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ORIGINAL REPORT

Association Between Results of a Gene Expression Signature Assay and Recurrence-Free Interval in Patients With Stage II Colon Cancer in Cancer and Leukemia Group B 9581 (Alliance)

Donna Niedzwiecki, Wendy L. Frankel, Alan P. Venook, Xing Ye, Paula N. Friedman, Richard M. Goldberg, Robert J. Mayer, Thomas Anthony Colacchio, Jude Marie Mulligan, Timothy S. Davison, Eamonn O'Brien, Peter Kerr, Patrick G. Johnston, Richard D. Kennedy, D. Paul Harkin, Richard L. Schilsky, Monica M. Bertagnolli, Robert S. Warren, and Federico Innocenti

СТ

BSTRA

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R.S.W. and F.I. contributed equally to this publication.

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The content of this manuscript is solely the responsibility of the authors and does not necessarily represent the official views of the National Cancer Institute.

Authors' disclosures of potential conflicts of interest are found in the article online at www.jco.org. Author contributions are found at the end of this article.

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Purpose

Conventional staging methods are inadequate to identify patients with stage II colon cancer (CC) who are at high risk of recurrence after surgery with curative intent. ColDx is a gene expression, microarray-based assay shown to be independently prognostic for recurrence-free interval (RFI) and overall survival in CC. The objective of this study was to further validate ColDx using formalin-fixed, paraffin-embedded specimens collected as part of the Alliance phase III trial, C9581.

Patients and Methods

C9581 evaluated edrecolomab versus observation in patients with stage II CC and reported no survival benefit. Under an initial case-cohort sampling design, a randomly selected subcohort (RS) comprised 514 patients from 901 eligible patients with available tissue. Forty-nine additional patients with recurrence events were included in the analysis. Final analysis comprised 393 patients: 360 RS (58 events) and 33 non-RS events. Risk status was determined for each patient by CoIDx. The Self-Prentice method was used to test the association between the resulting CoIDx risk score and RFI adjusting for standard prognostic variables.

Results

Fifty-five percent of patients (216 of 393) were classified as high risk. After adjustment for prognostic variables that included mismatch repair (MMR) deficiency, CoIDx high-risk patients exhibited significantly worse RFI (multivariable hazard ratio, 2.13; 95% CI, 1.3 to 3.5; P < .01). Age and MMR status were marginally significant. RFI at 5 years for patients classified as high risk was 82% (95% CI, 79% to 85%), compared with 91% (95% CI, 89% to 93%) for patients classified as low risk.

Conclusion

CoIDx is associated with RFI in the C9581 subsample in the presence of other prognostic factors, including MMR deficiency. CoIDx could be incorporated with the traditional clinical markers of risk to refine patient prognosis.

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INTRODUCTION

Stage I colon cancer (CC) is treated with surgery alone, and patients with stage III CC are routinely offered adjuvant chemotherapy; however, the optimal clinical management of patients with stage II CC is not clearly defined. Stage II CC accounts for approximately 25% of patients with CC, and the 5-year survival rate is approximately 75% to 80%, with 15% to 20% of patients experiencing disease recurrence.¹ Current guidelines^{2,3} recommend adjuvant chemotherapy for patients with stage II CC with high-risk features such as T4 stage, high tumor grade, inadequate number of nodes sampled, positive or unknown margins, lymphovascular and perineural invasion, bowel obstruction, and perforation. However, a number of studies have suggested that these histopathological and clinical markers are limited in their prognostic power.^{4,5} These limitations have prompted efforts to identify molecular markers to guide the clinical management of patients with stage II CC. Although some molecular markers, such as loss of heterozygosity and p53 mutational status, require more evidence to establish their

clinical utility as prognostic markers, assessment of microsatellite instability status has been adopted into clinical practice.³ Several gene signatures have also been shown to be independently associated with clinical outcome in patients with stage II CC.⁶⁻¹⁰

One such gene expression signature, the ColDx assay, can be applied to formalin-fixed paraffin-embedded (FFPE) tissue collected at the time of diagnosis and has previously been shown to be prognostic for recurrence-free interval (RFI) and overall survival (OS) both independent of, and superior to, standard pathologic features.⁶ The gene expression signature was developed in two stages. Stage one involved creation of a Colorectal Cancer Disease-Specific Array that included genes specifically expressed in human colorectal cancers.¹¹ The Colorectal Cancer Disease-Specific Array content (61,528 total probe sets) was generated by a combination

