Mismatch Repair-Deficient Rectal Cancer and Resistance to Neoadjuvant Chemotherapy

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ABSTRACT

Purpose: Evaluate response of mismatch repair-deficient (dMMR) rectal cancer to neoadjuvant chemotherapy.

Experimental Design: dMMR rectal tumors at Memorial Sloan Kettering Cancer Center (New York, NY) were retrospectively reviewed for characteristics, treatment, and outcomes. Fifty patients with dMMR rectal cancer were identified by IHC and/or microsatellite instability analysis, with initial treatment response compared with a matched MMR-proficient (pMMR) rectal cancer cohort. Germline and somatic mutation analyses were evaluated. Patient-derived dMMR rectal tumors were assessed for chemotherapy sensitivity.

Results: Of 21 patients receiving neoadjuvant chemotherapy (5-fluorouracil/oxaliplatin), six (29%) had progression of disease. In comparison, no progression was noted in 63 pMMR rectal tumors (P = 0.0001). Rectal cancer dMMR tumoroids reflected this resistance to chemotherapy. No genomic predictors of chemotherapy response were identified. Of 16 patients receiving chemoradiation, 13 (93%) experienced tumor downstaging; one patient had stable disease, comparable with 48 pMMR rectal cancers. Of 13 patients undergoing surgery, 12 (92%) had early-stage disease. Forty-two (84%) of the 50 patients tested positive for Lynch syndrome with enrichment of germline MSH2 and MSH6 mutations when compared with 193 patients with Lynch syndrome-associated colon cancer (MSH2, 57% vs 36%; MSH6, 17% vs 9%; P < 0.003).

Conclusions: Over one-fourth of dMMR rectal tumors treated with neoadjuvant chemotherapy exhibited disease progression. Conversely, dMMR rectal tumors were sensitive to chemoradiation. MMR status should be performed upfront in all locally advanced rectal tumors with careful monitoring for response on neoadjuvant chemotherapy and genetic testing for Lynch syndrome in patients with dMMR rectal cancer.

Introduction

Screening for deficient DNA mismatch repair (dMMR) is now recommended for all individuals with colorectal cancer (1). While in colon cancers dMMR occurs in 15%–20% of tumors, the prevalence of dMMR in rectal cancer is less frequent at approximately 10%. Microsatellite instability (MSI), the hallmark of dMMR tumors, may be caused by a germline mutation in one of the MMR genes (MLH1, MSH2, MSH6, PMS2, and deletion of EPCAM), consistent with Lynch syndrome, or by epigenetic silencing of the MLH1 promoter region (2). In colon cancer, dMMR tumors differ from MMR-proficient (pMMR) tumors in several aspects, including prognosis, response to treatment, and pattern of metastatic spread (3).

Initial clinical differences in dMMR tumors were described in adjuvant studies of resected early-stage colorectal cancers where tumors exhibiting MSI(dMMR) were noted to have a favorable prognosis but no improvement with adjuvant 5-fluorouracil (5-FU)-based chemotherapy (4–7). The prognostic impact of dMMR appears to diminish with advancing disease stage, and the addition of oxaliplatin to fluoropyrimidines seems to be associated with a survival benefit in stage III MSI/dMMR colorectal cancer (8, 9). In the metastatic setting, dMMR/MSI is considerably less common, occurring in 3%–5% of cases and, while data are sparse and inconsistent, dMMR tumors appear to be less responsive to chemotherapy (10–12), but have a similar response to chemoradiation (13). More recently, checkpoint inhibitors have demonstrated impressive responses in metastatic dMMR colorectal cancers, as well as in other dMMR tumors (14, 15).

The prognostic and predictive role of MSI/dMMR in rectal cancer is not well characterized. In one study, dMMR rectal cancers appeared to have excellent prognosis with traditional trimodality therapy that included neoadjuvant chemoradiation, surgery, and adjuvant systemic
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Translational Relevance

While treatment of locally advanced rectal cancer with neoadjuvant chemotherapy is efficacious and has become more commonplace, our study demonstrates resistance to standard neoadjuvant chemotherapy (fluorouracil/oxaliplatin), both clinically and ex vivo, in patients with locally advanced mismatch repair-deficient (dMMR) rectal cancer. Thus, induction systemic chemotherapy in dMMR locally advanced rectal cancer should be pursued with caution. In comparison, dMMR rectal tumors appear to be sensitive to chemoradiation. The vast majority of dMMR rectal tumors occur in the setting of Lynch syndrome, with an enrichment of MSH2 and MSH6 germline mutations. To optimize treatment selection, all locally advanced rectal tumors should be screened for dMMR with subsequent genetic testing for Lynch syndrome in those with dMMR tumors.

