**Name of journal:** **World Journal of Gastroenterology**

**ESPS Manuscript NO: 12053**

**Columns: OBSERVATIONAL STUDY**

**Prognostic value of *KRAS* and *BRAF* mutations in curatively resected colorectal cancer**

Kadowaki S *et al*. Prognostic maker in colorectal cancer

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**Supported by** Japanese Ministry of Health, Labor and Welfare

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**Received:** June 19, 2014 **Revised:** September 9, 2014

**Accepted:** October 14, 2014

**Published online:**

**Abstract**

**AIM:** To investigate the prognostic role of *KRAS* and *BRAF* mutations after adjustment for microsatellite instability (MSI) status in Japanese colorectal cancer (CRC) population.

**METHODS:** We assessed *KRAS* and *BRAF* mutations and MSI status in 813 Japanese patients with curatively resected, stage I–III CRC and examined associations of these mutations with disease-free survival (DFS) and overall survival (OS) using uni- and multivariate Cox proportional hazards models.

**RESULTS:** *KRAS* and *BRAF* mutations were detected in 312 (38%) of 812 and 40 (5%) of 811 tumors, respectively. *KRAS* mutations occurred more frequently in females than in males (*P* = 0.02), while the presence of *BRAF* mutations was significantly associated with the female gender (*P* = 0.006), proximal tumor location (*P* < 0.001), mucinous or poorly differentiated histology (*P* < 0.001), and MSI-high tumors (*P* < 0.001). After adjusting for relevant variables, including MSI status, *KRAS* mutations were associated with poorer DFS (HR = 1.35; 95%CI: 1.03–1.75) and OS (HR = 1.46; 95%CI: 1.09–1.97). *BRAF* mutations were poor prognostic factors for DFS (HR = 2.20; 95%CI: 1.19–4.06) and OS (HR = 2.30; 95%CI: 1.15–4.71). Neither the *BRAF* by MSI interaction test nor the *KRAS* by MSI interaction test yielded statistically significant results for DFS and OS.

**CONCLUSION:** *KRAS* and *BRAF* mutations are associated with inferior survival, independent of MSI status, in Japanese patients with curatively resected CRC.

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**Key words:** Colorectal cancer; *KRAS*; *BRAF*; microsatellite instability; Prognostic factor

**Core tip:** Although *KRAS* and *BRAF* mutations play a critical role in colorectal cancer development, little is known regarding the prognostic role of these genetic alterations after adjustment for microsatellite instability status in Asian populations. To the authors' knowledge, the current study is the first large-scale study to clarify the impact of *KRAS* and *BRAF* mutations on the survival outcomes of colorectal cancer in Asian populations. We found that *KRAS* and *BRAF* mutations were separately associated with inferior disease-free survival and overall survival, independent of microsatellite instability status, in patients with curatively resected colorectal cancer.

Kadowaki S, Kakuta M, Takahashi S, Takahashi A, Arai Y, Nishimura Y, Yatsuoka T, Ooki A, Yamaguchi K, Matsuo K, Muro K, Akagi K. Prognostic value of *KRAS* and *BRAF* mutations in curatively resected colorectal cancer. *World J Gastroenterol* 2014; In press

**INTRODUCTION**

Colorectal cancer (CRC) develops through diverse mechanisms such as chromosomal instability (CIN), microsatellite instability (MSI), and epigenetic DNA promoter methylation [CpG island methylator phenotype (CIMP)][[1](#_ENREF_1)]. CIMP and MSI-high (MSI-H) phenotypes are closely associated. Most sporadic MSI-H tumors develop through CIMP-associated methylation of *MLH1*, and *BRAF* mutations occur frequently in both phenotypes[[2](#_ENREF_2),[3](#_ENREF_3)]. *KRAS* mutations mainly occur in CIN and are partly associated with intermediate CIMP epigenotypes[[4](#_ENREF_4)]. *KRAS* and *BRAF* mutations are mutually exclusive; both cause RAS/RAF/MAPK signaling pathway upregulation and are crucial in CRC development.

