Gastric cancer: prevention and treatment of peritoneal metastases

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Abstract

Gastric cancer is an aggressive malignancy that may metastasize through the bloodstream to the liver, through lymphatics to regional lymph nodes, or by penetration of the peritoneal lining of the stomach to result in seeding of the abdominal and pelvic surfaces. Peritoneal metastases are the most common mode of cancer dissemination. Technologies to prevent or treat peritoneal metastases from advanced gastric cancer are presented in this manuscript. The world's literature, both recent and over the past three decades, was reviewed in order to identify publications that present information regarding gastric cancer peritoneal metastases. Over one dozen randomized controlled trials to test perioperative chemotherapy for prevention of peritoneal metastases were reviewed. All of the trials performed with regional chemotherapy during or shortly after gastrectomy were positive. The clinical data regarding the treatment of peritoneal metastases diagnosed at the time of primary cancer resection or in follow-up were reviewed. Neoadjuvant intraperitoneal and systemic chemotherapy shows that some long-term survivors occur after these treatments were combined with cytoreductive surgery and gastrectomy. Similar treatments are advocated for primary gastric cancer with cytology positive for gastric cancer but no visible implants. Surgery for gastric cancer should be combined with perioperative systemic and regional chemotherapy in order to maximally benefit patients with this disease by reducing the negative impact of peritoneal metastases on survival.

Keywords: Hyperthermic intraperitoneal chemotherapy, normothermic intraoperative intraperitoneal chemotherapy, early postoperative intraperitoneal chemotherapy, intraperitoneal chemotherapy, gastric cancer, peritoneal metastases, carcinomatosis
INTRODUCTION

Gastric cancer is the fourth most common cancer in the world with a 5-year survival rate of 25%\textsuperscript{[1,2]}. In follow-up, a large percentage of gastric cancer patients will develop peritoneal dissemination (up to 40%) which results in a less than 5% 5-year survival rate\textsuperscript{[3-5]}. In primary gastric cancer, peritoneal metastases are a common finding present in 5%-20% of patients undergoing gastrectomy\textsuperscript{[6]}. The peritoneum is the most common location of first recurrence in about half of patients\textsuperscript{[7]}. Although the standard of care for treatment of primary gastric cancer involves surgery, intravenous chemotherapy and radiotherapy, specific treatments for peritoneal metastases are poorly defined. Possible treatments include neoadjuvant systemic chemotherapy (NAC), neoadjuvant intraperitoneal and systemic chemotherapy (NIPS), cytoreductive surgery (CRS) and perioperative chemotherapy which may include hyperthermic intraperitoneal chemotherapy (HIPEC) and/or early postoperative intraperitoneal chemotherapy (EPIC)\textsuperscript{[8]}. CRS and HIPEC/EPIC is already considered standard of care for selected patients with appendiceal peritoneal metastases, peritoneal mesothelioma, and a limited extent of peritoneal metastases from colorectal carcinomatosis\textsuperscript{[9-11]}. For gastric cancer with peritoneal metastases, current treatment recommendations remain controversial. The following is an attempt to summarize the role and efficacy of NAC, NIPS, CRS and HIPEC and/or EPIC as prevention or treatment for peritoneal metastases of gastric cancer.

PREVENTION OF PERITONEAL METASTASES USING PERIOPERATIVE INTRAPERITONEAL CHEMOTHERAPY

Surgical treatment failure with resection site and intraabdominal tumors are the most common sites of first recurrence in gastric cancer after potentially curative resection\textsuperscript{[12-14]}. Regardless of neoadjuvant chemotherapy or postoperative adjuvant treatment, this local-regional progression occurs\textsuperscript{[15]}. The peritoneal surfaces and liver remain the major sites of recurrence with a reduced local progression when extended lymphadenectomy as compared to limited surgery is used\textsuperscript{[16-18]}. Although confined to the abdomen, peritoneal seeding has an adverse impact on survival\textsuperscript{[19-22]}. Sources of recurrence after curative resection are (1) spontaneous spreading from the primary tumor; and (2) surgical trauma causing scattering of cancer cells during the surgical procedure. If serosal surface invasion has occurred within the primary tumor, then spontaneous dissemination is more common and patients are frequently found to have viable intraperitoneal cancer cells (positive cytology)\textsuperscript{[19,21-23]}. Tumor cells can also seed the intraabdominal cavity during surgery according to the tumor cell entrapment hypothesis [Figure 1].

