Correlation between Choline Level and Gd-DTPA Enhancement in Patients with Brain Metastases of Mammary Carcinoma

Paul E. Sijens, Pieter van Dijk, Matthijs Oudkerk

Single voxel $^1$H double spin-echo MR spectroscopy was used to examine 15 cases of brain metastasis of mammary carcinoma (18 lesions) in relation to Gd-DTPA enhanced MR imaging. For lesions larger than 50% of MRS voxel size, there was significant correlation between Gd-DTPA-enhanced MRI signal and MRS-detected signal of choline (Cho) containing compounds ($r = 0.86$, $P < 0.01$; $n = 8$). The observed loss of correlation when including the smaller lesions was overcome by correcting for partial volume effects ($r = 0.69$, $P < 0.002$; $n = 18$). Metastasis spectra showed increased Cho compared with control spectra, except for those lesions showing detectable lactate (Lact) signal. The detection of Lact in four of the larger lesions coincided with comparatively low levels of creatine (Cr) and Cho and heterogeneous Gd-DTPA enhancement (ring-enhancement). It was concluded that in brain metastases of mammary carcinoma Lact represents a product of ischemia preceding/during tissue decay resulting in central necrosis, rather than tumor specific metabolism resulting in increased glycolysis.

Key words: brain metastases; Gd-DTPA contrast agent; magnetic resonance imaging; magnetic resonance spectroscopy.

INTRODUCTION

Gd-DTPA is the most frequently used contrast agent in brain MR imaging (1–3). The $T_1$ shortening metal chelate distributes primarily in the intravascular, extracellular space and quickly passes into the extravascular, interstitial space (4). Used intravenously it provides contrast enhancement in $T_1$-weighted MR images of intracranial lesions associated with an abnormal vascular bed, as is the case in metastatic lesions, or breakdown of the blood-brain barrier (BBB) as is generally the case in the more malignant primary brain tumors (5). Metastatic lesions possess non-CNS capillaries similar to their tissues of origin, so brain metastases virtually always enhance. The use of Gd-DTPA for CNS applications has become routine in distinguishing the extent of lesions from peritumor edema and in detecting residues, recurrence, and growth of primary tumors and metastases (2, 6). However, there has been confusion whether necrotic tissues are specifically enhanced with Gd-DTPA (4, 7, 8) or to the contrary that the necrotic regions in tumors show up as hypointense areas (9, 10).

Localized $^1$H MR spectra of normal brain tissue measured under suppression of the water signal show the resonances of Cho, Cr, and NAA. The Cr peak includes both creatine and phosphocreatine signal and there are still unanswered questions regarding what compounds contribute to the Cho and NAA peaks (11). Brain tumors have relatively high Cho levels and reduced NAA levels (12–19). Whereas Cho level generally increases with tumor size, it cannot be used as a discriminator of tumor grade because necrotic high grade tumors have low levels of Cho (20). Metabolic heterogeneity of human intracranial tumors has been shown (18, 20–22). Some authors have suggested that Lact is concentrated in necrotic tissue (13, 16) or peritumoral edema (22) as opposed to others who correlated the presence of Lact with increased glucose uptake (18).

$^1$H MRS publications to date have focused on the larger brain lesions such as (high grade) primary brain tumors. Brain metastases are generally small and therefore more difficult to characterize by MRS. Single volume MRS was used in this study because of the better quality lesion spectra obtained after shimming on small volumes compared with the results of spectroscopic imaging methods, which require shimming on larger volumes to obtain arrays of spectra including the lesion. To our knowledge $^1$H MRS studies including more than a single case of mammary carcinoma brain metastasis have not been published. New in this study is also the correlation of the $^1$H MRS measured metabolic profiles of brain lesions with Gd-DTPA-enhanced MR imaging.

MATERIALS AND METHODS

Fifteen patients with untreated brain metastases of mammary carcinoma, all showing Gd-DTPA-induced signal enhancement in $T_1$-weighted MRI, were examined by $^1$H MRS. MR examinations were performed at 1.5 T using a 1.5/2 T SP63/84 Helicon whole-body MR scanner (Siemens AG, Erlangen, Germany). The standard circular polarized transmit/receive head coil and 10 mT/m gradient system were used for both MRI and MRS.

