A retrospective analysis of plasma concentration monitoring of fluorouracil in patients with advanced colorectal cancer

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ABSTRACT

Objectives To analyse the results of fluorouracil (5-FU) plasma concentration monitoring in patients with advanced colorectal cancer after 5-FU treatment, and to provide a reference for the application prospect of 5-FU plasma concentration monitoring technology.

Methods A retrospective analysis was performed with advanced colorectal cancer patients treated with 5-FU from March 2015 to August 2018. The results of plasma concentration monitoring of 5-FU, severe adverse reactions, and anti-tumour efficacy were analysed.

Results Among 47 patients, 5-FU plasma concentration monitoring was carried out a total of 289 times. The area under the receiver operating characteristic (ROC) curve (AUC) reflecting 5-FU exposure in vivo was 2.8–158 mg·h/L (41±94.6 mg·h/L). Mean AUC range within the target range (20–30 mg·h/L) for each patient was observed in 28.8% of patients. The overall incidence of related severe adverse reactions in the AUC <30 mg·h/L group was lower than that in the >30 mg·h/L group (24.0% and 50.0%, respectively) (p=0.06), and the incidence of severe neutropenia was 12.0% and 40.9%, respectively (p=0.05). The disease control rate and overall response rate of the AUC <20 mg·h/L group was lower than that of the ≥20 mg·h/L group: 83.3% vs 97.1% (p=0.19) and 25.0% vs 51.4% (p=0.10), respectively.

Conclusions The 5-FU plasma concentration monitoring technique can improve the safety and efficacy of 5-FU administration to advanced colorectal cancer patients. It is expected to become an important means to individualise 5-FU use in the Chinese population.

INTRODUCTION

Colorectal cancer is one of the top five malignant tumours in China; its high incidence and mortality are second only to lung, gastric, oesophageal and liver cancer.1 FOLFOX/FOLFIRI consists of fluorouracil (5-FU), oxaliplatin and irinotecan combined with bevacizumab, and is a first-line anti-tumour therapeutic regimen for advanced colorectal cancer.2 Due to the general condition of a patient, severe adverse drug reactions, patient medication compliance, and so on, the chemotherapy drug dose cannot be calculated solely based on body surface area or be empirically reduced in most cases.3 4 The evaluation of indexes such as dose index (DI), time index (TI) and relative dose intensity (RDI) can, however, objectively reflect patient acceptance of a chemotherapy drug dose.3 5

Monitoring 5-FU plasma concentration has been used with some success in Europe, and serves as an important technical means to individualise 5-FU drug dose. A series of studies by Gamelin et al found that when 5-FU area under the receiver operating characteristic (ROC) curve (AUC) is within the target range, namely 20–25 or 20–30 mg·h/L, an optimal anti-tumour effect with a relatively small toxic and side effect profile could be achieved. However, when 5-FU AUC is lower than the target, subsequent 5-FU dosage should be increased; otherwise, 5-FU dosage should be reduced.6 9 Since March 2015, our hospital has been using 5-FU plasma concentration monitoring with the aim of exploring its reference value and comparing its clinical efficacy and safety for patients with advanced colorectal cancer in China. Herein, we reviewed all patients who underwent 5-FU plasma concentration monitoring from the initiation of this technology to August 2018. The 5-FU DI, plasma concentration monitoring results, severe adverse reactions and anti-tumour curative efficacy were retrospectively analysed to provide a reference for the potential application of the 5-FU plasma concentration monitoring technology in Chinese patients.

DATA AND METHODS

Clinical data

We selected patients with advanced colorectal cancer who received 5-FU therapy at our hospital...
from March 2015 to August 2018; basic data such as gender, age, body surface area, complicating diseases, chemotherapy regimen, and drug dosage were retrieved. The 5-FU was produced by Tianjin Jinyao Pharmaceutical Co., Ltd, State Food and Drug Administration approval number: H12020959 (specification: 10 mL, 0.25 g).

Main inclusion criteria included patients: treated with mFOLFOX6, FOLFOR4 or FOLFIRI as the first-line or second-line therapeutic regimen; who received ≥2 cycles of anti-tumour therapy and ≥1 5-FU plasma concentration monitoring procedure during hospitalisation; and evaluated with radiographic tumour assessment at least once. Patients or their legal representative must have also given written informed consent. Main exclusion criteria included patients with incomplete medical records; who developed severe adverse reactions or complications; and refused further therapy; and patients with visibly poor treatment compliance.

