Role of Chronic Inflammatory Ratios in Predicting Recurrence of Resected Patients with Stage I–III Mucinous Colorectal Adenocarcinoma

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Background: Cancer-related inflammation is the main cause of the progression of mucinous colorectal adenocarcinoma (MCA). Circulating fibrinogen-to-pre-albumin ratio (FPR) is associated with the clinical outcome in colorectal cancer (CRC). However, the prognostic role of FPR and which is the best inflammatory prognostic biomarker within MCA remain unknown.

Methods: We enrolled 157 patients with stage I–III MCA in this study. Kaplan-Meier curve, Cox regression, and time-dependent receiver operation characteristic curve analysis were performed to assess the prognostic value and efficacy of the neutrophil-to-albumin ratio (NAR), neutrophil-to-pre-albumin ratio (NPAR), albumin-to-alkaline phosphatase ratio (AAPR), albumin-to-globulin ratio (AGR), albumin-to-fibrinogen ratio (AFR), and FPR in these patients.

Results: We found that NAR, NPAR, and FPR were significantly associated with unsatisfactory recurrence-free survival (RFS) in patients with stage I–III MCA, and the predicted efficacy of FPR was superior to that of the other two inflammatory biomarkers. Moreover, patients with a high combined TNM-CA199-FPR score had worse outcomes, with a high predicted efficacy of up to 0.779 (0.703–0.856). Using FPR, the patient was monitored for the recurrence up to two months earlier than that achieved using the common imaging techniques (4 vs 6 median months) in stage I–III MCA patients undergoing radical resection.

Conclusion: FPR is the preferred inflammatory biomarker and commonly used for predicting and monitoring recurrence in stage I–III MCA patients. The combined TNM-CA199-FPR score is an economical, simple, effective, and independent prognostic factor for localized disease.

Keywords: mucinous colorectal carcinoma, fibrinogen-to-pre-albumin ratio, prognosis, inflammation

Introduction

Mucinous colorectal adenocarcinoma (MCA) is a distinct, rare, and fatal colorectal cancer (CRC) worldwide.1 It is commonly characterized by abundant extracellular mucin, which accounts for more than 50% of the tumor cells. In China, the reported incidence of the disease is only 8.17%,2 and the disease is mostly observed in women and younger patients, with the tumor located on the right side.1 MCA is associated with a poor response to chemotheraphy and poor prognosis.3 However, there is no factor that can effectively predict and monitor recurrence.

Cancer-elicited inflammation is an important hallmark of cancer.4 MCA is more commonly diagnosed in patients with inflammatory bowel diseases or Lynch syndrome,5
and emerging epidemiological evidence indicates that non-steroidal anti-inflammatory drugs are potent in preventing CRC. Thus, chronic inflammation is a critical characteristic in the onset and progression of MCA, fostering its proliferation, invasion, metastasis, and survival. Moreover, the degree of inflammation is presented as an alteration of inflammatory factors and cells within the cancer microenvironment and circulating peripheral blood, and these changes are possibly linked to clinical recurrence and disease progression. Previous studies have shown that the albumin-to-globulin ratio (AGR), neutrophil-to-albumin ratio (NAR), and albumin-to-globulin ratio (AGR) are significantly associated with the prognosis of solid malignancies. In addition, the albumin-to-fibrinogen ratio (AFR), and fibrinogen-to-pre-albumin ratio (FPR) are promising factors for predicting the clinical outcomes of solid malignancies. However, no study has reported the association between these inflammatory biomarkers and the clinical outcomes of MCA.

The present study aimed to investigate the prognostic roles of AAPR, NAR, NPAR, AGR, AFR, and FPR in localized MCA and assess the monitoring role of FPR in 250 patients. These findings would help clinicians tailor decision-making in MCA treatment.

Materials and Methods

In this study, we enrolled patients diagnosed with MCA at the Second Affiliated Hospital of Nanchang University between November 2010 and April 2017. The study was performed in accordance with the guidelines of the Declaration of Helsinki, and was approved by the Ethics Committee of the Second Affiliated Hospital of Nanchang University. Written informed consent was obtained from each participant before the study.