Gd-DTPA Enhanced MR Imaging

MRI studies, preceding the MRS by up to 2 weeks, were performed with a $T_1$-weighted sequence (TR/TE 600/15, 2 NEX, matrix size 192 × 256, 3/4 rectangular FOV) and
a $T_2$-weighted sequence ($TR/TE$ 2000/12–80), 1 NEX, matrix size 192 × 256, 3/4 rectangular FOV) in 6-mm slices. $T_2$-weighted MRI was repeated 5 min after administration of Gd-diethylenetriamine pentaacetic acid-dimeglumine (0.2 mmol Gd-DTPA/kg; Schering AG, Berlin, Germany). Brain metastases do not show significant (further) enhancement at later times post-Gd-DTPA contrast (P. van Dijk and M. Oudkerk, works in progress). Routines provided by the manufacturer of the MR system were used to determine the mean signal intensities and standard deviations for the largest transverse 6-mm thickness MRI cross section of each lesion.

**Localized $^1$H MR Spectroscopy**

The use of stimulated echo (STEAM) spectroscopy to select a VOI reliably as well as to suppress the unwanted water signal in a single experimental step has been described in detail (23). In this study the double spin-echo (DSE) sequence rather than the STEAM sequence was used because in our experience the gain in signal is close to the reported 90% (24). In phantom experiments, artifacts due to acetate or Lact signals from outside the VOI were not larger than expected from the calculated chemical shift artifacts of 0.074 and 0.037 ppm at the used slice selection gradients of 2 mT/m (for selecting volumes of 8–36 cm$^3$) and 4 mT/m (for volumes between 1 and 8 cm$^3$), respectively. Care was taken to ensure proper positioning of lesion MRS voxels in those regions that had shown signal enhancement in the Gd-DTPA-enhanced MRI studies (previous paragraph). Analysis of patient spectra was confined to Cho (chemical shift 3.21 ppm), Cr (3.02 ppm), NAA (2.01 ppm), and Lact (doublet centered at 1.32 ppm with coupling constant of 7 Hz) peak integrals. Lipid signals were not quantitated because in peripheral voxels contributions from nonspoiled subcutaneous fat signals were often significant. The presented MRS data and the spectra shown in the figures were all acquired with $TE = 135$ ms. At $TE = 135$ ms, due to $J$ modulation, the doublet signal of Lact is inverted (180° out of phase) relative to the singlets of Cho, Cr, and NAA (25). To confirm the identity of presumed Lact peaks, MRS measurements were repeated with $TE = 270$ ms (Lact doublet in phase). The whole procedure including sagittal and transverse MRI, global shimming, volume selection, local shimming, water suppression optimization and the measurement of two voxels took 60 min. The acquisition of a sagittal localized was followed by proton density and $T_2$-weighted imaging (1500/12–80) using a 55° flip angle, 15 slices of 6 mm without a slice gap, a 128 × 256 matrix and 1 acquisition (3.15 min). The field homogeneity achieved in global and local shimming resulted in water peak line widths of at most 17 Hz and 6 Hz, respectively. SE MR spectra were collected with 2.56-ms sinc-shaped RF pulses preceded by 25.6 ms Gaussian-shaped RF pulses for chemical shift selective excitation (CHESS) and subsequent spoiling of the resultant water signal. The second half of the spin echo was collected using 1024 data points and a spectral width of 500 Hz. All DSE measurements of voxels that varied in size between 1 and 36 cm$^3$, were the sum of 256 acquisitions with four prescans and $TR = 1600$ ms (acquisition time 6.57 min). Typically time domain data were zero-filled to 4096 points, multiplied with a Gaussian function (center 0 ms, half width 256 ms), Fourier transformed, phase corrected and fitted to Lorentzian curves. Because the gradient-induced artifacts in amplitude and phase of the localized RF pulses were minor, the spectra were already of good quality when not corrected for the effects of eddy currents. All MRS data were therefore processed without introducing an additional source of error in the form of eddy current compensation.