Drug dose

Standard drug dose mFOLFOX6 and FOLFIRI regimen, the 5-FU standard medication regimen, was 400 mg/m², intravenous (iv), d1, 5-FU 2400 mg/m², iv pump 46 hours. FOLFOX4 regimen was 5-FU 400 mg/m², iv, d1, d2, 5-FU 600 mg/m² iv pump 22 hours, d1, d2. One treatment cycle spanned 14 days.

Dose evaluation index

DI refers to the percentage of the actual drug dose in the standard drug dose. TI refers to the percentage of the standard drug use interval in the actual drug use interval. Dose intensity=DI×TI. RDI refers to the percentage of the actual dose intensity of a specific drug in the standard dose intensity. For instance, for mFOLFOX6, as the standard dose intensity is 1400 mg/m²/week, and the actual dose intensity is 1200 mg/m²/week, its RDI is 85.7%.

DI, TI and RDI

When determining initial DI, TI and RDI, factors such as general condition, age, body surface area, and willingness to accept treatment, and so on should be considered. Subsequent DI, TI and RDI should be based on 5-FU plasma concentration monitoring results, adverse reactions after medication and other factors; 5-FU drug dose was jointly decided by two experienced tumour physicians, and a clinical pharmacist assisted for some cases.

5-FU plasma concentration monitoring

Plasma sample collection

For the 5-FU pump, medication time was uniformly confirmed to start at 15:30-16:30 on day 1. At 8:30-9:30 on day 3 from the 5-FU continuous intravenous pumping, venous blood of the patient was sampled and placed into a polyethylene tube pretreated with stabiliser (gimeracil) for 10 min by a nurse, then stored for 2 hours at 2°C–8°C. Plasma was extracted by centrifugation, and cryopreserved at −80°C for 3 months by a pharmacist.

5-FU plasma concentration assay

The 5-FU plasma concentration monitoring method established in the previous stage was used. In brief, patient venous blood was sampled and centrifuged to extract plasma, and plasma sample pretreated by liquid-liquid extraction. For chromatographic conditions, an Agela Innoval NH2 chromatographic column (2.1 mm×50 mm, 5 μm) was used for isocratic elution in the fluid phase with methanol:ultrapure water (2:98) at a flow rate of 0.3 mL/min; column temperature was 40°C. Using mass spectrometry, 5-FU and 5-bromouracil (the internal standard) were detected under a multi-reaction monitoring scan mode using electrospray negative ion mode. Quantitative analysis of ion pairs yielded m/z 128.8→42.1 (5-FU) and m/z 188.6→42.1 (5-bromouracil). The 5-FU plasma concentration was measured using a liquid chromatography tandem mass spectrometer (Shimadzu LC System Tandem AB Sciex API Model 3200 Triple Quadrupole Mass Spectrometer).

5-FU plasma concentration monitoring results

5-FU plasma concentration monitoring results were expressed by AUC. The calculation formula was as follows: AUC (mg*h/L)=5-FU plasma concentration (ng/mL)×pump maintenance time (h)/1000. Based on reports by Kaldate and Denda for this subject group, we used AUC in the 20–30 mg*h/L group as the target range. Mean AUC represents average AUC obtained per patient per time.

Evaluation of adverse reactions

The Common Terminology Criteria for Adverse Events (CTCAE) Version 4.03 standard was used as an evaluation criteria, and severe adverse events associated with 5-FU (including...
neutropenia, anaemia, thrombocytopenia, diarrhoea, oral mucositis, hand-foot syndrome) were statistically analysed.13

Curative effect judgment criteria
The curative effect was evaluated using the Response Evaluation Criteria in Solid Tumours (RECIST) Version 1.1, divided into complete response (CR), partial response (PR), stable disease (SD) and progressive disease (PD). Disease control rate (DCR) = (CR+PR+SD) caseload/total caseload ×100%, overall response rate (ORR) = (CR+PR) caseload/total caseload ×100%.14

Statistical methods
Original case data were sorted using Microsoft Excel, and mean AUC, SD and median were calculated. Curve regression coefficient analysis was performed using Spearman. SPSS 19.0 software was used, and the χ² test employed to compare 5-FU AUC, severe adverse reactions, and efficacy evaluation. P<0.05 was considered statistically significant.

