

Levels of CEA and Ca 19 - 9 in the sera and peritoneal cavity in patients with gastric and pancreatic cancers¹

Níveis de CEA e Ca 19 - 9 em soro e cavidade peritoneal em pacientes com câncer do estômago e pâncreas

David Hoskovec^I, Jozef Varga^{II}, Ellen Konečná^{III}, František Antoš^{IV}

^IMD, Fellow PhD degree, Postgraduate Program of Experimental Surgery, Charles University Prague, Czech Republic. Assistant Professor of Surgery, 1st Medical Faculty Czech Rep. Main author; study concept and design; acquisition, analysis and interpretation of data; drafting and revision of the manuscript; intellectual content; study supervision; responsible for English language.

^{II}MD, Department of Surgery, 1st Medical Faculty Charles University, Czech Republic. Assistant Professor of Surgery. Concept, design, intellectual content, acquisition of data and critical revision.

^{III}MD, Fellow PhD degree, Postgraduate Program of Experimental Surgery, Charles University Prague, Czech Republic. Assistant Professor of Surgery, 1st Medical Faculty Czech Rep. Concept, design, intellectual content, acquisition of data and critical revision.

^{IV}Professor of Surgery, Charles University Prague, Czech Republic. MD, PhD, former Head of Department of Surgery, Hospital Bulovka, Prague, Czech Republic, Consultant and Supervisor of PhD programs. Concept, design, intellectual content, drafting and revision of the manuscript, responsible for English language.

ABSTRACT

PURPOSE: Tumor markers are substances found in blood and other biological fluids if tumor is present in the body. They can be produced by tumor itself or can be results of cancer – body relation. They may be used in the follow-up of cancer patients to identify tumor recurrence. Pre-treatment levels have prognostic tool and could signalize persistence of minimal residual disease despite radical surgery.

METHODS: We operated on 52 patients with upper GI malignancy (32 with gastric cancer and 20 with pancreatic cancer). Blood samples were taken before surgery and peritoneal samples immediately after laparotomy before any manipulation with tumor. All samples were examined by standard biochemical technique and the level was compared with a stage of the disease.

RESULTS: Patients suffering from gastric carcinoma of stage I and II had higher level of both markers in sera then in the peritoneal cavity, however most of them were within physiological range. Patients in stage III and IV had average marker levels in the peritoneal cavity higher than in sera. Number of positive findings was increasing according to the stage of the disease. The peritoneal levels of both markers varied extremely in higher stages. In patients suffering from pancreatic carcinoma the CEA levels both in sera and peritoneal cavity were parallel but peritoneal levels were slightly higher in stages III and IV. Ca 19 – 9 was more sensitive for pancreatic cancer. The percentage of positive findings was higher in sera but the level of Ca 19 – 9 was higher in the peritoneal cavity. The number of positive findings again correlated with the stage of the disease.

CONCLUSIONS: Levels of tumor markers in sera could signalize inoperability of tumor (Ca 19 – 9 in cases of pancreatic carcinoma); peritoneal levels could predict R1 resection especially in gastric cancer patients and risk of early peritoneal recurrence of the disease. Difference between the levels in the peritoneum and sera may signalize the route of dissemination (hematogenous and intraperitoneal).

Key words: Stomach Neoplasms. Pancreatic Neoplasms. Tumor Markers, Biological. Neoplasm, Residual.

RESUMO

OBJETIVO: Os marcadores tumorais são substâncias encontradas no sangue e outros fluidos biológicos em pacientes com doenças oncológicas. São produzidos pelo próprio tumor ou ser resultado da interação entre o tumor e o organismo. Podem ser usados no seguimento de pacientes com câncer para identificar recidiva tumoral. Os níveis pré-tratamento têm valor prognóstico e podem sinalizar persistência de doença residual mínima após cirurgia radical.

MÉTODOS: Foram operados 52 pacientes com tumores do trato gastrointestinal superior (32 com câncer do estômago e 20 do

apoptosis in physiologic state, so it is expressed in normal mucosal cells and over-expressed in adenocarcinoma (colorectal, gastric, pancreatic, breast, lung and others)^{8,10-13}.

