Predictive Values of the Selected Inflammatory Indexes in Colon Cancer

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Abstract

**Purpose:** Ample evidence has revealed that the lymphocyte-to-monocyte ratio (LMR), albumin-to-globulin ratio (AGR), and mean platelet volume (MPV) are cancer-related inflammatory markers. The present study aimed to combine these indicators to better assess the progression of colon cancer.

**Methods:** This retrospective study enrolled 251 patients with colon cancer, 171 patients with benign colon diseases, and 187 healthy control subjects. The receiver operating characteristic curve and area under the curve (AUC) were used to determine the diagnostic values of the selected inflammatory index.

**Results:** The levels of LMR, AGR, and MPV were decreased in the colon cancer group compared with the healthy control and benign colon disease groups. The LMR, AGR, and MPV were all correlated with tumor size. Moreover, LMR and AGR was associated with lymph node metastasis and clinical stage, AGR was related to distant metastasis. Both the LMR (P = .030) and AGR (P = .005) were negatively correlated with the concentration of carcinoembryonic antigen (CEA). The AUC value of MPV combined with CEA had a good diagnostic ability for distinguishing colon cancer cases (AUC = .950) and patients with benign colon diseases (AUC = .886) from controls. Meanwhile, the combination of LMR or AGR with CEA could enhance larger AUC (.746 for LMR + CEA, .737 for AGR + CEA) than CEA, LMR, or AGR alone in detecting colon cancer from benign colon diseases.

**Conclusions:** CEA combined with the LMR, AGR, or MPV may be used as better blood-based biomarkers in the progression of colon cancer patients.

**Keywords**
colon cancer, lymphocyte-to-monocyte ratio, albumin-to-globulin ratio, mean platelet volume, diagnosis

Introduction

Colon cancer comprised the highest incidence in tumors of the digestive system and represented a commonly diagnosed malignant tumor worldwide in 2020¹; it was the fourth most frequent cause of cancer morbidity and fifth leading cause of cancer mortality in China in 2015.² Due to the lack of obvious manifestations, most patients with colon cancer have silent symptoms for years. More than 50% of colon cancer cases are clinically diagnosed at the advanced cancer stage.³ Therefore, effective screening protocols are essential for colon cancer detection. As well known, fecal occult blood test (FOBT) is a cheap and convenient screening method for colon cancer.⁴

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Nevertheless, the result is frequently affected by many dietary factors, multiple drugs, and upper or lower gastrointestinal bleeding site, which may lead to false positives and subsequent unnecessary tests and panic. Other methods that colonoscopy and biopsy have been used as ideal methods for the diagnosis of early colon cancer, but these inspections greatly increase the physical and financial burden on patients harboring colonic diseases, resulting in poor compliance. Hence, low-cost, non-invasive and easily obtainable markers have important significance for the diagnosis and prevention of colon cancer.

