Introduction

Thrombotic microangiopathy (TMA) is a rare but potentially devastating disease. It is defined as microangiopathic hemolytic anemia (MAHA) in combination with thrombocytopenia and signs of organ damage. These pathological features are included in different clinical syndromes, such as thrombotic thrombocytopenic purpura (TTP), hemolytic uremic syndrome (HUS), and also the HELLP syndrome in pregnancy. Recently, the pathophysiology of primary TTP has been widely elucidated. A severe deficiency of the metalloprotease ADAMTS13, either due to mutations or to autoantibodies, is presumed to be the main pathophysiological mechanism [1].

Keywords
ADAMTS13 · Bone marrow infiltration · Microangiopathic anemia · Thrombotic microangiopathy · Thrombotic thrombocytopenic purpura
Secondary TMA can be triggered by many factors such as various drugs, bone marrow transplantation, infections, and malignant diseases. In cancer-associated TMA at least 2 different entities can be identified: chemotherapy-induced TMA and TMA induced directly by the cancer (non-chemotherapy-induced). Chemotherapy-induced TMA is described with a variety of known chemotherapeutic drugs such as mitomycin C, bleomycin and cisplatin [2], but also with newer agents such as gemcitabine [3]. The incidence of chemotherapy-induced TMA varies between 0.1 and 10%, depending on the drug. Only non-chemotherapy-induced TMA is the subject of this report.

Non-chemotherapy-induced TMA is mainly seen in late-stage metastasized carcinomas. The underlying cancers are predominantly adenocarcinomas. There is increasing evidence that bone marrow infiltration, frequently seen in prostate, lung, breast, and ovarian cancer, is associated with TMA. However, many questions concerning pathophysiological aspects as well as the best therapeutic options are as yet unresolved.

Therefore, we have retrospectively analyzed all consecutively diagnosed patients with breast cancer-associated TMA. Pathophysiological aspects and therapeutic options are discussed. We present a promising therapeutic approach with fractionated chemotherapy resulting in longer survival compared to previously published reports.

### Methods and Patients

We retrospectively analyzed all patients with breast cancer-associated TMA diagnosed at our institution between 2003 and 2008. Diagnosis was based on the presence of thrombocytopoenia and Coombs-negative hemolytic anemia, as previously reported [4–6].

The patients were consecutively diagnosed and treated at our institution. Information recorded for each patient included age, tumor characteristics (size, regional lymph node status, histological characteristics, grading, hormone receptor status, and HER2/neu status), characteristics of metastases (localization and number of sites), therapy (palliative systemic treatment), and disease progression (time to progression and overall survival). The response evaluation criteria in solid tumors guidelines were used to assess the response to therapy. Laboratory findings, clinical signs and symptoms, treatment effects, and toxicity data were derived directly from the patient charts.

The following parameters of clinical routine were analyzed in the study: normal blood count, differential blood count, parameters of hemolysis (bilirubin, lactate dehydrogenase (LDH), haptoglobin), Coombs' test, renal and hepatic parameters, basic coagulation parameters (prothrombin time (PT), activated partial thromboplastin time (aPTT)).

In order to generate more insight into possible pathological mechanisms, we also measured the levels of von Willebrand factor (vWF) antigen and the serum activity of the vWF-cleaving protease, ADAMTS13. ADAMTS13 activity was measured by a modified method described by Furlan et al. [7].

### Results

We identified 8 patients with breast cancer-associated TMA treated at our institution between 2003 and 2008, out of about 750 breast cancer patients. The baseline characteristics are presented in Table 1.

