Diagnostics, treatment, and follow-up in craniopharyngioma

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Craniohypophysis is a rare, non-glial, non-malignant intracranial tumor derived from a malformation of embryonal tissue. The pathogenesis of craniopharyngioma is currently debated between two primary hypotheses: one describes tumor origin as ectodermal remnants of Rathke’s pouch; the second argues craniopharyngioma represents a malformation of residual embryonal epithelium of the anterior pituitary gland and anterior infundibulum (Garré and Cama, 2007; Müller, 2010). Anywhere from 30 to 50% of the 0.5 to 2 cases per million persons per year manifest during childhood and adolescence (Bunin et al., 1998; Nielsen et al., 2011), representing 1.2–4% of all childhood intracranial tumors. In childhood and adolescence, histological type is usually adamantinomatous with cyst formation (Müller-Scholden et al., 2001; Rushing et al., 2007). Incidences of adult onset craniopharyngioma (usually between ages 50 and 75) most often present with a squamous-papillary histological type (Rushing et al., 2007). More than 70% of craniopharyngioma of the adamantinomatous type bear a mutation of the β-catenin gene, which is not detectable in the papillary type of craniopharyngioma (Hölsken et al., 2010).

The German Childhood Cancer Registry obtained data on 496 patients in whom a craniopharyngioma was diagnosed at age ≤18 years from 1980 to 2007. Of these, 451 patients (91%) were younger than 15 years-of-age at the time of diagnosis. The sex ratio was 1:1 and the median age at primary diagnosis was 8.6 years; survival rate (1980–2007) was 97% at 3 years, 96% at 5 years, and 93% at 10 years after diagnosis. Patients who were diagnosed and treated in the 1980s had a lower survival rate than those diagnosed in the 1990s. For example, the survival rate at 5 years was 91% for patients diagnosed in the 1980s versus 98% for those diagnosed in the 1990s (Müller et al., 2003c). This observation is supported by other reports (Sherlock et al., 2010), indicating that mortality is increased in childhood craniopharyngioma patients compared to general population.

Diagnosis – clinical manifestations and imaging methods

A background study of craniopharyngioma patients reveals initial symptoms often occur long before diagnosis is made (Müller, 2010, 2011), the non-specific manifestations of increased intracranial pressure such as headache and nausea often going unrecognized as primary symptoms of craniopharyngioma. Other manifestations of the craniopharyngioma diagnostic profile can include visual impairment (62–84%) and endocrine deficits (52–87%; Figure 1). Craniopharyngioma tumors that involve the hypothalamic–pituitary axis can affect the secretion of growth hormone (75%), gonadotropins (40%), adrenocorticotropic hormone (ACTH; 25%), and thyroid-stimulating hormone (TSH; 25%). Growth impediment caused by these hormonal disturbances often appears as early as 12 months of age; however, premature puberty and/or unmanageable weight gain are more often the precipitating factors preceding clinical diagnosis in older children.
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FIGURE 1 | Manifestations before diagnosis of craniopharyngioma in children and adolescents. Frequency of occurrence of each manifestation before diagnosis (blue) and frequency of occurrence as the initial manifestation (red). The median time (month) from the appearance of each initial manifestation until diagnosis is indicated above each red column. In the overall group, the median time from the initial manifestation of disease until diagnosis was 12 months, with a range of 0.01–96 months. (Modified from Müller et al., 2003c, with the kind permission of Springer).

(Müller, 2010). In summary, young children presenting symptoms of decreased growth rate, older children presenting symptoms of premature puberty and/or unmanageable weight gain, patients (adults or children) presenting headache, visual impairment, and/or unmanageable weight gain, and/or polydipsia/polyuria should arouse suspicion of craniopharyngioma (Müller, 2010).

The combination of solid, cystic, and calcified tumor components is an important radiological clue to craniopharyngioma diagnosis (Warmuth-Metz et al., 2004). The signal intensity of craniopharyngioma in magnetic resonance imaging (MRI) is highly variable, as it depends on the protein content of the cysts. Solid tumor portions and cyst membranes appear isointense in T1-weighted images, often with a mildly heterogeneous structure (Figure 2). MRI before and after gadolinium application is the standard imaging for detection of craniopharyngioma, but because CT is the only way to detect or exclude calcification, which is found in approximately 90% of tumors, (Warmuth-Metz et al., 2004), a sellar or parasellar mass detected on MRI should be further imaged by native CT for detection of calcifications.