Several serum tumor markers have been commonly used in colon diseases, such as carcinoembryonic antigen (CEA), carbohydrate antigen 19-9 (CA19-9), and carbohydrate antigen 242 (CA242). CEA is considered the primary marker, with almost all researches and clinical practices for the detection and monitoring of colon cancer used this indicator.9 However, several studies discovered that CEA possessed low sensitivity and could not appropriately present the complete potency for clinical diagnosis and treatment in colorectal cancer. Thus, more reliable and powerful biomarkers of identifying the colon cancer are expected to obtain. In recent years, many inflammatory parameters from peripheral blood and serum have been evaluated in the diagnosis and prognosis of multiple malignancies, including colorectal cancer, such as circulating neutrophils, lymphocytes, monocytes, platelets, mean platelet volume (MPV), and albumin. Due to the release of chemokines and cytokines, MPV is, as an early indicator of platelet activation, an inflammatory marker. And it is distorted and has been shown to have diagnostic/predictive value in some non-malignant inflammatory conditions. Moreover, several reports discovered that it was closely related to the occurrence, features, and outcomes of many neoplasms. As crucial components of host immunity, lymphocytes can infiltrate into the tumor microenvironment, and they play a vital role in cell-mediated immunity, preventing the proliferation and metastatic activity of colorectal cancer. Systemic inflammation can induce changes in the hematological system, leading to a significant decrease of lymphocytes. Lymphocytopenia is considered an insufficient immune response against tumor, resulting in hyperproliferation and tumorigenesis. Conversely, the excess circulating monocytes gather and settle in solid tumor tissues after being mediated by chemokines of inflammatory cytokines; they are then differentiated into tumor-associated macrophages with specific phenotypic characteristics. Increasing evidence has demonstrated that the accumulation of tumor-associated macrophages in the tumor sites contributes to the angiogenesis, tumorigenesis, and pathogenesis of colon cancer. As a result, the relatively lower number of lymphocytes is an indicator marker of weak immune response, and the elevated monocyte count is a microenvironment monitor of high tumor burden. Therefore, lymphocyte-to-monocyte ratio (LMR), as a reflection of systemic inflammation and immunological statuses, may have a crucial role in the progression of colon cancer. The albumin-to-globulin ratio (AGR) which combines serum albumin and globulin, is a routinely available and cost-effective marker and associated with the process of inflammatory and nourishment state. Accumulated evidence displayed that AGR was an independent and useful predictor in the prognosis of colon cancer by regulating cells and/or releasing several mediators; moreover, elevated AGR was a favorable factor for better clinical outcomes. Hence, we hypothesized that AGR may have a significant diagnostic value in colon cancer.

Up to now, to our knowledge, studies have rarely investigated the diagnostic role of these three inflammatory parameters (LMR, AGR, and MPV) in the progression of colon cancer, especially for persons with benign colon diseases. Therefore, this study investigated the value of LMR, AGR, and MPV combined or not with CEA in the progression of colon cancer.

Material and Methods

Patients

251 patients with colon cancer, 171 benign colon diseases cases, and 187 healthy controls with complete clinical data were consecutively included in this retrospective study from January 2012 to September 2020. In colon cancer participants with new diagnoses, the disease was confirmed by histology and treated with surgical resection. Clinical staging of colon cancer was conducted accorded to the seventh edition of the American Joint Committee on Cancer/TNM tumor staging criteria. Patients with other cancers, cardiovascular disease, diabetes mellitus, hematological disease, autoimmune disease, recent blood transfusion, liver cirrhosis, alcoholic fatty liver disease, nephropathy, or treatment with other therapies, such as radiotherapy, chemotherapy, and hormone therapy, were excluded. Colon polyps, colon adenomas, and colonitis were included as benign colon disease patients who were diagnosed by colonoscopy and histopathology. The healthy controls were healthy subjects without the above-mentioned diseases and clinical gastrointestinal symptoms during the same period of physical examination. Ever smokers were defined as participants who had smoked at least 100 cigarettes in their lifetime, and ever drinkers were defined as participants who had drunk alcoholic beverages once a week for at least a year. All patient details have been de-identified in this study.

Data Collection

All data were collected from the hospital’s electronic medical records, including gender, age, height, weight, smoking status, drinking status, white blood cells (WBC), platelets, hemoglobin, lymphocyte, monocyte, albumin, globulin, MPV, CA19-9, CA242, and CEA. Whole blood-cell parameters were detected with a Beckmann 780 device (Beckman Coulter, Brea, CA). The levels of albumin and total protein were analyzed by a Hitachi 7600 automatic biochemical
The concentration of serum CEA was tested by using a Roche E6000 analyzer (Roche Diagnostics, Basel, Switzerland). Architect i2000 and its reagents (Abbott GmBH Diagnostika, Wiesbaden, Germany) were used to evaluate the serum CA19-9 values. The level of serum CA242 was determined using Enzyme Linked Immunosorbent Assay (CanAg Company, Sweden). The ratios of interest were calculated as follows: body mass index (BMI) = weight/height,\(^2\) LMR = lymphocyte count/monocyte count, and AGR = albumin/(total protein – albumin).