A pilot study on six healthy volunteers (aged between 22 and 41 years) demonstrated a fairly constant ratio between the total metabolite signals detected in the occipital lobe (8 cm$^3$ voxel) and in caput nuclei caudati (also 8 cm$^3$), that is $1.09 \pm 0.29$ SD. To get semiquantitative data, metabolite levels have been presented as percentages of the NAA level measured in the contralateral occipital voxel. The same normalizations were made by others (14, 26). Ratios of Cho to NAA and Cr to NAA of 38 ± 6 and 51 ± 13 (percentages ± SD; occipital lobe) and 52 ± 10 and 47 ± 10 (corpus nuclei caudati) in the healthy volunteer study, confirmed the good reproducibility of the MR spectra. Lesion voxels and occipital control voxels had the same dimensions, except for lesion voxels larger than 8 cm$^3$ (the largest cubic volume not exceeding the dimensions of the occipital lobes of the brain). In the latter case differences in voxel size were accounted for by normalization.

**Statistics**

Correlation analyses were applied and Student's $t$ test, two-tailed, was used in the assessment of the significances of the presented linear correlations and of differences between group means.

**RESULTS**

The $^1$H MRS characteristics of the examined metastases of mammary carcinoma are summarized in Table 1. Voxels varying in size between 1 and 36 cm$^3$ were selected to include the whole abnormality. With respect to their $T_2$-weighted MRI behavior on addition of Gd-DTPA, lesions could be divided in two categories: (1) more or less homogeneously enhancing (Cases 1, 2, 3b, 4–9, 11–13, 15) and (2) ring-enhancing, that is nonenhancing areas surrounded by an enhancing shell (Cases 3a, 10, 14). In Table 1 lesion contrast is presented as percent difference between mean lesion signal and normal tissue signal at the largest MRI cross section (6-mm slice thickness), and lesion heterogeneity by the included standard deviations. There was a significant linear correlation between total lesion volume as estimated from postcontrast MRI series, and signal heterogeneity ($r = 0.59, P < 0.02; n = 18$).

MRS detected metabolite levels are presented in Table 1 as percentages of the integrated NAA signal in the contralateral occipital voxel measured in the same patient examination. Lesion-containing voxels generally revealed high Cho, normal Cr, and reduced NAA levels compared with the control voxels, which had Cho to
NAA and Cr to NAA ratios similar to those encountered in healthy volunteers (Cho = 38 ± 6 and Cr = 51 ± 13 SD relative to NAA = 100). The 1H MR spectrum of a voxel containing a typical brain metastasis of mammary carcinoma (Case 2) is shown in Fig. 1. The level of Cho is mildly increased, Cr within the normal range and NAA slightly decreased; Lact is not detected. The Gd-DTPA-enhanced MRI of the same lesion shows a small area of comparatively homogeneous signal enhancement. The mean MRI signal intensity in the whole enhanced area was 610 ± 60 SD arbitrary units (AU) compared with 472 ± 9 AU in contralateral brain tissue. The percent difference between both values, 29%, is listed in Table 1 ("contrast"). Lesion center signal was measured in a circular ROI covering half the total area (637 ± 28 AU). The quotient between the latter standard deviation, not influenced by the signal drop on the outer rim, and control signal was used as measure of lesion heterogeneity: 28/472 = 0.06 or 6% (Case 2, Table 1). The results for a large brain metastasis showing more heterogeneous Gd-DTPA enhancement are shown in Fig. 2 (Case 4a). The level of Cho in the voxel containing the metastasis is very high, 168% of control NAA level. Lact was not detected. The third example is a large brain metastasis showing highly heterogeneous Gd-DTPA enhancement, "ring-enhancement," in T1-weighted MRI (Fig. 3, Case 14). The Cho level, 55% of control NAA, is comparatively low and Lact signal is present. The 1H MR spectra of the whole lesion and its mostly hypointense center, upscaled with a factor of 4.5 to correct for the difference in MRS voxel size, are shown in Fig. 3. In the center of the lesion, the levels of Cr and NAA are very low and Lact (the inverted doublet at about 1.27 and 1.38 ppm) twice as high as in the lesion containing voxel. Alanine, reportedly present in meningiomas measured under comparable experimental conditions (16, 17), was detected in none of the 18 MRS examined brain metastases of mammary carcinoma.

