Oncology

Perioperative management of hypertensive neuroblastoma: A study from the Italian Group of Pediatric Surgical Oncologists (GICOP)

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Abstract

Background: Hypertension (HT) is rarely reported in patients affected by Neuroblastoma (NB), and management guidelines are lacking. Clinical features and perioperative medical treatment in such patients were reviewed to 1) ascertain whether a shared treatment strategy exists among centers and 2) if possible, propose some recommendations for the perioperative management of HT in NB patients.

Methods: A retrospective multicenter survey was conducted on patients affected by NB who presented HT symptoms.

Results: From 2006 to 2014, 1126 children were registered in the Italian Registry of Neuroblastoma (RINB). Of these, 21 with HT (1.8%) were included in our analysis. Pre- and intraoperative HT management was somewhat dissimilar among the participating centers, apart from a certain consistency in the use of alpha-1 blocker urapidil. Six of the 21 patients (28%) needed persistent antihypertensive treatment at a median follow-up of 36 months (range 4–96 months) despite tumor removal. Involvement of the renal pedicle was the only risk factor consistently associated with HT persistency following surgery. A correlation between the presence of HT and the secretion of specific catecholamines and/or compression of the renal vascular pedicle could not be demonstrated.
Neuroblastoma (NB) is one of the most frequent pediatric solid tumors. It often originates from the adrenal gland or from the adjacent paravertebral sympathetic ganglia and frequently involves the kidney.

Hypertension (HT) is a relatively rare symptom of NB, its reported incidence ranging between 10% and 19% of cases. Few studies have focused on the perioperative management of HT, and most of these have been case reports [1–9]. Both cardiac and neurological complications of HT have been described [10].

Both catecholamine secretion and renal vascular compression have been advocated as the possible pathophysiological mechanism of HT in NB [2,3]. Although the secretion of catecholamines may be increased in NB patients, HT and other symptoms of catecholamine excess frequently observed in pheochromocytoma have been reported in relatively few children with NB (1–9). This difference is probably owing to the presence of many storage granules in pheochromocytoma, which release large quantities of catecholamines, thereby causing HT. In neuroblastic cells, this storage mechanism seems to be inefficient, causing intracellular granule disruption, the inactivation of catecholamines and the release of their metabolites into the circulation. A small number of NB do exhibit abundant storage granules and patients can exhibit blood pressure patterns like those of patients with pheochromocytoma [11–14]. Chemotherapy may result in additional catecholamine release and HT owing to tumor lysis, thus warranting close observation during preoperative cycles [4].

Renal artery stenosis is a possible cause of HT and has been observed in about 3% to 10% of cases of pediatric HT [15,16]. NB causing renovascular HT owing to extrinsic aortic narrowing and/or stenosis of the renal artery has been reported [2,5].

Preoperative echocardiography is the cornerstone in the diagnostic workup of these patients, as catecholamine excess can induce several complications other than, or subsequent to, HT, including seizures and heart failure, cardiomyopathy, cardiomegaly and cardiacogenic shock [3,6,15,17–26]. All in all, preoperative HT control has lowered reported mortality from 13% to 45% to 0%–3% in patients with catecholamine-secreting tumors [27].

The anesthesiological intraoperative management of hyper- and hypotensive peaks has rarely been reported [3,4]. Whatever the cause of HT in NB, symptoms are generally well controlled by the preoperative administration of alpha-1 adrenergic blocking agents, such as phenoxybenzamine, phentolamine, prazosin and, more recently, doxazosin [3,4,6,9,17–26].

Despite effective preoperative alpha-1 adrenergic blockade, intraoperative HT may still occur, especially when the tumor is being manipulated. Therefore, the availability of a wide range of rapidly acting medications to treat intraoperative hypertensive peaks is mandatory; these include prostaglandin E1, diuretics, calcium channel antagonists, sodium nitroprusside, hydralazine, magnesium sulfate, phentolamine, adenosine, and fenoldopam [4,6,20–22]. Beta blockers, such as propranolol, labetalol, atenolol and esmolol, may be administrated intraoperatively in association with alpha-1 blockers to control catecholamine-induced arrhythmia and tachycardia [4,6]. Postresection hypotension may be successfully managed by means of continuous iv administration of dopamine in the first 24 postoperative hours [6].

In the present study, we retrospectively reviewed an Italian series of NB patients with HT, with a view to assessing the possible correlation between HT and the secretion of specific catecholamines and/or compression of the renal vascular pedicle. We also evaluated the perioperative management of HT to find out whether a possible treatment recommendation can be presented.

1. Materials and methods

The Italian Registry of Neuroblastoma (RINB) collects all newly diagnosed cases of NB in Italy, together with demographic, chemotherapeutic, surgical, and follow-up data.