Cancer antigen 19-9

Ca 19 – 9 is an intracellular adhesion molecule. The marker is 210 KD tumor associated glycoprotein antigen present as carbohydrate determinant on glycolipid and glycoprotein. Since the reference to antibody detection for Ca 19 – 9 published in 1979, it has been found to be the most useful tumor marker for pancreatic adenocarcinoma and gall bladder carcinoma too. Patients who are negative for Lewis antigen (a–, b–) do not synthesize Ca 19 – 9, and this constitutes 4 to 15 % of the population^{8,11-13}.

Methods

We examined 52 patients with malignant disease (32 with gastric cancer and 20 with pancreatic cancer). There were 19 men and 11 women in group with gastric cancer and 12 men and eight women in group with pancreatic cancer. The average age of the patients in the group with gastric cancer was 67 years (range 37 – 87 years), in group with pancreatic cancer was 63 years (range 48 – 79 years) (Table 1).

TABLE 1 - Patients according the disease and stage.

Stage	I	II	III	IV	Total
C 16	5	4	8	15	32
C 25	2	4	2	12	20

Comparison of the levels of CEA and Ca 19 – 9 in sera and peritoneal lavage in patients suffering from gastric and pancreatic carcinoma was performed. Blood samples were collected after diagnosis of the disease. Peritoneal samples were taken immediately after laparotomy. When ascites was present, 20 ml were sent to biochemistry laboratory. In cases without ascites the peritoneal cavity was lavaged with 100 ml saline solution and the samples were collected 5 minutes later. Levels of the CEA and Ca 19 – 9 were examined. Physiological range was the same for peritoneal fluid and sera. CEA ranges were 0 – 4, 6 µg/l, Ca 19 – 9 levels were normal within 0 – 37 U/l.

Patients were stratified according to the stage of the disease.

Results

Tumor markers sensitivity and specificity (Tables 2 and 3)

Sensitivity and specificity for microscopic intraperitoneal cancer dissemination were evaluated in group of patient without confirmation of the malignant tumors and in patients with serosal infiltration or serosal metastases (we expected 100% prevalence of tumor cells in the peritoneal fluid in these patients and 100% positivity of the tumor markers).

TABLE 2 - Tumor markers examination in group of benign disease.

	Benign disease (No)
CEA within normal ranges	5
CEA elevation	0
Ca19-9 within normal ranges	5
Ca19-9 elevation	0

TABLE 3 - Tumor markers in cases of peritoneal dissemination.

	Peritoneal infiltration		Total
	Local (Gastric wall)	Distant	
CEA within normal ranges	21	9	30
CEA elevation	2	16	18
Ca19-9 within normal ranges	18	10	28
Ca19-9 elevation	5	15	20

CEA sensitivity was 38% and Ca 19-9 42%, respectively. Combination of both markers achieved 53% (we repeatedly found elevation only ones of both markers). Specificity of the biochemistry examination was 100%.

Gastric cancer

CEA levels in sera were within normal ranges in nearly all patients in stages I and II (with only one exception in patients with chronic kidney failure) and in nearly 70% of patients in stage III and IV. When the elevation was present it was not any extremely high value. Peritoneal levels of CEA were normal in all cases of stages I and II. Patients in stage III displayed normal CEA levels in about 65%. More positive finding in the peritoneal cavity was in stage IV (9:6). The variance of the levels in stage IV

was high (0 - 5500 µg/l) (Figure 1).

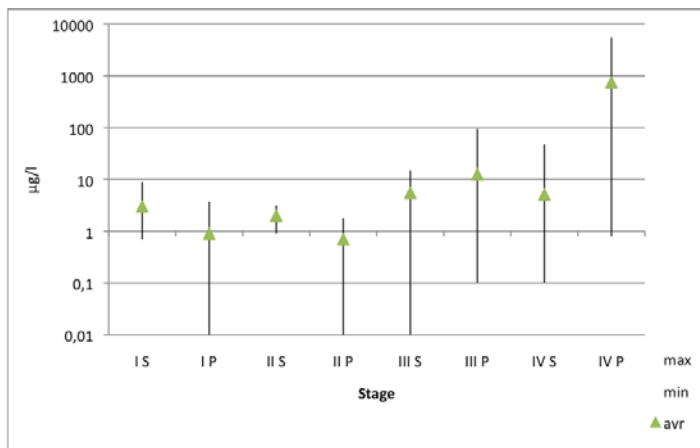


FIGURE 1 - Gastric cancer – CEA levels in sera (S) and peritoneal cavity (P) according to stage of the disease.

