Zusammenfassung


Material und Methoden: 14 Patienten zeigten nicht resezierbare hepatische Metastasen solider Tumoren; 13 Patienten wurden in 1–3 Therapiesitzungen mittels Chemosaturation-PHP behandelt. Melphalan 2,0 (n = 1) und 3,0 (n = 12) mg/kg wurde 30 Minuten in die hepatischen Arterien infundiert. Es gab geringere Nebenwirkungen und eine beschleunigte Erholung. In einem Fall musste das Verfahren aufgrund von einer heparininduzierten Thrombozytopenie abgebrochen werden.


Abstrak


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Conclusion: Chemosaturation-PHP for non-resectable liver metastases is a feasible treatment option when performed by an experienced mul-
vaginalen Blutung vorzeitig abgebrochen werden und ein Patient starb aufgrund retroperitonealer Blutung unter Heparinantikoagulation.

**Schlussfolgerung:** Chemosaturation-PHP nicht resektabler Lebermetastasen ist eine geeignete Behandlungsopion, die von einem erfahrenen, multidisziplinären Team durchgeführt werden kann. Es scheint ein aussichtsreiches regionales Verfahren für Patienten ohne andere effektive Behandlungsmöglichkeiten zu sein.

### Introduction

The liver has a unique dual blood supply which makes regional treatment possible. Whereas normal hepatocytes receive their blood primarily from the portal vein, liver tumors are supplied almost exclusively (up to 95%) by the hepatic artery [1]. This allows isolation of the hepatic arterial inflow and venous outflow, and selective delivery of cytotoxic drugs to unresectable liver metastases while sparing healthy liver tissue. Regional chemotherapy procedures include hepatic arterial infusion, percutaneous hepatic perfusion, transarterial (chemo)embolization (TACE) and selective internal radiation therapy [2].

Chemosaturation with percutaneous hepatic perfusion (chemosaturation-PHP) has been developed as a minimally invasive and repeatable regional therapy. It relies on placing a unique double-balloon catheter percutaneously into the inferior vena cava to isolate the hepatic venous blood. High doses of chemotherapy can then be infused directly into the hepatic artery. A fenestrated section in the double-balloon catheter allows the isolated hepatic blood to be filtered extra-corporeally before being returned to the systemic circulation. The feasibility of chemosaturation-PHP has been shown in several studies of patients with unresectable hepatic metastases or primary hepatic cancer [4–9].

Chemosaturation-PHP has been developed commercially (Hepatic CHEMOSAT® Delivery System; Delcath Systems Inc., New York, NY) to make the procedure simpler and more widely accessible. A formal clinical trial program for the Hepatic CHEMOSAT® Delivery System is ongoing; phase I [9], II [10, 11] and III studies [12, 13] have recently been completed. The phase I study established that melphalan 3.0 mg/kg was the maximum tolerated dose deliverable by chemosaturation-PHP [9]. An overall response rate (i.e., complete plus partial response) of 50% was also documented in patients with metastatic ocular melanoma [9]. These favorable results prompted a randomized multicenter phase III trial comparing chemosaturation-PHP delivery of melphalan with best alternative care (BAC) in 93 patients with unresectable hepatic metastases from ocular or cutaneous melanoma. When compared with BAC, chemosaturation-PHP was associated with a significant 6.5-month improvement in hepatic progression-free survival, the primary study endpoint (median 8.1 vs. 1.6 months with BAC; hazard ratio 0.34; p < 0.0001) [13].

Melphalan was selected as the chemotherapeutic agent for the formal clinical trial program of the Hepatic CHEMOSAT® Delivery System on the basis of several observations. Firstly, it does not cause significant liver toxicity even when given at myeloablative doses [14, 15]. Secondly, melphalan delivered by operative isolated hepatic perfusion has previously shown efficacy in patients with hepatic metastases from a variety of cancers, including melanoma [16–19], colorectal cancer [20–22], hepatocellular carcinoma [23], and neuroendocrine tumors [24]. These data show that the melphalan doses deliverable by hepatic perfusion are adequate for efficacy against a range of solid tumors. Lastly, melphalan is widely available and relatively inexpensive, making it an accessible choice for clinics around the world.

