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KRT17 as a prognostic biomarker in stage II colorectal cancer.

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論文内容要旨

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<th>学位論文題名</th>
<th>KRT17 as a prognostic biomarker in stage II colorectal cancer.（stage II大腸癌における予後予測バイオマーカーとしてのKRT17）</th>
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大腸癌治療において手術は主要な役割を担う。Stage I症例は手術単独で良好な予後を得られるが、stage IIIでは手術に加え術後補助化学療法を行うことで予後が改善することが明らかにされている。一方、stage IIでの術後補助化学療法の有用性は確立されておらず、臨床病理学的な再発高リスク群に対して化学療法を行うのが現時点での妥当な戦略とされているが、その適応について未だ明確なコンセンサスはない。本研究の目的は、stage II大腸癌術後の再発高リスク症例を判別するバイオマーカーの作成である。大規模かつ複数の大腸癌コホートを用いた網羅的発現解析と免疫組織染色を行い、再現性の高い分子マーカーの抽出と、簡便かつ日常的な手技への応用を試みた。9つの独立したコホート、いずれも根治手術を施行された総計997例のstage II大腸癌症例を用いた。計458症例、4つのマイクロアレイコホート（Microarray Discovery set）の網羅的発現解析により、各コホートに共通して再発の有無と有意に関連し発現変動する6つの遺伝子が抽出され、そのうちKRT17遺伝子を含む4つの遺伝子発現は無再発生存期間と有意に関連していた。なお4つの遺伝子のうちKRT17遺伝子のみが、89例の独立したコホート（Microarray Validation set 1）において有意に無再発生存期間と関連していた。さらに、93例のマイクロアレイコホート（Microarray Validation set 2）、および183例のRNAシークエンスコホート（RNA-seq Validation set）において、KRT17発現は無再発生存期間と有意に関連していた。多変量解析により、KRT17遺伝子の高発現は、臨床高リスク因子のデータを持つコホートにおいて、他の臨床高リスク因子に対し独立して不良な無再発生存期間と関連していた。これらの転写レベルでのKRT17発現と予後不良との関連を、免疫組織染色（IHC）を用いてタンパク発現レベルでの検出に応用した。110例および44例の2つの大腸癌FFPEコホートに対しKRT17の免疫組織染色を行った（IHC Validation set 1、IHC Validation set 2）。いずれのコホートでもKRT17染色陽性症例は有意に不良な無再発生存と関連していた。また多変量解析により、KRT17染色は既知の臨床病理学的な再発高リスク因子と独立して無再発生存と関連していた。以上より複数のコホートを用いた多段階の検証を経て、新規の再発関連遺伝子としてKRT17を同定し、極めて高い再現性を示した。したがってKRT17染色は、stage II大腸癌術後の再発リスクを層別化するための予後バイオマーカーとして、臨床におけるstage II大腸癌患者の個別化医療に有用であることが示唆される。
Abstract

The mainstay of treatment for patients with stage I-III (locoregional) colorectal cancer is surgical resection. Large randomized trials revealed that adjuvant treatment significantly improves survival in stage III patients. However, the role of adjuvant treatment for stage II CRC remains inconclusive, despite numerous clinical trials and meta-analyses. This study aimed to develop clinically useful biomarkers that can identify a subset of stage II patients with high risk of relapse after surgery.

We investigated comprehensive expression analysis using several cohorts of colorectal cancer and immunohistochemical staining and attempted to extract highly reproducible molecular markers and apply it conveniently to routine procedures.

We analyzed 997 stage II colorectal cancer patients in nine independent cohorts in this study. We initially analyzed four independent Affymetrix microarray cohorts of stage II patients as discovery datasets. Six genes were found as candidate genes which were associated with relapse. Furthermore, four genes were associated with Relapse free survival. KRT17 was found to be the only gene whose expression was significantly associated RFS in 89 stage II patients in Microarray Discovery set. Moreover, KRT17 expression was significantly associated with relapse-free survival in 93 patients in Microarray Validation cohort and 183 patients in RNA Sequence cohort.

To investigate whether the expression of KRT17 in protein levels could also identify stage II patients with poor RFS, survival analysis was conducted in two IHC Validation set. Kaplan-Meier analysis showed that patients with KRT17-positive tumors had significantly shorter RFS after surgery, compared to those of KRT17-negative. We identified and validated the expression of KRT17 transcript or KRT17 protein as a prognostic biomarker for RFS with high reproducibility. Our study suggests that KRT17 IHC provides additional prognostic stratification beyond the currently-available high-risk factors, highlighting the potential of its performance in precision management for stage II patients in the clinical setting.