Ca 19 – 9 levels in sera and in peritoneum was nearly the same as CEA in stage I and II – except for one patient the levels were within the normal ranges. Elevation of sera Ca 19 – 9 in stage III occurred in 25 % and in stage IV in 50% of patients. Peritoneal positivity was higher. 25% of patients in stage III and nearly 70 % in stage IV were positive. Value of Ca 19 – 9 levels varied extremely in stage IV (0 – 101 066 U/l) (Figure 2).

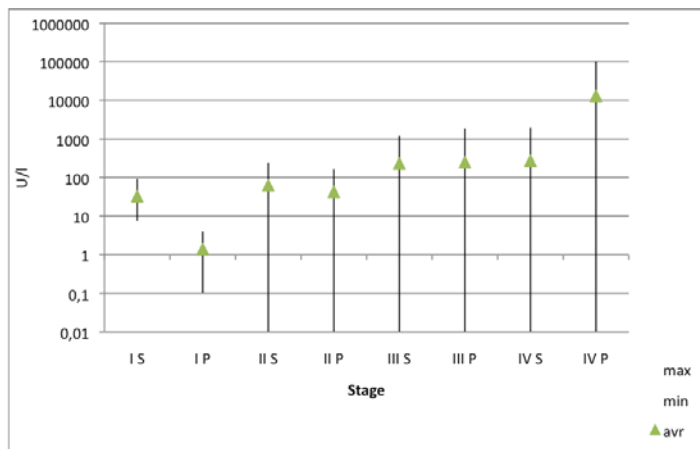


FIGURE 2 - Gastric cancer – Ca 19 – 9 levels in sera (S) and peritoneal cavity (P) according to stage of the disease.

The positivity of at least one tumor markers was increasing according to the stage of the disease (Figure 3).

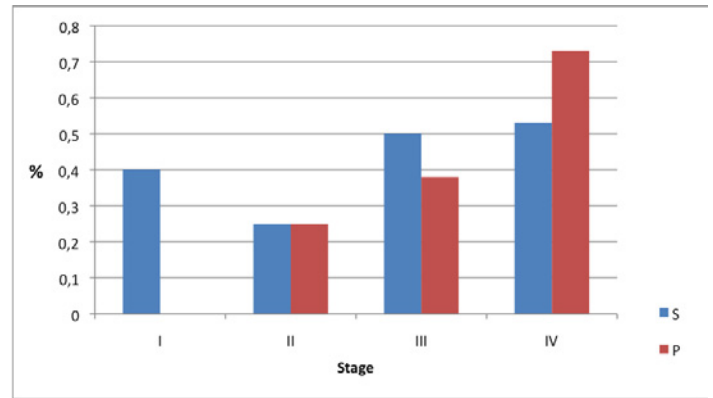


FIGURE 3 - Gastric cancer – Elevation of the tumor markers according to stage of the disease (S – serum, P – peritoneal cavity).

To sum-up, the percentage of negative findings of both markers in sera and peritoneal cavity was decreasing according to the stage of the disease. Positivity of both markers was increasing reversely and Ca 19 – 9 was probably more sensitive than CEA (Figure 4).

