Abstract. Background/Aim: Stomal metastases from a primary rectal adenocarcinoma are rare, therefore, clear guidelines on treatment options are limited. We performed a systematic review including a case report on this subject with the primary objective of identifying the total number of cases in the literature. The secondary objective was to assess median survival. Materials and Methods: A 59-year-old woman presented to our institution with anal incontinence to mucus leakage. Flexible sigmoidoscopy identified a carpet adenoma from the dentate line to the rectosigmoid junction. An abdomino-perineal resection (APR) was performed using the transanal total mesorectal excision technique (TaTME). No adjuvant chemotherapy was offered. Twenty-one months following the operation a stomal recurrence was identified. Palliative radiotherapy was commenced and the patient is alive 6 months later with no visible recurrence at the site of the stoma. A systematic review was performed in accordance with the Preferred Reporting Items for Systematic Review and Meta-Analysis (PRISMA) guidelines. Results: The systematic review identified 19 individual cases of either metachronous or true metastatic recurrence, including our own case. Median survival was 30 months in the 8 cases where further treatment was offered and accepted. Conclusion: Stomal metastases or metachronous colorectal cancer is uncommon. The causes for this pattern of spread are not clear. Long-term survival from cutaneous recurrence is generally poor. For carefully selected patients, redo surgery is an option with satisfactory results.

An estimated 704,000 new cases of rectal cancer were diagnosed worldwide in 2018 according to GLOBOCAN data, making it the eighth most common cancer of all cancers diagnosed globally (1). Low rectal cancers, defined as those “with its lower edge at, or below, the origin of the levators on the pelvic sidewall” (2) should be considered for an abdomino-perineal resection (APR), taking either an intersphincteric approach or, for more advanced tumours, using the extra-levator “ELAPE” technique (2). Although the incidence of APR has fallen in recent years as the use of the laparoscopic low anterior resection (LAR) has increased, it still accounts for about 25% of operations of rectal cancer (3). Local recurrence rates with APR are as high as 33% as opposed to 13% with LAR (4) and five-year survival rates are in the region of 38-60% and 57-76% respectively (4). Proven recurrence at a wound site is rare and sparsely reported in the literature, but an estimate of approximately 4% recurrence has been documented in low rectal cancer (5). There are very few cases in the literature. The latest technique employed to resect rectal cancers is the transanal total mesorectal excision (TaTME). This involves two surgical teams operating from the abdomen and anus with gas insufflation at both ends. The abdominal dissection takes place as for a standard laparoscopic anterior resection. The surgeon operating from the anus creates a rectotomy below the tumour and meets the abdominal surgeon’s resection as the dissection continues cranially, critically outside the mesorectal fascia. A full description of this novel technique is given by Arroyave et al. (6).

We have undertaken a systematic review, which includes our own case report written up below, to identify how many cases of either true metastatic disease or metachronous recurrence have been reported at a stoma site, where the original stoma was created as part of a primary resection for rectal cancer with curative intent. The possible reasons for recurrence and treatment options are discussed. This systematic review was undertaken in response to having had...
a proven stomal metastatic deposit at the site of a colostomy following trans-anal rectal surgery in our institution in 2019. Our case involves a 59 year old female who presented with incontinence to anal mucus leakage. The cause was identified as a large carpet adenoma within which an adenocarcinoma was identified, from the dentate line to the rectosigmoid junction. An intersphincteric APR was undertaken. 21 months following that operation, a stomal recurrence was identified. Palliative radiotherapy was commenced and the patient is still alive 6 months later.

