Molecular profiling of appendiceal adenocarcinoma and comparison with right-sided and left-sided colorectal cancer

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Figures/Tables: 2 figures and 3 tables, and 1 supplementary figure and 3 supplementary tables.
References: 49

Use of standardized official symbols:
In this study, HUGO (Human Genome Organisation)-approved official symbols for genes and gene products were used: all of which are described at www.genenames.org.

Abbreviations: AA, appendiceal adenocarcinoma; CRC, colorectal cancer; CRS, cytoreductive surgery; FFPE, formalin fixed paraffin-embedded; HIPEC, hyperthermic intraperitoneal chemotherapy; IHC, immunohistochemistry; IPMN, intraductal papillary mucinous neoplasm; L-CRC, left-sided colorectal cancer; MSI, microsatellite instability; MSI-H, MSI-high; MU, mucinous adenocarcinoma; NGS, next-generation sequencing; NOS, adenocarcinoma, not otherwise specified; PD-L1, programmed death-ligand 1; PMP, pseudomyxoma peritonei; R-CRC, right-sided colorectal cancer; SR, signet ring cell carcinoma; TMB, tumor mutational burden; TMB-H, TMB-high.
**Translational relevance:**

Compared to colorectal cancer (CRC), appendiceal adenocarcinoma (AA) is more associated with peritoneal dissemination and increased mucin production. However, AA patients typically receive therapies approved for CRC in spite of lower vulnerability. The molecular profiling of AA and to compare it with those of right-sided CRC (R-CRC) and left-sided CRC (L-CRC) is needed for personalized treatment strategy. We found that AA had higher mutation rates in *GNAS* and *KRAS*, and lower mutation rates in *TP53*, *APC*, and *PIK3CA* than CRC, and gene mutation rates differed between the histopathological subtypes. Although the immune profile results (microsatellite instability, tumor mutational burden, and PD-L1 expression status) of AA were similar to those of L-CRC (not to those of R-CRC), not otherwise specified AA (NOS) had higher overexpression of the markers than did other subtypes of AA and L-CRC. This molecular profiling may support developing a personalized treatment strategy in patients with AA.
Abstract

Purpose: The natural history and prognosis of appendiceal adenocarcinomas (AA) differ from those of adenocarcinomas arising in other large bowel sites. We aimed to compare the molecular profiles exhibited by AAs and CRCs, or between the histopathological subtypes of AA.

Experimental Design: A total of 183 samples from AA (46 adenocarcinoma, not otherwise specified (NOS), 66 pseudomyxoma peritonei (PMP), 44 mucinous adenocarcinoma (MU), and 27 signet ring cell carcinoma (SR)), 994 from right-sided colorectal cancer (R-CRC), and 1080 from left-sided CRC (L-CRC) were analyzed by next-generation sequencing (NGS) and immunohistochemical (IHC) markers. Microsatellite instability (MSI) and tumor mutational burden (TMB) were tested by NGS, and PD-L1 by IHC.

Results: We observed high mutation rates in AA samples for KRAS (55%), TP53 (40%), GNAS (31%), SMAD4 (16%), and APC (10%). AA exhibited higher mutation rates in KRAS and GNAS, and lower mutation rates in TP53, APC, and PIK3CA (6%) than CRCs. PMP exhibited much higher mutation rates in KRAS (74%) and GNAS (63%), and much lower mutation rates in TP53 (23%), APC (2%), and PIK3CA (2%) than NOS. Alterations associated with immune checkpoint inhibitor response (MSI-high, TMB-high, PD-L1 expression) showed similar frequency in AA compared to L-CRC, but not R-CRC, and those of NOS were higher than other subtypes of AA and L-CRC.

Conclusion: Molecular profiling of AA revealed different molecular characteristics than noted in R-CRC and L-CRC, and molecular heterogeneity between the histopathological subtypes of AA. Our findings may be critical to develop individualized approach for AA treatment.
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Introduction

The natural history and prognosis of appendiceal adenocarcinomas (AA) differs from those of adenocarcinomas arising in other large bowel sites (1, 2). Compared to colorectal adenocarcinoma, AA more commonly are associated with peritoneal dissemination and increased mucin production, and are distinguished by the diagnostic classification as pseudomyxoma peritonei (PMP) (3, 4). Histological variants of appendiceal epithelial neoplasia include low- and high-grade mucinous neoplasms, goblet cell tumors, neuroendocrine neoplasms, adenoma, and adenocarcinomas (common colonic type, mucinous type, and signet ring cell carcinoma) (2, 5). The primary treatment for these neoplasms is surgical resection (6). Also in PMP, cytoreductive surgery (CRS) and hyperthermic intraperitoneal chemotherapy (HIPEC) are associated with favorable outcome when conducted in specialized centers of excellence (6-8). However, both patient selection and the expertise of the treating team are critical to best outcomes. Furthermore, in patients with metastatic AA or recurrent PMP, the most effective chemotherapy and molecular targeted therapy is controversial. At present, AA patients typically receive therapies approved for colorectal cancer (CRC) (3, 9, 10), although their efficacy, especially in low grade tumors, suggests lower vulnerability of these slow growing tumors to conventional therapies as compared to CRCs (11, 12).

