INVITED REVIEW

Single isocenter stereotactic irradiation for multiple brain metastases: current situation and prospects

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Abstract
The prognosis of patients with brain metastases has dramatically improved, and long-term tumor control and reduction of the risk of late toxicities, including neurocognitive dysfunction, are important for patient quality of life. Stereotactic irradiation for multiple brain metastases, rather than whole-brain radiotherapy, can result in high local control rate with low incidence of neurocognitive deterioration and leukoencephalopathy. Recent advances in radiotherapy devices, treatment-planning systems, and image-guided radiotherapy can realize single isocenter stereotactic irradiation for multiple brain metastases (SI-STI-MBM), in which only one isocenter is sufficient to treat multiple brain metastases simultaneously. SI-STI-MBM has expanded the indications for linear accelerator-based stereotactic irradiation and considerably reduced patient burden. This review summarizes the background, methods, clinical outcomes, and specific consideration points of SI-STI-MBM. In addition, the prospects of SI-STI-MBM are addressed.

Keywords Single isocenter stereotactic irradiation · Multiple brain metastases · Linear accelerators

Introduction
Brain metastases are common in cancer patients, with an incidence of 9.6% of all cancer patients [1]. Brain metastases and peritumor edema often cause severe neurological symptoms, which directly affect the patients’ quality of life. Surgery, systemic therapy, radiotherapy, and the best supportive care are the treatment options for patients with brain metastases. Although surgery can be a treatment option for brain metastases generally larger than 3 cm in diameter, complications due to surgery are sometimes severe, and the risks of decline in performance status and neurological function should be considered. In contrast, radiotherapy has been used as a useful and less invasive local therapy for brain metastases [2].

Whole-brain radiotherapy (WBRT) used to be the sole standard radiotherapeutic approach for patients with brain metastases, and the intent and irradiation dose of WBRT are palliative rather than curative. This is because the whole brain is irradiated during WBRT, and ablative doses to the whole brain inevitably result in severe acute and late toxicities. Despite only palliative irradiation doses, neurocognitive dysfunction and leukoencephalopathy due to WBRT should be considered, because these late complications are sometimes observed and directly affect patient’s lives [3]. Recently, prognosis of patients with brain metastases has dramatically improved owing to advances in systemic therapy [4]. Therefore, long-term brain tumor control and reduction of the risks of late complications caused by radiotherapy are important for improving patient quality of life. Thus, ablative local radiotherapy for brain metastases has become increasingly important and is used in clinical practice.

Stereotactic irradiation was developed and clinically used as an alternative ablative radiotherapy option for intracranial lesions using the Leksell Gamma Knife System for gamma knife radiosurgery (GKS) [5]. In addition to the gamma knife, linear accelerators (LINAC) can deliver stereotactic irradiation for brain metastases using circular cones and multileaf collimators [6, 7]. In GKS- and conventional LINAC-based stereotactic irradiation, tumors are irradiated lesion-by-lesion, the number of brain metastases increases, and the total treatment time increases. Recently, owing to advances in image-guided radiotherapy and treatment-planning devices, LINAC-based stereotactic irradiation can deliver a precise dose distribution...
with excellent conformity and steep dose gradient in a shorter time and in a less invasive manner [8]. Only 1–4 brain metastases have been treated with LINAC so far, because of technical challenges and prolonged treatment time. However, sophisticated optimization in treatment-planning systems has realized stereotactic irradiation for multiple brain metastases with only one isocenter: single isocenter stereotactic irradiation for multiple brain metastases (SI-STI-MBM) [9]. Thus, more than four brain metastases become technically treatable using SI-STI-MBM. Here, the current situation and prospects of SI-STI-MBM are reviewed.

### Improvement of prognosis of patients with brain metastases

Prognosis of patients with brain metastases was limited before 2000s. In 1997, Gasper et al. reported the recursive partitioning analysis (RPA) classification, which showed the prognosis of patients with brain metastases treated with radiotherapy [10]. The survival rate at 20 months was close to 15% even in the best prognosis group (RPA class 1).

Thereafter, other prognostic classifications have been published: GPA [11], disease-specific GPA [12], lung-mol GPA [4], updated Lung-GPA [13], melanoma mol-GPA [14], GI-GPA [15], and breast GPA [16]. Lung-mol GPA was advocated in 2017, 20 years after publication of the RPA classification. In the lung-mol GPA group, the survival rate at 20 months was similar between the worst group (GPA 3.5–4) and RPA class 1. In contrast, the best group (GPA 0–1) showed a survival rate close to 80% at 20 months, and the median overall survival was 46.8 months.

