Patient with Carcinoid Syndrome and Severe Cardiac Disease with Both Tricuspid an Pulmonary Lesions

Ileana Nitu¹,²*, Iulia Florentina Burcea¹,², Daniela Greere¹,², Adina Croitoru¹,², Catalina Poiana¹,²

Abstract

Introduction: Carcinoid Heart Disease (CHD) is a rare cardiac manifestation occurring in patients with advanced Neuroendocrine Tumours (NET) and the carcinoid syndrome, usually involving the right-sided heart valves and eventually leading to right heart failure. The additional burden of cardiac dysfunction heralds a steep decline in quality of life and survival, the prompt recognition of this disease being therefore of the utmost importance.

Case summary: We present the case of a 57 year old female patient diagnosed in 2016 with carcinoid tumor of the proxymal ileum (Grade 2 NET ki 67 8%) and liver metastasis. She underwent surgery with both tumor and secondary determinations resection, followed by chemotherapy in the same year. The patient comes to our clinic in February 2019, complaining in the last 5 months of abdominal pain associated with flushing of the face, simetric peripheral edema, progressive limitation in effort, appearance of dyspnea on moderate exertion and dry caugh. The transthoracic echocardiography detected severe dilation of the right chambers, systolic dysfunction of the right ventricle, tricuspid disease with severe regurgitation and mild stenosis. The echocardiography also revealed double pulmonary lesion with moderate regurgitation and mild stenosis. Endocrine evaluation revealed high levels of serum markers for Neuroendocrine Tumor (NET): serotonin (1543 ng/ml), cromogranin A (291 ng/ml), urinary level of 5-hydroxyindolacetic acid (84 mg/24 h).

Discussion: The carcinoid syndrome was considered to be the cause of the right-heart disease, regarding the fact that echocardiographic features suggested primary valvular lesions. Somatostatin analogue treatment was started with significant improvement in symptoms and NET markers, thus allowing cardiac surgery a month later with remission of symptoms related to heart failure, giving us time for deciding the adequate treatment for the NET. Therefore, the patient received double bioprosthetic valves (tricuspid and pulmonary) in March 2019. Six months later, echocardiographic examination evidenced that both the tricuspid and pulmonary bioprosthesis were functional, with pressure gradients within the normal limits; her functional capacity improved significantly.

Conclusion: The particularities of this case are: the fact that the patient was diagnosed with advanced cardiac disease three years after primary tumor resection, tumor which represents the trigger of the right valvular disease and the fact that not only one valve was affected, but both tricuspid and pulmonary involvement, as it often happens in cases of carcinoid heart disease.
Introduction

Neuroendocrine Tumours (NETs) represent a rare class of neoplasms with an incidence between 2.5 and 5 cases per 100 000 people [1, 2]. It is also a rare cause of acquired valvular heart disease, although cardiac involvement has been described in up to 60% of patients with carcinoid syndrome. Without treatment, carcinoid heart disease has a poor prognosis with a 3 year survival rate as low as 31% (compared with 68% in patients without concomitant heart disease) [3]. Well-differentiated neuroendocrine tumors are rare. They are characterized by the secretion of a vasoactive substance (serotonin), responsible for the appearance of carcinoid syndrome symptoms (flushing, diarrhea, and bronchoconstriction) [2, 3]. The direct action of these vasoactive substances on the heart leads to carcinoid heart disease (CaHD; Hedinger’s syndrome), which is a frequent complication of carcinoid syndrome; CaHD can be the first presentation of carcinoid syndrome in up to 20% of patients. Right-sided heart failure is a prognostic factor of carcinoid syndrome [3]. Transthoracic echocardiography is indicated in all patients diagnosed with carcinoid syndrome, and it should be performed on a routine basis every 3-6 months depending on the clinical presentation of the patient and the severity of the disease [4]. Cardiac involvement can vary widely in patients. In fact, all heart valves may be affected, but the involvement of the tricuspid valve is most common [5]. Tricuspid leaflets show a specific involvement with thickened leaflets that have reduced mobility and are retracted. The concomitant regurgitation can vary from mild to severe [6-8]. Right ventricular volume overload can result in these two valves becoming dysfunctional and can lead to right heart failure [9].