Molecular profiling has been used effectively to identify novel treatment options for malignant diseases. In CRC, genetic profiling, such as RAS and BRAF alterations, has suggested phenotypic clustering with profiles that are prognostic as well as predictive of different susceptibilities to molecularly targeted therapeutics (13). In addition, assessment of the presence or absence of microsatellite instability (MSI), tumor mutational burden (TMB), and programmed death-ligand 1 (PD-L1) expression can be predictive of the likelihood that an individual patient
will see a tumor response to treatment with immune checkpoint inhibitors (14). With molecular profiling, differences in the genetic and immune characteristics between the right-sided (R-CRC) and left-sided CRC (L-CRC) have been reported and have developed a personalized treatment strategy in CRC (15). On the contrary, studies on molecular profiling of AA have been handicapped due to the rarity of AA and assay failures in available PMP samples because of low cellularity (2), and these analyses have provided limited genetic data (16, 17). Although the prognosis varies according to the histopathological subtypes of AA (18), studies are lacking that correlate molecular profiles with AA subtypes (19). Identification of molecular alterations of AA is critical for the development and selection of more effective therapeutic strategies.

We performed molecular profiling of AA and compared it with those of R-CRC and L-CRC, using integrated data within a total of 183 samples from AA, 994 from R-CRC, and 1,080 from L-CRC. Our analysis demonstrates that histopathological and molecular classification of AA could be a key step towards personalized treatment strategies of AA.

Materials and Methods

Tumor Samples

Figure 1 summarizes the workflow of this study. Consecutive appendiceal cancer (N = 224) and CRC (N = 4,600) cases submitted to a commercial CLIA-certified laboratory (Caris Life Sciences, Phoenix, AZ) from April, 2015 to January, 2018 were retrospectively analyzed for their molecular alterations. Formalin fixed paraffin-embedded (FFPE) samples were sent for analysis from treating physicians around the world. The tissue diagnoses were submitted based on pathologic assessment of physicians who requested the assays and were further verified by a board-certified oncological pathologist at the Caris laboratory. A total of 183 AAs were analyzed.
and 41 tumors of neuroendocrine/goblet histology were excluded from their analysis. Included in the AA cohort were 66 PMP, 44 mucinous adenocarcinoma (MU), and 27 signet ring cell carcinoma (SR). Forty-six tumors were determined to be adenocarcinoma, not otherwise specified (NOS) as no additional detailed histological features were noted (Supplementary Figure S1). R-CRC defined as tumors arising from the cecum to the hepatic flexure and transverse colon (N = 994) and L-CRC defined as those arising from the splenic flexure to the rectosigmoidal colon (N = 1,080) were analyzed; while tumors with origin unclearly annotated (N = 2,526) were excluded. Samples taken from original tumor sites were considered primary tumors and samples taken from organs other than the primary were considered metastases. Tissues were profiled by next-generation sequencing (NGS) and immunohistochemical (IHC) analysis using Caris Molecular Intelligence. Human subjects were anonymized prior to analysis. This study was conducted in accordance with guidelines of the Declaration of Helsinki, Belmont report and U.S. Common rule. In keeping with 45 CFR 46.101(b)(4), this study was performed utilizing retrospective, de-identified clinical data. Therefore this study is considered IRB exempt and no patient consent was necessary from the subject.

**Next-generation sequencing (NGS)**

NGS was performed on genomic DNA isolated from FFPE tumor samples using the NextSeq platform (Illumina, Inc., San Diego, CA). A custom-designed SureSelect XT assay was used to enrich 592 whole-gene targets (Agilent Technologies, Santa Clara, CA). All variants were detected with > 99% confidence based on allele frequency and amplicon coverage, with an average sequencing depth of coverage of 750 and an analytic sensitivity of 5%. Prior to molecular testing, tumor enrichment was achieved by harvesting targeted tissue using manual
microdissection techniques. Genetic variants identified were interpreted by board-certified molecular geneticists and categorized as ‘pathogenic,’ ‘presumed pathogenic,’ ‘variant of unknown significance,’ ‘presumed benign,’ or ‘benign,’ according to the American College of Medical Genetics and Genomics (ACMG) standards. When assessing mutation frequencies of individual genes, ‘pathogenic,’ and ‘presumed pathogenic’ were counted as mutations while ‘benign’, ‘presumed benign’ variants and ‘variants of unknown significance’ were excluded.