The Italian Group of Pediatric Surgical Oncologists (GICOP) conducted a retrospective survey of all the patients affected by NB with associated HT registered in the RINB from 2006 to 2014. HT was defined as blood pressure values higher than the 99th percentile + 5 mmHg, as stated by the National High Blood Pressure Education Program Working Group in 2004 [28]. All children who had undergone surgical resection were included in the present study.

The following data were retrospectively collected in an electronic database: series demographics, diagnostic workup, including assays of urinary and/or blood catecholamines, image-defined risk factors (IDRFs) present on preoperative imaging (CT or MRI), tumor histology according to the International Neuroblastoma Pathology Classification (INPC) [29], tumor staging according to the International Neuroblastoma Staging System (INSS) [30] and the International Neuroblastoma Risk Group Staging System (INRGSS) [31], and biomolecular features such as the presence or absence of MYCN amplification.

Pre-, peri- and postoperative antihypertensive treatment regimens were examined. Treatment duration, drugs administered and possible HT persistence during follow-up were recorded.

Anesthesiological treatments were collected and divided into premedication, epidural, induction, muscle relaxation, maintenance and the need for packed red blood cell transfusion.

Surgery was defined as complete if more than 95% of the tumor had been resected (microscopic residual disease), as leaving minimal residual disease (if more than 90% but less than 95% of the tumor had been removed, macroscopic minimal residual disease), or as incomplete (if less than 90% of the tumor had been removed).

Clinical follow-up examination was conducted by a pediatric nephrologist, an oncologist and a cardiologist, and included 24-h continuous blood pressure measurement and radiological reevaluation (CT, MRI and ultrasonography scan).

2. Results

From 2006 to 2014, 1126 children affected by NB were registered in the RINB, 25 of whom presented HT (2.2%). Four of these 25 children did not undergo surgical resection, and were therefore excluded from the analysis. Consequently, the study cohort comprised 21 patients. Patients’ demographics and tumor characteristics are detailed in Table 1.

Vanillylmandelic acid (VMA) and homovanillic acid (HVA) were tested in all patients and were found elevated in 9 (43%). Norepinephrine and epinephrine were assayed in only 3 patients, and proved to be elevated in all of them. Dopamine was never analyzed.

An age-comparable series of 28 patients presenting with non-HT NB and involvement of the renal vascular pedicle was examined to detect any significant differences. The only finding that emerged from this analysis was that a relatively higher proportion of patients in the nonhypertensive group displayed increased VMA/HVA (89% of normal-tension NB patients vs 43% of HT NB patients). No statistical analysis was conducted on these data.
Preoperative radiological imaging showed normal-sized kidneys (age- and sex-matched) in all patients. Involvement of the renal vascular pedicle was observed in 8 of the 21 patients (38%) in our study cohort (Fig. 1).

The four excluded patients had not undergone surgery owing to the persistence of 3 or more IDRFs after chemotherapy (Patient 1: bilateral renal pedicle, aorta, celiac artery, superior mesenteric artery; Patient 2: renal pedicle, hepatic pedicle, cava, aorta; Patient 3: renal pedicle, superior mesenteric artery, aorta; Patient 4: renal pedicle, cava, aorta). Only Patient 4 had increased VMA/HVA. Patients 1 and 2 are alive with active disease and are on dual anti-HT therapy. Patients 3 and 4 have died.

None of the patients presented with cardiac hypertensive-related complications on preoperative echocardiography.

Preoperative antihypertensive treatment was necessary in 18 patients (86%) for a median preoperative time of 90 days (range: 4–96 months). This consisted of single-drug therapy in 8 patients and multiple-drug therapy in 10 patients (Table 2).

The induction and maintenance of anesthesia were almost uniform in all centers, with midazolam being used in 19 patients (90%), propofol for the induction phase in 17 (81%), rocuronium for muscle relaxation in 18 (86%) and sevoflurane for anesthesia maintenance in all patients. Epidural anesthesia was performed in 9 patients (43%). Six patients developed intraoperative hypertensive peaks, which were successfully managed with alpha 1 blockers (urapidil) in 5 cases and with a combination of calcium antagonist (nifedipine) plus diuretic (furosemide) in the remaining case.

No hypertensive complications were documented during surgical procedures.

Surgical results are detailed in Table 1. None of the patients in this series underwent nephrectomy to achieve complete resection.

On follow-up examination conducted after a median of 36 months (range 4–96 months), 12 patients were in complete remission (57%), 6 presented with residual disease (28%) and 3 had died owing to disease progression (14%). Six patients (28%) presented persistent postoperative HT despite complete tumor resection in 4 of them and minimal residual in 1. Long-term antihypertensive treatment was therefore started in these patients. Four of these six patients had been treated with antihypertensive medications for more than 100 days preoperatively.