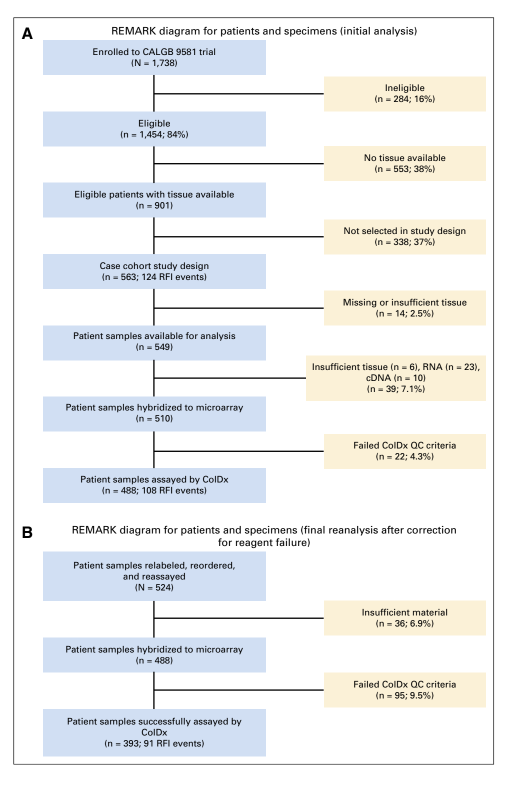


Fig 1. (A) REMARK diagram for patients and specimens (initial analysis). (B) REMARK diagram for patients and specimens (final analysis). CALGB, Cancer and Leukemia Group B; RFI, recurrence-free interval; QC, quality control; REMARK, REporting recommendations for tumor MARKer prognostic studies. of high-throughput sequencing, public database mining, and experimental investigation and was optimized to enable profiling of RNA extracted from fresh and FFPE samples. A set of 215 human, stage II colorectal cancers with known outcome data were identified as a training set and used to develop a 634-probe set, partial least squares prognostic signature, specific for high risk of recurrence. The probe sets and statistical methods used were previously described, and statistically significant pathways were identified.⁶ The analytical properties of this assay have also been reported.¹² The assay was developed and intended for use on FFPE, a commonly used, standard method of tumor tissue preservation.

The Cancer and Leukemia Group B (CALGB), now the Alliance for Clinical Trials in Oncology (Alliance), conducted a randomized phase III clinical trial (C9581) of 1,738 patients with stage II CC without high-risk features, which concluded that adjuvant edrecolomab did not significantly increase overall survival compared with observation only.¹³ The tissue and data from this clinical trial were archived for the purpose of biomarker analysis and thus provided a resource for validation of prognostic molecular markers in a large cohort of patients with stage II CC. The objective of this study was to use the C9581 trial samples to further assess the utility of the ColDx assay to classify patients with stage II CC as low or high risk of recurrence within 5 years postsurgery.

PATIENTS AND METHODS

Patients

Eligibility requirements for enrollment to the C9581 study have been reported previously.¹² Institutional review board approval and patient informed consent were required at each participating institution. The study excluded patients with high-risk features, and the overall risk of recurrence

observed in the study was 14.6% at 5 years.¹³ The eligibility requirements for the current study included: enrollment in C9581 with available FFPE tissue and patient consent for use of this tissue; a primary diagnosis of stage II adenocarcinoma of the colon; colon surgery with curative intent; negative surgical margins; no nonprotocol, postoperative therapy before recurrence; no preoperative therapy within 1 year of surgery; no local or regional recurrence only. Patients with rectal cancer were not eligible.

Of 1,454 patients who were eligible for the study, FFPE tissue was available for analysis from 901 patients. Among the 901 eligible patients with available tissue (Figs 1A and 1B), minimum follow-up time was 2.9 months and maximum follow-up time was 12.1 years. The incidence rate was 0.020 recurrences per person-year, with 124 (13.8%) RFI events and 777 (86.2%) censored observations. Median follow-up was 8.1 years (95% CI, 7.9 to 8.2 years) on the basis of the final clinical data set as of December 4, 2009.

Data from eligible patients treated on C9581 with available samples were uniquely coded and analyzed in a blinded fashion for gene expression. All statistical analyses involving the clinical data were conducted by Alliance statisticians.