Materials and Methods

A systematic review of the literature was performed in accordance with the Preferred Reporting Items for Systematic Review and Meta-Analysis (PRISMA) guidelines. PubMed, Embase (since 1980), Medline, Google Scholar, CENTRAL, CINNHAL and The Cochrane Library were searched using the following MeSH search terms “peristomal recurrence”, “peristomal metastasis”, “stomal recurrence”, “stomal metastasis”, “rectal cancer”, “colon cancer”, “colonic cancer”, “colorectal cancer”, “rectal adenocarcinoma”, “colon adenocarcinoma”, “colonic adenocarcinoma”, “colostomy recurrence”, “colostomy metastasis”, “colostomy site recurrence”, “colostomy site metastasis”; terms were combined using “and/or” as appropriate with pluralisation also used where indicated (metastasis/metastases). Variations in hyphenation were also included. Bibliographies of relevant papers were used to identify additional studies or reports. Our own case report also used where indicated (metastasis/metastases). Variations in spelling such as MINORs, was not applicable. No randomised control trials were used to identify additional studies or reports. Our own case report was added to the data. The most recent search was conducted on the 3rd of February, 2020. Data points extracted for analysis within the systematic review included: i) the location of cancer, ii) age at initial presentation, iii) type of operation originally undertaken, iv) sex of patient, v) interval between initial operation and stomal recurrence, vi) original histology, vii) overall survival and viii) treatment offered for the recurrence. The data was recorded within a simple spreadsheet (Microsoft Excel), which enabled direct assessment and comparison between the identified cases. Median survival was calculated using the standard formulas within Microsoft Excel. The extracted data points are presented in Tables I and II.

Inclusion criteria. Given the scarcity of these cases, all published articles, case reports, letters and conference abstracts were considered for inclusion where recurrence of a rectal or colonic cancer at the stoma site was mentioned.

Exclusion criteria. Papers published in languages other than English where a translation was not available were excluded.

Outcome measures. The primary outcome from this systematic review was to collate the number of cases reported worldwide to gain insight into how common a problem or not this pattern of spread has become. The secondary outcome was to report the median survival of the identified cases, once a metastatic deposit had been found. In addition, we were interested to report how common metachronous recurrences are, and suggest guidance on how to best treat this rare entity.

Study selection. The primary author (SRD) performed the search, undertaking a title screen followed by an abstract review, and then a full-text review of potentially eligible papers. Where doubt existed as to the relevance of a paper, this was escalated to the senior author for review (KMC). Publications without abstracts directly underwent full text review, if available.

Quality assessment. There was no formal assessment of the quality of the articles considered, since almost all were either case reports or limited case series, thus, applying quality assessment criteria, such as MINORS, was not applicable. No randomised control trials were identified from the search.

Case Report

Our case involves a 59-year-old lady presenting with anal incontinence to mucus leakage, from whom written consent was obtained, permitting the authors to include her case in this systematic review. She had no abdominal symptoms and no family history of gastrointestinal disorders. Flexible sigmoidoscopy revealed a large, circumferential carpet adenoma extending from the dentate line to the rectosigmoid junction. Biopsies of the polypl were villous adenoma with low grade dysplasia. It was agreed with the patient and at the colorectal multi-disciplinary team (MDT) meeting that an intersphincteric APR was the best surgical option as a piecemeal endoscopic excision conferred a high risk of polyp recurrence. The patient was not keen to face the high risk of faecal incontinence associated with a colo-anal anastomosis.

The TaTME operation proceeded uneventfully. At the time of the surgery the appendix appeared smoothly enlarged and was abnormal in appearance. An appendicectomy was performed. The pathology specimen reported a large polypoid tumour mass, 100 mm wide and 65 mm long protruding out of its distal resection margin to a distance of up to 10 mm beyond its wall. Within the polyp, there was a 20 mm, moderately differentiated adenocarcinoma, with high grade dysplasia. The final staging was T2N0M0, with the appendix reported as a low-grade mucinous neoplasm (LAMN), lined by a mostly flat mucinous epithelium with low grade dysplasia. The neoplasm was confined to the appendical wall with no extra-appendiceal cellular or acellular mucin and no invasive malignancy (R0). The decision of the colorectal MDT was to offer a standard post colorectal cancer surveillance. Three months later, staging Computed Tomography scans of the chest, abdomen and pelvis were performed, revealing a 16 mm lymph node located in the external iliac chain which was considered abnormal. The accepted short axis diameter upper size limit is less than one centimetre for oval nodes and less than 0.8 centimetres for round nodes (7). A subsequent Positron Emission Tomography scan revealed that this node was fluorodeoxyglucose (FDG)-avid. Biopsy analysis through fine needle aspiration was inconclusive and the patient went on to undergo a right groin exploration and lymph node dissection. Histopathological analysis of this tissue, using standard staining with haematoxylin and eosin, demonstrated reactive lymphoid tissue only with no malignant changes seen.
Table I. Reported cases in the literature.