Six out of 8 children with documented involvement of the renal pedicle (on IDRF evaluation) presented persistence of HT after surgery. As these 6 patients constituted the whole cohort of children with persistent postoperative HT, renal pedicle involvement seems to be the only risk factor predictive of persistent HT (owing to the small number, no statistical analysis was conducted on these data). These patients underwent a thorough postoperative radiological assessment by means of US-Doppler and CT scan to detect possible renal hypotrophy/atrophy possibly owing to persistent spasm of the renal artery owing to prolonged intraoperative vessel manipulation and/or impaired renal perfusion observed during surgery. This analysis revealed no pathological findings in patients with persistent HT, showing normal-shaped and normal-sized kidneys in all of them, regardless of the involvement of the renal pedicle.

3. Discussion

NB is one of the most common solid tumors in infancy, accounting for approximately 8% of childhood cancers. HT was documented in nearly 2.2% of the cases recorded in the RINB, thus constituting a rare condition associated to NB; its management varies widely according to the center involved.

To the best of our knowledge, the present study analyzed the largest series of children affected by NB with HT. Nevertheless, it is still limited,
and no definite conclusions can be drawn owing to the heterogeneity of diagnostic and therapeutic protocols within the different centers involved in the study. Unfortunately, only urinary metabolites of vasoactive amines, namely VMA and HVA, were routinely investigated in all patients, and few centers performed detailed assays of dopamine, epinephrine and norepinephrine.

Surprisingly, only 1 patient from our series showed 5-MC-N amplification. We could not find any clinical explanation for such an unusual distribution (considering that at least 16 patients had stage L2/M disease). Although none of the patients from our series presented with cardiac hypertensive-related complications, a complete cardiologic assessment, including echocardiography, is mandatory in the correct preoperative workup of these patients [3,6,11,25,26].

With regard to the antihypertensive medications used in our series, the wide heterogeneity in drug selection, both preoperatively and perioperatively, is immediately evident. This is mainly owing to the lack of a recognized and accepted guideline dedicated to cases of NB presenting with HT. However, a certain consistency was observed regarding the use of one rapidly acting alpha-1 blocker, namely urapidil, which proved to be quite effective in the management of intraoperative hypertensive peaks. Although described in the intraoperative management of pheochromocytoma in adult patients [23], to the best of our knowledge, urapidil administration has never been reported in children affected by NB presenting with HT.

In this retrospective study, preoperative anti-HT therapy was not homogeneous: 18/21 underwent preoperative anti-HT therapy; the remaining three patients were stage L1 on diagnosis and did not receive any anti-HT therapy. The three centers concerned were therefore contacted; it transpired that preoperative treatment had not been undertaken owing to the short time (<7 days) between the diagnosis of hypertension and surgical intervention. In all 18 patients, preoperative anti-HT therapy achieved HT control without any clinical complication. One patient underwent prolonged treatment (300 days) owing to the ‘unresectable’ nature of the tumor (stage L2 with multiple IDRFs). The patient was eventually transferred to another center and underwent surgery. As reported by Avanzini and colleagues [32], IDRFs remain unchanged in 50% of cases after chemotherapy, and this can delay surgical resection.

Nearly 30% (6/21) of the patients in our series showed persistently increased blood pressure levels following medical and surgical treatment. Of note, in only 1 of these 6 patients might this condition have been explained by persistent disease owing to incomplete surgery. We could not find any plausible reason for HT persistence in the remaining 5 patients, since all of them underwent thorough postoperative radiological assessment by means of US-Doppler and CT scan, without any renal hypertrophy/atrophy being documented. Similarly, the persistence of HT despite surgical treatment was documented in 1 out of 12 cases reported by Madre et al. [1], without any evident clinical reason.

Involvement of the renal pedicle, as detected on preoperative imaging, seems to be the only risk factor that strongly predicts the postoperative persistence of HT; this scenario was observed in 75% of patients with renal pedicle involvement, who constituted the whole cohort of children with persistent postoperative HT.

On the basis of this retrospective review of NB patients with HT, no definite therapeutic protocol can be recommended. However, we would highlight a certain agreement on the intraoperative use of an alpha-1 adrenergic blocker (namely urapidil) in association with a beta-adrenergic blocker and a calcium channel antagonist to control tachycardia and possible arrhythmias. An effective combination of these proposed medications may limit the risk of cardiac complications, which may affect anesthetic and surgical outcomes.

Similarly, postresection hypotension and hypovolemia should be carefully addressed by means of continuous infusion of dopamine and fluid replacement, preferably in a pediatric intensive care setting, at least for the first 24 h postoperatively.

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