*KRAS* encodes a guanosine triphosphate/guanosine diphosphate binding protein; *KRAS* mutations are observed in approximately 30%-40% CRCs[[5-8](#_ENREF_5)]. *KRAS* mutations are well known as predictive markers of resistance to epidermal growth factor receptor (EGFR)-targeted antibodies in metastatic CRC, but their prognostic value remains controversial. Some studies have shown that *KRAS* mutations are associated with poorer survival in CRC[[8](#_ENREF_8),[9](#_ENREF_9)], while others found no association[[6](#_ENREF_6),[7](#_ENREF_7)].

*BRAF* encodes a serine/threonine protein kinase, a downstream effector of the KRAS protein. Activating *BRAF* mutations occur in approximately 4%-20% CRCs[[6](#_ENREF_6),[10-14](#_ENREF_10)], with the vast majority being the V600E hotspot mutation. Although some previous studies have shown that *BRAF* mutations confer poorer prognosis in CRC[[10-12](#_ENREF_10)], others have not[[6](#_ENREF_6),[13](#_ENREF_13)], probably because of associations with favorable MSI-H CRC prognosis[[15-17](#_ENREF_15)].

Although genetic background and geographical factors may influence mutation frequency and prognosis, most reports are from Western countries; less data are available regarding the prognostic role of *KRAS* and *BRAF* mutations in Asian populations. Two independent studies from Taiwan and Japan have been published recently. However, both had a small sample size and heterogeneous cohorts including metastatic disease; the study from Taiwan did not examine MSI status[[14](#_ENREF_14),[18](#_ENREF_18)]. Hence, a large homogenous cohort with MSI status is essential for assessing the prognostic value of various clinical or molecular variables in CRC. Here, we clarified associations of *KRAS* and *BRAF* mutations and MSI status with survival outcomes in a larger Japanese cohort of patients with curatively resected CRC.

**MATERIALS AND METHODS**

***Patients and tissue samples***

A total of 813 consecutive stage I to III CRC patients undergoing curative resection at Saitama Cancer Center between July 1999 and May 2006 were included. Written informed consent was obtained from all patients. Patients with the following conditions were excluded: (1) history of radiotherapy or chemotherapy preoperatively; (2) inflammatory bowel disease; or (3) history of familial adenomatous polyposis. Pathological staging was performed according to the tumor, node, and metastasis (TNM) classification system (6th edition)[[19](#_ENREF_19)]. CRCs were typically divided into 3 types: rectum, distal colon (splenic flexure and descending and sigmoid colon), and proximal colon (cecum and ascending and transverse colon). Adjuvant chemotherapy was administered to 40% (129/322) and 76% (232/307) of stage II and III CRC patients, respectively. Among 361 patients treated with adjuvant chemotherapy, only 10 patients received combination chemotherapy with 5-fluorouracil, leucovorin, and oxaliplatin, while remaining were treated with single-agent fluoropyrimidines. Patients were followed-up until death or February 2012, whichever came first. We obtained approval from the Ethics Committee of Saitama Cancer Center.

***Genomic DNA extraction and KRAS and BRAF mutation analysis***

Primary CRCs and paired healthy colorectal mucosa obtained perioperatively were immediately frozen at −80 °C until analysis. Genomic DNA was extracted from fresh frozen specimens using the standard phenol–chloroform extraction method. Exons 2 and 3 of *KRAS* were examined for mutations by denaturing gradient gel electrophoresis, as described previously[[20](#_ENREF_20)]. The *BRAF* V600E mutation was detected using PCR and restriction enzyme digestion, as described previously[[21](#_ENREF_21)].

