Correlation of oncologic long-term results and genetic instability in soft tissue sarcomas

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Abstract: Purpose: Soft tissue sarcomas (STS) represent a heterotop group of tumours. Microsatellite instabilities (MSI) and loss of heterozygosity (LOH) as phenomena of a genetic instability should be analysed in STS and correlated with the long-term oncologic outcome. Methods: Patients treated for a STS with a follow-up of at least 10 years were included. Thus, 86 patients (mean age 50.5 years, range 16-86 years) treated for a STS between 1993 and 2000 were routinely controlled every 6 months. Incidence of local recurrences, distant metastases, and overall survival were analysed. Sixty-six tumour samples were available for microsatellite analysis using the former traditional method of PCR amplification at 6 loci in the neighbourhood of hMSH2, hMLH1, p53, p16, rb1, and hTR. Results: There were 30 low-grade and 56 high-grade sarcomas. The mean follow-up was 144 months (120-192 months). Twenty-nine patients died of their disease. Local recurrences were seen in 13 patients, whereas metastases were noticed in 23 patients. The overall survival was dependent on the tumour stage (p<0.05), whereas the local tumour control (incidence of local recurrence) was influenced by the surgical margin achieved (p<0.05). The molecular biologic findings revealed 67% of the investigated loci as informative. MSI was found in 6.8% of the informative loci, whereas LOH in 18.8%, respectively. LOH was present in high-grade tumours in 23.8%, whereas in 1.7% in low-grade tumours. In high-grade sarcomas, the 5-year and 10-year survival probabilities were significantly lower in LOH-positive tumours (48.6% and 38%) than in LOH-negative tumours (72.5% and 62%). Conclusion: The overall survival in soft tissue sarcoma is mainly influenced by the tumour stage. In high-grade sarcomas, the survival rate will drop even after 5 years. The detection of loss of heterozygosity represents a negative prognostic predictor in high-grade sarcomas. Microsatellite instability is a rare phenomenon supposing no relevance in the oncogenesis and tumour progression of soft tissue sarcomas.

Keywords: Soft Tissue Sarcoma, Oncologic Result, Genetic Instability, Microsatellite Instability

1. Introduction

Soft tissue sarcomas (STS) are rare with an incidence of 20 – 25 per million. There are more than 50 histologic subtypes of STS. Although there exists a line of differentiation in most STS, many of these tumours show heterogeneous morphology and multiple, often complex, cytogenic aberrations [1, 2].

There exist inconsistent molecular biological data concerning different types of STS [3 - 9].

Further classification and elucidation of the molecular pathogenesis of STS is highly desirable. Genetic instability is a paramount feature of cancer which leads to accumulation of genetic alterations that varies from subtle changes in DNA sequence to chromosomal abnormalities [1, 2, 10].

Microsatellite instability (MSI) is a particular type of genetic instability affecting short sequences of DNA repeats found throughout the genome [10]. It is considered as the appearance of replication errors which present mutations close to the genes of the mismatch repair (MMR) family [5, 6, 11 - 15]. The relevance of MSI is well known in different cancers, especially in the hereditary nonpolyposis colorectal cancer (HNPCC) [16, 17].

The loss of heterozygosity (LOH) of different tumour suppressor genes is considered to be a further well-
established marker of the genomic instability in different types of cancers [1, 2, 16, 18].

But there exist different statements in the literature about the relevance of MSI and LOH in sarcomas. Ewing’s sarcoma, rhabdomyosarcoma, liposarcoma, and leiomyosarcoma are the most frequently examined sarcomas [4, 9, 14, 15, 19 - 27]. Microsatellite analysis is mostly reported at the neighborhood of the tumour suppressor genes (p53, rb1, hTR, p16) and of the mismatch repair genes (hMLH1, hMSH2) [5, 6, 11 - 15]. Mostly due to small series the role of genetic instability in sarcomas remains unclear, especially concerning the clinical outcome and the biological behaviour [3, 7 - 9, 21].

Most of the clinical studies of patients with soft tissue sarcomas cannot be assessed concerning the oncologic long-term outcome due to the mostly limited follow-up’s of five years. Studies with real long-term results of oncological outcome are rare.