Pathology

During the conduct of C9581, FFPE primary tumor and normal colon were obtained for each case, with histology confirmed by central pathology review. Staging was conducted as previously described.¹³ Of the samples within the final study cohort for this investigation, 85% had a viable tumor volume of \geq 50%. Specimens with a tumor volume of < 50% were macrodissected for inclusion within the study. DNA mismatch repair (MMR) status was assessed by immunohistochemistry for mut L homolog 1 (MLH1) and mut S homolog 2 (MSH2), also as previously described.⁷

Gene Expression Profiling

Total RNA was extracted from two $10-\mu m$ sections obtained from a single representative FFPE tissue block, amplified and hybridized to the Almac Colorectal Cancer disease-specific microarray in

 Table 1. Patient and Tumor Characteristics for 393 Patients in the Study Cohort, 1,061 Eligible Patients Not in the Study Cohort, and Total 1,454 Eligible Patients From the Parent Trial

	Value	Eligible Included in Study (n = 393)* No. (%)†‡	Eligible Patients Not Included in Study (n = 1,061)* No. (%)†‡	Parent C9581 Trial Eligible Patients (N = 1,454)*
Characteristic				No. (%)†
Age	Years	64 (23)	64 (13)	64 (11)
Sex	Male	209 (53)	554 (52)	763 (52)
Race/ethnicity	White	357 (91)	971 (91)	1,328 (92)
		(n = 392)	(n = 1,057)	(n = 1,449)
Treatment arm	Edrecolomab	192 (49)	528 (50)	720 (50)
T stage	T4	18 (5)	35 (3)	53 (4)
Nodes examined	No.	14.5 (17)	14.7 (12)	14.6 (9)
Perineural invasion	Present	10 (2)	27 (3)	37 (3)
		(n = 391)	(n = 1,056)	(n = 1,447)
Lymphovascular invasion	Present	47 (11)	115 (11)	162 (11)
Tumor grade	High	67 (17)	148 (14)	215 (15)
			(n = 1,054)	(n = 1,447)
Obstruction or perforation	Present	6 (2)	20 (2)	26 (2)
			(n = 1,060)	(n = 1,453)
Tumor location	Proximal	244 (62)	634 (59)	878 (60)
			(n = 1,060)	(n = 1,453)
MMR status§	Deficient	94 (26)	84 (18)	178 (21)
		(n = 377)	(n = 456)	(n = 833)

Abbreviation: MMR, mismatch repair.

*Sample size unless otherwise specified.

†Denotes mean and standard deviation for age and nodes examined.

 \pm Percentages weighted to reflect those in the entire eligible cohort (N = 1,454).¹²

\$MMR as determined by immunohistochemistry on mut L homolog 1 (MLH1), mut S homolog 2 (MSH2).

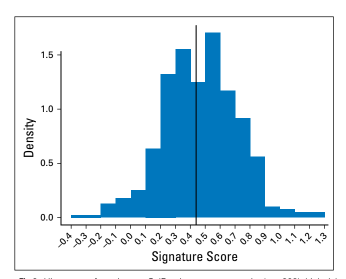


Fig 2. Histogram of continuous CoIDx signature score units (n = 393); high risk (≥ 0.4377) in 55% of patients. Data breaks occur between -0.4 and 1.3 at intervals of 0.1. For each interval, the density multiplied by 0.1 estimates the probability of the signature score being in that interval. The distribution mean and standard deviation are 0.46 and 0.24, respectively, with a median of 0.47 and range of -0.33 to 1.26.

a Clinical Laboratory Improvement Amendments (CLIA), New York State–approved laboratory (Helomics Corporation, Pittsburgh, PA). The ColDx gene expression signature was used to classify patients as low or high risk for recurrence by computing a signature score as described previously.⁶

Statistical Methods

The primary end point for this study was RFI measured from study entry (randomization) to distant recurrence or death related to colon cancer. Patients without distant recurrence or who died of other causes including second malignancies were censored. Overall survival (any cause of death) measured from study entry was studied as a secondary end point.

A case-cohort design was used to compare RFI between patients with ColDx-predicted low- and high-risk scores.^{14,15} Under this design, a randomized subcohort (RS) was selected from 901 eligible patients after stratification on treatment.¹⁶ The recurrent cases that were not selected randomly were included in the analysis data set. The case-cohort design was used to maximize cost efficiency by limiting the number of nonrecurrent patient samples to be assayed. Power estimates were determined on the basis of the methods proposed by Cai and Zeng.¹⁷ The initial study cohort comprised a subcohort of 514 patients randomly

Table 2. Univariable Analyses of Prognostic Factors					
Factor	HR (95% CI)	Р			
Age (continuous)	1.02 (1.00 to 1.04)	.03			
Sex (male v female)	0.97 (0.62 to 1.50)	.89			
T stage (T4 v T3)	1.15 (0.37 to 3.57)	.80			
Nodes examined (continuous)	0.97 (0.95 to 1.00)	.05			
Tumor grade (high <i>v</i> low)	1.30 (0.75 to 2.25)	.34			
Tumor location (proximal v distal)	1.11 (0.70 to 1.75)	.65			
MMR (deficient <i>v</i> intact)	0.56 (0.31 to 1.01)	.05			

NOTE. n = 393 with 91 RFI events for all variables except n = 377 with 91 RFI events for MMR.

