Impact of Nutritional Support in Patients with Gastrointestinal Malignancies - A Review

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Abstract

Cancer Cachexia-Anorexia Syndrome (CACS) is a common and often underdiagnosed syndrome in cancer population. If undiagnosed, this initially reversible syndrome leads to deterioration and is direct cause of death in 20% of cancer patients. Oppositely, with timely diagnosis, nutritional counseling can help to slow the progression and positively influence on quality of life, tolerance to chemotheraphy with ultimate goal of prolonging patient’s life. Colorectal and pancreatic cancers are very common tumours type worldwide. The prognosis for the survival in pancreatic cancer is poor as in colorectal after disease progression. Cancer anorexia-cachexia syndrome is highly prevalent among patients with colorectal and pancreatic cancer, and has a large impact on morbidity and mortality, and on patient quality of life. The etiology of primary CACS appears to be related to the pathological loss of inhibitory control of catabolic pathways, whose increased activities are not counterbalanced by the increased central and peripheral anabolic drive. Secondary CACS (related to gastrointestinal obstruction, vomiting due to chemotheraphy etc.) is contributing to bad patient’s condition. As a result of being complex and influencing a great number of metabolic pathways, cancer cachexia can be treated in multimodal manner. In this review we are presenting most promising targets and current opinions in ways to treat cachexia and our results with nutritional supplementation in colorectal and pancreatic cancer patients.

Keywords: Cancer Cachexia-Anorexia Syndrome (CACS); Proinflammatory cytokines; Colorectal cancer; Pancreatic cancer; Nutritional supplementation

Introduction

Cachexia is a clinical syndrome of a distressing and debilitating condition, affecting significant numbers of patients with advanced malignant disease and causing huge distress; it is the primary cause of death in about 20% of all patients with cancer. Although cachexia is most commonly associated with particular tumours, such as head and neck, gastrointestinal tract, pancreas, central nervous system and lungs, it may affect any patient with any tumour at any site [1].

Cancer anorexia-cachexia syndrome (CACS) [2,3], usually consists of a combination of anorexia, tissue wasting, malnutrition, weight loss and loss of compensatory increase in feeding. Anorexia represents the result of a failure of the usual appetite signals and is preceding cachexia (pre-cachexia). Biochemistry tests in cachetic patients are expressing anaemia, hypoalbuminemia, hypertriglyceridemia and glucose intolerance due to insulin resistance.

Primary cachexia represents the result of a complex interaction between cancer growth and host response, and is associated with a poor response to chemotheraphy with an increase in drug-related toxicity [4].

Other debilitating conditions which decrease food intake (gastrointestinal obstruction, nausea and vomiting due to chemo and radiotherapy, pain, emotional factors, renal impairment etc.) are causing secondary cachexia.

Weight loss can be unrelated to reduced nutritional intake and is mostly caused by elevated resting energy expenditures (REE) and proinflammatory cytokine expression [5].

One important mechanism is the activation of the acute phase response cascade [4]. Kemik et al. found significantly higher serum CRP, IL-1α, IL-1β, IL-6, IL-8, IL-10, TNF-α, VEGF-A, VEGF-C, VEGFR1, and leptin concentrations in patients with esophageal, gastric, pancreas, colon, and rectum cancers than controls and lower levels of the serum albumin, midkine, adiponectin, and ghrelin in patients with esophageal, gastric, pancreas, colon, and rectum cancers compared to control subjects [6].

Proinflammatory responses are stimulating the expression and release of leptin. This hormone, secreted by adipose tissue, plays a crucial role in the homeostasis of body weight as its high levels in the brain decrease the activity of the hypothalamic orexigenic mediators (ghrelin, neuropeptide Y, agoutin, orexin, melanocortin-releasing hormone) and increase anorexigenic signals (choleystatin, glucagon-like peptide, pro-opiomelanocortin, thyroid-releasing hormone, corticotropin-releasing hormone, oxytocin). Leptin levels regulate rest energy expenditure (REE) [7].

A range of novel drug targets relevant to the treatment of cachexia are discovered. Interventions may be either upstream (e.g. by antagonizing key mediators of systemic inflammation) or downstream (e.g. by blocking catabolic pathways or stimulating anabolic pathways in skeletal muscle). Upstream targets have the advantage of affecting multiple aspects of cachexia. For example, interleukin-6 is known to be the main mediator of the hepatic acute phase response in humans, but it may also play a role in anorexia, fatigue, anaemia, oedema and...
muscle loss. By contrast, myostatin acts as a physiological brake to the continued growth of skeletal muscle and is therefore a potential downstream target. Alternative targets include the melanocortin pathway in the CNS control of appetite. Therapies based on these pathways are currently in Phase I and Phase II clinical trials in cancer patients [8].

