect.26, 27 Thus, we carried out this study and our result is encouraging.

Compared with systemic chemotherapy, regional chemotherapy can have these advantages: (1) chemotherapy drug concentration in the tumors increases significantly and makes chemotherapy drug acts on the tumor tissues directly and efficiency, (2) Along with the increase of the drug concentration in the tumor, curative effect increase accordingly. However, it is a critical factor of finding effective chemotherapy drugs for the patients and administrates these drugs by regional catheters.

Seven anticancer drugs were chosen in our study for in vitro drug sensitivity test in 60 cases of patients with unresectable HCC. The results showed that the tumor inhibition rate is different in seven anticancer drugs, and different drug have different resistance rate. Chemotherapy guided by drug susceptibility test shows that the tumor response rates, including the changes of AFP, OS, and second-stage-resection, were superior in experimental group than that of control group, whereas the side effects were relatively small and the survival rate was higher.

In the dual channel infusion chemotherapy under the guidance of drug susceptibility test, regional chemotherapy has better clinical benefit than systemic chemotherapy, providing us a useful way for patients’ treatment with primary liver cancer. However, because there were cases in which we failed to detect sensitive chemotherapy drugs in the experimental group, more drugs should be tested in future studies.

Conclusion

The artery and portal vein pump transfusion chemotherapy guided by drug sensitive test was efficient for HCC treatment. The patients can get longer OS and lower complication incidence.

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Conflict of interest

The authors declare that they have no conflicts of interest. The authors alone are responsible for the content and writing of the paper.

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