Malignant soft tissue sarcoma of the shoulder treated by surface mould brachytherapy boost in an adjuvant setting

Ashutosh Mukherji, MD1, Mourougan Sinnatamby, PhD2
1Department of Radiotherapy, 2Department of Medical Physics, Regional Cancer Centre, JIPMER, Puducherry, India

Abstract

Purpose: Soft tissue sarcomas of the extremities account for half of all soft tissue sarcomas. Radiotherapy and surgery have been the standard modalities in the treatment of this type of cancer. Brachytherapy can be used as the sole therapy, if the target volume is localized and easily accessible. This work reports three cases of shoulder soft tissue sarcomas with positive deep resected margins, treated with a combination of external beam radiotherapy and surface mould brachytherapy boost technique.

Material and methods: Between January and June 2014, three patients received brachytherapy with sites close to the shoulder, and post-surgery involved deep resected margins. Each mould was made on a base of thermoplastic, over which dental wax was coated and catheters implanted. The target volume was defined as the tissue covering the tumor bed with lateral margins of 2-2.5 cm and depth of 1-1.5 cm. Treatment planning was computed tomography-based and dose prescribed was 85-100% isodose. Treatments has been delivered twice daily, six hours interval, and a review of reactions evaluated.

Results: Volume receiving more than 150% of the prescribed dose has been limited to less than 2%, and that above 200% to be inside the mould. Brachytherapy equivalent dose at 2 Gy per fraction (EQD2) of these patients was 24 and 28.6 Gy. Maximum dose to organ at risk (OAR) (2 cc of OAR) ranged between 55-87% of prescribed dose, with a median dose being 80%. All cases had only grade 1 post-radiotherapy skin immediate reactions, which resolved within four weeks. In all patients, no treatment failures were noted at nearly 2-years post-irradiation.

Conclusions: Surface mould brachytherapy in soft tissue sarcomas could be a useful alternative to interstitial brachytherapy, especially where the target volume is superficially extensive with underlying critical structures, and where catheter placement may be difficult, such as the shoulder.

Key words: brachytherapy, soft tissue sarcoma, surface mould.

Purpose

Soft tissue sarcomas (STSs) of the extremities are not commonly seen, and usually occurs in less than 1% of all adult tumors, but make up nearly half of all soft tissue sarcomas [1]. Many patients present with a progressively enlarging painless swelling [1,2]. Symptoms like local pain, neuralgias, or restriction of motor function can result later due to the involvement of nerve bundles, bone, or significant muscular compartment [2]. Radiotherapy and surgery have been the standard modalities in the treatment of extremity soft tissue sarcomas. Tumor grade is considered one of the main prognostic factors. Low-grade tumors predominantly have the risk of local recurrence, requiring only adjuvant radiotherapy. On the other hand, most patients with intermediate or high-grade tumors larger than 5 cm in size, develop distant metastases compared to less than 10% in low-grade lesions or less than 5 cm in size [3,4].

Adjuvant radiotherapy either used as interstitial brachytherapy or external beam radiotherapy (EBRT) has been found to improve local control rates (LC) in patients who have undergone limb-sparing resections with Level 1 evidence available [5,6]. For completely resected high-grade sarcomas of the extremities, brachytherapy can be used as the sole therapy if the target volume is localized and accessible, as it gives high doses to the tumor bed and extremely low doses to tissues outside the treated volume [7]. Data also suggests improved outcome with a combination of brachytherapy (interstitial) and EBRT for patients with positive margins [8,9,10].

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Clinical and treatment details

Case history (Table 1)

Case 1

A 48-year-old female patient was referred to the Radiotherapy Department for post-operative radiotherapy for a soft tissue sarcoma of the right shoulder. She presented with a history of a swelling developed over a six-month period (size and surgery details are not available).

Case 2

A 45-year-old female patient was referred to the Radiotherapy Department for post-operative radiotherapy for a soft tissue sarcoma of the left chest wall. She presented with a swelling over the left upper part of the chest in the infra-clavicular region developing over three years, and was operated at another hospital in January 2014 (size and surgery details are not available).