Abbreviations: HR, hazard ratio; MMR, mismatch repair; RFI, recurrence-free interval.

sampled, including 75 patients with RFI events and 49 patients with RFI events outside the RS, resulting in a total of 563 patients (124 patients with RFI events).

For reasons described in the Appendix (online only), the study was amended. Power computations were revised on the basis of the following assumptions: 449 of 901 patients sampled in the RS using a sampling fraction of 0.4983; a proportion of samples marker positive, p1, of 0.20; a proportion of events in the full cohort, pD, of 0.1199 (108 of 901). Under these assumptions, a hazard ratio of 2.05 was detectable with approximately 80% power. Data from 393 patients were included in the final analysis.

Patients were categorized as low or high risk coded on the basis of a prognostic score of 0.4377, fixed before analysis involving patient outcomes. This threshold was selected after migration of the ColDx assay from the Affymetrix GeneChip System 3000 7G scanner to the Affymetrix microarray platform GeneChip System 3000Dx v.2.

The primary hypothesis that the ColDx classification was significantly associated with RFI independently of known prognostic factors was tested using a weighted Cox proportional hazards model adjusting for standard prognostic variables.¹⁵ This multivariable analysis included the coded prognostic score and the following potentially prognostic variables: age at study entry (years); sex (male, female); T stage (T3, T4); tumor grade (I-II, III-IV); tumor location (distal, proximal); number of nodes examined (continuous measure); MMR status (MMR-deficient, MMR-intact) determined by immunohistochemistry for MLH1 and MSH2, in 377 patients with complete data and 91 RFI events. Patients with missing data on any variable were omitted from the multivariable analysis. The likelihood ratio test was used to test the significance of the addition of the prognostic score to the reduced multivariable model. The association between RFI and only the coded prognostic score was also tested. Analyses of Schoenfeld residuals and regression methods were used to assess the proportional hazards assumption for each variable.

Data analysis was conducted in R using the Survival package by Therneau (Self-Prentice method)^{18,19} and in SAS, version 9.2, using the programming for case-cohort analysis proposed by Barlow et al.²⁰ Two-sided *P* values less than .05 were considered significant.

The χ^2 test was used to test the associations between ColDx and categorical prognostic factors; the *t* test was used to test associations with continuous factors. No adjustment was made for multiple comparisons in these analyses.

RESULTS

Sample Cohort

An initial cohort of 563 patients, with 124 RFI events, was selected for the study (comprising 514 patients with 75 RFI events

Factor	HR (95% CI)	Ρ
CoIDx signature score (high risk v low risk)	2.13 (1.3 to 3.5)	< .01
Age (continuous)	1.02 (1.00 to 1.04)	.07
Sex (male <i>v</i> female)	0.94 (0.59 to 1.47)	.78
T stage (T4 v T3)	0.78 (0.20 to 3.0)	.72
Nodes examined (continuous)	0.98 (0.95 to 1.01)	.11
Tumor grade (high <i>v</i> low)	1.47 (0.82 to 2.6)	.19
Tumor location (proximal v distal)	1.24 (0.74 to 2.1)	.41
MMR (deficient v intact)	0.55 (0.29 to 1.02)	.05

Abbreviations: HR, hazard ratio; MMR, mismatch repair; RFI, recurrence-free interval.

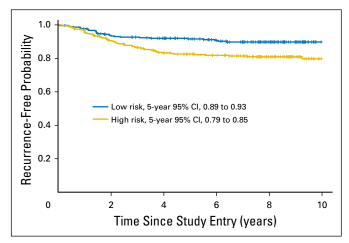


Fig 3. Weighted Kaplan-Meier plot of recurrence-free interval by CoIDx signature score dichotomized at the prespecified cut point, 0.4377; 62 and 29 events observed for high and low risk, respectively.

in the RS and 49 patients with RFI events outside the RS). Tissue was available from 549 of these patients for analysis. A quality control (QC) failure arose in a subset of patient samples because of a reagent fault during the initial analysis, and the study was subsequently repeated with qualified reagents (Data Supplement).