**Statistical Analysis**

The Kolmogorov–Smirnov test was used to detect the distribution of the continuous variables. The mean and standard deviation were applied for normally distributed data, and non-normally distributed data were expressed as the median and quartile. Differences between groups in laboratory parameters and clinical characteristics were calculated using the Student’s \(T\) test (normally distributed data), Mann–Whitney nonparametric \(U\) test (non-normally distributed data), or \(\chi^2\) test (categorical variable). The Spearman correlation coefficient was conducted to detect correlations between inflammatory index (LMR, AGR, or MPV) and CEA in the colon cancer group. The receiver operating characteristic (ROC) curve and area under the curve (AUC) were calculated by MedCalc statistical software (version 18.1.1). Data processing and analysis were determined by SPSS 16.0 statistical software package, using a significance level of .05.

**Results**

**Patient Characteristics**

Basic information and laboratory parameters are summarized in Table 1. The median age of the colon cancer cases, benign colon diseases patients, and control individuals were 56.00, 48.00, and 53.00 years, respectively. No intergroup difference was observed in gender, BMI, smoking status and drinking status among the three groups. Patients with colon cancer had higher levels of WBCs, platelets, monocytes, and CEA compared with healthy individuals and benign colon disease patients. Conversely, the levels of hemoglobin, MPV, albumin, LMR, and AGR were significantly lower in the colon cancer group than they were in the control and benign colon disease groups, and there were statistical differences in LMR (Figure 1A), AGR (Figure 1B) and MPV (Figure 1C) among the three groups. The values of CA19-9 were evident discrepancies between benign colon diseases and healthy controls, as well as in colon cancer and benign colon diseases groups. The concentrations of CA242 were not available in the benign colon disease group, and there was difference between the healthy control group and colon cancer group \((P < .001)\).

**Correlations Between LMR, AGR, MPV, and the Clinicopathological Characteristics in Patients With Colon Cancer**

There was a negative correlation presented between CEA and the LMR \((r = -.137, P = .030)\) (Figure 2A) and AGR \((r = -.178, P = .005)\) (Figure 2B) in the colon cancer group, respectively. Nevertheless, no correlation was observed between AGR and MPV \((r = .012, P = .846)\) (Figure 2C). According to the seventh edition of the American Joint Committee on Cancer/TNM tumor stage, the clinicopathological characteristics of the 251 patients carrying colon cancer are shown in Table 2. The levels of LMR, AGR, and MPV in the colon cancer group were all closely related to the tumor size, but not associated with tumor invasion. Moreover, the AGR and LMR were correlated with lymph node metastasis and clinical stage. The colon cancer subjects with stage M0 had significantly higher levels of AGR compared to the cases with stage M1 \((P = .013)\).

**Logistic Regression Used to Distinguish Colon Cancer From Controls**

The correlation between several potential risk factors and colorectal cancer was analyzed by binary logistic regression (Table 3), including gender (odds ratio [OR] = .892, 95% confidence interval [CI] = .610–1.034, \(P = .556)\), age (OR = 1.013, 95% CI = .994–1.031, \(P = .174)\), BMI (OR = .975, 95% CI = .918–1.035, \(P = .401)\), smoking status (OR = .785, 95% CI = .466–1.323, \(P = .363)\), drinking status (OR = .857, 95% CI = .443–1.659, \(P = .647)\), MPV (OR = .089, 95% CI = .055–.143, \(P < .001)\), CA242 (OR = 1.061, 95% CI = 1.033–1.089, \(P < .001)\), CA19-9 (OR = 1.012, 95% CI = 1.002–1.022, \(P = .019)\), CEA (OR = 2.855, 95% CI = 2.223–3.666, \(P < .001)\), LMR (OR = .547, 95% CI = .466–.644, \(P < .001)\), and AGR (OR = .036, 95% CI = .015–.088, \(P < .001)\). The above important indexes \((P < .05)\) were selected as potential independent predictors for further multivariate analysis. After multivariate analysis, MPV \((\beta = -2.352, P < .001)\), LMR \((\beta = -.306, P = .001)\), AGR \((\beta = -4.091, P < .001)\), and CEA \((\beta =...}
.967, \( P < .001 \) were also recognized as crucial markers in the occurrence of colon cancer. The optimal model (logit \( P = .967 \times \text{CEA} - .306 \times \text{LMR} - .4091 \times \text{AGR} - 2.352 \times \text{MPV} + 27.383 \)) was set up for differencing colon cancer cases from controls. The AUC, sensitivity, and specificity were reach up to .964, 90.84%, and 92.51%, respectively.

