Tumour oxygenation is known to enhance the efficacy of radiotherapy, because the presence of hypoxic tumour cells is considered to be one of the major reasons for failure to control tumours (Hall, 1994). Regarding hypoxic tumour cells, it is also known that ionizing radiation and some chemotherapeutic agents are less effective at low oxygen levels. Many studies on tumour oxygen tension levels using direct invasive measurements have been reported (Vaupel et al, 1984; Rampling et al, 1994; Brizel et al, 1996; Collingridge et al, 1997; Helmlinger et al, 1997; Al-Hallaq et al, 1998). Non-invasively, there has been increasing interest in measurements of changes in tissue oxygen tension using magnetic resonance imaging (MRI) methods. Semi-quantitative measurements of the tumour oxygen level have been discussed using oxygenation-sensitive 1H-MRI measurements during 100% oxygen inhalation (Karczmar et al, 1994; Kuperman et al, 1995; Edelman et al, 1996; Oikawa et al, 1997; Obata et al, 1998). These approaches have been used to increase tumour oxygenation sensitizing to radiotherapy and chemotherapy. Hyperbaric oxygenation (HBO) increases the oxygen supply to hypoxic tumour cells independent of its blood flow. Thus, HBO has been used clinically in combination with radiotherapy, but the previous combination method in which irradiation was administered during HBO exposure was both hazardous to patients and complex (Dische, 1978; Jain, 1990). As a result, HBO has not been routinely adopted with radiotherapy to treat patients with cancer. We irradiated human malignant gliomas 15 min after HBO exposure based on the hypothesis that elevated partial oxygen tension (P02) in the tumours was maintained for substantial periods after decompression (Kohshi et al, 1996). Using invasive measurements, it was reported that tissue P02 increased slowly during HBO exposure and that the decline in P02 after HBO was slower in subcutaneous tissues than in muscles (Wells et al, 1977), but no study on P02 change of tumours after HBO has been reported. The purpose of this study was to non-invasively monitor the tumour P02 changes produced by HBO exposure using MRI, and to clarify whether the elevated oxygen level in the tumours is maintained for substantial periods after decompression.

MATERIALS AND METHODS

Phantom preparation

To investigate the effect of oxygen dissolved in water, we measured the proton relaxation time of five phantoms. Small tubes were prepared with different oxygen concentrations. The water phantoms were as follows: (a) water without oxygen, (b) water with bubbling oxygen under 1.0 atmosphere absolute (ATA), (c) water with bubbling oxygen under 1.5 ATA, (d) water with bubbling oxygen under 2.0 ATA, and (e) water with bubbling oxygen under 2.5 ATA. MR spectroscopic measurements were performed using a Spectroscopy Imaging Systems Corporation (SISCO, Varian NMR Instruments, Palo Alto, CA, USA) 4.7 Tesla, 40 cm bore system. The hydrogen-1 resonant frequency was 200.43 MHz. The T1 relaxation time was measured by alteration of the inversion time (TI) using an inversion recovery pulse sequence. An exponential fitting was utilized to calculate T1 relaxation time.
Signal intensity = \( Mo \left[1 - 2 \exp \left(\frac{TI}{T1}\right)\right] \),

where \( Mo = \) longitudinal magnetization at equilibrium.

**Tumour model**

Ten- to 12-week-old female C3H/He mice were used. The research was conducted according to the principles described in the ‘Guiding Principles for the Care and Use of Animals approved by the Faculty Meeting of the University of Occupational Environmental Health’. SCCVII tumour cells, the hypoxic fraction of which was 9.1% (Shibamoto et al, 1994), were maintained in culture in RPMI-1640 (Gibco Laboratories, Grand Island, NY, USA) supplemented with 10% fetal bovine serum (Gibco Laboratories) and antibiotics, and trypsinized before making single cell suspensions. The mice were inoculated in the left leg with \(3 \times 10^5\) viable SCCVII tumour cells. Urethane-anaesthetized, spontaneously breathing mice were studied when the tumour size was about 1 cm in diameter. During MRI measurements, their legs and tumour were restrained in alginate impression material without occluding the blood supply on the table graduating scale. The mice were transferred with the table into a small experimental hyperbaric chamber. We attempted to set the same position, within 0.5 mm difference between pre- and post-HBO exposure. The temperature was maintained using warm oxygen forced through a hyperbaric chamber during HBO exposure.

