TEGAFIRI is an effective alternative regimen for the management of recurrent or metastatic colorectal cancer

TZU-CHI HSU

Department of Surgery, Mackay Memorial Hospital, Zhongshan, Taipei 104, Taiwan, R.O.C.

Received February 26, 2014; Accepted November 21, 2014

DOI: 10.3892/ol.2015.2855

Abstract. At present, the global incidence of colorectal cancer is increasing, with numerous individuals succumbing to the disease. The standard treatment strategy for colorectal cancer is curative resection. However, a cure is rarely achieved for metastatic colorectal cancer. Currently, chemotherapy is the main treatment for metastatic and recurrent colorectal cancer. The majority of metastases or recurrences have been found to respond well to chemotherapy. The present study evaluated the response rates of recurrent or metastatic colorectal cancer patients treated with a combination chemotherapy of irinotecan and oral uracil-tegafur (UFUR). In the pilot study, 33 patients with metastatic or recurrent colorectal cancer were treated with different regimens of irinotecan and UFUR with or without leucovorin; however, irinotecan (150 mg/m² every two weeks) with continuous UFUR and leucovorin without interruption resulted in improved survival compared with the other regimens evaluated and, thus, was employed for the present study of 113 patients. The patients that received irinotecan with UFUR and leucovorin without interruption exhibited similar efficacy in terms of overall survival and response rate to that of the pilot study. In addition, the incidences of diarrhea, alopecia and hematologic toxicity were acceptable, which was in agreement with the results of the pilot study. Therefore, combination chemotherapy with irinotecan, oral UFUR and leucovorin appears to be a satisfactory treatment strategy for recurrent or metastatic colorectal cancer.

Introduction

Currently, the preferred treatment strategy for metastatic and recurrent cancer is chemotherapy (1-4), as numerous types of metastatic and recurrent cancer appear to respond well to this mode of therapy (1-6). In addition, multidrug therapies result in improved responses compared with single agent therapies.
were monitored from January 2006 until December 2010 or mortality.

Patients eligible for the pilot study (15) received different regimens and doses of TEGAFIRI with or without leucovorin. The patients were randomized into three groups: Group I, 150 mg/m² irinotecan every two weeks and UFUR for one week every two weeks; group II, 100 mg/m² irinotecan for two weeks followed by a one week of rest, with continuous UFUR and leucovorin; and group III, 150 mg/m² irinotecan every two weeks with continuous UFUR and leucovorin. The UFUR dose was standardized at 300 mg/m²/day and leucovorin was administered at 45-60 mg/m²/day. Although the intention was to include a greater number of patients in the pilot study, the initial results demonstrated that group I patients exhibited lower response rates compared with the other two groups; therefore, it was unethical to proceed with the group I regimen and the enrollment was terminated after 33 patients had enrolled. From the pilot study it was determined that that the group III regimen was optimal for the treatment of metastatic or recurrent colorectal cancer; thus, this regimen was administered to patients eligible for the present study.

In the present study, all of the 113 enrolled patients were treated with irinotecan at a standard dose of 150 mg/m² every two weeks with continuous UFUR (300 mg/m²/day) and leucovorin (45-60 mg/m²/day). The Dukes' stage of colorectal cancer before recurrence was determined and evaluated for each patient (18). In addition, prior to irinotecan administration, patients were administered with 10 mg dexamethasone intravenously, 3 mg granisetron or 8 mg ondansetron intravenously, and 0.5 mg atropine subcutaneously. Supportive care included loperamide for diarrhea, antiemetic agents and oral cephradine for diarrhea lasting >48 h. Upon progression, patients were administered with an oxaliplatin-based salvage regimen in addition to UFUR and leucovorin (TEGAFOX); however, agents such as bevacizumab and cetuximab were not routinely used due to a lack of funding during the study period.

