Characterizing the Blood Oxygen Level-Dependent Fluctuations in Musculoskeletal Tumours Using Functional Magnetic Resonance Imaging

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This study characterized the blood oxygen level-dependent (BOLD) fluctuations in benign and malignant musculoskeletal tumours via power spectrum analyses in pre-established low-frequency bands. BOLD MRI and T1-weighted imaging (T1WI) were collected for 52 patients with musculoskeletal tumours. Three ROIs were drawn on the T1WI image in the tumours’ central regions, peripheral regions and neighbouring tissue. The power spectrum of the BOLD within each ROI was calculated and divided into the following four frequency bands: 0.01–0.027 Hz, 0.027–0.073 Hz, 0.073–0.198 Hz, and 0.198–0.25 Hz. ANOVA was conducted for each frequency band with the following two factors: the location of the region of interest (LoR, three levels: tumour “centre”, “peripheral” and “healthy tissue”) and tumour characteristic (TC, two levels: “malignant” and “benign”). There was a significant main effect of LoR in the frequencies of 0.073–0.198 Hz and 0.198–0.25 Hz. These data were further processed with post-hoc pair-wise comparisons. BOLD fluctuations at 0.073–0.198 Hz were stronger in the peripheral than central regions of the malignant tumours; however, no such difference was observed for the benign tumours. Our findings provide evidence that the BOLD signal fluctuates with spatial heterogeneity in malignant musculoskeletal tumours at the frequency band of 0.073–0.198 Hz.

Acute hypoxia (i.e., transient cycles of hypoxia–reoxygenation) is known to occur in solid tumours¹ and may be associated with resistance to radiation therapy²,³, impaired delivery of chemotherapeutic agents⁴ or metastasis development⁵. The origin of acute hypoxia is not firmly established; however, it is generally believed to result from tumour blood flow instabilities. Flow fluctuations may result from arteriolar vasomotion⁶,⁷, haemodynamic changes resulting from a disorganized vascular hierarchy⁸, and other factors.

Several methods can detect acute hypoxia. Oxygen microelectrodes⁹,¹⁰ and fibre-optic oxygen-sensing device¹¹,¹² can invasively detect the spontaneous oxygen tension fluctuations in animal tumour models by inserting a microprobe into the tumour. In addition to their invasive nature, another drawback of these techniques is the restriction of spatial resolution. Additionally, although phosphorescence lifetime imaging to detect the partial oxygen tension through a cutaneous window chamber has high spatial resolution, it is an invasive method and is not allowed in humans¹³. Additionally, hypoxia can be detected using positron emission tomography (PET) with a hypoxia-specific radio-tracer. Acute hypoxia may be identified through acute changes in the intensity

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of the tracer in human tumours with a mathematical model. However, detection of acute hypoxia using PET requires several injections of radiotracers over days. Moreover, in intratumoural injections of perfluorocarbon (PFC) compounds, the relaxation rate 1/T1 is proportional to the percentage of O2 acquired. This is also an invasive procedure and has limited temporal resolution.

Blood oxygen level-dependent (BOLD) functional MRI that depends on deoxyhaemoglobin as an endogenous contrast agent is sensitive to changes in blood flow and deoxyhaemoglobin content with high temporal and spatial resolution. The spatial and temporal heterogeneity of spontaneous T2* (BOLD) MR signal fluctuations was first observed in an implanted fibrosarcoma mouse model by Baudelet et al. and later confirmed in tumour xenograft models of colorectal carcinoma by Goncalves and colleagues.

However, the occurrence of spontaneous BOLD signal fluctuations in the tumours described above has never been demonstrated in human tumours. We examined patients with malignant or benign musculoskeletal (MSK) tumours using BOLD for 6 minutes per patient and analysed the BOLD fluctuation characteristic in the centre and periphery of each tumour via power spectrum analyses. The aim of this study was to assess the spatial heterogeneity of the BOLD signal fluctuations in human MSK tumours via power spectrum analyses in pre-established low-frequency bands.