Case 3

A 36-year-old male patient has been referred to the Radiotherapy Department for post-operative radiotherapy of soft tissue sarcoma of the anterior shoulder at left side. He had presented in another hospital with a swelling of the left anterior shoulder over ten months. Local examination reported an 8 x 6 cm mass lesion with no nodes or restriction of mobility. The patient underwent surgery. The post-surgery histopathological review reported the mass as a low-grade sarcoma with fibro sarcomatous pattern and with involved deep margin. The patient then received external beam radiotherapy with 6 MV, using AP-PA portals with the use of bolus of 0.5 cm over surgical scar to a prescribed dose of 40 Gy in 22 fractions. This was followed by a boost, with surface mould brachytherapy to a dose of 3.5 Gy per fraction, twice daily for 7 fractions (Figure 3). The patient completed treatment in April 2014.

Case 2

A 45-year-old female patient was referred to the Radiotherapy Department for post-operative radiotherapy for a soft tissue sarcoma of the left chest wall. She presented with a swelling over the left upper part of the chest in the infra-clavicular region developing over three years, and was operated at another hospital in January 2014 (size and surgery details are not available). A post-surgery histopathological review reported dermatofibrosarcoma protuberans with deep resected margin involved by tumor. The patient was subsequently treated by external beam photon irradiation with 6 MV photons, using conventional AP-PA portals with lung shielding and use of bolus (of 0.5 cm thickness) over surgical scar to a prescribed dose of 40 Gy in 20 fractions. After EBRT, a boost with surface mould brachytherapy using a base of thermoplastic immobilizer with wax mould was given (Figures 1 and 2). The patient received brachytherapy boost with dose of 3.5 Gy per fraction, twice daily, for seven fractions. Treatment was completed in February 2014.

Case 3

A 36-year-old male patient has been referred to the Radiotherapy Department for post-operative radiotherapy of soft tissue sarcoma of the anterior shoulder at left side. He had presented in another hospital with a swelling of the left anterior shoulder over ten months. Local examination reported an 8 x 6 cm mass lesion with no nodes or restriction of mobility. The patient underwent surgery. The post-surgery histopathological review reported the mass as a low-grade sarcoma with fibro sarcomatous pattern and with involved deep margin. The patient then received external beam radiotherapy with 6 MV, using AP-PA portals with the use of bolus of 0.5 cm over surgical scar to a dose of 44 Gy in 22 fractions, followed by a boost with surface mould brachytherapy to a dose of 3.5 Gy per fraction, twice daily for 7 fractions (Figure 3). The patient completed treatment in April 2014.

Table 1. Patient characteristics

| Total number of cases | Number |
|-----------------------|--------|
| Males                 | 1      |
| Females               | 2      |
| Age group (years)     |        |
| 31-40                 | 1      |
| 41-50                 | 2      |
| ≥ 51                  | 0      |
| Site involved         |        |
| Site involved         | 0      |
| Chest wall            | 1      |
| Shoulder              | 2      |
| Type of lesion        |        |
| Primary               | 2      |
| Recurrent             | 1      |

Fig. 1. Upper torso mould used in the first patient with extensive involvement of right shoulder tissue.
fraction twice daily for six fractions. The patient completed treatment in June 2014.

**Mould preparation**

Each mould was prepared from a framework of thermoplastic sheet molded to the patient’s anatomy, and then covered by thin layer of dental wax. After marking of target volume by the oncologist, the physicist helped to make an initial cut-out with thermoplastic sheet. Then the wax was heated in a water heater, and cast in the required shape over the thermoplastic and molded according to local anatomy. The shaped wax mould was approximated to the lesion in the treatment position, matched with lesion surface and local anatomy, edges were trimmed and smoothened. Catheters were fixed to the mould, which was then placed on the patient to check the feasibility of placement and target volume coverage (Figure 1). The lesion was then marked with lead wires and computed tomography (CT)-simulation of the patient in the treatment position was done with mould in place.

**Treatment planning and delivery**

After CT-simulation, image reconstruction, target volume contouring, and treatment planning was carried out on Eclipse (Varian Medical Systems, Palo Alto, CA, USA) TPS version 10.0, and treatment delivery was done on GammaMedplusiX (Varian Medical Systems, Palo Alto, CA, USA). The target volume was defined as the volume encompassing the tumor bed with lateral margins of 2-2.5 cm and depth of 1-1.5 cm. The tumor bed has been identified by comparing pre-operative and post-operative CT/magnetic resonance (MR) films, as well as taking into account the surgical scar. Dosimetry has been done using volume optimization techniques. The dose was prescribed to 80%, 85%, and 90% isodose lines, respectively, in the three cases in order to cover the depth of 1-1.5 cm (Figures 2, 3 and 4). Treatments were delivered twice daily, six hours interval, and review of reactions was done every third day and at the completion. Evaluation was done of planning details, response to therapy, and reactions encountered.