After the promising results of studying genetic instability in the defined entity of intraosseous leiomyosarcoma [9], the aim of the present study was to evaluate the occurrence of genetic instability in different STS. The correlation with the oncologic long-term outcome was established by using the findings of patients with a follow-up of at least 10 years.

2. Patients and Methods

2.1. Patients and Tumour Samples

From January 1993 to December 2010, 203 cases of soft tissue sarcomas located at the trunk or the extremities were treated surgically. To get information about the long-term prognosis, patients with a minimum follow-up of 10 years were selected. Thus, patients treated between 1993 and 2000 were included in this study.

There were 86 patients (51 men, 35 women) with an average age of 50.5 years (range, 16-86 years). The different histologies of STS include malignant fibrous histiocytoma (MFH) (n = 26), liposarcoma (n = 19), leiomyosarcoma (n = 12), fibrosarcoma (n = 8), rhabdomyosarcoma (n = 7), synvialsarcoma (n = 6), and others (n = 8).

Concerning to the Enneking’s classification [28], there were 30 (34.8%) low-grade (stage I) and 52 (60.4%) high-grade (stage II) tumours. In 4 cases distant metastases had to be observed (stage III).

The lower extremity was involved in 44 cases, the upper extremity in 26 cases, and the trunk in 16 cases, respectively.

In 35 (40.7%) of the patients the tumour was primarily resected under the suspicion of a benign soft tissue tumour, in 31 of these cases the tumour was localized extracompartimentary. The histopathological examination revealed a sarcoma with an intraslesional resection. These patients were admitted for further surgery to achieve a wide resection. The median duration between previous intraslesional surgery and admission was 5.2 weeks (range, 2 to 16 weeks). There were 19 stage-I tumours, and 16 stage-II tumours in this group.

In 20 out of these patients we did not found any tumour tissue after the required secondary surgical procedure to achieve a wide margin. Thus, tumour samples were available only in 66 cases for microsatellite analysis. All patients were informed that their data would be submitted for publication, and gave their consent.

2.2. Oncologic Follow-Up

All patients were routinely controlled clinically every 6 months up to five years postoperatively. Afterwards the patients were controlled yearly. The survival rate and the occurrence of local recurrence and metastases were determined. The survival probability was estimated by the method describe in [29]. The tumour stage, the tumour volume, the achieved surgical margin [28], and the fact of intraslesional previous surgery were also analysed as prognostic factors.

2.3. Microsatellite Analysis

Tumour tissue samples were available in 66 cases. Specimes of formalin-fixed, paraffin embedded 7µm tissue sections containing both, tumour and surrounding normal tissue, were deparaffinized and stained with haemalaun. DNA was extracted using a microdissection device as previously described in [9]. The precipitated DNA was used for polymerase chain reactin (PCR) analysis.

2.3.1. Analysis of Respective Microsatellites

The technique commonly used at the former times of the study period is extensively described in [9].

PCR amplification was done on the following panel on 6 highly polymorphic microsatellite markers. The chosen oligonucleotide repeat markers should located in the neighborhood of well-known tumour suppressor genes (p53, p16, rb1, hTR), or of the mismatch repair genes (hMSH2, hMLH1), respectively (genes linked with each marker are provided in parentheses):

- D2S136 (hMSH2), D3S1076 (hMLH1), D17S250 (p53), D9S942 (p16), D13S153 (rb1), and D3S1246 (hTR) (9).

PCR was performed in the Qiagen Hotstar Taq Mastermix kit (Fa. Qiagen, Hilden, Germany), and then products were separated using the ExcelGel™ DNA analysis kit (Fa. Amersham Biosciences, Munich, Germany). Products were silver-stained following the procedure outlined in [30].