**Microsatellite instability (MSI)**

MSI was examined using over 7,000 target microsatellite loci and compared to the reference genome hg19 from the University of California, Santa Cruz (UCSC) Genome Browser database, and the status was defined as MSI-high (MSI-H) or MSI-low/ microsatellite stable (MSS). The number of microsatellite loci that were altered by somatic insertion or deletion were counted for each sample. Only insertions or deletions that increased or decreased the number of repeats were considered. Genomic variants in the microsatellite loci were detected using the same depth and frequency criteria as used for mutation detection. MSI-NGS results were compared with results from over 2,000 matching clinical cases analyzed with traditional PCR-based methods. The threshold to determine MSI by NGS was determined to be 46 or more loci with insertions or deletions to generate a sensitivity of >95% and specificity of >90%.

**Tumor mutation burden (TMB)**

TMB was measured by counting all non-synonymous missense mutations found per tumor that had not been previously described as germline alterations (592 genes and 1.4 megabases [MB] sequenced per tumor). The threshold to define TMB-high (TMB-H) was
greater than or equal to 17 mutations/MB and was established by comparing TMB with MSI by fragment analysis in CRC cases, based on reports of TMB having high concordance with MSI-H in CRC.

**Immunohistochemical (IHC) analysis**

IHC analysis was performed on full slides of FFPE tumor specimens using automated staining techniques (Benchmark XT, Ventana, and Autostainer Link 48, Dako). The primary antibody and details of evaluation for analysis are shown in **Supplementary Table S1**. Staining was scored for intensity (0 = no staining; 1+ = weak staining; 2+ = moderate staining; 3+ = strong staining) and staining percentage (0-100%). Results were categorized as positive or negative by defined thresholds specific to each marker, based on published clinical literature.

**Statistical analysis**

All statistical analyses were performed with SPSS v23 (IBM SPSS Statistics, Cary, USA), and all tests were two-sided at a significant level of 0.05. The comparison of age was analyzed using Student’s t-test and that of molecular profile between groups were analyzed using Fisher’s exact test. Cases with missing information in any of the categorical data were not included in the analysis.

**Results**

**Patient and tumor characteristics**

Baseline patient and tumor characteristics in regard to age, gender, and location of tumor sampling are shown in **Supplementary Table S2**. Patients with AA were significantly
younger than patients with R-CRC ($P < 0.001$) and were more likely to be female than patients with L-CRC ($P = 0.011$). In addition, more AA samples were collected from metastatic sites than R-CRC ($P < 0.001$) and L-CRC ($P < 0.001$) samples. Between histopathological subtypes of AA, there is no significant difference in regard to age, gender, and location of tumor sampling.

**Common gene mutations in AA, and comparison with R-CRC and L-CRC**

The observed patterns of common gene mutations were totally different between AA, R-CRC, and L-CRC. (Table 1) The most prevalent mutations seen in AA were $KRAS$ (55%), $TP53$ (40%), $GNAS$ (31%), $SMAD4$ (16%), $APC$ (10%), $ARID1A$ (8%), $RNF43$ (7%), $PIK3CA$ (6%) and $BRAF$ (5%). Compared to both R- and L-CRCs, AA had significantly higher mutation rates in $GNAS$ (31% vs 2% vs 1%) and $SMAD4$ (16% vs 11% vs 10%), and lower mutation rates in $TP53$ (40% vs 66% vs 75%), $APC$ (10% vs 70% vs 83%), $PIK3CA$ (6% vs 22% vs 17%), $FBXW7$ (3% vs 11% vs 9%), $NRAS$ (1% vs 3% vs 5%), and $AMER1$ (0% vs 9% vs 2%). In addition, compared to R-CRC, AA had significantly lower mutation rates in $ARID1A$ (8% vs 26%), $BRAF$ (5% vs 17%), $ATM$ (2% vs 7%), $KMT2D$ (2% vs 7%), $PTEN$ (1% vs 8%), $MSH6$ (1% vs 5%), $HNF1A$ (1% vs 5%), $PITCH1$ (1% vs 5%), and $CTNNB1$ (0% vs 4%); and compared to L-CRC, AA had significantly higher mutation rates in $KRAS$ (55% vs 43%) and $RNF43$ (7% vs 2%). Moreover, the mutation rates in $TP53$, $GNAS$, $APC$, $PIK3CA$, and $AMER1$ had significant differences between AA, R-CRC, and L-CRC. $BRAF$ mutation rate was the highest in R-CRC (AA, R-CRC, L-CRC: 5%, 17%, 5%), and $KRAS$ mutation rate was the lowest in L-CRC (AA, R-CRC, L-CRC: 55%, 56%, 43%).

