Bcl-2 expression significantly correlates with thymidylate synthase expression in colorectal cancer patients

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Abstract

AIM: To examine the expression of thymidylate synthase (TS) and oncoprotein Bcl-2 in advanced colorectal cancer (CRC) patients, and to determine their mutual relationship, association to therapeutic response and impact on disease outcome.

METHODS: Tumor samples from 67 patients with CRC, who were treated at advanced stage with either irinotecan alone or in combination with 5-fluorouracil/leucovorin, were analyzed for expression of TS and Bcl-2 using immunohistochemistry.

RESULTS: A significant linear correlation between lower expression levels of Bcl-2 and lower levels of TS expression was found ($P = 0.033$). Patients with high levels of both TS and Bcl-2 expression had a significantly longer disease-free survival (DFS) ($42.6$ mo vs $5.4$ mo, $n = 25$) than those with low TS/Bcl-2 index ($P = 0.001$). Tumors with low levels of both TS and Bcl-2 were associated with a longer survival with metastasis (WMS) interval in the whole patients group ($n = 67$, $P = 0.035$). TS/Bcl-2 index was not significantly related to disease-specific survival.

CONCLUSION: The present data suggest that CRC patients with low TS/Bcl-2 demonstrate a significantly shorter DFS and longer WMS.

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Key words: Thymidylate synthase; Bcl-2; Colorectal cancer; Disease-free survival; Survival with metastases

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INTRODUCTION

Colorectal cancer (CRC) is the second most frequent cancer in Europe in 2004, responsible for 13% (376 400) of all incident cancer cases. It is also the second most frequent cause of cancer mortality in Europe, with an annual mortality of 11.9%, 203 700 annual deaths[1]. In the early stages, CRC is often a curable disease, but the overall prognosis is determined by the extent of local and particularly metastatic tumor spread. However, disease outlook is relatively poor for advanced disease and thus is a significant cause of worldwide cancer-related mortality[1]. For locally advanced and metastatic CRC, fluoropyrimidine, 5-fluorouracil (5-FU) has been the standard cytostatic drug for the last 50 years, in recent years used as modulated by leucovorin and in combination with irinotecan or oxaliplatin.

Fluoropyrimidine metabolites form a covalent complex with thymidylate synthase (TS). Formation of this complex prevents biosynthesis of intracellular thymidylate, which is essential for DNA biosynthesis. TS expression has been shown to be an independent prognostic factor in several other cancers. Higher TS levels in hepatic metastases and resection margin are independent predictors of disease progression and survival in patients with metastatic CRC[2]. Comparable results have been reported in other tumors e.g. gastric[3], cervical[4], ovarian, and head and neck cancers, for which TS+ tumors have demonstrated significantly worse outcome as compared to TS-negative tumors.

Increased expression of the proto-oncogene Bcl-2, a 24-kDa intracellular membrane protein that is able to inhibit programmed cell death without affecting cell proliferation, has been reported in gastrointestinal adenocarcinoma and its precursor lesions[5][6]. Bcl-2...
has been shown to prolong cell survival by inhibiting apoptosis in several cell types\[7\]. Abnormal activation of the Bel-2 gene appears to be an early event in colorectal tumorigenesis\[6\].

In this study, we examined the expression of TS and the oncoprotein Bcl-2 in locally advanced and metastatic CRC and determined their inter-relationships, as well as their impact on patient survival.

MATERIALS AND METHODS

Patients, treatment and follow-up
A series of 67 patients were diagnosed and treated for Stage II, III, IV CRC at the Department of Oncology and Radiotherapy, Turku University Hospital (TUH) and six other hospitals in the same hospital district, between January 1996 and August 2003. The key clinical characteristics of the patients are summarized in Table 1.

