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INTESTINAL RESEARCH

Development and validation of a scoring system for advanced colorectal neoplasm in young Korean subjects less than age 50 years

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Background/Aims: Colorectal cancer incidence among patients aged \leq 50 years is increasing. This study aimed to develop and validate an advanced colorectal neoplasm (ACRN) screening model for young adults aged <50 years in Korea. **Methods:** This retrospective cross-sectional study included 59,575 consecutive asymptomatic Koreans who underwent screening colonos-copy between 2003 and 2012 at a single comprehensive health care center. Young Adult Colorectal Screening (YCS) score was developed as an optimized risk stratification model for ACRN using multivariate analysis and was internally validated. The predictive power and diagnostic performance of YCS score was compared with those of Asia-Pacific Colorectal Screening (APCS) and Korean Colorectal Screening (KCS) scores. **Results:** 41,702 and 17,873 subjects were randomly allocated into the derivation and validation cohorts, respectively, by examination year. ACRN prevalence was 0.9% in both cohorts. YCS score comprised sex, age, alcohol, smoking, obesity, glucose metabolism abnormality, and family history of CRC, with score ranges of 0 to 10. In the validation cohort, ACRN prevalence was 0.6% in the low-risk tier (score, 0–4), 1.5% in the moderate-risk tier (score, 5–7), and 3.4% in the high-risk tier (score, 8–10). ACRN risk increased 2.5-fold (95% confidence interval [CI], 1.8–3.4) in the moderate-risk tier and 5.8-fold (95% CI, 3.4–9.8) in the high-risk tier compared with the low-risk tier. YCS score identified better balanced accuracy (53.9%) than APCS (51.5%) and KCS (50.7%) scores and had relatively good discriminative power (area under the curve = 0.660). **Conclusions:** YCS score based on clinical and laboratory risk factors was clinically effective and beneficial for predicting ACRN risk and targeting screening colonoscopy in adults aged <50 years. **(Intest Res 2019;17:253-264)**

Key Words: Colorectal neoplasm; Risk assessment; Early detection of cancer; Young adults

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INTRODUCTION

Colorectal cancer (CRC) is the one of the most common cancer and cause of cancer-related death worldwide, with 1.3 million new cases and 700,000 deaths annually.¹ The incidence and mortality of CRC have also been increasing in Korea, with agestandardized incidence of 37.3/100,000 and mortality rate of 9.7/100,000.² Most CRCs arise from premalignant lesions, such as colorectal adenoma (CRA) and especially, advanced adeno-

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ma, for 5 to 10 years.³ Therefore, if CRA is detected and treated by screening colonoscopy, most CRCs can be prevented.⁴⁵

In many countries, including Korea, fecal occult blood test (FOBT) and colonoscopy have been used for CRC screening starting at 50 years of age.⁶ Even with repeat FOBT, significant number of CRCs and premalignant adenomas may be missed owing to low sensitivity.⁷ Colonoscopy is considered the most sensitive and convenient method for CRC screening because it can treat CRA in one stage. However, it is an expensive and invasive procedure that can induce bleeding and perforation. Therefore, identifying high risk subjects with advanced colorectal neoplasm (ACRN) who need screening colonoscopy is very important.

Several risk scoring models have been recently developed for identifying high risk subjects who need screening colonoscopy.⁸⁹ However, these models targeted subjects aged >50 years. The incidence and prevalence of CRC have been increasing in subjects aged <50 years, and the need for developing models targeting this age group is being raised.¹⁰⁻¹² Therefore, this study aimed to develop and validate a scoring system for ACRN in a large cohort comprising Korean subjects aged <50 years who underwent screening colonoscopy.

METHODS

1. Study Population for Development of the Risk Score

Asymptomatic subjects aged <50 years who underwent a health examination between 2003 and 2012 at a single comprehensive health care center were enrolled. Exclusion criteria were as follows: incomplete colonoscopy, history of CRC or other cancers, history of IBD, history of previous colonoscopy, colorectal surgery, or missing clinical or laboratory data. Finally, 59,575 subjects were included with different characteristics by examination years. Thus, the dataset was randomly divided into the derivation (70%) or validation (30%) cohort by examination years. The derivation cohort included 41,702 subjects, and the validation cohort included 17,873 subjects.

This study was approved by the Institutional Review Board of Kangbuk Samsung Hospital (IRB No., 2013-01-119) and written informed consents were obtained.

2. Measurements and Diagnosis of Colorectal Neoplasm

Data on medical history, medication use, and health-related behaviors were collected using a self-administered questionnaire under the supervision of a well-trained interviewer. Data on alcohol consumption and smoking were noted. A heavy drinker was defined as a subject who drinks >4 times per week regularly. Family history of CRC was defined as CRC in 1 or more first-degree relatives at any age. Weekly frequency of moderate to vigorous physical activity was also assessed.

Physical measurements and serum biochemical parameters were measured by trained nurses. The Asia-Pacific criteria for obesity based on BMI guidelines were used to diagnose obesity (BMI ≥ 25 kg/m²).¹³ Blood pressure was measured using a standard mercury sphygmomanometer with subjects seated after at least 10 minutes of rest. Blood samples were obtained from the antecubital vein after at least 10 hours of fasting. Colonoscopies were performed by experienced colonoscopists who were unaware of the present study. Bowel preparations were performed using 4 L of polyethylene glycol solution. Histological assessment of all polyps was performed by experienced pathologists who were unaware of the subjects' clinical data. ACRN was defined as CRA ≥ 10 mm in diameter, CRA with any component of villous histology, high-grade dysplasia, or carcinoma.¹⁴

3. Development of the Risk Scoring System

The authors developed the Young Colorectal Screening (YCS) score to identify high risk subjects aged <50 years for ACRN. Univariable analysis was performed for the derivation cohort using the chi-square test or *t*-test to assess the association between clinical variables and ACRN. Variables associated with ACRN in the univariable analyses (P < 0.05) were included in the multivariable logistic regression analysis. Potential risk variables that were significant in the multivariable analysis with stepwise selection procedure based on Akaike information criterion were included in the risk score. We assigned weighted points to each risk factor. Each risk point was rescaled to designate the point of current obesity as one (e.g., the risk point for age between 40 and 50 years was 3, which was rounded from 0.676/0.485 = 2.808). The risk score for an individual was the summation of their individual risk factors. The validity of the score was assessed by receiver operating characteristic (ROC) curve analysis.