A locus was considered informative for the respective polymorphism when normal tissue DNA exhibited two different alleles (heterozygosity). The intensities of the two alleles in the corresponding tumour were compared. The complete absence of an allele was interpreted as loss of heterozygosity (LOH). Reduced intensity of one of the two alleles was considered to indicate allelic imbalance (AI). Appearance of new alleles in tumour samples was considered to indicate microsatellite instability (MSI). Two investigators evaluated the results of electrophoresis independently without any patient information.
2.4. Statistical Analysis

Analysis of variance (ANOVA) was used to determine the oncologic outcome regarding the variables tumour volume, tumour stage, surgical margin, and the fact of intralesional previous surgery.

The correlation of the results of the detected genomic instabilities and the oncologic outcome was also analysed. Wilcoxon-test was used to compare subgroups. The survival probability was calculated by the method described in [29].

Statistical analysis of the data obtained was performed using the software SPSS 9.0 (SPSS Inc., Chicago, IL, USA). P-values <0.05 were considered significant.

3. Results

3.1. Oncologic Results

In the 30 low-grade tumours, the surgical margins achieved were wide in 21 cases, and marginal in 9 cases. In the 52 high-grade tumours the margins were radial in 7, wide in 21, marginal in 19, and intralesional in 5 cases. In the 4 stage-III tumours the margins were marginal in two cases, wide in one, and intralesional in one case. Amputations had to be performed in 9 patients (10.4%): hemipelvectomy, the lower extremity, and the upper extremity, each in three cases, respectively.

The median tumour volume of the 66 tumour containing specimens was 451 ml (range, 3-4800 ml). The stage-I tumours were smaller (386 ml) than the stage-II tumours (442 ml), but the difference was not significant.

At the date of actual evaluation in January 2010, twenty-nine patients died of their disease, and 7 patients died of other causes.

The mean follow-up was 12 years (range, 10 - 16 years) or 144 months (range, 120 - 192 months), respectively. The median survival time was 58 months (range, 26-74 months). The patients presenting with stage III tumours had a median survival time of 15.25 months. Local recurrences were observed in 13 patients (15.1%) after an average of 14.3 months (range, 4-47 months). Distant metastases were seen in 23 patients (26.7%) after an average of 11.7 months (range, 3-21 months). In most cases (n=21) there were pulmonary metastases.

Table 1 gives a survey of the oncological results in relation to the tumour stage and the surgical margin achieved.

The incidence of a local recurrence significantly depended upon the achieved surgical margin. The tumour volume had no influence.

Concerning all tumour stages, local recurrences were significantly more frequent in cases of achieved marginal margin compared with those of wide or radical margin. These findings could be stated in both subgroups, (i.e. in the low-grade as well as in the high-grade tumours). The incidence of distant metastases was higher in high-grade tumours than in low-grade-tumours. But in both tumour subgroups the incidence of distant metastases did not depend upon the surgical margin.

Table 1. Oncologic results in relation to the tumour stage and the surgical margin

| Tumour stage | Surgical margin | LR | Met's | DOD | DOC |
|--------------|----------------|----|-------|-----|-----|
| I            | II             | III | r     | w   | m   | i   |
| 30           | 21             |     | 1     | 1   | 1   | 1   |
| 9            | 4              | 2   | 2     | 1   |     |     |
| 52           | 7              |     | 1     | 4   | 3   |     |
| 21           | 1              | 6   | 7     | 2   |     |     |
| 19           | 6              | 6   | 9     | 2   |     |     |
| 5            | 1              |     | 3     | 1   |     |     |
| 4            | 1              | 2   | 1     | 4   | 4   |     |
| 30           | 52             | 4   | 7     | 43  | 30  | 6   |
| 13           | 23             | 29  | 7     |     |     |     |

(Abbreviations: r: radical; w: wide; m: marginal; i: intralesional; LR: local recurrence; Met’s: metastases; DOD: died of disease; DOC: died of other causes)

Excluding the four patients presenting with stage-III tumours, two different groups were also analyzed, the one (n=35) with previous intralesional surgery, and the other (n=47) without previous intralesional surgery. We found even less local recurrences and less distant metastases in the patients with previous intralesional surgery compared with those patients without previous surgery, but the difference was not significant. However, the achieved surgical margin was valued as wide in 20 cases of the 35 patients with previous intralesional surgery, where no tumour could be found at the histological examination.

