Introduction

Malignant fibrous histiocytoma, liposarcoma, and rhabdomyosarcoma are the three most common types of sarcoma, and synovial sarcoma—a rare and aggressive soft tissue sarcoma—is the fourth most common, accounting for 5–8% of soft tissue sarcomas. It typically develops in young adults as well as adolescents. Despite its nomenclature, synovial sarcoma lesions are not usually intra-articular but instead develop near tendon sheaths, bursae, and joint capsules (1). Signs or symptoms may not be noticeable in the early stages; however, with an increase in tumor size over time, individuals with synovial sarcoma may discover that they have a lump or swelling. For some individuals, range of motion may be limited by the mass and numbness and/or pain may be caused due to the compression of nerves in the locality. The exact cause of synovial sarcoma has not yet been identified, but the disease’s development is believed to be affected by genetic factors. Translocation t(X;18) characterizes synovial sarcoma cytogenetically; it is specific to this cancer type and frequently employed for diagnostic purposes. Although surgery and radiation therapy can result in comprehensive local control, survival is still limited by the prevalence of distant metastasis. Among those who have ≥5 cm synovial sarcoma in a primary extremity, distant metastasis is discovered in more than half of patients, and mortality occurs within 5 years in nearly two-fifths of such patients (2,3). A combination treatment is generally employed that comprises surgery, radiation therapy, and/or chemotherapy (4). Even if initial treatment is aggressive, with a wide surgical resection and multiple adjuvant treatments, local recurrence and metastasis often occur and oncologists are cautious in their prognoses.

Case presentation

This case report relates to a 22-year-old male patient with synovial sarcoma, who had a painful mass in right thenar region that he had been aware of for approximately 3 years before he sought medical treatment. Magnetic resonance imaging (MRI) showed a nodular mass localized at the volar aspect of the palm between the first and second finger, with the mass measuring 5×3×3 cm (Figure 1). At that time, chest computed tomography (CT)-scan was negative. He was treated with en bloc resection on September 21, 2017. Histologically, epithelial and spindle-cell components were present in varying proportions. The epithelial cells were cuboidal and had glandular areas, the epithelial cells were cuboidal and had
ovoid vesicular nuclei and abundant, palely eosinophilic cytoplasm. The spindle cells had sparse cytoplasm, ovoid and bland but also hyperchromatic nuclei, and inconspicuous nucleoli (Figure 2). FNCLCC (Fédération Nationale des Centres de Lutte Contre le Cancer) Grade 2 with a total score of 5 was obtained. The surgical margin was considered to be positive due to fragmented tissue.

Immunohistochemical stains revealed strong positivity for transducer-like enhancer of split 1 (TLE1) and focal positivity for cytokeratin (AE1 + AE3) (Figure 3). Other stains revealed that the patient was negative for CD34, S-100 protein, myeloperoxidase (MPO), and desmin. Afterwards, in an attempt to gain local control, the patient was referred to the radiation oncology department for radiotherapy.

Thermoplastic cast was used for immobilization of the patient during radiotherapy. We outlined the index finger and the thumb of the patient for each CT slice scan; therefore, the compartment was assumed to be the clinical target volume (CTV). The CTV was isotropically extended by 1 cm to define the planning target volume (PTV) (Figure 4). Synovial sarcoma is known to sometimes invade skin tissues. For routine skin dose assessment, Gafchromic EBT3 film measuring 3×3 cm² was used for in vivo dosimetry to verify the dose to skin. In this case, the patient’s right hand was immersed in a water tank and subjected to irradiation using a bilateral photon beam of energy 6 MV (Figure 5). Thirty fractions of 2 Gy were administered, resulting in a

![Figure 1](image1.png)  
**Figure 1** Contrast-enhanced T1-weighted MRI image. MRI, magnetic resonance imaging.

![Figure 2](image2.png)  
**Figure 2** Hematoxylin and eosin staining. (A) ×200; (B) ×400.

![Figure 3](image3.png)  
**Figure 3** Tumor cells are (A) positive for TLE1 and (B) positive focally for cytokeratin (IHC staining, ×400). TLE1, transducer-like enhancer of split 1.
total dose of 60 Gy. The Philips pinnacle version 9.8 was the treatment planning system employed. We selected the palm (point A) and back of the hand (point B) as the two points for assessment and repeated the measurement once daily for 3 days. The average skin dose at points A and B was 220.9±2.2 and 219.6±1.2 cGy, respectively. The planned doses at points A and B were 216.2 and 217.0 cGy, respectively; thus, the actual delivered treatment dose were 1.9% and 2.2% higher than the planned amounts. This study demonstrated that employing a water tank to act as a tissue compensator results in skin tissues receiving sufficient doses. The patient tolerated the treatment very well, and grade I dermatitis was the only observed side effect. MRI performed 4 months after completion of radiotherapy indicated a 0.6 cm nodule at the volar aspect of the palm, which was not the original site of the disease. Consequently, further MRI study is indicated for confirmation of recurrence.

**Discussion**

The optimal treatment of synovial sarcoma is undefined at the present time, but *en bloc* tumor resection is the current first-choice treatment. A minimum requirement of surgical treatment is that the pathologic specimen must be tumor-free at the margin. The difference between complete local resection and wide tumor resection appears to significantly influence outcome (5). Noncurative surgery—for example, intralesional excisions or an incomplete gross resection—should have no part in treatment (6,7). Surgery is nonetheless the most commonly employed modality for treatment of locally recurrent disease, as it is for treatment of primary disease (8,9). Adjuvant radiation therapy is employed in synovial sarcoma under the same rationale as it is employed in soft tissue sarcomas (6,7). Adjuvant radiation therapy modalities include brachytherapy, conventional two-dimensional radiotherapy, and intensity-modulated radiation therapy. The local control rate in patients with high-grade sarcomas such as synovial sarcoma has been demonstrated to be higher when radiation therapy is employed, irrespective of the radiation therapy type (10,11). The various international guidelines have inconsistent recommendations regarding radiotherapy dose. In one study, 61.2 Gy (range, 45–66.6 Gy) was discovered to be a sufficient dose in adjuvant radiotherapy, with 5-year overall survival being 77% and 10-year overall survival being 65% (12). However, that local irradiation is associated with superior local control of the disease in patients with localized, completely resected tumor and negative surgical margin is yet to be adequately demonstrated. By contrast, radiotherapy appears to significantly reduce local recurrence rates in patients with gross residual tumor after surgery.

![Figure 4](image1.png)

**Figure 4** The PTV (cyan) is expanded from the CTV (red) and extends beyond the patient's body in places. PTV, planning target volume; CTV, clinical target volume.

![Figure 5](image2.png)

**Figure 5** Dose verification and patient positioning. (A) The 3 cm × 3 cm EBT3 film employed for surface dose measurement; (B) water tank for producing a uniform dose.
Regarding treatment order, preoperative radiotherapy appears to result in similar local control and long-term physical function as does postoperative radiotherapy. A crucial effect of timing does exist, however: postoperative wound complications are more likely to occur if radiotherapy is performed preoperatively. A more palliative approach when treating a patient with a nonresectable, localized tumor is hypofractionation—for example, 13 to 15 fractions of 3 Gy—although the suitability of this approach is dependent on factors such as patient age and comorbidities (13). The role of adjuvant chemotherapy in synovial sarcoma—a particularly chemoreactive soft tissue sarcoma—remains controversial (14,15).

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None.

Footnote

Conflicts of Interest: The authors have no conflicts of interest to declare.

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References


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