To improve the prognosis of patients with brain metastases, clinicians should pay more attention to late toxicities caused by radiotherapy. Neurocognitive dysfunction is a major and serious late complication of WBRT [17]. In contrast, stereotactic irradiation has little impact on neurocognitive dysfunction when compared with WBRT and can deliver an ablative irradiation dose to the tumor, which leads to long-term local control. Therefore, stereotactic irradiation is used more often than in the past, especially in patients with a good prognosis and a few lesions. From the point of view of the technical aspects of LINAC and clinical evidence on stereotactic irradiation in multiple brain metastases, up to four brain metastases are candidates for LINAC-based stereotactic irradiation.

### Stereotactic irradiation for up to ten brain metastases

In several randomized control trials involving patients with WBRT and stereotactic irradiation, the overall survival in patients with 1–4 brain metastases was similar regardless of whether the patients received WBRT and/or stereotactic irradiation [18–22]. Although 1–4 brain metastases are commonly treated with stereotactic irradiation for long-term tumor control and reduction of late toxicities, patients with four or more brain metastases are only treated with gamma knife radiosurgery in a limited number of institutions [21, 23, 24]. However, Yamamoto et al. reported a multi-institutional prospective observational study, termed JLGK0901, in which patients with up to ten brain metastases were treated with stereotactic radiosurgery (SRS) alone [25]. That study investigated GKS in patients with 1–10 brain metastases and showed a similar overall survival between patients with 2–4 brain metastases and patients with 5–10 brain metastases. The study included patients with the largest tumor with a volume of 10 cm³ or less and a longest diameter of 3.0 cm or less. In addition, the cumulative volume of all tumors was limited to 15.0 cm³ or less. Based on the JLGK0901 study, the NCCN guidelines developed the concepts of “limited brain metastases” and “extensive brain metastases” in 2018. “Limited brain metastases” is defined as a group of patients for whom SRS is equally effective and offers significant cognitive protection when compared with WBRT. SRS is preferred over WBRT in patients with limited brain metastases. Thus, up to ten brain metastases are now often treated with stereotactic irradiation.

### Development of SI-STI-MBM for multiple brain metastases

In LINAC-based stereotactic irradiation, brain metastases are conventionally treated with conformal arcs, one by one [6]. Accordingly, treatment time increases as the number of brain metastases increases. Considering the increased treatment time and burden in patients with frame-based fixation or oppressive frameless thermoplastic masks, only 1–4 brain metastases could be treated with LINAC-based stereotactic irradiation.

Owing to advances in image-guided radiotherapy and optimization methods for treatment-planning systems, SI-STI-MBM can be performed in clinical settings [26]. In SI-STI-MBM, only one isocenter was sufficient, and multiple brain metastases were irradiated simultaneously. Therefore, the treatment time was dramatically reduced in SI-STI-MBM when compared with GKS and dynamic conformal arc therapy lesion-by-lesion [26–29]. By shortening the treatment time, the burden on patients can be reduced.

Currently, two irradiation approaches are clinically used for SI-STI-MBM. The first is the volumetric-modulated arc therapy (VMAT) (Fig. 1). By using VMAT for SI-STI-MBM, dose distribution for multiple brain metastases with better conformity and steep dose fall-off can be made, even if tumors have irregular shapes or large maximum
If two brain metastases are close to each other, it is challenging to reduce the dose delivered to the normal brain between the two lesions. However, VMAT can decrease the irradiated dose of normal tissue as much as possible while maintaining conformity and a steep gradient using dose-intensity modulation and inverse-planning methods. Although VMAT plans can be manually generated by modifying the optimization objects for each lesion and index, knowledge-based planning (KBP) can help clinicians to generate SI-STI-MBM with VMAT plans [31]. In SI-STI-MBM with VMAT plans, inverse-planning methods and dose-intensity modulation can realize that all multiple targets are covered by the prescribed dose while maintaining high conformity, steep dose fall-off, and a high maximum dose for each site. However, making SI-STI-MBM plans places a large burden on radiation oncologists and medical physicists and may limit the chances of timely treatment initiation in patients with multiple brain metastases. RapidPlan™ (Varian Medical Systems, Palo Alto, CA, USA) is a KBP product that utilizes a machine learning system to establish a model to predict dose-volume histograms. RapidPlan™ enables less-experienced physicians to easily create high-quality plans in a short time [31, 32]. RapidPlan™ can be used to make SI-STI-MBM plans.