In the 15 cases presented primarily with a local recurrence, there were no worse oncological results; in the most cases (n=11) there were high-grade tumours. The local recurrences could be cured in 10 out of these 15 patients.

Thirteen patients developed local recurrences, in 10 of these cases within the first two post-operative years. The local recurrences could be cured in four cases, twice by amputation, and twice by a renewed resection.

Distant metastases developed in 23 patients; 6 out of them were treated surgically. Only two patients could be cured, one by resection of a peripheral soft tissue metastasis, the other is now for 148 months considered a complete responder after chemotherapy. All other patients with distant metastases died of their disease.

The Kaplan-Meier survival analyses in relation to the tumour stage and the achieved surgical margin are shown in table 2 and in table 3.
5.5-year and 10-year survival probability in relation to the tumour stage

Excluding the stage-III tumours and the intralesional resected tumours, the overall survival was significantly dependent upon the tumour stage after 5 years as well as after 10 years. The survival probability of 56.5% after 5 years decreased in patients with high-grade sarcomas to 49% after 10 years, whereas the reduction of the survival probability during this period was insignificant in patients with low-grade sarcomas (89.5% to 88%).

Patients underwent wide resection showed a better prognosis than patients with marginal resection, but the difference was not significant. In both groups, there was a similar reduction of the survival probability in the period of 5 years to 10 years postoperatively. The worse prognosis after radical resections can be explained by the fact that there were only high-grade tumours in this group.

There were also no differences in the 5-year and the 10-year survival rate concerning the tumour volume, the fact of previous intralesional surgery, the primary presentation with a local recurrence, or the fact of chemotherapy in high-grade sarcomas.

3.2. Molecular Biological Results

In 66 patients tumour tissue was available for microsatellite analysis. Thus, a total of 396 loci were investigated, 130 of them (32.8%) were non-informative. In the remaining 266 loci, 94 of them (35.4%) were without pathological findings, i.e. negative.

Overall, allelic changes could be found in 172 out of the informative loci (64.6%), in 79.6% of the high-grade sarcomas, and in 13.3% of the low-grade sarcomas.

MSI was detected only in 18 out of the 266 informative loci (6.8%); all of them were in high-grade sarcomas. LOH was seen in 50 loci (18.8%), in one out of the 60 informative loci (1.7%) of the low-grade sarcoams, whereas in 49 loci out of the 206 informative loci (23.8%) of the high-grade sarcoams. This difference was statistically significant. Allelic imbalances (AI) were found in 39% out of all loci, in 11.6% of the low-grade sarcomas, and in 47.1% of the high-grade sarcomas, respectively.

Table 4 summarizes the results of the genomic instability regarding the tumour stage.

Table 4. Microsatellite analysis in relation to the tumour stage

|                         | total | low-grade (stage I) | high-grade(stage II) |
|-------------------------|-------|---------------------|----------------------|
| n                       | 66    | 15                  | 51                   |
| loci                    | 396   | 90                  | 306                  |
| informative             | 266   | 60                  | 206                  |
| thereof                 |       |                     |                      |
| negative                | 94 (35.4%) | 52 (86.7%)         | 42 (20.4%)           |
| genomic instabilities   | 172 (64.6%) | 8 (13.3%)          | 164 (79.6%)          |
| MSI                     | 18 (6.4%) | -                   | 18 (8.7%)            |
| LOH                     | 50 (18.8%) | 1 (1.7%)*           | 49 (23.8%)*          |
| AI                      | 104 (39%)  | 7 (11.6%)           | 97 (47.1%)           |

(Abbreviations: MSI: microsatellite instability; LOH: loss of heterozygosity; AI: allelic imbalance; *: statistically significant (p<0.05))
Correlation of the presence of genomic instability and the patient’s age, the patient’s sex, the tumour volume, the localization of the tumour, or the histologic type of the tumour were not detected.

The number of each tumour type was too small to allow an inference about the survival analysis as well as about the role of genetic instabilities. Die descriptive results of the different tumour types concerning the survival rate and the molecular biological findings are shown in table 5.