Materials and Methods

Patients

After institutional review board (IRB) approval, we retrospectively reviewed our Center’s clinical cancer database to identify patients with a diagnosis of dMMR colorectal cancer from 2003 to 2018. This study was conducted in accordance with the Declaration of Helsinki and all applicable laws. Lynch syndrome was defined as patients with a likely pathogenic or pathogenic germline mutation in an MMR gene. dMMR was determined by IHC and/or MSI testing. MSI testing was performed either using a standard PCR-based method or using targeted next-generation sequencing (NGS) via MSISensor, a computational algorithm that analyzes sequencing reads at designated microsatellite regions in tumor–normal pairs (23). The distribution of germline mutations in the MMR genes was compared between patients with Lynch syndrome–associated colon and Lynch syndrome–associated rectal cancer.

Among dMMR or MSI-high colorectal cancers, we then identified all those with rectal cancer as defined in the Supplementary Data (22). Patients were divided into three groups based on initial treatment received. Treatment choice was dependent on staging, with clinical stage I patients proceeding directly to surgery. Patients with locally advanced rectal cancer underwent chemoradiotherapy prior to 2012 and TNT after 2012, reflecting the changes in institutional clinical practice after the acceptance of TNT as initial treatment.

Neoadjuvant regimens

The TNT regimen included mFOLFOX, which is described in the Supplementary Data (24–26). Chemoradiation was administered as initial neoadjuvant therapy or commenced 2–4 weeks after completing neoadjuvant chemotherapy. Details of chemoradiation administration and scheduling are in the Supplementary Data.

Response to neoadjuvant chemotherapy or neoadjuvant chemoradiation

To assess the efficacy of neoadjuvant chemotherapy, we evaluated patients who underwent pre- and postchemotherapy (before chemoradiation initiation) staging with rectal MRI and/or endorectal ultrasound and digital examination. Most, but not all, patients received all modalities of evaluation. For chemoradiation response, pathology specimens were evaluated for partial or complete pathologic response according to published criteria (27).

Germline analyses

Clinical germline genetic analyses included full gene sequencing and multiplex ligation-dependent probe amplification analysis for large genomic rearrangement of the implicated DNA MMR gene(s) according to IHC staining patterns. For MLH1/PMS2-deficient tumors, germline analysis included the PMS2 and MLH1 genes, while for MSH2/MSH6-deficient tumors, MSH2 and MSH6 were analyzed. For MSH2/MSH6-deficient tumors, once commercially available, 3’ deletions of the EPCAM gene and MSH2 exons 1–7 inversion analysis were included.

Molecular analyses

Archival formalin-fixed, paraffin-embedded samples of 16 patients (18 samples) were analyzed by using an on-site cancer-associated gene bait capture, NGS assay MSK-IMPACT, using a 341–468 gene panel. The assay is capable of detecting mutations, small insertions and deletions, copy-number alterations, and select structural rearrangements (23). Tumor tissue samples were obtained either pre- or post-treatment at the time of surgical resection or at development of metastatic disease. All subjects undergoing this molecular analysis via MSK-IMPACT provided written consent in the setting of a prospective IRB-approved institutional protocol.

Comparison with historic controls

Outcomes for the patients with dMMR who received upfront chemotherapy or upfront chemoradiation were compared with outcomes for patients with pMMR rectal cancer who received the same treatment modality matched for pretreatment tumor and lymph node staging, based on our previously published study (17), by using a 1:3 pairing (upfront chemotherapy, 21 dMMR:63 pMMR; upfront chemoradiation, 16 dMMR:48 pMMR). The pairings were selected chronologically from our existing database based upon previously published data (17). Baseline demographics were compared between the pMMR and dMMR cases. The spectrum of germline MMR mutations in Lynch syndrome–associated rectal cancer (n = 42) was compared with all patients with Lynch syndrome–associated colon cancer (n = 193) using the Clinical Genetics Lynch syndrome database, which captures patients undergoing germline testing for Lynch syndrome under an Memorial Sloan Kettering Cancer Center IRB-approved (New York, NY) protocol since 2004.

