Sudden Death Due to Sarcoma: An Unusual Case and Literature Review

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Abstract

Background: Sudden death has varied causes, and autopsy is usually required to determine cause of death. Sarcoma involvement is very infrequent, and detailed pathological evaluation is needed for correct diagnosis. When the medical history of the deceased is unavailable and autopsy is not performed, the true cause of death will not be established. Here we present an unusual case of sarcoma involving heart and lung. We also review 13 cases found in the literature with sarcoma involvement in sudden death.

Introduction

A natural but unexpected death happening within 1 hour of the onset of final symptoms is called sudden death [1], and almost 85% of all sudden deaths are of cardiac origin, a leading cause of death in Western countries that is responsible for around 30 to 200 deaths per 100,000 individuals every year [2]. Scene investigation for sudden death requires questioning of the witnesses and family members of the deceased [3]. However, about 40% of sudden death cases are unwitnessed [4], which makes it more difficult to collect pertinent evidence and information. In such cases, forensic autopsy is usually performed; it is also usually required in order to gain any information. Even with autopsy, however, a small proportion of cases will not reveal a cause of death (COD). In these cases, the COD is categorized as “sudden unexplained death” [5,6].

Soft tissue sarcomas are less common than carcinomas. Sarcomas can occur anywhere in the body; common primary sites are an extremity, the trunk, or the retroperitoneum [7]. The number of newly diagnosed cases of soft tissue sarcoma in the United States in 2004 was 8,680, accounting for only 0.63% of all malignant tumors [8]. The rate of sarcoma in all malignancy-related death ranges from 1.5% to 2% [8,9].

Sudden death caused by cardiac involvement in malignancy is uncommon, and primary cardiac sarcoma is also rare [10]. Therefore, rates of sarcoma involving the heart causing sudden death are even smaller [11]. The task of establishing a true COD is more difficult when the medical history of sarcoma is not available, or the autopsy is not performed.
Here we present a case of a 58-year-old man with a history of sarcoma who suffered sudden death, together with a literature review of related cases. The autopsy of the presented case revealed sarcoma involvement in the heart and lung, and immunohistochemical stainings were performed to determine the phenotypes of the sarcoma. The literature research collected 13 additional sudden death cases involving sarcoma.

Materials and Methods

Chemicals and Antibodies

Hematoxylin and eosin for histochemical staining were purchased from Leica Biosystems (Singapore). For immunohistochemical staining, the antibodies cytokeratin and CD31 were purchased from BioGenex (Fremont, CA, USA); CD34, vimentin, h-caldesmon, and myogenin were purchased from Dako (Carpinteria, CA, USA); and 3,3’-Diaminobenzidine Quanto was purchased from Thermo Scientific (Waltham, MA, USA).

Immunohistochemical Staining

Immunohistochemical staining was performed using the Ventana BenchMark autostainer (Tucson, AZ, USA). Antigen retrieval was processed by the autostainer with a routine CC1 course (pH 8.0, 1 h). Antibodies were incubated on the slides in iVIEW-labeled conjugate (Ventana) at 25 °C for 90 min. The slides were then incubated in a solution of hydrogen peroxide and 3,3’-Diaminobenzidine at 25 °C for 8 min, and the sections were counterstained with hematoxylin. The slides were reviewed by two pathologists.

Literature Review

The keywords “sudden death” and “sarcoma” were used for a PubMed search. Cases that were not written in English, did not meet the sudden death definition [1], or did not provide case descriptions were excluded. Thirteen cases were found and included for analysis.

Statistical Analysis

Fisher’s exact test and other statistical calculations, including means and standard deviations, were performed using SPSS 16.0 (IBM, New York, NY, U.S.A.)

Results

Case Presentation

A 58-year-old man living alone with no systemic diseases was found dead face down in his kitchen. His family stated that he had no discomfort in recent days. Four years before his death, he had a wound over his left ankle that did not heal spontaneously. Later, he noticed a protruding mass on the left ankle, but he had not sought medical assistance until 9 months before he died. He then received wide excision, and a pathology report showed high-grade sarcoma. He did not receive regular follow-up or any adjuvant therapy after excision, however. Five months before his death, he noted persistent bleeding over his left lower leg. Local recurrence of sarcoma was suspected and above-knee amputation was performed.