| Author          | Year | Cancer | Operation | Gender | Age at original surgery | Age at diagnosis of recurrence at colostomy site | Interval (years) | Original staging |
|-----------------|------|--------|-----------|--------|-------------------------|-----------------------------------------------|-----------------|-----------------|
| Takami et al.   | 1983 | Rectal | APR       | M      | 19                      | 38                                            | 19              | Not stated      |
| Saegusa         | 1986 | Rectal | APR       | M      | 55                      | 60                                            | 5               | Not available   |
| Nakano et al.   | 1987 | Rectal | APR       | F      | 31                      | 53                                            | 22              | Not available   |
| Takeyuchi et al.| 1990 | Rectal | APR       | M      | 44                      | 56                                            | 12              | Not available   |
| Ohta et al.     | 1991 | Rectal | APR       | F      | 68                      | 77                                            | 9               | Not available   |
| Ishikawa et al. | 1994 | Rectal | APR       | F      | 46                      | 76                                            | 30              | Not available   |
| Ohtsuka et al.  | 1996 | Rectal | APR       | M      | 77                      | 81                                            | 4               | Not available   |
| Shibuya et al.  | 1997 | Rectal | APR       | M      | 73                      | 81                                            | 8               | Duke’s A        |
| Mechet-Boillot et al. | 2000 | Rectosigmoid | Hartmans | F | 76 | 78 | 2 | Not available |
| Mechet-Boillot et al. | 2000 | Rectal | AR       | F      | 85                      | 87                                            | 2               | Not available   |
| Townley et al.  | 2005 | Rectal | APR       | F      | 52                      | 57                                            | 5               | Duke’s A        |
| Chintamani et al. | 2006 | Rectal | APR       | M      | 30                      | 36                                            | 6               | T4 N0 M0        |
| Greenberg et al.| 2006 | Rectal | APR       | F      | 44                      | 44                                            | 0.5             | T2 N0 M0        |
| Vijayasekar et al. | 2008 | Rectal | APR       | F      | 47                      | 61                                            | 14              | T4 N0 M0        |
| Varma           | 2011 | Rectal | AR        | M      | 34                      | 51                                            | 17              | Not available   |
| Kuo et al.      | 2012 | Rectal | LAR       | F      | 71                      | 71                                            | 0.5             | T4 N1 M0        |
| Cremades-Pérez et al. | 2015 | Rectal | APR       | F      | 31                      | 58                                            | 27              | Not available   |
| Sarma et al.    | 2015 | Sigmoid | Hartmans | F  | 72 | 74 | 2 | Not available |
| Davey and McCarthy | 2020 | Rectal | TaTME     | F      | 59                      | 61                                            | 2               | T2 N0 M0        |

APR: Abdominoperineal resection; AR: anterior resection; LAR: low anterior resection; TaTME: trans-anal total mesorectal excision.