***MSI analysis***

MSI analysis was performed using fluorescence-based PCR, as described previously[[22](#_ENREF_22)]. Five Bethesda markers BAT25, BAT26, D5S346, D2S123, and D17S250 were used to classify tumor MSI status. MSI status was graded as MSI-H with 2 or more unstable markers, MSI-low (MSI-L) with only 1 unstable marker, and microsatellite-stable (MSS) with no unstable marker. MSI-positive markers were re-examined at least twice to confirm the result.

***Statistical analysis***

The aim of this study was to evaluate the impact of *KRAS*/*BRAF* mutations on prognosis in patients with resected CRC. Prognosis was evaluated according to 2 measures: overall survival (OS) and disease-free survival (DFS). OS was defined as the interval from the date of resection until death due to any cause or until the censor date of February 1, 2012. DFS was defined as the time from the date of resection to tumor recurrence, occurrence of a new primary colorectal tumor, or death due to any cause. Survival probability was estimated using the Kaplan-Meier method and compared using the log-rank test. Cox proportional hazards models were used to estimate uni- and multivariate adjusted hazard ratios (HRs) for DFS and OS according to mutation status. Factors for which the multivariate models were adjusted are age (≥ 65 *vs* < 65), gender (male *vs* female), tumor stage (III *vs* II *vs* I), adjuvant chemotherapy (Yes *vs* No), and status of MSI and *BRAF* or *KRAS* mutations (Yes *vs* No). To further evaluate the potential heterogeneity of the impact of *KRAS* and *BRAF* mutations according to MSI status and other covariates [age (≥ 65 *vs* < 65), gender (male *vs* female), tumor location (distal/rectum *vs* proximal), and stage (III *vs* I/II)], we tested the models that included interaction terms, cross-products of gene mutation status, and another variable of interest in a multivariate Cox model. The likelihood ratio test was performed to determine the significance of the results.

 Clinicopathological factor distribution according to gene mutation status was assessed using the *χ*2 or Fisher’s exact tests for categorical variables, when appropriate, and Student’s *t*-test for continuous variables. All statistical analyses were performed using Dr. SPSS II software (SPSS Japan Inc., Tokyo, Japan); 2-sided *P* < 0.05 was considered statistically significant.

**RESULTS**

***Clinicopathological characteristics of KRAS and BRAF mutant tumors***

Patient characteristics according to *KRAS* or *BRAF* status are summarized in Table 1. MSI status was determined in all cases, whereas mutation status was not determined in 1 case for *KRAS* and 2 for *BRAF*. *KRAS* or *BRAF* mutations were detected in 38% (312/812) and 5% (40/811) of cases, respectively. Only 1 patient harbored *KRAS* or *BRAF* mutations. *KRAS* mutations were more frequent in females than in males (43% *vs* 35%; *P* = 0.02). *BRAF* mutations were significantly more frequent in females than in males (7% *vs* 3%; *P* = 0.006), proximal than in distal or rectal tumors (13% *vs* 1% *vs* 2%; *P* < 0.001), mucinous or poorly differentiated tumors than in moderately or well-differentiated tumors (17% *vs* 4%; *P* < 0.001), and MSI-H tumors than in MSS/MSI-L tumors (36% *vs* 2%; *P* < 0.001).

***Survival analysis***

The median follow-up time was 87.7 mo (range: 13–148 mo). Based on univariate Cox proportional hazard analysis results (Table 2), greater age (≥ 65), male gender, advanced TNM stage, and presence of *KRAS* mutations were significantly associated with poor prognosis for DFS and OS. For *KRAS* mutant *vs* *KRAS* wild-type tumors, 5-year DFS was 71% *vs* 77% (log-rank *P* = 0.02; Figure 1A); 5-year OS was 80% *vs* 84%, respectively (log-rank *P* = 0.01; Figure 1B). Presence of *BRAF* mutations was not significantly associated with poorer DFS and OS in the entire cohort. For *BRAF* mutant *vs* wild-type tumors, 5-year DFS was 70% *vs* 75% (log-rank *P* = 0.23; Figure 1C); 5-year OS was 77% *vs* 83% (log-rank *P* = 0.11; Figure 1D), respectively.