**Common gene mutations in the histopathological subtypes of AA**
We further evaluated the pattern of common gene mutations in the histopathological subtypes of AA (NOS, PMP, MU, and SR). (Table 2) Compared to NOS, PMP exhibited much higher mutation rates in \( KRAS \) (74% vs 44%) and \( GNAS \) (63% vs 7%), and much lower mutation rates in \( TP53 \) (23% vs 51%), \( APC \) (2% vs 22%), and \( PIK3CA \) (2% vs 15%); MU exhibited higher mutation rate in \( KRAS \) (64% vs 44%) and \( GNAS \) (25% vs 7%); and SR exhibited lower mutation rate in \( KRAS \) (15% vs 44%), \( GNAS \) (4% vs 7%), \( TP53 \) (33% vs 51%), \( APC \) (0% vs 22%), and \( PIK3CA \) (0% vs 15%). Notably, \( BRAF \) mutations were identified in NOS (7%), MU (9%), and SR (7%), but not identified in PMP.

**Immune profiling of AA, comparison with R-CRC and L-CRC, and differences between the histopathological subtypes of AA**

The immune profile of AA was similar to that of L-CRC but not that of R-CRC. (Figure 2) MSI-H, TMB-H, and PD-L1 high expression rates were much lower in AA (MSI-H, TMB-H, PD-L1 high: 2.2%, 2.2%, 2.8%) than in R-CRC (MSI-H, TMB-H, PD-L1 high: 14.5%, 14.9%, 6.7%), but similar to L-CRC (MSI-H, TMB-H, PD-L1 high: 3.4%, 4.6%, 2.7%). The positive ratio of any of immune check point inhibitor markers (MSI-H, TMB-H, PD-L1 high) is much higher in R-CRC (19.8%) than in AA (5.1%) and in L-CRC (6.7%). Notably, some differences were observed between the histopathological subtypes of AA: PMP had no MSI-H or TMB-H case; and the positive ratio of any one marker (MSI-H, TMB-H, PD-L1 high) was higher in NOS (11.4%) than in PMP (1.6%), in MU (4.8%), and in SR (3.9%).

**Protein expression of chemotherapeutic sensitivity markers in AA, comparison with R-CRC and L-CRC, and differences between the histopathological subtypes of AA**
As shown in Table 3, expression status of protein markers for chemotherapeutic sensitivity varied between AA, R-CRC, and L-CRC, and between the histopathological subtypes of AA. The positive ratios of ERCC1, TOPO1, PTEN, and MGMT were higher in AA than in R-CRC and L-CRC. Further, TS was overexpressed most frequently in R-CRC. Between the histopathological subtypes of AA, although the protein expressions were similar in NOS, MU and SR, those expressions were significantly different between NOS and PMP: ERCC1, TOPO1, PTEN, and MGMT were overexpressed in PMP, and TS was suppressed in PMP.

Comparison of molecular characteristics between the locations of tumor sampling

Finally, we analyzed the association of molecular characteristics with the location of tumor sampling (primary or metastatic sites). (Supplementary Table S3) In AA, the molecular characteristics had no difference between the locations of tumor sampling. On the other hand, in both R-CRC and L-CRC, the positive ratios of ARID1A mutation and MSI-H were significantly lower in metastatic sites.

Discussion

To the best of our knowledge, we performed so far the largest study to determine molecular profiling of AA and to compare it with R-CRC and L-CRC, using integrated data within a total of 183 samples from AA, 994 from R-CRC, and 1,080 from L-CRC. We found that AA had higher mutation rates in GNAS and KRAS, and lower mutation rates in TP53, APC, and PIK3CA than R-CRC and L-CRC; gene mutation rates differed between the histopathological subtypes of AA; and although the immune profile results (MSI, TMB, and PD-L1 expression status) of AA were similar to those of L-CRC (not to those of R-CRC), those of NOS had higher
likelihood of overexpression than did other subtypes of AA and L-CRC. Our results support developing a personalized treatment strategy in patients with AA that is tailored to the individual’s histopathological subtypes.