TREATMENT – CHALLENGES AND STRATEGIES

Disturbance of cerebrospinal fluid (CSF) flow often causes hydrocephalus, which, depending on severity, must be stabilized by surgical treatment. Tumor resection is the first-choice treatment for restoring normal CSF flow, but a pre-surgery shunt operation may also be required. For a craniopharyngioma with large cystic components, stereotactic, or open implantation of an intracystic catheter is a proven treatment both for the relief of pressure and, in some cases, for the instillation of sclerosing substances (bleomycin). An intracystic catheter with a subcutaneous reservoir can be effective for reducing cyst volume and is particularly appropriate for infants and toddlers where extending the interval until radiotherapy or surgical resection is advantageous in some cases. For patients experiencing preoperative visual impairment due to large cysts exerting pressure on the optic nerve, a two-staged treatment approach is proposed, with cyst drainage to relieve pressure and improve vision, followed by resection (Choux et al., 1991).

After preoperative assessment of calcifications by CT, the recommended therapy for favorably tumor localized tumors is complete resection while preserving visual, hypothalamic, and pituitary functions (Choux et al., 1991; Müller, 2006). However, radical resection, especially in infants and small children, is problematic due the heightened risk of surgically induced deficits (mainly hypothalamic) and the susceptibility of children to recurrence (23%), even in apparently complete resections. For topographical-anatomical reasons, transsphenoidal surgery is the proven neurosurgical technique for minimizing disturbance to these structures (Fahlbusch et al., 1999). Following postoperative MRI confirmation of complete resection, a post-surgical native CT of the sellar/parasellar area is recommended for reconfirmation of complete resection. A right frontotemporal operative approach is standard, although localization of craniopharyngioma should dictate the approach. Intrasellar craniopharyngioma can be approached via transsphenoidal route. However, childhood craniopharyngioma usually extends to the suprasellar area and must be removed through a transcranial approach. For topographical-anatomical reasons, transsphenoidal surgery has proven advantageous for minimizing disturbance to hypothalamic functions (Fahlbusch et al., 1999).

For unfavorably localized tumors (optic nerve and/or hypothalamic involvement), a limited resection followed by local irradiation is statistically favorable: the risk of progression is 71–90% without postoperative radiotherapy (Figure 3), but drops to 21% if combined with postoperative radiotherapy (Becker et al., 1999;
In addition to difficulties in deciding the degree of resection, the timing of postoperative residual tumor irradiation remains unclear and controversial (Regine and Kramer, 1992; Stripp et al., 2004; Moon et al., 2005; Tomita and Bowman, 2005; Merchant et al., 2006; Müller et al., 2006b). Some favor immediate irradiation following surgical procedure and some purport a wait-and-see approach to minimize both the necessity for irradiation and sequelae associated with radiation therapy. Proactive post-surgical irradiation to prevent tumor progression is favored by the above-mentioned North American Center (Merchant et al., 2006). But even though immediate postoperative irradiation significantly delays tumor progression, progression-contingent irradiation is effective, reflected in its unaffected overall survival rates. These contrasting approaches have been retrospectively compared by three series (Stripp et al., 2004; Moon et al., 2005; Tomita and Bowman, 2005). Moon et al., reported no overall survival rate differences between the two strategies. Tomita et al., reported that for those patients undergoing immediate postoperative irradiation after complete resection, 83% were relapse-free after 5 years and 70% were relapse-free after 10 years. For those undergoing incomplete resection followed by immediate irradiation, 71% were relapse-free after 5 years and 36% were relapse-free after 10 years. Only 9% were relapse-free after 5 years when incomplete resection was not immediately combined with radiation therapy. Progression-contingent irradiation achieved similar survival- and progression-free survival rates of 90 and 70%, respectively, meaning progression-contingent irradiation in this series also was highly effective. However, the interpretation of these studies is confounding by different indications for treatment. They also only reflect relapse-free survival rates and do not include consequences of craniopharyngioma QoL sequelae.

The target volume of irradiation is calculated based on CT and/or MRI images. Current imaging techniques with enhanced resolutions allow a safety margin ideally no greater than 5 mm, depending on the tumor location, size, composition (solid, cystic, calcified), and adjacent structure involvements (hypothalamic–pituitary axis and/or optic nerve or chiasm), requiring a larger safety margin if the hypothalamus is involved. Three-dimensional planning and multi-field techniques with individual field configurations (collimation) are recommended to protect radiosensitive structures and to provide a maximal dose fall-off between the tumor and the adjacent structures (Becker et al., 1999; Einhaus and Sanford, 2007).