In multivariate analysis, adjusting for potential prognostic variables, *KRAS* retained its prognostic impact on DFS (HR = 1.35; 95%CI: 1.03–1.75) and OS (HR = 1.46; 95%CI: 1.09–1.97; Table 3). Presence of *BRAF* mutations was significantly associated with poorer DFS (HR, 2.20; 95%CI: 1.19–4.06) and OS (HR = 2.30; 95%CI: 1.15–4.71) after adjustment (Table 3).

***Survival analysis stratified by MSI status***

Given the potential prognostic effect of MSI status, we evaluated interactions of *KRAS* or *BRAF* mutations with MSI status (Table 4). The effect of *KRAS* mutations on DFS and OS was limited to patients with MSS/MSI-L tumors (HR = 1.37; 95%CI: 1.05–1.80; HR = 1.49; 95%CI: 1.10–2.02, respectively); however, the *KRAS* by MSI interaction test was not significant (*P* = 0.95 and 0.70, respectively). *BRAF* mutations were significantly associated with reduced OS (HR = 2.74; 95%CI: 1.19–6.30) in MSS/MSI-L, but not MSI-H, tumors. However, the *BRAF* by MSI interaction test did not reach statistical significance (*P* = 0.44).

***Survival analysis stratified by other potential variables***

We also analyzed the prognostic value of *KRAS* and *BRAF* mutations for OS across strata of other potential prognostic factors (Figure 2). The prognostic effect of *KRAS* mutations appeared to be consistent across potential variables, and interactions between *KRAS* status and these factors were not significant. In contrast, *BRAF* mutations were significantly associated with poor OS in stage III, but not stage I–II, disease. Interactions between *BRAF* status and TNM stage showed suggestive statistical significance (*P* = 0.10).

**DISCUSSION**

To our knowledge, this is the largest study to assess the prognostic value of *KRAS* and *BRAF* mutations for survival outcomes in CRC patients in Asian populations. Tumor specimens were prospectively collected from patients with curatively resected CRC (stage I–III); *KRAS* and *BRAF* mutations and MSI status were analyzed using a consistent methodology at a single institution. *KRAS* and *BRAF* mutations were associated with poor prognosis, independent of MSI status.

Many studies have examined associations of *KRAS* mutations with various clinical features, with no consistent results[[5-8](#_ENREF_5)]. *KRAS* mutations were more frequent in females; however, these mutations were not associated with any other clinical variable. Similarly, Watanabe *et al*[[5](#_ENREF_5)] found relationships of *KRAS* mutations with the female gender and older age. In contrast, the Kirsten Ras Colorectal Cancer Collaborative Group study (RASCAL) demonstrated that *KRAS* mutations were associated with histological grade but no other variables[[8](#_ENREF_8)]. In analysis of the PETACC-3 trial, Roth *et al*[[6](#_ENREF_6)] reported associations of *KRAS* mutations with histological grade and tumor location but not gender. Such inconsistencies may be attributed to differences in the distribution of age, race, stage, or other factors among subject groups.