Tumoroids

Pretreatment tumoroids were derived as described previously (28, 29) from fresh rectal tumor samples and embedded in Matrigel. Tumoroids were validated by comparison with the rectal specimens from which they were derived via IHC, targeted gene sequencing, and MMR staining to verify retention of alterations present in the original tumor. Drug treatments with FOLFOX were conducted as reported previously (28) with cell viability assessed using the CellTiter Glo Assay (Promega) following kit protocol. Details on tumoroid treatment and analyses are provided in the Supplementary Materials and Methods and have been published previously (28). All subjects from...
whom tumor tissue was used for tumoroid analysis provided written consent in the setting of a prospective IRB-approved institutional protocol.

**Statistical analysis**

Continuous variables were compared using a two-tailed t test. Categorical variables were compared using X² statistics or Fisher exact test, for limited sample sizes. P < 0.05 was considered statistically significant. Tumoroid IC₅₀ values with 95% confidence intervals (CIs) were calculated as described in the Supplementary Materials and Methods and as described previously (28). Nonoverlapping 95% CIs were used to determine statistical significance with P < 0.05 (30).

**Results**

**Patient and tumor characteristics**

Fifty patients met criteria of dMMR and/or MSI rectal adenocarcinoma with 25 having had IHC alone, seven MSI alone, and 18 having both IHC and MSI analysis. At diagnosis, 38 (76%) had locally advanced disease (stage II or III), three (6%) had metastatic disease, and eight (16%) had stage I disease; one patient’s disease stage was unknown. Of the 50 patients, 21 (42%) received chemotherapy with fluorouracil and oxaliplatin as initial treatment, 16 (32%) received chemoradiation and surgery performed as initial treatment, and 13 (26%) were treated by surgery alone. Of the three patients (14%) who did not undergo surgery, one had local disease progression while on FOLFOX, and one had disease progression with metastasis staging, treatment, and outcomes are outlined in (65%).

**Initial treatment with neoadjuvant chemotherapy**

Twenty-one patients, all meeting criteria for evaluation of response to induction chemotherapy, underwent pre- and postchemotherapy staging with rectal MRI and/or endorectal ultrasound prior to chemoradiation (Fig. 2B). Nineteen had locally advanced and two had stage IV disease. Six (29%) of the 21 patients had local disease progression, with symptoms including change in stool caliber (4) and return of rectal pain (2). Progression was confirmed by rectal MRI (4), CT (1), or endoscopic evaluation (1). Five of the six patients had Lynch syndrome with three harboring an MSH6 mutation. In the remaining 15 patients, tumor response was detected by MRI or endorectal examination.

Seventeen of 21 patients who received initial chemotherapy were evaluable for response to subsequent neoadjuvant chemoradiation. Four (24%) of the 17 patients had a complete response (three with pathologic complete response, one clinical complete response), 11 (65%) had tumor downstaging (T, N, or both), one (6%) was upstaged. Of the four (19%) patients deemed not evaluable, one had inadequate baseline imaging, one had metastatic disease prior to treatment initiation, and two did not receive chemoradiation (one declined due to fertility concerns). Of the two patients who did not receive chemoradiation, one had a complete pathologic response to FOLFOX and the other had disease progression with upstaging on pathologic examination. Details of patient tumor–node–metastasis staging, treatment, and outcomes are outlined in Fig. 2B.

Eighteeneight-sixty-six (86%) of the 21 patients underwent total mesorectal excision. Of the three patients (14%) who did not undergo surgery, one had a clinical complete response and opted for watchful waiting, one had local disease progression while on FOLFOX, and one had metastatic progression in the peritoneum while on FOLFOX.

Of the six (29%) patients whose disease progressed during induction FOLFOX therapy, four received neoadjuvant chemoradiation resulting in downstaging (2) and stable disease (2). The fifth patient did not receive chemoradiation due to peritoneal disease progression. The sixth patient declined chemoradiation for fertility preservation; this patient received 3 months of adjuvant FOLFIRI but experienced rapid progression of disease in the liver and peritoneum with eventual, long-term response to a checkpoint inhibitor (CPI; Fig. 2B). None of these six patients had a prior cancer diagnosis nor received chemotherapy prior to the diagnosis of rectal cancer.