Due to the low histological malignancy of craniopharyngiomas and especially because they are located near essential structures regulating eyesight, growth, and energy homeostasis, proton beam therapy appears to be a promising therapeutic option, offering a more protective radio-oncological technique than conventional external irradiation (Baumert et al., 2004; Fitzek et al., 2006; Luu et al., 2006). Also reported as a promising treatment technique is the intracystic instillation of interferon alpha (Ierardi et al., 2007). The stereotactic instillation of radioisotopes is mainly applicable to monocular recurrences of craniopharyngioma, which is recommended for pre-surgical treatment of large cystic tumors and for tumors that recur after both surgery and percutaneous radiation therapy.
Other therapeutic options, including stereotactic gamma-radiotherapy (Gamma Knife), are less promising due to limited experience; and in the case of single-dose convergence irradiation, of little treatment value due to reasons related to radiation biology.

Beyond treatment success ratings of craniopharyngioma related to diagnostic and treatment strategies is the experience level of the neurosurgeon. A few studies have analyzed treatment outcome related to the neurosurgeons’ experience (Sanford, 1994; Boop, 2007; Müller et al., 2011b). Both Sanford and Boop report a marked difference in outcome according to the neurosurgeons’ experience with the condition. The most recent of the three reports (Müller et al., 2011b) analyzed craniopharyngioma outcome and prognosis relative to patient load of the treating neurosurgical centers, finding centers with lower craniopharyngioma patient load – and therefore relatively less experience treating craniopharyngioma – tended to use more radical surgical approaches, resulting in less favorable outcomes regarding obesity and other QoL considerations. Current literature advises a multidisciplinary team approach to craniopharyngioma treatment, enabling micro- and macro-examinations of pre-surgical, intraoperative surgical, and post-surgical radio-oncological options, as well as long-term QoL issues.

QoL MANAGEMENT – PRE- AND POST-TREATMENT CONSIDERATIONS

KRANIOPHARYNGEOM 2007, a prospective, European multinational trial (Müller et al., 2003c, 2006b; Müller and Sörensen, 2007; Müller, 2010, 2011), is currently evaluating craniopharyngioma patients’ prognoses (QoL, event-free, and overall survival rates) following defined therapeutic strategies. A stratified randomization of two treatment arms is conducted with respect to timing of postoperative irradiation (immediate irradiation versus irradiation at the time of progression) for the subgroup of patients ≥5 years-of-age at the time of incomplete resection (Figure 4). The schedule of prospective data collection and the set and definition of parameters in KRANIOPHARYNGEOM 2007 are based on a European consensus (Müller et al., 2006a,c). The trial is open for international recruitment (www.kraniopharyngeom.net). There are also other current prospective studies underway on national and multinational levels to adopt strategies tailored to risk factors for morbidity and QoL (Garré and Cama, 2007; Puget et al., 2007). A recent report (Müller et al., 2011b) shows that especially tumor involvement and surgical lesions of posterior parts of hypothalamic structures predispose to adverse late effects such as obesity and consecutively impaired quality of life (Figure 5). The primary craniopharyngioma sequelae affecting patients’ quality of life are visual impairment (Choux et al., 1991), hypothalamic lesions causing neuropsychological deficits (emotional instability, rage attacks, abnormal sexual behavior, and deficits of memory and intellectual capacities; Fischer et al., 1990; Fisher et al., 1998; Riva et al., 1998; Müller et al., 2005a,b; Pierre-Kahn et al., 2005), and endocrine deficits (including diabetes insipidus, hormonal deficiencies causing growth and puberty disturbances, hyperphagia and hypothalamic obesity, and eating disorders, which are observed in 40–50% of craniopharyngioma patients (DeVile et al., 1996b,c; Müller et al., 2001, 2003b,d, 2004a; Srinivasan et al., 2004).

Most patients (85–95%) suffer from multiple deficits of hypothalamic–pituitary function, ranging to complete pituitary insufficiency (DeVile et al., 1996a; Merchant et al., 2002a), and full restoration of preoperatively deficient hormonal function occurs.
only in rare cases (Honegger et al., 1999). Growth hormone substi-
tution therapy is reported as a well-documented effective and
safe treatment option in patients with growth hormone deficiency
(Lustig et al., 2003b, Müller et al., 2010). Nevertheless, the distur-
bance of hypothalamic structures by the tumor and/or its treat-
ment is considered to be the major pathogenic factor for hyper-
phagia and obesity, confirmed by an imaging study assessment
of the extent of hypothalamic involvement and consequent
sequelae (Devile et al., 1996b). Another study performed by Roth
et al. (1998) measured serum leptin levels in craniopharyngioma
patients and found significantly elevated leptin concentrations in
relation to BMI in patients with a suprasellar tumor components,
finding that the body mass index (BMI) of affected patients posi-
tively correlated with the degree of hypothalamic damage. Normal
inhibition of appetite after eating fails to occur because of disrup-
tion of the negative feedback loop involving leptin, formed in
adipocytes, which binds to hypothalamic leptin receptors. Weight
gain occurs in craniopharyngioma patients, even when caloric
intake is similar in craniopharyngioma patients compared to
BMI-matched controls (Harz et al., 2003).

