ABDOMINAL HEMANGIOSARCOMA – AN UNUSUAL CASE OF ACUTE ABDOMEN.

CASE REPORT

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Abstract
In our case report and literature review we report on a rare case of acute abdomen – bleeding angiosarcoma of the stomach. Our patient was a 26-year-old woman. We present the current classification of tumours of the gastrointestinal tract, including peroperative and histological evidence.

In the discussion we present two-year results after the surgery. Conclusions: Malignant vascular tumours are an infrequent disease, but are very aggressive if occurred. Complete curative removal can be feasible in patients with localized lesions.

Keywords
Acute abdomen, Angiosarcoma, Hemangiosarcoma

INTRODUCTION
In our case report, we present one of the very infrequent but possible causes of acute abdomen – abdominal hemangiosarcoma. The patient presented is a young woman who underwent surgery in our department.

We would like to skip the general and well-known causes of acute abdomen and their classification in the present publication.

We present a classification of the gastrointestinal tumours, mainly because in the current time it is absent in the Czech literature (1, 3).
GASTROINTESTINAL TUMOURS – CURRENT CLASSIFICATION (1)

Robbins, 1999:
A: benign polyps
   - hyperplastic polyps
   - hamartoma polyps (juvenile - Peutz-Jeghers)
   - inflammatory polyps
   - lymphoid polyps

B: tumorous epithelioid lesions
   - benign: adenoma
   - malignant: adenocarcinoma, carcinoïd (can be semimalignant)

C: mesenchymal tumorous lesions
   - gastrointestinal stromal tumours (benign and malignant)
   - next: lipoma, angiomia, Kaposi’s sarcoma

D: lymphomas (more frequent than carcinomas)

Classification of vascular tumours of soft tissue (Enzinger (1))
I. Benign vascular tumours
A: Localized hemangioma
   - capillary hemangioma (including juvenile type)
   - cavernous hemangioma
   - venous hemangioma
   - arteriovenous hemangioma (racemose hemangioma)
   - epithelioid hemangioma (angiolympoid hyperplasia, Kimura’s disease)
   - hemangioma of granulation tissue type (pyogenic granuloma)
   - miscellaneous hemangiomas of deep soft tissue (synovial, intramuscular, neural)
B: Angiomatosis (diffuse hemangioma)

II. Vascular tumours of intermediate malignancy (hemangioendothelioma)
   A: Epithelioid hemangioendothelioma
   B: Spindle cell hemangioendothelioma
   C: Malignant endovascular papillary angioendothelioma

III. Malignant vascular tumours
   A: Angiosarcoma (including lymphangiosarcoma)
   B: Kaposi’s sarcoma
Hemangiosarcoma is a malignant mesenchymal tumour that has its origin in blood vessels. It develops lymphogenous and hematogenous metastases. In most cases it is localized to the liver, then (in descending order) spleen, breast, lungs, thyroid gland, and skeletal muscles. The incidence can, at least partially, be influenced by the environment - for example, liver hemangiosarcoma is a professional disease of those who work in PVC (polyvinyl chloride) production. From the point of epidemiology, it is rather a rare tumour (5,6,7,11,15) (Fig.1).

CASE

The patient presented was a 26-year-old woman, previously completely healthy. Clinical history included light abdominal pain, which worsened two days before the admission. Clinical presentation at admission was diffuse abdominal pain, without peritoneal signs or peritonism (9); large painful resistance in the left mesogastrium and hypogastrium was palpable. Other physical findings on the abdomen were negative. Blood pressure and pulse frequency were normal. Pathological laboratory findings included: hemoglobin 98 g/l, leucocytes 13.6 x 10^9/l, bilirubin 37.4 mmol/l. Other laboratory parameters were normal.

Ultrasonography verified solid mass in the epigastrium, more on the left side, and ascites. CT findings: non-homogenous mass 12x10x18 cm in the left mesogastrium and epigastrium, during the post-contrast phase some parts of the stroma changed density from 30 to 100 HU. Relations to the surrounding tissues are unclear. Gynaecology examination was negative.

Because of progressing symptomatology few hours after admission the surgery was indicated (23.10.2002). The abdominal cavity was opened via middle laparotomy. A hemoperitoneum (700 ml) was found, along with a tumour of 10x12x20 cm in size, with many blood vessels on its surface, invading the posterior wall of the stomach (Fig. 2, Fig. 3).