Introduction

The number of deaths caused by colorectal cancer (CRC) ranked second in all cancers in 2015 worldwide [1, 2]. CRC is a highly heterogeneous disease associated with complex molecular mechanisms, thus clinicopathologically similar patients can strikingly differ in survival outcomes. The mainstay of treatment for patients with stage I-III (locoregional) disease is surgical resection [3]. Stage I CRC patients have good prognosis treated by surgery alone. It is recommended that all stage III patients receive adjuvant chemotherapy after surgery, as large randomized trials revealed that adjuvant treatment significantly improves survival in these patients [3-5]. However, the role of adjuvant treatment for stage II CRC remains inconclusive, despite numerous clinical trials and meta-analyses [3, 6-9]. Given the clear benefits of adjuvant treatment seen in stage III patients, recent efforts have
been focused on discriminating stage II patients with poor prognosis who might benefit from adjuvant chemotherapy or intensive postoperative management, since approximately 15% of stage II patients develop recurrence even after curative surgery [4].

Expert groups, including the American Society of Clinical Oncology (ASCO) [10], the National Comprehensive Cancer Network (NCCN) [11], the European Society of Medical Oncology (ESMO) [12], and the Japanese Society for Cancer of the Colon and Rectum (JSCCR) [13], published separate guidelines for adjuvant treatment of stage II CRC with considerable variations. Despite the lack of direct evidence from randomized trials to support the benefit for adjuvant therapy in those patients [6], guidelines recommend consideration of adjuvant chemotherapy for stage II patients with high-risk clinicopathologic features, including T4 tumors, poorly differentiated histology, lymphatic/vascular/perineural invasion, obstruction/perforation, inadequately sampled lymph nodes, although the definition of "high-risk" is not standardized [6]. NCCN guidelines further stratified stage II patients based on microsatellite instability (MSI)/mismatch repair (MMR) status, in which MSI-high/deficient MMR patients do not require adjuvant treatment [11]. However, a study of approximately 25,000 patients with stage II disease in the Surveillance, Epidemiology, and End Results (SEER) database revealed that high-risk clinicopathologic features alone are insufficient for treatment decision [14]. Therefore, there is a clear clinical need for prognostic assays to facilitate the selection of stage II patients for additional treatment.

To address this question, many studies have exploited gene expression profiling for stratifying CRC patients to provide an accurate and personalized assessment of the risk of relapse, and multi-gene assays have been developed based on dozens or hundreds of genes [6]. Many assays have been shown to be promising, however, they are often difficult to translate into routine diagnostic tests, possibly because of low technical reproducibility, the requirement of fresh-frozen tissues, or the lack of validation in large, prospective trials [15-21]. Moreover, many of them were not specifically designed to target stage II disease [18, 20]. As a result, none of the previously reported prognostic gene signatures have been adopted in clinical practice.

From these points of view, we consider that molecular biomarkers are needed to be developed with clear focus on the intended clinical populations, to have high reproducibility across multiple cohorts, and to be easily applicable in clinical practice by simple methods. Therefore, this study aimed to develop clinically useful biomarkers that can identify a subset of stage II patients with high risk of relapse after surgery. Using a total of 997 patients with stage II CRC obtained from 9 independent cohorts, we conducted a step-wise strategy with high-throughput technology-based screening and validation, followed by immunohistochemistry (IHC) using formalin-fixed paraffin-embedded (FFPE) tissue samples, resulting in the identification of KRT17 protein (keratin 17, cytokeratin 17, CK17, K17) expression as a robust prognostic biomarker for postoperative recurrence in stage II CRC.
Materials and Methods

High-throughput data analysis

All microarray data are publicly available from the Gene Expression Omnibus (GEO) database (http://www.ncbi.nlm.nih.gov/geo), and the normalized expression values were obtained from each dataset and were not processed further. This study utilized multiple microarray datasets of stage II CRC tumor samples determined on Affymetrix HG-U133 Plus 2.0 platform. We initially collected 4 datasets of stage II CRC, including GSE17536 (n=55), GSE39582 (n=260), GSE26906 (n=90) and GSE18088 (n=53), as Microarray Discovery set for genome-wide screening of genes that were differentially expressed between tumors with postoperative recurrence and those without recurrence. Those 4 datasets contained relapse information, and the former two datasets also had RFS data. To confirm the survival impact of genes found in the discovery steps, an independent stage II CRC dataset (GSE33113) with Relapse-free-survival (RFS) data was employed as Microarray Validation set 1 (n=89). We further utilized two additional datasets of stage II CRC, including GSE37892 (n=73) and GSE31595 (n=20). Because of their relatively small sample size, those two datasets were combined to increase the statistical power, designated as Microarray Validation set 2 (n=93).