During cancer resection, there is transection of lymphatic channels, close margins of resection, and tumor-contaminated blood spillage. Marutsuka et al.\textsuperscript{[24]} identified free cancer cells in peritoneal lavage samples in patients’ initial cytology negative approximately 70 min after dissection of lymph node metastases. Takebayashi et al.\textsuperscript{[25]} showed that gastrectomy spilled viable cancer cells into the peritoneal space in 24 of 57 patients. They concluded that surgery induces peritoneal metastases. Arita et al.\textsuperscript{[26]} determined that large amounts of intraoperative hemorrhage increased the risk of peritoneal recurrence. This may support the contention that cancer cells are present in large numbers within blood lost from the gastric cancer specimen. These iatrogenically disseminated tumor cells adhere spontaneously within minutes and vascularization is facilitated by fibrin entrapment and the wound healing process. Cytokines, such as growth factors important for wound healing, may also propel tumor progression. The tumor cell entrapment hypothesis explains part of the pathogenesis of resection site and distant peritoneal metastases. It is the theoretical basis for adjuvant perioperative intraperitoneal chemotherapy\textsuperscript{[27]}.

Perioperative timing of intraperitoneal chemotherapy

The tumor cell entrapment hypothesis suggests that intraperitoneal chemotherapy must be administered perioperatively in order to access the tumor cells prior to entrapment within fibrin and conversion into cancer implants within adhesive scar tissue. If intraperitoneal chemotherapy is delayed until after the formation of adhesive scars, it will have uneven distribution and lack direct contact with viable cancer cells.
PREVENTION PROTOCOLS USING PERIOPERATIVE CHEMOTHERAPY WITH GASTRECTOMY

Perioperative intraoperative chemotherapy can eliminate progression of peritoneal implantation after curative surgery, however, it cannot treat residual disease within lymph nodes. Therefore, an adequate lymphadenectomy is essential. Intraperitoneal chemotherapy enters the peritoneal nodule by simple diffusion so it only penetrates to 1 or 2 mm\(^2\). It is not effective in lymph nodes. Also, peritoneal nodules larger than 1 or 2 mm have ineffective drug delivery and all visible nodules must be removed prior to treatment.

LITERATURE REGARDING PERIOPERATIVE INTRAPERITONEAL CHEMOTHERAPY FOR ADVANCED T-STAGE PRIMARY GASTRIC CANCER

There have been randomized and non-randomized trials about adjuvant perioperative intraperitoneal chemotherapy compared to surgery alone for resectable primary gastric cancer without peritoneal spread. Sugarbaker \textit{et al.}\(^{[7]}\) published a meta-analysis in 2003 of articles published in English. Xu \textit{et al.}\(^{[29]}\) published a similar study in 2004. Yan \textit{et al.}\(^{[30]}\) published a summary of randomized control trials about adjuvant intraperitoneal chemotherapy for resectable gastric cancer in 2007. Feingold \textit{et al.}\(^{[31]}\) published the most recent summary of non-randomized and randomized studies in English of CRS and HIPEC and/or EPIC in gastric cancer.

Yan \textit{et al.}\(^{[30]}\) selected 10 of 13 randomized controlled trials that were judged to be of fair quality to be used in the meta-analysis. There was a survival benefit associated with HIPEC [hazard ration (HR) 0.060; 95% CI 0.43-0.83; \(P = 0.002\)] or HIPEC with EPIC (HR 0.45; 95% CI 0.29-0.68; \(P = 0.0002\)). There was a marginal benefit with normothermic intraoperative intraperitoneal chemotherapy but no significant improvement in survival with EPIC alone or delayed postoperative intraperitoneal chemotherapy \[\text{Figure 2}\].
Although there may be a survival benefit, perioperative intraperitoneal chemotherapy can increase morbidities. Even the most experienced peritonectomy centers that remove all macroscopic disease and then administer intraperitoneal chemotherapy have a higher morbidity and cost [32-34]. Yan et al. [30] discussed an association of improved overall survival with HIPEC with or without EPIC after resection of advanced gastric primary cancer, however, with EPIC there was an associated greater risk for intraabdominal abscess (P = 0.003) and neutropenia (P = 0.007). Yu et al. [35] also saw an increased risk of intra-abdominal abscess with the use of EPIC compared to the control arm. Intraperitoneal chemotherapy does have less systemic toxicity as compared to systemic chemotherapy. Although individual studies did not show a significant difference in neutropenia between treatment arms, the meta-analysis demonstrated a significantly higher risk of neutropenia in the intraperitoneal chemotherapy arm [30].

