Multiple Intestinal Lymphoma

B. MASTALIER B1, VIOLETA DEACONESCU1, W. ELAIHAHC DRĂGHICI2, CRISTIANA POPP3, SABINA ZURAC3, M.BALEA4, MIHAELA TEVET4, C. BOTEZATU1

1Surgical Clinic, “Colentina” Hospital, Bucharest, Romania
2ICU, “Colentina” Hospital, Bucharest, Romania
3Pathology Unit, “Colentina” Hospital, Bucharest, Romania
4Haematology Unit, “Colentina” Hospital, Bucharest, Romania

Gastrointestinal tract is the most common location for extralymphonodular lymphomas. The small intestine is affected only in 9% of the cases. Intestinal lymphoma may have single or multiple location. This paper describes a case of multiple location in the small intestine of a non-Hodgkin B-cell in a 53 years old patient, who was initially diagnosed with bilateral pneumonia with pleurisy with E. coli, steeper on the right side, but the persistence of symptoms as fever, malaise, despite appropriate treatment, required further investigation. The CT exam observed fluid collection in the hypogastrium around a digestive loop. The patient underwent surgery, the intraoperative findings being: a large mesenteric tumor ~ 5 cm in diameter, a terminal ileal mesenteric tumor, a mesenteric tumor ~ 6 cm in diameter, omentum with nodular formations, a tumor ~ 3.3/2.5.1 cm in the abdominal wall, pseudotumoral appendix. Segmental enterectomy with entero-enterostomy, excision of mesenteric tumors, appendectomy and omentectomy were performed. Pathological diagnosis was non-Hodgkin marginal zone B-cell MALT type lymphoma of the small intestine with extension to the appendix, meso, omentum and abdominal wall. Postoperatively, the patient received chemotherapy for remission.

Key words: lymphoma, small intestine, mesenteric tumors, enterectomy.

Lymphoma is a cancer with the lymphocytes or lymphoblasts as a starting point. The disease can only affect the lymphatic system or can locate extralymphonodular primary level. It represents approximately 4% of cancers and it is more common in developed countries. The etiology is unknown but possible risk factors are:
- viral infection (ex: Epstein-Barr virus, HIV, HCV)
- bacterial infections (ex: H. pylori)
- chronic state of immunosuppression (ex: post-transplant)
- chemotherapy in history (especially with alkalizing agents) or certain medications (ex: digoxin).

According to current WHO classification of tumors of hematopoietic and lymphoid tissue, lymphomas are divided into:
- Hodgkin’s lymphoma (40%)
- non-Hodgkin’s lymphoma (60%) B cell, T cell, post-transplant lymphoproliferative disorders. 85% of them are represented by B cell lymphomas [1].

According to Ann-Arbor modified staging, Hodgkin lymphomas are classified into:
I: tumor confined to the intestine;
II: regional lymphonodular extension;
III: unresectable extra-regional lymph node extension;
IV: extension to other organs intra or extra-abdominal.

In children, lymphomas represent 10-15% of cancers, accounting for the third form of malignancies. It may sometimes appear as Burkitt lymphoma localized ileocecal [2].

Regarding the location of the small intestine lymphoma, it represents 25% of primary malignancies at this level and 40% of primary gastrointestinal lymphomas location. In most cases, lymphoma occurs in the small intestine as an extranodular location of the systemic disease. As a primary malignancy, individuals with AIDS, celiac disease, organ transplantation, infection with Helicobacter pylori may be involved [3].

In cases with small bowel localisation, the disease sets insidious, symptoms are unspecific, like abdominal cramps, nausea, vomiting, fatigue. Diagnosis is difficult, because imaging examinations such as barium swallow or computer tomography have low sensitivity and specificity [4]. Some cases may present as acute intestinal obstruction, perforation or intestinal invagination, but still complications are rare [5].

ROM. J. INTERN. MED., 2015, 53, 1, 75–80
DOI: 10.1515/rjim-2015-0010
CASE REPORT

Patient S.G., 53 years old, from urban environment, known with dermatitis herpetiformis since 1992, ocular trauma by foreign body in 02.2011, diagnosed in 11.2011 (in another hospital) with bilateral pneumonia with pleurisy *E. coli* stronger on the right, accompanied by fever, malaise, inflammatory syndrome, hypochromic microcytic anemia. Given the persistence of symptoms, the patient continued investigation, abdominal ultrasound showing pericecal and periileum pseudotumoral infiltration, hepato-splenomegaly. Abdominal CT 30.11.2012: fluid collection in hypogastrium around a digestive loop, suggesting abscess or blood; liver and spleen may suggest cirrhosis; bilateral pleural effusion (Fig. 1).

On 5/12/2012 the patient was transferred in the “Colentina” Surgical Clinic, presenting right iliac fossa and hypogastrium pain, fever, pale skin and mucous, weakness. Biological: inflammatory syndrome, mild microcytic hypochromic anemia. Echo reaffirms abdomen CT appearance. Persistence pleurisy certified by Rx heart-lung, right pleural puncture externalises 150 ml serocitrin liquid: sterile culture, normal cytology, erythrocytes, lymphocytes and rare mesothelial cells.