Of the initial cohort of 549 tissue samples available for analysis, a total of 156 samples (28.4%) were excluded from the analysis because of insufficient tissue, total RNA, or poor quality. Median (range) time between resection and analysis was 13.2 years (11.2 to 16.0 years). The final evaluable data set in the repeated analysis contained 393 patients, of whom 91 experienced an RFI event (360 patients with 58 RFI events within the RS; 33 patients with RFI events outside the RS). Figures 1A and 1B depict the flow of patients through the study and reasons for exclusion.

The demographic characteristics of the C9581 cohort have been described.^{7,13} Patient and tumor characteristics for patients evaluated in the study were similar to those not included and to those in the subset of all eligible C9581 patients (Table 1).

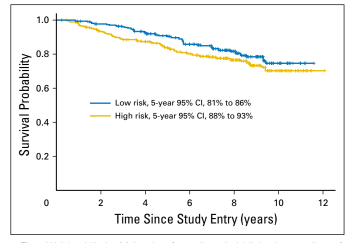


Fig 4. Weighted Kaplan-Meier plot of overall survival (all deaths regardless of cause) by CoIDx signature score dichotomized at the prespecified cut point, 0.4377; 64 and 40 events observed for high and low risk, respectively.

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Association Between ColDx and Prognostic Factors

Results of preprocessing and derivation of the ColDx signature score were confirmed by the Alliance statisticians. Figure 2 provides a histogram of the distribution of signature scores. At the prespecified threshold, ColDx classified 55% of patients (216 of 393) as high risk and 45% (177 of 393) as low risk.

Associations between the dichotomized signature score (high risk, low risk) and the potential prognostic variables in Table 1 were explored. ColDx was significantly associated with sex (P = .003), race (P < .001), and MMR status (P < .001) with higher proportions of men (57% v 49%), nonwhite race (73% v 52%), and MMR intact tumors (56% v 45%) classified as high risk, respectively. Associations with T stage, lymphovascular invasion, and tumor location were marginally significant (Appendix Table A1).

Association Between ColDx and Recurrence Risk

Results of univariable analyses of individual prognostic factors and RFI are given in Table 2. The primary test of hypothesis was the assessment of the ColDx score in the presence of conventional prognostic factors (Table 3). ColDx score was significantly associated with RFI after adjustment for other prognostic factors (hazard ratio [HR], 2.13; 95% CI, 1.3 to 3.5; P < .01) in multivariable analysis and was the prognostic factor most strongly associated with RFI (P < .01) in this data set. Age and MMR status were of borderline significance. The ColDx signature score contributed significantly to the reduced (prognostic variables only) model (likelihood ratio test, P < .001). No major deviations from proportional hazards were observed on the basis of the Schoenfeld residuals and regression models methods.

ColDx high-risk patients also had a significantly shorter RFI relative to ColDx low-risk patients in an unadjusted analysis, with an HR of 2.03 (95% CI, 1.3 to 3.3; P < .01). The recurrence-free probability at 5 years for patients classified as high risk by the ColDx score was 82%, with a 95% CI of 79% to 85%, compared with 91% with a 95% CI of 89% to 93% for patients predicted as low risk (Fig 3).

OS by ColDx score is illustrated in Figure 4. A marginally significant association was observed after adjustment for other prognostic factors in multivariable analysis (OS HR, 1.74; 95% CI, 0.97 to 3.1; P = .06).

DISCUSSION

Given the lower rates of recurrence among patients with stage II CC, there is reluctance to subject patients to potential toxicity and other treatment complications with little evidence of treatment benefit from adjuvant therapy. Current guidelines indicate that adjuvant therapy may be most appropriate for patients with high-risk disease characteristics.³ Substantial effort has been devoted to identifying a higher-risk patient subset under the assumption that patients at high risk of recurrence would be the most likely to benefit from adjuvant therapy. CoIDx is one of several assays that have been reported to reliably stratify patients with colon cancer into groups at high and low risk of disease recurrence.

The ColDx assay has previously been shown to be an independent prognostic factor for RFI in stage II CC.⁶ In this second independent prospectively designed study, the ColDx assay was again demonstrated to be significantly associated with RFI in the stage II colon cancer C9581 cohort and to be the most significant prognostic factor in multivariable analysis, in the presence of other prognostic factors, inclusive of MMR status. The data and samples collected as part of the C9581 study protocol were obtained from patients who were uniformly treated and well characterized. As such, this cohort provided an invaluable resource for objective assessment of novel prognostic markers such as the ColDx assay.

