Malignant transformation of osteoblastoma: Study using image analysis microdensitometry

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Abstract

Aim—To determine if the malignant transformation, as perceived histologically, in a case of osteoblastoma from the right femur, was also expressed as a quantitative change in nuclear DNA during tumour progression over five months.

Methods—Nuclear DNA microdensitometry by computer image analysis was used to acquire relative DNA distribution patterns. Tissue had been removed on four separate occasions from a lesion in the right femur of an 18 year old man. Retrospective DNA analysis was performed on formalin fixed, paraffin wax-embedded tissue.

Results—The DNA profile of the initial biopsy specimen, which was histologically diagnosed as osteoblastoma, was euploid with a near diploid (2c) modal DNA. The second biopsy specimen taken one month later also resembled osteoblastoma but showed an aneuploid DNA profile with a diploid modal DNA and some nuclei with ploidy greater than 5c. The third biopsy specimen taken four months later showed histological evidence of osteosarcoma and a near pentaploid (5c) modal DNA with large number of nuclei exceeding 5c.

Conclusions—DNA microdensitometry confirmed the initial and final diagnosis. The technique also seems to be capable of detecting aneuploidy before malignancy is morphologically evident.

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Osteoblastoma is a rare osteoid producing bone neoplasm of osteoblastic origin. This is regarded by some as the benign counterpart of osteosarcoma, which in many cases is a highly malignant rapidly metastasising bone tumour. Most tumours diagnosed initially as osteoblastoma are cured by either intrale- sional or en bloc excision. About 10–15%, however, behave aggressively with local recurrence. Some of these either undergo malignant transformation and metastasise or are reappraised as osteoblastoma-like osteosarcomas. Inadequate DNA microdensitometry has been used as a diagnostic adjunct for neoplasms in which the diagnosis is equivocal. It may also have a more important and fundamental role in the investigation of the quantitative changes in nuclear DNA distribution associated with the process of malignant transformation.

Case report

An 18 year old man experienced worsening intermittent pain and swelling in his right thigh for a period of 14 months. The pain was temporarily relieved by mild analgesics and physiotherapy. Conventional radiography of his femur showed a prominent zone of pronounced cortical thickening and periosteal reaction involving the upper shaft. A zone of lucency measuring 35 × 6 mm was present within the sclerotic area (fig 1).

On admission to hospital the patient’s serum alkaline phosphatase was raised (202

Figure 1  Initial radiograph of right femur showing obvious fusiform new bone formation mainly along the anteromedial aspect of the proximal third of the shaft. There is also a well defined central translucency 4 cm in length in the medial cortex abutting the endosteal surface.
Osteoblastoma DNA microdensitometry

U/l: range <160 U/l) while the serum calcium was normal (2.46 mmol/l). Further x ray pictures did not show any abnormalities in the patient's lungs, mediastinum, right tibia and fibula. An open biopsy specimen produced gritty, bloody tissue 10 mm and 8 mm across and several fragments of bone, including two cores. Histological examination showed vascular cellular osteoid tissue and with adjacent dense bone resembling osteoid osteoma (fig 2). Because the nidus radiologically exceeded 1-5 cm, the lesion was classified as a benign osteoblastoma.12

After the biopsy there was a continuous ache in his right leg which was easily tolerated for one month. Then some dense cortical bone was removed and the nidus thoroughly curetted and filled with bone cement. Histological examination of the cortical bone specimens showed densely woven bone. The curetted tissue was similar to that seen in the first biopsy specimen showing highly vascular osteoid containing numerous osteoblasts and scattered osteoclasts. On review some of this osteoblastic tissue seems to have permeated the dense bone (fig 3).

Four months later, after experiencing pain of greater intensity than ever before, the patient was re-admitted for surgery a third time. X ray pictures showed an area of lucency around the upper half of the bone cement suggestive of a local recurrence (fig 4). The lesion was then curetted even more widely and treated with liquid nitrogen before packing it with bone chips from the iliac crest. Several pieces of cortical bone and curetted material were examined histologically. Some of the latter included fragments of abnormal osteoid with permeation of bone and moderate cellular pleomorphism, with up to 6 mitoses/mm² and an occasional abnormal mitosis (figs 5A, B, 6A and B). Voluntary muscle and plasma cells were found within cortical bone in some sections. High grade osteoblastic osteosarcoma was diagnosed. This was later confirmed by Professor KK Unni from the Mayo Clinic, and after a three cycle regimen of intra-arterial chemotherapy en bloc diaphyseal resection with intercalary reconstruction was performed.

New bone formation was found in the medullary cavity of a resected specimen which measured 180 mm in length. Histological examination showed no evidence of residual neoplasm. There was no clinical evidence of recurrence of metastatic disease 14 months after the resection.

Figure 2 Initial biopsy specimen. Histological features of osteoblastoma showing minimal invasion of adjacent bone trabeculae.