4. Calculation and Validation of the Risk Scoring System

Each individual had a personal risk score calculated by the R software that summed the points attributed based on the presence of a risk factor in the individual. The score was calculated using the R software at the data center after data were sent from individual clinical study sites. The performance of the risk scoring system was evaluated by comparing the OR of the high-risk

Table 1. Characteristics of Participants in the Derivation and Validation Cohorts

Characteristic	Derivation cohort (n=41,702)	Validation cohort (n = 17,873)	<i>P</i> -value
Age (yr)	38.9±5.3	38.9±5.4	0.878
< 30	345 (0.8)	163 (0.9)	
30-39	21,776 (52.2)	9,397 (52.6)	
40-49	19,581 (47.0)	8,313 (46.5)	
Sex			0.679
Female	11,933 (28.6)	5,145 (28.8)	
Male	29,769 (71.4)	12,728 (71.2)	
Alcohol drinking			
No	12,878 (30.9)	5,563 (31.1)	
Non-heavy drinker	27,548 (66.0)	11,741 (65.7)	0.495
Heavy drinker ^a	1,276 (3.1)	569 (3.2)	0.566
Smoking			
Never	23,091 (55.4)	9,864 (55.2)	
Former smoker	6,646 (15.9)	2,927 (16.4)	0.231
Current smoker	11,965 (28.7)	5,082 (28.4)	0.789
Family history of colorectal	cancer		0.445
No	40,242 (96.5)	17,224 (96.4)	
Yes	1,460 (3.5)	649 (3.6)	
Exercise			0.608
No	18,436 (44.2)	7,860 (44.0)	
Yes ^b	23,266 (55.8)	10,013 (56.0)	
BMI (kg/m²)	23.8±3.2	23.8 ± 3.2	0.820
Obesity			0.578
No	27,757 (66.6)	11,939 (66.8)	
Yes ^c	13,945 (33.4)	5,934 (33.2)	
Diabetes			0.247
No	32,605 (78.2)	14,051 (78.6)	
Yes	9,097 (21.8)	3,822 (21.4)	
Insulin (µIU/mL)	5.1 ± 3.3	5.1±3.3	0.952
HbA1c (%)	5.6 ± 0.5	5.6±0.4	0.363
< 6.5	40,681 (97.5)	17,413 (97.4)	
≥ 6.5	1,021 (2.5)	460 (2.6)	
Glucose (mg/dL)	92.9±13.7	92.9±13.7	0.565
< 100	33,472 (80.3)	14,383 (80.5)	
≥ 100	8,230 (19.7)	3,490 (19.5)	
Total cholesterol (mg/dL)	199.8±34.2	199.0 ± 34.5	0.053
< 200	19,189 (46.0)	9,465 (53.0)	
≥ 200	22,513 (54.0)	8,408 (47.0)	

(Continued to the next)

Table 1. Continued

Characteristic	Derivation cohort (n = 41,702)	Validation cohort (n = 17,873)	P-value
HDL-cholesterol (mg/dL)	55.1 ± 13.7	55.3 ± 13.9	0.065
\geq 40 (male), \geq 50 (female)	35,602 (85.4)	15,287 (85.5)	
<40 (male), <50 (female)	6,100 (14.6)	2,586 (14.5)	
LDL-cholesterol (mg/dL)	124.0±31.7	124.3±31.82	0.296
< 100	9,441 (22.6)	4,071 (22.8)	
≥ 100	32,261 (77.4)	13,802 (77.2)	
Triglyceride (mg/dL)	115.1±76.2	115.7±78.1	0.375
≤ 150	32,712 (78.4)	13,962 (78.1)	
>150	8,990 (21.6)	3,911 (21.9)	
SBP (mmHg)	113.0±12.9	112.9±12.9	0.799
DBP (mmHg)	72.3 ± 9.6	72.3 ± 9.5	0.943
ACRN			0.999
No	41,309 (99.1)	17,704 (99.1)	
Yes	393 (0.9)	169 (0.9)	
Colorectal cancer			0.999
No	41,684 (99.96)	17,866 (99.96)	
Yes	18 (0.04)	7 (0.04)	

Values are presented as mean ± SD or number (%).

^aMore than 4 times per week as heavy.

^bMore than once per week.

^cBMI ≥ 25 kg/m².

HbA1c, hemoglobin A1c; SBP, systolic blood pressure; DBP, diastolic blood pressure; ACRN, advanced colorectal neoplasm.

(HR) and moderate-risk (MR) tiers versus the low-risk (LR) tier or HR tier versus MR and LR tiers from 3 categories.

5. Statistical Analyses

In the derivation cohort, univariable analysis was performed to assess the associations between clinical variables and ACRN using the chi-square test for categorical variables, namely sex, alcohol consumption, smoking, family history of CRC, exercise, obesity, and diabetes, and the *t*-test for continuous variables, namely age, HDL-cholesterol, low-density lipoproteincholesterol, total cholesterol, systolic blood pressure, diastolic blood pressure, insulin, triglyceride, hemoglobin A1c (HbA1c), and glucose. Multivariable logistic regression analysis with stepwise selection was used to select ACRN predictors. Initial candidate variables were those with *P*-value < 0.05 in the univariable analyses. To predict the presence of ACRN, we designed a prediction rule with the clinical variables selected in the multivariable analyses. The Hosmer-Lemeshow statistic