As serotonin produced by the carcinoid tumour is inactivated in the lungs, right-sided carcinoid heart disease is more frequent than left-sided heart involvement (>90% of the cases) [5, 10]. Therefore, the presence or absence of an atrial septum defect or a Patent Foramen Ovale (PFO) needs to be thoroughly evaluated, because in the presence of PFO, left heart valve involvement is more frequent. Cardiac magnetic resonance imaging and Computed Tomography (CT) can be helpful tools to assess damage of heart structures, as well as to evaluate cardiac metastases. Nuclear medicine imaging may be useful to detect the primary tumour; however, it may be of limited use when diagnosing carcinoid heart disease, except for depicting cardiac metastases [11]. In addition to transthoracic echocardiography, biomarkers such as N-terminal pro-brain natriuretic peptide (NT-proBNP), which is a well-established marker for heart failure, has been shown to also have diagnostic and prognostic relevance in patients with carcinoid tumours [12].

However, confounders may lead to false results. Foods and medications that may elevate the levels of 5-hydroxyindoleacetic acid (5-HIAA) should be avoided, and a 24 h urine sample should be taken [6]. Another biomarker is chromogranin A, which is a glycoprotein that is released from NET cells. Its levels are also associated with cardiac involvement [12]. The combination of NT-proBNP and chromogranin A has also been shown to determine survival probability, as patients with elevated levels of both parameters have only a survival probability of 16% compared with 81% in patients with normal levels of both parameters [12]. The management of CaHD is based on medical therapy (somatostatin analogues), resection of the primary tumor, and cardiac interventions. Surgical valve replacement is an effective treatment option for symptomatic patients, since it improves the symptoms and life quality [12].

Case report

In February 2019, a 57-year-old woman presented with heart failure with increasing dyspnea, lower limb edema, and cyanosis of her lips and extremities with refractory hypoxia. She was complaining in the last 5 months of abdominal pain associated with flushing of the face accompanied by night sweats, hepatalgia and asthenia, distension of the abdomen, simetrical peripheral edema, progressive limitation in effort, appearance of dyspnea on moderate exertion and dry cough. We noted that she was diagnosed in 2016 with carcinoid tumor of the proxymal ileum (Grade 2 NET ki 67% and liver metastasis. She underwent surgery with both tumor and secondary determinations resection, followed by chemotherapy in the same year. Clinical examination revealed blood pressure of 120/70 mmHg, oxygen saturation at 85%, and a holosystolic murmur (grade 4/6) suggestive of tricuspid valve regurgitation. Lower limb edema up to her knee joint was also present. We also detected flushing, pale and dry mucous membranes, rales in pulmonary bases, hepatomegaly (liver palpable 4cm below the right costal margin), and positive fluid wave sign. A neurological examination was normal; in particular, there was no sensory or motor deficit and no language disorder. A 12-lead Electrocardiogram (ECG) showed sinus rhythm and a heart rate of 66 beats per minute, QRS axis +30 degrees, inverted T waves in D III and flattened T waves in aVF. She underwent transthoracic echocardiography, which showed dilated right heart cavities with Right Ventricular End Diastolic Diameter (RVEDD)/Left Ventricular End Diastolic Diameter (LVEDD) of 0.8, fractional area change of 30%, Right Atrial (RA) area of 25 cm², and tricuspid ring of 39 mm; Tricuspid Annular Plane

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Systolic Excursion (TAPSE) was 12 mm and Right Ventricular (RV) velocity (S’) was 5 cm/second. Tricuspid valves appeared thickened and retracted with restricted mobility and poor coaptation responsible for a severe tricuspid regurgitation-vena contracta of 8 mm, proximal radius of 7 mm and mild stenosis (Figure 1). Pulmonary valves were thickened with a moderate Pulmonary (IP) insufficiency (Figure 2). The left-sided valves were within normal limits. Our patient had a minimal mitral insufficiency. The function of the left ventricle was good with Ejection Fraction (EF) of 55%.