RESULTS
General condition of patients
This study included 47 cases with 28 men and 19 women (average age, 60±10 years). The number of patients with colon, rectal-sigmoid junctional, and rectal cancer was 24, 5 and 23, respectively, and included five cases that successively received both chemotherapy regimens. The complicating diseases included diabetes, hypertension, sinus tachycardia, atrial fibrillation, lacunar infarction, and allergic dermatitis with caseloads of 12, 14, 2, 2, 1 and 1, respectively. There were 33 and 19 cases of first-line and second-line chemotherapy, respectively. The number of patients treated with mFOLFOX6, FOLFOX4 and FOLFIRI was 20, 7 and 15, respectively, and 32 cases were treated via combination with bevacizumab.

DI, TI, RDI and AUC of 5-FU
Chemotherapy was administered to the 47 cases and cumulatively, plasma concentration monitoring was performed 289 times; the average number of measurements per case was 5.9±3.1 times. Mean DI, TI and RDI reflecting 5-FU dose were 83.3%±16.5%, 76.1%±20.4% and 63.1%±21.1%, respectively. The AUC range was 2.8–158 mg·h/L, and mean AUC for each patient was 41±94.6 mg·h/L, of which AUC of 60 cases (20.8%) and average AUC of 13 patients (27.7%) were within the target range (table 1). A Spearman linear regression analysis was performed on the average AUC of 5-FU for each patient and DI of 5-FU. The results suggested that 5-FU AUC increased with an increase in DI (figure 1).

Correlation between AUC and RDI of 5-FU, and severe adverse reactions and curative effect of anti-tumour therapy
Severe adverse reaction and effect of anti-tumour therapy for the AUC <20 mg·h/L, 20–30 mg·h/L and >30 mg·h/L patient groups are shown in table 2. All differences in the incidence of 5-FU-related severe adverse reactions (neutropenia, anaemia, thrombocytopenia, diarrhoea, oral mucositis, hand-foot syndrome), overall incidence of severe adverse reaction, DCR and ORR between the three groups were not statistically significant (p>0.05). Overall incidence of severe adverse events was lower in the AUC ≤30 mg·h/L group (combination of the AUC <20 and 20–30 mg·h/L groups; 24.0%) than in the AUC >30 mg·h/L group (50.0%); no statistical difference was found (p=0.06). The incidence of severe neutropenia was 12.0% and 40.9%, respectively, and similarly, no statistical difference was found (p=0.05). The DCR of the AUC ≥20 mg·h/L group (combination of the AUC 20–30 and >30 mg·h/L groups; 97.1%) was higher than that of the AUC <20 mg·h/L group (83.3%); no statistical difference was found (p=0.09). The ORR of the two groups was 51.4% and 25.0%, respectively, and no statistical difference was found (p=0.11).

The limit 68% is the median RDI value of overall level. Table 3 shows the severe adverse reaction and anti-tumour effect of 5-FU RDI <68% group and ≥68% group. DCR between the two groups was 86.4% and 100.0%, respectively, with no statistical difference found (p=0.19). The 5-FU RDI <68% group had a lower ORR than the 5-FU RDI ≥68% group (31.8% and 56.0%, respectively); we found no statistical difference (p=0.10).

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<tr>
<th>Table 2</th>
<th>The relationship between area under the curve (AUC) and treatment outcomes</th>
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<td>AUC level</td>
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<td>&gt;30</td>
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AUC, area under the receiver operating characteristic (ROC) curve.

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<th>Table 3</th>
<th>The relationship between relative dose intensity and treatment outcomes</th>
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<tr>
<td>RDI level</td>
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RDI, relative dose intensity.
DISCUSSION

Technical points and clinical significance of 5-FU plasma concentration monitoring

The techniques used to detect 5-FU TDM include high-performance liquid chromatography, liquid chromatography tandem mass spectrometry, and immunoassay, with each having their respective advantages and disadvantages. In this study, 5-FU plasma concentration was determined via liquid chromatography tandem mass spectrometry that was strictly validated using a previous methodology. This technique is accurate and reliable for detection. The 5-FU plasma concentration presents chronological characteristics, and fluctuation in plasma concentration appears during a single intravenous pumping of 5-FU. In this study, the time of blood sample collection was uniformly controlled between 8:30 and 9:30 (4.5–5.5 hours before the end of intravenous pumping) to reduce the fluctuation in plasma concentration caused by the chronological characteristics of 5-FU. The premature end of venous pumping was avoided, and it was convenient for staff members to collect plasma samples during regular working hours.