Table II. Secondary outcomes and treatment undertaken.

| Author          | Year | Original Staging | Overall Survival (after stomal recurrence) | Treatment offered |
|-----------------|------|------------------|--------------------------------------------|------------------|
| Takami et al.   | 1983 | Not stated       | 2 years (alive)                           | 1. En bloc resection of colostomy and 5 cm of distal bowel |
| Saegusa         | 1986 | Not available    | Not available                             | Not available    |
| Nakano et al.   | 1987 | Not available    | Not available                             | Not available    |
| Takeyuchi et al.| 1987 | Not available    | Not available                             | Not available    |
| Ohta et al.     | 1991 | Not available    | Not available                             | Not available    |
| Ishikawa et al. | 1994 | Not available    | Not available                             | Not available    |
| Ohtsuka et al.  | 1996 | Not available    | Not available                             | Not available    |
| Shibuya et al.  | 1997 | Duke’s A         | Not available                             | Not available    |
| Mechet-Boillot et al. | 2000 | Not available | 7 years (died – unrelated – at age 94) | 1. Resection and revision of stoma with adjacent colon |
| Mechan-Boillot et al. | 2000 | Not available | (alive at time of report) | 1. Resection and revision of stoma with adjacent colon |
| Townley et al.  | 2005 | Duke’s A         | 4 years (died)                            | 1. Resection and revision of original colostomy |
| Chintamani et al. | 2006 | T4 N0 M0         | 4 years (died)                            | 1. Wide local excision of stoma, with resection of descending and 2/3rds transverse colon |
| Greenberg et al. | 2006 | T2 N0 M0         | Not clearly stated “months”                | 2. Para and pre-aortic lymphadenectomy |
| Vijayasekar et al. | 2008 | T4 N0 M0         | Not clearly stated                         | 3. 12 cycles - Levamisole and 5 Fluorouracil |
| Varna           | 2011 | Not available    | Not stated                                 | 1. Resection and revision of original colostomy |
| Kuo et al.      | 2012 | T4 N1 M0         | 7 years (alive)                            | 2. Leucovorin and oxaliplatin for further metastatic disease |
| Cremades-Pérez et al. | 2015 | Not available    | 2 years (alive)                           | 1. En bloc resection of mass and completion colectomy |
| Sarma et al.    | 2015 | Not available    | Not stated                                 | 2. Patient declined adjuvant chemotherapy |
| Davey and McCarthy | 2020 | T2 N0 M0         | 6 months (alive)                           | 1. Joint colostomy resection with bariatric operation at the same time |
|                 |      |                  |                                            | 1. 21 cycles of 5 Fluorouracil |
|                 |      |                  |                                            | 1. Resection and revision of original colostomy |
|                 |      |                  |                                            | 1. No details given on treatment plan accepted by patient |
|                 |      |                  |                                            | 1. Palliative intent radiotherapy – 20 Gray in 5 fractions |
Twenty-one months following TaTME surgery a suspicious mass appeared at the muco-cutaneous junction of the colostomy. It measured 15 mm and felt woody underneath the skin. Excisional biopsy confirmed a mucinous adenocarcinoma thought to be related to the rectal cancer rather than the appendiceal neoplasm. Staging investigations revealed a 5.6 cm mass at the site of the end colostomy invading the left rectus muscle. New nodules were felt in the ventral abdominal wall fat, with the largest one measuring 2 cm at the site of the stoma. Additional peritoneal nodules had formed at the midline. Palliative radiotherapy with 20 Gray in 5 fractions was commenced. Having had a good response to treatment with no regression visible at the stoma, the patient was still alive 6 months later. Additional chemotherapy is still an option, however, at the time of writing serious consideration had to be given concerning concurrent chemoradiotherapy due to the Covid-19 pandemic.

**Systematic Review.** The literature search returned 11 articles of which 7 abstracts were identified for further review (see Figure 1: PRISMA flowchart). Two papers were identified through a manual search of the “related articles” function in Pubmed. One further case was added to the dataset from our own institution. We thus identified a total of 9 papers from the systematic review, which, including our case, yielded 19 individual incidences of either metastases or metachronous disease at a stoma site reported since 1983, where the original operation had been undertaken for a rectal or a sigmoid malignancy.