An autopsy was executed to investigate the COD. Height and body weight were 172 centimeters and 70 kilograms. No brain hemorrhage or external traumatic lesions were noted. The weights of the heart and right and left lungs were 590, 750, and 900 grams respectively. Tumor masses were noted in the right ventricle (Figure 1a) and lung (Figure 1b). The outflow of the right ventricle was completely blocked by the tumor mass. The cut surface of the mass was soft, and areas of necrosis were seen. Other organs were not remarkable. The toxicological panels did not indicate use of recreational drugs or poisoning. The blood alcohol level was below the limit of detection. Microscopic findings revealed a spindle cell tumor arranged in a sheet-like pattern. Nuclear pleomorphism was conspicuous (Figure 2). An immunohistochemical study showed positivity for vimentin in the spindle cells (Figure 3a), while other markers, including CD31, CD34, cytokeratin, h-caldesmon, and myogenin, were negative (Figures 3b-f). The COD was sudden death caused by high-grade sarcoma blocking the outflow of the right ventricle.

Literature Review

Thirteen cases of sudden death involving sarcoma were found in the literature. These cases and the presented case are summarized in Table 1. The mean age at death was 30 years (30 ± 8.3), although seven cases happened before adulthood. Of the seven juvenile cases, four were younger than 1 year old, two were between 1 and 6 years old, and one (case 12) was 19 years old. The six adult cases and the presented case (case 14) were all more than 35 years old at time of death.

Regarding medical history, 10 of the 14 cases did not have a history of sarcoma prior to death. The other 4 cases, on the contrary, were diagnosed with sarcoma prior to the sudden death. In the 10 cases without previous sarcoma history, 7 showed primary sarcoma sites in vital organs, including heart, lung, or brain. In the 4 cases with sarcoma history, only 1 case showed a primary site of sarcoma developing in a vital organ. However, the differences in history between these groups were not statistically significant (p = .145).
Figure 1: Macroscopic pictures of tumors in the heart and lung. (a) A grayish-white tumor resided in the right ventricle with infiltration in the myocardium. (b) One of the tumors was located in the right middle lobe of the lung

Figure 2: Microscopic pictures of tumors in the heart and lung (a) Spindle cells with prominent necrosis infiltrates in the myocardium, magnification rate: 40X (b) Prominent nuclear atypia, magnification rate: 100X

Figure 3: Immunohistochemical studies of the sarcoma (a) Vimentin (b) CD31 (c) CD34 (d) Cytokeratin (e) H-caldesmon (f) Myogenin
All cases demonstrated either heart or lung involvement of sarcoma except for case 13, whose primary and only sarcoma involvement was in the brain. Nine out of 14 cases showed cardiac involvement of sarcoma, and 4 out of 14 cases showed lung involvement. The presented case is the only case who manifested both cardiac and lung involvement.

The final diagnoses and grading of sarcoma in these cases were varied. The results suggested that differences in type or grade do not play an important role in sudden death. In diagnosing the types of sarcoma, cases 9 through 13 did not use any immunohistochemical staining. The cases reported after 1994, including our case, all used immunohistochemical panels to achieve the final diagnoses.

### Discussion

Cases of sudden death usually require autopsy to determine COD [5]. However, less than 50% of sudden death cases actually receive autopsy [15]. Among those that do receive autopsy, findings of sarcoma have been reported [12-14] but are very unusual. Moreover, our results show that in most cases of sarcoma causing sudden death, the deceased did not have a medical history of sarcoma, meaning that scene investigation and questioning of family members alone may not provide sufficient information to determine COD. Therefore, promoting an increased rate of autopsy is necessary to establish a correct COD in these cases.

Six out of 14 cases reviewed in our study were younger than 6 years old at time of death. Trutmann and colleagues found that incidence rates of soft-tissue sarcoma peak in two age groups, 10 to 19 years and 70 to 79 years [16], yet both of these age groups are much older than the cases reviewed here. It may be that younger cases with sarcoma, especially children less than 1 year old, have less-favorable survival rates [17]. The idea that these infants or children with sarcoma, therefore, may present as sudden death, is supported by the fact that almost half of the sarcoma-induced sudden deaths reviewed in this study were quite young. These findings may also contribute to future studies in sudden infant death that include malignancy, especially sarcoma, in the differential diagnoses and etiologies [18].

Seven of the 10 cases without medical history of sarcoma had vital organs, including heart, lung, and brain, as the primary site of sarcoma, whereas only 1 of the 4 cases with a history (case 12) had vital organs as primary site. However, the differences between the groups were not statistically significant, most likely due to the small number of cases (p = .145 in the results, but p = .006 by duplicating the cases).