The second approach is to use a treatment-planning device specialized for SI-STI-MBM with conformal arcs [33] (Fig. 2). The specialized treatment-planning system adopts an inverse-planning method and optimizes the collimator angles, couch angles, number of arcs, selection of the arc that irradiates the lesions, and the leaf motion for each target. Because of the utilization of dynamic conformal arc therapy for each lesion, leaf motion is not as complex as that of VMAT, and there is no need to conduct quality assurance (QA) for VMAT. As a result, SI-STI-MBM using dynamic conformal arcs can be performed quickly after computed tomography simulation, and the burden for medical physicists to conduct QA can be reduced. As this exclusive treatment-planning system of SI-STI-MBM with dynamic conformal arcs, Multiple Brain Mets SRS (BrainLab AG, Munich, Germany) has been released and is clinically used worldwide [34].
Clinical outcomes of SI-STI-MBM

Several publications have examined the clinical outcomes of SI-STI-MBM for multiple brain metastases [26, 34–39]. Table 1 summarizes clinical outcomes of SI-STI-MBM. The local control rate at 12 months ranged from 82 to 97.5%, and toxicities above grade 2 occurred in 8% of patients. Although most reports were from retrospective studies, Kim et al. conducted a prospective study examining patients with 4–10 brain metastases [38]. They reported that the local recurrence rate at 12 months was 5% for all lesions, with no grade 3–5 treatment-related adverse events [38]. Therefore, SI-STI-MBM seems to achieve high local control with a low incidence of severe complications when compared with conventional LINAC-based stereotactic irradiation. Although the low-dose irradiated volume of the normal brain was larger in SI-STI-MBM than in GKS, target conformity and high-dose irradiated volume of the normal brain were similar to GKS [27, 28, 40]. The clinical effects of a larger low-dose irradiated volume when compared with that of GKS, remain unknown.

While only brain metastases are usually irradiated in SI-STI-MBM, a new approach that combines SI-STI-MBM and WBRT using a simultaneous integrated boost (SIB) method is applied in some institutions [41, 42]. In a phase 2 trial conducted in Canada, 47.5 Gy in 5 fractions was delivered to each brain metastasis, and the whole brain was covered with 20 Gy in 5 fractions simultaneously using the SIB method [42]. The local control rate at 12 months was 88%, and the severe radionecrosis above grade 2 was only 1.9% in nondeep lesions. Although these outcomes appeared tolerable, grade 3–5 radionecrosis occurred in 25% of patients with deep brain metastases, including basal ganglia and thalamus. In this study, the planning target volume (PTV) was created by adding 2 mm margin to the gross tumor volume (GTV) for brain and brainstem metastases. When brain metastases are located in eloquent or deep areas, radionecrosis and local recurrence directly affect neurological symptoms, and it is better to pay more attention to delineate contours and add margins to create PTV. Zhong et al. reported the clinical outcomes and quality of life of patients with brain metastases who underwent WBRT with SIB boost [41]. Although only 13 patients were included, the local control rate per lesion at 12 months was 98.6% and no adverse events above grade 2 were observed. In addition, there was no significant cognitive decline among the included patients during the median follow-up period of 11 months. Hippocampal-avoidance WBRT (HA-WBRT) can be delivered to patients with brain metastases, and a clinical trial investigating the benefits of delivering higher doses to brain metastases in HA-VMAT using the SIB method is ongoing in Singapore [43]. Although it is unclear whether WBRT combined with SI-STI-MBM is suitable, the advancement of radiation oncology may offer more options to treat brain metastases in a less invasive manner.