A flowchart of the protocol for screening and identifying eligible patients in this study is shown in Figure 1. We initially recruited 250 diagnosed stage I–III MCA patients for this study. The eligible patients were identified according to the following inclusion criteria: a) first diagnosed MCA patients were clinically confirmed by histopathological examination of the resected biopsies; b) radical resection was performed in stage I–III patients with tumor-negative resection margins; and c) without any emergency or neoadjuvant chemoradiotherapy, and other malignancies within the patients. In contrast, the following criteria were used to exclude unsuitable participants: a) recent diarrhea, infection, hereditary polyposis, ulcerative colitis, autoimmune or chronic kidney disease, hematopath, hepatopathy, or cardiovascular and cerebrovascular disease was clinically confirmed in the patients; b) non-steroidal anti-inflammatory drug or intravenous albumin supplement was undertaken in the past three months; and c) clinical characteristics, and baseline information not provided by the participants or lost to follow-up within three months.

We collected the clinical characteristics, baseline information, and pathological results of each eligible patient. Abundant peripheral blood, plasma, and serum samples were collected after admission with or two days before the surgical operation. Circulating neutrophil count was detected by flow cytometry, laser scattering, and cytochemical staining combined techniques using a Sysmex HST-302 machine (Sysmex, Tokyo, Japan). OLYMPUS AU5400 (Beckman Coulter, Tokyo, Japan) was used to determine the levels of serum alkaline phosphatase (ALP), albumin, and pre-albumin, and the Clauss assay was used to measure plasma fibrinogen level using a SYMEX CA-7000 machine (Sysmex, Tokyo, Japan). Meanwhile, both carcinoembryonic antigen (CEA) and carbohydrate antigen 19–9 (CA19-9) were quantified through the chemiluminescence immunoassay by a SIEMENS ADVIA Centaur XP machine (Siemens, Erlangen, Germany). The inter- and intra-batch coefficients of these detections were less than 10%. AAPR, NAR, NPAR, AGR, AFR, and FPR were calculated according to the formulas shown in Table 1.

The primary survival endpoint in our study was recurrence-free survival (RFS) in the cohort. We performed a three-years follow-up with a frequency of three months in the first two years, and six months subsequently until recurrence or metastasis in the third year or the deadline (April 1st, 2020). The time from resection to recurrence within the localized region or death was considered as RFS. Physical examination, tests for detecting common tumor biomarkers (CEA and CA19-9), and common imaging tests (abdominal computed tomography scan or magnetic resonance imaging) or colonoscopy were performed during follow-up. Contrast-enhanced chest abdominal computed tomography and bone scans were performed to detect lung and bone metastases, respectively. Recurrence or distal metastasis of the disease was diagnosed according to one of the following criteria: a) colonoscopy examination; and b) typical appearances in common imaging detection.

The cut-off values for AAPR, NAR, NPAR, AGR, AFR, and FPR were calculated using X-tile software according to RFS. Binary variables are summarized as numbers and frequencies. The chi-square test and Fisher’s exact tests were used to analyze the significant difference in the comparisons. Continuous variables are expressed as the median and
Results
Baseline Characteristics of Eligible Patients
As shown in Figure 1, 157 patients with stage I–III MCA patients were included in our study according to the inclusion and exclusion criteria. The baseline characteristics and laboratory detection results were described in Table 2. The majority of patients were aged <60 years old (61.78%), and 42.04% of the patients had tumor on the right side. The proportions of patients with stage I, II, and III MCA were 7 (4.46%), 71 (45.22%), and 79 (50.32%), respectively. All localized lesions were resected, 77.71% and 11.46% of the included patients received adjuvant chemotherapy and radiotherapy, respectively. The recurrence rate was 38.85% after the follow-up, and the median RFS was 36 months.
Table 1 Definition and Cut-Off Values of the Included Inflammatory Ratio in Predicting Recurrence in the Localized Patients