Diagnostic efficacy of LMR, AGR, CEA, and MPV alone or in combination to differentiate patients with colon cancer from other subjects.

The results of the ROC curve analysis are presented in Table 4 and Figure 3. The AUC value of the combination of MPV and CEA was .950 (95% CI = .925–.968, positive likelihood ratio [PLR] = 9.48, negative likelihood ratio [NLR] = .097, positive predictive value [PPV] = 92.7%, negative predictive value [NPV] = 88.5%), which possessed a good diagnostic ability for distinguishing colon cancer cases from healthy controls. Meanwhile, the sensitivity and specificity of the combination of MPV and CEA were
increased to 91.24% and 90.37%, respectively. Compared to the subjects with benign colon disease, LMR combined with CEA produced larger AUC (.746) than other combined indicators in subjects with colon cancer. However, the sensitivity (75.30%) of combination for LMR and CEA was lower than LMR (77.29%) and AGR (83.27%) alone. To predict colon cancer, the optimal cut-offs of LMR, AGR, MPV, and CEA in the benign colon disease subjects were 4.58, 1.71, 7.80, and 3.44, respectively. For the diagnosis of benign colon disease, the combined use of MPV and CEA resulted in better AUC (.886) and sensitivity (84.80%) than any single index or combination of other indicators. However, the specificity (84.49%) of the combination of MPV and CEA was inferior to MPV (96.79%) or AGR (89.84%) alone.

**Table 2.** Correlation between LMR, AGR, and MPV and clinicopathological features in colon cancer.

|                     | N   | LMR             | p   | AGR             | p   | MPV             | p   |
|---------------------|-----|-----------------|-----|-----------------|-----|-----------------|-----|
| **Tumor invasion (T stage)** |     |                 |     |                 |     |                 |     |
| T1 + T2             | 86  | 3.63 (2.83–4.87)| .513| 1.46 (1.33–1.62)| .602| 8.10 (7.53–8.73)| .973|
| T3 + T4             | 165 | 3.63 (2.89–4.36)| 1.50 (1.31–1.66)| 8.14 (7.40–8.69)| |
| **Lymph node metastasis (N stage)** |     |                 |     |                 |     |                 |     |
| N0                  | 177 | 3.75 (2.85–4.87)| .033| 1.52 (1.33–1.68)| .025| 8.10 (7.34–8.69)| .300|
| N1–N3               | 74  | 3.44 (2.90–4.12)| 1.43 (1.31–1.58)| 8.17 (7.60–8.71)| |
| **Distant metastasis (M stage)** |     |                 |     |                 |     |                 |     |
| M0                  | 242 | 3.63 (2.94–4.52)| .366| 1.50 (1.32–1.65)| .013| 8.11 (7.44–8.70)| .900|
| M1                  | 9   | 3.09 (2.48–4.28)| 1.19 (1.04–1.44)| 8.16 (6.90–8.97)| |
| **Tumor size (cm)** |     |                 |     |                 |     |                 |     |
| <5                  | 159 | 3.85 (3.00–4.65)| .004| 1.52 (1.34–1.67)| .002| 8.20 (7.60–8.80)| .026|
| ≥5                  | 92  | 3.26 (2.70–4.17)| 1.43 (1.15–1.60)| 7.90 (7.14–8.60)| |
| **Clinical stage**  |     |                 |     |                 |     |                 |     |
| I + II              | 179 | 3.77 (2.95–4.86)| .013| 1.52 (1.33–1.69)| .002| 8.10 (7.40–8.70)| .833|
| III + IV            | 72  | 3.43 (2.80–4.03)| 1.44 (1.23–1.57)| 8.17 (7.49–8.69)| |

LMR, lymphocyte-to-monocyte ratio; AGR, albumin-to-globulin ratio; MPV, mean platelet volume.