Due to these findings the tumour was excised completely (after dissection of the greater omentum); (Fig.4) excision of the posterior wall of the stomach with biopsy of the resection margin (peroperative cryo-cut histology was negative, definitive histology was also negative); (Fig. 5) complete revision of abdominal cavity, drainage.

To prevent thrombembolia, low-molecular heparin (LWMH) was given (higher risk of thrombembolia vs. risk of bleeding (20)). After surgery, the patient was afebrile, vegetatively stabilized, the wound was healed without any complications, per primam intentionem.

Definitive histology showed that it was a “low-grade” angiosarcoma of the abdominal cavity (M – 9120/31), invading gastric wall through serosa to the smooth muscle layer (Fig. 6, Fig. 7).

There were some doubts about this diagnosis due to the partly controversial immunohistochemistry results (negative factor VIII, CD34 and thrombomodulin); therefore differential diagnostics should also include the possibility of epithelioid leiomyosarcoma or gastrointestinal stromal tumour (Fig. 8).

Post-operative adjuvant therapy included four series of chemotherapy (Adrimycin + Isososfamid + Mesna) with good tolerance. So far (by 30.5.2004), the patient is completely without any subjective problems and without signs of generalization or relapse - ultrasonography of the abdominal cavity is negative, CT scans of lungs, mediastinimum, abdomen and pelvis minor are also negative, oncological markers (NSE) are negative, vaginal ultrasonography is negative, gastrofibroscopy and gastric endosonography are negative; she has no weight loss, either.
Fig. 1
Age distribution of patients with sarcomas of soft tissues (AFIP, 1966–1976) (89 cases) (11)

Fig. 2
Abdominal angiosarcoma – exclusion of the tumour after middle laparotomy
Fig. 3
Visible blood vessels on the tumour surface and the direction of invasion (into the posterior gastric wall)

Fig. 4
Tumour excision after the greater omentum was dissected; signs of malignant angiogenesis are visible.
Fig. 5
Detail of the site where the tumour invaded into the posterior gastric wall (after resection and suture)

Fig. 6
Cavernous vessel spaces are filled with groups of erythrocytes. Between them there are tumour cells with a pink cytoplasm and purple nuclei.
Fig. 7
Tumour tissue with a big amount of blood vessels

Fig. 8
Immunohistochemistry for the positivity of vimentin, which is a component that proves the mesenchymal origin of the tumour. The brownish areas of the cytoplasm mean positiveness.
DISCUSSION

Hemangiosarcoma is a malignant mesenchymal tumour that has its origin in blood vessels. It develops lymphogenous and hematogenous metastases. In most cases it is localized to the liver, then (in descending order) to the spleen, breast, lungs, thyroid gland, and skeletal muscles. Liver hemangiosarcoma is also a professional disease of those who work in PVC production. From the point of epidemiology, it is rather a rare tumour. Even less frequently, hemangiosarcoma is a cause of intraabdominal bleeding (12). In some very rare cases, malignant mesenchymal tumours of the retroperitoneum (4) or pelvis minor (22) are found, which spontaneously perforate, causing bleeding into the abdominal cavity.

Analysis of the current literature shows that many surgical departments had similar experience with angiosarcomas (mostly in the abdominal cavity) (12, 15–19, 21). What is important is that the conclusions were also similar – angiosarcomas and vascular malignant tumours as a whole are very rare but, if occurred, have also a very aggressive growth pattern and symptomatology, including the tendency to bleeding (12, 15, 17, 19). A positive fact is that if the tumour is localized, its complete resection is often curable (16, 18). Our case supports this practice (though we should note that our patient had a “low-grade” angiosarcoma).

CONCLUSIONS

Therapy of hemangiosarcomas is very difficult, as reported in the literature (8, 11, 13). A multidisciplinary approach should be always used, as the therapy is multimodal: surgery + adjuvant chemotherapy + radiotherapy (in some exceptional cases). Exact diagnosis (especially immunohistochemistry) is very important (13, 14).

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HEMANGIOSARKOM DUTINY BŘIŠNÍ – VZÁCNÁ PŘÍČINA NÁHLÉ PŘÍHODY BŘIŠNÍ

S o u h r n

REFERENCES