Exacerbating the management of weight gain caused by cranio-
pharyngioma and its treatment is that children with craniopharyn-
gioma have a markedly lower than normal level of physical activity
caused by daytime and disturbances of circadian rhythms (Müller
et al., 2002). Initial trials with melatonin substitution in child-
hood craniopharyngioma patients have proved promising (Müller
et al., 2006c). Polysomnographic studies in craniopharyngioma
patients with severe daytime sleepiness reveal sleeping patterns
typical for secondary narcolepsy (Müller et al., 2006d), confirming
this sequela to be a major contributor to depreciated QoL. Mason
et al. (2002) reported a significant weight loss in childhood cranio-
pharyngioma patients treated with a central stimulating agent
dextroamphetamine).

Both the cause and treatment of weight gain in craniopharyn-
gioma patients has been approached on several fronts. Lustig et al.
(2003a,b) postulated hypothalamic disinhibition of vagal output
as a cause of increased beta-cell stimulation, leading to hyper-
insulinism and obesity. In a randomized, double-blinded study,
treatment with a somatostatin analog (octreotide) to suppress
beta-cell activity resulted in weight reduction. Due to reduced con-
centrations of catecholamine metabolites in the urine of patients
with childhood craniopharyngioma correlating with the degree
of obesity and the level of physical activity, Roth et al. (2007)
hypothesized that craniopharyngioma patients’ decreased phys-
ical activity and severe obesity might be related to decreased
central sympathetic output. In another study, Roth et al. (2011)
analyzed the gastrointestinal hormones ghrelin and peptide YY
and their effect on satiety in patients with childhood craniopharyngioma and obesity, supporting that reduced ghrelin secre-
tion coupled with reduced postprandial suppression of ghrelin
leads to disturbed regulation of appetite in these patients. Pept-
ide YY levels did not differ between normal weight, obese, and
very obese patients with childhood craniopharyngioma. Recently
reported (Roth et al., 2010) is a possible pathogenic role of
peripheral alpha-melanocyte-stimulating hormone in childhood
craniopharyngioma obesity. Even though initial experiences with
bariatric surgery (laparoscopic adjustable gastric banding, LAGB)
to treat severe craniopharyngioma obesity have been encouraging,
long-term follow-up will be necessary to analyze its efficacy, safety,
and long-term effect on weight development (Müller et al., 2007,
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CONCLUSION
A retrospective study of functional capacity using the FMH instrument that quantifies patients’ abilities to perform everyday psychomotor tasks (Wolff et al., 1996) found significantly lower rankings for craniopharyngioma patients compared to age-matched controls (Müller et al., 2003a, 2004b), with craniopharyngioma patients’ morbid hypothalamic obesity cited as responsible for their low functional capacity self-rating. Age-dependent differences between childhood and adult onset craniopharyngioma are related to histological diagnosis, biological behavior, clinical manifestations, treatment options and follow-up (Koranyi et al., 2001; Attanasio et al., 2002; Kendall-Taylor et al., 2005). The Kendall-Taylor et al. (2005) study compared childhood craniopharyngioma with adult onset craniopharyngioma and reported a poor state of health and QoL in both cohorts. The majority of childhood and adult craniopharyngioma patients displayed pituitary insufficiency, with 60% suffering from diabetes insipidus. Nearly all patients were overweight or obese, reporting a poor QoL. What we can take away from this and other reports of both childhood and adult onset craniopharyngioma is that craniopharyngioma involves a series of chronic morbidities ranging from visual impairment to life-long, seemingly untreatable, unmanageable obesity that require on-going patient monitoring to optimize their QoL. Such monitoring provides a two-way communication platform that serves as both a data collection medium on how sequelae unfolds after treatment, as well as a patient-reporting medium of new treatment options that may be revealed as continuing research develops on this all-too confounding disease.

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