The molecular feature of AA is different from CRC. As in the past reports (16, 20-22), our current study showed that AA had higher GNAS and KRAS mutation rates in comparison with CRC. The mutations of GNAS, a member of the G-protein family, cause constitutive activation of the protein kinase A (PKA) pathway via elevated levels of cyclic AMP (cAMP), and affected the MAPK and Wnt signaling pathways (20). GNAS mutation is common in benign tumors such as villous adenomas of the stomach (23) and colon (24), and intraductal papillary mucinous neoplasm (IPMN) of the pancreas (25, 26) and bile duct (27). Although the cAMP-PKA pathway can stimulate cell proliferation via MAPK or Wnt signalings, exogenous GNAS mutation in CRC (28) and pancreatic ductal adenocarcinoma (29) did not promote cell proliferation but increased expressions of MUC2 and MUC5AC (which induce mucin production). Additionally, Taki et al. demonstrated that transgenic mice with pancreas-specific GNAS and KRAS mutations developed a cystic pancreatic tumor (30). However, intraperitoneal injection of CRC cells with exogenous GNAS mutation in mice did not produce PMP but resulted in the formation of solid tumors (28). These data suggest that both GNAS and KRAS may contribute to the oncogenesis of PMP. Further, the mutation rates in GNAS and KRAS increased from NOS to MU and to PMP in our data, implying a functional interaction of these two oncogenes on mucin production. Moreover, several studies showed that GNAS and KRAS mutations were independent from pathological grade which is related to PMP activity (31-35). In support of these findings, both GNAS and KRAS mutations were not reported to be prognostic in patients with PMP (32, 36). GNAS and KRAS mutations might be a genetic feature of PMP but not be easy therapeutic targets.
Our data for the first time demonstrated that all the subtypes of AA had much lower TP53, APC, and PIK3CA mutation rates in comparison with CRC, and the mutation rates were further much lower in PMP and SR than in NOS and MU. In CRC, TP53, APC, and PIK3CA are key driver genes for “adenoma-to-carcinoma sequence”, and the mutation rates are high in any stage or tumor status (37, 38). Our findings suggest that carcinogenic mechanisms of AA may be different from that of CRC. Wilson et al. showed that GNAS mutation cooperated with inactivation of APC leading to colorectal tumorigenesis, but not carcinogenesis (39). In addition, Noguchi et al. demonstrated that TP53, PIK3CA, and AKT1 mutations were detected in peritoneal mucinous adenocarcinoma but not in PMP (21). Although GNAS mutation might contribute to APC-driven tumorigenesis, mutations in TP53 and/or genes related to the PI3K-AKT pathway may be necessary for malignant transformation in PMP. On the other hand, SR exhibited lower mutation rates not only in TP53, APC, and PIK3CA, but also in KRAS and GNAS. In addition, the mutation rate of BRAF was much lower (7%) compared to SR of CRC (about 40%) (40, 41), suggesting that SR might differ from both other histopathological subtypes of AA and SR of CRC in both development and treatment strategies.

Recently, immune checkpoint based therapy has demonstrated better survival and tolerance in subsets of patients with both solid and hematological malignancies. Studies further showed benefit in immune checkpoint blockade in patients whose tumors with MSI-H, TMB-H, and PD-L1 high expression (14). There is a need for molecular markers which can identify patients who are likely to benefit from immune checkpoint inhibitors. Therefore, overall immune profiling is an important emerging biomarker for cancer treatment. The appendix originated from the cecum, has many lymphoid clusters, and regulates IgA production in the large bowel sites (42), suggesting that AA may be subject to lymphocytic regulation more than R-CRC and
L-CRC. However, interestingly, our study showed that immune profile of AA was similar to L-CRC but not R-CRC. In addition, NOS had a higher positive likelihood of expressing any of the evaluated immune check point inhibitor markers (11.4%) than other subtypes of AA (PMP: 1.6%, MU: 4.8%, SR: 3.9%) and L-CRC (6.7%). Although the carcinogenic mechanism of NOS is different from that of R-CRC and L-CRC, the immune characteristics might be closer to R-CRC. Further experimental studies are necessary to better understand this mechanism.

The efficacy of systematic chemotherapy and molecular targeted therapy for AA has not been well studied so far. Although a phase II trial in unresectable PMP suggested capecitabine combined with mitomycin C, the combination of fluorouracil (5-FU) and oxaliplatin/irinotecan is commonly used for the treatment of metastatic AA and PMP despite the paucity of efficacy data (3, 43). Our data showed that the positive ratios of ERCC1 and TOPO1 were higher in AA, especially in PMP, than in R-CRC and L-CRC, and that of TS was lower in AA, especially in PMP. ERCC1, a protein of the nucleotide excision repair (NER) complex, is essential for repairing platinum-DNA adducts and is involved in drug resistance to oxaliplatin (44); TOPO1, a molecular target of SN38, is a plausible positive predictive marker for irinotecan (45); and TS, a rate-limiting enzyme in the synthesis of pyrimidine nucleotides, is required for DNA synthesis and the activity is a negative predictive marker for 5-FU (44, 46). Thus, compared to CRCs, combination therapy with 5-FU and irinotecan (due to lower TS and higher ERCC1 and TOPO1) may be more effective for AA, especially for PMP treatment. However, the predictive protein expressions for chemosensitivities are still controversial even in CRC. As increased mucin production is responsible for major complications and fatal outcome in patients with PMP, GNAS-related pathways (cAMP-PKA, MAPK, and Wnt signaling pathways) might be a potential therapeutic target: PKA inhibitor (28), BIM-46174 (inhibitor of heterotrimeric
G-protein complex) (47), and a MEK-inhibitor (48) were reported to cause mucin production reduction in tumors with GNAS mutation. Biomarker studies according to the histopathological and molecular subtypes are needed to determine individualized treatment strategies for AA.