However, a limitation of this study was the age of the tissue samples, which may have contributed to the high proportion of assay failures. (A maximum attrition rate of 10% was originally expected on the basis of experience in a number of prior studies.) On average, the length of time between patient resection and assay analysis was 13.2 years. In addition, results of the first assay series were affected by a reagent failure in a subset of patient samples, which was discovered only after the first statistical analysis was conducted. A plan to salvage the validation and maintain the integrity of the study was developed in conjunction with the National Cancer Institute Cancer Therapy Evaluation Program and incorporated into the protocol. Under this plan, specimens were returned to the Alliance Pathology Coordinating Office, relabeled, and reanalyzed by Helomics Corporation. The final statistical analysis was conducted on the resulting expression data. This process was transparent, and we are confident in the final results.

The study demonstrated that patients within the C9581 cohort predicted as high risk by the ColDx assay have a recurrencefree probability of 82% (18% probability of recurrence within 5 years) compared with 91% (9% probability of recurrence within 5 years) for patients predicted as low risk with an HR after multivariable analysis of 2.13. This compares favorably with the previously reported clinical performance of the assay, where it was demonstrated to predict high-risk patients with an HR of recurrence of 2.53 and an HR of cancer-related death of 2.21 in an independent validation data set.⁶ The ColDx assay was significantly associated with RFI in this study (P < .01), providing greater prognostic value than standard clinical markers, including age (P = .07), number of nodes examined (0.11), and, notably, MMR status (P = .05). Analysis of these clinical markers is currently the primary means by which clinicians determine the clinical management of patients with stage II colon cancer.

The prognostic performance of ColDx is likely derived from its relationship to biologic pathways previously reported to differentiate between good and poor prognosis in colon cancer via promotion of tumor growth, invasion, and metastasis that are not accounted for by traditional prognostic factors.⁶ In addition, although ColDx is a significant predictor of RFI in its own right, the addition of the ColDx score significantly contributes to a prognostic model including all clinical covariates (likelihood ratio test, P < .001), which demonstrates its additive contribution.

There have been several prognostic gene expression assays for stage II colon cancer.⁷⁻¹⁰ Only the 12-gene colon cancer recurrence score is comparable to ColDx, because it has also been developed to work with FFPE tissue and has undergone independent validation using the stage II C9581 sample cohort.⁷ The 12-gene colon cancer recurrence score was also found to be the most significant prognostic factor among patients studied from C9581 (HR, 1.68; 95% CI, 1.18 to 2.38; P = .004), with the HR calculated on a continuous scale for a 25-unit increase in the score. In the subset of 271 patients for whom data are available from both assays, there is low correlation between the continuous scores (R = 0.18; 95% CI, 0.06 to 0.29). Although there is some overlap, it does not seem that the signatures are measuring the same thing or identifying the same patients as high risk. Further analysis is being conducted in this patient subset.

Prognostic assays, such as ColDx, would have enhanced clinical utility if they were also predictive of response to therapy in colon cancer.²¹ Other than a marginal treatment effect in the Quick and Simple and Reliable (QUASAR) trial,²² there have never been convincing data showing benefit for adjuvant therapy in any study of stage II disease for CC, including C9581. A retrospective subset analysis of the stage II cohort in Intergroup 0035 suggested that fluorouracil therapy might be beneficial in patients with clinical high-risk factors, such as T4 lesions, obstruction, inadequate nodal sampling, and perforation.²³ Although that hypothesis-generating observation has never been validated in a prospective manner, the practice has been embraced and incorporated into guidelines and clinical care. To support clinical utility, a direction for future research would be to show that prognostic gene expression signatures are also predictive for treatment efficacy.

In summary, the ColDx assay has now been prospectively validated for a second time as a prognostic marker for patients with stage II colon cancer and in this patient subset superior to current prognostic markers such as T stage, nodal sampling, and MMR status. This study is then an external validation of the prognostic value of the ColDX assay. Thus, ColDx assay results could be incorporated with the traditional clinical markers of risk to refine prognosis. Future studies could be designed to demonstrate the potential for treatment benefit among high-risk patients defined by the ColDx risk score.

AUTHORS' DISCLOSURES OF POTENTIAL CONFLICTS OF INTEREST

Disclosures provided by the authors are available with this article at www.jco.org.