### Choline and Creatine

For lesions larger than 50% of the MRS voxel size (eight out of 18) significant correlation was found between (total) Cho level and the lesion MRI signal after administration of Gd-DTPA ($r = 0.86$; $P < 0.01$). Inclusion of smaller lesions leads to loss of correlation which may be attributed to dilution of results due to higher normal tissue contributions. Correlations between Cr level and either Gd-DTPA enhancement, lesion size, or Cho level were not found.

#### Table 1

<table>
<thead>
<tr>
<th>Case</th>
<th>Gd-DTPA enhancement MRI</th>
<th>MRS detected metabolite levels</th>
<th>Lesion metabolite levels divided by filling factor</th>
<th>Lipid</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Lesion volume (cm$^3$)$^a$</td>
<td>Contrast$^b$ ± SD$^c$</td>
<td>Cho Cr NAA Lact</td>
<td>Cho Cr Lact</td>
</tr>
<tr>
<td>1</td>
<td>6.2 25 ± 8</td>
<td>65 20 18 0</td>
<td>56 11 0</td>
<td>72 14 0</td>
</tr>
<tr>
<td>2</td>
<td>1.3 29 ± 6</td>
<td>73 47 75 0</td>
<td>35.5 9.5</td>
<td>218 58 0</td>
</tr>
<tr>
<td>3a</td>
<td>7    -13 ± 24</td>
<td>48 25 33 61</td>
<td>31.5 6.5</td>
<td>26 10 70</td>
</tr>
<tr>
<td>3b</td>
<td>1.1 2 ± 5</td>
<td>40 28 42 0</td>
<td>19 7 0</td>
<td>59 22 0</td>
</tr>
<tr>
<td>4a</td>
<td>12    46 ± 12</td>
<td>168 53 68 0</td>
<td>134 19 0</td>
<td>201 28 0</td>
</tr>
<tr>
<td>4b</td>
<td>0.4    28 ± 9</td>
<td>129 64 112 0</td>
<td>73 8 0</td>
<td>3285 360 0</td>
</tr>
<tr>
<td>5</td>
<td>0.07   82 ± 6</td>
<td>48 59 67 0</td>
<td>14.5 25.5</td>
<td>1657 2914 0</td>
</tr>
<tr>
<td>6</td>
<td>2.6    47 ± 12</td>
<td>84 89 100 0</td>
<td>34 39 0</td>
<td>353 143 0</td>
</tr>
<tr>
<td>7</td>
<td>3.1    96 ± 10</td>
<td>66 10 0 0</td>
<td>66 10 0</td>
<td>170 3 0</td>
</tr>
<tr>
<td>8</td>
<td>4.2    87 ± 27</td>
<td>199 19 25 0</td>
<td>186.5 6.5</td>
<td>355 12 0</td>
</tr>
<tr>
<td>9</td>
<td>3.6    90 ± 26</td>
<td>103 48 74 0</td>
<td>66 11 0</td>
<td>147 24 0</td>
</tr>
<tr>
<td>10</td>
<td>12    -6 ± 51</td>
<td>19 17 29 15</td>
<td>4.5 2.5 15</td>
<td>8 5 28</td>
</tr>
<tr>
<td>11</td>
<td>1.4    26 ± 7</td>
<td>78 37 17 0</td>
<td>69.5 28.5</td>
<td>169 69 0</td>
</tr>
<tr>
<td>12</td>
<td>19    65 ± 18</td>
<td>97 10 13 35</td>
<td>92 5 35</td>
<td>115 6 44</td>
</tr>
<tr>
<td>13a</td>
<td>8.6    71 ± 28</td>
<td>43 67 34 0</td>
<td>26 50 0</td>
<td>82 157 0</td>
</tr>
<tr>
<td>13b</td>
<td>5.0    34 ± 18</td>
<td>76 47 53 0</td>
<td>49.5 20.5</td>
<td>79 33 0</td>
</tr>
<tr>
<td>14</td>
<td>22    17 ± 35</td>
<td>55 50 48 37</td>
<td>31 26 37</td>
<td>51 43 61</td>
</tr>
<tr>
<td>15</td>
<td>0.3    49 ± 13</td>
<td>45 49 58 0</td>
<td>16 20 0</td>
<td>53 67 0</td>
</tr>
</tbody>
</table>