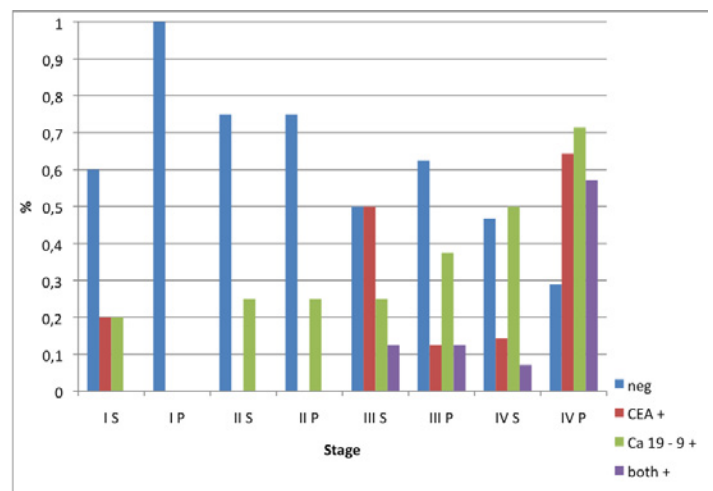


FIGURE 4 - Gastric cancer – Summary of the results in sera (S) a peritoneum (P) according to stage of the disease.

Pancreatic carcinoma

Patients with pancreatic carcinoma had normal levels of CEA in stages I and II (except for one patient). Positivity in stage III and IV is much higher. 20 % of patients had elevated levels of CEA in sera and 43 % displayed elevation in the peritoneal cavity (Figure 5).

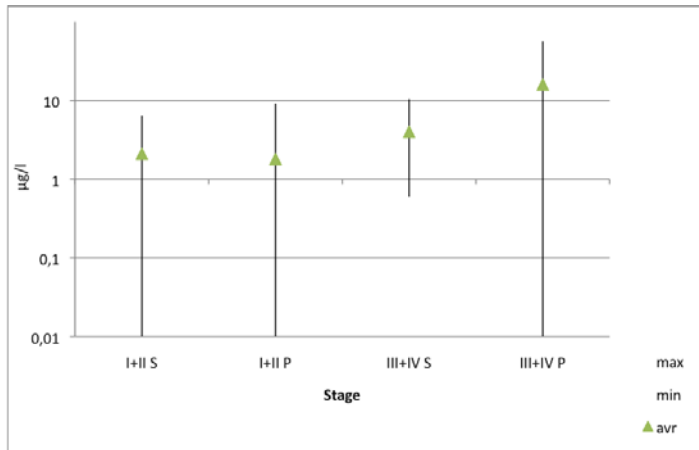


FIGURE 5 - Pancreatic cancer – CEA levels in sera (S) and peritoneal cavity (P) according to stage of the disease.

Ca 19 – 9 was more sensitive in cases of pancreatic cancer. Elevation in sera was found in 66 % of cases in stage I and II and in the peritoneal cavity it was in 33%. In stages III and IV the findings of Ca 19 – 9 were positive in 85% in sera and in 60% in the peritoneal cavity (Figure 6).

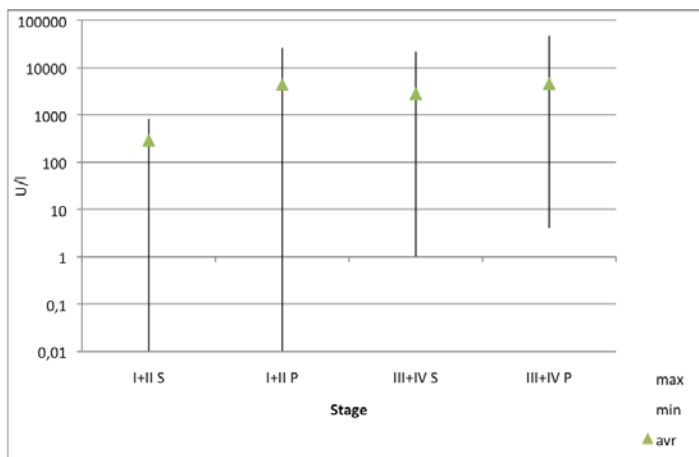


FIGURE 6 - Pancreatic cancer – Ca 19 – 9 levels in sera (S) and peritoneal cavity (P) according to stage of the disease

Similarly as in gastric cancer patients, the positive finding of at least one tumor marker is higher with the higher stage of the disease (Figure 7).