Currently, no convincing evidence exists that *KRAS* mutations are independent prognostic factors in CRC. In a Taiwanese study by Liou *et al*[[14](#_ENREF_14)], KRAS mutations were not associated with inferior OS; however, the magnitude of multivariate HR (HR = 1.61; 95%CI: 0.91–2.84) was of the same order as that in the present study. A study from Japan revealed that the prognostic impact of *KRAS* mutations on recurrence-free survival was limited in patients with stage II CRC, and the association of *KRAS* mutations with OS was not observed[[18](#_ENREF_18)]. Both studies had a small sample size and heterogeneous cohorts, including stage IV disease. In the large homogeneous cohort in this study, we found significant association of *KRAS* mutations with inferior DFS and OS. Because we previously found no difference in survival outcomes among different *KRAS* mutations, including those in exons 2, 3, and 4[[23](#_ENREF_23)], prognostic analyses of specific codons for these mutations were not performed in the present study. Similarly, the RASCAL study indicated that *KRAS* mutations resulted in overall poorer prognosis[[8](#_ENREF_8)], whereas subsequent analysis (RASCAL II) showed that only the glycine to valine substitution in codon 12 (G12V) was associated with poor prognosis in patients with Dukes’ C disease[[24](#_ENREF_24)]. Furthermore, recent randomized phase III trial results supported *KRAS* mutations as prognostic factors; 3-year DFS ranged from 72% to 75% across treatments for *KRAS* wild-type tumors, with 65% to 67% for *KRAS* mutant tumors[[25](#_ENREF_25)]. In contrast, in the PETACC-3 trial, no association was found between *KRAS* mutations and poorer relapse-free survival (RFS) or OS[[6](#_ENREF_6)]. Although further research of the prognostic effect of *KRAS* mutations is needed, the influence of these mutations seems to be mild across previously reported studies.

The frequency of *BRAF* mutations (5%) and MSI-H (8%) in our cohort was lower than that in Western populations (*BRAF*: 8%-20%, MSI-H: 11%-17%)[[6](#_ENREF_6),[9](#_ENREF_9),[11-13](#_ENREF_11),[15](#_ENREF_15),[16](#_ENREF_16" \o "Ribic, 2003 #903)] and comparable with that in Asian populations (*BRAF*: 4%-7%, MSI-H: 6%-12%)[[14](#_ENREF_14),[18](#_ENREF_18),[26](#_ENREF_26)]. Generally, *BRAF* mutations and MSI-H are frequently observed in females, proximal tumors, and poorly differentiated tumors. In a systematic review including 9,885 CRC patients, a *BRAF* mutation was associated with a proximal tumor location, poor differentiation, and female sex[[27](#_ENREF_27)]. Consistent with this observation, *BRAF* mutations were more frequent in proximal tumors, poorly differentiated tumors, and females. Previous Western cohorts showed more patients with proximal and poorly differentiated tumors compared with Asian cohorts, including the current cohort. Thus, the discrepancy in *BRAF* mutations and MSI-H status between Western and Asian populations may be attributed to the different distribution of patients’ characteristics such as gender, tumor location, histological grade, or racial and/or environmental differences.

Most previous studies found associations of *BRAF* mutations with poorer survival[[6](#_ENREF_6),[10-12](#_ENREF_10)]. In meta-analysis of 26 independent studies (11,773 patients), *BRAF* mutations increased the risk of mortality in CRC patients (HR = 2.25; 95%CI: 1.82–2.83)[[28](#_ENREF_28)]. However, this evidence is mainly based on studies in Western populations; little is known regarding the prognostic role of *BRAF* mutations in Asian populations. In a Taiwanese study[[14](#_ENREF_14)], *BRAF* mutations were associated with reduced OS, but MSI status was not estimated. In a Japanese study, Nakanishi *et al*[[18](#_ENREF_18)] found no such association because of the insufficient number of patients with *BRAF* mutations. In the present study with larger sample size and homogeneous cohorts, we found associations of *BRAF* mutations with poorer DFS and OS in CRC patients with stage I–III disease, with the same order of magnitude of HR for OS as in the above meta-analysis. The prognostic effect of *BRAF* mutations on survival seems to be even stronger than that of *KRAS* mutations.