### Results

#### Participants and Histopathology.

Of the 52 MSK tumour patients, four had excessive body motions (translation >3 mm or rotation >3°) that could have introduced severe artefacts in BOLD signals; these patients were removed from further analyses. The data from the remaining 48 patients were entered into further analyses (Table 1). Histological results showed that 35 patients had malignant tumours; the others had benign tumours. The malignant MSK tumours included osteosarcoma (12 cases), malignant fibrous histiocytoma, synovial sarcoma, alveolar sarcoma, Ewing sarcoma, liposarcoma, chondrosarcoma, metastatic adenocarcinoma, metastatic renal clear cell carcinoma, primitive neuroectodermal tumour, malignant chondroblastoma, epidermoid leiomyosarcoma, fibrosarcoma, malignant tenosynovial giant cell tumour.

### Statistical Analyses.

The power spectrum of the BOLD fMRI signal was divided into the following frequency bands (Table 1): band-1, 0.01–0.027 Hz; band-2, 0.027–0.073 Hz; band-3, 0.073–0.198 Hz; and band-4, 0.198–0.25 Hz. The reason for adopting these cut-offs is that the BOLD fluctuation power within the four bands was approximately equal. These results are statistically shown in our 2-by-3 ANOVA.
analysis, which used tumour characteristic (TC, consisting of two levels: "malignant" and "benign") and the location of ROI (LoR, consisting of three levels: tumour "centre/c" and "peripheral/p", and "healthy tissue") as two factors. As shown in Table 2, there was no significant main effect for the TC or the interaction effect between TC and LoR in all frequency bands. Notably, Table 2 also showed a significant main effect for LoR in the frequency of band-3 (0.073–0.198 Hz) ($P = 0.0167$) and band-4 (0.198–0.25 Hz) ($P = 0.0243$).

After the ANOVA analyses, the data with a significant main effect of LoR was further processed with post-hoc pair-wise comparison, as shown in Table 3. For malignant tumours, there was a statistically significant difference ($P < 0.05$, corrected) in BOLD fluctuation power in band-3 and a trend towards a significant difference ($P < 0.1$, corrected) in band-4 between the tumour periphery and the centre (i.e., $p > c$); the benign tumours showed no such difference (Fig. 2).

**Intra- and Inter-rater Reliability in ROI definition.** To determine whether these results were affected by the ROI selection, the intra- and inter-rater reliability of the BOLD fluctuation power between ROI selections

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**Figure 1.** BOLD fluctuation magnitude within and around a malignant MSK tumour from a randomly selected patient. The first row shows the 3DT1-weighted image of the MSK tumour in the femur and the BOLD-fMRI field-of-view (within the red rectangle). The second to the last row shows the BOLD fluctuation magnitude (normalized power) within frequency bands 1–4, which were rendered onto the T1-weighted image.
were there significantly (band-3, \(P\)-rounding normal muscle (Table 2). The post-hoc simple effect analysis showed that only in the malignant tumour associated with the BOLD fluctuation 25,26. Blood flow fluctuation may induce acute hypoxia in tumours. Thus,

Table 2. The ANOVA Table of Power Difference in 48 Cases. Note: *\(P\) value lower than 0.05; **\(P\) value lower than 0.005; *\(P\) value lower than 0.1 after correction, which is a \(P\) value less than 0.0174. *\(P\) value lower than 0.05, corrected) or a trend towards significantly (band-4, \(P\) < 0.05, corrected) or a trend towards significantly (band-4, \(P\) < 0.1, corrected) higher BOLD fluctuations in the peripheral tumour region than the central tumour region; there was no such difference for benign MSK tumours (Table 3).