The purpose of the present article is to describe our experiences with the Hepatic CHEMOSAT® Delivery System in patients with unresectable hepatic metastases treated at two European centers. It describes the first patients treated in Europe with this delivery system. In addition, a second-generation high-efficiency filter was approved for use in conjunction with the Hepatic CHEMOSAT® Delivery System in Europe in 2012. The phase I, II and III clinical trials were performed using the first-generation filter, so the present report includes the first clinical data with the new filter.

### Patients and Methods

#### Patients

Between January 2012 and February 2013, 14 consecutive patients with unresectable hepatic metastases from various solid tumors underwent chemosaturation-PHP with melphalan at the Frankfurt University Hospital (n = 7) and the European Institute of Oncology in Milan (n = 7). Both centers adhered to the Hepatic CHEMOSAT® Delivery System product instructions with regard to contraindications and precautions. Data evaluation was performed retrospectively.

#### Pre-procedural assessments

Prior to treatment, a physical examination, laboratory tests, and comprehensive imaging including computed tomography (CT) of the thorax and abdomen, upper abdominal magnetic resonance imaging (MRI) and, if clinically indicated, a brain MRI, bone scintigraphy or positron emission tomography (PET) scan were performed. The day before (Frankfurt) or the same day (Milan) of treatment, a complete visceral angiogram was performed to identify potential variant vascular anatomy. Embolization of selected arterial branches supplying the gastrointestinal tract was performed as needed to avoid inadvertent administration of chemotherapy into gastrointestinal or visceral arterial branches.

#### Treatment

Patients received melphalan delivered using the Hepatic CHEMOSAT® Delivery System (Fig. 1). The procedure was performed under general anesthesia in an interventional radiology suite. Melphalan was given at a dose of 3.0 mg/kg ideal body weight (maximum 220 mg/treatment) as a 30-minute infusion into the hepatic artery. Venous effluent blood was filtered via the extracorporeal hemofiltration circuit during and for 30 minutes after each infusion. Percutaneous venous access was performed under ultrasound guidance in order to reduce the number of puncture attempts and any possible cause of bleeding. Heparin (400 IU/kg body weight) was administered during the procedure to ensure free extracorporeal flow and filtration.
Response and toxicity
CT, MRI and/or PET scans of the liver were performed at 4- to 8-week intervals. Tumor response of liver lesions was assessed using RECIST criteria [25]. Systemic and local adverse events were classified by the National Cancer Institute Common Terminology Criteria for Adverse Events, version 3.0. Only systemic events (which did not normalize within 24 hours) and elevations of hepatic transaminases (which did not normalize within 7 days) were reported.

Results

Patients
Patient demographics and tumor characteristics are shown in Table 1. Patients had ocular (n = 8) or cutaneous melanoma (n = 3), breast cancer (n = 1), gastric cancer (n = 1) and cholangiocarcinoma (n = 1). All patients, except for 1, had metastases confined to the liver. All patients had disease progression despite receiving a range of other treatments for hepatic metastases.

Chemosaturation-PHP
13 patients received treatment; melphalan was not administered in 1 patient because of vaginal bleeding. Patients treated in Frankfurt received a single chemosaturation-PHP treatment, and patients treated in Milan received 1 – 3 treatments. A total of 18 (Frankfurt, n = 6; Milan n = 12) chemosaturation-PHP procedures were performed. The interval between repeat treatments ranged from 57 – 177 days. The recommended melphalan dose of 3 mg/kg was given in all patients but one who received a 2 mg/kg dose because of aberrant hepatic vascularization.

The first-generation filter only was used in 3 patients (Frankfurt, n = 2; Milan n = 1), the second-generation filter only was used in 7 patients (Frankfurt, n = 4, Milan, n = 3), and 3 patients (Milan, n = 3) were treated using the first-generation filter for their first treatment and the second-generation filter for repeat treatment(s).

Procedural care
Frankfurt experience
Preprocedural embolization was necessary in 6 patients because gastroduodenal artery (GDA) branches were close to the proper hepatic artery. In one patient with metastases in both liver lobes, chemotherapy was injected into the right hepatic artery to avoid unwanted infusion of

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the gastrointestinal vessels. One patient developed hepatic arterial spasm during the angiogram which was treated successfully with intra-arterial glyceryl trinitrate 0.6 mg. On inflation of the occlusion balloons and establishment of the hemofiltration circulation, transient hypotension was seen in all patients. The median total procedure time was 225 (range, 145–270) minutes. The average venovenous bypass time was 74 (range, 68–81) minutes which included positioning and inflation of the occlusion balloons, the melphalan infusion and washout period.