$^a$Total lesion volume estimated from postcontrast MRI series.

$^b$Lesion contrast, i.e., percent difference between postcontrast lesion signal (at largest MRI cross-section) and the signal of contralateral brain tissue.

$^c$Lesion heterogeneity, standard deviation in postcontrast lesion MRI signal in percentage of contralateral brain tissue signal.

$^d$Calculated by subtracting the normal brain tissue contributions from the overall 1H MRS detected signals shown in previous columns.

$^e$filling factor, lesion volume/MRS voxel size.

$^f$Absent (-), low (+), high (++).

$^g$Major part of lesion located extracerebral.

$^h$Center of the same lesion.
Correcting for Partial Volume Effects

Partial volume effects, the lesions being smaller than the voxels measured by $^1$H MRS, obscure the characterization of metastases of mammary carcinoma. Because NAA is generally accepted to be localized in neurons (11, 27), it is reasonable to assume that brain metastases do not contain NAA. When a second assumption is made, that the NAA in the spectra comes with normal brain tissue Cho and Cr signals in a ratio of 2:1:1 (roughly the proportions obtained in occipital brain lobe and in corpus nuclei caudati of volunteers), the net lesion metabolite levels can be calculated: “Lesion Cho” = Cho − $\frac{1}{2}$NAA, “Lesion Cr” = Cr − $\frac{1}{2}$NAA, “lesion Lact” = Lact. Post-contrast MRI signal is corrected for partial volume effects by multiplication of the contrast with the filling factor:

$$\text{Filling factor \cdot Contrast} = \frac{(\text{Lesion volume})}{(\text{MRS voxel size})} \cdot \frac{(\text{lesion MRI signal} - \text{control MRI signal})}{(\text{control MRI signal})}$$

with control MRI signal set at 100.

After applying the above corrections on all lesions, a significant positive correlation between the lesion Cho signal (“Lesion metabolite level = Cho” in Table 1) and the postcontrast MRI signal is found. The resulting linear regression is shown in Fig. 4 ($r = 0.69, P < 0.002; n = 18$). Lesion metabolite signals per unit of MRI visualized lesion volume are shown in the right-most columns of Table 1, “lesion metabolite level/filling factor.” In the four lesions showing the weakest MRI enhancement (to intensities of −13% to +17% relative to normal brain tissue), the thus calculated Cho levels are not significantly higher than contralateral Cho (Cases 3a, 3b, 10, 14).
**FIG. 3.** $^1$H MR spectra of 36 cm$^3$ voxel containing large brain metastasis of mammary carcinoma and of 8 cm$^3$ voxel positioned in the largely hypointense center of the same lesion (the latter spectrum upscaled by a factor of 4.5) and axial $T_1$-weighted image after Gd-DTPA administration (SE 600/10) showing a profile through the R-parietal lesion (Case 14). In addition to Cho, Cr, and NAA, there are Lact doublets at 1.27 and 1.38 ppm.