Table 3. Post-hoc pairwise comparisons analysis with Sidak adjustment in 0.073–0.198 Hz and 0.198–0.25 Hz. Note: **\(P\) value lower than 0.005. *\(P\) value lower than 0.1 after correction, which is a \(P\) value less than 0.0174. \(p\) > c: power of the periphery is higher than that of the centre. \(p\) > m: power of the periphery is higher than that of the normal muscle.

Discussion
This is the first time that human MSK tumours have been characterized using a frequency spectrum analysis of BOLD in specific frequency bands. Taking our observations together, ANOVA showed that in the frequency bands 0.073–0.198 Hz (band-3, \(P\) = 0.0167) and 0.198–0.25 Hz (band-4, \(P\) = 0.0243), there was a significant difference in the BOLD fluctuation power between the centre and the periphery of the MSK tumour and the surrounding normal muscle (Table 2). The post-hoc simple effect analysis showed that only in the malignant tumour were there significantly (band-3, \(P\) < 0.05, corrected) or a trend towards significantly (band-4, \(P\) < 0.1, corrected) higher BOLD fluctuations in the peripheral tumour region than the central tumour region; there was no such difference for benign MSK tumours (Table 3).

According to Ogawa’s group, the source of BOLD fMRI signal displays the ratio of oxyhaemoglobin and deoxygenated haemoglobin in the arterioles, capillaries and post-capillary venules. The BOLD signal has also been shown to be positively related to blood flow or the partial pressure of oxygen in normal tissue or tumour tissue. Flow fluctuation is also called flowmotion. Flowmotion as detected via perfusion imaging is closely associated with the BOLD fluctuation. Blood flow fluctuation may induce acute hypoxia in tumours. Thus, spontaneous BOLD fluctuations are likely related to acute hypoxia in tumours.

The BOLD fluctuations in band-3 in the periphery of the malignant MSK tumours were larger than those in the centre in our series, whereas for benign tumours there was no significant difference in all frequency bands between the centre and periphery. Häfner detected the perfusion of human cutaneous malignant melanomas, benign melanocytic nevi and normal skin. The wavelet analysis reveals that the mean vasomotion scale variance values obtained from healthy skin significantly differed from the margins and centres of cutaneous malignant melanomas but did not significantly differ between the margin and centre of benign melanocytic nevi. These findings support our results. However, the amplitude difference of BOLD fluctuation between the centre and periphery in malignant MSK tumours was only at the band of 0.073–0.198 Hz; the cause for this remains to be determined.
Interestingly, the frequency band of 0.073–0.198 Hz in tumours has been suggested to be caused primarily by vascular myogenic activity in vasomotion\(^{28,29}\). Further study is needed to determine whether this frequency band detected via BOLD has the same implications.

The frequency of oxygen fluctuation (hypoxia-reoxygenation) detected via microprobe\(^9\),\(^{11,12}\) or \(^{19}\)F MRI methods\(^9\) ranges widely from 0.00035–0.0167 Hz, with an average of approximately 0.001 Hz.

Our MR scanning protocol precluded the analysis of frequency bands higher than 0.25 Hz (due to the limited temporal sampling rate, 2 s) and less than 0.00278 Hz (due to the 6-minute signal acquisition time). To elucidate the spatial heterogeneity of the amplitude of the BOLD signal at some fluctuation frequencies, we tested the differences between the benign and malignant MSK tumours as well as between the centre and periphery of the MSK tumours. Only an amplitude of 0.073–0.198 Hz was significantly higher for the peripheral tumour region in the malignant tumour. The significant frequency band (0.073–0.198 Hz) detected in our study may provide new pathological and physiological significance in characterizing the acute hypoxia fluctuations in the tumours.