**MRI measurements of tumours**

MRI measurements were performed using the same MR system described above. MR images were taken with a bird cage-type resonator (inner diameter, 8.9 cm) in a magnet fitted with an actively shielded gradient coil (1.8 G cm\(^{-1}\)). The molecular oxygen-enhanced MR images were acquired using an inversion recovery-preparation fast low angle shot (IR-FLASH) sequence. The acquisition parameters for the IR-FLASH sequence were as follows: repetition time (TR), 30 ms; echo time (TE), 8 ms; flip angle, 30°; field of view (FOV), 80 \(\times\) 80 mm; matrix, 128 \(\times\) 128; one excitation; slice thickness, 2.0 mm. The inversion time was 1000 ms to sensitize the acquisition to the paramagnetic effects of molecular oxygen. Each image of IR-FLASH took about 5 s to acquire. The acquisition parameters for the spin echo (SE) sequence were as follows: FOV, 80 \(\times\) 80 mm; matrix, 256 \(\times\) 128; two excitation average; slice thickness, 2.0 mm. For T1-weighted images and gadolinium (Gd)-enhanced images, TR was 300 ms and TE was 20 ms; for the T2-weighted images TR was 2000 ms and TE was 80 ms. The resonance frequency and shimming did not change between the pre- and post-HBO exposure.

**Experimental schedule**

For T1-weighted images, a slice was selected through the centre of the tumour and two baseline IR-FLASH images were initially acquired while the mice were breathing air. For the HBO-treated group \((n = 6)\), HBO exposure was given in a small experimental hyperbaric chamber according to the following schedule: 10 min of compression with oxygen, 60 min of 100% oxygen inhalation at 2.0 ATA, and 10 min of decompression with oxygen inhalation. For the normobaric group \((n = 5)\), oxygen inhalation was given in the same schedule as above but without compression. With air inhalation, the acquisition of IR-FLASH images was started 5 min after decompression, and images in both groups were obtained every 2.5 min. Finally, T1- and T2-weighted SE images were obtained. Subsequently, after gadopentetate dimeglumine (Gd-DTPA, 0.4 ml kg\(^{-1}\)) (Magnevist, Berlex Laboratories, Wayne, NJ, USA) was administered intravenously, the Gd-enhanced image was taken. Tumours with a haemorrhagic lesion on the T1-weighted SE image or necrotic tissue on the T2-weighted SE image were excluded from this study. A region of interest encompassing the tumour image was chosen and the average pixel intensity was calculated.

**RESULTS**

**Phantom study**

T1 relaxation time of water protons was related to the presence of paramagnetic molecular oxygen dissolved in water. The T1 relaxation time for each water phantom at 25°C was as follows: 3.12 ± 0.06 s without oxygen, 2.83 ± 0.04, 2.36 ± 0.01, 2.20 ± 0.02, 2.09 ± 0.02 s with oxygen under 1.0, 1.5, 2.0 and 2.5 ATA respectively. The relaxation rate \((R1 = 1/T1)\) of pure water without dissolved oxygen was 0.32, and a non-linear relationship \((r^2 = 0.981)\) was observed between \(R1\) and ATA (Figure 1). The phantom study indicated that the T1 relaxation time was shortened by dissolved molecular oxygen under the high-pressure environment.

**Tumour study**

T1-weighted SE images of pre- and post-treatment of HBO exposure revealed the same registration (Figure 2 A,B), and T2-weighted SE images demonstrated no necrotic lesions (Figure 2C). The Gd-enhanced image showed a homogeneous enhanced tumour of 1 cm in diameter (Figure 2D). The pathological specimen stained haematoxylin and eosin (Figure 2E) showed no evidence of haemorrhage or necrosis. Immediately after HBO exposure, IR-FLASH signals from the tumours of HBO-treated mice showed a signal increase in the tumour compared with the pre-HBO image (Figure 2 F–K). Compared to the two baseline IR-FLASH images, the average signals of
tumours exposed to HBO showed 20%, 18%, 15%, 13% and 10% increases and those of the muscles demonstrated 18%, 11%, 5%, 0% and –2% in each image intensity at 5, 15, 30, 60 and 90 min after decompression respectively. There was a logarithmic relationship ($r^2 = 0.929$) between the MR signal intensity of tumours and time. Similarly, the average signals from muscles exposed to HBO showed a logarithmic relationship ($r^2 = 0.946$). In contrast, the signals from the tumours in the normobaric group showed no significant change during the course of measurement with air breathing. It is also noteworthy that the MR signal increase of the tumours lasted over 60 min after decompression in the HBO-treatment group, unlike that of the muscle tissues (Figure 3).

**DISCUSSION**

Non-invasively, we detected that the decline of MR signal intensity of the tumour was slower than that of muscle using T1-weighted imaging during air-inhalation after HBO decompression. The first study using this non-invasive method examined the effect of hyperoxia on T2*-weighted images of rat R3230AC mammary adenocarcinomas (Karczmar et al, 1994). The same group reported that T2*-weighted images differentiated tumours from normal tissue (Kuperman et al, 1995). They reported that significant signal increases were observed within the tumour centre and rim, while little change was observed in muscle during hyperoxia. Using the same T2*-weighted gradient echo images, another study on the responses of six rodent tumours to carbogen (95% oxygen/5% carbon dioxide) suggested that the MR signals were consistent with an increase in oxygen content of blood, tumour cell oxygenation and tumour blood flow (Robinson et al, 1997). On the other hand, using T1-weighted images instead of T2*-weighted images, semi-quantitative measurements of the tumour oxygen level have been discussed (Edelman et al, 1996; Tadamura et al, 1997; Obata et al, 1998). Tadamura (1997) reported that there was no significant change in the T2 value during oxygen inhalation in the tissues, including the spleen and myocardium, in which T1 shortening was observed. These results indicate that T1-weighted imaging is more useful to evaluate the effect of tissue oxygenation compared to T2*-weighted imaging which was affected by blood oxygenation, blood flow and tissue oxygenation.