For TCGA data analysis, level 3 Illumina RNA-seq data for both colon and rectal adenocarcinoma (COADREAD) with clinicopathological and RFS data were downloaded through cBioPortal (http://www.cbioportal.org/) [22, 23]. MSI status were further obtained as described previously [24]. We found two different versions of RNA-seq data normalized either by RPKM or RSEM methods, namely, RNA-Seq V1 and RNA-Seq V2, containing 60 and 123 stage II patients, respectively, after removing 3 redundant samples from the former dataset. Those two datasets were combined and designated as RNA-seq validation set (n=183).

For Kaplan-Meier survival analyses, stage II patients in each dataset were dichotomized into clearly-defined high and low groups on the basis of the median expression value for each genes within each datasets. We also used transcriptomic profiles of human cancer cell lines, including Affymetrix dataset GSE36133 from The Cancer Cell Line Encyclopedia (CCLE) and Illumina microarray dataset GSE59857 for 151 CRC cell lines.

Patient samples

We have previously described a large set of FFPE tissue samples obtained from 368 consecutive patients with stage 0-IV primary CRC treated in Fukushima Medical University (FMU) Hospital [24]. Among them, we used 110 tissues with available FFPE sections from stage II patients, who underwent curative resection (R0) between 2000 and 2009 without preoperative chemotherapy
or radiotherapy, designated as IHC validation set 1. For additional validation, we further enrolled 141 consecutive patients with stage I-IV CRC, who underwent surgery between 2010 and 2013, and then 44 stage II patients were utilized as IHC validation set 2. All tumors were classified according to the TNM classification of malignant tumors [25]. Clinical and pathological information, as well as high-risk characteristics for stage II disease were retrospectively obtained by review of medical records, with the last follow-up in December 2017. The primary endpoint of interest was RFS, which was defined as time from the date of surgery to the date of first relapse. The study was conducted in accordance with the Declaration of Helsinki and was approved by the Institutional Review Board of Fukushima Medical University.

**Immunohistochemistry (IHC)**

Suitable antibodies were identified using the Human Protein Atlas database, in which antibody-based comprehensive proteomic data are available (www.proteinatlas.org) [24, 26]. Primary mouse monoclonal anti-KRT17 antibody (M7046, Cytokeratin17 [E3], Dako, Glostrup, Denmark) was used for KRT17 staining. Four-μm thick sections were deparaffinized in xylene and rehydrated in a graded ethanol series. Endogenous peroxidases were blocked with 0.3% hydrogen peroxide in methanol. Antigens were retrieved by autoclave for 5 min in 10 mM citrate buffer solution (105°C, pH 6.0). Primary anti-KRT17 antibody was incubated in a 1:40 dilution of 10mM phosphate-buffered saline containing Tween 20 (Sigma–Aldrich) at 4°C overnight, and subsequently detected by a horseradish peroxidase (HRP)-coupled anti-mouse polymer (Envision+System-HRP, Dako, CA, USA) followed by incubation with diaminobenzidine (Dako). All sections were counterstained with Carrazzi’s hematoxylin. Normal mammary gland specimens were used as positive controls for KRT17 staining. We defined positive when at least a few tumor cells or tumor cell clusters was stained in a tumor specimen. Sections were evaluated independently by two investigators who had no prior knowledge of the clinical data.

**Determination of mismatch repair (MMR) status**

IHC for MMR proteins, including MLH1, MSH2, MSH6, and PMS2 was performed using Dako EnVision+ System with mouse or rabbit monoclonal antibodies against MLH1 (clone ES05, 1:50; Dako), MSH2 (clone FE11, 1:50; Dako), MSH6 (clone EP49, 1:200; Dako) and PMS2 (clone EP51, 1:50; Dako), as described previously [24]. Loss of MMR protein was defined as the absence of nuclear staining of tumor cells in the presence of positive nuclear staining in internal controls. Loss of at least one MMR protein was designated as deficient MMR (dMMR), and tumors with intact MMR protein expression as proficient MMR (pMMR).