We also compared the molecular profiles of samples from primary and metastatic tumors to identify features that are associated with distant metastasis in AA. In consensus with the data from past reports (38), a high level of genomic concordance was detected. However, ARID1A mutation and MSI-H were specifically enriched in primary tumors compared with distant metastases in both R-CRC and L-CRC, suggesting their potential protective effects. Recently, ARID1A, a subunit of the chromatin remodeling complex SWI/SNF, mutation was reported to contribute to impaired mismatch repair and mutator phenotype in cancers (49). ARID1A could be a promising target for novel treatment strategies for immune checkpoint based therapy.

Limitations in our study need to be mentioned. First, the retrospective study design could not exclude selection bias. Second, due to the loss of clinical data, such as precise TNM stage, treatment, and patient outcome, the direct effect of our findings in clinical perspectives is unclear, as well as the protective effect of ARID1A mutation and MSI-H for CRC patients. However, using the biggest dataset (183 samples from AA, 994 from R-CRC, and 1,080 from L-CRC), our results may support the past findings and compare molecular profiles between AA, R-CRC, and L-CRC, and also between minor histopathological subtypes of AA. In addition, our dataset included immune profile and protein expressions, which could lead to selection of treatment strategies. Further large-scale prospective studies with detailed clinical data may be warranted to validate our findings and provide us more information for treatment strategies of AA.
In conclusion, molecular profiling of AA revealed different characteristics from R-CRC and L-CRC, and heterogeneity between the histopathological subtypes of AA. Our data suggests that these molecular differences should be recognized in treating the patients with AA. Upon validation with clinical features, our findings may provide novel insight to develop individualized approach for AA treatment.

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Author contributions:

Conception and design: R. Tokunaga, J. Xiu, W. M. Korn, H. J. Lenz

Development of methodology: R. Tokunaga, J. Xiu, W. M. Korn, H. J. Lenz

Acquisition of data: J. Xiu, C. Johnston, W. M. Korn

Analysis and interpretation of data: R. Tokunaga, J. Xiu, H. J. Lenz

Writing, review, and/or revision of the manuscript: All authors

Administrative, technical, or material support: J. Xiu, C. Johnston, W. M. Korn

Study supervision: Heinz-Josef Lenz
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Figure legends

Figure 1. CONSORT diagram.
Flow chart showing the inclusion/exclusion criteria in this study.

Figure 2. Immune profiling (the status of MSI, TMB, and PD-L1 expression) of AA, R-CRC and L-CRC, and the histopathological subtypes of AA.
A: The characteristics of immune profiles in AA, R-CRC, and L-CRC. B: The characteristics of immune profiles in the histopathological subtypes of AA.
Abbreviations: AA, appendiceal adenocarcinoma; L-CRC, left-sided colorectal cancer; MSI, microsatellite instability; PD-L1, programmed death-ligand 1; R-CRC, right-sided colorectal cancer; TMB, tumor mutational burden.
Table 1. Frequency of gene mutations in AA, R-CRC, and L-CRC.

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<td>0.034</td>
<td></td>
</tr>
<tr>
<td>BRAF</td>
<td>5</td>
<td>17*</td>
<td>5</td>
<td>&lt;0.001</td>
<td>&lt;0.001</td>
<td></td>
</tr>
<tr>
<td>ATM</td>
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<td>7*</td>
<td>5</td>
<td>0.012</td>
<td>0.041</td>
<td></td>
</tr>
<tr>
<td>KMT2D</td>
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<td>7*</td>
<td>2</td>
<td>0.006</td>
<td>&lt;0.001</td>
<td></td>
</tr>
<tr>
<td>PTEN</td>
<td>1</td>
<td>8*</td>
<td>3</td>
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<td>&lt;0.001</td>
<td></td>
</tr>
<tr>
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<td>5*</td>
<td>2</td>
<td>0.016</td>
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<tr>
<td>HNF1A</td>
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<td>1</td>
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<td>&lt;0.001</td>
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<tr>
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<td>5*</td>
<td>1</td>
<td>0.009</td>
<td>&lt;0.001</td>
<td></td>
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<tr>
<td>CTNNB1</td>
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<td>4*</td>
<td>1</td>
<td>0.004</td>
<td>&lt;0.001</td>
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<tr>
<td>KRAS</td>
<td>55</td>
<td>56</td>
<td>43*</td>
<td>0.004</td>
<td>&lt;0.001</td>
<td></td>
</tr>
<tr>
<td>RNF43</td>
<td>7</td>
<td>11</td>
<td>2*</td>
<td>0.004</td>
<td>&lt;0.001</td>
<td></td>
</tr>
</tbody>
</table>