Cardiac Magnetic Resonance (CMR) imaging revealed dilated right ventricle, with decreased systolic function, an ejection fraction of 44% (Figure 3), severe tricuspid regurgitation- regurgitant fraction of 51%, dilated tricuspid ring- area of 4.3cm2 and moderate pulmonary regurgitation-regurgitant fraction of 23%.

Laboratory findings revealed increased serum NT-proBNP (2248 ng/ml), serotonin (1543 ng/ml), cromogranin A (291 ng/ml), urinary level of 5-hydroxyindolacetic acid (84 mg/24 hours, normal < 10 mg/24hours). An octreotide scan showed an intense radiotracer accumulation in her liver as well as normal distribution of the radiotracer in her kidneys and spleen (Figure 3). Paralleling her diagnostic workup by the patient’s cardiologist and endocrinologist, a comprehensive evaluation was undertaken in the structural heart valve clinic including discussions with oncology and cardiothoracic surgery as to the potential and feasibility of repair options. Medical therapy regarding right heart failure was initiated: loop diuretics, beta blocker, aldosterone receptor antagonist-furosemide (80mg/day), carvedilolum (50mg/day) and spironolactone (50mg/day). Somatostatin analogue treatment was started with significant improvement in symptoms and NET markers, thus allowing cardiac surgery a month later with remission of symptoms related to heart failure, giving us time for deciding the adequate treatment for the NET. As valve repair was not feasible owing to severe fibrosis of the valve leaflets, valve replacement (Edwards St. Jude Epic, 33 mm bioprosthesis) was performed. Therefore,
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Discussion

Carcinoid syndrome was considered to be the cause of the right-heart disease, regarding the fact that echocardiographic features suggested primary valvular lesions and the initial diagnosis in 2016 didn’t reveal any hemodynamically significant valvulopathy. Carcinoid syndrome is a paraneoplastic syndrome mediated by humoral factors released by some carcinoid tumors. The majority of neuroendocrine tumors arise from the small intestine, particularly in the ileum, and they release into the systemic circulation a variety of vasoactive substances: serotonin, 5-hydroxytryptamine, 5-hydroxytryptophan, histamine, tachykinins, bradykinin, and prostaglandins, which explain the clinical manifestations of carcinoid syndrome [2]. Carcinoid syndrome is rare and once it is developed, more than 50% of the patients develop CaHD that can inaugurate this syndrome in 20% of cases. It is associated with a poor prognosis [3]. Symptoms of carcinoid syndrome (facial flushing, hypermotility of the gastrointestinal system, bronchoconstriction, and hypotension) usually occur in patients with hepatic metastatic lesions, due to the lack of hepatic inactivation of these released hormones [5]. These vasoactive substances act directly on the valvular endocardium and activate a fibrotic process. Histopathology reveals fibrous plaque-like and endocardial thickening leading to thickening and retraction of the valves. In this disease, the involvement of the right-sided heart valves is predominant and may be in the form of regurgitation and/or stenosis [6, 7]. Right-sided heart failure remains a major factor of morbidity in patients with CaHD [8]. Left-sided cardiac involvement is rarely observed (<10% of patients), particularly in the case of right-to-left shunt (a PFO) or bronchial carcinoids due to the bypassing of inactivation of serotonin within the lung [9].