Most of the randomized studies were completed in Asia and it is unknown if they can be compared with disease in Western areas. Perioperative chemotherapy may be of greater benefit in Western patients with more advanced disease and less lymph nodes dissected. Data does suggest a role of HIPEC with or without EPIC to improve overall survival for advanced primary gastric cancer with advanced T-stage and no peritoneal metastases. A prospective multi-institutional randomized controlled trial with well-defined eligibility criteria, interventions and end-points is currently in progress in France (D2 resection ± HIPEC) in locally advanced gastric carcinoma, GASTRICHIP, ClinicalTrials.gov Identifier: NCT01882933.

**TREATMENT PROTOCOLS FOR GASTRIC CANCER WITH PERITONEAL METASTASES**

Gastric cancer with peritoneal metastases has been considered a terminal condition. Prospective studies had a median survival of less than 6 months [36]. Although response rates to systemic chemotherapy regimens have improved, there has not been a similar reflection in survival rates [37]. There may be some effective palliation of gastric cancer resections in patients with peritoneal metastases, however there is no long-term improvement in survival.

**CYTOREDUCTIVE SURGERY AND HIPEC AS AN EFFECTIVE STRATEGY**

There is potential for long-term survival for patients with gastric cancer and peritoneal metastases with the combined use of CRS and HIPEC. There are single institutional data and phase II studies that support use of this strategy [Table 1] [31-34,38-41]. Glehen et al. [33] studied 159 patients with a median follow-up of 20.4 months. There was a median overall survival of 9.2 months but the 5-year survival rate was 13%. Although CRS and HIPEC in gastric cancer with peritoneal metastases is less effective than with other...
gastrointestinal malignancies, there were a few long-term survivors. Gastric cancer is a more aggressive
disease. Gastric cancer patients with peritoneal metastases treated with CRS and HIPEC were the only
patients that reported a 5-year survival [39-43]. These studies have shown that strict patient selection criteria are necessary. The extent of peritoneal metastases as measured by Sugarbaker's peritoneal cancer index (PCI) significantly influences survival and is correlated with the completeness of cytoreduction [44]. Cytoreductive surgery must reduce the residual disease to a minimum for intraperitoneal chemotherapy to be effective (due to minimal chemotherapy penetration). Glehen et al. [33] demonstrated a 5-year survival of 23% with median survival of 15 months in patients after a complete macroscopic resection [Figure 3]. Yonemura et al. [45] demonstrated a similar 27% 5-year survival rate and 15.5-month median survival. Hall et al. [34] reported a 11.2-month overall survival after CRS and HIPEC with mitomycin C, however there was no patient alive after 2 years who had residual disease at CRS. CRS with no residual disease burden is essential for effective HIPEC. HIPEC with macroscopic disease burden does not improve survival more than 6 to 8 months. HIPEC can have morbidity and therefore should not be used for patients with bulky residual disease [46]. Palliative use for ascites may always be considered [45,47].

Even if cytoreduction is incomplete, HIPEC is less useful for patients with high burden of peritoneal metastatic disease as measured by PCI. Glehen et al. [33] showed that one of the strongest prognostic factors was extent of carcinomatosis. When the PCI was greater than 12, despite a complete cytoreduction there were no survivors greater than 3 years [Figure 4]. Fujimoto et al. [20] reported 40%-50% 5-year survival for limited peritoneal metastases but only an 18% 1-year survival for patients with extensive peritoneal metastases. Cytoreduction with HIPEC in gastric cancer patients with a greater than 12 PCI score may be contraindicated.

