ABSTRACT

Neuroblastoma is the most frequent solid extra-cranial malignant tumor in children. A proper staging of the disease allows the selection of the appropriate treatment, including the surgical approach. The present review aims to discuss and illustrate the role of computed tomography (CT) and magnetic resonance (MRI) in the differential diagnosis of the retroperitoneal or mediastinal masses, prior discovered by ultrasound or X-ray examination. Furthermore, it is discussed the role of CT and MRI for staging the patients with a histological proof of neuroblastoma. CT scan is very useful to detect and characterize the thoracic and abdominal tumors and organ metastases, with a special quality in detecting structural microcalcifications. MRI is sensitive for cranio-spinal primary tumors, marrow-bone metastases and mediastinal or pelvic primary or secondary tumors. In conclusion, CT and MRI imaging are mandatory diagnostic tools in staging neuroblastomas, finding the optimal moment for the surgical treatment and establish the risk categories of the children.

Key words: neuroblastoma, staging, computed tomography, magnetic resonance

INTRODUCTION

Neuroblastoma is a very redoubtable pathology: the most frequent solid extra-cranial malignant tumor in children (1). Neuroblastoma accounts for 10/100 000 children annually (under 15 years of age) in Europe (2), with approximately 3/100 000 in Romania (3). In the United States 8-10% of all children’s cancers are neuroblastomas (4). The median age at diagnosis is about 2 years (2), with 70% of the patients discovered in stage 4 disease. Thus, the 5-year survival rate is only 30%.

A proper diagnosis of neuroblastoma is crucial for primary surgical treatment and precise staging of the disease allows correct inclusion of the patient in one of the multiple treatment choices for neuroblastoma.

The present review aims to discuss and illustrate the role of computed tomography (CT) and magnetic resonance (MRI) in the differential diagnosis of the retroperitoneal or mediastinal masses, prior discovered by ultrasound or
X-ray examination. Furthermore, it is discussed the role of CT and MRI for staging the patients with a histological proof of neuroblastoma.

**The natural history and clinical course of a neuroblastoma**

The origin of a neuroblastoma can follow tree different pathways:

- most of them arises in the adrenal medulla (medulla glandulae suprarenalis);
- periarterial sympathetic nervous system represented by cervical, abdominal or lateral pelvic (iliac arteries) ganglionar chain;
- ararhidian and presacrat sympathetic system represented by sympathetic chain of the cervical, thoracic, lumbar paravertebral or median pelvic ganglions.

According to these origins, multiple locations of the disease can appear, as follow (2):

I. Abdominal in 60-70% of cases, from which 32-35% originate in adrenal glands;
II. Thoracic (15-20%);
III. Pelvic (5%);
IV. Cervical (5%);
V. Cerebral (1%);
VI. Unknown (0.5-1%).

Distant spread can lead to multiple organ metastases which are generally present at the time of diagnosis; their frequencies, attained at the moment of the diagnosis are, according to some authors (2): sponge bone marrow (70.8%), bone (56%), ganglionar (31%), liver (30%), cranio-orbito-epidural (18%), skin (4%), pleural (3.5%), lung (3%), peritoneal (2%), cerebro-meningeal (0.6%), para-testicular (0.5%), ovarian.

Although clinical diagnosis might be difficult and deceiving, there are clinical warning signs, such as: abdominal or limb mass, with or without pain, weight loss, fever, weakness, or, with some specificity - proptosis or the complex of signs named “raccoon eye”, “blueberry muffin” syndrome in skin metastases (6, 7).

The biological criteria may include increased levels of: serum or urine catecholamines (95% of patients), lactate dehydrogenase, ferritin, neuron-specific enolase (6, 7).

**Staging of neuroblastomas**

Nowadays, neuroblastomas are staged according to the International Neuroblastoma Staging System, as shown in **table 1**.

All the patients with a histological proof of a neural crest tumor can be classified as follows:

- neuroblastoma or poor stromal neuroblastoma (more than 50% neuroblasts);
- ganglioneuroblastoma or rich stromal neuroblastoma with more than 50% stromal tissue (2).

Other authors classify the neuroblastomas in undifferentiated, poorly differentiated, or differentiated (4, 5).

Staging strategy is based on: CT or MRI scan, bone scintigraphy and histological proof by biopsy, including bone marrow biopsy (6).

**Assessment of the neuroblastomas by CT and MRI**

Most of the patients with supposed abdomino-pelvic mass are referred to a first line imaging exam, like ultrasound, while patients with suspected mediastinal mass are referred for a thoracic X-ray examination.