At the time of diagnosis, 11 patients had stage II, 14 had stage III and 42 had stage IV disease. When patients developed metastases or inoperable local recurrence, they were entered into the chemotherapy protocol. In the protocol, patients received one of two treatment regimens; 18 received irinotecan alone and 49 received a combination of irinotecan, 5-FU and folinic acid (FA) as first line treatment for metastatic disease. Irinotecan (350 mg/m$^2$) was administered as a 60-90 min intravenous (i.v.) infusion every 3 wk. In the combination regimen, irinotecan (180-210 mg/m$^2$) was administered as 60-90 min intravenous infusion and 5-FU (500 mg/m$^2$, i.v. bolus) modulated with folinic acid (FA) (60 mg/m$^2$, i.v. bolus). The 5-FU/FA administrations were repeated again on the following day. The cycle was repeated every 2 wk\[9\]. The mean duration of chemotherapy was 6.3 mo (SD, 3.4 mo). Treatment was continued until disease progression, or occurrence of unacceptable toxicity.

The patients were prospectively followed-up until the end of March 2007; mean follow-up time from diagnosis was 34.4 mo (± 26.2 mo). We used three endpoints to calculate the patient survival: (1) disease-free survival (DFS), which was calculated in 26 patients with stage II or III disease at diagnosis; (2) overall disease-specific survival (DSS); and (3) survival with metastases (WMS). DFS is the time from diagnosis to the appearance of metastatic disease and relevant only for those patients with radically operated stage II and III patients at the time of diagnosis ($n = 25$). DSS is the time from diagnosis to death or to the time point when last seen alive at the clinic, and was calculated for all patients in the study. WMS was calculated from the date of recording the appearance of disease recurrence/metastases at the clinical visit, until to death or to the time point when last seen alive.

The study was approved by the Ethical Committee and was conducted in accordance with the Declaration of Helsinki. Samples were collected with the endorsement of the National Authority for Medico-legal Affairs.

Immunohistochemical detection of TS and Bcl-2 expression
Sixty-seven formalin-fixed, paraffin-embedded primary tumors were obtained from 67 patients. Sections were cut serially at 5 μm for routine hematoxylin and eosin staining and for immunohistochemical analysis. An experienced pathologist confirmed all histological diagnoses.

TS expression was studied immunohistochemically using monoclonal antibody (Mouse Clone TS 106) from Zymed Laboratory. Bcl-2 protein expression was studied using anti-Bcl-2 monoclonal mouse antibody, which recognizes a peptide comprising amino acids 41-54 of the human Bcl-2 protein (Clone 124, DAKO A/S, Glostrup, Denmark). Signal detection was performed using the streptavidin-biotin method (Vectastain ABC kit). Formalin-fixed, paraffin-embedded sections were deparaffinized in xylene, rehydrated in graded alcohol, immersed in 0.01 mol/L citrate buffer (pH 6.0), heated in a domestic microwave oven at full power for 2 × 5 min, and left in the buffer to cool to room temperature. The sections were incubated in 0.3% hydrogen peroxide for 20 min to block endogenous peroxidase activity. Incubation with the primary antibody diluted in 1%
bovine serum albumin/Tris-buffered saline, TS (1:25)
and Bcl-2 (1:50), were carried out overnight in a
humid chamber at 4°C. The following day, the slides
were washed and incubated first with the biotinylated
secondary antibody (30 min, 20°C), then with avidin-
biotin-peroxidase complex (30 min, 20°C). Positive
staining was visualized with 3,3’ diamino benzidine (DAB)
substrate solution and the sections were counterstained
with Mayer’s hematoxylin. As negative controls, slides
were processed with the omission of the primary antibody.

**Evaluation of TS and Bcl-2 expression**

Expression of TS and Bcl-2 was assessed by an
observer blinded to the clinical data. The slides were
first screened for an overview of the general staining
pattern. Four pictures of each slide, covering most of
the tumor area, were taken with a light microscope
(4 × magnification) connected to a camera and AnalySIS
v 3.00 software (Soft Imaging System GmbH, Munster,
Germany). Expression of TS in the four pictures was
analyzed using Imaging Research Inc., St. Catharine’s,
Ontario, Canada), which detected the brown color of
the positively stained tumor cells and counted the area
of those cells in pixels, and also counted in pixels the
total tumor area. The percentage of positively stained
tumor cells from the whole tumor area was counted and
used in further analysis. This method of evaluating TS
expression was able to distinguish between the presence
of many cells expressing low amounts and a few cells
expressing high amounts of TS, such that the percentage
of TS expression reflected total TS expression in the
tumor, which may be more relevant biologically.