**Initial treatment with neoadjuvant chemoradiation**

Of 16 patients treated with initial neoadjuvant chemoradiation, 14 were evaluable for response and 13 (93%) experienced tumor downstaging, 12 by pathologic findings. Two patients (14%) experienced a pathologic complete response. Two patients did not undergo surgery due to complete clinical response. One patient did not undergo surgery due to synchronous liver disease. Two patients developed disease syndrome, the prevalence of germline MSH2 and MSH6 mutations was significantly higher in the patients with rectal (n = 42) versus colon (n = 193) cancer (rectal vs colon: MSH2: 57% vs 36%; MSH6: 17% vs 9%; P <0.003; Fig. 1B). The mean age at rectal cancer diagnosis in the patients with Lynch syndrome (40.1 years) was equivalent to those without a germline mutation (45.6 years). Eleven (26%) patients with Lynch syndrome had a prior primary cancer diagnosis and 13 (31%) developed metachronous cancers. No prior or metachronous cancers were observed in the patients with non-Lynch syndrome. The mechanism of dMMR in the eight patients with non-Lynch syndrome was found to be MLH1-promoter hypermethylation in two cases, double somatic mutation in one case, and unknown in five cases.

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**Table 1. Patient characteristics.**

<table>
<thead>
<tr>
<th>Characteristic</th>
<th>No. (%) of patients (n = 50)</th>
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<tbody>
<tr>
<td>Age, median (range)</td>
<td>39 (23–83)</td>
</tr>
<tr>
<td>Sex</td>
<td>Male 34 (68)</td>
</tr>
<tr>
<td></td>
<td>Female 16 (32)</td>
</tr>
<tr>
<td>Race</td>
<td>White 39 (78)</td>
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<tr>
<td></td>
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<tr>
<td></td>
<td>Black 1 (2)</td>
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<td>Ashkenazi Jewish 9 (18)</td>
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<td>Not declared 2 (4)</td>
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<tr>
<td>Histology</td>
<td>Well differentiated 0 (0)</td>
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<td></td>
<td>Moderately differentiated 37 (74)</td>
</tr>
<tr>
<td></td>
<td>Poorly differentiated 10 (20)</td>
</tr>
<tr>
<td></td>
<td>Not available 3 (6)</td>
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<tr>
<td>Clinical stage at diagnosis</td>
<td>I 8 (16)</td>
</tr>
<tr>
<td></td>
<td>II 8 (16)</td>
</tr>
<tr>
<td></td>
<td>III 30 (60)</td>
</tr>
<tr>
<td></td>
<td>IV 3 (6)</td>
</tr>
<tr>
<td></td>
<td>Not available 1 (2)</td>
</tr>
<tr>
<td>First treatment for rectal cancer</td>
<td>Surgery 13 (26)</td>
</tr>
<tr>
<td></td>
<td>Chemoradiation 16 (32)</td>
</tr>
<tr>
<td></td>
<td>FOLFOX chemotherapy 21 (42)</td>
</tr>
<tr>
<td>Patients with dMMR who have Lynch syndrome</td>
<td>42 (84)</td>
</tr>
</tbody>
</table>
recurrence and received immunotherapy with checkpoint inhibitors. The treatment course and outcome of these patients is outlined in Fig. 2C.

Initial treatment with surgical resection

Thirteen patients underwent upfront surgical resection; eight due to clinical stage I disease and five for borderline stage I–II disease. The 12 lymph node-negative patients remain disease free, while one patient developed disease recurrence. Three received adjuvant chemoradiation. All patients are alive with a median follow-up of 60 months (range, 25–304).

Comparison of neoadjuvant therapies with pMMR historic controls

We next compared our patients with dMMR rectal cancer who received upfront chemotherapy with our recently published pMMR rectal cancer cohort receiving the same neoadjuvant treatment (17). We sequentially evaluated 63 (3:1 pairing) patients with pMMR and tabulated clinical and pathologic characteristics and evaluated for response using the criteria described above. Demographics for both the pMMR and dMMR cohorts are outlined in Supplementary Table S1. As expected, the patients with dMMR rectal cancer reflect a younger cohort due to the presence of Lynch syndrome. However, no significant differences were noted in other baseline demographics including gender, race, or tumor grade. In contrast to dMMR patients where 29% progressed on FOLFOX, none of the patients with pMMR progressed on neoadjuvant chemotherapy and 89% had tumor downstaging ($P = 0.0001$; Table 2). In a similar fashion, we compared patients with dMMR rectal cancer receiving upfront chemoradiation to 48 (3:1 pairing) pMMR published historic controls. Of baseline demographics, again, only age was different in the pMMR and dMMR upfront chemoradiation groups (Supplementary Table S1). The pathologic response rate of 13% in the dMMR group was similar to the 17% in the pMMR cohort (Table 2).