**Milan experience**
Preprocedural embolization was performed in 6 patients. GDA coiling was necessary in 2 patients because the proper hepatic artery was very short. Right gastric artery embolization was also necessary in 1 of these patients. 3 patients underwent embolization of the right phrenic arteries. In all patients but one, superselective drug injection was achieved by separately cannulating the left and right arteries. One of the patients in whom the hepatic infusion system had previously been implanted was treated separately through the left and right hepatic arteries because of the presence of new tiny vessels from a previously coiled GDA, feeding the duodenum and pancreas.

**Hospitalization**

**Frankfurt experience**
After chemosaturation-PHP, patients were kept in the intensive care unit (ICU) or recovery room and transferred to the ward within a maximum of 24 hours when no complications occurred. Patients remained in the hospital for 4–8 days.

**Milan experience**
After treatment, all patients were admitted to the ICU for an average of 12 hours, except for one patient who died in the ICU 30 hours post-treatment. There was no difference between the first- and second-generation filters in terms of hospital stay (average 5.5 days).

**Tumor response**
1 complete response (Milan, n = 1) and 6 partial responses (Frankfurt, n = 2; Milan, n = 4) according to RECIST criteria were observed. Stable disease was documented in a further 5 patients (Frankfurt, n = 4; Milan, n = 1). 2 patients were not evaluable for tumor response (procedure abandoned because of vaginal bleeding, Frankfurt, n = 1; patient died shortly after treatment, Milan, n = 1).

**Frankfurt experience**
One patient with cutaneous melanoma and multiple liver metastases (diameter ≤ 20 mm) had a tumor response (tumor volume decrease 95 %) after one chemosaturation-PHP treatment (Fig. 2). The residual tumor was treated successfully with vemurafenib plus laser-induced thermotherapy (LITT). On the MRI scan 3 months after chemosaturation-PHP, no more perfused tumor was visible. The patient remained tumor-free for 10 months; disease recurrence (> 50 hepatic lesions) was documented 13 months after treatment. Another patient with multiple liver metastases from ocular melanoma had a partial response (Fig. 3). Stable disease (tumor volume decrease < 5 %) was observed in a patient with hepatic and bone metastases from breast cancer. Two months after chemosaturation-PHP, the follow-up MRI scan
showed disease progression in the liver. Ten months after treatment, the patient died from progression of liver metastases. The remaining 3 patients with stable disease had ocular melanoma. Two months after chemosaturation-PHP, stable disease was maintained in 1 patient. Without any further therapy, the follow-up 6 months post-treatment imaging showed a partial response which is still ongoing 2 months later (Fig. 4). In another patient with bilobar disease, melphalan was infused only into the right hepatic artery, and complete hepatic perfusion was not achieved. Tumor response was evaluated only in the perfused right hepatic lobe (Fig. 5). The first follow-up after treatment showed stable disease in the right liver lobe and disease progres-

**Fig. 3** Magnetic resonance images (T2 haste) pre-treatment a and 8 weeks post-treatment b of a patient with metastatic ocular melanoma. The liver metastases showed a reduction in size of over 30% (partial response).

**Abb. 3** MRT-Bilder (T2 haste) vor a und 8 Wochen nach der Behandlung b eines Patienten mit metastasiertem Aderhautmelanom. Die Lebermetastasen zeigen eine Größenreduktion von über 30% (partielle Remission).
sion in the untreated left lobe. The left lobe was subsequently treated with TACE. However, disease progression in both liver lobes was documented after 2 months and the patient died shortly afterwards. In the other patient with metastatic ocular melanoma, the left and right hepatic arteries were cannulated separately and the right liver lobe was perfused with 2/3 of the melphalan dose and the left one with the remaining 1/3. One month after chemosaturation-PHP, stable disease was seen in the right liver lobe, whereas disease progression occurred in the left lobe. A follow-up scan 2 months post-treatment showed progressive disease in the whole liver. The patient died 6 months after treatment from massive progression of liver metastases.