Yang et al. [46] has provided the first and only phase III study regarding CRS and HIPEC in gastric cancer presenting with peritoneal metastases. They used cisplatin (120 mg) and mitomycin C (30 mg) in 6000 mL of normal saline at 43°C for 60-90 min. Median follow-up was 32 months and 97.1% (33 of 34) of patients

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**Figure 2.** Forest plot of the relative risk (RR) of the local-regional recurrence with adjuvant intraperitoneal (IP) chemotherapy versus controls for advanced gastric cancer. The studies were analyzed according to the regimens of intraperitoneal chemotherapy used. The estimate of the RR of each individual trial corresponds to the middle of the squares and horizontal line gives the 95% confidence interval (CI). On each line, the numbers of events, expressed as a fraction of the total number randomized, are shown for both treatment groups. For each subgroup the sum of the statistics, along with the summary RR, is represented by the middle of the solid diamonds. A test of heterogeneity between the trials within a subgroup is given below the summary statistics. (From Yan et al. [30] with permission).
after CRS died as compared to 85.3% (29 of 34) of CRS with HIPEC patients who died. Median survival was 6.5 months (95% CI 4.8-8.2 months) after CRS as compared to 11 months (95% CI 10.0-11.9 months) in CRS with HIPEC group ($P = 0.046$). There was similar morbidity between the groups. The independent predictors in a multivariate analysis for improved survival were synchronous peritoneal metastases, CC 0-1 cytoreduction, more than 6 cycles of systemic chemotherapy, and no adverse events. Glehen et al.\textsuperscript{[48]} suggested that HIPEC should be reserved for patients with limited peritoneal carcinomatosis. Also, the prognostic factors analyzed by Yang et al.\textsuperscript{[46]} suggests that it should be restricted to a limited patient population.

**Role of laparoscopy in patient selection for CRS and HIPEC**

Laparoscopy has been suggested as an important modality for selecting patients for aggressive treatments with CRS and HIPEC. Even today with high technology radiology studies, diagnostic laparoscopy remains
as an important tool to detect disease below a size threshold of approximately 1 cm\textsuperscript{[49]}. If a gastric cancer patient is found to have macronodular small bowel disease or would otherwise not be able to be completely cytoreduced, HIPEC would not be warranted, and the morbidity of exploratory laparotomy could be avoided. Laparoscopy may establish that patients have very limited peritoneal metastases (PCI $\leq 6$) and should be considered for CRS and HIPEC\textsuperscript{[50,51]}. Recent randomized trials suggest that neoadjuvant chemotherapy should be used for gastric cancer patients free of peritoneal disease\textsuperscript{[52]}. Laparoscopy may exclude patients with peritoneal metastases who would not benefit from aggressive neoadjuvant chemotherapy that is unlikely to improve their survival.

**NEOADJUVANT INTRAPERITONEAL AND SYSTEMIC CHEMOTHERAPY**

In medically fit patients with gastric cancer with peritoneal metastases systemic chemotherapy may be recommended. Chemotherapy can provide palliation, improve survival, and improve quality of life compared to best supportive care in patients with metastatic disease. However, the benefits of systemic chemotherapy in gastric cancer patients with peritoneal metastases may be reduced when compared to metastatic disease at other sites. Preusser \textit{et al.}\textsuperscript{[53]} demonstrated that an aggressive systemic chemotherapy regimen can have a 50% response rate in advanced gastric cancer, however this less effective in patients with peritoneal metastases. Ajani \textit{et al.}\textsuperscript{[54]} used neoadjuvant chemotherapy and reported the failure of the regimen was most common in patients with peritoneal metastases. Systemic chemotherapy alone for primary gastric cancer with peritoneal metastases is a disappointing plan of management.

Neoadjuvant chemotherapy for gastric cancer can be modified for patients with peritoneal seeding by combining systemic and intraperitoneal chemotherapy. Chemotherapy may gain access to small peritoneal cancer nodules via the systemic circulation and by diffusion from a chemotherapy solution within the peritoneal cavity. Yonemura \textit{et al.}\textsuperscript{[55]} proposed a prospective phase II study to establish the efficacy and assess toxicities of NIPS chemotherapy in patients with gastric cancer with peritoneal metastases. They identified patients with peritoneal metastases by laparoscopy, laparotomy with biopsy or cytology from ascites. To qualify for NIPS, patients must have: (1) proven peritoneal seeding by histology or cytology; (2) no hematogenous or remote lymph node metastases; (3) be less than or equal to 65 years; (4) have an Eastern Clinical Oncology Group score of 2 or less; (5) adequate bone marrow, liver, cardiac, and renal function; and (6) no other severe medical comorbidities or synchronous malignancies.

Qualifying patients had a peritoneal port system (Bard Port, C.R. Bard Inc., USA) inserted into the abdominal cavity under local anesthesia with the tip placed within the cul-de-sac of Douglas.