Recurrent and persistent disease

Of the 50 patients, 12 experienced disease progression or recurrent disease, nine patients in the neoadjuvant chemotherapy cohort, two in the chemoradiation cohort, and one in the surgical cohort. Nine patients were treated with CPIs for persistent (four), locally recurrent (one), or metastatic disease (four). Remarkably, eight (88%) had significant response radiographically and all but one are alive with a mean follow-up time from initiation of immunotherapy of 13 months (9–44 months). One patient with local disease recurrence had significant tumor response on a CPI, which has been sustained for >1 year (Fig. 3A). One patient developed rapidly progressive disease recurrence and died 3 weeks after receiving the first cycle of CPI (Fig. 2A). One patient who was treated with upfront surgery developed a recurrence in the liver and underwent curative liver resection.

Molecular analyses

Sixteen patients who received neoadjuvant chemotherapy as initial treatment had tissue samples available for genomic analyses. All but one patient had tumor sequencing performed on samples obtained after treatment from the resected tumor ($n = 11$), a posttreatment biopsy ($n = 1$) or from a site of metastatic recurrence ($n = 4$). No significant genomic differences, including tumor mutational burden, were found between responders to chemotherapy and nonresponders. There were no BRAF mutations in the eight patients with non-Lynch syndrome (Fig. 3B).
To investigate the association between MMR status and chemoresistance in a biological model, pretreatment rectal cancer tumoroids were derived from pMMR and dMMR rectal tumors as described previously (28). We tested the tumoroids for ex vivo sensitivity to FOLFOX to determine whether our clinical observations would be reflected in the patient-derived tumoroids (28). Indeed, in this proof-of-principle assay, the dMMR rectal cancer tumoroids demonstrated significant resistance to FOLFOX compared with pMMR tumoroids.

**Figure 2.** Overall study flow is demonstrated in A. Patient characteristics, treatments, and outcomes for patients who received neoadjuvant FOLFOX chemotherapy (B) or chemoradiotherapy (ChemOxRT; C) as initial treatment. CR, complete response; F/U, follow-up; NED, no evidence of disease; pCR, pathologic complete response; PD, progression of disease; PR, partial response; SD, stable disease.
Outcomes in patients with rectal cancer with dMMR or pMMR.

<table>
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<tr>
<th>Outcome</th>
<th>dMMR</th>
<th>pMMR</th>
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<tbody>
<tr>
<td>FOLFOX as initial treatment</td>
<td>n = 21</td>
<td>n = 63</td>
</tr>
<tr>
<td>Progression of disease</td>
<td>6 (29)</td>
<td>0</td>
</tr>
<tr>
<td>Response or stable disease</td>
<td>15 (71)</td>
<td>63 (100)</td>
</tr>
<tr>
<td>Chemoradiation as initial treatment</td>
<td>n = 16</td>
<td>n = 48</td>
</tr>
<tr>
<td>Progression of disease</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Complete pathologic response</td>
<td>2 (13)</td>
<td>8 (17)</td>
</tr>
</tbody>
</table>

DICussion

Colon tumors exhibiting dMMR/MSI are biologically and clinically distinct; however, differences resulting from dMMR/MSI status specific to rectal cancers are not well defined. In our study, 29% of dMMR/MSI locally advanced rectal tumors had disease progression on neoadjuvant chemotherapy, an alarmingly high rate compared with no progression in our large cohort of pMMR rectal tumors as well as published data from prospective studies of neoadjuvant therapy in locally advanced rectal cancer (31–33). With the recent shift in the delivery of systematic chemotherapy in locally advanced rectal cancers from the adjuvant to the neoadjuvant setting (18–20, 22), this observation has important clinical implications.

Historically, dMMR/MSI has been both a prognostic and a predictive marker with dMMR/MSI portending a better prognosis in early-stage colorectal tumors, yet limited benefit from fluoropyrimidines (6, 34, 35). As the backbone of systemic therapy, fluoropyrimidine resistance raises concern for primary chemotherapy resistance in patients with dMMR colorectal cancer. In fact, in the metastatic setting, small but well analyzed retrospective data consistently suggest that dMMR colorectal cancers achieve inferior progression-free survival and response rates to chemotherapy compared with pMMR tumors (13). Further supporting our clinical findings are emerging data from the FoxTrot neoadjuvant colon cancer study, demonstrating a significantly worse response to neoadjuvant FOLFOX in dMMR versus pMMR colon tumors, with 75% of dMMR colon tumors exhibiting no pathologic response to chemotherapy. As in our cohort, one dMMR tumor had a complete pathologic response indicating that resistance is not universal (36).

