Modeling Efficacy of Bevacizumab Treatment for Metastatic Colon Cancer

Rezwan Islam1, Po-Huang Chyou2, and James K Burmester3

1. Department of Oncology/Hematology, Marshfield Clinic Weston Center, Marshfield, WI, USA; 2. Biomedical Informatics Research Center, Marshfield Clinic Research Foundation, Marshfield, WI, USA; 3. Clinical Research Center, Marshfield Clinic Research Foundation, Marshfield, WI, USA.

Corresponding author: Rezwan Islam, MD. Marshfield Clinic Weston Center, Department of Oncology/Hematology, 3501 Cranberry Blvd, Weston, WI 54476. Phone: 715-393-1400 Fax: 715-389-5757 Email: islam.rezwan@marshfieldclinic.org.

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Abstract

Purpose: Bevacizumab, an FDA-approved adjuvant treatment for metastatic colon cancer, has extended survival for many patients. However, factors predicting response to treatment remain undefined.

Patients and Methods: Relevant clinical and environmental data were abstracted from medical records of 149 evaluable patients treated with bevacizumab for metastatic colon cancer at a multi-specialty clinic. Tumor response was calculated from radiologic reports using Response Evaluation Criteria in Solid Tumors (RECIST) criteria and verified by oncologist review. Patients with at least one occurrence of complete or partial response or stable disease were classified as responders; those exhibiting progressive disease were classified as non-responders.

Results: Univariate analysis demonstrated that blood in stool (P<0.05), unexplained weight loss (P<0.05), primary colon cancer site (P<0.05), chemotherapy treatment of primary tumor site (P<0.05), and adenocarcinoma versus adenoma subtype (P<0.05) was associated with tumor responsiveness. Factors remaining statistically significant following multivariate modeling included adenocarcinoma as tumor cell type versus other adenocarcinoma subtypes (OR=6.35, 95% CI: 1.08-37.18), chemotherapy treatment applied to primary tumor (OR=0.07, 95% CI: 0.0-0.76), tumor localization to cecal/ascending colon (OR=0.061, 95% CI: 0.006-0.588), and unexplained weight loss (OR=0.1, 95% CI: 0.02-0.56). Chemotherapy treatment of primary tumor, unexplained weight loss, and cecal/ascending localization of the tumor were associated with poorer outcomes. Adenocarcinoma as cell type compared to other adenocarcinoma subtypes was associated with better response to bevacizumab treatment.

Conclusion: Results suggest that response to bevacizumab therapy may be predicted by modeling clinical factors including symptomology on presentation, tumor location and type, and initial response to chemotherapy.

Key words: Bevacizumab; Colon cancer; Monoclonal antibodies; Prognosis; Vascular endothelial growth factor/VEGF.

INTRODUCTION

Colon cancer is the third most common cancer diagnosed in the United States (excluding skin cancer), with about 101,000 new cases diagnosed during 2011. Colon cancer is the second leading cause of cancer death in the US with about 50,000 deaths occurring annually.
In colon cancer increased expression of vascular endothelial growth factor (VEGF) occurs concurrently with increasing vascular density of an invasive tumor in the context of metastasis, and recurrence. Bevacizumab is a recombinant monoclonal antibody that specifically blocks the activity of human VEGF. VEGF is a potent, specific stimulator of angiogenesis supporting vascularization of growing tumors. In the absence of VEGF tumors become hypoxic. Bevacizumab treatment causes prolonged hypoxia resulting in vascular cell death primarily through apoptosis and decreased blood supply to the tumor.

Clinical trials demonstrated an increase in median survival for colon cancer patients when treated with bevacizumab as an adjunct to standard chemotherapy. The addition of intravenous bevacizumab to bolus fluorouracil/folinic acid (FU/FA), irinotecan/bolus fluorouracil/folinic acid (IFL), irinotecan/infusional fluorouracil/folinic acid (FOLFIRI), infusional fluorouracil/folinic acid/oxaliplatin (FOLFOX), capcitabine/oxaliplatin (XELOX), or bolus fluorouracil/folinic acid/oxaliplatin has shown efficacy as a first-line therapy for metastatic colorectal cancer. In most randomized controlled trials, the addition of bevacizumab to standard treatment increased median overall survival by about 5 months for first-line therapy. The increase in survival in second-line therapy was about 2 months.