**Chemotherapy regimen**

Prior to administration of chemotherapy, 500 mL of saline was instilled into the peritoneal cavity and fluid was removed for cytology. Taxotere 40 mg and carboplatin 150 mg were used for intraperitoneal chemotherapy in addition to 1000 mL of saline over 30 min. Methotrexate 100 mg/m$^2$ and 5-fluorouracil 600 mg/m$^2$ in 100 mL of saline over 15 min were administered intravenously the same day. This regimen was administered weekly for two cycles. After the second cycle, peritoneal wash cytology was again performed. If cytology was positive, neoadjuvant chemotherapy was continued for 2 more cycles. Peritoneal cytology testing is repeating after the fourth cycle and the process is continued as long as cytology is positive.

If cytology became negative, upper endoscopy, laparoscopy and computed tomography scan was performed. If tumors showed no demonstrable change, then 2 more cycles were administered. The number of NIPS chemotherapy cycles was controlled by the effect on the primary cancer and peritoneal cytology. Complete cytoreduction was required for prolonged survival in prior studies that examined peritoneal metastases. Therefore, the goal of the NIPS regimen was complete or near complete response of metastases on small bowel surfaces \[Figure 5\].
SUGERY FOR GASTRIC CANCER WITH PERITONEAL METASTASES AFTER NIPS

Gastrectomy and peritoneectomy were performed if peritoneal wash cytology became negative or there was a partial response to neoadjuvant chemotherapy. If peritoneal metastases on small bowel surfaces were eliminated by NIPS, there was a possibility that gastrectomy and parietal peritonectomy could achieve a complete cytoreduction. Patients with progressive disease or who continue to have positive cytology despite 4 to 6 cycles of NIPS were not candidates for surgery.

Sugarbaker[56] and Yonemura et al.[57] reported the use of peritonectomy for peritoneal metastases to cytoreduce the peritoneal surface and facilitate total resection of all disease associated with the primary gastric cancer. Peritonectomies required for gastric cancer have been described[9]. The epigastric peritonectomy includes any prior midline abdominal scar with the preperitoneal epigastric fat pad, xiphoid process, round and falciform ligaments. The anterolateral peritonectomy removes the greater omentum with the anterior layer of peritoneum from the transverse mesocolon, peritoneum of the right paracolic gutter along the appendix, and the peritoneum in the right subhepatic space. Sometimes the peritoneum of the right and left paracolic gutter must also be removed [Figure 6]. The subphrenic peritonectomy takes the peritoneal surfaces from the medial half of the right and left hemidiaphragm as well as the left triangular ligament [Figure 7]. The omental bursa peritonectomy starts with cholecystectomy and then removes the peritoneal covering of the porta hepatitis, hepatoduodenal ligament, and floor of the omental bursa including the peritoneum overlying the pancreas [Figure 8]. If tumor was within the cul-de-sac, a pelvic peritonectomy was also performed and electroevaporative surgery strips the peritoneum from the pouch of Douglas. Sometimes, the pelvic peritonectomy will necessitate removal of the rectosigmoid colon [Figure 9]. Some or all of these visceral resections and parietal peritonectomies were performed to completely remove visible disease.

RESULTS AFTER NIPS

Canbay et al.[58] analyzed 194 patients treated by NIPS. Average age was 51.5 ± 12.6 years. One hundred and four patients had primary gastric cancer and 90 patients had recurrent PM. Peritoneal fluid cytology was
positive in 137 patients prior to NIPS chemotherapy and 152 patients were negative cytology after treatment. Complete or near-complete response was seen in 51 patients (26%). After induction treatment, 152 (78.3%) patients who showed negative cytology underwent CRS and HIPEC.
Operative interventions were total gastrectomy \((n = 94)\), subtotal colectomy \((n = 68)\), small bowel resection \((n = 44)\). Left and right subdiaphragmatic peritonectomy and pelvic peritonectomy was complete in 44, 31, and 61 patients, respectively. In 102 (67.7\%) patients, of the 152 cytoreductions a complete cytoreduction was recorded.