| Patient | Age | TNM at initial diagnosis | Grading | Histology | Mucinous HER2/neu | Visceral metastases | Bone marrow infiltration | Bone metastases | Bone marrow infiltration | Bone metastases | Bone marrow infiltration | Bone metastases | Bone marrow infiltration | Bone metastases | Bone marrow infiltration | Bone metastases | Bone marrow infiltration | Bone metastases | Bone marrow infiltration | Bone metastases | Bone marrow infiltration | Bone metastases | Bone marrow infiltration | Bone metastases | Bone marrow infiltration | Bone metastases | Bone marrow infiltration | Bone metastases | Bone marrow infiltration |
|---------|-----|-------------------------|---------|-----------|------------------|-------------------|----------------------|---------------------|----------------------|---------------------|---------------------|---------------------|---------------------|---------------------|---------------------|---------------------|---------------------|---------------------|---------------------|---------------------|---------------------|---------------------|---------------------|---------------------|---------------------|---------------------|---------------------|---------------------|---------------------|---------------------|
| 1       | 62  | T3N0M0                  | G3      | ILC       | yes              | neg.              | no                   | yes                 | no                   | yes                 | no                   | yes                 | no                   | yes                 | no                   | yes                 | no                   | yes                 | no                   | yes                 | no                   | yes                 | no                   | yes                 | no                   | yes                 | no                   | yes                 | no                   |
| 2       | 78  | T4N3M0                  | G3      | IDC       | no               | neg.              | no                   | yes                 | no                   | no                  | no                   | yes                 | yes                 | no                   | yes                 | yes                 | no                   | yes                 | no                   | yes                 | no                   | yes                 | no                   | yes                 | no                   | yes                 | no                   | yes                 | no                   |
| 3       | 67  | T3N1M0                  | G2      | ILC       | yes              | pos.              | no                   | yes                 | no                   | no                  | yes                 | yes                 | yes                 | yes                 | yes                 | yes                 | yes                 | yes                 | yes                 | yes                 | yes                 | yes                 | yes                 | yes                 | yes                 | yes                 | yes                 |
| 4       | 60  | T1N3M1                  | G1      | ILC       | yes              | pos.              | neg.                 | yes                 | no                   | yes                 | yes                 | yes                 | yes                 | yes                 | yes                 | yes                 | yes                 | yes                 | yes                 | yes                 | yes                 | yes                 | yes                 | yes                 | yes                 | yes                 |
| 5       | 35  | T1N2M0                  | G3      | IDC       | unknown          | neg.              | no                   | yes                 | yes                 | yes                 | yes                 | yes                 | yes                 | yes                 | yes                 | yes                 | yes                 | yes                 | yes                 | yes                 | yes                 | yes                 | yes                 | yes                 | yes                 | yes                 |
| 6       | 56  | T3N2M0                  | G2      | IDC       | no               | pos.              | neg.                 | no                   | no                   | yes                 | yes                 | yes                 | yes                 | yes                 | yes                 | yes                 | yes                 | yes                 | yes                 | yes                 | yes                 | yes                 | yes                 | yes                 | yes                 | yes                 |
| 7       | 33  | T2NxM1                  | G3      | ILC       | no               | pos.              | neg.                 | yes                 | no                   | yes                 | no                   | yes                 | yes                 | yes                 | yes                 | yes                 | yes                 | yes                 | yes                 | yes                 | yes                 | yes                 | yes                 | yes                 | yes                 | yes                 |
| 8       | 51  | TxNxMx                  | G2      | IDC       | unknown          | pos.              | neg.                 | yes                 | yes                 | yes                 | yes                 | yes                 | yes                 | yes                 | yes                 | yes                 | yes                 | yes                 | yes                 | yes                 | yes                 | yes                 | yes                 | yes                 | yes                 | yes                 |

TNM = Tumor, node, metastasis, ILC = invasive lobular carcinoma, IDC = invasive ductal carcinoma.

Bone marrow infiltration was diagnosed by bone marrow biopsy in all cases.*

*Bone marrow infiltration was diagnosed by bone marrow biopsy in all cases.

The following parameters of clinical routine were analyzed in the study: normal blood count, differential blood count, parameters of hemolysis (bilirubin, lactate dehydrogenase (LDH), haptoglobin), Coombs' test, renal and hepatic parameters, basic coagulation parameters (prothrombin time (PT), activated partial thromboplastin time (aPTT)).

In order to generate more insight into possible pathological mechanisms, we also measured the levels of von Willebrand factor (vWF) antigen and the serum activity of the vWF-cleaving protease, ADAMTS13. ADAMTS13 activity was measured by a modified method described by Furlan et al. [7].
Breast Cancer-Associated Thrombotic Microangiopathy

Interestingly, 50% of the patients were diagnosed with invasive lobular carcinoma, which is a higher proportion than in general breast cancer cohorts (10–20%). We found evidence of mucine-producing carcinoma in 3 patients. All patients were widely metastasized, with visceral involvement in 5/8 patients, and all showed bone marrow infiltration diagnosed by bone marrow biopsy. In 5 patients, the diagnosis of TMA was simultaneous to the diagnosis of distant metastases.