Progestogens, particularly megestrol acetate, are commonly used to treat CACS. The mechanism of action of megestrol is believed to involve the stimulation of appetite by both direct (neuropeptide Y) and indirect pathways.

Omega-3 polyunsaturated fatty acids have been shown to modulate levels of proinflammatory cytokines, hepatic acute-phase proteins, eicosanoids, and tumor-derived factors in animal models of cancer and may reverse some aspects of the process of cachexia. The metabolites of EPA and DHA have less inflammatory and immunosuppressant potency than the substances derived from arachidonic acid. The competitive metabolism of EPA and DHA with arachidonic acid is occurring at the cyclooxygenase and 5-lipoxygenase levels [7]. Oral EPA has been found to stabilize the body weight of cancer patients and together with an energy- and protein-rich nutritional supplement, can enhance weight gain by increasing lean body mass.

Some other drugs also showed positive influence in CACS. Thalidomide, which is an inhibitor of tumor necrosis factor-alpha synthesis, may represent a rational therapeutic approach [9] also as inexpensive oral supplementation of L-Carnitin [10].

In postoperative setting, early nutritional support was shown to reduce the incidence of complications and to shorten the hospital stay. For patients with functional bowel, early enteral nutrition (EEN) is method of choice, and total parenteral nutrition (TPN) is reserved for highly selected cases [11,12]. If patient’s condition cannot provide normal feeding (but gastrointestinal tract is functional) or anticipated time for feeding is longer than 4 weeks, trans nasal way can be used (nasogastric, nasoduodenal and nasojejunal tubes). Percutaneous endoscopic gastrostomy tubes or percutaneous endoscopic jejunostomy may be placed (endoscopic or surgical way) if longer period of feeding is anticipated.

Pancreatic and colorectal cancer are most common gastrointestinal malignancies. Pancreatic cancer (PCa) is a very aggressive, invasive cancer whose prognosis remains very poor and represents the fourth leading cause of cancer-related mortality. Only 5% of patients is living longer than 5 years. There are approximately 277,000 new cases of pancreatic cancer and 266,000 deaths from pancreatic cancer annually in Europe, indicating a mortality rate of 96% of the cases diagnosed. Etiology of pancreatic carcinoma remains largely unknown but consistent evidence of a positive association was found for family history and cigarette smoking. Also, some studies showed a positive association with diabetes mellitus and chronic pancreatitis.

Cancers of the colon and rectum (CRC) are the third most common forms of cancer worldwide. In the developed countries CRC is the significant cause of morbidity and mortality (412 000 new diagnosed patients in Europe every year). The overall five-year survival for colon cancer is varied from 43% (Europe) to 62% (USA) [13-20].

There are four risk factor categories for CRC: epidemiological, intestinal, dietetic, and mixed. CRC is a disease in which genetic mutations of somatic cells are the molecular base of the disease. About 25-30% of CRC are diagnosed in the advanced stage and another 30% of patient will develop metastatic or locally advanced disease in next three years. Despite advances in therapeutic methods, the five-year survival rate for advanced disease is still poor (15%).

For patients with advanced stage of CRC and PCa, the therapeutic goal is quality of life (QoL). Early intervention with nutritional supplementation has been shown to halt malnutrition, reduce the consequences of CACS, extend patient survival and improve quality of life.

Aim of our Studies

In our studies we tried to assess the influence of nutritional support (counseling, enteral supplementation liquids, megestrol acetate) on nutritional status and symptoms prevalence in patients with pancreatic and colorectal cancer during chemotherapy.

Methods and Patients

During our routine clinical work nutritional status of our patients is evaluating according to changes in body weight. Body Mass Index (BMI) is calculated for all patients (on chemo/biotherapy procedures or achieving nutritional support) using the standard procedure of dividing weight in kg by height in m². We also use following questionnaires for evaluation of patient’s nutritional status: Nottingham Screening Tool Score (NST score 0-7) (Table 1), Appetite Loss Scale (0-10; where 0 is no appetite at all, and 10 is the best possible appetite), and for Performance Status or Eastern Cooperative Oncology Group (ECOG) (Table 2) or Karnofsky Performance Status (KPS) (Table 3).

Nutritional and pharmacological support is consisting of nutritional counseling, and if according to NST patients are in nutritive risk, administration of 10 ml (400 mg) per day Megostat (megestrol acetate) and enteral nutrition supplements with commercially available products.