The diagnosis is essentially based on biological examinations and echocardiography which remains the principal imaging modality in assessment of CaHD [6-10]. Biological examinations are helpful in the diagnosis of CaHD. High levels of N-terminal pro-brain natriuretic peptide (NT-proBNP), chromogranin-A (a neuroendocrine secretory protein), and urinary 5-HIAA (a metabolite of serotonin) are correlated with the progression of CaHD [11, 12]. Echocardiography shows typical valvular involvement: the tricuspid valve is constantly thickened, rigid, and retracted with restricted mobility and poor coaptation which lead to tricuspid regurgitation. Stenosing character is rarer (25% of cases) but may be associated. The pulmonary valve, which is more difficult to study, is reached in 30% of cases with regurgitation or pulmonary stenosis [10-13]. The right cavities and the inferior vena cava are dilated with or without right ventricular dysfunction [14]. Myocardial strain allows detection of an early right ventricular dysfunction in patients with CaHD independently of valvular involvement [15]. Of all patients with CaHD, ≤10% have lesions of the left-side valve. CaHD in the left side of the heart is less severe than in the right side of the heart. Left-sided heart involvement in carcinoid syndrome happens in patients with persistent foramen ovale or bronchial carcinoid [16].
Patients with CaHD have a decreased life expectancy compared with patients without cardiac lesions [17]. The treatment of patients with carcinoid syndrome is complex and involves multidisciplinary management. It is based on medical therapy, surgery of the tumor, and cardiac surgery [2]. Medical management consists of controlling the heart failure (diuretics agents and aldosterone antagonist) and symptoms of carcinoid syndrome. Therapy by somatostatin analogues allows improvement of symptoms and quality of life and decreases the incidence of CaHD from 50 to 20% [4]. New agents (telotristat and pasireotide) have shown promising results in patients with carcinoid syndrome refractory to somatostatin analogues [18]. Telotristat etiprate is a potent inhibitor of the synthesis of serotonin. The phase III TELESTAR clinical trial has shown that Telotristat may control the bowel movements in patients with carcinoid syndrome. It represents a new option for the treatment of patients with refractory carcinoid syndrome. However, other research is required to verify the safety and the benefit to control symptoms of that new drug [19]. Interferon alpha can be used as a complementary treatment for somatostatin analogues in refractory carcinoid syndrome. Because of its side effects, it should be initiated by 3 MU thrice weekly and then an individual titration [20]. Surgical resection of the primary tumor and resection of liver metastases appears to decrease the cardiac progression in CaHD and improve prognosis [16]. Since hepatic surgery exposes a patient to the risk of perioperative bleeding, hepatic intra-arterial treatment (transarterial chemoembolization, selective internal radiotherapy) is well suited to patients with hepatic metastases [21]. Valve replacement surgery or valvuloplasty is the only effective treatment for symptomatic CaHD; it improves the symptoms and increases the life expectancy of these patients [22]. The optimal moment for valve replacement surgery is not established. However, valvular surgery is proposed when patients become symptomatic or develop ventricular dysfunction provided they have a life expectancy of at least 1 year [20]. Bioprosthetic valves are generally preferred over a mechanical prosthesis, which requires anticoagulation for life and then exposes patients with liver metastases to the risk of bleeding in addition to the risk of prosthesis thrombosis in tricuspid position [23]. Percutaneous valve implantation is a novel option in high-risk patients with severe CaHD, who have poor performance status and comorbidities that do not allow open surgery [24].

Conclusion

The recognition, diagnosis, and treatment of neuroendocrine tumours and their associated syndromes are a rare but important subset of malignancies capable of affecting the cardiovascular system. Rapid diagnosis utilizing serum studies, CT scans, and echocardiography can help expedite the diagnosis and treatment of such rare conditions, and assist in the avoidance of complications due to the effects of circulating vasoactive substances. Despite its relatively well-recognized clinical symptoms, carcinoid syndrome and its associated heart disease still remains a challenging condition to manage and treat, often requiring the input of several subspecialties to treat the condition appropriately.

Declarations

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Conflicts of interest/Competing interests

The authors declare that there is no conflict of interest.

Ethics approval

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Availability of data and material

All data generated or analyzed during this presentation are included in this published article.

Code of availability

Not applicable.

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