Concerning the different markers, the highest rate of genomic instability was seen at the microsatellite nearby hMLH1 (78%), hTR (75.5%), and hMSH2 (70.2%). The lowest validity was seen at the marker nearby p16 (40%).

It was particularly noticeable and statistically significant, that 15 out of the 18 cases of MSI were localized at the marker in the neighborhood of the mismatch repair genes, i.e. hMSH2 and hMLH1. The frequency of different genomic instabilities in relation of the different markers is demonstrated in table 6.

### Table 5. Survival rate and genetic instability categorized by different types of STS

| histology  | n   | 5-year survival | 10-year survival | specimen available for MSI analysis | informative loci | thereof genetic instabilities |
|-----------|-----|----------------|-----------------|-------------------------------------|-----------------|-------------------------------|
| MFH       | 25  | 70.5%          | 66%             | 20                                  | 80              | MSI                           |
| liposarcoma | 18  | 67.5%          | 62%             | 14                                  | 61              | 5                             |
| leiomyosarcoma | 11  | 64%            | 60.5%           | 9                                   | 44              | 5                             |
| fibrosarcoma | 7   | 68%            | 63%             | 6                                   | 28              | 2                             |
| rhabdomyosarcoma | 7   | 68.5%          | 61%             | 6                                   | 22              | 2                             |
| others    | 14  | 70.4%          | 62%             | 11                                  | 31              | 3                             |
| total     | 82  | Ø68.5%         | Ø63.2%          | total 66                            | 266             | 18                            |

(Abbreviations: MFH: malignant fibrous histiocytoma; MSI: microsatellite instability; LOH: loss of heterozygosity; AI: Allelic imbalance; stage III tumours were excluded)

Correlation of the presence of genomic instability and the patient’s age, the patient’s sex, the tumour volume, the localization of the tumour, or the histologic type of the tumour were not detected.

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### Table 6. Frequency of different genomic instabilities in relation to the individual markers

| markers (linked genes) | genomic instabilities | | | | | |
|------------------------|-----------------------|---|---|---|---|---|
| D2S136 (hMSH2)         | total 26              | 12* | 11 | 3 | 2 | |
| D3S1076 (hMLH1)        | total 32              | 3  | 12 | 17 | 0 | 0 |
| D17S250 (p53)          | total 34              | 3  | 13 | 18 | 0 | 0 |
| D9S942 (p16)           | total 18              | 18 | 18 | 0 | 0 | 0 |
| D13S153 (rb1)          | total 28              | 3  | 25 | 0 | 0 | 0 |
| D3S1246 (hTR)          | total 34              | 11 | 23 | 0 | 0 | 0 |
| total                  | total 172             | 18 | 50 | 104 | | |

(Abbreviations: MSI: microsatellite instability; LOH: loss of heterozygosity; AI: allelic imbalance; *: statistically significant (p<0.05))

### 3.3. Correlation of Oncologic and Molecular Biological Results

As expected, the low-grade sarcomas presented significantly better oncologic outcome. There was a low rate of genomic instability in these cases, only 8 out of the 60 informative loci (13.3%) were positive. Excluding the stage III sarcomas, the remaining 47 high-grade sarcomas presented genomic instabilities in 79.6% of the informative loci. Only in one of these cases a solitary alteration could be found showing LOH at the marker nearby hMSH2. In all other cases there were multiple genetic alterations at different corresponding markers, in 7 cases at 2 markers, in 15 at 3, in 20 at 4, and in two cases at 5 markers, respectively.

The low frequency of MSI in high-grade sarcomas (8.7%) had no influence on the overall survival. The appearance of allelic imbalance was also considered to be non-specific.

But there was a significant influence of the LOH concerning the survival probability in the high-grade sarcomas. When LOH was detected at least at one marker, the 5-year and 10-year survival probabilities were 48.6%, in cases without LOH detectable.

There was no significant difference concerning the detection of LOH at the different markers or the frequency of LOH at several markers. Excluding the markers nearby p16 and rb1, there was no preference of any marker, where LOH could be detected.