For statistical purposes, expression profiles of each
marker were treated as dichotomous variables, where
tumors with negative or weak expression of Bcl-2 was
one category (reduced expression), and all those with
moderate or strong expression were grouped into the
second category. For TS, we used median values as cut-
off to build up the dichotomous variable of low- and
high TS expression. In addition, combined TS-Bcl-2
indices were created, using the dichotomous Bcl-2
variables and TS variables (median cut-offs), resulting in
four possible combinations of TS/Bcl-2: low/low (L/L); low/high (L/H); high/low (H/L); and high/high (H/H).
Finally, these were converted to a 3-class index as
follows: class 1, TS/Bcl-2, L/L; class 2, TS/Bcl-2, L/H
or H/L; and class 3, TS/Bcl-2 H/H.

**Statistical analysis**

Statistical analyses were performed using the SPSS*
(SPSS, Inc., Chicago, IL, USA) and STATA (Stata Corp.,
TX, USA) software packages (SPSS for Windows,
version 14.0.1 and STATA/SE 10.1). Frequency tables
were analyzed using the \( \chi^2 \) test, with likelihood ratio (LR)
or Fisher’s exact test being used to assess the significance
of the correlation between categorical variables.
Differences in the means of continuous variables were
analyzed using non-parametric tests (Mann-Whitney or
Kruskal-Wallis tests) for two and multiple independent
samples, respectively. ANOVA was only used for deriving
the mean values in each stratum. Univariate survival
(life-table) analysis for the outcome measure, DSS, DFS
and WMS was based on Kaplan-Meier method, and the
groups were compared with the log-rank (Mantel-Cox)
test. In all tests, \( P < 0.05 \) was regarded as statistically
significant.

**RESULTS**

Expression of Bcl-2 and staining of TS are shown in
Figure 1. Bcl-2 index and TS index (absolute values)
were significantly correlated \( (r = 0.286, \text{Spearman rho,}
\( P = 0.019) \). Similarly, using the median-cut off
values, there was a significant correlation between Bcl-2
and TS expression profile \( (P = 0.039, \text{Spearman rho or } P =
0.037, \chi^2) \). In pair-wise comparison, individual samples
did not significantly deviate in their Bcl-2 and TS
expression profile (median cut-off; \( P = 0.841, \text{Wilcoxon}
signed ranks test) indicating that in individual tumors,
co-detection of H/H and L/L of both markers was a
frequent occurrence.

The combined TS-Bcl-2 index was evaluated in
relation to all clinical variables recorded, including the
response to treatment and survival (DSS, DFS and
WMS). Interestingly, for the first time, a significant
correlation between TS/Bcl-2 index and DFS was
observed. Among stage II and III disease \( (n = 25) \),
tumors with H/H profile of TS and Bcl-2 (class 3) were
associated with substantially longer DFS (42.6 mo) than
those with L/L (5.4 mo) or those with H/L or L/H
profile (20.7 mo; \( P = 0.031, \text{Kruskal-Wallis} \)), and this
was even more evident in life-table analysis \( (P = 0.001, \text{Mantel-Cox} \); Figure 2A).

There was a close correlation between TS/Bcl-2
expression and WMS; tumors with L/L profile of TS
and Bcl-2 (class 1) were associated with a longer WMS
(31.9 mo) than those with H/H (25.7 mo) or H/L and
L/H (18.9 mo) in the whole series \( (n = 67, P = 0.076, \text{Kruskal-Wallis}
and \( P = 0.092 \text{Mantel-Cox} \); Figure 2B). There was no such
difference in DSS among the three TS/Bcl-2 expression profiles, however. TS/Bcl-2
expression was not significantly associated with any
other clinicopathological variables, including age, sex,
TNM status, grade, stage or carcinoembryonic antigen
(CEA) levels.

**DISCUSSION**

The treatment of CRC has become increasingly
complex over recent years. With the emergence of new
chemotherapy drugs and targeted agents, there has been
a major improvement in the prognosis of patients with
metastatic CRC. The identification of prognostic and
predictive markers is clinically important, because CRC
is a heterogeneous disease with various biological and
clinical characteristics.