Patient monitoring and follow-up. In the pilot study, patients in each group were evaluated by determining the serum carcinoembryonic antigen (CEA) and performing a chest X-ray, abdominal ultrasound and computed tomography scan of the chest or abdomen every three months. In addition, the Response Evaluation Criteria In Solid Tumors (RECIST) was used to assess the efficacy of each chemotherapy regimen by categorizing the response into the following four grades: Progression, stable disease, partial response and complete response (19). Progressive disease was defined as an increase in CEA levels, a ≥25% increase in the number or size of the metastatic lesions or the development of new lesions; a partial response was defined as a decrease in CEA levels or a ≥25% decrease in the number or size of the metastatic lesions; and a complete response was defined CEA levels with the normal range or as the disappearance of metastatic lesions.

Dose modification. The severity of the adverse effects was evaluated using the National Cancer Institute Toxicity Criteria (version 2.0) (20). Upon the initial appearance of grade II toxicity, no dose reduction was required. In addition, upon the appearance of reactions, which were determined as unlikely to become serious or life-threatening, no treatment interruption or dose reduction was implemented. However, in cases of grade III or greater toxicity, TEGAFIRI/leucovorin treatment was interrupted and was not resumed until the toxicity had resolved or had improved to grade I. When treatment was resumed, the dose of leucovorin was as before; however the doses of irinotecan and UFUR were reduced as follows: Irinotecan and UFUR doses were reduced by 25% in patients who exhibited a second occurrence of grade II toxicity or any occurrence of grade III toxicity; and irinotecan and UFUR were reduced by 50% in patients who experienced a third occurrence of grade II toxicity or a second occurrence of grade III toxicity. Treatment was discontinued if, despite dose reduction, grade II toxicity occurred for a fourth time, grade III toxicity...
occurred for a third time or if grade IV toxicity occurred at all. If granulocytes decreased to <500/mm^3, grade III-IV diarrhea developed or granulocytes decreased to <1,000/mm^3 with concomitant fever, irinotecan and UFUR doses were reduced by 20% for one cycle.

Statistical methods. The baseline characteristics of the patients were quantified using descriptive statistics (median, percentile, and range) and the principle results were overall survival and progression-free survival. Overall survival was defined as the period of time from the commencement of irinotecan therapy to mortality. Progression-free survival was defined as the period of time from group randomization to disease progression, or mortality from disease progression or an unknown cause. For multivariate analysis, factors associated with the time to progression were identified by performing Cox’s regression analysis with forward stepwise conditional analysis.

Results

Pilot study. In the pilot study (15), no significant differences were identified between the three groups in terms of dose intensity and dose delivery. Groups II and III exhibited improved response rates compared with group I (Table I). The overall survival times of groups I, II and III were 15, 21.5 and 19.8 months, respectively (Fig. 1). No statistically significant differences were identified between the three groups in terms of overall survival; however, group III patients exhibited fewer non-hematological side effects and improved tolerance to the regimen compared with groups I and II. Therefore, the group III regimen from the pilot study was selected for use in the present study (n=113).

Present study. Table II indicates the demographic data of the 113 patients enrolled in the present study. The median age of the patients was 61 years (range, 21-81 years). According to the RECIST criteria, the response rate of the present study was 52.2%, which is similar to that of the pilot study (54.5%). In

| Group I (n=11) | Group II (n=11) | Group III (n=11) |
|---------------|----------------|-----------------|
| Complete response | 0 | 2 | 2 |
| Partial response | 3 | 4 | 4 |
| Mortalities | 9 | 9 | 8 |
| Survival range, months | 6-30 | 5-31 | 5-33 |

Table III. Adverse effects experienced by patients in the present study exhibiting an overall toxicity grade of III-IV.

| Adverse effect | Patients (n=113) |
|---------------|-----------------|
| Non-hematological | n | % |
| Diarrhea | 22 | 19.5 |
| Vomiting | 28 | 24.8 |
| Alopecia | 79 | 69.9 |
| Nausea | 45 | 39.8 |
| Hematological | n | % |
| Anemia | 34 | 30.1 |
| Neutropenia | 67 | 59.3 |
| Febrile neutropenia | 5 | 4.4 |
| Thrombocytopenia | 12 | 10.6 |