![Figure 2](image)

**Figure 2.** Post-hoc pairwise comparison analysis with Sidak adjustment in 2 frequency bands between benign and malignant tumours. **P < 0.05 after correction; *P < 0.1 after correction.**

| Frequency band (Hz) | ROIs in the centre of the tumour | ROIs in the periphery of the tumour | ROIs in the normal muscle |
|--------------------|---------------------------------|-----------------------------------|--------------------------|
| Band-1: 0.01–0.027 | 0.62\(^a\) | 0.68 | 0.80 |
|                    | 0.72/0.59\(^b\) | 0.37/0.54 | 0.79/0.86 |
| Band-2: 0.027–0.073 | 0.64 | 0.60 | 0.65 |
|                    | 0.80/0.71 | 0.59/0.64 | 0.45/0.66 |
| Band-3: 0.073–0.198 | 0.81 | 0.49 | 0.58 |
|                    | 0.70/0.75 | 0.52/0.63 | 0.43/0.78 |
| Band-4: 0.198–0.25 | 0.93 | 0.52 | 0.68 |
|                    | 0.81/0.83 | 0.63/0.64 | 0.66/0.87 |

Table 4. Values of Intra- (M.J.W-1 and M.J.W-2) or Inter- (M.J.W-1–Z.J.Z/M.J.W-2–Z.J.Z) Rater Correlation Coefficient for the Reproducibility Analyses. Note: a = ICC values between M.J.W-1 and M.J.W-2; b = ICC values between M.J.W-1 and Z.J.Z; c = ICC values between M.J.W-2 and Z.J.Z.
In the present study, 48 patients with malignant or benign MSK tumours were studied; this represents a relatively small sample size. The substantially smaller sample size of benign tumours compared with malignant tumours may induce statistical bias.

In conclusion, we found that in the frequency band of 0.073–0.198 Hz, the fMRI BOLD fluctuation power of the central is lower than the peripheral regions in human malignant MSK tumours; however, no significant difference was found for benign tumours. The meaning of this difference remains under investigation.

Materials and Methods
Participants. Fifty-two patients (34 males and 18 females; age 39.9 ± 17.8 years, 14–75 years) who were hospitalized consecutively from February 2009 to June 2011 with primary MSK tumours were included in this study. All patients were drug-naive and without any treatment prior to MR imaging. The demographic and clinical information are summarized in Table 1. The inclusion criteria were the following: (1) the tumour had to be predominantly solid; (2) the tumour dimension had to be larger than 3 cm in each of the axial, sagittal and coronal plains to facilitate ROI drawing; and (3) the tumour had no extensive intra tumour necrosis or bleeding. Tumour characteristics were determined via histological testing after resection or needle biopsy. This study was approved by the institutional research ethics board in the Third Hospital of Hebei Medical University, China. All experiments were performed in accordance with relevant guidelines and regulations, and all patients signed written informed consent before the study was carried out.

MR Imaging. MR imaging was performed on a 1.5 T Siemens MR scanning system (Avanto, Siemens, Erlangen, Germany). Scanning was performed with patients lying in a supine position and being asked to relax throughout the scan. Depending on which body part had tumours and was scanned, image acquisition used one of the following coils: a large circularly polarized flexible coil, an eight-channel knee coil or an eight-channel body array coil. The scanning protocol included three-dimensional turbo fast low angle shot (FLASH) T1-weighted images (3D T1WI) (repetition time/echo time, 1900/2.97 ms; flip angle, 15°; number of slices, 176; slice thickness/gap, 1/0.5 mm; acquisition matrix, 256 × 246; field of view, 220 mm × 220 mm). Two-dimensional echo planar imaging (EPI) BOLD fMRI with 20 axial slices was also taken (repetition time/echo time, 2000/40 ms; slice thickness/gap, 5/1 mm; field of view, 220 mm × 220 mm; acquisition matrix, 64 × 64; voxel size, 3.44 mm × 3.44 mm × 6.0 mm; number of dummy scans, 3; scanning time, 6 min, 177 frames total). 3DT1WI images were taken to identify the tumour locations; the BOLD fMRI scan was taken to detect BOLD fluctuations in corresponding body tissues.