In contrast to previous reports[[6](#_ENREF_6),[9](#_ENREF_9),[15-17](#_ENREF_15)], our analysis did not show that patients with MSI-H tumors exhibited better survival than those with MSS/MSI-L tumors. However, the number of patients with MSI-H tumors was too small to draw meaningful conclusions regarding the prognostic effect of MSI status. Therefore, additional larger studies are needed to clarify the prognostic impact of MSI status. Inconsistent results were reported regarding the prognostic effect of *BRAF* mutations according to MSI status[[6](#_ENREF_6),[10](#_ENREF_10),[13](#_ENREF_13)]. Samowitz *et al*[[10](#_ENREF_10)] found associations of BRAF mutations with poor survival in MSS/MSI-L, but not MSI-H tumors. Meanwhile, French *et al*[[13](#_ENREF_13)] reported associations of *BRAF* mutations with poor survival in MSI-H tumors. In our analysis, associations of *BRAF* mutations with reduced OS were limited in MSS/MSI-L tumors. However, the *BRAF* by MSI interaction test was not significant; statistical power was considerably limited due to the small number of patients with MSI-H and *BRAF* mutant tumors. Larger studies are needed to clarify the modifying effect on the relation between *BRAF* mutations and survival outcome according to MSI status. Advantages of this study include comprehensive analysis of molecular markers using consistent methodology at a single institution, large sample size, and homogeneous cohort of Japanese patients. These results suggest that constitutive activation of the RAS/RAF/MAPK signaling pathway may be closely associated with clinical prognosis in CRC. Prognostic effects of *KRAS* and *BRAF* mutations seem to be consistent across most strata of clinical variables, while the adverse effect of *BRAF* mutations on OS may be attenuated in stage I to II CRC patients, with marginal statistical significance. The interaction of *BRAF* mutations with tumor stage warrants further research.

In conclusion, we found that Japanese CRC patients with *KRAS* or *BRAF* mutations have poorer survival, independent of MSI status. Additional investigations are warranted to clarify the interaction between these mutations and potential relevant factors, such as MSI status and tumor stage.

**COMMENTS**

***Background***

*KRAS* and *BRAF* mutations occur in 30%–40% and 4%–20% of colorectal cancers (CRCs), respectively. Microsatellite instability (MSI) is characterized by inactivation of the DNA mismatch repair system and is observed in 5%–15% of CRCs. MSI-high tumors are less likely to metastasize compared with the other phenotypes and have favorable survival outcomes. *KRAS* mutations are well known as predictive markers of resistance to epidermal growth factor receptor-targeted antibodies, and *BRAF* mutations are of current interest as a therapeutic target in metastatic CRCs. However, their prognostic value remains controversial for patients with curatively resected CRCs.

***Research frontiers***

Most previous studies investigating the prognostic role of *KRAS* and *BRAF* mutations in CRCs are from Western countries. Genetic background and geographical factors may influence mutation frequency and prognosis; however, few data are available regarding the prognostic role of these genetic alterations in Asian populations. Thus, clinical implications will be obtained by assessing the prognostic value of these mutations in a large cohort of CRCs in Japan, after adjustment for MSI status.

***Innovations and breakthroughs***

This study is the first large-scale study to demonstrate the prognostic impact of *KRAS* and *BRAF* mutations in Asian populations. After adjustment for relevant factors, including MSI, *KRAS* and *BRAF* mutations were independently associated with inferior disease-free survival and overall survival in patients with curatively resected CRCs. These findings will offer new insight into prognostic role of *KRAS* and *BRAF* mutations in CRCs.

***Applications***

*BRAF* and *KRAS* mutations may be useful as molecular markers for stratification of the clinical prognosis of curatively resected CRCs. Further investigation on whether the prognostic impact of *KRAS* and *BRAF* mutations could be modified by MSI status may provide more precise stratification of clinical outcomes in CRC.

***Terminology***

The protein product of the *KRAS* gene is a guanosine triphosphate/guanosine diphosphate-binding protein, and *KRAS* mutations play a key role in the development of various malignancies, including lung cancer, pancreatic cancer, and CRC. The protein product of the *BRAF* gene, a protein called B-Raf, is a serine/threonine protein kinase serving as downstream effector of the KRAS protein. *BRAF* mutations are involved in the development of many malignancies, *e.g.*, malignant melanoma, papillary thyroid cancer, and CRC.

