A Case of Extra-adrenal Retroperitoneal Ganglioneuroma in a 9-year-old Female: A New Case Report with a Comprehensive Literature Review

Zlatan Zvizdic¹, Nedzad Rustempasic², Irmina Sefic Pasic³, Faruk Skenderi¹, Semir Vranic⁵

¹ Clinic of Pediatric Surgery, University Clinical Center Sarajevo, Sarajevo, Bosnia and Herzegovina, ² Clinic of Cardiovascular Surgery, University Clinical Center Sarajevo, Sarajevo, Bosnia and Herzegovina, ³ Department of Radiology, University Clinical Center Sarajevo, Sarajevo, Bosnia and Herzegovina, ⁴ Department of Pathology, University Clinical Center Sarajevo, Sarajevo, Bosnia and Herzegovina and ⁵ College of Medicine, QU Health, Qatar University, Doha, Qatar

Correspondence: semir.vranic@gmail.com or svranic@qu.edu.qa: Tel.: + 974 4403 7873; Fax.: + 974 4403 3344; ORCID: http://orcid.org/0000-0001-9743-7265

Received: April 25, 2020 Accepted: May 21, 2020

Abstract

Objective – We present herein a new case and survey comprehensively literature on this rare condition. Case report – A 9-year-old girl with a medical history of surgical correction of clubfoot three months earlier presented to our department with an incidentally detected abdominal mass during diagnostic workup for orthopedic surgery. A CT scan revealed a solid right extra-adrenal mass measuring 7×6 cm. It was compressing/involving the infrahepatic part of inferior vena cava, right renal vein and artery with an incomplete encasing of the abdominal aorta. The surgery was successfully performed. The histopathological analysis confirmed GN. Conclusions – Although pediatric extra-adrenal retroperitoneal ganglioneuroma (GN) are rare, their propensity for enveloping major blood vessels is not. GNs should be considered in the differential diagnosis of any circumscribed retroperitoneal mass. These tumors can be successfully treated with surgery that leads to an excellent outcome, even in case of incomplete resections with tumor residuals <2 cm.

Key Words: Ganglioneuroma ● Retroperitoneum ● Children ● Management.

Introduction

Ganglioneuromas (GNs) are rare (one per million population), typically slow-growing, benign tumors arising from the autonomic nervous system. GNs are composed of mature Schwann cells, ganglion cells and nerve fibers, arising from the sympathetic ganglia. The GNs are most commonly diagnosed in children, with the female sex predominance (1). The retroperitoneum is a common site for GNs, and approximately half of the reported cases are extra-adrenal (2). However, GN can arise in any sympathetic tissue, such as neck, posterior mediastinum, adrenal glands, retroperitoneum and pelvis (3).

These tumors are generally asymptomatic because they grow slowly and are usually detected incidentally at a late stage as a large mass (4). GNs may occasionally present with abdominal pain, palpation of an abdominal mass or patients may have symptoms related to adjacent organs compression. A minority of GNs exhibit metabolic activity caused by vasoactive intestinal peptides, catecholamines, and testosterone and present with hypertension, diarrhea, and virilization (5). Although the diagnosis of GN cannot be reliably established on radiological findings alone, magnetic resonance imaging (MRI) and computed tomography (CT) scanning are the preferred imaging methods to assess these tumors (6). A complete surgical removal
is the treatment of choice whenever possible and it usually gives an excellent outcome (7). Unlike smaller GNs that are commonly diagnosed in childhood, large GNs tumors are mainly diagnosed after the age of ten. To date, only a few cases of giant extra-adrenal retroperitoneal GNs were reported in pediatric patients, especially below 10 years of age.

We present herein a case affecting a 9-year-old girl, discuss the therapeutic approach and briefly review literature on this rare pediatric condition.