**Discussion**

Colon cancer is closely associated with inflammation which has been unraveled as a crucial hallmark in all the steps of colon tumorigenesis, including initiation, invasion, progression, and metastasis. Recent work has elucidated that cancer-associated inflammatory markers are increasingly used in the early diagnosis and prognosis of malignant tumors. However, rarely studies have investigated the diagnostic role of these inflammatory parameters (LMR, AGR, and MPV) in distinguish colon cancer from benign colon diseases. Therefore, this study assessed the diagnostic efficacy of the common inflammatory indexes and detected whether they could be used as surrogate markers in the progression of colon cancer.
In this retrospective analysis, we discovered that the healthy controls had significantly higher MPV compared to the benign colon disease patients and colon cancer cases, which was consistent with the result of Stojkovic et al.\(^33\) but contrary to the findings of Li et al.\(^34\) and Kilincalp et al.\(^35\) We found that the MPV values of the control group in the first two articles were 9.06 and 10.7, which were similar to the results of this study, but the wide gulf was obtained from the Kilincalp et al.’s article in the control group. The differences may be due to health groups included criteria, different sample size, and ethnic variation in different study. As stated in the previous research that MPV was connected with obesity, smoking, and so on,\(^36\) which may also differ among the studies population, resulting in the inconsistent outcomes. Although no association was demonstrated between clinical stage and MPV in our current finding, the results revealed that the MPV levels in patients with stage III and IV were slightly higher than that in cases with stage I and II, which was similar to Li et al.’s study.\(^34\) And the present study was the first research revealing the correlation between MPV and the tumor size.

In concordance with previous results, the patients with colon cancer had a lower LMR level than the benign colon disease patients and healthy controls did. For example, Aksoy et al.\(^37\) observed that the level of monocyte-to-lymphocyte ratio (MLR) was higher in the gastric cancer group than it was in the intestinal metaplasia and healthy control groups. A study by Luo et al.\(^38\) found that MLR was significantly elevated in patients carrying urothelial carcinoma of the bladder relative to healthy controls. Moreover, our study disclosed that the level of LMR was significantly correlated with the features of colon cancer, such as lymph node metastasis, tumor size, and clinical stage. Indeed, Ozawa et al.\(^39\) demonstrated that cancer-specific survival was significantly worse in patients with low LMR levels than in high-LMR patients, and LMR may be an independent prognostic marker for stage III and IV colon cancer cases.\(^40\) Peng et al.\(^41\) assessed the prognosis of patients harboring colorectal liver-only metastases and elucidated that elevated LMR predicted a favorable outcome in both 5-year recurrence-free survival and overall survival of patients with lymph node metastases and liver tumor up to a diameter of less than 5 cm. Furthermore, several meta-analyses have demonstrated that malignant patients with high preoperative LMR have better predicted clinical outcomes compared with patients with low LMR in populations comprising Asians, digestive system carcinomas, non-metastatic diseases, and early disease stages,\(^42,43\) which confirmed our findings.