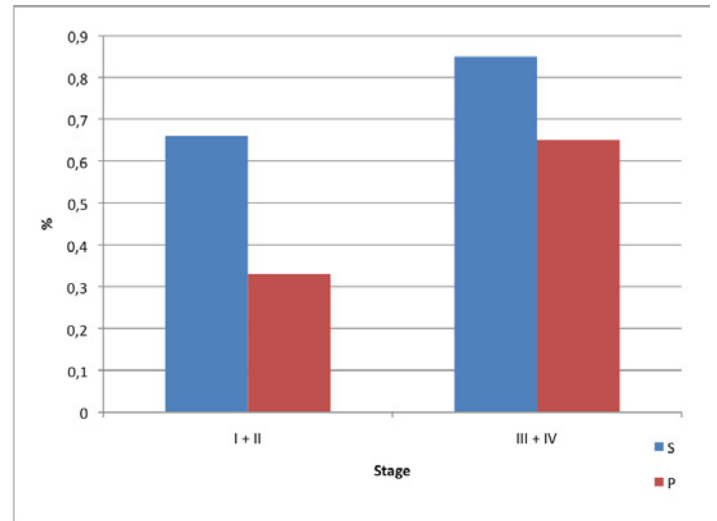


FIGURE 7 - Pancreatic cancer – Elevation of the tumor markers according to stage of the disease (S – serum, P – peritoneal cavity).

In summary, physiological ranges of both markers were less frequent than in cases of gastric carcinoma. Only 30% of patients had negative finding in sera in stages I and II. Negativity of both markers in higher stages was about 15%. The intraperitoneal finding was similar, but there were more negative patients in stages I and II (Figure 8).

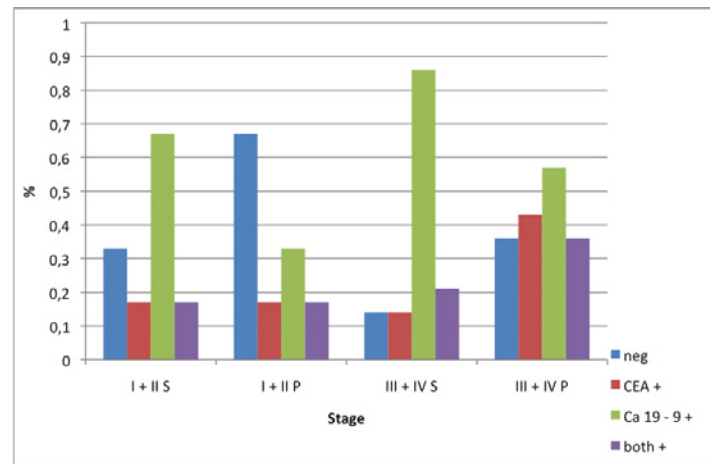


FIGURE 8 - Pancreatic cancer – Summary of the results in sera (S) a peritoneum (P) according to stage of the disease.

Discussion

There are few publications focusing on comparison of levels of various tumor markers in sera and in peritoneal cavity. All demonstrate that higher level of tumor markers is associated with advanced disease, poor prognosis and early recurrence. In cases of gastric cancer peritoneal recurrence is one of the most important reasons of the treatment failure. Examination of peritoneal fluid by cytology, biochemistry or RT PCR could predict this relapse

of the disease. The cytological examination is widely used in East Asian countries for diagnosis of microscopic peritoneal dissemination which is undetectable by other diagnostic tools in time of laparotomy or laparoscopy. Patients with positive cytology are classified as advanced disease with distant metastasis. All reports confirm worsening of the patient's prognosis with positive cytology despite various detection rates¹⁴⁻¹⁸.

Positive peritoneal cytology could predict the site of recurrence. It is an independent prognostic factor of the intraperitoneal dissemination (sensitivity 56%, specificity 97%)¹⁹.

However, the sensitivity and specificity of cytology is very high in Asian countries, nonetheless in the western countries the results are worse. In our previous study we published that cytology had sensitivity only 34% and specificity 80%. Biochemistry for combination of CEA and Ca 19 – 9 proved sensitivity of 53% and specificity 100%²⁰. Similar data were published by other western centres. Our data show that gastric cancers with infiltration of the serosal layer or beyond associate with increasing frequency of positivity of tumor markers. CEA and Ca 19 – 9 levels can show the amount of tumor cells in the peritoneal cavity. Early gastric carcinoma may reveal increased temporal marker in the peritoneal washing, suggesting the presence of penetration pathways in the stomach wall by cancer cells, without invasion through continuation into the serosa^{1,21}. Lymphatic channels or metastases in lymph nodes are examples of these possible pathways. This notion is partially supported by the fact that peritoneal recurrence takes place in around 1% of patients with early gastric carcinoma¹⁵.