### 4. Discussion

Soft tissue sarcomas are very rare tumours accounting for less than 1% of all malignancies in the adult [30, 31, 32]. Basically, they can be divided into low- and high-grade tumours by the means of histopathological criteria resulting in the surgical classification according to [28].

In all malignancies, every treatment modality has to be measured by the achieved oncologic outcome. In sarcomas, surgery is considered to play the most significant role. Different prognostic factors are reported in STS [31, 32].

The overall-survival mostly depends upon the tumour stage. The 5-year-survival probabilities range from 50-60% in stage-II tumours up to 80-90% in stage-I tumours [31, 32]. These reports are confirmed by our findings, 89.5% in stage-I tumours, and 56.5% in stage-II tumours. Most studies reported an oncologic follow-up of 5 years [31 - 34]. But we could demonstrate in our series, that the survival probability will decrease furthermore. Low-grade sarcomas from 89.5% after 5 years to 88% after 10 years, but high-grade sarcomas from 56.5% after 5 years to 49% after 10 years, respectively.

There are some other prognostic factors in soft tissue sarcomas reported in the literature, such as tumour type, tumour volume, and localization [31, 32]. We could not
confirm these observations, maybe due to the limited number of patients in our series. The overall survival in our series only depended upon the tumour stage. The low-grade sarcomas were on average smaller than the high-grade sarcomas (386 ml versus 442 ml), but the difference was not significant. The thought can be suggested, that low-grade sarcomas were smaller in our series and therefore resectable more frequently with a wide margin. However, we could not find a statistical significance. The 5-year survival rate was 60.7%, when a marginal margin was achieved, and it was 80%, when a wide margin was feasible, the 10-year survival rate was 58% and 78%, respectively.

But we agree with other authors [31, 35], that the achieved surgical margin is the most important prognostic factor for local tumour control. The incidence of a local recurrence was significantly higher in tumours resected with a marginal margin compared with those resected with a wide margin. We observed significantly more local recurrences, when only a marginal margin was feasible.

Some studies [31, 34] demonstrated that the occurrence of local recurrence promotes the development of distant metastases resulting in a poor overall survival. In our series, we saw 15 patients presented with a local recurrence without concomitant distant metastases. Ten of them could be cured, and we could not find a worse oncologic outcome. But the small number of cases may not allow any statistical predication.

Despite intensive search of clinical and histomorphologic criteria individual prognostic statements are often not feasible in STS due to their considerable variability. Therefore, further predictive markers are intensively searched to allow a more precise prediction of the prognosis, or to define patient groups, who may benefit from specific treatment modalities.

The role of genetic instability is well known in different cancers [2, 16 - 18]. The two phenomena, microsatellite instability (MSI) and loss of heterozygosity (LOH), are considered to be of great importance [1 - 3, 8, 10 - 13, 24].

MSI is considered as a dysfunction of the mismatch repair (MMR) genes [5, 10 - 12, 14], whereas LOH seems to correspond to genetic defects of tumour suppressor genes [7, 16, 21, 22, 24, 26, 27].

Thus, MSI is reported as a negative prognostic factor in different cancers, such as lung cancer, or colorectal cancer [16 - 18]. It is considered to be a phenomenon of the early phase of tumour progression [10]. LOH is also accepted to be a negative predictor in many cancers. The colorectal cancer is the most frequently investigated tumour [16, 17].

The role of MSI and LOH in STS is still unclear. The most molecular-biological and cytogenetic data are reported in osteosarcomas, in Ewing’s sarcomas, or in uterine sarcomas [3, 4, 6, 24, 26, 27].

As described in [4], there was no MSI in 29 bone sarcomas at 6 different loci. MSI was also not found in 28 liposarcomas at 9 different loci, but LOH was present in 11 tumours (39%) [7]. MSI was found in only 3 cases out of 39 STS examined for 12 different loci, whereas LOH for at least one marker was seen in 14% to 85% depending upon the tumour type [3].