**Cell culture, siRNA transfection, quantitative reverse-transcription (qRT)-PCR and western
Authenticated human CRC cell lines, including SW480, SW620, RKO, WiDr, SW837, LoVo, HCT116, SNU503, SNU81, and SNU407 were purchased from American Type Culture Collection (ATCC, Manassas, VA, USA), JCRB Cell Bank (Osaka, Japan), Korean Cell Line Bank (Seoul, Korea) or RIKEN Cell bank (Ibaraki, Japan). A human gastric cancer cell line, NCI-N87 was obtained from ATCC. LS180 was obtained as previously described and authenticated by STR analysis [24]. RKO, WiDr and LS180 cells were maintained with DMEM (ThermoFisher Scientific); others with RPMI-1640 (ThermoFisher Scientific) containing 10% fetal bovine serum (FBS) and penicillin/streptomycin (100 IU/ml) (ThermoFisher Scientific) at 37°C in a humidified atmosphere of 5% CO₂. For knockdown of KRT17, cells were plated and transfected with siRNA oligonucleotides for KRT17 or scramble control (Ambion® Silencer Select; s7992, s7993 and negative control #1) using Lipofectamine RNAiMAX Reagent (ThermoFisher Scientific), as described previously [24]. Total RNA extraction, reverse transcription and quantitative PCR were carried out as previously described [24, 27], with TaqMan assays, including KRT17 (Assay ID Hs00356958_m1) and ACTB (Hs99999903_m1) (ThermoFisher Scientific). Western blotting was performed as previously described [24]. Briefly, protein lysate was separated by 4-20% Tris-Glycine gel (ThermoFisher Scientific) and transferred onto PVDF membrane. Primary antibodies used were mouse monoclonal anti-KRT17 (M7046, Cytokeratin17 [E3], 1:600), mouse monoclonal anti-vinculin (7F9) (#SC-73614, 1:1000; Santa Cruz Biotechnology) or mouse monoclonal anti-β-actin (#SC-69879, 1:2000, Santa Cruz Biotechnology).

Statistical analysis

Student’s t-test, Mann–Whitney U test and Fisher’s exact test were used to determine differences in clinicopathological variables between two groups. Pearson’s correlation was used to evaluate the correlations between expression levels of two groups. Cumulative survival was estimated by the Kaplan–Meier method, and differences between two groups were analyzed by the log-rank test. Univariate and multivariate models were computed using Cox proportional hazards regression. All statistical analyses were two-sided and were conducted using Graphpad Prism v7.0 (Graphpad Software, Inc., La Jolla, CA, USA) or SPSS Statistics version 24 (IBM Corporation, NY, USA). P-values less than 0.05 were considered statistically significant.

Results

Genome-wide screening identified 6 differentially expressed genes associated with recurrence that were common in 4 cohorts of stage II CRC

The workflow of this study is illustrated in Figure 1A. We initially analyzed four
independent Affymetrix microarray cohorts of stage II patients as discovery datasets (Table 1), including GSE17536 (n=55), GSE39582 (n=260), GSE26906 (n=90) and GSE18088 (n=53). Each of these cohorts were observed more than five years. This allowed us to comprehensively search for relapse-related genes that were significantly differentially expressed between patients experienced recurrence after surgery and those who remained disease free (P<0.05). Seven probes, which corresponded to six genes, were found, as listed in Table 2 and showed in Figure1B. This indicated that in all four cohorts of stage II CRC in the discovery dataset, the expression of KRT17, HCFC1IR1, CERCAM, PACS1 and HTRA1 was increased and the expression of ORC5 was decreased significantly in patients with postoperative recurrence, compared to those without recurrence.

**Four genes were associated with RFS in the discovery set**

To evaluate the prognostic impact of the 6 candidate genes on RFS, the former two datasets, including GSE17536 and GSE39582 in the discovery set were combined (n=315) because of their availability of RFS information. Kaplan-Meier survival analyses were conducted, in which patients were dichotomized into high or low groups based on the median expression of each genes. As demonstrated in Figure 2A-D, high levels of KRT17, HCFC1IR1, CERCAM and low levels of ORC5 expression were each significantly associated with poor RFS (P<0.05). However, PACS1 or HTRA1 expression did not show statistically significant impact on RFS at least in this cohort (Figure 2E-F), thus these two genes were excluded from further analyses.

**KRT17 expression was associated with RFS in a microarray validation cohort**

Given that the 4 genes were chosen in the discovery set, we next sought to evaluate the reproducibility of their prognostic performance in an additional independent microarray cohort of stage II CRC. Same as above, Kaplan-Meier survival analyses were conducted, in which patients were dichotomized into high or low groups based on the median expression of each genes. As shown in Figure 3A, KRT17 was found to be the only gene whose expression was significantly associated RFS in Microarray Validation set 1 (P=0.0087, GSE33113, n=89). Although high expression of HCFC1IR1 and CERCAM demonstrated a clear trend towards poor RFS (Figure 3B-C), at least partly confirming their predictive power of RFS, they did not reach statistical significance. Whereas, ORC5 expression had no survival impact in this cohort (Figure 3D). Those analyses finally led to the identification of KRT17 expression as a promising candidate for prognostic biomarker in stage II CRC.