The possible correlation between medical history and vital organs as primary sites may be explained by the fact that cases with primary sites of sarcoma in vital organs are more vulnerable to sudden death, and thus are less likely to be diagnosed before death. Sudden death caused by carcinoma among people without a medical history has been reported, and these were initially presented with vital organ involvement as well [19-21]. Future studies are needed to clarify the correlation between medical history and primary sites of sarcoma.

Five cases in this review, all published before 1994, solely relied on traditional hematoxylin-eosin staining and did not perform immunohistochemical staining to achieve the final diagnoses [22-26]. The other nine cases reviewed here all used immunohistochemical methods to make diagnoses. The adaptation of immunologic methods to histochemical techniques has revolutionized diagnostic pathology, especially for soft-tissue malignancy [27]. Newer immunohistochemical markers facilitate the progress of defining the tumor cell lineage and origin much more accurately than the traditional hematoxylin-eosin method [28-30]. Moreover, molecular diagnostic tools, such as fluorescence in situ hybridization (FISH) and DNA sequencing, are now prevalent in diagnosing sarcoma [31,32]. Using these molecular tools to determine COD is now accepted and can be expected in future studies [33].

<table>
<thead>
<tr>
<th>Case</th>
<th>Age (years)</th>
<th>Gender</th>
<th>Known History</th>
<th>Heart Involvement</th>
<th>Lung Involvement</th>
<th>Diagnosis</th>
<th>Primary Site</th>
<th>IHC*</th>
<th>Ref.</th>
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<tr>
<td>1</td>
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<td>Male</td>
<td>-</td>
<td>+</td>
<td>-</td>
<td>Liposarcoma</td>
<td>Heart</td>
<td>+</td>
<td>[34]</td>
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<tr>
<td>2</td>
<td>&lt; 1</td>
<td>Female</td>
<td>-</td>
<td>+</td>
<td>-</td>
<td>Sarcoma, low grade</td>
<td>Heart</td>
<td>+</td>
<td>[35]</td>
</tr>
<tr>
<td>3</td>
<td>42</td>
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<td>-</td>
<td>+</td>
<td>-</td>
<td>Intimal sarcoma</td>
<td>Heart</td>
<td>+</td>
<td>[36]</td>
</tr>
<tr>
<td>4</td>
<td>77</td>
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<td>+</td>
<td>-</td>
<td>-</td>
<td>Sarcoma, undifferentiated</td>
<td>Urinary bladder</td>
<td>+</td>
<td>[37]</td>
</tr>
<tr>
<td>5</td>
<td>5</td>
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<td>-</td>
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<td>&lt; 1</td>
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<td>-</td>
<td>-</td>
<td>+</td>
<td>Malignant hemangioendothelioma</td>
<td>Thoracic cavity**</td>
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<td>[39]</td>
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<tr>
<td>7</td>
<td>36</td>
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<td>+</td>
<td>-</td>
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<td>Malignant mesenchymoma</td>
<td>Retroperitoneum**</td>
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<td>-</td>
<td>-</td>
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<td>Prostate</td>
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<tr>
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<td>75</td>
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<td>-</td>
<td>+</td>
<td>-</td>
<td>Sarcoma, undifferentiated</td>
<td>Heart</td>
<td>-</td>
<td>[26]</td>
</tr>
</tbody>
</table>
Here we have presented an unusual case of sarcoma causing sudden death. We reviewed 13 cases in the literature. The results showed that no specific type of sarcoma diagnosis occurred more frequently in these cases, neither did the grading of sarcoma have a significant effect. It seemed that in the cases without a medical history of sarcoma prior to the sudden death, the origin of the sarcoma may be more likely to reside in vital organs, but again, the result was not statistically significant. More studies and an increase in the rate of autopsy will help shed light on the relationship between the history and primary site of sarcoma in the sudden death cases.

**Conclusion**

Here we have presented an unusual case of sarcoma causing sudden death. We reviewed 13 cases in the literature. The results showed that no specific type of sarcoma diagnosis occurred more frequently in these cases, neither did the grading of sarcoma have a significant effect. It seemed that in the cases without a medical history of sarcoma prior to the sudden death, the origin of the sarcoma may be more likely to reside in vital organs, but again, the result was not statistically significant. More studies and an increase in the rate of autopsy will help shed light on the relationship between the history and primary site of sarcoma in the sudden death cases.

**References**


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