Studies of dMMR/MSI as a biomarker of radiation response have failed to demonstrate its predictive utility (37–39). Our finding, of a 93% response to neoadjuvant chemoradiation, is analogous to previously published dMMR rectal tumors (13) and similar to pMMR rectal cohorts (17). Notably, some patients with progression of disease on neoadjuvant chemotherapy achieved a subsequent response to chemoradiation, suggesting that at least a fraction of such patients may be salvaged with chemoradiation. However, as many patients with locally advanced rectal cancer have bulky and clinically symptomatic tumors, prompt local control with selection of the optimal first-line treatment is imperative.

While our somatic genomic analysis of dMMR rectal tumors did not identify any significant differences in genomic profiles among chemotherapy responders and nonresponders, utilizing tumoroids derived from Lynch syndrome–associated dMMR rectal tumors, we were able to replicate the observed clinical resistance to FOLFOX. In addition, our prior work has shown that dMMR rectal cancer tumors are exquisitely sensitive to radiotherapy (28). These patient-derived biological models of rectal cancer provide the means to study mechanisms of resistance to chemotherapy in real-time and could also help inform whether an early change in therapy is indicated.

Lynch syndrome comprised a very high proportion (84%) of our patients with dMMR rectal cancer, similar to the series by de Rosa and colleagues (13). In contrast, approximately two-thirds of dMMR/MSI colon cancers are driven by epigenetic factors, most commonly MLH1 promoter hypermethylation (40, 41). We also found that the distribution of germline MMR mutations varies in rectal versus colon cancer, with MSH2 and MSH6 mutations accounting for nearly 75% versus only 45% patients with Lynch syndrome–associated rectal cancer and colon cancer, respectively. Most studies have consistently demonstrated that colorectal cancer is more prevalent in MLH1 than MSH2 mutations carriers (2, 16, 17, 25). As the majority of dMMR/MSI cancers arise in the proximal colon, and most prior studies have analyzed dMMR colorectal cancers without distinguishing tumor location (42, 43), the high prevalence of Lynch syndrome in dMMR rectal cancer may have inadvertently been masked. With increasing recognition of the biological importance of colorectal cancer subite stratification (44, 45), whether dMMR rectal tumors portend a different tumor biology from right-sided dMMR tumors remains to be elucidated. Nonetheless, while a rare entity, the diagnosis of a dMMR rectal cancer in a patient should prompt referral for genetic testing and counseling given implications for cancer surveillance and risk-reduction measures.

In our analysis, patient-derived organoid (PDO) models replicated the clinical resistance to FOLFOX chemotherapy in select dMMR/MSI cases. Our group, as well as others, has demonstrated the feasibility of establishing PDOs in evaluating sensitivity to therapy (28, 46, 47). In locally advanced rectal cancer, PDOs may serve as an important test of sensitivity to other agents in select patients including chemoradiation (28) and irinotecan-based therapies (46). Use of organoid models as “oracles of response prediction” in dMMR and pMMR rectal cancers still requires additional validation and prospective testing, but our data suggest that it is possible and provides the first evidence of this potential utility in patients with dMMR rectal cancer.

Universal tumor testing for markers of MMR deficiency is now endorsed for all patients with colorectal cancer (1). However, our study also suggests that, in rectal cancer, MMR status should be determined prior to treatment initiation to allow for optimal selection of first-line treatment. Studies have demonstrated excellent concordance of IHC staining in matched biopsy and colorectal cancer resection samples (48). Moreover, there is also concern that neoadjuvant treatment, especially chemoradiation, may cause inaccuracies in IHC analysis (49). As such, if tissue sample is inadequate from baseline colonoscopy biopsy, a repeat sampling of the rectal tumor prior to treatment initiation is appropriate.