**Prognostic values of KRT17 expression was validated in an additional validation cohort**

To further validate the robustness of our findings, we tested whether the prognostic value of KRT17 expression in stage II CRC could be reproduced in additional independent cohorts. We used Microarray Validation set 2 (n=93) and RNA-seq validation set (n=183) obtained from TCGA,
demonstrating a significant association between high expression of \textit{KRT17} and worse RFS in both cohorts (P=0.0158 and P=0.0227, respectively, Figure 4A-B). Furthermore, as shown in Table 3, univariate and multivariate Cox analyses revealed that high expression \textit{KRT17} mRNA was significantly associated poor RFS, independent of other clinical factors in all 4 transcriptomic cohorts we analyzed, including Microarray Discovery set (hazard ratio [HR] 2.31; 95% confidence interval [CI] 1.38–3.89; P = 0.002), Microarray Validation set 1 (HR 3.99; 95%CI 1.29-12.28; P=0.016), Microarray Validation set 2 (HR 6.19; 95%CI 1.29-29.83; P=0.023), and RNA-seq Validation set (HR 2.51; 95%CI 1.09-5.78; P=0.030).

\textbf{The expression of KRT17 in protein levels}

We next sought to evaluate the expression of KRT17 protein levels. Primary mouse monoclonal anti-KRT17 antibody was identified using the Human Protein Atlas database [26]. We initially analyzed two microarray datasets of human cancer cell lines on different platforms (GSE36133 and GSE59857), consistently demonstrating multiple cell lines with various levels of \textit{KRT17} mRNA expression (Figure 5A). Subsequently, \textit{KRT17} mRNA and KRT17 protein expression levels in a panell of CRC cell lines were detected by qRT-PCR and western blot, respectively (Figure 5B). We also used a gastric cancer cell line, NCI-N87, as it was reported to express high levels of \textit{KRT17} [28]. We found significant and strong correlation between mRNA and protein expression (r$^2$=0.9041, P<0.0001, Figure 5C). Knock-down of \textit{KRT17} in CRC cells analyzed with western blot also supported the specificity of this antibody (Figure 5D). KRT17 protein expression was lower in both of two si-RNA transfection cancer cells.

To evaluate KRT17 protein expression on FFPE specimens by IHC, normal mammary gland tissues were used as a positive control, since it is known that KRT17 is expressed in breast ductal myoepithelial cells (Figure 6A) [29, 30]. We then evaluated the expression of KRT17 using IHC in 110 FFPE tumor samples (designated as IHC Validation set 1) obtained from stage II patients who underwent curative surgery between 2000 and 2009 in our hospital. Representative images are shown in Figure 6B-D. The cytoplasmic expression of KRT17 protein was found in cancer cells and cancer cell clusters, partly or sparsely in the tissue. Of 110 tumors, 19 (17.2%) were determined positive for KRT17 expression.

\textbf{Prognostic values of KRT17 protein expression in multiple cohorts}

To investigate whether the expression of KRT17 in protein levels could also identify stage II patients with poor RFS, survival analysis was conducted in IHC Validation set 1. Kaplan-Meier analysis showed that patients with KRT17-positive tumors had significantly shorter RFS after surgery, compared to those of KRT17-negative (P=0.0294, Figure 7A), which is highly consistent with the prognostic significance of KRT17 expression in mRNA levels. To further validate KRT17 staining as
a prognostic tool, an additional independent set of FFPE tissues were newly obtained from 44 stage II patients who underwent surgical resection between 2010 and 2013 (designated as IHC Validation set 2). As demonstrated in Figure 7B, KRT17-positive tumors were significantly associated with worse RFS than those of KRT17-negative in IHC Validation set 2 (P=0.0006).

**KRT17 as an independent prognostic factor in stage II CRC**

For Cox regression analyses, IHC Validation set 1 and 2 were combined to increase the statistical power. In this combined IHC cohort, no significant association was found between KRT17 expression and clinicopathological parameters, including age, gender, tumor location, histological differentiation, MMR status, or high-risk features (Table 4). Univariate Cox analysis showed that the expression of KRT17 protein significantly associated with RFS (HR 6.61; 95%CI 2.34-18.643; P<0.001). Moreover, the prognostic significance of high KRT17 protein expression on RFS was independent of conventional clinical factors and known high-risk stage II features by multivariate analysis (HR 10.01; 95%CI 2.74-36.56; P<0.001) (Table 5).

**Discussion**

In this study, we extensively collected 9 independent cohorts of stage II CRC consisting of 977 patients who underwent surgery. Since our aim was to develop prognostic biomarkers that can stratify stage II patients who are at high risk of relapse, all analysis in this study only used tumor samples from stage II patients. The strength of this study is that it is based on the analysis of large number of samples and cohorts from homogenous patient population, providing highly consistent results across multiple platforms, where the reproducibility of our findings was clearly demonstrated in the intended clinical context. Although genome-wide approaches might yield numerous false discoveries, our step-wise strategy of integrating currently available gene expression data based on microarray and RNA-seq from many laboratories in various countries successfully minimized the frequency of such false positive results.