**Mechanisms affecting MR signal intensity**

Two major mechanisms affect MR signal changes in tissue oxygenation. The first mechanism is blood oxygenation level-dependent (BOLD) contrast based on paramagnetic deoxyhaemoglobin and the second is paramagnetic molecular oxygen itself containing two unpaired electrons. Paramagnetic deoxyhaemoglobin in blood creates magnetic susceptibility gradients near blood vessels that produce phase dispersion of water proton magnetization in the surrounding tissue, so the gradient recalled echo-type sequences are very sensitive to the BOLD effect (Ogawa et al, 1990). This BOLD contrast has been utilized to evaluate regional blood flow and/or tissue oxygenation on functional MR imaging. During oxygen inhalation, the mean enhancement on T2*-weighted brain images in the grey matter and the white matter were 4.23% and 1.92% respectively, but T1-weighted
Tumour oxygen after HBO monitored by MRI

Tumour oxygenation method

Many studies to improve tumour oxygenation have been performed. Some investigators have reintroduced the clinical use of carbogen to improve tumour oxygenation. Carbogen breathing may improve tumour blood oxygenation in two ways: (1) the 95% oxygen may simply increase the arterial blood PO$_2$; (2) 5% carbon dioxide may induce vasodilation of afferent tumour vessels (Robinson et al, 1995). Carbogen breathing caused increases of up to 100% in normalized MR image intensity in GH3 prolactinomas grown in rats, and reversion to air breathing caused a subsequent fall in MR image intensity. These changes in signal intensity are consistent with an increase in oxygen content of the blood, tumour cell oxygenation and blood flow of the tumour. Using the Eppendorf PO$_2$ histography, however, normobaric oxygen and carbogen caused no significant change in tumour oxygenation, whereas HBO and hyperbaric carbogen led to improvement of oxygenation (Brizel et al, 1995). Moreover, hyperbaric carbogen was less effective than HBO in increasing the tumour because of the result of adrenergic stimulation from the inspired carbon dioxide. HBO might be the most effective method to reduce tumour hypoxia by increasing the amount of dissolved oxygen in the plasma and tumour cells.

The changes in tissue PO$_2$ reduction after HBO exposure depend on blood flow and/or oxygen consumption in tissues. Wells (1977), using a mass spectrometer probe that quantified the duration and magnitude of the HBO effect, found that tissue PO$_2$ changed slowly during and after HBO exposure and that the decline in PO$_2$ was slower in subcutaneous tissue than in muscle. They concluded that the different PO$_2$ changes in tissues were affected by differences in tissue perfusion. On the other hand, Hall (1994) emphasized poor tissue perfusion in the presence of hypoxic tumour cells (Thomlinson and Gray, 1955). In this study, an increased oxygen level in the tumour continued for a substantial period despite fast oxygen reduction in the muscle. Although the oxygen metabolism in tumours and muscles was not measured in this condition, one of the reasons for the slower PO$_2$ decline in tumours was probably the lower blood flow in tumours.

Clinical application of HBO

Most malignant tumours appear to have a hypoxic core, and the elimination of this core may destroy cells which are not killed by usual radiation procedures and which may be responsible for recurrence. It is well known that hypoxic tumour cells are resistant to some types of chemotherapy and radiotherapy. Tumour oxygenation is a critical determinant of many forms of cancer therapy. Clinical trials of radiotherapy during HBO showed improvements in the local cure and survival rates of cancers in the head, neck, and uterine cervix and evidence of HBO benefits has been obtained in carcinomas of the bronchus, but not of the bladder (Dische, 1978). Since HBO has an enhancing radiation effect on normal tissues as well as tumours, controversy remains concerning the actual improvement in the therapeutic efficacy of HBO. In addition, the dose correction of radiation absorbed in the chamber wall is complex. But our new protocol that involved irradiation immediately after HBO exposure was simple and safe compared to radiotherapy during HBO exposure (Kohshi et al, 1996). The results of the present non-invasive study supported the theory that elevated MR signal intensity indicates tissue PO$_2$ in the tumours was maintained for substantial periods after decompression. Therefore, irradiation immediately after decompression is considered to be effective for malignant tumours without exerting the influence of radiation on normal tissue. Furthermore, multivariate analysis in our small series revealed that combination with HBO was a good predictive prognostic factor for survival (Kohshi et al, 1999).

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