Data Preprocessing. The fMRI data were preprocessed using DPARSFA v2.430 and REST v1.831 based on SPM8 (www.fil.ion.ucl.ac.uk/spm) and Matlab 2010a (MathWorks, Inc., Natick, MA, USA). See Fig. 3 for a flowchart of the entire analysis procedure. The preprocessing steps included the following: (1) the fMRI data were converted to Neuroimaging Informatics Technology Initiative (NIITI) format; (2) the first five frames of each patient’s fMRI data were discarded to allow for MR machine equilibrium and the patients’ adaption; (3) within-frame slice acquisition timing was corrected; (4) patient’s body motion was corrected using a 6-parameter rigid-body transformation, and the patients found to have excessive body motion (>3 mm or >3 degrees) were excluded from further analyses; (5) the three-dimensional T1 image of each patient was co-registered to the patient’s own averaged BOLD fMRI image to spatially match the two modalities. Note that we did not perform spatial re-sampling or smoothing because we intended to keep the original BOLD signal and to minimize spatial blurring influence and signal contamination caused by neighbouring voxels.

ROI Drawing. BOLD fMRI images provide critical information about blood flow and oxygen levels but do not readily allow visual distinction between tumour and non-tumour regions due to the limited spatial resolution. Therefore, we defined ROIs based on 3D T1WI and then transformed the coordinates of the ROIs to match the BOLD fMRI images according to the projection derived from the co-registration. Prior to ROI definition, the tumour tissue was divided into three nearly concentric regions (centre, transition and periphery regions) from the tumour centre to periphery as equally as possible by an experienced radiologist (M.J.W) according to previous studies32. As the transition area within a tumour was difficult to be differentiated from the tumour centre and peripheral regions32, we intended not to use it as an ROI. Within tumours, ROIs were only chosen from the tumour centre and periphery. The normal muscle surrounding the tumour but without abnormal signal was also defined as one ROI type. Therefore nine ROIs, three in the centre (c), three in the periphery (p) of the tumour and three in the surrounding normal muscle (m), with a radius of 4 mm each, were chosen for each patient. The three ROIs belonging to the same tissue type (c, p, or m) were chosen to spread as far as possible33 from each other to avoid systemic noise contamination. The regions with necrosis, calcification or that included large arteries or veins were avoided when defining the ROIs. For ten patients, the fMRI data did not include adequate normal muscle tissue because of the limited imaging field of view; we did not define the ROIs of the normal muscle for these patient (i.e., the muscle ROIs were available for 38 of 48 patients). Each ROI contained five voxels of the normal muscle for veins were avoided when defining the ROIs. For ten patients, the fMRI data did not include adequate normal muscle tissue because of the limited imaging field of view; we did not define the ROIs of the normal muscle for these patient (i.e., the muscle ROIs were available for 38 of 48 patients). Each ROI contained five voxels of the normal muscle for veins were avoided when defining the ROIs. For ten patients, the fMRI data did not include adequate normal muscle tissue because of the limited imaging field of view; we did not define the ROIs of the normal muscle for these patient (i.e., the muscle ROIs were available for 38 of 48 patients). Each ROI contained five voxels of the normal muscle for veins were avoided when defining the ROIs. For ten patients, the fMRI data did not include adequate normal muscle tissue because of the limited imaging field of view; we did not define the ROIs of the normal muscle for these patient (i.e., the muscle ROIs were available for 38 of 48 patients). Each ROI contained five voxels of the normal muscle for veins were avoided when defining the ROIs. For ten patients, the fMRI data did not include adequate normal muscle tissue because of the limited imaging field of view; we did not define the ROIs of the normal muscle for these patient (i.e., the muscle ROIs were available for 38 of 48 patients). Each ROI contained five voxels of the normal muscle for veins were avoided when defining the ROIs.