Rather than attempt to generate new multigene prognostic signatures, this study focused particularly on the identification of single genes whose expression levels are predictive of disease recurrence, with robust reproducibility across multiple independent cohorts. Indeed, in all 7 datasets we analyzed, KRT17 mRNA expression was positively associated with postoperative recurrence. The identification and validation of the prognostic impact of KRT17 transcript was recapitulated by further validation at protein levels using two independent FFPE series with well-characterized information on high-risk stage II features. We consider that the use of FFPE tissues could extend the clinical utility of prognostic tools, as archived FFPE tumor specimens remain the most available and convenient tissue source for biomarker studies and for practical purpose. Indeed, IHC for KRT17 protein is simple that
relies on only a single molecule and can be analyzed on readily-available FFPE tissues. In view of that, KRT17 IHC may have potential to be used as a prognostic biomarker for the management of stage II CRC after surgery in the current clinical setting. KRT17 staining demonstrated significant prognostic impact on poor RFS, independent of previously-known clinicopathological features of high-risk stage II disease. Those findings suggest that KRT17 can add additional discriminative value to identify stage II patients at high risk for relapse, beyond conventional clinicopathological characteristics. Therefore, for patients with positive KRT17 expression, intensive postoperative intervention, including adjuvant chemotherapy with more appropriate postoperative surveillance, may be considered. Since combination of multiple biomarkers can provide more accurate risk stratification than single markers, KRT17 IHC combined with other genetic, epigenetic and clinicopathological modalities would further contribute to precision management for patients with stage II CRC after surgery.

Keratins belong to the superfamily of intermediate filament proteins primarily expressed in epithelial cells, which are encoded by 54 distinct keratin (KRT) genes [31]. The expression of keratins is tightly regulated in a tissue-specific and context-dependent manner, contributing to the maintenance of cellular integrity. Keratins typically retain their epithelial cell type-specific expression during tumorigenesis, and de novo expression also occurs in some types of tumor. Therefore, immunohistochemical detection of keratins has been widely established in clinical cancer pathology, particularly, with the routine use of keratin 7 (CK7) and keratin 20 (CK20) IHC for determining the tissue of origin in cancer of unknown primary [29, 31]. In breast cancer, keratin 19 (CK19) mRNA expression is currently used in identifying tumor cells within sentinel lymph nodes [31].

KRT17 (keratin 17, CK17, K17) is known to be expressed in basal and myoepithelial cells of various glands, including mammary gland, respiratory epithelium, and urothelium [29, 30]. Although KRT17 is not expressed in healthy skin, it is rapidly inducible in skin epithelial cells under stressful condition, such as skin injury, resulting in cell growth [32] and inflammatory reaction [33]. KRT17 leads positive regulation of epithelial cell size and growth by binding to the adaptor protein 14-3-3σ by stimulating Akt/mTOR pathway at the wound. In tumor tissue, likewise, several previous studies reported KRT17 affect the function of protein synthesis and cell growth and is considered to be great importance of cell viability and function through the same way [32] . KRT17 expression is also found in tumor cells, and a number of studies have consistently demonstrated that KRT17 accelerates oncogenic transformation and tumor progression in vitro and in vivo in many types of cancer [28, 33-39]. In gastric cancer, knockdown of KRT17 by RNA interference was able to significantly suppress cell proliferation, migration and invasion in vitro [28] . In addition, silencing of KRT17 induces apoptosis and cell-cycle arrest at G1/S phase of gastric cancer cell. They say that is why high KRT17 expression leads poor prognosis. Moreover, in cervical cancer, KRT17 has been identified and used as a marker of cancer
stem cells [40-42]. It is worth noting that the expression of KRT17 is associated with adverse prognosis in several cancer types, including cervical cancer [34, 43], gastric cancer [38, 44], and breast cancer [45-47]. Collectively, all of those studies clearly implicate the oncogenic function of KRT17 in human cancer, correlated with poor prognostic phenotypes. Although the present study did not address the functional role of KRT17, it is therefore conceivable that KRT17 also plays a tumor-promoting role in CRC.

Our study had several limitations, including its exploratory and retrospective nature. One limitation for the analysis of public cohorts is that there was incomplete or unavailable information on high-risk clinicopathological features for stage II patients. Thus, we were not able to include those covariates in the models. In addition, the predictive value of KRT17 expression for the response to adjuvant treatment remains largely unknown, despite its robust prognostic significance on RFS. This is mostly due to the lack of information on the administration of adjuvant chemotherapy in the majority of the transcriptomic cohorts used in this study. Since stage II patients are usually treated by surgery alone in the current practice, we were not able to directly address the predictive role of KRT17 IHC, owing to relatively small number of patients who received postoperative chemotherapy even in our FFPE cohorts.