**Study characteristics and primary outcome.** Our primary outcome was to identify the number of cases of cutaneous stomal recurrence after curative rectal cancer. We identified 19 such cases (Table I). Only one non-English written study was identified, for which a translation was obtained. Of the 19 cases, 7 involved males and 12 involved females, with a median age of 52 years (range=19-85) at the time of original surgery. The median time from the original operation to recurrence was 6 years (range=6 months-30 years). In terms of the location of the cancers, two were labelled as either sigmoid or rectosigmoid, while the remaining 17 were located in the rectum.

Staging was either incompletely reported or not available for 12 of the 19 cases. Cases reported before 2000 used the
Duke’s staging criteria (8), while most of the more recent cases (post 2000) used the current standard tumour, node, metastasis criteria, abbreviated to TNM (9). In cases where the original pathology was reported (see Table I), there were two Duke’s A, with two T2 and three T4 cases. Only one case was N1 at the time of original diagnosis. None had metastatic disease on original presentation.

Of the 19 reported cases, 12 (63%) underwent APR and one (5.2%) went through a laparoscopic low anterior resection. Two (10.5%) of the anterior resections were not specified further (24, 27). A Hartman’s procedure was performed in two other cases (10.5%) (24, 28), while in our institution, the case was treated using TaTME (6).

Without a reported, detailed histology of the new deposit, we cannot clearly state which of these cases represents a true metastasis or a metachronous disease. However, looking at the timeframe from the original operation to the diagnosis of the new deposit, 9 recurrences occurred within 5 years or less, while the remaining ones appeared after 5 years. This may indicate that these 9 cases involved metastases, whereas the remaining ones are more correctly categorised as metachronous disease.

Secondary outcomes. Overall survival (OS) of patients was not available, or not clear for 11 of the 19 reported cases (Table II). Of the identified cases where OS was reported, the median survival was 30 months (range=6-84 months), calculated using the standard formulas within Microsoft Excel.

Table II outlines the treatment modalities used in each case. A combination of surgery and adjuvant chemotherapy or radiotherapy is reported with varying results.

Given the relatively few cases in the literature, no further attempt was made to estimate the prevalence of stomal recurrence. Our systematic review suggests that purely stomal recurrence is rare.

Discussion

Whilst there is a high volume of published literature surrounding the cutaneous presentation of colorectal adenocarcinoma, only a few cases of metachronous recurrence and even more rare true cases of metastases at the stoma site have been reported. This systematic review identified a total of 19 cases, out of which only four can be considered as confirmed metastases with five more cases likely to represent metastases. The remaining 11 cases should be considered metachronous disease.

One might argue that the distinction may not seem relevant. However, as discussed further below, categorisation is of vital importance since the emergence of increasing numbers of early recurrence at the stoma site in the literature may indicate that we should re-examine our surgical techniques. Five of the nine cases representing likely metastatic recurrence have occurred since 2005. The reasons for this are currently unclear and may simply reflect greater reporting of such cases in the literature.

At present, there is no consensus as to why stomal recurrence occurs. There is an association with local recurrence and distance metastases (10); however, in the 19 cases identified in this systematic review, no patient was noted to have distant metastases at the time of reoperation. Aside from the established and well-understood patterns of spread in colorectal cancer through lymphatics, haematogenous or direct extension (11), there is clearly a concern that iatrogenic implantation or exfoliation of cancer cells during tissue manipulation in the operation is a possible route for metastatic spread (12). The concern regarding implantation has also been considered for the removal of synchronous polyps at colonoscopy. However there is little evidence thus far to support implantation of malignant cells into the colon by removing multiple polyps (13). Another school of thought is that the spread may arise from micro-metastasis left behind in the lymph nodes along the inferior mesenteric artery pedicle at the time of the APR. In our case, from initial presentation, there was concern that the patient may have micro-metastases within the external iliac chain; however, we were unable to formally diagnose micro-metastatic disease using standard approaches to aspirate and biopsy the nodal chain.