Table 2. Univariate and Multivariate Analyses in the Derivation Cohort

Variable	Total cohort	ACRN (n = 393)	Non-ACRN (n = 41,309)	Univariate analysis <i>P</i> -value	Multivariate analysis <i>P</i> -value
Age (yr)		41.1±5.0	38.9 ± 5.3	< 0.001	< 0.001
Sex				0.016	0.164
Female	11,933	91 (0.8)	11,842 (99.2)		
Male	29,769	302 (1.0)	29,467 (99.0)		
Alcohol					
No	12,878	141 (1.1)	12,737 (98.9)		
Non-heavy drinker	27,548	240 (0.9)	27,308 (99.1)	0.030	0.005
Heavy drinker ^a	1,276	12 (0.9)	1,264 (99.1)	0.611	0.209
Smoking					
Never	23,091	204 (0.9)	22,887 (99.1)		
Former smoker	6,646	49 (0.7)	6,597 (99.3)	0.254	0.330
Current smoker	11,965	140 (1.2)	11,825 (98.8)	0.010	0.009
Family history of colorectal cancer				0.537	0.991
No	40,242	377 (0.9)	39,865 (99.1)		
Yes	1,460	16 (1.1)	1,444 (98.9)		
Exercise				0.592	0.798
No	18,436	179 (1.0)	18,257 (99.0)		
Yes ^b	23,266	214 (0.9)	23,052 (99.1)		
Obesity				< 0.001	0.114
No	27,757	227 (0.8)	27,530 (99.2)		
Yes ^c	13,945	166 (1.2)	13,779 (98.8)		
Diabetes				< 0.001	0.678
No	32,605	269 (0.8)	32,336 (99.2)		
Yes	9,097	124 (1.4)	8,973 (98.6)		
Insulin (FB)		5.6 ± 3.4	5.1±3.3	0.003	0.750
HbA1c ≥ 6.5%				< 0.001	0.015
No	40,681	367 (0.9)	40,314 (99.1)		
Yes	1,021	26 (2.5)	995 (97.5)		
Glucose ≥ 100 (mg/dL)				< 0.001	0.262
No	33,472	277 (0.8)	33,195 (99.2)		
Yes	8,230	116 (1.4)	8,114 (98.6)		
Total cholesterol (mg/dL)		204.4±35.4	198.4±34.2	< 0.001	0.413
HDL-cholesterol (mg/dL)		53.3±12.9	55.1 ± 13.7	0.008	0.501
LDL-cholesterol (mg/dL)		129.4±31.8	124.0±31.7	< 0.001	0.242
Triglyceride (mg/dL)		130.6±85.9	115.0±76.1	< 0.001	0.255
SBP (mmHg)		113.0 ± 13.4	113.0±12.9	0.890	0.698
DBP (mmHa)		72.2±9.7	72.3±9.6	0.325	0.469

Values are presented as mean \pm SD or number (%).

^aMore than 4 times per week as heavy.

^bMore than once per week.

^cBMI ≥ 25 kg/m².

ACRN, advanced colorectal neoplasm; FB, fasting blood; HbA1c, hemoglobin A1c; SBP, systolic blood pressure; DBP, diastolic blood pressure.

was used to determine the goodness-of-fit of the logistic regression model, a low *P*-value (<0.05) indicating a lack of fit of the model. A new logistic regression model was implemented with transformed variables, which were obtained by converting continuous variables. The risk prediction rule was developed from the new logistic regression equations using the beta coefficient-based scoring method. We assigned weighted points to each risk factor. On the basis of the lowest beta coefficient, other factors were rescaled and rounded to the nearest integer. The total score for each individual was calculated by summing the scores of all individual risk factors. The area under the ROC curve was then computed to assess the ability of the risk scoring system.

In the derivation cohort, we chose the score that was divided into 3 tiers (HR, MR, and LR) as the cutoff value and then calculated the sensitivity, specificity, and balanced accuracy of the prediction rule to evaluate predictive accuracy. The prediction rule was also validated in the validation cohort. We compared the performance of the risk prediction rule, as measured by sensitivity, specificity, and balanced accuracy, against existing methods such as Asia-Pacific colorectal Screening (APCS) score⁹ and Korean Colorectal Screening (KCS) score.¹⁵ To compare the statistical ability of the YCS, APCS, and KCS scores in predicting the risk of ACRN in the validation cohort, ROC analysis and McNemar test were performed. Statistical analyses were performed using R and MedCalc for Windows, version 16.2 (MedCalc Software, Ostend, Belgium).

RESULTS

1. Characteristics of Subjects in the Derivation and Validation Cohorts

The deviation cohort included 41,702 asymptomatic subjects (mean age, 38.9 ± 5.3 years; males, 71.4%), of which 393 (0.9%)

Table 3. Predictors of Advanced Colorectal Neoplasm in the New Logistic Regression Model and the Associated Prediction Rule in the Derivation Cohort (n = 41,702)

Variable	Coefficients	OR (95% CI)	<i>P</i> -value	Points assigned
Sex				
Female		1 (Reference)		0
Male	0.1996	1.22 (0.95–1.56)	0.113	1
Age (yr)				
< 40		1 (Reference)		0
40-49	0.7142	2.04 (1.66–2.52)	< 0.001	3
Alcohol				
Never or non-heavy drinker		1 (Reference)		0
Heavy drinker ^a	0.3394	1.40 (1.13–1.74)	0.002	1
Smoking				
Never or former smoker		1 (Reference)		0
Current smoker	0.3443	1.41 (1.14–1.75)	0.002	1
Obesity				
No		1 (Reference)		0
Yes ^b	0.2543	1.29 (1.04–1.59)	0.018	1
HbA1c \geq 6.5% or glucose \geq 100 mg/dL				
No		1 (Reference)		0
Yes	0.4377	1.55 (1.24–1.93)	< 0.001	2
Family history of colorectal cancer				
No		1 (Reference)		0
Yes	0.0843	1.09 (0.66–1.80)	0.743	1

^aMore than 4 times per week as heavy.

^bBMI ≥25 kg/m².

HbA1c, hemoglobin A1c.

had ACRN and 18 (0.04%) had CRC. The baseline characteristics of the patients are described in Table 1. The validation cohort included 17,873 asymptomatic subjects (mean age, 38.9 ± 5.4 years; males, 71.2%), of which 169 (0.9%) had ACRN and among them, 7 (0.04%) had CRC (Table 1).