In conclusion, with the use of 9 independent cohorts consisting of nearly 1000 patients with stage II CRC who underwent surgery, we extensively conducted a genome-wide screening of single genes, combined with step-wise validation strategies based on multiple platforms. We identified and validated the expression of KRT17 transcript or KRT17 protein as a prognostic biomarker for RFS with high reproducibility. To our knowledge, this is the first study reporting the prognostic role of KRT17 expression in CRC, especially in postoperative stage II patients who are at high probability of disease recurrence. Our study suggests that KRT17 IHC provides additional prognostic stratification beyond the currently-available high-risk factors, highlighting the potential of its performance in precision management for stage II patients in the clinical setting.

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Figure Legends

**Figure 1.** (A) Workflow for the stepwise identification and validation of prognostic genes in patients with stage II colorectal cancer. (Step 1) Genome-wide screening for genes
differentially expressed between patients experienced recurrence and those who remained disease free using four microarray cohorts, followed by confirmation in relapse-free survival (RFS) analysis for candidate genes. (Step 2) Validation of candidate genes by RFS analysis in three independent microarray and RNA-seq cohorts. (Step 3) Validation of candidate genes at protein levels by RFS analysis in two independent immunohistochemistry cohorts. (B) Venn diagram of 4 cohorts in step 1. Many probes were detected in each cohort and only seven probes were commonly identified in all 4 cohorts.

**Figure 2.** Kaplan-Meier curves depicting relapse-free survival for patients with stage II colorectal cancer in the microarray discovery cohorts (n=315). Patients were dichotomized as high or low based on the median expression of candidate genes, including KRT17 (A), HCFC1R1 (B), CERCAM (C), ORC5 (D), PACS1 (E), and HTRA1 (F).

**Figure 3.** Kaplan-Meier curves showing relapse-free survival for patients with stage II colorectal cancer in Microarray validation set 1 (n=89). Patients were dichotomized as high or low based on the median expression of candidate genes, including KRT17 (A), HCFC1R1 (B), CERCAM (C), and ORC5 (D).

**Figure 4.** Kaplan-Meier curves showing relapse-free survival for KRT17 expression in additional independent cohorts of stage II colorectal cancer, including Microarray validation set 2 (n=93) (A) and RNA-seq validation set (n=183) (B). In each cohort, patients were dichotomized based on the median expression of KRT17.

**Figure 5.** KRT17 mRNA and KRT17 protein expression in human cancer cell lines. (A) Two microarray datasets of cell lines derived from different types of cancer (GSE36133, left panel) and from colorectal cancer (GSE59857, right panel), showing various expression levels of KRT17 transcript. (B) Western blotting (upper panel) and qRT-PCR (lower panel) analysis for KRT17 protein and mRNA levels, respectively, in a gastric cancer cell line NCI-N87, and a panel of 11 colorectal cancer cell lines. (C) Correlation between the expression of KRT17 mRNA by qRT-PCR and KRT17 protein by western blotting in cancer cell lines. (D) Knockdown of KRT17 in two colorectal cancer cell lines, confirmed by western blotting.

**Figure 6.** Immunohistochemistry for KRT17 protein expression. (A) KRT17 expressed by breast ductal myoepithelial cells in normal mammary gland. (B) Non-tumor colon mucosa showing negative for KRT17 staining. (C,D) Colorectal cancer tissues showing positive-KRT17 expression in the cytoplasm of cancer cells (C) and negative-KRT17 expression (D).
**Figure 7.** Kaplan-Meier curves displaying relapse-free survival for KRT17 protein expression by immunohistochemistry in two independent cohorts of stage II colorectal cancer, including IHC validation set 1 (n=110) (A) and IHC validation set 2 (n=44) (B).
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Table 3. Univariate and Multivariate Cox regression of KRT17 mRNA expression for relapse-free survival in microarray or RNA-seq cohorts of stage II colorectal cancer

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Univariate and multivariate analysis were each adjusted for cohort membership.
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Table 5. Univariate and multivariate Cox regression of KRT17 protein expression for relapse-free survival in stage II colorectal cancer by immunohistochemistry (n=154)