On 13/12/2012 the patient underwent the surgical intervention: adhesion intestinal block with an ileal dilated segment, fixed in the pelvis, corresponding to a large mesenteric 5 cm in diameter tumor, another mesenteric tumor of the terminal ileum, another mesenteric 6 cm tumor, omentum with nodular formations, another tumor (~ 3.3/2.5.1 cm) in the abdominal wall, pseudotumoral appendix. Extended enterectomy including the 2 tumors with latero-lateral entero-enterostomy, double-layer closure of bowel ends with TA 60 stapler, excision of the mesenteric tumors, appendectomy and omentectomy were performed (Figs. 1-18).

![Figure 1.](image1.png)

![Figure 2. Terminal ileon tumor decollated from the pelvis.](image2.png)

![Figure 3. Mesoleium tumor (intraoperative).](image3.png)
Figure 4. Enterectomy sample.

Figure 5. Enterectomy sample – sectioned.

Figure 6. Omentectomy sample.

Figure 7. Appendectomy sample.

Figure 8. Mesenteric tumor – sectioned.

Figure 9. Small intestine glands; muscular mucosa; c. lymphoid proliferation.

Figure 10. Intestinal gland lymphoid infiltrate and intraepithelial glandular structure dezorganization (lymphoepitelial lesion).

Figure 11. Monomorphic lymphoid infiltration with small cells with hypertrophic nuclei with minimal pleomorphism.

Figure 12. Lymphoid proliferation in mesoappendice, seizing small adipocyte lobules remaining.
FAVORABLE POSTOPERATIVE COURSE

The patient was admitted in the Clinic of Hematology and he received chemo-therapy initiation – cure CHOP, well tolerated and Rithuximab 02/06/2012 for remission. The last admission in the hematologic unit was in January 2015 and found normal clinical aspects, with minimal hepato-splenomegaly and without any superficial and internal adenopathy. He will continue his bimonthly therapy with Rithuximab 700 mg.

DISCUSSION

Intestinal localization of the lymphomas represents one of the most frequent situation met in the current practice. If we have the chance to find the illness in an initial stage, we can try to do a radical type of surgical intervention. But, in most cases, the surgeon is facing a situation when the lymphoma is in an advanced stage that does not allow a radical intervention, the operation having the goal to permit the patient to have a normal life, considering the digestive function, and the sampling
of pathologic tissue for the histopathological exam. This is the one who will give us a positive diagnosis that will require the hematologic expertise for establishing the chemotherapy, including the use of monoclonal antibodies substances, for post-operative treatment that has the chance, sometimes, for a practically complete remission of the disease. This case presentation proves that, despite the advanced stage of the lymphoma and the patient’s much altered general state at the moment of admission in our surgical clinic, the evolution after the proper treatment, including surgery and chemotherapy, was good and today, 4 years after the moment of diagnoses, the disease is practically in full remission.

CONCLUSIONS

Intestinal lymphomas can occur as primary tumors or in a systemic disease (6). They are usually associated with immunoproliferative diseases of the small intestine. It generally manifests by infiltrative and diffuse lesions, being most commonly localized on the proximal intestine [7]. The lymphomas can be Hodgkin or non-Hodgkin, the last type being classified in B cell or T cell lymphomas. It affects twice more often males, being met with a higher incidence in white people. The disease has two peaks of age, one under the age of 15 years and another in patients over 50 years [8].

The malignant tumors must be resected with oncological safety margin along the edge of the opposite side of the mesentery in order to remove the lymph locally, regional lymph nodes, to restore the continuity of the digestion. NHL nodular have a better prognosis compared to diffuse NHL. Type B lymphomas have better prognosis compared to type T lymphomas. Low malignacy lymphomas (B cells) have a longer evolution, but they rarely enter in complete remission.

Surgical treatment with excision of macroscopic lesions in conjunction with chemotherapy allows better control of the disease, and the patient has the chance even of healing ad integrum.

Tractul gastro-intestinal reprezintă cea mai frecventă localizare pentru limfoamele extralimfomialor, intestinul subțire fiind afectat doar în 9% din cazuri. Limfoamele intestinale pot avea localizare unică sau multiplă. Articolul de față descrie un caz de localizare multiplă la nivelul intestinului subțire a unui limfom non-Hodgkin cu celule B, la un pacient în vârstă de 53 de ani, care inițial a fost diagnosticat cu pneumonie cu pleurezie bilaterală cu E. coli mai accentuată pe partea dreaptă, dar la care persistența simptomelor sub formă de febră, stare generală alterată, în ciuda tratamentului adecvat, a împus continuarea investigațiilor, examenul CT decoatând colecție fluidă în hipogastru cloazonată în jurul unei anse digestive. Pacientul a fost supus intervenției chirurgicale, intraoperator constatându-se prezența unei mari tumori mezenterice cu diametrul ~ 5 cm, a altei tumori mezenterice de ileon terminal, a încă unei tumori mezenterice cu diametrul ~ 6 cm, oment cu formațiuni nodulare, formațiune tumorală ~ 3,3/2,5,1 cm la nivelul peretelui abdominal, apendice pseudotumoral. S-a practicat enterectomie segmentară cu entero-enterostomie, excizia tumorilor mezenterice, appendicectomie și omentectomie. Diagnosticul histopatologic a fost de limfom non-Hodgkin cu celule B de zonă marginală tip MALT de intestin subțire cu extensie la nivelul apendicelui, mezoului, epiplooonului și peretelui abdominal. Postoperator, pacientul a beneficiat de tratament chimioterapic pentru obținerea remisiunii.

Corresponding author:
REFERENCES


Received January 10, 2015