Stereotactic biopsies of brainstem lesions: which approach?

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Abstract
Background Stereotactic biopsies for brainstem lesions are frequently performed to yield an accurate diagnosis and help guide subsequent management. In this study, we summarize our experience with different stereotactic approaches to brainstem lesions of different locations and discuss possible implications for safety and diagnostic yield.

Methods We retrospectively analyzed 23 adult patients who underwent a stereotactic biopsy for brainstem lesions between October 2011 and December 2019. Depending on the location supra- or infratentorial, trajectories were planned. We assessed the postoperative complications during the hospital stay as well as the diagnostic yield.

Results A supratentorial transfrontal approach was used in 16 (70%) cases, predominantly for lesions in the midbrain, upper pons, and medulla oblongata. An infratentorial, transcerebellar-transpeduncular approach was used in 7 (30%) cases mainly for lesions within the lower pons. All biopsies were confirmed to represent pathological tissue and a definitive diagnosis was achieved in 21 cases (91%). Three patients (13%) had transient weakness in the contralateral part of the body in the immediate postoperative period, which improved spontaneously. There was no permanent morbidity or mortality in this series of patients.

Conclusion Lesions of various locations within the brainstem can be successfully targeted via either a supratentorial transfrontal or an infratentorial transcerebellar transpeduncular approach. Our high diagnostic yield of over 90% and the low rate of complications underlines the diagnostic importance of this procedure in order to guide the medical management of these patients.

Keywords Brainstem · Stereotactic biopsy · Frame-guided · Stereotactic trajectory

Introduction

Brainstem lesions comprise 10–15% of all space occupying lesions of the brain in children and 2% in adult patients [13, 15]. Many diffuse brainstem lesions in the pediatric population do not warrant a biopsy due to the typical appearance on MR imaging suggesting a diffuse brainstem glioma with no relevant differential diagnoses [19]. In adults, however, pure radiological findings will often fail to make the correct diagnosis, due to the broader spectrum of underlying diseases including gliomas, tumors of other origin, infectious diseases, infarctions, vasculitis, demyelinating diseases, and gliosis [1, 6]. In fact, MRI-based diagnosis is reported to be erroneous in 10–20% of the cases [12]. Furthermore, the MRI accuracy to assess the tumor classification and grading was estimated to be correct in only 35% of low-grade gliomas and 27% of high-grade gliomas [17]. With growing importance of molecular markers for tumor classification and guidance of treatment, together with increasing safety and diagnostic yield of stereotactic biopsy of brainstem lesions, the justification of specimen-based diagnosis seems ever more important even in pediatrics population, as the availability of biopsy material will allow for molecular biology analysis including whole genome sequencing, thus potentially allowing for the development of future therapies [15, 18].

Stereotactic biopsies of brainstem lesions can be performed along different trajectories depending mainly on the location of the lesion. Both supratentorial and transcerebellar approaches have been described with similar outcomes [22]. In this study, we report our experience in 23 brainstem biopsies in an adult population, including the spectrum of diagnosis, the chosen trajectory, and the complications. Secondarily, we aim to propose surgical biopsy trajectories (i.e., supratentorial transfrontal or infratentorial transcerebellar, transpeduncular) according to the anatomical location of the lesion.
Material and method

Patient population

A consecutive series of twenty-three adult patients (16 males, 7 females) who underwent a stereotactic biopsy for a brainstem lesion at the University Hospital Bern between October 2011 and December 2019 are included in this retrospective study. All adult patients who present with a brainstem lesions without clear radiological or previously known diagnosis and/or which were not amenable to surgical resection were selected for a stereotactic biopsy to confirm the diagnosis in order to direct multidisciplinary treatment plans. The patients’ mean age is 58 years (ranging 28 to 82 years) All cases of radiologically demonstrated lesions localized to brainstem were included in the study. In patients with larger lesions involving other regions of brain, in addition to brainstem, or multiple lesions, only the cases where the target of biopsy was brainstem were included. All cases with a target outside the brainstem were excluded. The mean duration of symptoms was 4.2 months. Symptoms consisted of cranial nerve deficits in 6, limb weakness in 5, gait disturbance in 3 patients, headache and signs of increase intracranial pressure in 4, sensory disturbances in 3, and 2 patients were not documented. Patient’s demographic, clinical presentation, chosen trajectory, histological diagnosis, and complications are summarized in Table 1. This study has been approved by the local ethics committee (KEK number 2020-00440).