2. Univariate and Multivariate Analyses of Risk Factors for ACRN in Derivation Cohort

The risk factors for ACRN were assessed by univariable and multivariable analyses (Table 2). In the multivariable analysis, age (P < 0.001), non-heavy drinker (P = 0.005), current smoker (P = 0.009), and HbA1c level $\ge 6.5\%$ (P = 0.015) were significantly associated with the risk of ACRN (Table 2).

3. New Logistic Regression Model and Development of the Risk Score

In the present study, the risk factors associated with ACRN were assessed originally using the new logistic regression model to develop a precise and practical risk scoring model. Male sex (OR, 1.22; 95% CI, 0.95–1.56), age 40 to 49 years (OR, 2.04; 95% CI, 1.66–2.52), heavy drinker (OR, 1.40; 95% CI, 1.13–1.74), current smoking (OR, 1.41; 95% CI, 1.14–1.75), obesity (OR, 1.29; 95% CI, 1.04–1.59), glucose level \geq 100 mg/dL or HbA1c \geq 6.5% (OR, 1.55; 95% CI, 1.24–1.93), and family history of CRC (OR, 1.09; 95% CI, 0.66–1.80) were included in the YCS scoring model for ACRN. Male sex and family history of CRC were not statistically significant factor in the present study; however, these factors were revealed to be important risk factors in previous studies. Therefore, 1 point was assigned for each factor.

Table 4. Distribution of Participants for Each Advanced Colorectal Neoplasm Risk Score

Pick cooro	APC	S score	KCS	score	YCS	score
NISK SCOLE	Total	ACRN	Total	ACRN	Total	ACRN
Derivation cohort						
0	9,151	71 (0.8)	7,935	58 (0.7)	2,088	14 (0.7)
1	21,840	188 (0.9)	15,092	100 (0.7)	6,344	20 (0.3)
2	9,629	122 (1.3)	14,120	183 (1.3)	6,776	38 (0.6)
3	772	6 (0.8)	4,413	50 (1.1)	5,794	54 (0.9)
4	310	6 (1.9)	142	2 (1.4)	7,049	50 (0.7)
5					6,710	90 (1.3)
6					3,609	55 (1.5)
7					2,108	50 (2.4)
8					1,094	18 (1.6)
9					128	4 (3.1)
10					2	0
Validation cohort						
0	3,943	26 (0.7)	3,406	22 (0.6)	931	4 (0.4)
1	9,360	72 (0.8)	6,476	37 (0.6)	2,745	8 (0.3)
2	4,080	57 (1.4)	6,111	80 (1.3)	2,984	16 (0.5)
3	372	8 (2.2)	1,825	27 (1.5)	2,428	23 (0.9)
4	118	6 (5.1)	55	3 (5.5)	3,011	22 (0.7)
5					2,804	39 (1.4)
6					1,508	25 (1.7)
7					933	14 (1.5)
8					456	16 (3.5)
9					71	2 (2.8)
10					2	0

Values are presented as number (%).

APCS, Asia-Pacific Colorectal Screening; KCS, Korean Colorectal Screening; YCS, Young adult Colorectal Screening; ACRN, advanced colorectal neoplasm.

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A point for the YCS score of each risk factor was assigned as follows: female sex (0), male sex (1), age <40 years (0), age 40-49 years (3), never or non-heavy drinker (0), heavy drinker (1), never or former smoker (0), current smoker (1), non-obesity (0), obesity (1), glucose level <100 mg/dL and HbA1c <6.5% (0), HbA1c $\geq 6.5\%$ or glucose level ≥ 100 mg/dL (1), absent of family history of CRC (0), and present of family history of CRC (1) (Table 3).

The YCS score was represented with the sum of points for risk factors present in an individual in the present study. The score was composed of a range from 0 to 10 points. Subjects with ACRN showed an increasing tendency as the risk score increased proximately. However, no subject with a score of 10 had ACRN (Table 4). The score was classified into 3 tiers according to the risk of ACRN. Scores from 0 to 4 were assigned as LR tier, that from 5 to 7 were assigned as MR tier, and that from 8 to 10 were assigned as HR tier. There were 28,051 subjects (67.3%) in the LR tier, 23,427 subjects (29.8%) in the MR tier, and 1,224 subjects (2.9%) in the HR tier. Subjects with scores of MR and HR had increased risks of ACRN (OR, 2.5; 95% CI, 2.1-3.1 and OR, 2.9; 95% CI, 1.9-4.5, respectively) (Table 5). The prevalence of ACRN was 0.6% in the LR tier, 1.6% in the MR tier, and 1.8% in the HR tier. The area under the curve (AUC) for the ACRN risk score in the derivation cohort was 0.641, indicating good discrimination.

4. Validation of the Risk Scoring Model for ACRN in **Young Adults**

Among the 17,873 subjects in the validation group, 12,099 (67.7%) were classified into the LR tier, 5,245 (29.3%) into the MR tier, and 529 (3.0%) into the HR tier. In the derivation cohort, 176 subjects (0.6%) had ACRN in the LR tier, 195 (1.6%) had ACRN in the MR tier, and 22 (1.8%) had ACRN in the HR tier. In the validation cohort, 78 subjects (0.6%) had ACRN in the LR tier, 78 (1.5%) had ACRN in the MR tier, and 18 (3.4%) had ACRN in the HR tier. In the validation cohort, the risk of ACRN was higher in the MR tier (OR, 2.5; 95% CI, 1.8-3.4) and HR tier (OR, 5.8; 95% CI, 3.4–9.8) than in the LR tier (Table 5). The Hosmer-Lemeshow goodness-of-fit statistic was used to test the validation cohort, and a P-value of 0.261 indicated a fair match of predicted risk over observed risk. The AUC value was 0.660, indicating good discrimination.