**Milan experience**

Of the 6 patients evaluable for response, 1 patient with biliary tract adenocarcinoma achieved complete remission 6 months after chemosaturation-PHP. 2 patients with ocular melanoma had partial responses (MRI and PET) after the first of 3 sessions of chemosaturation-PHP. One of these patients remained stable with minimal disease after 2 further sessions (MRI/PET-negative but some small nodules visible). The other patient showed disease progression in the liver and bones after the third procedure, 1 year after the first session. One patient with cutaneous melanoma showed a partial response (MRI and PET) after the first chemosaturation-PHP session. The patient then received 4 courses of ipilimumab and recently had a second chemosaturation-PHP session. One patient with gastric cancer had a mixed response due to parasitic hepatic arteries that were not included during chemoperfusion. He had 3 consecutive liver resections before chemoperfusion. A huge portion of liver parenchyma, close to the previous resections, was fed by several arteries from the intercostal and mammary arteries. Lesions located within the non-perfused hepatic parenchyma grew after treatment, while metastases within the chemoperfused parenchyma showed a partial response. Because it was not possible to manage this kind of hepatic vascular supply, a second chemosaturation-PHP session was not scheduled. One patient with cutaneous melanoma had a mixed response after the first perfusion, due to the growth of some lesions deep within segment 7, located under the dome. A phrenic artery feeding that area was occluded immediately before the second session and a partial response was documented in all target liver lesions (tumor volume reduction > 40%). Unfortunately, this patient subsequently developed multiple lung metastases.

**Toxicity**

**Frankfurt experience**

In the 2 patients treated with the first-generation filter, both developed grade 1/2 increases in liver transaminases in the first 2 days after chemosaturation-PHP; all values returned to baseline levels within 1 week. Grade 1/2 fatigue, nausea and fever within the first 5 days after treatment were also reported. In both patients, grade 1/2 bone marrow suppression was observed initially. However, nadir blood cell counts were reached after 11–12 days and both patients were hospitalized for 8–9 days because of grade 3/4 pancytopenia. Grade 3/4 leukocytopenia was treated with filgrastim for 8–9 days in both patients, and antibiotics were required in 1 patient. Platelet transusions were required for grade 3/4 thrombocytopenia (n=2), and 2 units of packed red blood cells (RBC) were required for grade 3 anemia (n=1). Of the 4 patients treated with the new filter system, 1 patient developed grade 1/2 pancytopenia after treatment which worsened to grade 3/4 pancytopenia after 1 week necessitating hospitalization for 6 days. However, only the leukocytopenia required treatment (filgrastim for 3 days plus an antibiotic). The second patient received a lower dose of melphalan and, with the exception of grade 1 anemia, did not experience any systemic adverse events. The last 2 patients both developed grade 1 anemia, grade 3/4 leukocytopenia and grade 2 or 4 thrombocytopenia and were treated as outpatients with platelet concentrates. Granulocyte colony-stimulating factor (G-CSF) was given for 4 days. No significant increases in transaminases occurred.

A premenopausal patient developed vaginal bleeding after systemic heparinization. Treatment was stopped before melphalan
administration. A gynecological examination showed that the event was most likely caused by heparin-induced bleeding of the endometrium. The patient recovered without sequelae.

Milan experience

Four patients were treated with the first-generation filter and all of them required multiple platelet and/or RBC transfusions after the procedure. One patient had grade 4 leukocytopenia, grade 3 anemia and grade 4 thrombocytopenia; the patient developed febrile neutropenia was admitted to the hospital to receive intravenous antibiotics and other support therapies. Three patients developed grade 4 leukocytopenia, grade 2/3 anemia and grade 1 or 4 thrombocytopenia which required transfusions without hospitalization. All of these patients had grade 2/3 fatigue and received filgrastim for at least 10 days. 3 of these 4 patients underwent 1–2 more sessions of chemoperfusion with the second-generation filter. These procedures were associated with markedly reduced hematological toxicity (grade 1 anemia) without any need for blood or platelet transfusions; G-CSF was required for 4 days only. All 3 patients experienced milder fatigue (grade 1). Of the 3 patients treated with the second-generation filter only, a similar toxicity profile was evident in 1 patient, while another patient developed febrile pancytopenia requiring transfusions because of a hepatic vascular shunt that avoided drug filtering. No patient had significant liver dysfunction, and grade 1–2 liver toxicity related to drug exposure was not influenced by the type of filter used. The remaining patient, who had a large fast-growing liver metastasis from ocular melanoma, died of a retroperitoneal giant hematoma 30 hours after chemosaturation-PHP. A post-mortem necropsy revealed multisite vascular bleeding with no damage to the inner surface of the abdominal veins and arteries. This unusual complication was most likely related to heparin which was needed for extracorporeal circulation.