Due to the poor stability of 5-FU in whole blood, it is necessary to centrifuge whole blood to extract plasma at the earliest convenience, or to place whole blood into a polyethylene tube with a stabiliser (5-FU metabolic enzyme dihydrouracil dehydrogenase inhibitor, gimeracil) after whole blood collection. Strict control was employed by the staff during plasma sample collection to prevent an overly-low result in the detection of 5-FU plasma concentration owing to its instability in whole blood.

Gamelin et al proposed a programme to adjust 5-FU drug dose based on its AUC, as depicted in table 4. When AUC is lower than the target value, the subsequent 5-FU dose is increased, and when it is higher, the subsequent 5-FU dose is decreased. This group of patients possesses a relatively broad target AUC range of 20–30 mg*h/L, which is consistent with relevant Chinese publications. AUC in multiple chemotherapy cycles was used to analyse the correlation between AUC and severe adverse reactions, and anti-tumour efficacy. We concluded that AUC levels were positively correlated with the incidence of severe adverse reactions and anti-tumour efficacy. The results obtained for this subject group are inherently consistent with the study by Ju et al. No statistical significance was found in this research group, which may be attributed to the overly-low number of cases included.

For most patients, 5-FU dose remained unchanged while AUC displayed a certain fluctuation range each time, suggesting that it is inappropriate to adjust 5-FU dose based on AUC level each time; this is consistent with the results of Matsumoto et al. Referring to the average value of 5-FU plasma concentration measured by Ju et al during multiple chemotherapy cycles, we combined 5-FU AUC with DI, TI, RDI and severe adverse reactions. The 5-FU dose adjustment scheme recommended by Denda in table 4 suggests the need for a comprehensive judgement.

We recommend: for patients in the AUC <20 mg*h/L group, if a severe adverse reaction does not occur, the upper limit of 5-FU drug dose can be increased until RDI reaches 100%, to improve anti-tumour efficacy; for patients in the AUC >30 mg*h/L group, if a severe adverse reaction occurs, the subsequent 5-FU dose can be reduced to improve medication safety; and for patients in the AUC ≤30 mg*h/L group, if a severe adverse reaction occurs, oxaliplatin or irinotecan dose can be preferentially reduced, and 5-FU dose may not be reduced temporarily to ensure 5-FU anti-tumour efficacy, while improving medication safety.

Role of DI, TI and RDI in 5-FU plasma concentration monitoring

DI can visually reflect each medication intensity. Mean DI, TI and RDI of 5-FU in this group were 83.3%±16.5%, 76.1%±20.4% and 63.1%±21.1%, respectively, suggesting that the under-dosing of anti-tumour drugs should be addressed. RDI is related to DI and TI numerically, and from a clinical view it is associated with the efficacy of anti-tumour therapy, and is related to drug resistance of cancer cells. Munker et al studied the effect of RDI on chemotherapy efficacy, and it was agreed that a relatively high RDI should be applied if adverse reactions are tolerable. In this study, the proportion of AUC <20 mg*h/L was 40.5%, which may be due to the relative low DI and RDI. The authors recommend that DI, TI and RDI be close to 100% when the patient’s adverse reactions are tolerable. When severe adverse reactions occur, 5-FU plasma concentration monitoring results should be considered, and chemotherapy dose may be adjusted to reduce the risk of severe adverse reactions and ensure anti-tumour efficacy.

In conclusion, 5-FU plasma concentration monitoring may improve the safety and efficacy of treating patients with advanced colorectal cancer in China. It is expected to become important for clinical individualised 5-FU use.

### Table 4: Adjusting scheme of fluorouracil dosage according to the area under the receiver operating characteristic curve

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AUC, area under the receiver operating characteristic (ROC) curve.

Refe

Competing interests

Plan of Medical Devices (No. 20160422) and funding for the research-oriented clinician plan of Tongji Medical College, Huazhong University of Science and Technology (No. 5001540076).

Patient consent for publication

None declared.

Provenance and peer review

Not commissioned; externally peer reviewed.

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REFERENCES