5. Diagnostic Performance of ACRN by Risk Category in Young Adults

Diagnostic performance of the YCS score for detecting ACRN

		APCS score			KCS score			YCS score	
KISK	Total	ACRN	OR (95% CI)	Total	ACRN	OR (95% CI)	Total	ACRN	OR (95% CI)
Derivation cohort									
Average	30,991 (74.4)	259 (0.8)	Reference	23,027 (55.3)	158 (0.7)	Reference	28,051 (67.3)	176 (0.6)	Reference
Moderate	10,401 (24.9)	128 (1.2)	1.5 (1.2–1.8)	18,533 (44.4)	233 (1.3)	1.8 (1.5–2.3)	12,427 (29.8)	195 (1.6)	2.5 (2.1–3.1)
High	310 (0.7)	6 (1.9)	2.3 (1.0–5.3)	142 (0.3)	2 (1.4)	2.1 (0.5–8.4)	1,224 (2.9)	22 (1.8)	2.9 (1.9–4.5)
Total	41,702 (100.0)	393 (0.9)		41,702 (100.0)	393 (0.9)		41,702 (100.0)	393 (0.9)	
Validation cohort									
Low	13,303 (74.4)	98 (0.7)	Reference	9,882 (55.3)	59 (0.6)	Reference	12,099 (67.7)	73 (0.6)	Reference
Moderate	4,452 (24.9)	65 (1.5)	2.0 (1.5–2.7)	7,936 (44.4)	107 (1.3)	2.3 (1.7–3.1)	5,245 (29.3)	78 (1.5)	2.5 (1.8–3.4)
High	118 (0.7)	6 (5.1)	7.2 (3.1–16.8)	55 (0.3)	3 (5.5)	9.6 (2.9–31.6)	529 (3.0)	18 (3.4)	5.8 (3.4–9.8)
Total	17,873 (100.0)	169 (0.9)		17,873 (100.0)	169 (0.9)		17,873 (100.0)	169 (0.9)	

APCS score risk category: low (0–1), moderate (2–3), high (4–7); KCS score risk category: low (0–1), moderate (2–3), high (4–8); YCS score risk category: low (0–4), moderate (5–7), high (8–10). Guid ۰ ה Val AP

in the validation cohort was also assessed. The YCS score was compared with the APCS and KCS scores. Participants for the APCS score were enrolled in 11 Asian cites and had a mean age of 54.4 ± 11.6 years. Points were assigned to each risk factor for ACRN as follows: age \geq 70 years (3), age 50–69 years inclusive (2), age < 50 years (0), male sex (1), female sex (0), familv history of CRC in a first-degree relative present (2) or absent (0), non-smoking (0), and smoking (1). The score ranged from 0 to 7 and was divided into 3 risk tiers: score 0-1, low risk (LR); score 2-3, MR; and score 4-7, HR. The mean age of patients enrolled for the KCS score was 51.3 ± 9.0 years, and the points of KCS score were assigned as follows: age \geq 70 years (4), age 50–69 yeas inclusive (2), age < 50 years (0), male sex (1), female sex (0), family history of CRC in a first-degree relative present (1) or absent (0), current or past smoker (1), nonsmoker (0), BMI $\ge 25 \text{ kg/m}^2$ (1), and BMI $< 25 \text{ kg/m}^2$ (0). This score ranged from 0 to 8 and was divided into 3 risk tiers: score 0-1, LR; score 2-3, MR; and score 4-8, HR. On comparing between HR tier and MR+LR tier, the newly proposed YCS score showed slightly lower specificity (97.1%) than the APCS

Table 6. Diagnostic Performance of the Risk Score for Detection

 of Advanced Colorectal Neoplasm in the Validation Cohort

	APCS score	KCS score	YCS score
HR vs. MR+LR			
Sensitivity	3.6	1.8	10.7
Specificity	99.4	99.7	97.1
Balanced accuracy	51.5	50.7	53.9
MR+HR vs. LR			
Sensitivity	42.0	65.1	56.8
Specificity	74.6	55.5	67.9
Balanced accuracy	58.3	60.3	62.4

Values are presented as percentage.

APCS, Asia-Pacific Colorectal Screening; KCS, Korean Colorectal Screening; YCS, Young adult Colorectal Screening; HR, high risk; MR, moderate risk; LR, low risk.

Table 7. Results of the Comparison of Methods by McNemar Te	est
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and KCS scores (99.4% and 99.7%, respectively). However, sensitivity (10.7%) and balanced accuracy (53.9%) were more precise than those of the APCS score (3.6% and 51.5%) and KCS score (1.8% and 50.7%). On comparison between MR+HR tier and LR tier, balanced accuracy was also higher in the YCS score (62.4%) than in the APCS score (58.3%) and KCS score (60.3%) (Table 6). McNemar test results showed that the YCS score (10.7%) had a higher sensitivity than the APCS (3.6%, P=0.014) and KCS (1.8%, P=0.001) scores when the HR tier was compared with MR+LR tier. Specificity in the YCS score (97.1%) was lower than that in the APCS (99.4%, P<0.001) and KCS (99.7%, P<0.001) scores. When HR+MR tier was compared with LR tier, YCS score showed higher sensitivity (56.8%) than



Fig. 1. ROC curve for YCS, APCS, and KCS score models. AUC for the risk score was significantly higher in the YCS score (0.660; 95% Cl, 0.65–0.77) than APCS score (0.595; 95% Cl, 0.59–0.60; P=0.007) and KCS score (0.606; 95% Cl, 0.60–0.61; P=0.012), when results of APCS and KCS score were deducted by using the validation cohort of the present study. ROC, receiver operating characteristic; AUC, area under the curve; YCS, Young Adult Colorectal Screening; APCS, Asia-Pacific Colorectal Screening; KCS, Korean Colorectal Screening.

HR vs. MR+LR				HR+M	R vs. LR		
Method ₁	$Method_2$	Sensitivity	Specificity	Method ₁	Method ₂	Sensitivity	Specificity
APCS	KCS	0.248	< 0.001	APCS	KCS	< 0.001	< 0.001
APCS	YCS	0.014	< 0.001	APCS	YCS	0.002	< 0.001
KCS	YCS	0.001	< 0.001	KCS	YCS	0.052	< 0.001

HR, high risk; MR, moderate risk; LR, low risk; APCS, Asia-Pacific Colorectal Screening; KCS, Korean Colorectal Screening; YCS, Young adult Colorectal Screening.