**Lactate**

Quantitation of Lact was straightforward in the majority of lesions where lipid signals were absent or low (13 out of 18). The MR spectra of the other five lesions showed intense lipid signals in phase with the Cho, Cr, and NAA signals (i.e., Case 4a, Fig. 2) without a trace of inverted doublet Lact signal (1.32 ppm) interfering with the comparatively broad -CH$_2$- lipid resonance (1.3 ppm). The Lact coupling constant of 7 Hz and the water line width of at most 6 Hz achieved in the local shimming of any metastasis containing voxel (see Materials and Methods section), make the possibility of significant amounts of Lact not being recognized and confused with lipid signal highly unlikely. Compromising of any Lact level listed in Table 1, also the zeros in Cases 4a, 8, 11, and 13ab, by the presence of lipid is therefore negligible. Lact was detected in heterogeneously enhancing lesions (Cases 3a, 10, 12, 14), most of which had comparatively low Cho levels (Cases 3a, 10, 14) and very low Cr levels (Cases 3a, 10, 12).

**DISCUSSION**

For lesions larger than 50% of MRS voxel size (8/18), there was significant correlation between Gd-DTPA-enhanced MRI signal and MRS-detected choline (Cho) signal ($r = 0.86, P < 0.01$). The observed loss of correlation when including the smaller lesions was overcome by correcting for partial volume effects ($r = 0.69, P < 0.002, n = 18$; Fig. 4). The knowledge that Cho and other metabolites are not detected in fully necrotic tissue, warrants the conclusion that our data do not support the hypothesis that necrotic tissues are specifically enhanced with Gd-DTPA (4, 7, 8), but to the contrary confirm that the necrotic regions in tumors show up as hypointense areas (9, 10).

Significant linear correlation between total lesion volume, as determined from postcontrast MRI, and signal heterogeneity was shown ($r = 0.59, P < 0.02; n = 18$). Lact was only observed in large heterogeneously enhancing (ring-enhancing) metastases (Cases 3a, 10, 12, and 14) and with preference for the lesion center (Case 14; Fig. 3). This implies that Lact is only present in late stage lesions that contain large necrotic regions. The comparatively low lesion Cr and Cho levels at that stage are in line with
reduced Cho levels in necrotic high-grade gliomas compared with solid high-grade gliomas (20). We conclude that in brain metastases of mammary carcinoma, the cancer cells do not preferentially ferment glucose to Lact, in other words the "Warburg effect" (28, 29) does not apply. The detected Lact is a product of ischemia preceding/during tissue decay and the formation of central necrosis and does not reflect tumor-specific metabolism resulting in increased glycolysis.

These results are in agreement with suggestions that Lact might be concentrated in necrotic tissue (28, 29) and contradict a reported correlation of the present of Lact in intracranial tumors with high Cho levels and increased glucose uptake (18). In our experience edema is characterized by normal Cho and Cr levels and NAA reduced by up to 50% (compared with normal brain tissue) and, in contrast to an assumption published by others (22), by absence of Lact signal. The presented lesion metabolite signals have not been corrected for the (not easily quantifiable) amount of edema rather than normal brain tissue included in the MRS voxels. In the volume-corrected lesion metabolite levels, it has been assumed that the whole lesion is visualized with Gd-DTPA-enhanced MRI (last columns in Table 1). The unlikely high Cho and Cr levels in two of the smallest lesions might be interpreted as evidence that early stage brain metastases of mammary carcinoma are only partially visualized with Gd-DTPA-enhanced MRI (Cases 4b, 5).

This study has contributed both to understanding of tumor pathophysiology and to the evolving science of metabolic fingerprinting of pathophysiological processes by MRS. In characterizing metastatic brain tumors, generally small lesions when detected for the first time, single voxel 1H MRS is a perfectly feasible method if not the method of choice. Only in four out of 18 lesions studied, those exceeding 2.5 cm diameter (Cases 4a, 10, 12, 14), spectroscopic imaging with a volume resolution of 1 cm³ would have yielded additional information regarding tumor heterogeneity.

ACKNOWLEDGMENTS

Some of the presented data are part of the "Clinical Trial Proton Spectroscopy" initiated by Dr. R. Sauter, Siemens AG, Erlangen, Germany. The authors thank D.J.A. Kraus for his participation in the collection of the data.

REFERENCES