In cases of pancreatic cancer free tumor cells in the peritoneal fluid predict unresectable, aggressive disease with early metastatic seeding and short survival time^{22,23}. Positive peritoneal cytology is classified as metastatic disease according to the 6th edition of the Cancer staging manual AJCC³. It is presumed that positive cytology indicates ventral spreading of the tumor and it is a sign of tumor penetration through the peritoneum of the omental burse²¹. Free cancer cells in the peritoneal cavity are the only sign of generalization in 12 – 30% patients^{3,22,24}. Positive cytology is more often found in tumors larger than 4 cm and tumors of the pancreatic body and tail²³⁻²⁵. Free tumor cells are diagnosed more often after fine needle biopsy (17: 28%)²². Similar to our previous published study, here again the biochemistry of peritoneal fluid is more sensitive to the presence of cancer cells in peritoneal cavity and could signal more reliably the advanced stages of the disease than conventional examinations (CT, EUS). We found lymph nodes metastases in all cases in stage II with elevation of the peritoneal Ca 19 – 9 lymph nodes metastasis.

Conclusions

Elevated levels of tumor markers both in peritoneal cavity and sera could signalize advanced stage of the disease than it is established by CT and EUS. Patients with a higher level of the markers in sera are usually treated as high risk patients for treatment failure. But the same signalize higher level of tumor markers in the peritoneal fluid. These patients are in risk of peritoneal and locoregional recurrence and further treatment should be focused to prevent this recurrence. Unfortunately there is no satisfactory procedure how to prevent and treat peritoneal cancer dissemination caused by gastrointestinal cancers. According to contemporary knowledge the best possibility now is probably the combination of surgery (peritonectomy) and some type of chemotherapy.

References

1. Aiko T, Sasako M. The new Japanese classification of gastric carcinoma: points to be revised. *Gastric Cancer*. 1998;1:25-30.
2. Wells Jr SA. Gastric cancer. *Curr Probl Surg*. 2006;43:558-669.
3. Stefanidis D, Grove KD, Schwesinger WH, Thomas CR Jr. The current role of staging laparoscopy for adenocarcinoma of the pancreas: a review. *Ann Oncol*. 2006;17:189-99.
4. Goggins M, Koopmann J, Yang D, Canto MI, Hruban R. National Academy of Clinical Biochemistry (NACB) Guidelines for the Use of Tumor Markers in Pancreatic Ductal Adenocarcinoma, Available from http://www.aacc.org/SiteCollectionDocuments/NACB/LMPG/tumor/chp3i_pancreatic.pdf.
5. Liu RC, Traverso LW. Laparoscopic staging should be used routinely for locally extensive cancer of the pancreatic head. *J Gastrointest Surg*. 2004;8:923-4.
6. Greene FL, Fleming ID, Fritz A, Balch ChM, Haller DG, Morrow M. *AJCC Cancer Staging Manual*. 6ed. Available from www.cancerstaging.org.
7. Li JK, Zheng M, Miao CW, Zhang JH, Ding GH, Wu WS. Peritoneal lavage cytology and carcinoembryonic antigen determination in predicting peritoneal metastasis and prognosis of gastric cancer. *World J Gastroenterol*. 2005;46:7374-7.
8. Malati T. Tumour markers: an overview. *Ind J Clin Biochem*. 2007;2:17-31.
9. Urban D, Catane R. Serum tumor markers in oncology. *Isr Med Assoc J*. 2009;11:103-4.
10. Bonfrer JMG, Louhimo L. National Academy of Clinical Biochemistry Guidelines for the Use of Tumor Markers in Gastric Cancer, http://www.aacc.org/SiteCollectionDocuments/NACB/LMPG/tumor/chp3g_gastric.pdf.
11. Perkins GL, Evan D, Slater D, Sanders GK, Prichard JG. Serum tumor markers. *Am Fam Physician*. 2003;68:1075-82.
12. Scientific Committee of the Association of Clinical Biochemists in Ireland (ACBI) Guidelines for the Use of Tumour Markers, <http://www.acbi.ie/Downloads/Guidelines-for-the-Use-of-Tumour-Markers-2005.pdf>. accessed Juni 15, 2011.
13. Schlieman MG, Hung H, Bold RJ. Utility of tumor markers in determining resectability of pancreatic cancer. *Arch Surg*. 2003;138:951-6.
14. Bold JR, Ota MD, Ajani JA, Mansfield PF. Peritoneal and serum tumor markers predict recurrence and survival of patients with resectable gastric cancer. *Gastric Cancer*. 1999;2:1-7.

