Some hospitals in Germany offer HIPEC (hyperthermic intraperitoneal chemotherapy) to treat patients with a primary diagnosis of ovarian cancer or recurrent ovarian cancer. As this procedure is currently not indicated for the treatment of patients with ovarian cancer, there is a risk that patients will be denied established procedures that have been proven to be effective, putting patients at risk. We therefore formulated the following statement after analysing the currently available data.

The standard therapy to treat advanced ovarian cancer consists of initial surgery with the goal of achieving macroscopic complete resection, followed by platinum-based intravenous chemotherapy [1–4]. This concept was established based on data from several prospective studies and 10 000 patients and it is the accepted standard worldwide [5]. Recently, the therapy options for patients with FIGO stage IIIB, IIIC and IV cancer have been expanded with the postoperative intravenous administration of the combination carboplatin-paclitaxel with the angiogenesis inhibitor bevacizumab. The preclinical rationale for hyperthermic chemotherapy is based on studies which reported an increased cytotoxicity of cisplatin and other cytostatic drugs in human cell lines and animal models [6–10]. To explain this increased cytotoxicity it was suggested that higher temperatures could overcome cisplatin resistance [11]. Moreover, increased penetration of cisplatin administered intraperitoneally was described when combined with hyperthermia [12]. These theoretical approaches and preclinical observations are the basis for the clinical application of HIPEC. There are currently 3 published randomised studies on the use of HIPEC to treat advanced colon and gastric cancer. A randomised phase III trial for recurrent colorectal cancer in 105 patients investigated the efficacy of systemic therapy alone compared to a combination of cytoreductive surgery and HIPEC (mitomycin C), followed by systemic therapy. The trial demonstrated a significant benefit.

Zusammenfassung

of combined therapy with respect to mean progression-free survival (7.7 vs. 12.6 months; p = 0.020) and mean disease-specific survival (12.6 vs. 22.2 months; p = 0.028). The strongest prognostic factor in this trial was postoperative residual tumour. Only patients with complete tumour resection benefited, while patients with residual tumour intraoperatively showed no benefit from HIPEC. Postoperative mortality in the experimental HIPEC arm was 8% [13, 14]. A further randomised study investigated the efficacy of surgery combined with HIPEC (cisplatin and mitomycin C) compared to surgery alone for peritoneal carcinomatosis of gastric cancer (n = 68). The rate of complete resections was 58% in both study arms. A significant benefit with regard to disease-specific survival (6.5 vs. 11.0 months; p = 0.046) was observed for surgery combined with HIPEC. No data was provided for overall survival. The strongest predictors in this study were the presence or absence of postoperative complications and completion of 6 cycles of postoperative chemotherapy. In absolute terms, patients who only underwent surgery with complete or almost complete resection (max. residual tumour 2.5 mm) had the best prognosis with a mean disease-specific survival of 31 months, while the mean disease-specific survival for patients with complete resection and HIPEC was 12 months [15]. A three-arm randomised study of 139 patients with a primary diagnosis of locally advanced gastric cancer stage T2–T4 compared surgery alone vs. surgery + HIPEC vs. surgery + intraperitoneal chemotherapy. Peritoneal carcinomatosis was not a prerequisite condition for inclusion in the study. Subgroup analysis (serosa invasion or lymph node metastasis) found a benefit for HIPEC [16]. The use of HIPEC is primarily discussed for the therapy of peritoneal carcinomatosis. The different tumour biology and therapeutic concepts make a highly differentiated approach necessary to take account of the different diagnoses. Studies have shown that peritoneal carcinomatosis from primary ovarian cancer has a different tumour biology and a significantly better overall prognosis compared to metastasised gastrointestinal tumours [17]. As the choice of systemic therapies to treat peritoneal metastases of gastrointestinal tumours is limited, HIPEC offers an additional option. But all studies to date have also highlighted an increase in postoperative complications such as infections, and the current S3 guideline on the treatment of gastric cancer therefore only recommends using HIPEC as part of a study (GoR A, LoE I) [18]. An update of the S3 guideline on colorectal cancer which includes an evaluation of HIPEC for this disease entity is still lacking. Early peritoneal metastasis often occurs with ovarian cancer, and most patients are only diagnosed at an advanced stage of disease [19]. For primary surgery, postoperative residual tumour is the strongest independent predictor in addition to tumour stage [4]. The presence of peritoneal carcinomatosis often limits the efficacy of complete resection [20,21]. However, it has not been shown that peritoneal carcinomatosis is in itself an independent predictor [22]. Peritoneal carcinomatosis has been shown to be a negative predictor for complete resection in recurrent ovarian cancer. However, if complete resection of the tumour is achieved, then peritoneal carcinomatosis no longer serves as a prognostic factor [23,24]. Thus, peritoneal carcinomatosis of ovarian, fallopian tube or primary peritoneal cancer appears to be a technical obstacle to complete resection (and in this context has prognostic importance), but by itself it does not appear to be of biological importance and therefore does not require specific treatment – with the exception of the appropriate surgical technique. To date, there are no randomised studies on HIPEC in the context of primary surgery for ovarian cancer or surgery for recurrent ovarian cancer. Some retrospective data has been published as well as a few phase I/II trials with different, mostly platinum-based regimens, dosages and administration times [25]. There are no systematic studies on dosages. None of the studies to date have demonstrated a benefit of HIPEC with regard to overall survival times compared to surgery alone [26], and many studies reported significantly increased complication rates. In contrast to the limited data currently available on HIPEC, results of randomised phase III trials are available for normothermic intraperitoneal chemotherapy. A somewhat higher efficacy was observed for normothermic intraperitoneal chemotherapy regimes but this was accompanied by significantly increased side-effects, particularly for doses repeatedly administered intraperitoneally [27]. Due to the increased side-effects and the lower associated benefit as measured by the much lower rates of therapy completion, intraperitoneal therapy is not currently recommended as a standard option [28,29].

In summary, there is currently no data available which shows an improvement in progression-free survival or overall survival after the use of HIPEC combined with cytoreductive surgery. Based on the available data, the increased rate of surgical complications means that HIPEC cannot be classified as practicable and safe. HIPEC should therefore not be used to treat ovarian, fallopian tube or primary peritoneal cancer outside prospective controlled studies, neither for primary therapy or to treat recurrence. This clear recommendation against the use of HIPEC has also been included in the most recent S3 guideline on the diagnosis and therapy of ovarian cancer and is based on an interdisciplinary consensus.

**Conflict of Interest**

No conflict of interest.

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