Adverse drug events (ADE) such as gastrointestinal perforation, wound healing complications, and hemorrhage are increased in patients treated with bevacizumab. Further, the cost of bevacizumab is very high due to the extensive effort required to produce an injectable product. Thus, methods are needed to characterize which patients will derive the greatest benefit from bevacizumab treatment and distinguish them from non-responders and/or patients at ADE risk following exposure.

To investigate personal and medical factors that may influence which patients will have a positive response to bevacizumab, we collected data on 149 patients treated with bevacizumab for metastatic colon cancer.

PATIENTS AND METHODS

Subject Recruitment

The study was approved by the Marshfield Clinic Research Foundation’s Institutional Review Board. An electronic search of the Marshfield Clinic electronic medical record identified 229 patients who had been treated with bevacizumab as part of first-line or second-line chemotherapy for metastatic colon cancer since 2002. Further screening of subjects to confirm eligibility for inclusion in the study based on inclusion criteria (metastatic colon cancer treated with bevacizumab) and verification of sufficient evaluable data (radiological data, and physician notes) determined 149/229 electronic met criteria for inclusion in the analysis.

Data Abstraction

Data abstracted from the medical records of patients validated as meeting inclusion criteria following screening (n=149) included: height and weight (to determine body mass index), primary site of colon cancer, co-morbidities (including other cancers, heart disease, cardiovascular disease, diabetes mellitus, Crohn’s disease, diverticulitis, and irritable bowel syndrome), unexplained or severe weight loss (defined as loss of 10% or greater of body weight), current age, age at colon cancer diagnosis, family history of colon cancer, stage and grade of primary tumor, number, site, stage, and grade of metastases, imaging scan results, treatment of the primary tumor including surgery and/or chemotherapy, and/or radiation, treatment of the primary metastasis including surgery, and/or chemotherapy, and/or radiation, dates and doses of chemotherapy, dates and doses of bevacizumab therapy, and all ADE (gastrointestinal perforation, wound healing complications, hemorrhage). Complete data were not consistently available for all 229 patients initially identified by electronic screening for all variables since each patient presented as an individual case and only patients who met inclusion criteria and had sufficient evaluable data were included in the analysis (n=149). Tumor response was calculated from radiologic images using Response Evaluation Criteria in Solid Tumors (RECIST) guidelines for evaluating tumor response in oncology trials and compared to oncologists’ assessment captured in the patients’ medical records. Records were systematically checked following each bevacizumab treatment to determine if new radiographs had been taken and were available for RECIST evaluation. Response criteria were classified in compliance with RECIST categorization as follows: complete response (disappearance of all lesions), partial response (30% decrease of tumor dimension from baseline measures), stable disease (neither response nor progression), and progressive disease (20% increase). Patients were categorized as responders if they exhibited complete or partial response or exhibited stable disease at least once following response to any treatment that included bevacizumab administration. For analytical purposes, complete response, partial response, and stable disease were grouped together as responders reflecting disease control, while patients...
exhibiting progressive disease were grouped as non-responders (poorer outcome).\textsuperscript{12}

**Statistical Analysis**

Colon cancer patients were divided into two groups based on the positive response status (yes/no) to bevacizumab. In the univariate analysis, the descriptive statistics (for any continuous measurements: mean, standard deviation, median, and the range; for any categorical measurements: frequency and percentage) were reported for each of the patient’s attributes such as body mass index, age at diagnosis, family history of colon cancer, and other clinical characteristics. These risk factors were compared between the two groups using the t-test for normally distributed data, the Wilcoxon rank sum test for skewed data (the adverse event count), and the Fisher’s exact test for cells with a value <5 or the Chi-square test for categorical data. Odds ratio (OR) and corresponding 95% confidence interval (CI) for the status of positive response to bevacizumab treatment were derived from unconditional logistic regression modeling approach using each of the above-named risk factors as the predictor variable. A \( P \)-value of <0.05 was considered statistically significant. In multivariate analysis, a stepwise (forward selection and backward elimination) logistic regression modeling was performed to determine the set of important predictors, using the status of positive response to bevacizumab treatment as the dependent variable and risk factors that showed a significant \( P \)-value in the univariate analysis as the independent variables. All statistical analyses were carried out using commercially available statistical software package (SAS).