The inherent limitations of our study include its retrospective design, and the heterogeneity of disease stages and subsequent treatment approaches. However, we were able to separate treatment modalities and evaluate response independently with comparison to unselected patients with locally advanced pMMR rectal cancer who were receiving treatment at the same time, utilizing identical modalities, at the same institution. As dMMR rectal cancer is rare, prospective randomized studies are not feasible; however, large national studies like the PROSPECT trial (NCI 2012-00234) in locally advanced rectal cancer may help shed further light on the impact of MMR status at
Figure 3.
A, Endoscopic imaging of a recurrent dMMR rectal adenocarcinoma pre- and 5 and 8 months posttreatment with pembrolizumab. B, Oncoprint and tumor mutational burden in FOLFOX responders and nonresponders with dMMR rectal adenocarcinoma. C, Tumoroid ex vivo sensitivity to FOLFOX demonstrating relative sensitivity to FOLFOX in pMMR tumoroids and resistance in dMMR tumoroids with calculated IC50 values reported as 95% CIs (IC50 = 1.97 (95% CI, 1.49–2.53) vs 5.02 (95%, CI 3.86–6.63), respectively; left). Representative brightfield microscopy of tumoroids in 3D culture both pre- (top micrographs) and post- (bottom micrographs) treatment with FOLFOX chemotherapy from which the IC50 data was derived (right). Decreased confluence demonstrates relative, qualitative resistance to FOLFOX treatment in dMMR tumoroids compared with pMMR tumoroids.
on treatment response. Further studies of the etiology of chemoresistance via DNA analysis and epigenetic modification using dMMR tumors, organoids, and/or cell lines is clearly warranted. Furthermore, we noted a strong sustained response to CPIs in our patients with disease progression or recurrence, in-line with the success of immunotherapy in advanced dMMR solid tumors. Trials of immunotherapy in early-stage dMMR colon cancer (ATOMIC NCT02912559) and in locally advanced rectal cancers are ongoing or planned.

In conclusion, our analyses demonstrate that induction chemotherapy is far less efficacious in dMMR than in pMMR rectal cancer, with over one-fourth of patients progressing while on induction chemotherapy. As such, our data strongly suggests that in patients with dMMR rectal cancer, induction systemic chemotherapy should be undertaken with careful monitoring for tumor response, with close assessment of clinical symptoms. If concern is raised, prompt proctosigmoidoscopy or rectal MRI should be performed to evaluate for response and ensure a timely transition to chemoradiation. In addition, given the importance of MMR status in the optimal selection of first-line treatment for patients with locally advanced rectal cancer, MMR assessment should be performed at the time of rectal cancer diagnosis using biopsy samples, via IHC staining or MSI analysis, with subsequent germline genetic analysis in all those identified to harbor a dMMR rectal tumor.

Disclosure of Potential Conflicts of Interest

A. Cercek is an employee/paid consultant for Bayer and Abbvie Pharma, and reports receiving commercial research grants from Tesaro, Seattle Genetics, and RoGenix. G. Dos Santos Fernandes is an employee/paid consultant for Roche and MSD, and reports receiving speakers bureau honoraria from Roche, Bayer, and BMS. R. Yaeger is an employee/paid consultant for Array BioPharma, and reports receiving commercial research grants from Array BioPharma, Novartis and Boehringer Ingelheim. J.J. Smith is an advisory board member/unpaid consultant for Array BioPharma. N. H. Segal is an employee/paid consultant for PistoOxus, Immunocore PureTech Ventures, Amgen, GSK, CSXone Pharmaceuticals, Synlogic, Pierre, Astrazeneca, Giststone Oncology, TRM oncology, Roche/Gentech, Kyn Therapeutics, Aduaro, Boehringer Ingelheim, Pfizer, Horizon Pharma, and IFM Therapeutics, and reports receiving other commercial research support from Roche/Gentech, Pfizer, Merck, BMS, Astrazeneca, Incyte, and Immunocore. G.M. Nash reports receiving other remuneration from Intuitive. J. Guilleum reports receiving speakers bureau honoraria from Roche. L.A. Diaz is an employee/paid consultant for Personal Genome Diagnostics, Neophere, and 4Paws, and reports receiving commercial research grants from Merck, holds ownership interest (including patents) in Personal Genome Diagnostics, Thrive Detect, 4Paws, and Neophere, and is an advisory board member/unpaid consultant for Merck. J.J. Smith reports receiving other remuneration from Guardant Health Inc. and Intuitive Surgical Inc. Z.K. Stadler’s immediate family member is an employee/paid consultant for RegenxBio, Genentech/Roche, Spark Therapeutics, Adverum, Allergan, Regeneron, BioMarin, and Optos. No potential conflicts of interest were disclosed by the other authors.

References


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