| Inflammation-Based Biomarkers | The Localized Cohort                  | Cut-Off Value | Score |
|-------------------------------|--------------------------------------|---------------|-------|
| NAR                           |                                      |               |       |
| Neutrophil/albumin ratio×100   |                                      | 5.60          | 0=NAR≤5.60 | 1=NAR>5.60 |
| NP                        |                                      | 12.80         | 0=NP≤12.80 | 1=NP>12.80 |
| AGR                           |                                      | 0.70          | 0=AAPR≤0.70 | 1=AAPR>0.70 |
| AGR                           |                                      | 1.50          | 0=AGR≤1.50 | 1=AGR>1.50 |
| AFR                           |                                      | 10.80         | 0=AFR≤10.80 | 1=AFR>10.80 |
| FPR                           |                                      | 19.50         | 0=FPR≤19.50 | 1=FPR>19.50 |

Note: The localized disease includes stage I–III colorectal mucinous carcinoma; the cut-off values of these inflammatory biomarkers are calculated using X-tile software according to RFS.

Prognosis of Inflammatory Scores

The optimal cut-off values of NAR, NP, AAPR, AGR, AFR, and FPR were 5.60, 12.80, 0.70, 1.50, 10.80, and 19.50, respectively (Table 1). The patients were stratified into high- (score 1) and low-score (score 0) subgroups according to the cut-off values of the six inflammatory ratios. However, only TNM stage (p<0.001, adjusted HR=3.664, 95% CI=1.813–7.404), CEA (p=0.018, adjusted HR=2.346, 95% CI=1.161–4.742), CA199 (p=0.005, adjusted HR=2.848, 95% CI=1.377–5.890), NAR (p=0.025, adjusted HR=2.280, 95% CI=1.111–4.678), NP (p=0.004, adjusted HR=2.818, 95% CI=1.399–5.674), and FPR (p=0.025, adjusted HR=2.359, 95% CI=1.115–4.990) were found to be significantly associated with poor RFS in the Kaplan-Meier curve, univariate and multivariate Cox regression (Figure 2 and Table 3). Nonetheless, there was no association between AAPR, AGR, or AFR, and the RFS of the patients.

We also evaluated the predictive efficacy of the significant inflammatory ratios, CEA, CA199, and TNM in predicting 3-years RFS. Areas under time-dependent receiver operation curves (AUC) were 0.576 (AUC=0.576, 95% CI=0.481–0.670), 0.586 (AUC=0.586, 95% CI=0.493–0.680), 0.601 (AUC=0.601, 95% CI=0.508–0.694), 0.606 (AUC=0.606, 95% CI=0.513–0.700), 0.642 (AUC=0.642, 95% CI=0.548–0.735) and 0.684 (AUC=0.684, 95% CI=0.597–0.770) for NAR, NP, AAPR, CEA, CA19-9, and TNM, respectively. The TNM stage had the highest AUC in predicting RFS. The AUC of FPR was higher than that of the other inflammatory biomarkers. The AUC of CEA was lower than that of CA19-9 (Figure 2D and Table 4).

Prognosis and the Predicted Efficacy of the Combined Score

According to the prognostic roles of TNM (stages I–II and III defined as negative and positive TNM stages, respectively), CA19-9, and FPR in the patients, we constructed a combined score based on the three biomarkers. We considered score 0 for patients with the three low biomarkers or a single positive FPR, score 1 for those with stage III or high CA19-9 and score 2 for those with two or three positive biomarkers. Sixty-five (41.40%), 43 (27.39%), and 49 (31.21%) patients had scores of 0, 1, and 2, respectively. As shown in Figure 2E, the RFS of the patients with a combined score of 1 (plog-rank=0.002, adjusted HR=3.191, 95% CI=0.964–10.599) and 2 (plog-rank =0.003, adjusted HR=4.548, 95% CI=1.671–12.377) was
Table 2 The Baseline and Clinicopathological Characteristics of Eligible Patients in the Localized Cohort