Furthermore, the progression-free and overall survival curves were calculated according to the Kaplan-Meier method and compared using a log-rank test. All statistical analyses were performed using SPSS software version 17.0.1 (SPSS, Inc., Chicago, IL, USA). P<0.05 was considered to indicate a statistically significant difference.
addition, the incidence of diarrhea, alopecia and hematologic toxicities (Table III), as well as the necessity of delaying or decreasing the dosage were similar to those of the pilot study. For Dukes’ stages B, C and D, the overall patient survival time was 19.9, 23.7 and 27.4 months, respectively (Figs. 2 and 3), and the progression-free survival time was 18.7, 16.8 and 18.7 months, respectively (Figs. 4 and 5).

Discussion

As the worldwide incidence rate of colorectal cancer increases, increasing numbers of patients are succumbing to the disease (21,22). In 2012, there were approximately 694,000 mortalities as a result of colorectal cancer (23), and the American Cancer Society have estimated that 136,830 novel cases of colorectal cancer will be diagnosed and 50,310 mortalities as a result of the disease will occur in 2014 (24). In Taiwan, colorectal cancer is the third leading cause of cancer-related mortality and the second most common cause of malignancy. The principal treatment strategy for colorectal cancer is curative resection; however, remission of metastatic colorectal cancer is rarely achieved (25). Therefore, chemotherapy is currently employed as the preferred treatment strategy for metastatic disease (1-6).

A novel inhibitor of the DNA enzyme topoisomerase I, irinotecan exerts cytotoxic activity by interrupting DNA replication and transcription. In studies conducted in Western countries, TEGAFIRI with leucovorin administration resulted in response rates of 25-40% (4,6). Furthermore, good response or survival rates have been observed in a number of Taiwanese studies of first- and second-line irinotecan therapy. However, the addition of 5-FU or its precursors plus leucovorin were crucial for achieving satisfactory response rates.

Tegafur is an oral fluoropyrimidine, which is metabolized to 5-FU in vivo (26). In the management of metastatic colorectal cancer, tegafur appears to be an active and minimally toxic alternative to other types of fluoropyrimidine (27). Additionally, uracil is a naturally occurring pyrimidine, which is able to incorporate into nucleic acids (28). Together, these agents may be administered as oral UFUR, which consists of tegafur combined with uracil in 4:1 molar ratio. Preclinical studies have demonstrated that this combination of tegafur and uracil is associated with higher plasma levels of 5-FU compared
with tegafur treatment alone (29,30). Furthermore, this difference in 5-FU plasma levels was associated with greater antitumor activity (29,30). Two phase III studies comparing UFUR/leucovorin with 5-FU/leucovorin demonstrated that the response rate, time to progression and overall survival time were similar between the two regimens, with an overall survival of 12-13 months. However, diarrhea, nausea and vomiting, stomatitis and mucositis, and myelosuppression occurred significantly less frequently in the UFUR/leucovorin compared with the 5-FU/leucovorin group (27,31).

In a number of Japanese and Taiwanese studies, UFUR was administered in favor of 5-FU as an agent in combination chemotherapy; for example, in the FOLFIRI regimen, UFUR replaced 5-FU in combination with irinotecan (9,32,33). In these previous studies, the dose of irinotecan (70, 80, 100, 150, 180, 220 or 350 mg/m²) and the interval between the doses (for example, once weekly, once every 2 weeks and once every three weeks) varied broadly.

Although leucovorin appears to enhance the antitumor efficacy of 5-FU in the treatment of metastatic colorectal cancer, the healthcare worker may select UFUR administration with or without leucovorin. Thus, we aimed to determine the optimal dosing schedule and dosage for the TEGAFIRI regimen in the metastatic colorectal setting. Previous studies indicated that the TEGAFIRI regimen was well-tolerated (11,14) and, by modulating with leucovorin, the TEGAFIRI and TEGAFOX regimens demonstrated comparable efficiency and safety (10,12,13).