a P value was based on Fisher's exact test. Blanks are P > 0.05.

* P < 0.05 compared to AA.

AA, appendiceal adenocarcinoma; L-CRC, left-sided colorectal cancer; R-CRC, right-sided colorectal cancer.
Table 2. Frequency of gene mutations in the histopathological subtypes of AA.

<table>
<thead>
<tr>
<th>Gene</th>
<th>NOS n = 46</th>
<th>PMP n = 66</th>
<th>MU n = 44</th>
<th>SR n = 27</th>
<th>P value a</th>
<th>P value a</th>
<th>P value a</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>%</td>
<td>%</td>
<td>%</td>
<td>%</td>
<td>NOS vs. PMP</td>
<td>NOS vs. MU</td>
<td>NOS vs. SR</td>
</tr>
<tr>
<td>KRAS</td>
<td>44</td>
<td>74*</td>
<td>64</td>
<td>15*</td>
<td>0.002</td>
<td>0.019</td>
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<tr>
<td>GNAS</td>
<td>7</td>
<td>63*</td>
<td>25*</td>
<td>4</td>
<td>&lt;0.001</td>
<td>0.020</td>
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</tr>
<tr>
<td>TP53</td>
<td>51</td>
<td>23*</td>
<td>57</td>
<td>33</td>
<td>0.004</td>
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<td></td>
</tr>
<tr>
<td>SMAD4</td>
<td>15</td>
<td>15</td>
<td>20</td>
<td>11</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>RNF43</td>
<td>9</td>
<td>6</td>
<td>7</td>
<td>4</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>APC</td>
<td>22</td>
<td>2*</td>
<td>16</td>
<td>0*</td>
<td>&lt;0.001</td>
<td>0.011</td>
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<tr>
<td>PIK3CA</td>
<td>15</td>
<td>2*</td>
<td>7</td>
<td>0*</td>
<td>0.009</td>
<td>0.042</td>
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</tr>
<tr>
<td>BRAF</td>
<td>7</td>
<td>0</td>
<td>9</td>
<td>7</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>ARID1A</td>
<td>11</td>
<td>0</td>
<td>15</td>
<td>11</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

* P value was based on Fisher’s exact test. Blanks are P > 0.05.

* P < 0.05 compared to NOS.

AA, appendiceal adenocarcinoma; L-CRC, left-sided colorectal cancer; MU, mucinous adenocarcinoma; NOS, adenocarcinoma, not otherwise specified; PMP, pseudomyxoma peritonei; R-CRC, right-sided colorectal cancer; SR, signet ring cell carcinoma.
<table>
<thead>
<tr>
<th>Protein</th>
<th>AA N = 183</th>
<th>R-CRC N = 994</th>
<th>L-CRC N = 1080</th>
<th>P value&lt;sup&gt;a&lt;/sup&gt; AA vs. R-CRC</th>
<th>P value&lt;sup&gt;a&lt;/sup&gt; AA vs. L-CRC</th>
<th>P value&lt;sup&gt;a&lt;/sup&gt; R-CRC vs. L-CRC</th>
</tr>
</thead>
<tbody>
<tr>
<td>TS</td>
<td>19%</td>
<td>31*%</td>
<td>18%</td>
<td>&lt;0.001</td>
<td>&lt;0.001</td>
<td>&lt;0.001</td>
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<tr>
<td>ERCC1</td>
<td>32%</td>
<td>16*%</td>
<td>15%</td>
<td>&lt;0.001</td>
<td>&lt;0.001</td>
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</tr>
<tr>
<td>TOPO1</td>
<td>65%</td>
<td>52*%</td>
<td>53%</td>
<td>0.001</td>
<td>0.003</td>
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<tr>
<td>PTEN</td>
<td>88%</td>
<td>65*%</td>
<td>66%</td>
<td>&lt;0.001</td>
<td>&lt;0.001</td>
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</tr>
<tr>
<td>MGMT</td>
<td>69%</td>
<td>52*%</td>
<td>56%</td>
<td></td>
<td></td>
<td>0.047</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Protein</th>
<th>NOS n = 46</th>
<th>PMP n = 66</th>
<th>MU n = 44</th>
<th>SR n = 27</th>
<th>P value&lt;sup&gt;a&lt;/sup&gt; NOS vs. PMP</th>
<th>P value&lt;sup&gt;a&lt;/sup&gt; NOS vs. MU</th>
<th>P value&lt;sup&gt;a&lt;/sup&gt; NOS vs. SR</th>
</tr>
</thead>
<tbody>
<tr>
<td>TS</td>
<td>26%</td>
<td>8*%</td>
<td>23%</td>
<td>27%</td>
<td>0.015</td>
<td></td>
<td></td>
</tr>
<tr>
<td>ERCC1</td>
<td>22%</td>
<td>44*%</td>
<td>27%</td>
<td>27%</td>
<td>0.034</td>
<td></td>
<td></td>
</tr>
<tr>
<td>TOPO1</td>
<td>53%</td>
<td>76%</td>
<td>70%</td>
<td>52%</td>
<td>0.022</td>
<td></td>
<td></td>
</tr>
<tr>
<td>PTEN</td>
<td>83%</td>
<td>98*%</td>
<td>88%</td>
<td>76%</td>
<td>0.010</td>
<td></td>
<td></td>
</tr>
<tr>
<td>MGMT</td>
<td>50%</td>
<td>95*%</td>
<td>60%</td>
<td>38%</td>
<td>0.006</td>
<td></td>
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</tr>
</tbody>
</table>