***Peer review***

This is well written and illustrated paper. The authors investigate the prognostic role of KRAS and BRAF mutations after adjustment for MSI status. And they demonstrated that KRAS and BRAF mutations are associated with inferior survival, independent of MSI status in Asian colorectal cancer population. As the authors mentioned, in contrast to previous reports, their analysis did not show that patients with MSI-H tumors exhibited better survival than those with MSS/MSI-L tumors.

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**P-Reviewer:** Paoluzi oa, Sakakura C, Tajika M, Wang jy **S-Editor:** Ma YJ **L-Editor:** **E-Editor:**



**Figure 1 Kaplan–Meier curves for disease-free survival and overall survival according to *KRAS* or *BRAF* status.** A: disease-free survival (DFS) according to *KRAS* status; B: overall survival (OS) according to *KRAS* status; C: DFS according to *BRAF* status; D: OS according to *BRAF* status.



**Figure 2** **Stratified analysis of *KRAS* or *BRAF* status and overall survival.** Loge [adjusted hazard ratio (HR)] and 95% confidence interval (CI) for *BRAF* and *KRAS* mutant tumors (*vs* wild-type tumors) in various strata are shown. A: *KRAS* mutant tumors; B: *BRAF* mutant tumors.

**Table 1 Patient characteristics according to *BRAF* or *KRAS* status *n* (%)**

|  |  |  |
| --- | --- | --- |
| **Characteristics** | ***KRAS* status** | ***BRAF* status** |
| **Wild-type*****n* = 500** | **Mutant*****n* = 312** | ***P-*value** | **Wild-type*****n* = 771** | **Mutant*****n* = 40** | ***P-*value** |
| Age (yr) Mean ± SD | 63.5 ± 10.3 | 64.7 ± 10.3 | 0.11 | 63.9 ± 10.3 | 65.4 ± 11.6 | 0.40 |
| Gender Male Female | 308 (62)192 (38) | 166 (53)146 (47) | 0.02 | 459 (60)312 (40) | 15 (38)25 (63) | 0.006 |
| Tumor location Proximal Distal Rectum | 134 (27)213 (43)153 (31) | 98 (31)125 (40)89 (29) | 0.37 | 201 (26)332 (43)238 (31) | 31 (78)5 (13)4 (10) | < 0.001 |
| Histological grade Well/moderate Poor/mucinous | 472 (94)28 (6) | 288 (92)24 (8) | 0.24 | 728 (94)43 (6) | 31 (78)9 (23) | < 0.001 |
| T stage 1 2 3 4 | 52 (10)106 (21)286 (57)56 (11) | 31 (10)46 (15)200 (64)35 (11) | 0.12 | 79 (10)145 (19)462 (60)85 (11) | 4 (10)7 (18)23 (58)6 (15) | 0.89 |
| LN metastasis Yes No | 180 (36)320 (64) | 127 (41)185 (59) | 0.18 | 292 (38)479 (62) | 15 (38)25 (63) | 0.96 |
| TNM stage I II III | 125 (25)195 (39)180 (36) | 58 (19)127 (41)127 (41) | 0.09 | 173 (22)306 (40)292 (38) | 10 (25)15 (38)15 (38) | 0.92 |
| Adjuvant chemotherapy Yes No | 217 (43)283 (57) | 144 (46)168 (54) | 0.44 | 344 (45)427 (55) | 16 (40)24 (60) | 0.57 |
| MSI status MSS/MSI-L MSI-H | 455 (91)45 (9) | 290 (93)22 (7) | 0.33 | 728 (94)43 (6) | 16 (40)24 (60) | < 0.001 |

SD: Standard deviation; LN: Lymph node; TNM: Tumor–Node–Metastasis; MSI: Microsatellite instability; MSS: Microsatellite stable; MSI-L: Microsatellite instability-low; MSI-H: Microsatellite instability-high.