**RESULTS**

A total of 229 patients who had been treated with bevacizumab for metastatic colon cancer were evaluated for inclusion in the study. Of these, 149 met inclusion criteria, and had sufficient evaluable data available to qualify them for inclusion in the analyses (see Figure 1.) Table 1 summarizes demographic data for subjects grouped in accordance with response status of the subjects (i.e., responder, defined as having at least one full or partial response characterized as \( \geq 30\% \) reduction in tumor size from baseline or stable disease in response to bevacizumab treatment, versus non-responder). No statistically significant differences between individuals meeting criteria as responders and non-responders were observed for racial background, ethnic background, gender, or smoking status. At the time of enrollment into the study, 135/149 patients were living and provided informed consent, and 14/149 patients were deceased.

![Figure 1. Flow diagram of study enrollment.](http://www.jcancer.org)
Table 1. Positive tumor response related to patient demographics.

| Characteristics          | Positive Tumor Response | P-value* |
|--------------------------|-------------------------|----------|
|                          | No. (%)                 | Yes (%)  |          |
| Racial background        |                         |          |          |
| Asian                    | 1 (7)                   | 0 (0)    | .1044    |
| White or Caucasian       | 13 (93)                 | 129 (96) |          |
| Other                    | 0 (0)                   | 6 (4)    |          |
| Hispanic or Latino       | 0 (0)                   | 3 (2)    | 1.0000   |
| Not Hispanic or Latino   | 11 (100)                | 119 (98) |          |
| Gender                   |                         |          | .2657    |
| Male                     | 5 (36)                  | 72 (53)  |          |
| Female                   | 9 (64)                  | 63 (47)  |          |
| Smoking Status           |                         |          | .6448    |
| Current                  | 1 (8)                   | 27 (20)  |          |
| Ex-Smoker                | 6 (46)                  | 56 (42)  |          |
| Never Smoker             | 6 (46)                  | 49 (37)  |          |

*P-value was derived from Fisher's Exact Test.

To determine whether existing medical conditions, family history of cancer, treatment for colon cancer (surgery, radiation, or chemotherapy), and adverse drug responses are associated with a positive tumor response, we collected this information from the patients’ medical records in sequential fashion corresponding to each drug treatment. No differences between responders and non-responders were noted for any other factors evaluated including symptomology, comorbidities, tumor characteristics, or genetic factors, with the exception of those shown in Table 2. Significant differences between responders and non-responders detected in univariate analysis included blood in the stool (P<0.05), unexplained weight loss (P<0.05), primary site of colon cancer (cecal/ascending) (P<0.05), adenocarcinoma type vs. all other adenocarcinoma subtypes (P<0.05), and chemotherapy treatment of the primary tumor (P<0.05) (see Table 2). Blood in the stool correlated with response in the subset of patients with adenocarcinoma as tumor type (P<0.05). No statistical difference was noted between responders and non-responders for other treatment approaches to primary tumor including radiation therapy or surgery.

Multivariate stepwise logistic regression analysis was next applied to determine the set of important predictors for positive response to bevacizumab treatment. Among the variables modeled by multivariate analysis were those that had achieved statistical significance by univariate analysis with the exception of blood in stool. Inclusion of blood in stool was precluded because no patients among the non-responders exhibited this characteristic and thus comparison was not possible. Blood in stool was defined as observation of blood in stool by the patient, and in many cases, this was the symptom that brought the patient to the clinic for further evaluation. Variables that remained significant following multivariate regression analysis included adenocarcinoma as tumor cell type compared to other adenocarcinoma subtypes (OR=6.35, 95% CI:1.08-37.18), chemotherapy treatment approach to the primary tumor (OR=0.07, 95% CI: 0.00-0.76), tumor localization to cecal/ascending colon (OR=0.61, 95% CI: 0.06-0.588), and unexplained weight loss (OR=0.1, 95% CI: 0.02-0.56).