Figure 5 demonstrates survival of the 194 patients. Median survival was 18 months for all 194 patients. For those who had received surgical intervention, median survival was 15.8 months \(\text{vs.} 9.7\) months for patients who did not have an operation. There was a significant survival difference \((P < 0.001, Z = 20.98)\) between patients who underwent operative intervention \(\text{vs.}\) those who did not. There was a higher median survival of 20.5 months for patients who received a complete cytoreduction \(\text{vs.} 10.9\) months for those who did not have a complete cytoreduction. There was no difference between primary and recurrent disease after cytoreduction.
with a median survival of 18.0 vs. 17.4 months, respectively. If patients did not receive an operation, median survival was similar for primary and recurrent disease as well, 9.6 vs. 8.2 months, respectively.

Another effort to use neoadjuvant intraperitoneal chemotherapy to control peritoneal metastases prior to gastrectomy was presented by Kitayama et al. They used a combination of intraperitoneal and intravenous paclitaxel along with S-1. Repeated laparoscopy was performed to assess response and gastrectomy was used selectively on patients who showed shrinkage of their peritoneal nodules as well as negative peritoneal cytology at a repeat laparoscopy. After a median number of chemotherapy cycles of 5, gastrectomy was performed in 34 of the 64 patients. Sixty-five percent of these patients had an R0 resection. Median survival time and 1-year overall survival of the gastrectomized patients was 26.4 months and 82%, respectively. Those 30 patients who did not receive gastrectomy had a median survival of 12.1 months and a 26% 1-year survival. Kitayama et al. concluded that salvage gastrectomy after intravenous and intraperitoneal paclitaxel was promising even for patients with gastric cancer and peritoneal metastases with ascites.

Fujiwara et al. reported on 18 patients with primary gastric cancer and peritoneal metastases treated with NIPS. After combined intraperitoneal and systemic chemotherapy, 14 patients showed negative peritoneal cytology and no macroscopic peritoneal metastases. The median survival time of his entire group was 24.6 months and there was no treatment-related mortality.

**Neoadjuvant systemic chemotherapy vs. NIPS to date**

Clinical trials comparing the beneficial effects of systemic chemotherapy using modern regimens versus NIPS chemotherapy have not occurred. No doubt, in both treatments, those patients who have a resolution of their peritoneal metastases and then go on to have a successful R0 gastrectomy have a superior outcome. Al-Batran et al. used neoadjuvant systemic chemotherapy followed by surgical resection in patients with limited metastatic gastric or gastroesophageal junction cancer. A small number (4 of 60, 6.7%) had peritoneal metastases as an isolated site of metastatic disease. Nevertheless, the strategy of neoadjuvant systemic chemotherapy prior to resection of all clinical evidence of disease was similar to the NIPS strategy. In their arm B, 36 of 60 (60%) of patients proceeded to surgery. Overall survival of the patients who proceeded to surgery was 31.3 months and 15.9 months for the other patients. These results are similar to the benefits of NIPS followed by cytoreductive surgery. Comparative studies at some time in the future are indicated.

**Adverse events from NIPS and cytoreductive surgery**

The adverse events related to combined therapies NIPS, cytoreductive surgery and then HIPEC may be less than that anticipated for a complex treatment that requires up to 6 months for completion. Problems with the intraperitoneal port are much less than in prior reports of long-term intraperitoneal chemotherapy for ovarian cancer. In this report there were many catheter-related complications, most of which were caused by the extensive peritoneal adhesions. The intraperitoneal ports were placed after a major surgical intervention and only 42% of patients completed all 6 cycles of intraperitoneal chemotherapy. In contrast, catheter-related complications were rare in patients having NIPES because the ports were placed prior to any surgical intervention. Adverse effects grade 3 and 4 were reported in 9% of patients in the multi-institutional study reported by Yonemura et al. in 2012. All of these side effects were from chemotherapy and not catheter-related.

In the 194 patients reported by Canbay et al. in 2014, the most common chemotherapy-related grade 3 or 4 adverse events were bone marrow suppression and diarrhea. Bone marrow suppression occurred after 3 courses in 3 patients, after 5 courses in 3 patients, and after 6 courses in 4 patients. Less common adverse events were port site infection ($n = 2$) and renal failure ($n = 1$).

Prior reports of extensive cytoreductive surgery plus HIPEC following multiple cycles of intraoperative chemotherapy showed an increased morbidity primarily a result of fistula. In the multi-institution report
The grade 3 and 4 complications were 21% and mortality was 3.7%. These morbidity and mortality statistics are approximately the same as reported for cytoreductive surgery plus HIPEC in the absence of NIPS\(^{65,66}\).