Nutritional counseling include interviews with a physician, with purpose of learning how to prepare and ingest food during chemotherapy and to change eventually bad eating habits.

We analysed impact of nutritional support in patients with histologically confirmed pancreatic cancer (PCa) patients during 18 months period (from 1st July 2005 to 31st December 2006 at Gastroenterology department, Clinical Hospital Centre Rijeka). We followed up 44 patients with pancreatic cancer – 26 males (mean age 69 years ± 2.4 years) and 18 females (mean age 63 ± 3.2 years). All patients were with metastatic or locally advanced disease curing with standard
The effect was most pronounced 4-6 weeks after beginning the supportive pharmacological support PCa patients stopped to lose weight. This to the oncology unit.

Patients were counseled and re-evaluated by the same physician before surgery if performed or one week before chemotherapy initiation.

As side effects patients in Group I comparing with patients in Group II expressed more diarrhea 17.5% vs. 13% of patients and more edemas 13% of patients and more cachexia 13% vs. 12% of patients. Edemas, the main side effect of megestrol acetate was experienced in 29.2% of patients.

In all patients we used the same procedure for establish nutritional status as already described above. Nutritional and pharmacological support consisted of nutritional counseling, administration of 10 ml (400 mg) per day Megostal® (megestrol acetate) and enteral nutrition supplements with commercially available products Ensure® (400 mL daily with 600 kcal) and Prosure® product containing eicosapentaenoic acid (EPA) (480 ml daily, containing 2.2 g of EPA and 600 kcal). In the study with CRC patients at initial visit, upon evaluating the risk measurements according to BMI, decrease in weight, and NST, we did not find any significant differences between the prospective and retrospective group. After completion of chemotherapy and nutritional support cycle, comparing these two groups we noticed weight gain in those with a BMI <20 who received counseling and nutrition and an opposite effect was observed in the group without nutritional support. After 4 weeks of supplementation in Group I, 73.34% patients had an increase in body weight, with an average weight gain of 1.5 kg (0.6-3.3 kg) versus Group II where increase in weight gain was observed in only 19.49% of patients. Patients who achieved nutritional supplementation also expressed appetite improvement from 3.3 to 4.6 on Appetite Loss Scale. On week 12 there was a significantly smaller proportion of patients with BMI<20 and NST ≥ 5 in the group with nutritional counseling. A greater proportion of patients in the same group had a better appetite according to Appetite Loss Scale (Table 5).

As side effects patients in Group I comparing with patients in Group II expressed more diarrhea 17.5% vs.13% of patients and more abdominal pain in 15.9% vs. 12% of patients. Edemas, the main side effect of megestrol acetate was experienced in 29.2% of patients.

We also analyzed survival in the two groups and we determine that patients with nutritional support had a significantly longer median survival than patients in the control group (19.1 vs. 12.4 months).

**Discussion**

At the time of diagnosis patients with PCa mostly have disseminated disease; 20% of patients who does not have macroscopic disseminated disease have positive intra-abdominal lymph nodes when using Polymerase Chain Reaction (PCR) method. Therefore, despite the clear advances in surgical treatment of pancreatic carcinoma, 5 years survival rates are still low, ranging from 5-30%. Early metastasizing and
angiogenesis in gastroesophageal cancers and decreased survival [24].

Acute-phase response proteins are related to cachexia, accelerated cytokine (LIF) which significantly induced cell proliferation [17].

Enhanced the expression of Leukemia inhibitory factor-pleiotropic in the organism, not all of which are known until now [23].

institute treatment [12,22].

[12]. It is essential to have a validated and universally accepted [12]. Therefore, cachexia is infrequently identified, and rarely treated [12].

[12]. Substantial proportion of CRC patients develops CACS. Depending of complex series of molecular changes which are only partly known.

CRC spreading, antitumor treatments and its complications, we can see that CRC is second in global cancer incidence with increased risk in industrialized nations. There is no incidence difference between genders for CRC which is the most common cause of cancer death among non-smokers. Despite vast achievements in surgery, chemo/bio and radiotherapy, the percentage of 5-year-survivals is still poor and reaches 15%.

More than 80% of PCa cases develop CACS; and it is among highest incidence of CACS compared to other tumors. Therefore the PCa is paradigm for investigation and treatment of CACS.

Muscle wasting in cancer patients is consequence from activation of the ubiquitin proteasome pathway by proteolysis inducing factor (PIF) which is independent of nutrient intake, and therefore nutritional supplementation alone is unable to reverse the process of cachexia. EPA prevents muscle wasting by down-regulating the increased expression of this pathway [26].