| Variables                  | The Localized Cohort |
|---------------------------|----------------------|
| N(157)                    | %                    |
| Gender(male)              | 91                   | 57.96                |
| Age(>60 year)             | 60                   | 38.22                |
| Smoking(Yes)              | 14                   | 8.92                 |
| Drinking(Yes)             | 12                   | 7.64                 |
| Diabetes(Yes)             | 15                   | 9.55                 |
| Hypertension(Yes)         | 28                   | 17.83                |
| TNM stage(III)            | 79                   | 50.32                |
| T stage(T3-4)             | 143                  | 91.08                |
| LN status(N1-2)           | 79                   | 50.32                |
| Differentiation(G1-2)     | 78                   | 49.68                |
| Cancer bulk(>5cm)         | 67                   | 42.68                |
| Primary location (Right)  | 66                   | 42.04                |
| Radical/palliative surgery(Yes) | 157         | 100.00               |
| Chemotherapy(Yes)         | 122                  | 77.71                |
| Radiotherapy(Yes)         | 18                   | 11.46                |
| CEA(>5ng/mL)              | 50                   | 31.82                |
| CA199(>37U/mL)            | 33                   | 21.02                |
| NAR(score=1)              | 109                  | 69.43                |
| NPAR(score=1)             | 97                   | 61.78                |
| AAPR(score=1)             | 13                   | 8.28                 |
| AGR(score=1)              | 86                   | 54.78                |
| AFR(score=1)              | 100                  | 63.69                |
| FPR(score=1)              | 66                   | 42.04                |
| Number of recurrence      | 61                   | 38.85                |
| Median RFS (months)       | 36(9-36)             | –                    |

Notes: Tumors located at the caecum, ascending colon and transverse colon were defined as right-sided, and those located within the splenic flexure, and beyond were defined as left-sided.

Abbreviations: LN, lymph node; NAR, neutrophil/lymphocyte/albumin × 100; score=1 means NAR>5.60; NPAR, neutrophil/lymphocyte/pre-albumin ratio × 1000; score=1 means NPAR>12.80; AAPR, albumin/alkaline phosphatase ratio, score=1 means AAPR>0.70; AGR, albumin/globulin ratio, score=1 means AGR>1.50; AFR, albumin/fibrinogen ratio, score=1 means AFR>10.80; FPR, fibrinogen/pre-albumin ratio × 1000, score=1 means FPR>19.50; RFS, recurrence-free survival.

extremely inferior to that of patients with score 0, and their predicted AUC was high (0.779), which was superior to that of the single biomarkers such as TNM, CA19-9, and FPR. Moreover, the sensitivity and specificity of the combined score were 84.75% and 64.55%, respectively, in predicting the 3-year RFS (Figure 2F; Tables 4 and 5).

Predicting the Role of FPR in Monitoring Progression

To investigate the monitor role of FPR in predicting progression, we determined FPR during each follow-up. However, we obtained the complete data from only 21 patients during the follow-up period. Among them, 5 cases did not recur after surgical resection, and negative FPR was examined in each follow-up time. Sixteen patients were found to recurrence, and the first positive FPR detected time within 6 and 9 patients were earlier than or equal to the imaging detection after the operation, respectively. In only one patient, we found that the first positive FPR detection time was later than that in imaging detection (Figure 3).

Discussion

MCA accounts for approximately 1.6%–25.4% of all colorectal cancers, and is associated with a high recurrence rate and unsatisfactory clinical outcome. In this study, we found that NAR, NPAR, and FPR were significantly associated with unsatisfactory RFS in the cohort, and the predictive AUC of FPR was superior to that of the other two inflammatory biomarkers. Moreover, the patients with a high combined TNM-CA199-FPR score showed worse outcomes than that of the low score patients, and the predicted efficacy was high up to 0.779, which was significantly improved comparing to that of single biomarkers. In addition, FPR aided in monitoring recurrence up to two months earlier than that achieved using the common imaging techniques in patients with localized lesions.