The present study analyzed the combined TS and
Bcl-2 expression in CRC, with particular reference
to disease outlook. An increase in TS levels has been
suggested to be an important mechanism of resistance to fluoropyrimidine-based chemotherapy. In several studies, TS is reported to be a predictor of both survival in CRC and response to 5-FU therapy, with higher levels being associated with poorer prognosis and response to therapy. This suggests that TS levels are not only of importance in predicting the natural course of the disease, but may also predict the sensitivity of cancer cells to 5-FU, which is widely used both in the adjuvant setting and in the treatment of metastases.

The role of the Bcl-2 family of proteins in chemoresponse has been evaluated extensively in vitro models. Over-expression of Bcl-2 and Bcl-XL has been shown to induce drug resistance. In relation to treatment, it has been demonstrated that elevated levels of Bcl-2 protein confers cytotoxic drug resistance to tumor cell lines. As Bcl-2 blocks apoptosis in vitro and thus contributes to malignant cell accumulation, its over-expression is expected to be associated with more aggressive tumor biology. Indeed, genetic alteration of the Bcl-2 gene located on chromosome 18 is considered to be a key process in the pathogenesis and chemoresistance of human tumors, such as follicular lymphoma.

In our previous analysis of Bcl-2 expression in a subset of these tumors, we found a weak association of lower levels of Bcl-2 expression with longer overall survival. In another small series, we described that lower levels of Bcl-2 expression were significantly associated with lower levels of TS expression.

The present study clearly confirmed this observation in a larger series of CRC patients, for which a significant linear correlation was established between the two markers. High expression of
TS (above the median) significantly correlated with higher Bcl-2 expression levels. In our previous study, some evidence has suggested that higher levels of TS and Bcl-2 are associated with shorter overall survival, implicating that elevated levels of TS and Bcl-2 may confer cytotoxic drug resistance among these patients.

To the best of our knowledge, our study shows for the first time the relationship between these two key molecules in a group of patients with locally advanced or metastatic CRC receiving similar treatment. As to the patient survival, there was a significant correlation of TS/Bcl-2 expression with DFS, in that the patients with high TS/Bcl-2 index had a longer DFS (Figure 2A). No such effect was shown for DSS, which was not significantly different among the patients with low- and high TS/Bcl-2 indices.

To conclude, the present data suggest that patients with CRC whose tumors have high TS and Bcl-2 demonstrate a significantly longer DFS and shorter WMS, as compared to patients with low expression of these markers.

COMMENTS

Background
Thymidylate synthase (TS) is reported to be a predictor of both survival in colorectal cancer (CRC) and response to 5-FU therapy, with higher levels being associated with poorer prognosis and response to therapy. Elevated levels of Bcl-2 protein also confer cytotoxic drug resistance to tumor cell lines. We here examined the expression of TS and oncoprotein Bcl-2 in advanced CRC patients, and determined their mutual relationship, association to therapeutic response and impact on disease outcome.

Research frontiers
This study represents a translational study in which a clinical series of CRC samples was analyzed for two important molecular markers: TS and Bcl-2. Accordingly, this study represents a combined molecular analysis and a clinical study, whereby some key molecular pathways were analyzed and their relevance to clinical data was assessed in a series of 67 CRC patients with well-characterized treatment history and long-term follow-up data.

Innovations and breakthroughs
To the best of our knowledge, this is the first study to show the relationship between these two key molecules in a group of patients with locally advanced or metastatic CRC receiving similar treatment. As to the patient survival, there was a significant correlation of TS/Bcl-2 expression with DFS, in that the patients with high TS/Bcl-2 index had a longer DFS.

Applications
The present data suggest that patients with CRC whose tumors have high TS and Bcl-2 demonstrate a significantly longer DFS and shorter WMS, as compared to patients with low expression of these markers. This information should have potential clinical implications in the treatment of these patients.

Peer review
The importance of this paper to the reader resides in the fact that this study is believed to be the first to suggest that CRC patients with low TS/Bcl-2 demonstrate a significantly shorter DFS, but keep alive longer with a metastatic disease. In relevant cases, one might consider utilizing this type of molecular marker data in more individualized tailoring of the appropriate therapies.

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