<table>
<thead>
<tr>
<th>Variable</th>
<th>Univariate</th>
<th>Multivariate</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>HR (95% CI)</td>
<td>P</td>
</tr>
<tr>
<td>KRT17 Positive vs Negative</td>
<td>6.60 (2.34 - 18.64)</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>Age Continuous</td>
<td>1.04 (0.99 - 1.09)</td>
<td>0.159</td>
</tr>
<tr>
<td>Gender Female vs Male</td>
<td>1.94 (0.72 - 5.24)</td>
<td>0.194</td>
</tr>
<tr>
<td>Location Left vs Right</td>
<td>2.15 (0.67 - 6.89)</td>
<td>0.200</td>
</tr>
<tr>
<td>Histological differentiation Poor-Mucinous vs Well-Moderate</td>
<td>NA</td>
<td></td>
</tr>
<tr>
<td>Tumor invasion pT4 vs pT3</td>
<td>3.42 (1.26 - 9.16)</td>
<td><strong>0.014</strong></td>
</tr>
<tr>
<td>Lymphatic invasion Positive vs Negative</td>
<td>1.06 (0.33 - 3.47)</td>
<td>0.397</td>
</tr>
<tr>
<td>Vascular invasion Positive vs Negative</td>
<td>1.36 (0.29 - 6.47)</td>
<td>0.698</td>
</tr>
<tr>
<td>Lymph nodes examined &lt;12 vs ≥12</td>
<td>2.90 (1.07 - 7.90)</td>
<td><strong>0.037</strong></td>
</tr>
<tr>
<td>Perforation or Obstruction Yes vs No</td>
<td>2.41 (0.69 - 8.39)</td>
<td>0.166</td>
</tr>
<tr>
<td>Adjuvant chemotherapy Received vs Not received</td>
<td>0.93 (0.58 - 1.51)</td>
<td>0.774</td>
</tr>
<tr>
<td>MMR status dMMR vs pMMR</td>
<td>NA</td>
<td>NA</td>
</tr>
</tbody>
</table>

*Univariate and multivariate analysis were each adjusted for cohort membership.*
**Figure 1A**

**Step 1. Microarray Discovery**
- Cohort 1: GSE17536 (Stage II, n=55)
- Cohort 2: GSE39582 (Stage II, n=260)
- Cohort 3: GSE26906 (Stage II, n=90)
- Cohort 4: GSE18088 (Stage II, n=53)

GSE17536 + GSE39582 (Stage II, n=315)

Comprehensive screening of genes associated with postoperative recurrence

**Step 2. Microarray / RNA-seq Validation**
- Cohort 5: Microarray Validation set 1 GSE33113 (Stage II, n=89)
- Cohort 6: Microarray Validation set 2 GSE37892+GSE31595 (Stage II, n=93)
- Cohort 7: RNA-seq Validation set TCGA RNA-seq V1/V2 (Stage II, n=183)

Independent validation at transcript levels

**Step 3. Immunohistochemistry Validation**
- Cohort 8: IHC Validation set 1 FMU 2000-2009 (Stage II, n=110)
- Cohort 9: IHC Validation set 2 FMU 2010-2013 (Stage II, n=44)

Independent validation at protein levels
Figure 1B

Cohort 1
GSE17538
(n=55)

Cohort 2
GSE39582
(n=260)

Cohort 3
GSE26906
(n=90)

Cohort 4
GSE18088
(n=53)

3552
66
25
167
1978

518
7
32
169

3030
105
9
312

349
4002

169
Figure 2
Figure 3

Microarray validation set 1
(stage II, n=89)

A

\[ P = 0.0087 \]

- KRT17 Low (n=45)
- KRT17 High (n=44)

B

\[ P = 0.1231 \]

- HCFO1R1 Low (n=45)
- HCFO1R1 High (n=44)

C

\[ P = 0.0644 \]

- CERCAM Low (n=45)
- CERCAM High (n=44)

D

\[ P = 0.6916 \]

- ORCS Low (n=45)
- ORCS High (n=44)
Figure 4

A. Microarray validation set 2
   Stage II (n=93)
   P = 0.0158
   
   - KRT17 Low (n=47)
   - KRT17 High (n=46)

B. RNA-seq validation set
   Stage II (n=183)
   P = 0.0227
   
   - KRT17 Low (n=92)
   - KRT17 High (n=91)
Figure 5

A

B

C

D

GOLE (GSE38133) vs. GSE59857

KRT17 mRNA expression

KRT17 protein expression

vinculin

KRT17 mRNA expression

β-actin

r^2 = 0.9041
p < 0.0001

SNU07

SNU07

SNU03

SNU03

SNU03

SNU03

SNU03

SNU03

SNU03

SNU03
Figure 7

A  IHC validation set 1
Stage II (n=110)

\[ p = 0.0294 \]

- KRT17 negative (n=91)
- KRT17 positive (n=19)

B  IHC validation set 2
Stage II (n=44)

\[ p = 0.0006 \]

- KRT17 negative (n=20)
- KRT17 positive (n=15)