Power Spectrum Analyses. For each patient, averaged BOLD time series across the five voxels in each ROI were calculated, and a linear trend was removed from them. The averaged BOLD signal for each ROI was transformed from the temporal to frequency domain with a fast Fourier transformation using Matlab (i.e., frequency power spectrum in which the “power” is the square of the BOLD fluctuation amplitude, Fig. 1). This power spectrum was then divided into the following four frequency bands according to a previous study37: band-1, 0.01–0.027 Hz; band-2, 0.027–0.073 Hz; band-3, 0.073–0.198 Hz; and band-4, 0.198–0.25 Hz. The power values...
for each ROI were then normalized by dividing the total power value of the entire frequency band (0–0.25 Hz) to facilitate inter-patient comparisons (similar to the calculation of fractional amplitude of low-frequency fluctuation in Zang et al. 33). The normalized power values for the three ROIs belonging to the same tissue type were further averaged for each frequency band for the Analysis of Variance (ANOVA).

Statistical Analyses. We investigated whether there was significant difference in BOLD fluctuation power between different locations of ROIs or between malignant and benign tumours. Specifically, there were two main factors in the ANOVA model as follows: (1) tumour characteristics (TC), which consisted of two levels (i.e., either benign or malignant tumour) and (2) the location of the ROI (LoR), which has three levels (i.e., the tumour centre (c), the tumour periphery (p), and the normal muscle (m)). The 2 × 3 mixed-designed ANOVA (the two fixed effect factors stated above and one random effect factor, “subject”) was performed on the averaged BOLD fluctuation power values on each frequency band. Two main effects (i.e., the TC and LoR main effects) and one interaction effect (i.e., TC-by-LoR) were assessed. For any significant (P < 0.05, uncorrected) main effect derived from the ANOVA in any frequency band, simple effect analyses using post-hoc pair-wise comparisons were conducted to further explore the direction of the difference in BOLD fluctuation power. The significance level was set to P < 0.05 and corrected for multiple comparisons with Sidak adjustment.

Intra-Rater and Inter-Rater Reliability Assessment. We chose relatively objective ROI selection criteria (described above) to reduce the introduction of errors due to human subjectivity. To assess the intra-rater reliability of our ROI selection criteria, we performed the ROI definition twice with our primary radiologist (M.I.W.). The first and second round of ROI definition were separated by one week according to the same criteria but fully independent from each other. To determine inter-rater reliability, all of the ROIs were also independently drawn by a second experienced radiologist (Z.J.Z.) who was blinded to all previous results. The averaged BOLD fluctuation power value for each tissue type for each patient was calculated and fed into intra-class correlation coefficients (ICC) calculation. We calculated the intra-rater and inter-rater ICCs for each of the frequency bands. All ICC values were evaluated with the criterion reported in Zhang et al. 34. An ICC value larger than 0.75 indicates "excellent" reliability, between 0.59–0.75 indicates "good", 0.40–0.58 indicates "fair", and below 0.40 indicates "poor" reliability.

Figure 3. Schematic illustration of the course to data processing and analysis. The selected patient is a 19-year-old man with osteosarcoma in his left femur. (a) Functional image. (b) 3D T1-weighted image (c) A ROI in the centre of the tumour with a 4-mm radius. Its centre coordinate is (97,114,132). (d) Time course after preprocessing. (e) Power spectrum using FFT. ROI = region of interest; FFT = fast Fourier transformation; ANOVA = analysis of variance.
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Acknowledgements
This work was supported by the Medical Science Research Foundation of Hebei Province (Grant Number: 162777060) and partially supported by the National Natural Science Foundation of China (No. 81201156).

Author Contributions
J.-L.C. and H.Z. conceived the experiment, designed the statistical methods, and revised the manuscript for important intellectual content. L.-S.D. and M.-J.W. conducted the experiment, analysed the results and drafted the manuscript. E.S., Z.-J.Z. and M.X. conducted the experiment. Y.-F.Z., S.L. and S.-J.C. revised the manuscript. All authors reviewed the manuscript.

Additional Information
Competing financial interests: The authors declare no competing financial interests.