<sup>a</sup> P value was based on Fisher’s exact test. Blanks are P > 0.05.

* P < 0.05 compared to AA or NOS.

AA, appendiceal adenocarcinoma; L-CRC, left-sided colorectal cancer; MU, mucinous adenocarcinoma; NOS, adenocarcinoma, not otherwise specified; PMP, pseudomyxoma peritonei; R-CRC, right-sided colorectal cancer; SR, signet ring cell carcinoma.
Appendiceal cancer profiled between April, 2015 and January, 2018
N = 224

Neuroendocrine/Goblet cell excluded
N = 41

Appendiceal adenocarcinoma (AA) analyzed
N = 183

Adenocarcinoma, not otherwise specified (NOS)
N = 46

Pseudomyxoma peritonei (PMP)
N = 66

Mucinous adenocarcinoma (MU)
N = 44

Signet ring cell carcinoma (SR)
N = 27

Colorectal cancer profiled between April, 2015 and January, 2018
N = 4,600

Tumors with origin unclearly annotated excluded
N = 2,526

Colorectal cancer (CRC) analyzed
N = 2,074

Right-sided Colorectal cancer (R-CRC)
N = 994

Left-sided Colorectal cancer (L-CRC)
N = 1080
Figure 2

A

Positive ratio

<table>
<thead>
<tr>
<th></th>
<th>AA</th>
<th>R-CRC</th>
<th>L-CRC</th>
</tr>
</thead>
<tbody>
<tr>
<td>MSI-H</td>
<td>2.2</td>
<td>3.4</td>
<td>14.5</td>
</tr>
<tr>
<td>TMB-H</td>
<td>2.2</td>
<td>2.2</td>
<td>14.9</td>
</tr>
<tr>
<td>PD-L1 high</td>
<td>2.8</td>
<td>2.8</td>
<td>4.6</td>
</tr>
<tr>
<td>Any one</td>
<td>5.1</td>
<td>6.7</td>
<td>6.7</td>
</tr>
</tbody>
</table>

\( P < 0.001 \)

B

Positive ratio

<table>
<thead>
<tr>
<th></th>
<th>NOS</th>
<th>PMP</th>
<th>MU</th>
<th>SR</th>
</tr>
</thead>
<tbody>
<tr>
<td>MSI-H</td>
<td>4.4</td>
<td>0</td>
<td>2.4</td>
<td>0</td>
</tr>
<tr>
<td>TMB-H</td>
<td>4.4</td>
<td>0</td>
<td>2.4</td>
<td>0</td>
</tr>
<tr>
<td>PD-L1 high</td>
<td>6.7</td>
<td>1.6</td>
<td>2.3</td>
<td>1.6</td>
</tr>
<tr>
<td>Any one</td>
<td>11.4</td>
<td>4.8</td>
<td>3.9</td>
<td>1.6</td>
</tr>
</tbody>
</table>

\( P = 0.079 \)
Molecular profiling of appendiceal adenocarcinoma and comparison with right-sided and left-sided colorectal cancer

Ryuma Tokunaga, Joanne Xiu, Curtis Johnston, et al.

Clin Cancer Res  Published OnlineFirst January 28, 2019.

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