Figure 3 Residual osteoblastoma curetted before insertion of bone cement.

Figure 4 Radiograph four months after curettage and insertion of bone cement. The lucency around the proximal end of the cement had been increasing. Vascularity had also increased (angiogram).
Methods

Fresh tissue was fixed for 24 hours in 10% neutral buffered formalin, then processed routinely, and embedded in paraffin wax. Blocks containing bone fragments were de-mineralised in RDO (Du Page Kinetic Laboratories, Illinois). Paraffin wax sections (6 µm thick) were stained by a modified Feulgen method14 using acid hydrolysis for 1 hour in 5N HCl at 22°C before staining with Schiff reagent for 15 minutes, prepared according to the method of de Tomasi.15 Sections were selected for DNA analysis based on areas that exhibited the most malignant pattern.16

The integrated optical density (IOD) of at least 100 diagnostic cells from each biopsy specimen was measured using the MD20 microcomputer system (Leica) for video image analysis and microdensitometry.17 This system consists of a high resolution solid state video microscope mounted on a standard light microscope (Olympus BH2). The acquired images are digitised by an IBM/PC computer and IOD measurements are made on individually selected cells by grey level intensity thresholding and microdensitometry. Images can be edited to obtain precise threshold nuclear masking and corrections can be made which take into account irregularities in background illumination and inherent glare from the microscope optics.

Tumour associated lymphocytes were used as internal staining controls of the diploid DNA content. A diploid (2c) nuclear IOD value for each biopsy specimen was established from the lymphocyte measurements and used to convert the raw nuclear IOD values of the tumour cells to relative DNA values which are then displayed as ploidy frequency distributions. The raw measurement values and the diploid nuclear IOD values were also used to calculate various numeric indices of cell ploidy abnormality, including the 2c deviation index (2cDI) and the 5c exceeding rate (5cER). 2cDI is defined as the mean square deviation from the diploid
ploidy was evident, as reflected by the gradual increased frequency of nuclei with DNA content greater than the tetraploid (4c) level. A gradual increase in the variance of the tumour cell population about the normal diploid (2c) value was also evident. Both biopsy specimens I and II showed near diploid (2c) modal DNA values, but nuclei with DNA content greater than 5c were detected in the latter. Morphologically and radiologically, the lesion in both these instances looked benign and was diagnosed as osteoblastoma. The Böcking algorithm, however, classifies biopsy specimen I as benign (2cDI < bbl0 and 5cER < 0.1) and biopsy specimen II as malignant (5cER > 0.1) (table). Four months later (biopsy specimen III) morphological evidence of a transformation was apparent. The DNA distribution pattern showed a near pentaploid (5c) modal DNA value and a significantly increased population of tumour cells with a DNA content greater than 5c.

An analysis of biopsy specimen IV was not performed. The exposure of tumour cells to cytostatic agents disqualified the specimen from submission to the algorithm.

The malignancy grading of biopsy specimen II more than doubled in four months (biopsy specimen III). The results suggest that the tumour progressed from benign to high grade malignancy. The high number of mitotic figures noted in the second and third biopsy specimens compared with the low number in the initial biopsy specimen support the results.

A moderate reduction in Feulgen DNA stainability was noticed with the use of a commercially produced EDTA based decalcifying agent. Some authors advocate the use of EDTA-TRIS at pH 7.0 over other decalcifying agents on tissue requiring microspectrophotometric DNA analysis, reporting no modifications to the stoichiometric Feulgen reaction.

Lymphocytes were chosen as internal staining controls in preference to fibroblasts or chondrocytes on the basis of greater uniformity in size and shape, and due to their small size the increased probability of measuring unsectioned lymphocytes. The modal control IOD value for the lymphocyte population, however, multiplied by a factor of 1.15 to take into account the condensed chromatin normally associated with Feulgen stained lymphocytes.

Results
A profile of the nuclear DNA distribution through each stage of the transformation is shown in fig 7. A distinct change in tumour

<table>
<thead>
<tr>
<th>Biopsy</th>
<th>Time after initial biopsy (months)</th>
<th>2cDI</th>
<th>5cER</th>
<th>DI</th>
<th>MG</th>
<th>Classification by Böcking algorithm</th>
<th>Initial diagnosis*</th>
</tr>
</thead>
<tbody>
<tr>
<td>I</td>
<td>0</td>
<td>0.21</td>
<td>0.00</td>
<td>1.23</td>
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<td>Benign</td>
<td>Osteoblastoma</td>
</tr>
<tr>
<td>II</td>
<td>1</td>
<td>1.77</td>
<td>4.83</td>
<td>1.35</td>
<td>0.78</td>
<td>Malignant</td>
<td>Osteoblastoma</td>
</tr>
<tr>
<td>III</td>
<td>5</td>
<td>2.80</td>
<td>2.80</td>
<td>1.80</td>
<td>Malignant</td>
<td>Osteosarcoma</td>
<td>Malignant</td>
</tr>
</tbody>
</table>