**Observations**

All patients tolerated the mould boost well with skin darkening as observed towards the end of treatment, and grade 2 skin changes seen in the second and third patients. Grade 1 skin changes were seen in all three cases (Figure 5). Brachytherapy dose ranged from 21-24.5 Gy in frac-

![Fig. 2. A) Site of mould in the first patient. B) Computed tomography image with mould in place in the first patient](image1)

![Fig. 3. A) Dose color wash and B) 3D dose reconstruction of treated area in the second patient](image2)
Fig. 4. A) Dose color wash in the second patient. B) Isodose distribution with overlying catheters in the third patient

Fig. 5. Immediate post-radiotherapy skin changes in the third patient

Fig. 6. Biologically equivalent dose (BED) of treated cases

Table 2. Dose characteristics

| Age/sex | Brachytherapy dose | EBRT dose | No. of channels | EQD2 cGy | Brachytherapy dose |
|---------|--------------------|-----------|----------------|----------|-------------------|
|         | Total dose (Gy)    | Fractions | Total dose (Gy) | Fractions | BED3Gy                  |
|         |                    |           |                |          | 53.17                   |
| 48/F    | 24.5 (3.5)         | 7         | 40 Gy         | 20       | 53.17                   |
| 45/F    | 24.5 (3.5)         | 7         | 40 Gy         | 20       | 53.17                   |
| 36/M    | 21.0 (3.5)         | 6         | 44 Gy         | 22       | 56.07                   |

EQD2 – equivalent dose at 2 Gy per fraction, BED3Gy/BED10Gy – biologically equivalent dose for late effects (BED3Gy) and for early effects (BED10Gy).

Table 3. Dose volume comparisons

| Case number | Age/sex | Prescription isodose to PTV | V200% | V150% | Depth of PTV coverage | Lateral margins for PTV | % PTV covered by prescription isodose |
|-------------|---------|-----------------------------|-------|-------|-----------------------|------------------------|--------------------------------------|
| 1           | 48/F    | 80%                         | 0.001%| 1.0%  | 1.1                   | 2.0 cm                 | 100                                  |
| 2           | 45/F    | 85%                         | 0.001%| 0.5%  | 1.2                   | 2.5 cm                 | 80                                   |
| 3           | 36/M    | 90%                         | 0.001%| 0.6%  | 1.6                   | 2.0 cm                 | 70                                   |

PTV – planning target volume, V200% – volume receiving 200% of prescribed dose, V150% – volume receiving 150% of prescribed dose.
tumor volume coverage, while at the same time, keeping volume of tissue receiving more than 150% ($V_{150}$) to below 2% and volume of tissue receiving more than 200% ($V_{200}$) of prescribed dose to below 0.001%. This meant that areas receiving more than 200% of the prescribed dose were limited to the substance of the mould (Table 3 and Figure 7). The underlying bone in all cases was the organ at risk (OAR). Maximum dose to OAR was assessed as dose to 2 cc of OAR and ranged between 55-87% of the prescribed dose with a median dose being 80% (Figure 8).

**Sequelae and survival**

There were no severe skin reactions like ulcers or necrosis (Table 4). At six months of follow-up, none of the patients showed any local or distal recurrence both clinically as well by imaging (CT thorax and local MRI). All the cases again reported for follow-up in March 2015, in August 2015, in January 2016, July 2016, October 2016, and January 2017 (Figures 9 and 10). By then, patients had completed 36, 34, and 32 months uneventful follow-up, respectively. Local examination and imaging showed no local recurrence or distant metastases or even no local site late treatment sequelae (Figure 11, Table 4).

**Discussion**

Post-operative or adjuvant radiotherapy in soft tissue sarcomas are indicated in situations with increased risk of local recurrence after surgery. These include cases with narrow or positive surgical margins, local recurrence after prior surgery, tumor size of more than 5 cm, as well as lesions deep to or invading the superficial fascia, high grade, or age of patient younger than 50 years [3].