Pre-operative imaging

All patients underwent preoperative 3 Tesla MRI sequences including 3D T1-weighted MPRAGE with contrast and 3D T2-weighted sequences (voxel size 1 mm3) (Somaton, Siemens, Erlangen, Germany). Magnetic resonance spectroscopy (MRS) could be performed in only 11 patients due to

| Patient | Age/Gender | Localization                  | Presenting symptoms    | Approach   | Complications  | Histopathological diagnosis                     |
|---------|------------|-------------------------------|------------------------|------------|----------------|-----------------------------------------------|
| 1       | 39/F       | Diffuse pontomedullary       | Double image           | Transfrontal | None           | Astrocytoma WHO grade II                      |
| 2       | 76/F       | Pons                          | hemiparesis            | Transcerebellar | None | EBV-associated CNS lymphoma                   |
| 3       | 56/M       | Pontomesencephalalne          | Hemiparesis, gait instability | Transcerebellar | None | GBM, WHO grade IV                             |
| 4       | 71/M       | Midbrain tegmentum           | headache               | Transfrontal | None           | GBM, WHO grade IV                             |
| 5       | 78/M       | Pontomedullary                | Double image           | Transcerebellar | None | Lymphoma                                      |
| 6       | 28/M       | Pontomedullary                | Paresthesia, dizziness  | Transcerebellar | None | Anaplastic astrocytoma WHO grade III          |
| 7       | 75/M       | Pontomesencephalalne          | Double image           | Transfrontal | Right sided weakness | Anaplastic astrocytoma WHO grade III |
| 8       | 58/M       | Pontomesencephalalne          | Headache               | Transfrontal | None           | Anaplastic astrocytoma WHO grade III          |
| 9       | 50/M       | Diffuse brainstem            | Gait instability        | Transfrontal | None           | Anaplastic astrocytoma WHO grade III          |
| 10      | 70/M       | Midbrain                      | Double image           | Transfrontal | upper limb weakness | CNS lymphoma                                |
| 11      | 63/M       | Midbrain tegmentum           | Hemiparesis            | Transfrontal | None           | GBM, WHO grade IV                             |
| 12      | 82/F       | Midbrain                      | Double image, gait instability | Transfrontal | None | B-cell lymphoma                               |
| 13      | 73/M       | Midbrain tectum              | Headache               | Transfrontal | upper limb weakness | GBM, WHO grade IV                             |
| 14      | 66/F       | Midbrain                      | Headache, double vision, | Transfrontal | None           | Metastatic neuroendocrine carcinoma (colon CA) |
| 15      | 67/M       | Pontomedullary                | Paresthesia            | Transcerebellar | None | B cell lymphoma                               |
| 16      | 31/F       | Midbrain tegmentum           | Dizziness              | Transfrontal | None           | Hypercellularity (Rosenthal fibers)           |
| 17      | 66/M       | Midbrain                      | Not documented          | Transfrontal | None           | Anaplastic astrocytoma WHO grade III          |
| 18      | 55/M       | Pontomedullary                | Dysphagia              | Transcerebellar | None | GBM WHO grade IV                              |
| 19      | 45/M       | Medulla oblongata             | Headache               | Transcerebellar | None | Ependymoma WHO grade II                       |
| 20      | 37/F       | Pons                          | Headache, double image | Transfrontal | None           | Pilocytic astrocytoma                          |
| 21      | 40/M       | Midbrain                      | Gait disturbances       | Transfrontal | None           | Nonspecific gliosis with microhemorrhage      |
| 22      | 60/M       | Pontomesencephalalne          | Not documented          | Transfrontal | None           | GBM WHO grade IV                              |
| 23      | 68/M       | Pontomesencephalalne          | Paresthesia            | Transfrontal | None           | GBM WHO grade IV                              |
uncooperativeness of the patients. FDG-PET scan has been performed in 5 cases where there was no clear enhancement and/or because of reimbursement issues. Figure 1 illustrates examples of brainstem lesions that were biopsied.

A stereotactic CT with contrast (voxel size 1 mm$^3$) with the Leksell stereotactic frame (Elekta, Stockholm, Sweden) was performed on the day of surgery.

**Trajectory planning**

Surgical planning was performed with iPlan software (BrainLab, Germany). The target was chosen in the center of the contrast enhancing tumor or—in case of no enhancement—in the center of the brightest T2 signal abnormality of the lesion. We chose a supratentorial transfrontal trajectory in cases where the lesion was in the midbrain, upper pons, and medulla oblongata. Lesions in the lower pons that were accessible via the cerebellar peduncles were targeted via an infratentorial, transcerebellar, and transpeduncular approach (Fig. 2). All trajectories were planned to avoid blood vessels, sulci, and the ventricles.

**Surgical procedure**

The surgical procedure was performed under general anesthesia. Minimal shaving is limited to the entry point area. DIXI guides tubes (DIXI, Besançon, France) were used to perform a 2.5-mm diameter burr hole strictly following the axis of the defined trajectory and then insert the stereotactic needle used for the biopsy.