Clinical signs and symptoms at the onset of TMA are shown in table 2. 3 patients presented in a poor performance status (PS ≤ 50), whereas 2 patients were only slightly impaired. In contrast to classical TTP/HUS, most of our patients showed only comparatively few signs of organ damage, although mild cerebral symptoms were found in the majority.

Table 3 outlines the laboratory findings. All patients had severe changes in their normal blood count. Normochrome anemia with a median hemoglobin level of 5.19 mmol/l was detected. Anemia was proven to be Coombs-negative hemolytic anemia with schistocytes in all cases. Thrombocytopenia was also present in all cases, ranging from only a slight (95 × 10⁹/l) to a massive reduction (4 × 10⁹/l). The median platelet count at the time of diagnosis was 30 × 10⁹/l. Also, a severe elevation of the LDH level was found in 7/8 patients. Surprisingly, creatinine was within normal limits in all but 1 patient.

The basic coagulation parameters were only slightly altered (median PT 68% (normal value 70–120%), median aPTT 34.05 (normal limits 26–40)) whereas the vWF antigen was severely elevated in 6/7 patients.

Table 4. Therapeutic and survival data

A = Adriamycin, C = cyclophosphamide, F = 5-fluorouracil, PR = partial response, SD = stable disease.
20 mg/m² weekly was started after 2 days. Although a short stabilization was achieved, the patient died 39 days after the diagnosis of TMA. Patient 2 presented in a very poor condition. After the diagnosis of TMA, plasmapheresis was considered. However, the patient deteriorated rapidly with an evolution towards multi-organ failure. Therefore, no specific therapy against TMA was initiated. The patient died 4 days after diagnosis.

Because of a growing body of evidence that patients with cancer-associated TMA benefit only slightly from plasmapheresis, we changed our therapeutic algorithm to initiate chemotherapy as early as possible and to withdraw plasmapheresis whenever possible (from patient 3 onwards).

In order to minimize the hematologic toxicity of standard chemotherapy, we used fractionated regimens according to the individual patient history. As anthracyclines are highly effective in breast cancer, we used adriamycin (A) either as monotherapy or in combination with cyclophosphamide (C) and 5-fluorouracil (F). The following dosages were used: A mono: 20 mg/m²/d1 weekly; AC: 20/200 mg/m²/d1 weekly; FAC: 200/20/200 mg/m²/d1 weekly.

The response rate with partial remission in 5/6 patients was surprisingly high. The responses included a clinically significant improvement of laboratory parameters (hemoglobin, thrombocytes, and LDH) in all and even normalization in 4 of the responding patients. The median duration of response was 8.5 months. The overall survival with a median of 13 months was higher than previously reported.

Inevitably, progression occurred in all patients. In 2 patients, the peripheral blood count deteriorated again, but the progression of the systemic disease, including mainly the progression in visceral metastatic sites, was the predominant aspect of the clinical deterioration. In 3 patients, a devastating leptomeningeal metastasis occurred at the time of progression, leading to death.

Discussion

Cancer-associated TMA is considered a rare but fatal condition in advanced cancer. The diagnosis of TMA is based on the findings of microangiopathic hemolytic anemia and thrombocytopenia. The combination of low erythrocyte and platelet counts is a common phenomenon in cancer patients, mostly due to the hematologic toxicity of chemotherapy. However, rare and serious conditions such as TMA should be taken into account, especially if the Coombs’ test results are negative, and a thorough laboratory work-up should be undertaken to rule out the most obvious differential diagnoses. The most relevant sign of TMA in the differential blood count is the occurrence of schistocytes. However, these are not always present at the beginning of microangiopathy.

In contrast to primary TTP, in which a severe ADAMTS13 deficiency (< 5%) is the known pathophysiological mechanism [1, 7, 8], the pathophysiology of cancer-associated TMA remains largely unknown. A variety of pathophysiological considerations have been proposed, including tumor cell emboli, neoangiogenesis with fragile endothelium, mucin in mucin-producing tumors, and collision of disseminated carcinoma cells with erythrocytes. Also, injury to the endothelial cells of vessels in the bone marrow by direct tumor invasion has been discussed. We found very high vWF antigen levels in 6/7 patients, in accordance with Fontana et al. [9]. This can reflect endothelial damage, which might be of pathophysiological significance.