Interstitial brachytherapy has been more commonly used and can achieve good dose distribution especially in limited target volumes, and helps to achieve good control rates [11,12]. Early results from Memorial Sloan-Kettering Cancer Center (MSKCC) have shown better results when the brachytherapy is added to radiotherapy as adjuvant therapy, especially in cases with positive margins [8] as well as in recurrent disease [9,13]. In France, the Gustav Roussy Institute reported 80% local control rates with interstitial brachytherapy, but 30% margin failures and a 35% necrosis rate [14]. However, the use of interstitial brachytherapy in the treatment of sarcomas is associated with certain constraints. These include difficulty in covering a larger target volume, limits in catheter patterns, achieving good geometry because of the presence of bone or visceral organs or difficult anatomic sites...
such as around the shoulder, and risk of radiation injury to nerves that are in direct contact with the BT catheters. The MSKCC group noticed that shoulder location was an independent prognostic factor for poor local control with interstitial brachytherapy [7]. Therefore, in many of these situations, particularly when the target volume is superficially located, a customized mould using acrylic or thermoplastic casts can be used for catheter placement, geometry, and even the immobilization. The brachytherapy catheters are laid directly onto the mould or can be used with a spacer such as in Freiburg flap to ensure uniform catheter distribution.

Compared to interstitial brachytherapy, surface mould brachytherapy though extensively studied in skin cancers, head and neck, scalp, or even intra-ocular regions, has not been widely used for treating extremity soft tissue sarcomas, and dose guidelines have to be extrapolated from interstitial brachytherapy. These custom-made external molds can result in dose distribution with a sharp gradient in the edges of the applicator, especially over curved or irregular surfaces, and are easy to use with highly accurate daily treatment reproduction. They enable one to obtain excellent local control with minimum treatment-related sequelae or late complications. In our case series, the position of the tumor bed in all the three cases was very close to the head of humerus and with deep involved margins, and the need was not only to adequate cover the area at risk, but also to achieve sharp dose fall-off to avoid the joint or bone. It is better to prescribe to isodose of 85-100% to obtain an acceptable coverage of the target volume to limit the amount of tissue irradiated by 150% isodose. Any attempt to pull the isodose outward to cover a bigger target volume altogether could result in more tissue receiving doses above 200% and cause severe skin reactions. In all our cases, the authors managed to keep the area receiving dose of 200% and above to within the substance of the mould, while area receiving above 150% is less than 2%. Thus, the use of surface moulds can result in dose distributions more homogeneous than even electron beam therapy [15], and may even rival IMRT with the added benefits of minimal surrounding tissue dose and much less integral body dose [16]. In our series, we noticed grade 1 skin changes in all cases and grade 2 reactions in 2 cases. Furthermore, all reactions healed within six weeks.

Conclusions and learning points

Treatment of soft tissue sarcomas of the extremities requires usually combined modality therapy. Brachytherapy, usually interstitial, plays an useful role in the treatment of soft tissue sarcomas of the extremities helping achieve improved local control with both organ and function preservation.

Surface mould brachytherapy can be a valuable alternative to interstitial brachytherapy, especially in cases where the target volume is extensively superficial, underlying critical structures, and where catheter placement may be difficult, such as the shoulder, with excellent medium term local control and mild toxicity.

Disclosure

Authors report no conflict of interest.

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Table 4. Skin reactions and survival

| Case number | Skin reactions grade at 4 weeks post-brachytherapy (acute reactions) | Skin reactions/scar/fibrosis at 30 months post-brachytherapy (late effects) | Follow-up period (in months) | Disease free survival (in months) |
|-------------|-------------------------------------------------------------|-------------------------------------------------|-----------------------------|-------------------------------|
|             | Grade 1 | Grade 2 | Grade 3 | Grade 4 | Grade 1 | Grade 2 | Grade 3 | Grade 4 |                             |                             |
| 1           | Yes     | No      | No      | No      | No      | No      | No      | No      | 36                           | 36                           |
| 2           | Yes     | No      | No      | No      | No      | No      | No      | No      | 34                           | 34                           |
| 3           | Yes     | Yes     | No      | No      | No      | No      | No      | No      | 32                           | 32                           |

Fig. 11. Disease free survival (DFS) of treated cases
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