Emerging evidence suggests that AGR is mainly used as a clinical indicator for several kinds of cancers. Growing tumors induce hypoalbuminemia via secreting inflammatory cytokines, which may inhibit albumin synthesis and promote albumin loss, resulting in weak systemic response. Rasouli et al.\(^44\) reported that patients with malignant tumors had a decreased concentration of albumin, which were measured by colorimetric methods, compared with healthy controls, which was accordant with the present study results. Globulin, as a reflector for most proinflammation protein, was increased by the accumulation of acute-phase proteins and immunoglobulins.\(^45\) The AGR, which is compatible with hypoalbuminemia and hyperglobulinemia, may be able to more accurately reflect the nutritional and inflammatory state, and thus is associated with the progression of neoplasia. The electrophoretic data of serum proteins showed that the AGR was significantly decreased in 85 patients harboring cancer relative to controls.\(^44\) Cheng et al.\(^46\) confirmed that the globulin-to-albumin ratio (GAR), was significantly higher in the subjects with liver disease compared with individuals with no evidence of liver disease. Quite a few studies revealed that patients with lower pretreatment AGR were related to worse survival than higher AGR subjects in colorectal cancer,\(^47\) gastric cancer,\(^48\) pancreatic cancer,\(^49\) nasopharyngeal

### Table 3. Screening for significant predictors that distinguished colon cancer from healthy controls by using univariate and multivariate analyses.

| Variables | Univariate analysis | Multivariate analysis |
|-----------|---------------------|----------------------|
|           | OR                  | 95%CI                | P        |
|           | OR                  | 95%CI                | P        |
| Gender    | .892                | .610—1.304           | .556     |
| Age(years)| 1.013               | .994—1.031           | .174     |
| BMI       | .975                | .918—1.035           | .401     |
| Smoking status | .785                | .466—1.323           | .363     |
| Drinking status | .857                | .443—1.659           | .647     |
| MPV       | .089                | .055—.143            | <.001    |
| CA1242    | 1.061               | 1.033—1.089          | <.001    |
| CA19-9    | 1.012               | 1.002—1.022          | .019     |
| CEA       | 2.855               | 2.223—3.666          | <.001    |
| LMR       | .547                | .466—.644            | <.001    |
| AGR       | .036                | .015—.088            | <.001    |

BMI, body mass index; MPV, mean platelet volume; CA1242, carbohydrate antigen 242; CA19-9, carbohydrate antigen 19-9; CEA, carcinoembryonic antigen; LMR, lymphocyte-to-monocyte ratio; AGR, albumin-to-globulin ratio. CI, confidence interval; OR, odd ratio.
Table 4. Diagnostic efficacy of LMR, AGR, and CEA used alone or in combination to differentiate colon cancer from benign colon diseases.