Table 1 Clinical outcomes of SI-STI-MBM with LINAC

| Authors (publication year) | Number of patients | Number of metastases | Median dose | Overall survival | Local control | Toxicities |
|---------------------------|--------------------|----------------------|-------------|-----------------|--------------|------------|
| Nath et al. [26] (2010)   | 26                 | 138 (range 2–13)     | Median 18 Gy/1 fr (range 14–25 Gy/1–5 fr.) | 6 m: 50%      | 12 m: 97%  | Grade 3: 8% |
| Lau et al. [33] (2015)    | 15                 | Median 3 (range 2–13) | 20 Gy/1 fr 3 cases: SRT | 6 m: 60%      | 12 m: 92%  | No grade 3 or 4 |
| Serna et al. [34] (2015)  | 52                 | Total 87 (range 1–3) | 12–20 Gy/1 fr | Median 7.2 m    | 82%         | NA         |
| Palmer et al. [37] (2020) | 173                | 1014 (median 3, range 3–20) | 18–24 Gy/1 fr 21–27 Gy/3fr 25–30 Gy/5fr | Median 13.4 m | 12 m 99%  | Grade 2: 1.4%, Grade 3: 0.9% |
| Kraft et al. [35] (2021)  | 140                | Total 708            | SRS 18–20 Gy SRT 30 Gy/5 fr | Median 15.8 m | 12 m 94%  | NA         |
| Bodensohn et al. [32] (2021) | 65              | 254 (range 2–12)     | SRS 15–20 Gy | Median 15 m     | 12 m: 97.5% | Grade 2: 6.2%, Grade 3: 4.6% |
| Kim et al. [36] (2021)    | 40                 | 252 (range 4–10)     | SRS: 22.7 Gy/1 fr. (24 pt.) SRT: 29.0 Gy/5 fr. (16 pt.) | Median 8.5 m  | 12 m: 95%  | No grade 3–5 |

SI-STI-MBM single isocenter stereotactic irradiation for multiple brain metastases, LINAC linear accelerators, fr. fractions, SRT stereotactic radiosurgery, SRS stereotactic radiosurgery, NA not available, pt. patients
Specific consideration points in SI-STI-MBM; offset and small fields

There are two major physics issues in SI-STI-MBM. First, all lesions or all except one target must be offset in SI-STI-MBM. Translational and rotational errors in offset lesions result in larger positional displacement and dose coverage impairment as the distance between the isocenter and center of the target increases [44–46]. Therefore, correction of these errors should be considered in offset lesions in SI-STI-MBM. To minimize these errors, 6D positioning correction using ExacTrac X-ray system after every couch rotation can contribute to highly accurate positioning within a short time. Tsuruta et al. reported that there was no significant difference in GTV D_{99.5%} and D_{0.5%} despite the distance between the target and the isocenter in SI-STI-MBM with VMAT when 6D positioning correction was performed using the ExacTrac X-ray system [47]. With regard to adding a margin to the GTV to compensate for these errors, 1 mm seems to be sufficient from the point of view of physical aspects [46, 48]. Kraft et al. reported that clinical outcomes of SI-STI-MBM using 6D positioning correction with a 1 mm margin added to create PTV [37]. In their report, the local control rate at 12 months was 94%, and the distance between the isocenter and tumors had little effect on local control. These data imply that the 6D positioning correction and a 1 mm margin added to PTV made the translational and rotational errors clinically acceptable. Based on these reports, there is little need to give a larger margin in SI-STI-MBM when compared with conventional STI, as long as proper setup and position correction are performed.

Second, the inaccuracy of small fields generated by the optimization of SI-STI-MBM with VMAT should be considered. The leaf motion in SI-STI-MBM with VMAT is complex, and many small fields are generated for multiple targets. Research on small fields of stereotactic irradiation is ongoing, and information including ICUR 91 supports the QA of stereotactic irradiation for small tumors [49, 50]. Although much effort has been made to establish QA in SI-STI-MBM with VMAT, the uncertainty in small-field irradiation cannot be ignored. Therefore, clinical outcomes should be analyzed and confirmed in short- and long-term follow-ups at each institution.

Prospects in SI-STI-MBM

Additional evidence is required to evaluate the usefulness of stereotactic irradiation, including SI-STI-MBM. Here, we addressed the prospects of stereotactic irradiation, not limited to SI-STI-MBM, but also including conventional LINAC-based stereotactic irradiation and GKS for multiple brain metastases.