AUTHOR CONTRIBUTIONS

Conception and design: Donna Niedzwiecki, Wendy L. Frankel, Alan P. Venook, Paula N. Friedman, Richard M. Goldberg, Robert J. Mayer, Jude Marie Mulligan, Timothy S. Davison, Peter Kerr, Patrick G. Johnston, Richard D. Kennedy, D. Paul Harkin, Monica M. Bertagnolli, Robert S. Warren

Administrative support: Thomas Anthony Colacchio

Provision of study materials or patients: Richard M. Goldberg, Thomas Anthony Colacchio

Collection and assembly of data: Donna Niedzwiecki, Richard M. Goldberg, Thomas Anthony Colacchio, Jude Marie Mulligan, Peter Kerr **Data analysis and interpretation:** Donna Niedzwiecki, Alan P. Venook, Xing Ye, Richard M. Goldberg, Robert J. Mayer, Jude Marie Mulligan, Timothy S. Davison, Eamonn O'Brien, Richard D. Kennedy, Richard L. Schilsky, Robert S. Warren, Federico Innocenti **Manuscript writing:** All authors

Final approval of manuscript: All authors

REFERENCES

1. Edge S, Byrd D, Compton C, et al. AJCC Cancer Staging Manual (ed 7). New York, NY, Springer, 2010

2. Benson AB III, Schrag D, Somerfield MR, et al: American Society of Clinical Oncology recommendations on adjuvant chemotherapy for stage II colon cancer. J Clin Oncol 22:3408-3419, 2004

3. National Comprehensive Cancer Network: NCCN Clinical Practice Guidelines in Oncology, Colon Cancer v2.2015. http://www.nccn.org

4. Van Cutsem E, Oliveria J, ESMO Guidelines Working Group: Primary colon cancer: ESMO clinical recommendations for diagnosis, adjuvant treatment and follow-up. Ann Oncol 20:49-50, 2009 (suppl 4)

 Vicuna B, Benson AB III: Adjuvant therapy for stage II colon cancer: Prognostic and predictive markers. J Natl Compr Canc Netw 5:927-936, 2007

 Kennedy RD, Bylesjo M, Kerr P, et al: Development and independent validation of a prognostic assay for stage II colon cancer using formalin-fixed paraffinembedded tissue. J Clin Oncol 29:4620-4626, 2011

7. Venook AP, Niedzwiecki D, Lopatin M, et al: Biologic determinants of tumor recurrence in stage II colon cancer: Validation study of the 12-gene recurrence score in cancer and leukemia group B (CALGB) 9581. J Clin Oncol 31:1775-1781, 2013 8. Salazar R, Roepman P, Capella G, et al: Gene expression signature to improve prognosis prediction of stage II and III colorectal cancer. J Clin Oncol 29: 17-24, 2011

9. Agesen TH, Sveen A, Merok MA, et al: ColoGuideEx: A robust gene classifier specific for stage II colorectal cancer prognosis. Gut 61:1560-1567, 2012

10. Zhang JX, Song W, Chen ZH, et al: Prognostic and predictive value of a microRNA signature in stage II colon cancer: A microRNA expression analysis. Lancet Oncol 14:1295-1306, 2013

11. Kerr P, Black J, von Frese J, et al: Molecular profiling from FFPE tissue: Stratification of the stage II colorectal cancer population. Mol Cancer Ther 6: 3751S, 2007 (abstr C138)

 Plamadeala V, Huang S, McCreary SM, et al: Analytical performance of a formalin-fixed paraffinembedded tissue-based 634-probe prognostic assay for predicting outcome of patients with stage II colon cancer. Appl Immunohistochem Mol Morphol 22: 308-316, 2014

 Niedzwiecki D, Bertagnolli MM, Warren RS, et al: Documenting the natural history of patients with resected stage II adenocarcinoma of the colon after randomization of adjuvant treatment with edrecolomab or observation: results from CALGB 9581. J Clin Oncol 29:3146-3152, 2011

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14. Prentice RL: A case-cohort design for epide-

15. Gray RJ: Weighted analyses for cohort sam-

16. Borgan O, Langholz B, Samuelsen SO, et al:

17. Cai J, Zeng D: Sample size and power calculation

18. Therneau TR: Package "survival." Repository

19. Self S, Prentice RL: Asymptotic distribution

20. Barlow WE, Ichikawa L, Rosner D, et al: Analysis of

21. Benson AB 3rd, Hamilton SR. Path toward

22. Quasar Collaborative Group, Gray R, Barnwell

23. Moertel CG, Fleming TR, Macdonald JS, et al:

theory and efficiency results for case-cohort studies.

case-cohort designs. J Clin Epidemiol 52:1165-1172, 1999

prognostication and prediction: An evolving matrix. J

J, et al: Adjuvant chemotherapy versus observation in

patients with colorectal cancer: A randomised study.

Intergroup study of fluorouracil plus levamisole as adjuvant therapy for stage II/Dukes' B2 colon cancer.