Case Report

A 9-year-old girl, with a medical history of surgical correction of clubfoot three months earlier, presented to our department with an incidentally detected abdominal mass during diagnostic workup for orthopedic surgery. The physical examination revealed a painless palpable mass on the right side of her abdomen. Blood pressure was within the normal range. Family history was negative for neurofibromatosis and other cancer syndromes. Laboratory investigations (complete blood count, serum electrolytes, immunoglobulins, and urine) and endocrine tests (cortisol, adrenocorticotropic hormone levels, and 24-hour urinary catecholamines) were all within normal ranges. Routine tumor markers were also negative. Abdominal ultrasound examination revealed a right sided extra-adrenal hypoechogenic, somewhat heterogeneous solid mass, occupying the right aspect of the retroperitoneum, measuring 7 cm in its largest diameter.

Subsequent contrast-enhanced CT scan confirmed a vividly enhancing solid right-sided extra-adrenal mass measuring 7.4×6.6×4.2 cm that was compressing/involving the infrahepatic part of inferior vena cava (IVC), right renal vein and artery with an incomplete encasing of the abdominal aorta (Fig. 1A-B). IVC was markedly dilated. A provisional diagnosis of non-functioning, probably benign, right extra-adrenal tumor was made. MRI was also performed to further characterize the mass (not shown). An ultrasonography-guided percutaneous fine-needle biopsy (FNAB) of the tumor was performed revealing a cellular neoplasm composed of ganglion cells and spindle-shaped cells with “cigar-shaped nuclei”. Morphological findings indicated a ganglioneuroma (GN).

After obtaining parental consent, the patient underwent surgery. The lesion was approached transperitoneally through an extended transverse supra-umbilical incision. After mobilization of the right colon, an encapsulated right-sided extra-adrenal retroperitoneal tumor was carefully dissected away from the surrounding adherent structures including IVC, right renal vein and artery, abdominal aorta, and the right kidney, but patency of the compressed vessels were maintained (Fig. 2A-B).

Fig. 1A-B. Contrast Enhanced Coronal (a) and Axial (b) CT scan Showing a Large Heterodense Mass in the Right Upper Abdomen (white Arrows), Compressing Inferior Vena Cava with Incomplete Encasing of Abdominal Aorta.
Enlarged retroperitoneal lymph nodes were not detected. The small cuff of tumor capsule about 0.5 cm was left behind the fascia transversalis, as it was not possible to safely dissect it from the paravertebral space and the underlying spinal nerves. The tumor specimen was submitted to histopathology analysis that confirmed the mass to be a GN (Fig. 3A-B). The patient had an uneventful postoperative recovery and was discharged on the seventh day. One-year follow-up consisted of visits every 3 months revealing no recurrence of the tumor.

Discussion

Ganglioneuromas (GNs), well-differentiated, slow-growing, usually non-secreting and asymptomatic benign tumors, belong to the group of neuroblastic tumors originating from neural crest cells that comprise a spectrum of both benign tumors like GN and malignant tumors such as (ganglio)neuroblastomas (8). Although GNs are often localized in the retroperitoneum, they account for only 1% of primary retroperitoneal tumors and approximately
half of the reported cases are extra-adrenal (2). The differential diagnosis of solid retroperitoneal mass includes various tumors with neurogenic/neuroblastic origins (ganglioneuroma, ganglioneuroblastoma intermixed, neuroblastoma, neurofibroma, schwannoma, and pheochromocytoma), adrenal gland tumors (adenocortical adenoma and carcinoma), various lymphomas (usually non-Hodgkin type), soft tissue sarcomas (e.g. liposarcoma), and germ cell tumors (e.g. teratoma) (9).

Extra-adrenal retroperitoneal ganglioneuromas in pediatric population are very rare. Our literature survey (PubMed/MEDLINE, Scopus, and Web of Science) revealed a few reported cases and small case series (1, 4, 7, 10-20) with very few cases reported in the patients <10 years (summarized in Table 1). Scherer et al. (2001) reported the largest case series of 5 patients (4 patients under the age of 10) depicting the radiological (CT and MRI) features of retroperitoneal GNs. The tumor size varied between 4 and 11 cm and girls were predominantly affected (4/5). Although GNs may present as an isolated finding as confirmed in our case, these tumors may be occasionally seen in association with neurofibromatosis type 1 (NF1) (21-23).