Table 2. Positive tumor response related to patient medical condition with statistical differences between responders and non-responders: univariate analysis.

| Characteristics                      | Positive Tumor Response | P-value* |
|---------------------------------------|-------------------------|----------|
| Blood in the stool                    |                         |          |
| Yes                                   | 0 (0)                   | 62 (100) | .0007    |
| No†                                   | 11 (16)                 | 58 (84)  |          |
| Unexplained weight loss               |                         |          |
| Yes                                   | 7 (17)                  | 35 (83)  | .0371    |
| No†                                   | 4 (4)                   | 85 (96)  |          |
| Cecal/ascending                       |                         |          |
| Yes                                   | 10 (17)                 | 50 (83)  | .0072    |
| No†                                   | 3 (3)                   | 85 (97)  |          |
| Chemotherapy treatment of primary tumor|                         |          |
| Yes                                   | 12 (13)                 | 77 (87)  | .0456    |
| No†                                   | 2 (3)                   | 58 (97)  |          |

*P-value was derived from Fisher’s Exact Test.

Table 3. Multivariate Stepwise Logistic Regression Analysis for Selected Predictor Clinical Characteristics in Association with Positive Tumor Response among Colon Cancer Patients.

| Selected clinical characteristics* | OR‡ | CI§ | P-value¶ |
|-----------------------------------|-----|-----|----------|
| Adenocarcinoma                    |     |     |          |
| Yes                               | 6.35| 1.08-37.18 | 0.03    |
| No†                               | 1.00|     |          |
| Chemotherapy                      |     |     |          |
| Yes                               | 0.069| 0.006-0.757 | 0.0176  |
| No†                               | 1.00|     |          |
| Tumor site: Cecal/ascending colon |     |     |          |
| Yes                               | 0.06|     |          |
| No†                               | 1.00| 0.006-0.588 | 0.0008  |
| Unexplained weight loss           |     |     |          |
| Yes                               | 0.10| 0.02-0.56 | 0.006   |
| No†                               |     |     |          |

*Other clinical characteristics included in the multivariate stepwise logistic regression analysis were: blood in stool (yes vs. no), losing weight for no reason (yes vs. no), and primary site of colon cancer (cecal/ascending, yes vs. no). †Referent group. ‡OR = odds ratio. §CI = 95% confidence interval. ¶P-value was derived based on Wald chi-square test.
DISCUSSION

Efficacy of first-line/second-line bevacizumab treatment in combination with chemotherapy to extend progression-free survival and overall survival for many patients with metastatic colon cancer has been established. At the same time, an important subset of patients do not benefit from bevacizumab therapy. To date, few markers have been explored to distinguish potential responders from non-responders with respect to treatment with bevacizumab. Markers of response to bevacizumab that have previously been reported for metastatic colon cancer include carbonic anhydrase, visceral fat area, and hypertension.

Studies also suggest that heritable genetic polymorphisms may contribute to bevacizumab responsiveness in breast cancer, since single nucleotide polymorphisms (SNPs) in VEGF were associated with better overall survival. It is unknown whether these same DNA variants are also predictive of response to bevacizumab in colon cancer patients. In contrast, tumor mutations in k-ras, b-raf, or p53 did not predict response to bevacizumab, and tumor levels of VEGF, thrombospondin-2, and micro-vessel density did not predict response to bevacizumab.

