In vivo measurement of phosphorous markers of disease

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Abstract. Phosphorus Magnetic Resonance Spectroscopy (31P-MRS) has been utilized to study energy, carbohydrate, and phospholipid metabolism in vitro and in vivo in live tissues non-invasively. Despite its lack of sensitivity, its application has extended to in situ human tissues and organs since proper signal localization was devised. Follow-up of phosphocreatine in neuromuscular diseases and schizophrenia and follow-up of phospholipid-related molecules in tumors are described here to demonstrate the value of 31P-MRS as an imaging technique to determine in vivo markers of disease and in the diagnosis, prognosis, and follow-up of human diseases.

Keywords: 31P MR spectroscopy, chemical shift imaging, in vivo metabolism, neuromuscular disease, mitochondrial cytopathy, schizophrenia, non-Hodgkin’s lymphoma, chronic lymphocytic leukemia

Abbreviations: NMR, nuclear magnetic resonance; MRI, magnetic resonance imaging; MRS, magnetic resonance spectroscopy; CSI, chemical shift imaging; Etn-P phosphoethanolamine; Cho-P, phosphocholine; Pi, inorganic phosphate; Gro-P-Etn, glycerophosphoethanolamine; Gro-P-Cho, glycerophosphocholine; NTP, nucleotide triphosphates; NDP, nucleotide diphosphates; UDP, uridine diphosphate; PCr, phosphocreatine; PL, phospholipids; SNR, signal to noise ratio; NOE, nuclear Overhauser enhancement; RF, radiofrequency; ND, neuromuscular diseases; MC, mitochondrial cytopathy; ATP, adenosine triphosphate; CoQ10, Coenzyme Q10; BN, below-normal; NN near-normal (memory scores); NHL, non-Hodgkin’s lymphoma; IPI, international prognostic index; TTF, time to treatment failure; CCL, chronic lymphocytic leukemia

1. Historical background

In the past three decades, nuclear magnetic resonance (NMR), an analytical technique initially utilized to elucidate the structure and composition of molecules, has become a methodology that allows the study of live samples. The basic principle of NMR is that nuclei with a nuclear spin in a magnetic field show resonances at various frequencies proportional to the magnetic field. The strengths of these signals are proportional to the amount of their originating nuclei, and their frequencies depend on the molecule in which they are located. Many of these signals can be observed from living systems because the microenvironment inside cells and tissues resembles free solutions on a microscopic scale. Initial observations of living systems consisted of cells and isolated organs in high field, small bore magnets [1, 2]. With the development of magnetic resonance imaging (MRI) technology, large magnets at intermediate magnetic fields have become available, and these studies have been extended to humans. Paralleling the developments in magnetic technology were similar improvements in spectral localization techniques. Localization began with MRI, which primarily uses the strong 1H signal of water in biological tissues and occasionally the 1H signal from the methylenes of fatty acids. This development led to localized in vivo magnetic resonance spectroscopy (MRS) a few years later with the development of single- and double-volume localization techniques. The early work was well re-
viewed by Bottomley and colleagues [2]. More complex multiple-volume methodology for spectroscopy, chemical shift imaging (CSI), capable of isolating spectra in three dimensions, was developed by Brown and colleagues in 1983 [3]. Recently, even higher magnetic fields (4–7 T) have become available, which suggests substantial improvement in spectral and image resolution in vivo in the future.

The most extensively used nuclei in the study of live samples by MRS have been phosphorus-31 (31P), carbon-13 (13C), and hydrogen-1 (1H). In every case, the nuclear spins that can be observed are those in relatively small molecules (<5–10 kD) that can tumble in solution sufficiently quickly to produce a sharp resonance. The natural abundance of 31P is 100% of the available pool of phosphorus molecules, but its spectroscopic sensitivity is lower than 1H, which allows visualization of phosphorus-containing molecules with a concentration in the millimolar range. However, the molecules in living matter that can be visualized by 31P MRS are very important because they are linked to bioenergetics, the metabolism of carbohydrates, and the turnover of phospholipids and membranes. Note that, though phospholipids, membranes, polynucleotide molecules (i.e., DNA, RNA), and even bone constituents have a large amount of phosphorus, they are not observed as sharp lines because they are not able to tumble rapidly enough. On the other hand, 13C has a natural abundance of only 1.1% and its sensitivity is slightly lower than 31P. The natural abundance signals of 13C in living matter show only storage molecules with repetitive units (i.e., glycoprotein and fatty acids). However, experiments with 13C-enriched molecules have demonstrated intermediates and end products of metabolic pathways from which many inferences about metabolic regulation of such pathways can be drawn. Finally, 1H is the most sensitive of the resonant nuclei. Its natural abundance is almost 100%, and almost all biological molecules have hydrogen. The high sensitivity and abundance issues could allow visualizing many biologically important molecules but a small range of resonant frequencies makes it harder to distinguish signals from different compounds. The very large signals from hydrogen in water and lipids in living samples that obscure visualization of other 1H signals from metabolites of much lower concentration are another drawback. Nevertheless, the fact that MRI is based on 1H signals from water and lipids has made the clinical imagers accessible to collect 1H-MRS. This is why proton spectroscopy has been more available for in vivo applications in humans. Although the nuclei listed above (i.e., 31P, 13C, 1H) have been the dominant ones applied to studies in vivo, other nuclei also have demonstrated potential to be used in vivo (i.e., nitrogen-15, fluorine-19, and sodium-23) [4]. In this review, we focus on the applications and advancements of 31P MR spectroscopy in the study of biological samples in vivo, with special regard for the possible role of this technique in identifying markers of disease in humans.