The role of ADAMTS13 deficiency in cancer-associated TMA remains controversial. Although a severe deficiency has been described in some cases [6, 10], in most published cases a normal or only minimally reduced activity of the protease has been found [5, 6, 9, 11, 12], which corresponds with our findings. Interestingly, there is evidence that cancer itself [13–15], as well as other severe medical conditions [16, 17], is associated with a moderate ADAMTS13 deficiency. However, in all of these cases the ADAMTS13 deficiency was mild, not reaching a rate as low as in TTP. These previous findings are in accordance with our data. It can be hypothesized that a mild ADAMTS13 deficiency is an epiphenomenon of the underlying cancer rather than a crucial pathophysiological mechanism of cancer-associated TMA. Another hint reinforcing this hypothesis is the low efficacy of plasma exchange therapy in cancer-associated TMA compared to TTP, in which the efficacy of this therapy depends on substituting the missing ADAMTS13 and/or on removing inhibitory antibodies against ADAMTS13.

It is worth noting that all our patients exhibited bone marrow infiltration, which is in concordance with the majority of published cases [4, 9–11, 18–26]. We therefore agree with the hypothesis of Chang and Naqvi [4] that bone marrow infiltration is highly associated with cancer-associated TMA. Whether bone marrow infiltration is a cause of TMA or an epiphenomenon needs further elucidation.

The optimal therapy for cancer-associated TMA is unknown. However, there is a growing body of evidence that immediate initiation of an effective antineoplastic treatment is of utmost importance. While plasma exchange therapy is the treatment of first choice for primary TTP, its benefit for cancer-associated TMA remains highly controversial [4–6, 11, 12, 19, 25–28]. It is essential that plasma exchange therapy is further evaluated. However, there is increasing data that patients with cancer-associated TMA might not benefit from this invasive and expensive therapy.

The most important treatment option appears to be an efficient therapy of the underlying neoplastic disease. In line with this, several long-term survivors have been described, all of them with a complete or partial remission of the carcinoma [4, 6, 18–20, 22–25]. Due to this growing body of evidence, we have changed the treatment algorithm in our clinic. The initiation of chemotherapy became the primary goal, and omission...
of plasma exchange therapy was tolerated or even enforced. Our results with fractionated chemotherapy are surprisingly favorable and are in contrast to previous reports [4–6, 12]. The following reasons might explain these differences:

- In contrast to others, we describe only breast cancer patients, for whom various highly efficient chemotherapy options are known. Patients with metastatic breast cancer, although with a palliative disease, have a longer overall survival than patients with many other metastatic malignancies that have been included in previous case series on cancer-associated TMA.

- In 5/8 patients the diagnosis of TMA was simultaneous to the diagnosis of distant metastases, and hence the fractionated chemotherapy induced for TMA was their first-line therapy. It is well known that efficacy is higher in the first-line setting than in the following lines.

- The fractionated chemotherapy regimens used in our patients were generally well tolerated. Therefore, dose reductions and delay of administration were uncommon. This might also explain the high effectiveness of chemotherapy in our patients.

Although the outcome of our patients was higher than previously reported, all patients inevitably experienced a progression of breast cancer leading to death. Interestingly, while in 2 patients the peripheral blood count deteriorated again, in none of our patients was the recurrence of TMA the proposed cause of death. To the best of our knowledge, an association of TMA and leptomeningeal metastasis, as seen in 3/8 of our patients, has not been described before.

In conclusion, cancer-associated TMA is a devastating complication of malignant diseases. While a severe ADAMTS13 deficiency seems not to be the underlying pathophysiological mechanism, bone marrow infiltration by carcinoma cells is strongly associated with TMA. The optimal management of the patients is not yet clear. We believe, however, that plasma exchange therapy has no or only minor therapeutic efficiency in cancer-associated TMA. In our opinion, it is crucial to induce effective antineoplastic therapy as soon as possible. Fractionated regimes seem a possible therapeutic way in order to minimize hematologic toxicity and to enable regular administration of chemotherapy.

Disclosure Statement

The authors declare no conflict of interest.

References


Breast Cancer-Associated Thrombotic Microangiopathy