Exposure stratified case-cohort designs. Lifetime

for case-cohort studies. Biometrics 60:1015-1024, 2004

miologic cohort studies and disease prevention trials.

pling designs. Lifetime Data Anal 15:24-40, 2009

Biometrika 73:1-11, 1986

Data Anal 6:39-58, 2000

Ann Stat 16:64-81, 1988

CRAN. http://r-forge.r-project.org

Clin Oncol 29:4599-4601, 2011

Lancet 370:2020-2029, 2007

J Clin Oncol 13:2936-2943, 1995

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Appendix

Administrative Summary of Study Progress

Selection of initial study sample. Five hundred fourteen patients were randomly selected from the available 901 patients meeting the entry criteria. Seventy-five patients meeting the recurrence end point were included in the random component of the study sample. Under the case-cohort design, the remaining 49 patients meeting the recurrence end point were also part of the study sample. A total of 563 patients were identified for potential inclusion in the study. Five hundred forty-nine patient samples were provided to Helomics Corporation. There were 14 blocks that were either missing or unable to be cut because they were insufficient.

Microarray analysis results. Of the 549 samples provided by the Alliance, sufficient total RNA was extracted to allow for gene expression profiling of 520 samples. Among the 29 samples that were not profiled by gene expression, six were unable to be extracted because of lack of tissue availability and 23 because of insufficient RNA. Of these 520 samples, 510 samples were hybridized to a microarray generating raw data (CEL files). Ten samples were not hybridized to a microarray because of insufficient cDNA. Of these 510 samples with microarray data generated, 488 passed ColDx quality control (QC) metrics (a total of 22 QC failures).

An Excel file containing the following data fields was provided to the Alliance for the 510 samples that generated microarray data, including the 22 QC failures: Alliance Pathology Coordinating Office Number; Almac ID assigned to each sample by Almac (for correlation to raw CEL files); ColDx signature score; dichotomized clinical call (low risk, high risk, QC failure). The subgroup of 488 samples passing ColDx QC metrics comprised 449 samples, including 69 events in the randomized sample and 39 events outside of the randomized sample.

Statistical analysis of ColDx data. Results of the statistical analysis were not expected. To verify that the results were correct, the laboratory and clinical data merge, recurrence-free interval end points, and statistical analyses were reviewed. Patient-level data were provided to Almac for exploration of potential issues related to QC, macrodissection, and other laboratory procedures. In this process, a strong batch effect related to a specific reagent lot number (1201176-B) was noted in a subset of samples. It was concluded that a lower-than-expected hazard ratio was observed overall because of the reagent failure in this subset. The reagent lot batch effect was observed at the global gene expression level (principal component analysis), and this effect persisted through to the signature score. Other technical factors were also evaluated and eliminated as causes.

A plan to salvage the validation and maintain the integrity of the study was developed in conjunction with National Cancer Institute-Cancer Therapy Evaluation Program and incorporated into the protocol. Under this plan, all specimens were to be returned to the Alliance PCO, relabeled, and reanalyzed by Helomics Corporation, in a blinded fashion. It was agreed that initial assessments of patients (514 in the randomly selected subcohort [RS], stratified by treatment) would not be used, and the final statistical analysis would be conducted on the expression data resulting from reanalysis of the specimens.

Residual material was identified for 524 samples, which were subsequently relabeled, reordered, and reassayed with reagents that passed QC; 36 had insufficient material and 95 failed ColDx QC. The final set of samples with new ColDx assay values comprised 393 patients (360 in the randomly selected subcohort, with 58 recurrence-free interval events and 33 events outside the RS).

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Factor	Percent High Risk*	P†
Age (mean age; high <i>v</i> low risk)	64 v 63 years	.43
Sex (male <i>v</i> female)	57% v 49%	.003
Race (white v other)	52% v 73%	< .001
Treatment (edrecolomab v observation)	53% v 54%	.66
T stage (T4 v T3)	65% <i>v</i> 53%	.04
Nodes examined (mean No. of nodes; high <i>v</i> low risk)	14 v 14 nodes	.89
Perineural invasion (absent <i>v</i> present)	53% v 60%	.46
Lymphovascular invasion (absent v present)	52% v 63%	.01
Tumor grade (high v low)	53% v 53%	.95
Tumor location (proximal <i>v</i> distal)	51% v 57%	.03
Obstruction perforation (absent v present)	53% v 67%	.17
MMR (deficient v intact)	45% v 56%	< .001

Abbreviation: MMR, mismatch repair. *Except for continuous variables age and number of nodes examined, where means are given. $^{+}\chi^{2}$ test except for continuous variables, where the *t* test was used.