First, there is currently limited high-level evidence that directly compares WBRT and stereotactic irradiation in patients with 5–10 brain metastases. Although JLGK0901 showed the usefulness of stereotactic irradiation in patients with 5–10 brain metastases, there were no randomized controlled trials (RCTs) comparing WBRT versus stereotactic irradiation for more than three metastases, before the publication of the Dutch phase III RCT reported by Hartgerink et al. [51]. In a Dutch Phase III RCT, WBRT and stereotactic irradiation were compared in patients with 4–10 brain metastases, and the primary endpoint was quality of life (QOL) 3 months after completion of radiotherapy. Although the planned sample size was 230 (115 patients per group), patient recruitment was poor, and the study included 29 patients. The primary endpoint was not met, and there was no difference between the two groups, while the statistical power was weak due to poor accrual. The preference of radiation oncologists, patients, and referrers negatively affected patient recruitment in this clinical trial. Therefore, it may be challenging to conduct RCTs comparing WBRT and stereotactic irradiation for limited brain metastases. However, two RCTs are currently underway to investigate the feasibility and merits of stereotactic irradiation over WBRT in patients with more than four brain metastases. One RCT was the CE.7 study conducted by the Canadian Cancer Trials Group, which compared stereotactic irradiation with hippocampus-avoidance WBRT at 30 Gy in 10 fractions (ClinicalTrials.gov number NCT03550391). The number of brain metastases ranged from five to 15, and stereotactic irradiation was performed in a single fraction. The primary endpoint is overall survival (OS). Another clinical trial, which also compared stereotactic irradiation and WBRT, is now being conducted at the Dana-Farber Cancer Institute and Brigham and Women’s Hospital, and patients with 5–20 brain metastases are registered in this trial (ClinicalTrials.gov number NCT03075072). The primary endpoint was QOL 6 months after the completion of radiotherapy.

Second, the adaptation of SI-STI-MBM for more than 10 brain metastases is unclear. From a technical perspective, SI-STI-MBM can be adapted for more than 10 brain metastases. However, the superiority or noninferiority of stereotactic irradiation over WBRT in patients with more than 10 brain metastases remains unknown. These two RCTs included patients with more than 10 brain metastases. If these two trials are completed with sufficient statistical power, the treatment strategy for more than 10 brain metastases may change.

Finally, brain metastases from small-cell lung carcinoma are controversial in terms of the benefits from stereotactic irradiation. This is because brain metastases from small-cell lung carcinomas tend to occur more frequently than those
from nonsmall-cell lung carcinomas. Therefore, WBRT is the standard treatment for brain metastases from cell lung carcinoma. Recently, a multicenter retrospective cohort study named the FIRE-SCLC cohort study was published [52]. In this study, 710 patients treated with stereotactic irradiation without WBRT or prophylactic cranial irradiation from 1994 to 2018 were included. The median OS was 8.5 months, and it was similar between patients with 2–4 brain metastases and patients with 5–10 brain metastases, consistent with the JLGK0901 study [25]. Although the FIRE-SCLC cohort study is just a retrospective study, the clinical results of stereotactic irradiation seem clinically acceptable and suggest the feasibility of stereotactic irradiation as a reasonable treatment option for brain metastases from small-cell lung carcinoma. Moreover, two ongoing phase II studies investigate the feasibility of stereotactic irradiation in patients with brain metastases from small-cell lung carcinoma in Germany and the USA (ClinicalTrials.gov numbers NCT03297788 and NCT03391362). These clinical cases include patients with 1–10 brain metastases. In addition, Gondi et al. conducted a phase III trial comparing hippocampal-avoidance WBRT and stereotactic irradiation for 10 or fewer brain metastases from small-cell lung cancer (NRC CC009, ClinicalTrials.gov number NCT04804644). If the benefit of stereotactic irradiation is shown from these clinical trials, not only conventional stereotactic irradiation but also SI-STI-MBM can be a useful treatment in patients with multiple brain metastases from small-cell lung cancer.

Conclusion

As the prognosis of patients with brain metastases improves, stereotactic irradiation, rather than WBRT, has been performed in many patients to achieve long-term local control and avoid neurocognitive dysfunction due to radiotherapy. Recent advances in radiotherapy devices, treatment-planning systems, and image-guided radiotherapy have enabled SI-STI-MBM to deliver ablative high doses to multiple brain metastases within a short treatment time. In SI-STI-MBM, local control is excellent and the rate of adverse events, including radionecrosis, is tolerable. As SI-STI-MBM uses only one isocenter and many targets are offset, clinicians, physicists, and radiotherapists should reduce translational and rotational errors as much as possible by using a highly precise image-guided technique, including 6D positional correction. High-level evidence regarding stereotactic irradiation for more than 11 brain metastases and brain metastases from small-cell lung cancers will be shown in ongoing clinical trials.

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Declarations

Conflict of interest The authors declare that they have no conflict of interest.

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