**Palliative benefits to all patients with cancerous ascites**

In the publication by Canbay et al.\(^{58}\), there was improvement in symptoms for the 78 patients who had ascites. These benefits occurred in patients with primary gastric cancer and also in patients with recurrent disease. Cunliffe\(^{67}\) hypothesized that peritoneal metastases are nourished via ascites as well as blood supply. Therefore, peritoneal implants should be treated via a combined intraperitoneal and intravenous approach. Intravenous chemotherapy has minimal effects on peritoneal metastases and intraperitoneal chemotherapy alone has a less than 30% effect on ascites\(^{34-36,53,54}\). The bidirectional chemotherapy (intraperitoneal and intravenous) have a response rate of 57% with 100% resolution of ascites.

**Chemotherapy agents selected for NIPS**

According to the study by Morgan et al.\(^{68}\), the maximum tolerated dose (MTD) of intraperitoneal taxotere is 125 mg/m\(^2\) with no grade 3 or 4 toxicities at doses below 80 mg/m\(^2\). Fushida et al.\(^{69}\) showed an absence of hematological toxicities after intraperitoneal taxotere at 45 mg/m\(^2\) given once per week. The MTD of intraperitoneal carboplatin is 500 mg/m\(^2\) and 300 mg/m\(^2\) dose was reported as safe in Japanese ovarian cancer patients\(^{70,71}\). This study safely used taxotere 40 mg/m\(^2\) and carboplatin 150 mg/m\(^2\) combined. The combined use of systemic and intraperitoneal chemotherapy had no deaths and reasonable morbidity and was effective for ascites.

In summary, NIPS should be considered in gastric cancer patients with peritoneal metastases. It has maximal benefits for small volumes of peritoneal surface metastases and is reliable treatment for symptomatic ascites. Bidirectional chemotherapy may be the preferred strategy for preoperative chemotherapy of gastric carcinomatosis.

**Management of primary gastric cancer with positive peritoneal cytology**

The survival of primary gastric cancer patients with positive peritoneal cytology in the absence of macroscopic peritoneal dissemination is very nearly the same\(^{72}\). The 5-year survival rate of cytology-positive but peritoneal metastases negative patients is 2%. Coccolini et al.\(^{73}\) performed a systematic review and meta-analysis concerning the effects of intraperitoneal chemotherapy and peritoneal lavage on this group of patients. Coccolini et al.\(^{73}\) concluded that 2- and 5-year overall survival in patients with free cancer cells without carcinosis is increased by intraperitoneal chemotherapy. Peritoneal lavage further increases these survival rates and also it further decreases the peritoneal recurrence rate.

Shimada et al.\(^{74}\) in 2001 reported in a comparative non-randomized trial about the effects of intraoperative peritoneal lavage either associated or not with intraperitoneal chemotherapy for gastric cancer with free peritoneal cancer cells. Patients treated by intraoperative peritoneal lavage followed by intraperitoneal chemotherapy showed improved survival compared to patients treated by surgery alone or by surgery plus intraperitoneal lavage.

Kuramoto et al.\(^{75}\) reported the results of a comparison between three groups of patients (total 88 patients) with advanced gastric cancer with positive cytology at peritoneal lavage but without peritoneal metastases undergone to surgical resection alone or surgical resection associated either to intraperitoneal chemotherapy or to intraperitoneal chemotherapy plus peritoneal lavage. The 5-year survival rate of the patients who had intraperitoneal chemotherapy plus peritoneal lavage was 43.8%, in the intraperitoneal chemotherapy group was 4.6% and in the surgery alone group was 0% \((P = 0.0001)\). The median survival time of the intraperitoneal chemotherapy plus peritoneal lavage group, intraperitoneal chemotherapy group, and surgery alone group were 35, 16, and 15 months, respectively. The multivariate analysis showed that the peritoneal lavage is the only significant factor affecting the prognosis.
FUTURE PROSPECTS

Beyond the scope of this view are many promising new directions for prevention and treatment of peritoneal metastases from gastric cancer. Targeted therapies may be of great value for subsets of patients, such as those who are HER2-positive (ToGA Study)\[74\]. Also, pressurized intraperitoneal aerosol chemotherapy may substitute for NIPS in preparing patients for subsequent potentially curative resection of all clinically evident disease\[77\]. As these new treatments develop controlled trials comparing the new strategies will be necessary.

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