The aim of the pilot study was to evaluate the response rates of different regimens of combination chemotherapy employing TEGAFIRI with or without leucovorin for patients with recurrent or metastatic colorectal cancer. To decrease bias, all of the patients in the present study were analyzed by a single health worker. The results indicated that groups II and III exhibited similar response rates and were preferable to the regimen employed in group I. Furthermore, the response rates in groups II and III were similar to those of previous studies, which employed the FOLFIRI regimen with leucovorin (11,13,34). Grade III/IV diarrhea, alopecia and hematologic side effects were acceptable and similar in the three groups. Subsequently, the present study evaluated the response of a larger cohort to 150 mg/m² irinotecan every two weeks with continuous UFUR and leucovorin without interruption, and demonstrated similar side-effect and survival benefits, such that continuous UFUR and leucovorin without interruption appeared to be essential for improved patient survival.

This type of combination chemotherapy is advantageous as it requires no hospital admission, has a shorter injection time, and, by modulating with leucovorin, the TEGAFIRI and TEGAFOX regimens demonstrated comparable efficiency and safety (10,12,13).

Despite the possibility of the abovementioned disadvantages occurring, the pilot and the present study indicated similar response rates to the TEGAFIRI regimen compared with previous reports in the literature. For example, TEGAFIRI results in satisfactory response rates and patients report tolerable side effects. Furthermore, continuous UFUR administration without interruption appeared to result in improved outcomes compared with the intermittent administration of UFUR, and leucovorin is an essential component of the treatment regimen.

In conclusion, TEGAFIRI combination chemotherapy is a satisfactory alternative therapy to the FOLFIRI regimen, producing acceptable response rates for recurrent or metastatic colorectal cancer patients. The present study recommends that TEGAFIRI should be administered in combination with leucovorin and oral UFUR administration should not be interrupted during treatment.