On the other hand, another study [21] reported on 20 sarcomas demonstrating MSI in 5 out of 7 high-grade tumours using 8 different markers. The region of chromosome 12 was especially affected. The oncogene mdm2 is localized in the neighborhood of the described microsatellite, it is considered to be the main antagonist of the tumour suppressor gene p53. All low-grade sarcomas did not show any MSI. LOH was seen in 75% of the high-grade sarcomas at the chromosome 2 in the region of the locus nearby the hMSH2 gene.

The detection of LOH was described as a negative predictor in 31 patients with thoracic sarcomas [8].

As confirmed by the findings in our series presenting MSI in only 8% of the informative loci without any prognostic value, MSI seems to play no prominent role in STS.

But the rate of LOH in our series corresponds with the findings of other authors [7, 21, 24, 26]. The incidence of LOH correlated to the tumour stage. There was no relationship to any histological subtype of STS, but a statistical valid statement is not feasible due to the small number of patient in each histological subtype. In our series, the oncologic outcome was worse in the high-grade sarcomas, when LOH was presented.

Many studies describe a correlation between MSI and alterations of the mismatch-repair-genes hMSH2 and hMLH1 [6, 11, 12, 15, 19]. As reported in [11], MSI at 5 different markers were seen in 25% of 40 STS. In 50% of these cases, loss of expression of the products of the hMLH1- and hMSH2-genes had to be observed. Another study [19] revealed similar results. MSI was visible in 37.5% of 16 alveolar sarcomas, whereas in two thirds of these cases loss of expression of the gene products of hMLH1 and hMSH2 were notified.

In our series, the important role of the mismatch repair genes (hMLH1 and hMSH2) was also confirmed showing 15 of the 18 MSI, and 23 of the 50 LOH, respectively, detected at the loci nearby these genes.

The role of the tumour suppressor gene p53 is also extensively described in STS. The frequency of mutation is reported to amount 15% to 20 % [23]. In the balance of oncoproteins and tumour suppressor genes, p53 is considered the antagonist of mdm2 [2, 22, 23, 25]. The high significance of the co-expression of p53- and of mdm2-gene products as an independent molecular prognostic factor is reported in [25]. Thus, the thesis is confirmed, that a medium transcription rate of oncogenes and tumour suppressor genes is wanted for a physiologic cellular regulation process. The imbalance of this transcription rate seems to have an oncogenic potential [25].

There exist only a few reports on genomic instabilities of the microsatellites located nearby the rb1- and the p16-genes. In our series, we found no MSI for both markers, and only 10 % LOH at the locus nearby the rb1 gene. As reported in [26] about 47 osteosarcomas, the 5-year
survival was significantly reduced in cases of detected LOH nearby the rb1 gene locus. These findings were recently confirmed by another study [24]. Allelic loss at the p16 gene locus was detected in 5.1% and MSI was reported in 25% of cases in the large series of 135 sarcomas [36]. These results are confirmed by the findings in our series. Alterations of the microsatellite nearby the p16 gene locus were observed in 40%, but there were only non-specific allelic imbalances, LOH and MSI were not detected.

Thus, we assume that genetic alterations at the p16 gene locus seem to be not relevant to the oncogenesis or to the tumour progression in sarcomas.

The role of the hTR gene in STS is unclear. Related LOH was found in 50% of osteosarcomas [27] and in 20% of leiomyosarcomas [9]. In our series, we observed genomic alterations in 75% at the microsatellite located nearby the hTR gene, in one third of these cases presented as MSI.

5. Conclusions

The surgical margin achieved after resection of a sarcoma is the most important factor for local tumour control, whereas the overall survival mainly depends upon the tumour stage. In high-grade sarcomas the survival rate will drop even after 5 years.

Therefore, long-term studies with prolonged follow-up will be helpful to define the exact oncologic outcome.

The role of the phenomena of the genetic alteration, microsatellite instability and loss of heterozygosity, are still unclear. MSI seems to be a rare phenomenon in STS without any prognostic value, but LOH indicates a worse prognosis in high-grade sarcomas.

The value of the different tumour suppressor genes has to be investigated by further multicentric studies. To assess the relevance of MSI and LOH for tumour progression and oncogenesis in sarcomas, larger numbers, modern detection techniques, and an appropriate microsatellite loci panel will be essential.

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