After a median of at least 3-month supplementation with EPA, positive changes in weight, significant reduction of acute-phase protein production and the stabilization of resting energy expenditure were registered in many studies [27,28]. Studies suggest that analogized progestational agents (megestrol acetate and medroxyprogesterone), showed that these agents improve appetite and increase weight in advanced cancer patients with slight increase in risk of thrombophlebitis. Also, eicosapentaenoic acid (EPA) in phases I and II produce a decrease in proinflammatory interleukins and TNFαs and open a possibility of reverse cancer cachexia in early stages.

Although reported results are inconclusive in studies phase III, in every-day clinical practice, faced with patients with CACS (or with possibility to develop CACS) we are offering nutritional counseling and nutritional support with EPA and megestrol acetate to our patients who want to reduce their risk of cachexia in the future. Differences were found significant after 3-month supplementation for BMI, KPS, and appetite numeric scale (P<0.05) with EPA alone and EPA + megestrol acetate treatment. EPA produced a decrease in proinflammatory interleukins and TNFαs which are related to muscle wasting in cancer patients. EPA also produced a decrease in leptin and other acute-phase proteins which are related to cachexia. Moreover, EPA improved quality of life for cancer patients treated with EPA and megestrol acetate treatments. Therefore, we recommend EPA as a nutritional support in cancer patients with CACS.

Table 5: Number of patients in Group I and Group II with BMI<20, NST ≥ 5, loss of appetite and decrease in weight gain (>2 kg/month).

Table 4: Karnofsky performance status, weight increasing and appetite numeric range scale in follow up of patients with pca.
adipose tissue before weight loss. There is general failure to recognize patients in nutritive risk early enough and to implement effective nutritive intervention.

Multimodal approach (megestrol acetate, EPA, L-carnitine and thalidomide) has shown better treatment outcome than giving single preparation [29]. Thalidomide also causes increased appetite and body weight, but at a dose of 200 mg caused severe side effects, which make his use controversial. Nonsteroidal anti-inflammatory drugs (NSAIDs) are also interesting in their application to cancer cachexia because they can reduce systemic inflammation. Open-label studies with celecoxib and L-carnitine in preliminary experiments have shown an increase in body weight.

The infusion of ghrelin in patients with breast cancer increased appetite, food intake and feeding satisfaction, with no apparent side effects. Oral ghrelin mimetics increase body mass [30-32]. But as ghrelin and its mimetics have the ability to cause increased levels of IGF-1, it is necessary to examine the safety of their use in larger clinical studies.

Nonsteroid selective androgen receptor modulators (SARM) can have a positive anabolic effect while avoiding virilization and hypertrophy of the prostate gland in men. Application of SARM in postmenopausal women and older men with a tumor cachexia showed weight gain and better functioning of patients with relatively few side effects [33]. In studies of phases I and II by Baylis et al. it was applied a monoclonal antibody to IL-6 in patients with NSCLC. The results showed improvement in body weight and reduced disease symptoms level [34].

As some other authors, we concluded that nutritional counseling and nutritional support can temporarly stop weight loss and improve appetite, social life and quality of life in gastrointestinal cancer patients. In study with CRC patients we even demonstrated impact on survival rate. However, due to a relatively small number of patients and short follow up, those results has to be considered with great caution.

Today we have gap between necessity of preventing and treating CACS and paucity of clear recommendations and evidence for supportive nutritive therapy for oncology patients. We have ethical problems of supplemental nutrition in patients with advanced tumor illness and also problems of cost-benefit ratio of nutritional supplementation in view of cost development in the public health system. Although we can expect new drugs from outgoing studies, it will take a time for their implementation in clinical practice. Symptom control, counseling on nutrition and appropriate physical activity levels are still base for the good treatment of cancer cachexia. Other anabolic preparations such as ghrelin, thalidomide, SARM and monoclonal antibodies require further clinical testing in order to distinguish their performance, but they represent a new frontier in the multimodal treatment of cancer cachexia. Although we can expect new drugs from outgoing studies, it will take a time for their implementation in clinical practice.

Therefore, we still need more phase III studies of already existing nutrition supplements with large number of patients. We can conclude that the role of nutrition therapy is still assumed to be less important than tumor response on antitumor therapy as outcomes are less clear in literature.

No single preparation can be considered as a standard in the treatment of cancer cachexia. Currently, in the treatment of cancer cachexia, are well-established preparations containing EPA, megestrol acetate and corticosteroids.

References