Subsequently, the children with lesions detected by ultrasound or thoracic X-ray, according with the clinical and biological data, with a supposed diagnosis of neuroblastoma are furthermore referred to a sectional exam like CT or MRI scan in order to evaluate the local spread or systemic involvement of the disease, before starting the treatment.

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**Table 1 - The International Neuroblastoma Staging System**

<table>
<thead>
<tr>
<th>Stage</th>
<th>Description</th>
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<tbody>
<tr>
<td>1</td>
<td>localized tumor confined to the area of origin with complete gross resection or without microscopic residual disease; identifiable ipsilateral or contralateral lymph nodes negative macroscopically</td>
</tr>
<tr>
<td>2A</td>
<td>localized tumor with complete gross excision; identifiable ipsilateral or contralateral lymph nodes negative microscopically</td>
</tr>
<tr>
<td>2B</td>
<td>unilateral tumor with complete or incomplete gross excision with positive ipsilateral regional lymph nodes; contralateral lymph nodes microscopically</td>
</tr>
<tr>
<td>3</td>
<td>tumor infiltrating across the midline with or without regional lymph node involvement; unilateral tumor with contralateral lymph node involvement; midline tumor with bilateral regional lymph node involvement</td>
</tr>
<tr>
<td>4</td>
<td>dissemination of tumor to distant lymph nodes, bone, bone marrow, liver or other organs</td>
</tr>
<tr>
<td>4S</td>
<td>localized primary tumor (1, 2A or 2B) with dissemination limited to skin, liver or bone marrow (&lt; 10% tumor cells and MIBG scan negative in the marrow); limited to infants &lt; 1 year of age</td>
</tr>
</tbody>
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For initial staging, CT or MRI imaging are mandatory in correlation with other exams like bone scintigraphy and the histopathological confirmation of one of the neuroblastoma’s histological types.

A CT scan is very efficient in detecting and staging thoracic and abdominal neuroblastoma because of the excellent ability to characterize calcified tumors and to visualize after contrast administration the vascular structures usually involved in this multiregional spread tumor (fig. 1a). CT allows also the diagnosis of organ metastases, like the liver (fig. 1b), or, in some cases, cancellous bone, especially if there are multiples and confluent lesions that destroy cortical margin and invades adjacent structures (fig. 1c).

The MRI is very useful particularly in patients with central nervous system involvement, because of its special capacity to detect peri-vertebral or peri-spinal lesions, which sometimes develop in the foraminal space (fig. 2a); MRI imaging with multiplanar acquisitions allows a precise diagnosis, visualizing foraminal masses in all the tree dimensions (fig. 2b and c).

Unlike the CT, MRI can provide all body imaging that can detect and describe multiple neuroblastoma locations, without irradiating the patient and even without contrast administration: multiple abdominal and pelvic extensive tumoral lesions and numerous peritoneal metastases with consequent large quantity of ascites (fig. 3a and b). The coronal space can give global vision of the tumoral spread (fig. 3c).

The CT gives solid data for:
- detection of the primary site and his local spread in thoraco-abdominal and pelvic lesions;
- proper evaluation of secondary dissemination for cranial and thoraco-abdomino-pelvic regions.

The CT is especially useful in detection of structural microcalcifications (79% of neuroblastomas contain calcifications) (2) and recognition and permeability evaluation of the vascular structures that are surrounded and engulfed by the tumoral nodules, like celiac artery, superior mesenteric artery, aorta, renal vessels, in which case the patient became inoperable in the premier line (2, 7).
Abdominal CT scan can be the gold standard for detection and local development of the neuroblastoma (6).

The MRI is useful especially if:
- there are primary or secondary lesions in cranio-spinal and cervical regions, with a particular sensitivity in spinal extension (6), because the characteristic tendency of neuroblastoma to invade spinal canal via neuroforamina (7);
- there are pelvic primary or secondary lesions, or small parts invasion from a primary lesion;
- there are mediastinal primary tumors with typical posterior mediastinal location;
- there are bone or bone marrow metastases (4).

For thoracic (concerning lungs) or abdominal lesions, MRI is less effective than CT scan for local spread or anatomical details, because of the artifacts developed if there are patient respiratory or body movements.

Finally, the role of CT or MRI imaging is also important in establishing the prognosis: the “Children’s Oncology Group” classifies neuroblastoma patients into low, intermediate and high risk categories based on the age of the patient, the stage of the disease, histological type and genetic characters of the tumor (8).

CONCLUSIONS

The CT and MRI imaging are mandatory diagnostic tools in staging neuroblastoma, finding the optimal moment for the surgical treatment and establish the risk categories of the children.

REFERENCES