*Based on clinical, radiological, and histopathological findings. N/A = Not applicable to benign tumours.
and characterisation of oncogenes and the role they have in malignant transformation.24,25

The activation and subsequent amplification of oncogenes or their inactivation and loss may result in structural or numeric chromosomal changes that can be detected by cytogentic analysis.26 The procedure is technically difficult on solid tumours, however, and only produces a small percentage of interpretable results. Chromosomal changes accompanying the malignant transformation of a cell may be detected if large enough to procure a change in the total DNA content, resolvable as a ploidy aberration by quantitative nuclear cytophotometric or microdensitometric methods.10,25 Microdensitometry in this study was used to investigate changes in nuclear DNA content of a bone tumour during its course through a morphologically perceived malignant transformation.

A review of published findings showed only three cases in which DNA analysis had been performed on osteoblastomas with malignant termination, none of which reported a progressive change in nuclear DNA content. Bauer et al10 describe an osteoblastoma from the tibia of a 14 year old that recurred after open biopsy, reappearing six months later as an osteosarcoma. The patient showed no evidence of disease during 10 years of follow up after wide excision of the lesion. A similar case from the sacrum of a 31 year old was reported to have recurred nine months later as a high grade osteosarcoma. The patient died two years after intrallesional excision of the tumour. The primary lesions from both patients were hyperploid by microspectrophotometric DNA analysis. Heliö et al27 described a bone tumour that was biopsied on three separate occasions over a period of 17 months, the first two biopsy specimens were diagnosed as osteoblastoma and osteosarcoma was diagnosed on the third and final biopsy specimen. Flow cytometric (FCM) analysis of the lesion showed an aneuploid peak in the DNA histograms of all three specimens.

A comparative analysis of results from independent studies should attempt to take the definitions used by each investigator into account.16 In several investigations using microspectrophotometric analysis,10,11,22,28-31 tumours were simply defined as diploid or hyperploid by placing an upper limit of diploidy based on DNA measurements made on internal control cell populations such as fibroblasts. Cells with DNA values exceeding those of proliferating diploid cells were not analysed and a definition or measure of aneuploidy was usually not attempted, which may explain why our results are at variance with those of other reports.10,11,27

Our observations, however, cannot go unqualified without some reference to tumour representativity, considering that different biopsy procedures were used at various times, each of which yielded varying sample volumes from a morphologically heterogeneous bone tumour. Profuse bleeding from the lesion at investigative surgery largely accounted for the small volume of tissue procured during open biopsy in our study. The results and experience of other workers support our belief that the tumour was adequately biopsied on this occasion. Kreiebergs et al23 produced highly reproducible FCM DNA distribution curves of tumours sampled on different occasions—open biopsy and curative surgery. Bauer et al10 concluded that regardless of whether flow cytometry or cytophotometry is used, osteosarcomas express a uniform DNA content despite apparent morphological heterogeneity.

In most cases the ploidy profile of a bone tumour is strongly related to its histopathological diagnosis.11 Benign bone lesions are almost always diploid10,11,16,27,32 and classic osteosarcomas are predominantly hyperploid or aneuploid, with the exception of parosteal variants which are invariably diploid.10,11,28,29,31-33 Among the classic osteosarcomas studied by Bauer et al,10,29 modal DNA values in the triploid region predominated, while Kreiebergs et al23,30 reported modal DNA values ranging from 2-6c—5-4c in one study and between 3c—7c in another. The near penta- ploid modal DNA value obtained on the third biopsy specimen of the tumour in this study is within the range of values obtained by Kreiebergs et al.

The exceptional result obtained for the second biopsy specimen of our case study may prompt one to postulate the existence of a transitional stage where changes in nuclear DNA content precede a morphological change, to explain this observation. Experimental evidence exists, albeit indirect, to support this theory from a study on the pathogenesis of squamous cell carcinoma of the uterine cervix in mice treated with 3,4-Benzpyrene.34 Morphological evidence of invasive osteosarcoma in this study was also preceded by the existence of atypical cells with increased nuclear DNA. A progressively increased and scattered DNA content was noted during the transformation to invasive carcinoma of the cervix.

Nuclear DNA microdensitometry was a helpful adjunct for the diagnosis of this bone tumour. It remains to be determined whether the technique has the potential to have an investigative role in the biology of malignant transformation and further our understanding of carcinogenesis.

3 Pietters As, Vernon-Roberts B, Paterson DC, Cornish BL, Lewis FR. Osteoid osteoma transforming to aggressive (low grade malignant) osteoblastoma: a case report and literature review. Histopathology 1983;7:789-800.
7 Jackson JR, Bell MEA. Spurious "benign osteoblastoma".
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