Discussion

Chemosaturation-PHP is a minimally invasive, repeatable technique which delivers high doses of chemotherapy directly to tumors in the liver while limiting systemic toxicity through hemofiltration of the hepatic venous blood. Recently completed clinical trials performed in the US confirm the efficacy of chemosaturation-PHP delivery of melphalan in the treatment of patients with hepatic metastases from melanoma [12, 13] and neuroendocrine tumors [10, 11].

Our experience with chemosaturation-PHP at the University Hospital Frankfurt and the European Institute of Oncology is supportive of the findings of the clinical trial program, although it is too early to make definitive statements about tumor response rates and patient survival. Still, the data offer a number of insights. A complete response was documented in 1 patient with cholangiocarcinoma and partial responses were observed in a further 6 patients with ocular or cutaneous melanoma, giving an overall response rate of 50% in the total patient sample (n = 14). Of note, one of the melanoma patients became tumor-free after further treatment with LITT plus vemurafenib. The response lasted for 10 months before a recurrence in the liver was detected. At the time of writing, this patient is still alive with progression of liver metastases. 4 of the other 5 melanoma patients are also alive 7 to 18 months after their first chemosaturation-PHP session; all of them had disease restricted to the liver but were in progression after previous treatments and had no other therapeutic options (BRAF and C-kit wild type). The last melanoma patient died after 9 months. These observations compare favorably with the shorter median overall survival times typically reported in patients with liver metastases from cutaneous [26] or ocular melanoma [27, 28]. In addition, 5 other patients with ocular melanoma, gastric cancer or breast cancer experienced clear clinical benefit from the procedure, opening up the possibility of combining chemosaturation-PHP with other systemic treatments.

The toxicities observed in our patients were consistent with the profile of events documented in clinical trials [9, 11]. Mild to moderate filter-related toxicity, i.e. thrombocytopenia and anemia resulting from the removal of platelets and RBC by the hemofiltration system, was observed immediately after the procedure. Only patients treated with the first-generation filter needed platelet and RBC transfusions. Persistent and more severe melphalan-related pancytopenia tended to emerge later. Although these events were generally grade 3/4 in severity, they were predictable and were managed effectively in all patients with supportive measures. Transient increases in liver transaminases (grade 1/2) were also observed during the post-procedural period in some patients resulting from procedural manipulation of the liver rather than melphalan. In all cases, values returned to baseline levels within 1 week. Other systemic events were mild (nausea, vomiting, fever), apart from fatigue which significantly influenced patient performance after treatment with the first-generation filter. Two events were attributed to heparin which is routinely administered during chemosaturation-PHP to facilitate extracorporeal blood flow and hemofiltration. In one case the procedure was stopped prematurely due to vaginal bleeding, and in another case a perioperative death resulting from a retroperitoneal hematoma occurred.

In April 2012, a second-generation high-efficiency filter (98% bench-testing efficiency; Delcath Systems Inc., data on file) for use with the chemosaturation-PHP delivery system became commercially available in Europe. In Milan, three patients were treated initially with the first-generation filter and then switched to the new filter for repeated treatments, providing a unique opportunity to compare toxicities with the two systems. In these patients, toxicity with the new filter system was less severe and patients required fewer supportive measures (i.e., no transfusions, shorter courses of colony-stimulating factors, reduced fatigue). Further studies are required to confirm this positive trend. A phase III study to evaluate chemosaturation-PHP in the treatment of hepatocellular carcinoma is planned in Frankfurt, and the new filter system will also be investigated further.

In summary, chemosaturation-PHP for the treatment of non-resectable liver metastases is a feasible treatment option when performed by an experienced multi-disciplinary team. Hematological events, which are the predominant toxicities associated with chemosaturation-PHP, are predictable and manageable with appropriate supportive care. Based on our findings, we believe that chemosaturation-PHP is a promising technique for patients for whom there are no effective treatments.
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