MCA is a unique subtype of CRC, as it has distinct clinical and histological characteristics and genetic features. A previous study showed that MCA was mainly observed in female and younger populations; our study showed that 57.96% of the patients were male, and the middle-younger patients accounted for 61.78% of all eligible patients, consistent with findings reported by Song et al. Meanwhile, 42.04% of patients with localized lesions had a high FPR. Accumulating evidence showed that a high FPR within the patients indicated a high-degree of chronic inflammation, and severe cancer-related inflammation could weaken or lead to chemoradioresistance, which leads to unsatisfactory clinical outcomes in CRC patients. Thus, the recurrence rate in the cohort was high (38.85%).

Previous studies have shown that the common prognostic factors, such as TNM stage, venous and lymphoid invasion, microsatellite instability (MSI) status, CEA, and CA19-9 are associated with the prognosis of MCA. However, there was a need for an improved predicted efficacy. In our study, we found that stage III, high CA19-9 and high CEA were related to the worse survival. Recent studies have shown that cancer-elicited inflammatory biomarkers (neutrophil-to-lymphocyte ratio, AFR, and FPR) are promising prognostic biomarkers for the
Figure 2 Prognostic values of the significant inflammatory biomarkers in the study. (A) Kaplan-Meier (K-M) curve of NAR; (B) K-M curve of NPAR; (C) K-M curve of FPR; (D) time-dependent receiver operating characteristics curve (tdROC) of the independent prognostic factors; (E) K-M curve of TNM-CA199-FPR score; (F) tdROC of TNM-CA199-FPR score.
Table 3 The Relationship Between the Baseline and Pathological Variables, Inflammatory Ratios and Recurrence-Free Survival in the Localized Cohort

| Variables                  | Cox Regression | P_{log-rank} | Value   | Crude HR (95% CI) | Adjusted HR (95% CI) |
|----------------------------|----------------|--------------|---------|-------------------|----------------------|
| Gender(male)               | 0.644          | 0.887(0.534–1.474) | 0.015(0.478–2.156) |
| Age(>60 year)              | 0.770          | 1.080(0.646–1.805) | 0.872(0.414–1.838) |
| Smoking(Yes)               | 0.493          | 1.317(0.599–2.895) | 1.450(0.416–5.059) |
| Drinking(Yes)              | 0.737          | 1.169(0.468–2.920) | 0.616(0.074–5.153) |
| Diabetes(Yes)              | 0.175          | 1.673(0.795–3.523) | 1.056(0.268–4.157) |
| Hypertension(Yes)          | 0.934          | 1.028(0.535–1.974) | 1.451(0.577–3.646) |
| TNM stage(III/IV)          | <0.001         | 3.790(2.136–6.725) | 3.664(1.813–7.404) |
| T stage(T3–4)              | 0.081          | 5.805(0.804–4.192) | 3.775(0.347–32.24) |
| LN status(N1-2)            | <0.001         | 2.759(1.633–4.662) | 2.06(0.671–6.329) |
| Differentiation(G1-2)       | 0.048          | 2.165(1.006–4.656) | 1.263(0.525–3.037) |
| Cancer bulk(>5cm)          | 0.356          | 1.275(0.761–2.138) | 0.890(0.422–1.877) |
| Primary location(right)    | 0.068          | 0.610(0.359–1.038) | 0.943(0.424–2.100) |
| Chemotherapy(Yes)          | 0.425          | 0.773(0.411–1.454) | 0.933(0.381–2.289) |
| Radiotherapy(Yes)          | 0.162          | 0.603(0.296–1.226) | 0.512(0.178–1.474) |
| CEA(>5ng/mL)               | 0.005          | 2.081(1.245–3.478) | 2.346(1.161–4.742) |
| CA199(>37U/mL)             | <0.001         | 2.992(1.765–5.072) | 2.848(1.377–5.890) |
| NAR(score=1)               | 0.019          | 1.852(1.107–3.100) | 2.280(1.111–4.678) |
| NPAR(score=1)              | 0.020          | 1.817(1.098–3.006) | 2.818(1.399–5.674) |
| AAPR(score=1)              | 0.731          | 1.174(0.470–2.936) | 0.411(0.054–3.103) |
| AGR(score=1)               | 0.939          | 0.980(0.593–1.622) | 0.909(0.435–1.900) |
| AFPR(score=1)              | 0.154          | 0.693(0.418–1.148) | 0.707(0.338–1.480) |
| FPR(score=1)               | 0.028          | 1.764(0.604–2.924) | 2.359(1.15–4.990) |

Notes: Tumors located at the caecum, ascending colon and transverse colon were defined as right-sided, and those located within the splenic flexure, and beyond were defined as left-sided; multivariable Cox regression was adjusted by gender, age, tobacco, alcohol, diabetes, hypertension, chemotherapy, radiotherapy, and primary location.