The high cost of bevacizumab treatment, coupled with substantial adverse drug events, makes it important to identify clinical markers that can be used to predict patient response to bevacizumab treatment. In this study, clinical variables that may be informative with respect to predicting response to treatment were explored. Variables that exhibited a statistically significant difference between responders (defined as having at least one full or partial response characterized as a ≥50% reduction in tumor size from baseline, or stable disease in response to bevacizumab treatment) versus non-responders included (1) cancer site (celiac/ascending colon; 37% of responders vs 77% of non-responders, compared to tumor localization in the transverse, descending/sigmoid or other site in the colon); (2) cancer cell type (adenocarcinoma, 87% of responders vs 62% non-responders, compared to adenocarcinoma subtypes including mucinous adenocarcinoma, adenocarcinoma/mucinous adenocarcinoma, adenocarcinoma/signet ring cell carcinoma, mucinous adenocarcinoma/signet ring cell carcinoma, adenocarcinoma/mucinous adenocarcinoma/signet ring cell carcinoma); (3) chemotherapy treatment of primary tumor (57% of responders vs 86% of non-responders); (4) unexplained weight loss (29% of responders vs 64% of non-responders); and (5) blood in stool (52% of responders and zero non-responders). P-values are shown in table 2. When variables exhibiting statistically significant differences between responders and non-responders (excluding blood in stool due to lack of comparison group) were modeled in multivariate analysis, the following profile persisted relative to response to treatment: (1) patients with celiac/ascending colon site had worse outcomes (OR=0.06, CI=0.02-0.56); (2) patients with adenocarcinoma versus other subtypes showed better response (OR=6.35, CI=1.08-37.18); (3) patients initially treated with chemotherapy had worse response (OR=0.07, CI=0.01-0.76); and (4) patients who presented with unexplained weight loss had worse response to treatment (OR=0.10, CI=0.02-0.56). These data suggest that differential response to bevacizumab may be due to inherent differences in tumor gene expression between adenocarcinoma cell type and other colon cancer cell types. The difference noted between primary celiac/ascending tumors and transverse/descending tumors may be due to both anatomical and gene expression differences inherent between ascending colon consistent with differences in function. The ascending colon actively resorbs water, whereas the descending colon is more actively involved in bacterially-mediated breakdown of nutrients and resorption of vitamins and trace minerals. Anatomical differences may also impact on drug resorption and distribution along the length of the colon and may contribute to lower efficacy of bevacizumab treatment observed in non-responders who were treated with chemotherapy as the initial approach to primary tumor treatment. Blood in the stool as the symptom that brought patients in for evaluation before diagnosis of colon cancer was also a strong marker of response to bevacizumab in univariate analysis. However, because we did not observe any non-responders with blood in the stool, this variable could not be modeled in the multivariate analysis and will require a larger trial to obtain an accurate estimate of its effect. This marker as a positive predictor of response to bevacizumab may be useful clinically, since it occurs early in the trajectory of colon cancer diagnosis and treatment. Similarly, unexplained sudden weight loss was more prevalent among non-responders and could be an important early clinical predictor of response to treatment.

A limitation of this study was the small number of non-responders identified. Thus, further studies to validate these findings and enhance modeling through addition of further factors predictive of patient response to bevacizumab treatment are warranted.

Statistically significant outcomes stemming from univariate and multivariate analysis and modeling of clinical data related to clinical factors suggest that profiling patients with respect to their clinical status...
may have important predictive value for anticipating response to bevacizumab treatment. In summary, multivariate analysis demonstrated that patients with pure adenocarcinoma are more likely to respond to bevacizumab treatment compared to other types of tumors. Patients initially treated with chemotherapy treatment of the primary tumor are less likely to response compared to the patients who did not have chemotherapy. Patients who have unexplained weight loss are less likely to respond compared to patients without unexplained weight loss, and patients with cancer localized to the cecal/ascending colon were likely to have worse response to bevacizumab treatment. Univariate analysis suggested that patients presenting with blood in stool were more likely to respond to bevacizumab treatment, and no non-responders presented with blood in stool. Validation of these data in further studies is warranted since predictive value of these clinical observations may be important in informing treatment strategies of patients with colon cancer. This study underlines strong inherent value in advancing such modeling approaches in predicting patient response to treatment.

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Author Contributions

Conception and design: All authors

Provision of study materials or patients: Rezwan Islam

Collection and assembly of data: Po-Huang Chyou and James K Burmester

Data analysis and interpretation: Po-Huang Chyou and James K Burmester

Manuscript writing: All authors

Final approval of manuscript: All authors

CONFLICTS OF INTEREST

All authors declare no financial conflicts of interest.

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