| Cut off | Sensitivity (%) | Specificity (%) | PLR  | NLR  | PPV (%) | NPV (%) | AUC (95% CI) | P      |
|---------|-----------------|-----------------|------|------|---------|---------|--------------|--------|
| LMRa    | 3.78            | 55.38           | 88.77| .50  | 86.9    | 59.7    | .778(.736—.816) | <.001  |
| AGRa    | 1.59            | 66.53           | 71.12| .47  | 75.6    | 61.3    | .756(.713—.796) | <.001  |
| MPVa    | 8.93            | 83.27           | 96.79| .17  | 97.2    | 81.2    | .894(.862—.922) | <.001  |
| CEAa    | 1.35            | 80.08           | 78.07| .26  | 83.1    | 74.5    | .870(.835—.900) | <.001  |
| LMRa+AGRa| .64      | 65.74           | 88.24| .39  | 88.2    | 65.7    | .819(.780—.854) | <.001  |
| LMRa+MPVa| .60      | 82.47           | 94.65| .19  | 95.4    | 80.1    | .916(.886—.940) | <.001  |
| AGRa+MPVa| .65      | 78.49           | 96.26| .22  | 96.6    | 76.9    | .923(.894—.946) | <.001  |
| LMRa+CEAa| .47      | 86.45           | 81.82| .17  | 86.5    | 81.8    | .895(.863—.922) | <.001  |
| AGRa+CEAa| .46      | 84.86           | 80.75| .19  | 85.5    | 79.9    | .889(.856—.917) | <.001  |
| MPVa+CEAa| .45      | 91.24           | 90.37| .97  | 92.7    | 88.5    | .950(.925—.968) | <.001  |
| LMRb    | 4.58            | 77.29           | 53.22| .43  | 70.8    | 61.5    | .688(.642—.732) | <.001  |
| AGRb    | 1.71            | 83.27           | 42.11| .40  | 67.9    | 63.2    | .655(.607—.700) | <.001  |
| MPVb    | 7.80            | 38.25           | 74.85| .82  | 69.1    | 45.2    | .571(.523—.619) | <.011  |
| CEB     | 3.44            | 43.43           | 87.72| .64  | 83.8    | 51.4    | .686(.639—.730) | <.001  |
| LMRb+AGRb| .66     | 53.39           | 78.95| .59  | 78.8    | 53.6    | .717(.671—.759) | <.001  |
| LMRb+MPVb| .65     | 51.39           | 80.70| .60  | 79.6    | 53.1    | .698(.652—.742) | <.001  |
| AGRb+MPVb| .61     | 62.15           | 69.01| .55  | 74.6    | 55.4    | .680(.633—.724) | <.001  |
| LMRb+CEAb| .54     | 75.30           | 63.74| .39  | 75.3    | 63.7    | .746(.702—.787) | <.001  |
| AGRb+CEAb| .58     | 58.57           | 76.02| .55  | 78.2    | 55.6    | .737(.693—.779) | <.001  |
| MPVb+CEAb| .59     | 51.39           | 83.63| .58  | 82.2    | 54.0    | .715(.669—.758) | <.001  |
| LMRC    | 5.02            | 63.16           | 53.48| .69  | 55.4    | 61.3    | .595(.542—.646) | <.002  |
| AGRC    | 1.46            | 30.41           | 89.84| .77  | 73.2    | 58.5    | .567(.514—.619) | <.032  |
| MPVC    | 8.98            | 77.78           | 96.79| .23  | 95.7    | 82.6    | .856(.815—.891) | <.001  |
| CEAc    | 1.05            | 77.19           | 66.31| .29  | 64.7    | 76.1    | .776(.729—.818) | <.001  |
| LMRc+AGRC| .54     | 40.35           | 83.42| .72  | 69.0    | 60.5    | .610(.558—.661) | <.000  |
| LMRc+MPVc| .55     | 74.85           | 95.19| .26  | 93.4    | 80.5    | .864(.824—.898) | <.001  |
| AGRC+MPVc| .55     | 71.35           | 96.79| .30  | 95.3    | 78.7    | .851(.810—.887) | <.001  |
| LMRc+CEAc| .42     | 73.68           | 74.33| .35  | 72.4    | 75.5    | .779(.733—.821) | <.001  |
| AGRC+CEAc| .43     | 70.18           | 72.19| .41  | 69.8    | 72.6    | .773(.726—.816) | <.001  |
| MPVc+CEAc| .38     | 84.80           | 84.49| .18  | 83.3    | 85.9    | .886(.848—.917) | <.001  |

Note. CEA, carcinoembryonic antigen; LMR, lymphocyte-to-monocyte ratio; AGR, albumin-to-globulin ratio; MPV, mean platelet volume; PLR, positive likelihood ratio; NLR, negative likelihood ratio; PPV, positive predictive value; NPV, negative predictive value; CI, confidence interval; AUC, area under curve.

*colon cancer vs healthy controls.

bcolon cancer vs benign colon diseases.

*benign colon diseases vs healthy controls.

Figure 3. The diagnostic value of LMR, AGR, and MPV used alone or in combination with CEA in the progression of colon cancer. (A). colon cancer vs healthy controls, (B). colon cancer vs benign colon diseases, (C). benign colon diseases vs healthy controls. Note. LMR: lymphocyte-to-monocyte ratio, AGR: albumin to globulin ratio, MPV: mean platelet volume, CEA: carcinoembryonic antigen.
carcinoma,\textsuperscript{50} and esophageal cancer.\textsuperscript{51} Moreover, a significant correlation based on the above-mentioned researches was observed between clinical characteristics and the level of AGR, such as lymph node metastasis, tumor size, distant metastasis, and tumor stage. In agreement with previous studies, this study found that the value of AGR in the colon cancer subjects was lower than that in the benign colon diseases and healthy individuals; furthermore, it showed that the AGR was associated with lymph node metastasis, distant metastasis, tumor size, and clinical stage.