APCS score (42.0%, P=0.002) and higher specificity (67.9%) than KCS score (55.5%, P<0.001) (Table 7). In the ROC analysis, AUC for the risk score was significantly higher in the YCS score (0.660; 95% CI, 0.65–0.67) than the APCS (0.595; 95% CI, 0.59–0.60; P=0.007) and KCS (0.606; 95% CI, 0.60–0.61; P=0.012) scores when the results of APCS and KCS score were obtained using the validation cohort of the present study (Fig. 1).

DISCUSSION

Early detection and treatment of CRC is associated with a reduction in CRC mortality. Screening colonoscopy and polypectomy have decreased the incidence and mortality of CRC.^{5,16} CRC and precancerous ACRN should be targeted by CRC screening. The present study developed and validated the YCS score with multiple clinical and laboratory factors, including age, sex, social behaviors, metabolic factors, and family history of CRC. This is the first study to assess the ACRN screening score foe patients aged <50 years in a large Asian cohort.

Many studies have evaluated age as an important risk factor for the development of CRA and CRC.^{8,17} Most studies revealed that the prevalence of CRC increased in subjects aged ≥ 50 years; therefore, these subjects were considered as targets for CRC screening in most of countries.^{6,18} However, there is a chance of CRC developing in adults aged <50 years, and 1% of asymptomatic subjects had ACRN, including precancerous lesions, in the present study. A previous study reported that ACRN was found in 0.7% of patients in the 30 to 39-year age group, which increased to 2.7% of patients in the 40 to 49-year age group.¹⁹ It is difficult to screen all young adults due to the lower prevalence of CRC in young adults than that in older adults considering the cost-benefit problems and complications. Therefore, it is important to identify high risk subjects and also include young adults in the screening. In the present study, the risk of ACRN increased as the age increased, even in young adults aged < 50 years. Therefore, screening of ACRN should be considered for young adults as they age, especially for subjects with other risk factors.

Previous studies evaluated risk factors of ACRN, including age,^{8,14} male sex,²⁰ smoking,²¹ and family history of CRC in firstdegree relatives.²² Several CRC screening scoring models including those factors were developed.²³⁻²⁵ Recently, metabolic factors, including obesity,²⁶ diabetes mellitus, and dyslipidemia,²⁷ were reported as a risk factor of CRA and CRC. However, a small number of scoring models included metabolic factors, and few models used quantitative laboratory results. In the present study, metabolic factors, including obesity, glucose metabolism abnormality, dyslipidemia, and blood pressure, were quantitatively evaluated, and using these factors, the YCS score was developed. The risk of ACRN increased in subjects with obesity and HbA1c \geq 6.5% or glucose level \geq 100 mg/dL. Obesity influences the development of CRC,²⁸ and BMI was included as a risk factor in previous CRC screening scoring systems.^{8,23,29} In Asia, obesity is less prevalent, and the mean BMI is also lower than that in Western countries.³⁰ In our study, the overall mean BMI was 23.8 ± 3.2 kg/m², and extremely obese subjects with BMI > 30 kg/m^2 were rare. Obesity increased the risk of ACRN through insulin resistance and chronic inflammation.³¹ BMI was not included in the APCS score; however, it was included in the KCS score. In the YCS score, BMI was assigned 1 point, and this more clearly reflected the increased risk of ACRN according to metabolic abnormality. Glucose metabolism is also related to CRC. Insulin-like growth factor-1 (IGF-1) level increases due to hyperinsulinemia, resulting from insulin resistance, and elevated IGF-1 levels affect the incidence of CRA and CRC, inducing proliferation and dysplasia of normal and carcinoma cells owing to insulin resistance abnormality.³² Meta-analysis of published studies supported a protective association between antidiabetic medications and CRC risk in patients with diabetes mellitus.³³ Strict regulation of HbA1c and fasting glucose levels might be more important in the development of CRA and CRC. The risk of ACRN might be increased with glucose metabolism abnormality before the development of diabetes mellitus. Some previous studies reported that abdominal obesity and metabolic syndrome seemed to be related to the risk of colorectal neoplasm in younger subjects but not in older individuals.^{31,34} Therefore, in subject aged < 50 years, the YCS score, which included obesity and HbA1c and fasting glucose levels, could be more accurate and valuable for CRC screening.

In the present study, multiple factors, which were previously known as risk factors for ACRN, were accurately evaluated by questionnaires and laboratory tests. Laboratory tests used in the YCS score were generally used in the clinical practice. They could be quantitatively measured and represented the present patient's clinical state. The YCS score was constructed using multivariate analysis with multiple risk factors, so that it was valuable in the practical clinical field. The YCS score comprised factors that could be obtained relatively easily with precise information from subjects. It was an easily calculated score, and the stratification of risk groups was also simple. All study subjects were asymptomatic, of low risk, had first colonosco-

py, enrolled in a screening setting, and were representative of our general screening target population. Therefore, the YCS score can be helpful for CRC screening that targets subjects who need a colonoscopy and are aged < 50 years.