Abbreviations: LN, lymph node; NAR, neutrophil/lymphocyte/albumin ×100; score=1 means NAR>5.60; NPAR, neutrophil/lymphocyte/pre-albumin ratio ×1000; score=1 means NPAR>13.80; AAPR, albumin/alkaline phosphatase ratio; score=1 means AAPR>0.70; AGR, albumin/globulin ratio; score=1 means AGR>1.50; AFPR, fibrinogen/pre-albumin ratio; score=1 means AFPR>10.80; FPR, fibrinogen/pre-albumin ratio ×1000; score=1 means FPR>19.50; HR, hazard ratio; CI, confidence interval.

Table 4 Comparison of the Performance Discriminative Ability Between the Significant Prognostic Factors

| Biomarkers | AUROC(95% CI) | 36 Months Recurrence-Free Survival |
|------------|--------------|-----------------------------------|
|            | Sensitivity  | Specificity                       |
| NAR        | 0.576(0.481–0.670) | 60.66% | 25.00% |
| NPAR       | 0.586(0.493–0.680) | 52.46% | 32.29% |
| FPR        | 0.601(0.508–0.694) | 55.74% | 66.67% |
| CEA        | 0.606(0.513–0.700) | 54.24% | 24.47% |
| CA199      | 0.642(0.548–0.735) | 38.98% | 87.50% |
| TNM        | 0.684(0.597–0.770) | 73.77% | 64.58% |
| TNM-CA199-FPR score | 0.779(0.703–0.856) | 84.75% | 64.55% |

Abbreviations: NAR, neutrophil/lymphocyte/albumin ×100; NPAR, neutrophil/lymphocyte/pre-albumin ratio ×1000; FPR, fibrinogen/pre-albumin ratio ×1000; AUROC, area under time-dependent receiver operating characteristic curve; CI, confidence interval.

localized and metastatic CRC. Moreover, circulating FPR was the preferred inflammatory biomarkers in predicting recurrence in patients with stage II–III CRC patients in terms of prognostic ability. Furthermore, NAR, NPAR, and FPR were associated with short RFS adjusted by common characteristics and other confounders. This suggests that these variables could be considered as independent prognostic factors to predict disease progression. In addition, the predicted efficacy and sensitivity of the combined TNM-CA199-FPR score were high.
up to 0.779 and 84.75%, respectively; these values were significantly higher than those of individual factors. The revealed that the combined score improves the predicted efficacy of the individual factors, and it is economical, simple, and effective in predicting disease recurrence. Moreover, FPR is a better index that aids in monitoring
Table 5 Cox Analysis of TNM-CA199-FPR Score in the Localized Cohort

| Combined Score | The Localized Cohort |
|----------------|----------------------|
|                | $\rho_{log-rank}$-value | Crude HR (95% CI) | Adjusted HR (95% CI) |
| Score 0        | 0.002                | 3.349 (1.478–7.590) | 1.910 (0.964–10.599) |
| Score 1        | <0.001               | 8.560 (4.089–17.921) | 6.079 (2.898–15.528) |

Notes: Multivariable Cox regression was adjusted by gender, age, tobacco, alcohol, diabetes, hypertension, chemotherapy, radiotherapy, T and LN status, cell differentiation, cancer size, and primary location.

Abbreviation: FPR, fibrinogen/pre-albumin ratio ×1000.

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All authors contributed to data analysis, drafting or revising the article, have agreed on the journal to which the article will be submitted, gave final approval of the version to be published, and agree to be accountable for all aspects of the work.

Disclosure
The authors have no conflicts of interest to declare.

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