The case we presented here from our institution using the TaTME approach is concerning as there have been other, unusual patterns of recurrence using this approach (14). Previously reported cases have been characterised by rapid, multifocal growth in the pelvic cavity and sidewalls. Norway called a moratorium on the TaTME platform for excision of rectal cancers in 2018 (14), precisely due to the rising concern over these new patterns of recurrence. It should be noted that these concerns surrounding the oncological safety on the TaTME platform are not universally agreed upon (15). Evidence is currently lacking, but the TaTME technique using gas insufflation into the anus in order to facilitate the trans-anal dissection (6), may be part of the explanation as to why these unusual patterns of recurrence have been seen using this relatively new surgical approach. The colorectal community will need to watch closely for the long-term outcomes from this technique.

There is also no reported literature or studies on whether cleaning the stoma using either Betadine or Tisept, prior to bringing it through the abdominal wall, reduces the risk of cutaneous recurrence.

In the papers reviewed, there is little consensus on how to best treat these recurrences. In general, the outcomes are poor since a cutaneous recurrence suggests underlying, disseminated disease. Reports of stomal recurrence prior to 2008 suggest that the median survival after the appearance of
a cutaneous metastatic lesion is between 18 to 20 months (16) with a range from 1 to 34 months (17). However, in this systematic review, which includes more recently reported cases, the median survival is 30 months with a range from 6 to 84 months. It is important to note that in the cases identified for this review, no other sites of metastatic disease or cutaneous disease had been observed prior to the reoperation or instigation of adjuvant therapy. Our systematic review does suggest that if a metastatic deposit is identified at the stoma site, a sensible discussion about redo surgery with a willing patient is a viable option. Interestingly, where involved inguinal lymph nodes are still present after neoadjuvant therapy, separate surgical excision of such nodes, in addition to the resection of the primary tumour, appears to confer a survival benefit (18). This may suggest that when isolated disease recurrence is present, as in our systematic review, curative surgery is an achievable outcome.

Unfortunately, long-term survival when presenting with cutaneous recurrence is rare. Current treatment options involve wide local excisions plus resiting of a stoma, as well as use of adjuvant therapy. When disease is widespread throughout the pelvis, exenteration is an option with reasonable reported outcomes, preferably in a high-volume centre (19). Currently, there is no consensus from the reported cases regarding the best adjuvant protocol for the rare cases of isolated stomal recurrence. Vijayasekar et al. (17) have reported that their patients are offered chemotherapy under the established FOLFOX and FOLFIRI protocols (20, 21); however, patients have turned them down. Sarma and Alvi have stated that their patient was offered palliative chemotherapy with radiation, but no further comment or analysis was made (28). Interestingly, Kuo et al. (22) offered their patient 5-fluorouracil bi-weekly for 21 weeks and the patient was still alive 7 years later. In our own case, the patient received targeted radiotherapy with palliative intent at a dose of 20 Gray in 5 fractions, and is still alive 6 months later. The option of chemotherapy at a later date has not been excluded at this point.

Stomal metastases or metachronous colorectal cancer is uncommon, but they do occur. This systematic review has demonstrated that from the 19 cases we could access worldwide to date, 9 could be interpreted as true metastatic disease.

The cause for this particular pattern of spread is not clear, but given that more cases have been reported in recent years, all units undertaking colorectal resections with newer technology may need to be mindful of this route of spread.

Survival from cutaneous recurrence is generally poor with little understanding of the best way to manage these cases. However, for carefully selected patients, redo surgery can certainly be an option with satisfactory results. For patients with uncertain abdominal lymphadenopathy at the time of rectal cancer diagnosis, we would recommend referral to a specialist centre to consider advanced lymphadenectomy.

Conflicts of Interest

The Authors declare no conflicts of interest.

Authors’ Contributions

SRD undertook the initial searches and prepared the first draft of the manuscript. KMC was significantly involved in the revisions of the manuscript. Both Authors were involved in the preparation and finalisation of the manuscript to be eligible for authorship.

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