In the present study, around 33% subjects belonged to the MR+HR tier, and the prevalence of ACRN was 1.5% among those subjects. In case of the HR tier, the proportion of subjects was 3%, and the prevalence of ACRN was around 2% to 3%. In the study of APCS score, the prevalence of ACRN in subjects aged >50 years was around 3.5% to 5%,9 and in another study of our group, which comprised 10,000 asymptomatic subjects aged >50 years, the total prevalence of ACRN was around 3.7%, that of male patients was around 4.7%, and that of female patients was around 2.3%.³⁵ This prevalence of ACRN was similar to that of the HR tier in the present study. Considering the prevalence of ACRN and medical cost-effectiveness in asymptomatic subjects aged <50 years, it was effective that colonoscopy screening was recommended in the HR tier of YCS score, and FOBT might be recommended first in the MR tier. When results of the APCS and KCS scores were obtained using the validation cohort of the present study, only 0.3% to 0.7% of total subjects were classified into the HR tier, which represented a relatively small proportion for colonoscopic screening. The sensitivity was only lower than 4%. A previous study using the KCS score recommended that adults aged <50 years undergo CRC screening if their scores place them in either the MR or HR tier.¹⁵ However, the proportion of subjects in the MR and HR tiers in the KCS score was around 45%. The targets of screening colonoscopy will increase when KCS scores are used to screen subjects aged < 50 years. When we considered cost-effectiveness, KCS scoring could be ineffective in screening for ACRN because of the broad target population. When we compared between HR and MR+LR (or MR+ LR) among the 3 scores, the specificity of YCS score was slightly lower than those of APCS and KCS scores. However, sensitivity and balanced accuracy were significantly higher of YCS score than of APCS and KCS scores. Sensitivity is more important for screening than specificity. Specificity was only 2% lower for YCS score than for APCS or KCS score. Specificity is also less crucial than sensitivity because of the low prevalence of ACRN in subjects aged < 50 years. The YCS score was also more discriminative than the APCS and YCS scores in the ROC analysis. Therefore, YCS score would be more effective and valuable for assessment of high risk you adult subjects for ACRN screening.

The present study had several limitations. It was a cross-sec-

tional study with a single ethnic group. The prevalence of ACRN in Asian countries is lower than that in Western countries, and it is definitely low in subjects aged <50 years.^{34,36} The absolute value and prevalence of obesity are higher in Western countries than in Asian countries, and the metabolic profile is also probably different. The results of our study cannot be generally applied. However, the YCS score, which included metabolic factors, might be useful in Western subjects because of the high prevalence of ACRN and metabolic abnormality in them. Validation of the YCS score with various subjects are needed. Data on smoking, alcohol intake, and exercise were evaluated by simple questionnaires and not quantitatively. Data collection was difficult because this was a large-scale populationbased study. However, we collected various anthropometric measurements and metabolic laboratory factors, which are frequently used in clinical practice for all study participants. This provided valuable risk factors, which are crucial for determining screening strategies. Considering the distribution of participants for the YCS score, 2 subjects each had a score of 10 in the derivation and validation cohorts; however, ACRN was not detected in subjects with a score of 10. This could be a result of the small sample size of the score group; however, those subjects might not have developed ACRN yet because they were younger than 50 years and there could be a great possibility of developing ACRN after 50 years of age. AUC of the YCS score increased statistically compared with that of the APCS and KCS scores. However, the increment in AUC was only about 0.05, and AUC of 0.660 itself did not seem very high. The prediction of the risk stratification of ACRN was not easy, so AUC of previous scoring systems, including APCS and KCS scores, were only 0.6. However, AUC of YCS score increased to about 0.05 using simple clinical factors that could be easy to identify. Some of the false positive and negative results were reduced through increased AUC, the cost-effectiveness of the ACRN screening could be increased.

In conclusion, the YCS score was developed and validated for screening risks of ACRN using age, sex, alcohol consumption, smoking state, obesity, glucose metabolism abnormality, and family history of CRC in a large cohort of asymptomatic subjects aged <50 years. The prevalence of ACRN was lower in subjects aged <50 years than in those aged \geq 50 years. However, in the HR tier of the YCS score, risks of ACRN increased even in subjects aged <50 years. Therefore, the YCS score could be valuable for colonoscopic screening among young adults.

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CONFLICT OF INTEREST

No potential conflict of interest relevant to this article was reported.

AUTHOR CONTRIBUTION

Kim JY and Choi S analyzed the data and wrote the manuscript as first authors. Park DI organized and supervised the study as corresponding author. Jung YS, Park JH, Kim SK, and Kim HJ collected and arranged the data. Cho YK, Sohn CI, and Jeon WK attended to analyze the data. Park T supported the statistical analysis. Kim BI attended to revise the manuscript. Choi KY attended to make study concept and design. Approval of final manuscript: all authors.

REFERENCES

- 1. Ferlay J, Soerjomataram I, Dikshit R, et al. Cancer incidence and mortality worldwide: sources, methods and major patterns in GLOBOCAN 2012. Int J Cancer 2015;136:E359-E386.
- 2. Jung KW, Won YJ, Kong HJ, et al. Cancer statistics in Korea: incidence, mortality, survival, and prevalence in 2012. Cancer Res Treat 2015;47:127-141.
- St-Onge MP, Janssen I, Heymsfield SB. Metabolic syndrome in normal-weight Americans: new definition of the metabolically obese, normal-weight individual. Diabetes Care 2004;27: 2222-2228.
- Winawer SJ, Zauber AG, O'Brien MJ, et al. Randomized comparison of surveillance intervals after colonoscopic removal of newly diagnosed adenomatous polyps: the National Polyp Study Workgroup. N Engl J Med 1993;328:901-906.
- 5. Nishihara R, Wu K, Lochhead P, et al. Long-term colorectalcancer incidence and mortality after lower endoscopy. N Engl J Med 2013;369:1095-1105.
- 6. Levin B, Lieberman DA, McFarland B, et al. Screening and surveillance for the early detection of colorectal cancer and adenomatous polyps, 2008: a joint guideline from the American Cancer Society, the US Multi-Society Task Force on Colorectal Cancer, and the American College of Radiology. Gastroenterology 2008;134:1570-1595.