In case of supratentorial transfrontal approach, the patient was placed in a standard supine position, whereas semi-sitting position was provided for transcerebellar approach. In some of our transcerebellar cases, we removed one of the rear bars of the frame in order to be able to perform the surgically planned approach. Usually, three samples were collected for each patient by rotating the opening of the biopsy needle into different direction and taking the subsequent samples but staying within the same initially chosen target point. The first specimen was used for frozen section processing to ensure positive sampling. The second or the third specimen was used for the definitive histological and molecular analysis or microbiology.
if an infectious process was suspected. The average time for the surgical procedure including the time needed for the frozen section assessment by the pathologist was 45 min.

Routine postoperative imaging was not applied but reserved for cases presenting a new neurological deficit after the procedure or confirming that the tissue sample was collected according to the planned target in case of diagnostic failure.

Results

Lesion location and trajectory

The location of the lesion according to epicenter of contrast enhancement or the T2-hyperintensity is shown in Table 2.

We applied a supratentorial transfrontal approach in all our cases of midbrain lesions, as the transcerebellar transpeduncular approach is limited because of the narrow superior cerebellar peduncle and the very low entry point beyond the limits of the frame ring angle (Fig. 3a, b, c). For pontine lesions, we used a supratentorial transfrontal approach in 30% of the cases and infratentorial transcerebellar transpeduncular approach in 70% (Fig. 3d, e, f). The supratentorial transfrontal approach was used in one case of medullary lesion as a transcerebellar transpeduncular approach was technically not feasible due to a resulting entry point above the tentorium (Fig. 3g, h, i).

Considering the whole cohort of patients, the transfrontal approach was used in 16 (70%) cases and the transcerebellar transpeduncular approach was used in 7 (30%) cases.

Surgical outcome

There was no mortality in this series. Three patients developed transient neurological worsening which recovered spontaneously a few days later without any further neurosurgical intervention. One patient developed a contralateral upper and lower limb weakness after a biopsy of an astrocytoma WHO grade III located in the pontomesencephalic region. In this case, the biopsy was at cerebral peduncle taken via a supratentorial transfrontal approach. The other two patients developed an upper limb weakness after a biopsy of a lymphoma and a high-grade glioma (WHO grade IV). Both lesions were located in the midbrain tegmentum and the biopsy was taken via a supratentorial transfrontal approach. The immediate postoperative CT scan did not show any signs of hemorrhage, edema, or ischemia. No complications were found with the infratentorial, transcerebellar-transpeduncular approach.

Histological diagnosis

In all patients, the initial frozen sample confirmed pathological tissue sampling. A conclusive definitive diagnosis was
made in 21 patients (91%). Confirmed histological diagnoses are shown in Table 3.

Of the remaining two patients, one showed increased cellularity with Rosenthal fibers and the other nonspecific gliosis with microhemorrhage. In both cases, we performed a postoperative CT scan which confirmed that the biopsy was performed in the targeted area. Both patients were followed up closely with repeated radiological and clinical assessment without changes of the clinical status in both.

Table 3  Histopathological diagnosis of the biopsied brainstem lesions

| Histopathology results          | Number of patients (%) |
|---------------------------------|------------------------|
| Astrocytoma WHO II              | 1 (4.3)                |
| Astrocytoma WHO III             | 4 (17)                 |
| Glioblastoma multiforme WHO IV  | 8 (35)                 |
| Lymphoma                        | 5 (22)                 |
| Metastasis                      | 1 (4.3)                |
| Ependymoma                      | 1 (4.3)                |
| Pilocytic astrocytoma           | 1 (4.3)                |
| Uncertain diagnosis             | 2 (9)                  |

Fig. 3  Schematic coronal (left), sagittal (middle), and axial (right) representation of the cases and the used trajectories, midbrain cases (a, b, c), pons cases (d, e, f), and medulla cases (g, h, i)
Discussion

In this study, we present a systematic anatomical and clinical analysis of different trajectories for brainstem biopsies and propose an illustrative scheme to approach these technically demanding lesions. In this series of 23 patients, both the supratentorial and infratentorial approaches yielded a definitive pathological diagnosis in over 90% of the cases. Furthermore, both techniques were not associated with long-term morbidity or mortality. Transient neurological deterioration occurred in 13% of our patients who underwent a biopsy of midbrain lesions via a supratentorial transfrontal approach.

We used both supratentorial (transfrontal or transparietal) and transcerebellar approaches. Both approaches have been described and used by several groups [4, 5, 11, 15, 16, 22]. However, information about the choice of the trajectory related to their precise location in the brainstem has not been specifically addressed in previous studies.

Depending on the exact location of the lesion in the brainstem, we recommend the following approaches:

1. Midbrain lesions can be approached through a supratentorial transfrontal approach. Choosing a gyrus near the coronal suture as an entry point is feasible for anteriorly located lesions (crus cerebri, as well as anterior tegmentum) and may need to be adjusted more anteriorly (up to 2 cm anterior to the coronal suture) for lesions located in the tectum (Fig. 4).