The widespread use of imaging techniques in clinical practice has contributed to an increase in the number of GNs detected incidentally. However, it is difficult to make an accurate preoperative diagnosis of GN and definitive diagnosis is based on histopathological analysis after surgical excision of the tumor. In our case, the preoperative diagnosis of GN was obtained by FNAB and was further confirmed by histopathological analysis. In the current literature, reports of GN that are reliably diagnosed by FNAB and its cytological appearance are scarce. A literature search revealed only a few cases diagnosed by FNAB at this location (24, 25). GNs on FNAB can be difficult to differentiate from other neurogenic/neuroblastic tumors, peripheral nerve sheath tumors, such as schwannomas or neurofibromas and a meticulous search for mature ganglion cells is critical for making an accurate diagnosis. Taken together, a broad differential diagnosis and subtle morphological details give FNAB a limited clinical utility (26).

Retroperitoneal GNs may clinically remain silent for a long time and the diagnosis is often incidental. Despite their benign nature, retroperitoneal GNs show the tendency to partially or completely surround large blood vessels without compromising the lumen in most cases, making their surgical excision extremely challenging (27).

The optimal treatment approach for most patients with retroperitoneal GN is a complete surgical excision (7). However, this approach has certain limitations, particularly in the case in which the large GN surrounds and compresses large blood vessels making its dissection extremely difficult. A mitigating circumstance is a fact that the enclosed blood vessels commonly have no narrow lumen or filling defect (28). In this case, GN enclosed IVC

<table>
<thead>
<tr>
<th>Table 1. A Summary of the Previous Studies that Reported Extra-adrenal Retroperitoneal Ganglioneuromas in Children &lt;10 Years*</th>
</tr>
</thead>
<tbody>
<tr>
<td>References (year)</td>
</tr>
<tr>
<td>Menon et al. (2020) (10)</td>
</tr>
<tr>
<td>Maher et al. (2014) (17)</td>
</tr>
<tr>
<td>Noguchi et al. (2010) (18)</td>
</tr>
<tr>
<td>Zugor et al. (2009) (19)</td>
</tr>
<tr>
<td>Scherer et al. (2001) (4)</td>
</tr>
<tr>
<td>Vara Castrodeza et al. (2000) (20)</td>
</tr>
<tr>
<td>Jain et al. (1999) (14)</td>
</tr>
<tr>
<td>Lin et al. (1997) (15)</td>
</tr>
</tbody>
</table>

*The literature survey included PubMed/MEDLINE, Scopus and Web of Science databases using the following keywords: "Ganglioneuroma", "Retroperitoneum", "Extra-adrenal", "Children", and "Pediatrics". The studies were then filtered based on the patients' age (<10 years) and anatomic location (retroperitoneal but extra-adrenal); †Years; ‡Largest diameter (cm); §Extra-adrenal.
and right renal artery and vein and compressed the anterior side of the right kidney. The dissection of GN away from IVC was the crucial step in the surgery. In our case, all large blood vessels were protected effectively.

Although the postoperative prognosis is usually excellent, long-term follow-up is recommended because of sporadically reported cases of local recurrences (29). However, Decarolis et al. suggested that even incomplete resection of GN is not associated with increased risk of progression if the tumor residuals are <2 cm (29). Following the suggestion of other authors, we also performed a subtotal resection without endangering adjacent vital structures. This approach appears to be sufficient for the successful treatment of GN (30) as the disease did not recur within one-year follow-up period.

Conclusions

Although pediatric extra-adrenal retroperitoneal GNs are rare, their propensity for enveloping major blood vessels is not. GNs should be considered in the differential diagnosis of any circumscribed retroperitoneal mass in pediatric population.

Authors’ Contributions: Conception and design: ZZ and SV; Acquisition, analysis and interpretation of data: ZZ, NR, ISP, FS and SV; Drafting the article: ZZ and SV; Revising the article critically for intellectual content: ZZ, NR, ISP, FS, and SV; Approved final version of the manuscript: ZZ, NR, ISP, FS, and SV.

Conflict of Interest: The authors declare no conflict of interest.

Research Ethics and Patient’s Consent: The study was conducted according to the World Medical Association Declaration of Helsinki. The authors appreciate the patient’s family for consenting to the report of this instructive case.

References