**Table 2 Univariate prognostic analysis of disease-free survival and overall survival**

|  |  |  |
| --- | --- | --- |
| **Characteristics** | **Disease-free survival** | **Overall survival** |
| **HR** | **95%CI** | **HR** | **95%CI** |
| Age (yr) < 65 ≥ 65 | 11.73 | Reference1.35–2.28 | 12.21 | Reference1.64–2.98 |
| Gender Female Male | 11.57 | Reference1.20–2.06 | 11.57 | Reference1.16–2.13 |
| Tumor location Proximal Distal Rectum | 10.921.17 | Reference0.67–1.250.85–1.62 | 10.900.97 | Reference0.64–1.260.67–1.40 |
| Histological grade Well/moderate Poor/mucinous | 11.53 | Reference0.97–2.42 | 11.43 | Reference0.84–2.42 |
| AJCC stage I II III | 12.604.68 | Reference1.61–4.192.95–7.42 | 12.263.49 | Reference1.36–3.752.14–5.70 |
| Adjuvant chemotherapy No Yes | 11.24 | Reference0.96–1.60 | 11.29 | Reference1.10–1.51 |
| MSI MSS/MSI-L MSI-H | 10.71 | Reference0.42–1.20 | 10.92 | Reference0.54–1.59 |
| *KRAS* Wild-type Mutant | 11.35 | Reference1.04–1.74 | 11.44 | Reference1.08–1.92 |
| *BRAF* Wild-type Mutant | 11.38 | Reference0.82–2.32 | 11.57 | Reference0.90–2.76 |

HR: Hazard ratio; CI: Confidence interval; MSI: Microsatellite instability; MSS: Microsatellite stable; MSI-L: Microsatellite instability-low; MSI-H: Microsatellite instability-high.

**Table 3 Prognostic effects of microsatellite instability, *KRAS*, and *BRAF* status in Cox proportional models**

|  |  |  |
| --- | --- | --- |
|  | **Disease-free survival1** | **Overall survival1** |
| **HR****(95%CI)** | ***P-*value** | **HR** **(95%CI)** | ***P-*value** |
| MSI MSS/MSI-LMSI-H | 1 (Reference)0.64 (0.35–1.16) | 0.14 | 1 (Reference)0.81 (0.42–1.56) | 0.53 |
| *KRAS* Wild-type Mutant | 1 (Reference)1.35 (1.03–1.75) | 0.03 | 1 (Reference)1.46 (1.09–1.97) | 0.01 |
| *BRAF* Wild-typeMutant | 1 (Reference)2.20 (1.19–4.06) | 0.01 | 1 (Reference)2.30 (1.15–4.71) | 0.02 |

1Covariates include age (< 65 or ≥ 65), gender, AJCC stage (I/II/III), adjuvant chemotherapy (Yes/No), and MSI, *KRAS*,and *BRAF* status*.* CI: Confidence interval; MSI: Microsatellite instability; MSS: Microsatellite stable; MSI-L: Microsatellite instability-low; MSI-H, Microsatellite instability-high.

**Table 4 Prognostic Effects of *KRAS* and *BRAF* mutations according to microsatellite instability status**

|  |  |  |
| --- | --- | --- |
|  | ***KRAS*** | ***BRAF*** |
| **HR (95%CI)** | ***P-*value** | **HR (95%CI)** | ***P-*value** |
| DFS1 MSS/MSI-L MSI-H | 1.37 (1.05-1.80)1.34 (0.34-5.24) | 0.95 | 2.06 (0.96-4.43)2.46 (0.49-12.4)  | 0.91 |
| OS1 MSS/MSI-L MSI-H | 1.49 (1.10–2.02)1.39 (0.33-5.78) | 0.70 | 2.74 (1.19-6.30)1.18 (0.23-6.02) | 0.44 |

1Covariates include age, gender, AJCC stage (I-II/III), adjuvant chemotherapy, and *KRAS* and *BRAF* status. HR: Hazard ratio; CI: Confidence interval; DFS: Disease-free survival; OS: Overall survival; MSI: Microsatellite instability; MSS: Microsatellite stable; MSI-L: Microsatellite instability-low; MSI-H: Microsatellite instability-high.