References

1. Yeh KH, Cheng AL, Lin MT, et al: A phase II study of weekly 24-hour infusion of high-dose 5-fluorouracil and leucovorin (HDFL) in the treatment of recurrent or metastatic colorectal cancers. Anticancer Res 17: 3867-3871, 1997.
2. Merenmies PH, Lavrenkova K and Cohen Y: Phase II study of weekly high-dose fluorouracil in previously treated patients with metastatic colorectal cancer. J Chemother 12: 183-185, 2000.
3. Cunningham D, Pye S, James RD, et al: Randomized trial of irinotecan plus supportive care versus supportive care alone after fluorouracil failure for patients with metastatic colorectal cancer. Lancet 352: 1413-1418, 1998.
4. Douillard JY, Cunningham D, Roth AD, et al: Irinotecan combined with fluorouracil compared with fluorouracil alone as first-line treatment for metastatic colorectal cancer: a multicenter randomized trial. Lancet 355: 1041-1047, 2000.
5. Rothenberg ML, Cox JV, Devore RF, et al: A multicenter phase II trial of weekly irinotecan (CPT-11) in patients with previously treated colorectal carcinoma. Cancer 85: 786-795, 1999.
6. Van Cutsem E, Pozzo C, Kikuchi S, et al: A phase II study of irinotecan plus leucovorin combined with five days bolus of 5-fluorouracil and leucovorin in first-line chemotherapy of metastatic colorectal cancer. Ann Oncol 9: 1199-1204, 1998.
7. Ota K, Taguchi T and Kimura K: Report on nationwide-pooled data and cohort investigation in UFT phase II study. Cancer Chemother Pharmacol 22: 333-338, 1988.
8. Mallet-Martino M and Martino R: Clinical studies of three oral prodrugs of 5-fluorouracil (capecitabine, UFT, S-1): a review. Oncologist 7: 288-321, 2002.
9. Kusunoki M, Yanagisawa K, Noda M, et al: Results of pharmacokinetic and toxicological studies of tegafur, leucovorin and oral UFUR in mice. Cancer 69: 1371-1377, 1992.
10. Mackay HJ, Hili M, Travers W, et al: Phase I/III study of oral uracil/tegafur (UFT), leucovorin, and irinotecan in patients with advanced colorectal cancer. Ann Oncol 14: 1264-1269, 2003.
11. Mibaru T, Tanaka S, Futami K, et al: Phase I/I study of irinotecan and UFT for advanced or metastatic colorectal cancer. Anticancer Res 27: 2673-2678, 2007.
12. Bajetta E, Bartolomeo MD, Buzzoni R, et al: Uracil/tegafur/leucovorin combined with irinotecan (TEGAFIRI) or oxaliplatin (TEGAFOX) as first-line treatment for metastatic colorectal cancer patient: results of randomized phase II study. Br J Cancer 95: 439-444, 2007.
13. Delord JP, Bennouna J, Artru P, et al: Phase III study of UFT with leucovorin and irinotecan (TEGAFIRI): first-line therapy for metastatic colorectal cancer. Br J Cancer 97: 297-301, 2007.
14. Ogata Y, Mori S, Ishibashi Y, et al: Metronomic chemotherapy using weekly low-dosage CPT-11 and UFT as postoperative adjuvant therapy in colorectal cancer at high risk to recurrence. J Exp Clin Cancer Res 26: 475-482, 2007.
15. Hsu TC: Combination chemotherapy with irinotecan and oral uracil-tegafur for recurrent or metastatic colorectal cancer - results of different regimens. J Chinese Oncol Soc 25: 254-261, 2009 (In Chinese).
16. Oken MM, Creech RH, Tormey DC, et al: Toxicity and response criteria of the Eastern Cooperative Oncology Group. Am J Clin Oncol 5: 649-655, 1982.
17. Schag CC, Heinrich RL and Ganz PA: Karnofsky performance status revisited: reliability, validity, and guidelines. J Clin Oncology 2: 187-193, 1984.

18. Dukes CE: The surgical pathology of rectal cancer. J Clin Pathol 2: 95-98, 1949.

19. Eisenhauer EA, Therasse P, Bogaerts J, et al: New response evaluation criteria in solid tumours: revised RECIST guideline (version 1.1). Eur J Cancer 45: 228-247, 2009.

20. Cancer Therapy Evaluation Program: Common Toxicity Criteria, Version 2.0, April 30, 1999. http://prevention.cancer.gov/files/clinical-trials/common-toxicity-criteria.pdf. Accessed November 22, 2014.

21. Jemal A, Tiwari RC, Murray T, et al; American Cancer Society: Cancer statistics, 2004. CA Cancer J Clin 54: 8-29, 2004.

22. DeVita VT Jr, Hellman S and Rosenberg SA (eds): Cancer Principles and Practice of Oncology. 5th edition. Lippincott-Raven, Philadelphia, PA, pp 437-452, 1997.

23. Colorectal Cancer: Estimated Incidence, Mortality and Prevalence Worldwide in 2012. GLOBOCAN 2012. http://globocan.iarc.fr/old/FactSheets/cancers/colorectal-new.asp. Accessed November 22, 2014.

24. Colorectal Cancer Facts & Figures 2014-2016. American Cancer Society 2014. http://www.cancer.org/acs/groups/content/documents/document/acspc-042280.pdf. Accessed November 22, 2014.

25. Hsiao SC, Lin JF, Chuang MT, Lee YA and Wu DL: Retrospectively comparative evaluation of the first- and second-line chemotherapy with campto and oxaliplatin combined with oral tegafur/uracil (UFT)/leucovorin (LV) in patients with metastatic colorectal cancer. Int Surg 94: 298-303, 2009.