15. Crepaldi-Filho R, Palma RT, Marcelo Franchini Giusti MF, de Assis Galveo Bueno M, da Silva PSL, Waisberg J. Levels of carcinoembryonic antigen and CA 19-9 in the sera and peritoneal washing of patients undergoing surgical treatment for gastric carcinoma. *Arq Gastroenterol.* 2008;45:219-24.
16. Majima T, Ichikura T, Mochizuki H. Prognostic significance of the cytologic features of free cancer cells in the peritoneal cavity of patients with gastric cancer. *Surg Today.* 2002;32:35-9.
17. Manzoni G, Verlato G, Di Leo A, Tomezzoli A, Pedrazzani C, Pasini F, Piubello Q, Cordiano C. Peritoneal cytology does not increase the prognostic information provided by TNM in gastric cancer. *World J Surg.* 2006;30:579-84.
18. Ribeiro U Jr, Gama-Rodrigues JJ, Safatle-Ribeiro AV, Bitelman B, Ibrahim RE, Ferreira MB, Laudanna AA, Pinotti HW. Prognostic significance of intraperitoneal free cancer cells obtained by laparoscopic peritoneal lavage in patients with gastric cancer. *J Gastrointest Surg.* 1998;2:244-9.
19. Bando E, Yonemura Y, Takeshita Y, Taniguchi K, Yasui T, Yoshimitsu Y, Fushida S, Fujimura T, Nishimura G, Miwa K. Intraoperative lavage for cytological examination in 1297 patients with gastric carcinoma. *Am J Surg.* 1999;178:256-62.
20. Hoskovec D, Varga J, Antoš F, Dytrych P, Benková K. Peritoneal lavage cytology and biochemistry in cases of gastric and biliopancreatic tumors. *Gastroent hepatol.* 2008;62(Suppl 3):15. Available from www.medvik.cz/link/bmc09002002
21. Hermanek P. Pathology and biology of pancreatic ductal adenocarcinoma. *Langenbeck's Arch Surg.* 1998;383:116-20.
22. Jimenez RE, Warshaw AL, Fernandez-del Castillo C. Laparoscopy and peritoneal cytology in the staging of pancreatic cancer. *J Hepatobiliary Pancreat Surg.* 2000;7:15-20.
23. Jimenez RE, Warshaw AL, Ratner DW, Willett CG, McGrath D, Fernandez-del Castillo C. Impact of laparoscopic staging in the treatment of pancreatic cancer. *Arch Surg.* 2000;135:409-15.
24. Warshaw AL. Implications of peritoneal cytology for staging of early pancreatic cancer. *Am J Surg.* 1991;161:26-30.
25. Traverso LW. Pancreatic cancer: surgery alone is not sufficient. *Surg Endosc.* 2006;20:446-

Correspondence:

David Hoskovec, MD
1st Department of Surgery
General University Hospital
U nemocnice 2
12000 Prague 2
Czech Republic
Phone/Fax: (00 420)224963377
david.hoskovec@vfn.cz

Received: January 18, 2012

Review: March 14, 2012

Accepted: April 16, 2012

Conflict of interest: none

Financial source: none

¹Research performed at Department of Surgery, University Hospital Bulovka, Charles University Prague, Czech Republic.