- Sheng JQ, Li SR, Wu ZT, et al. Transferrin dipstick as a potential novel test for colon cancer screening: a comparative study with immuno fecal occult blood test. Cancer Epidemiol Biomarkers Prev 2009;18:2182-2185.
- 8. Freedman AN, Slattery ML, Ballard-Barbash R, et al. Colorectal cancer risk prediction tool for white men and women without known susceptibility. J Clin Oncol 2009;27:686-693.
- 9. Yeoh KG, Ho KY, Chiu HM, et al. The Asia-Pacific Colorectal Screening score: a validated tool that stratifies risk for colorectal advanced neoplasia in asymptomatic Asian subjects. Gut 2011;60:1236-1241.
- Kim JK, Choi YS, Suh JP, Lee IT, Youk EG, Lee DS. Results of screening colonoscopy in asymptomatic average-risk Koreans at a community-based secondary hospital. Korean J Gastrointest Endosc 2010;41:266-272.
- 11. Austin H, Henley SJ, King J, Richardson LC, Eheman C. Changes in colorectal cancer incidence rates in young and older adults in the United States: what does it tell us about screening. Cancer Causes Control 2014;25:191-201.
- Siegel RL, Fedewa SA, Anderson WF, et al. Colorectal cancer incidence patterns in the United States, 1974-2013. J Natl Cancer Inst 2017;109.
- 13. World Health Organization, Regional Office for the Western Pacific; International Diabetes Institute; International Association for the Study of Obesity; International Obesity Task Force. The Asia-pacific perspective: redefining obesity and its treatment. Sydney: Health Communications Australia, 2000.
- Heitman SJ, Ronksley PE, Hilsden RJ, Manns BJ, Rostom A, Hemmelgarn BR. Prevalence of adenomas and colorectal cancer in average risk individuals: a systematic review and meta-analysis. Clin Gastroenterol Hepatol 2009;7:1272-1278.
- 15. Kim DH, Cha JM, Shin HP, Joo KR, Lee JI, Park DI. Development and validation of a risk stratification-based screening model for predicting colorectal advanced neoplasia in Korea. J Clin Gastroenterol 2015;49:41-49.
- Kahi CJ, Imperiale TF, Juliar BE, Rex DK. Effect of screening colonoscopy on colorectal cancer incidence and mortality. Clin Gastroenterol Hepatol 2009;7:770-775.
- Rundle AG, Lebwohl B, Vogel R, Levine S, Neugut AI. Colonoscopic screening in average-risk individuals ages 40 to 49 vs 50 to 59 years. Gastroenterology 2008;134:1311-1315.
- Sung JJ, Lau JY, Young GP, et al. Asia Pacific consensus recommendations for colorectal cancer screening. Gut 2008;57:1166-1176.
- 19. Chung SJ, Kim YS, Yang SY, et al. Prevalence and risk of colorectal adenoma in asymptomatic Koreans aged 40-49 years un-

dergoing screening colonoscopy. J Gastroenterol Hepatol 2010; 25:519-525.

- 20. Wong MC, Lam TY, Tsoi KK, et al. A validated tool to predict colorectal neoplasia and inform screening choice for asymptomatic subjects. Gut 2014;63:1130-1136.
- 21. Shin A, Hong CW, Sohn DK, et al. Associations of cigarette smoking and alcohol consumption with advanced or multiple colorectal adenoma risks: a colonoscopy-based case-control study in Korea. Am J Epidemiol 2011;174:552-562.
- 22. Regula J, Rupinski M, Kraszewska E, et al. Colonoscopy in colorectal-cancer screening for detection of advanced neoplasia. N Engl J Med 2006;355:1863-1872.
- Tao S, Hoffmeister M, Brenner H. Development and validation of a scoring system to identify individuals at high risk for advanced colorectal neoplasms who should undergo colonoscopy screening. Clin Gastroenterol Hepatol 2014;12:478-485.
- 24. Park HW, Han S, Lee JS, et al. Risk stratification for advanced proximal colon neoplasm and individualized endoscopic screening for colorectal cancer by a risk-scoring model. Gastrointest Endosc 2012;76:818-828.
- Cai QC, Yu ED, Xiao Y, et al. Derivation and validation of a prediction rule for estimating advanced colorectal neoplasm risk in average-risk Chinese. Am J Epidemiol 2012;175:584-593.
- Ben Q, An W, Jiang Y, et al. Body mass index increases risk for colorectal adenomas based on meta-analysis. Gastroenterology 2012;142:762-772.
- 27. Jinjuvadia R, Lohia P, Jinjuvadia C, Montoya S, Liangpunsakul S. The association between metabolic syndrome and colorectal neoplasm: systemic review and meta-analysis. J Clin Gas-

troenterol 2013;47:33-44.

- 28. Harriss DJ, Atkinson G, George K, et al. Lifestyle factors and colorectal cancer risk (1): systematic review and meta-analysis of associations with body mass index. Colorectal Dis 2009; 11:547-563.
- 29. Kim KS, Moon HJ, Choi CH, et al. The frequency and risk factors of colorectal adenoma in health-check-up subjects in South Korea: relationship to abdominal obesity and age. Gut Liver 2010;4:36-42.
- 30. WHO Expert Consultation. Appropriate body-mass index for Asian populations and its implications for policy and intervention strategies. Lancet 2004;363:157-163.
- Gunter MJ, Leitzmann MF. Obesity and colorectal cancer: epidemiology, mechanisms and candidate genes. J Nutr Biochem 2006;17:145-156.
- 32. Eddi R, Karki A, Shah A, DeBari VA, DePasquale JR. Association of type 2 diabetes and colon adenomas. J Gastrointest Cancer 2012;43:87-92.
- 33. Singh S, Singh H, Singh PP, Murad MH, Limburg PJ. Antidiabetic medications and the risk of colorectal cancer in patients with diabetes mellitus: a systematic review and meta-analysis. Cancer Epidemiol Biomarkers Prev 2013;22:2258-2268.
- 34. Hong SN, Kim JH, Choe WH, et al. Prevalence and risk of colorectal neoplasms in asymptomatic, average-risk screenees 40 to 49 years of age. Gastrointest Endosc 2010;72:480-489.
- 35. Kim JY, Jung YS, Park JH, et al. Different risk factors for advanced colorectal neoplasm in young adults. World J Gastroenterol 2016;22:3611-3620.
- 36. Choe JW, Chang HS, Yang SK, et al. Screening colonoscopy in asymptomatic average-risk Koreans: analysis in relation to age and sex. J Gastroenterol Hepatol 2007;22:1003-1008.