2. Lesions in the pons accessible through an infratentorial, transcerebellar-transpeduncular approach should preferentially be targeted by this approach due to the short trajectory and paucity of any eloquent tracts or nuclei along this trajectory. Superior pontine lesions as well as lesions in the midline of the pons are often not accessible through a transcerebellar-transpeduncular trajectory and a supratentorial transfrontal approach can be performed in these cases (Fig. 5).

3. Medullary lesions can be approached through a supratentorial transfrontal approach. Alternatively, in cases of a lesion located at the ponto-medullary junction, a transcerebellar-transpeduncular approach can be evaluated (Fig. 6). Selected superficial medullary lesions could also be biopsied through an open transcisternal approach.

We performed all our procedures under general anesthesia. Of note, most of the procedures described in literature were performed under local anesthesia and intermittent intravenous sedation agent. Based on our results, we think that performing brainstem biopsies under general anesthesia is safe and offers a more tolerable atmosphere for the patients. Furthermore, when performing procedures within the brainstem, respiratory compromise is unpredictable and sometimes catastrophic. Emergent intubation and resuscitation are occasionally required. We also strictly control the intraoperative blood pressure. It is impossible to compare our values with other data obtained from patients operated under local anesthesia, as these values are not reported. Therefore, we are not able to make any correlation about the intraoperative blood pressure value and the bleeding risk.

Some previous studies have reported on complication rates exceeding 10% [7, 10]. Furthermore, the diagnostic yield of stereotactic biopsy for intracranial lesions shows wide variations among different previous studies ranging from 2 to 30% in some studies [8, 14] and more than 90% in others. A recent meta-analysis of 38 studies reported a diagnostic yield of 96.2% by weighted average proportions analysis, which is comparable to our results [9]. Nevertheless, based on these observation, some authors have even questioned the use of stereotactic biopsy for these lesions [2, 3]. Interestingly, most of the lesions in the present study were located in the midbrain or upper pons and targeted with a supratentorial transfrontal approach and—consequently—a long trajectory tract. In contrast, in many previous studies, the majority of lesions were located in pons followed by medulla [7, 10, 20] but no details on the chosen trajectories are provided. We speculate that by using an infratentorial, transcerebellar-transpeduncular approach for mid-pontine lesion that come along with a shorter trajectory length within the

Fig. 4 Coronal (left), sagittal (middle), and axial (right) T2-weighted MRI images showing the proposed midbrain regions of interest (ROI) and the preferred trajectory (red, crus cerebri; green, tegmentum; blue, tectum)
brainstem should decrease the risk of bleeding or the risk of induced microlesion in that very eloquent area and could be accountable for the variability of observed complication rates.

Molecular markers of brain primary tumors such as O6-methylguanin-DNS-methyltransferase (MGMT), isocitrate dehydrogenase-1 (IDH-1), and alpha-thalassemia/mental retardation syndrome X-linked (ATRX) have recently been shown to be correlated with chemotherapy response and prognosis [21]. These markers can only be determined by obtaining a tissue sample. This strongly argues for the important role of brainstem biopsies to not only confirm a suspected diagnosis but specify molecular subtypes of tumors with implications on subsequent therapeutic options.

There are some limitations for this study. The main limitation is the small sample size of patients of our cohort compromising a better estimate of complication rate and the diagnostic yield of brain stem biopsies in general. However, our results are in line with previous studies including larger numbers of patients [5, 11, 15, 16]. Furthermore, the current results are based only on observations reflecting our targeting strategy and no conclusions can be drawn if one of the two suggested approaches would yield lower complication rates. To answer this question, a prospective study or a retrospective case-control study of similar lesions targeted via different approaches and adequate sample size would be needed. On the other hand, we can demonstrate the feasibility of the two approaches with similar outcome. Although we would propose to apply a transcerebellar-transpeduncular approach whenever technically possible due to the shorter trajectory length and less eloquent brain tissue along the tract, of course, it remains the individual surgeon’s choice, which approach to apply.

Conclusions

Stereotactic biopsies of brainstem lesions play an important role in the diagnosis and subsequent treatment of patients, especially in the current era of fine molecular diagnosis and targeted chemotherapy. Supratentorial transfrontal and transcerebellar-transpeduncular approaches can be applied and yield a high diagnostic accuracy and low morbidity. A specific trajectory according to the specific location of the lesion should be considered when approaching brainstem lesions. For midbrain lesion, we suggest a supratentorial transfrontal approach; for pontine lesion, a transcerebellar-transpeduncular trajectory should be the first choice if the lesion is reachable from that approach; and for medullary lesion, a transfrontal approach should be preferred unless the lesion is in the upper part of the medulla where transcerebellar transpeduncular approach can still be used.
Conflict of interest  The authors declare no competing interests.

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