CEA is a serum glycoprotein that is mainly secreted by cells of the large intestine, and it has been widely applied as a tumor marker for the malignant characteristics of colorectal cancer. Unfortunately, high levels of CEA are not present in about 15\% of large intestine cancers.\textsuperscript{52} Although CEA has a high specificity in colorectal cancer,\textsuperscript{53,54} elevated CEA is also revealed in other malignant tumors. Moreover, increased circulating levels of CEA are observed not only in cancer patients but also in some benign intestinal lesions, and the role was poor in differentiating colorectal cancer patients from those with benign colorectal diseases.\textsuperscript{35} Therefore, the sensitivity and effectiveness of CEA are not sufficient for clinical diagnosis and treatment. Consistent with previous studies, the sensitivity (43.43\%) and diagnostic value (686) of CEA were not noticeable in identifying colon cancer from benign colon diseases in this study, while the specificity was up to 87.72\%. Whereas we found that CEA combined with LMR or AGR generated a significantly better sensitivity and AUC than CEA used alone in discriminating patients with colon cancer and benign colon diseases cases. Similar to this pilot study, a previous report displayed that LMR possessed a moderate ability (AUC = .71) and could contribute to distinguishing patients carrying gastric cancer from those with intestinal metaplasia,\textsuperscript{37} the diagnostic efficiency was similar to that of our study in colon cancer, and our research have improved diagnostic efficiency through the combination of indicators. Meanwhile, the AUC value of MPV combined with CEA had a good diagnostic ability and sensitivity for distinguishing colon cancer cases (AUC = .950 and 91.24\%) and patients with benign colon diseases (AUC = .886 and 84.80\%) from controls, was superior compared with individual indexes or combination of other indicators. Similarly, Stojkovic et al.\textsuperscript{33} revealed that ROC curve analysis showed high diagnostic efficacy of NLR, PLR and MPV in CRC patients compared with individual markers (AUC = .904). In many malignancies, AGR exhibited good diagnostic efficacy (AUC = .81) in differentiating cancer patients from healthy controls,\textsuperscript{54} which squared with our results. All these findings suggest that CEA combined with the selected inflammatory index could not only be used as a colon cancer diagnostic biomarker but may also improve the diagnostic efficiency of detecting patients harboring colon cancer from benign colon diseases cases.

There are certain potential limitations in the current research. On the one hand, this is a retrospective analysis of a relatively small sample size from a single center, so selection bias and statistical validity should be noted, which may affect the final results about the associations between the LMR or AGR and colon cancer. We failed to stratify benign colon diseases due to the relatively small sample size. On the other hand, confounding factors cannot be completely ruled out, including age, race, dietary habits and family histories, which may prevent us from drawing any firm conclusions. For example, median age of included patients was quite low in the three groups. Moreover, different populations may show different levels of “inflammatory conditions.” Probably, results would be different including elderly patients and patients with chronic/inflammatory conditions or Western populations. Therefore, a large-scale, prospective study with multiple centers is still needed to validate these results.

Conclusion
This study first described that the LMR and AGR were correlated with CEA and colon cancer, as well as lymph node metastasis, tumor size, distant metastasis, and clinical stage. Moreover, the combination of the LMR, AGR, or MPV with CEA may enhance sensitivity and diagnostic efficacy and may be a helpful diagnostic marker for differentiating colon cancer from benign colon diseases and healthy controls.

Declaration of Conflicting Interests
The author(s) declared no potential conflicts of interest with respect to the research, authorship, and/or publication of this article.

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Ethics Approval
This research was approved by the Ethics Committee of the First Affiliated Hospital of Guangxi Medical University (No.2022-KY-E-024), and all participants verbally agreed that their data could be used for scientific research under the protection of privacy.

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