2. Spectral information contained in phosphorus MRS of live samples

Phosphorus MRS (31P-MRS) initially was believed to have the best potential for use in biological research due to the nature of the biological information obtained. An example of this is shown in the spectrum at the bottom of Fig. 1, which is an 1H-decoupled 31P MR spectrum of a perchloric acid extract of human lymphocytes from a normal volunteer acquired at 9.4 T (see figure caption for acquisition parameters) [5]. Four expanded regions of the same spectral area are shown in the lower portions of insets A–D. This well-resolved spectrum is due to the higher magnetic field strength and the highly homogeneous aqueous solution of the extract. In the lower spectrum of the main panel of Fig. 1, the signals of the phospholipid-related phosphomonesters, phosphoethanolamine (EtN-P) and phosphocholine (Cho-P), are shown in extracted normal lymphocytes. The larger signal downfield from EtN-P (3.0 ppm) in the spectrum corresponds to inorganic phosphate (Pi). The Pi concentration in this particular spectrum is biologically meaningless due to contamination by the phosphate-based buffers utilized in the extraction procedures [5]. The smaller signals between Cho-P and Pi present in the spectrum have not been identified. Inset A shows the region where phosphodiesters (mainly glycerophosphoethanolamine [Gro-P-Etn] and glycerophosphocholine [Gro-P-Chol]) resonate. Inset B shows the resonant area of the terminal phosphates of nucleotide polyphosphates. In living tissues, these signals mainly are the resonances of the γ phosphate of nucleotide triphosphates (γ-NTP) and the β phosphate of nucleotide diphosphates (β-NDP). These signals are split in two peaks (doublets) due to their interaction with the neighboring phosphorus in the anhydride chain. Inset C in Fig. 1 shows the doublet signals of the initial phosphates (α) of the anhydride chain in NDP and NTP and the phosphates of several diphosphodiesters (i.e., UDP-sugars, nicotine adenine dinucleotide) not fully identified (region X),
Finally, inset D shows the triplet signal of the middle phosphate of the anhydride chain in NTP (β-NTP). In contrast, spectrum A of Fig. 2 (from [6]) shows the $^{31}$P MR signals from a whole human head in vivo at 1.5 T (see figure caption for acquisition parameters). As expected, the more inhomogeneous conditions of study and the lower magnetic field strength resulted in lower spectral resolution. Although in different ratios, the signals in Fig. 1 also are present in Fig. 2. Conversely, spectrum A in Fig. 2 shows signals that are not present in Fig. 1. Phosphocreatine (PCr) is selectively present, depending on the particular tissues involved (i.e., brain, muscle, and heart), but is not present in lymphocytes.

The broad phospholipid (PL) signal under the sharp signals from spectrum A in Fig. 2, although present in variable concentrations in all tissues, is lost in an acid extract such as the one used to obtain the spectra shown in Fig. 1. Furthermore, spectrum B in Fig. 2, acquired from the same human head, does not show the PL signal because an off-center Gaussian pulse as described elsewhere [6] was used to saturate this broad signal, which in turn has improved the baseline of the sharp signals.

The signals from the normal tissue spectra shown in Figs 1 and 2 exemplify the information that $^{31}$P MR spectroscopy can provide: Cho-P, Etn-P, Gro-P-Cho, Gro-P-Etn, and PL are metabolites in the phospholipid and membrane turnover; UDP-sugars are intermediates in the carbohydrate metabolism; and NTP, NDP, Pi, and PCr participate in bioenergetics. Furthermore, due to the interaction of Pi with free protons ($\text{H}^+$) and NTP with $\text{H}^+$ and magnesium ($\text{Mg}^{2+}$) and the near-equilibrium of these interactions in biological tissues, the Pi and NTP signals modify their resonant frequencies according to the concentrations of $\text{H}^+$ (pH) and $\text{Mg}^{2+}$ in vivo [7,8]. However, these interactions are lost when a tissue is extracted to obtain an aqueous solution.

3. Recent technical advances for human applications of in vivo $^{31}$P-MRS

3.1. Three-dimensional spectroscopic localization

Since its original publication in 1982 [3], chemical shift imaging (CSI) has been the technique of choice to carry out multivolume MRS localization irrespective of the nucleus studied. Since then, more than 250 MRS works have been published that based their spatial localization on CSI techniques. CSI is a phase-encoding technique that avoids the spatial misregistration of spectral signals that affect localization techniques based on frequency selective radiofrequency (RF) pulses in a constant gradient [9]. Aside from the improved signal localization, the CSI experiment can be designed to obtain the spatial distribution of spectra in localized volumes (voxels) in one, two, or three dimensions [3,9,10]. When $^{31}$P MRS is acquired by means of three-dimensional (3D) CSI, the spectroscopic data can be prescribed and aligned in the three dimensions with anatomic features present in MR images by means of the voxel-shifting capability of the Fourier transform [10]. This means that exact preselection of the volume of interest is not necessary, which in turn simplifies data acquisition and reduces machine usage.

An example of localized $^{31}$P MRS of the human brain using 3D-CSI is shown in Fig. 3 (from [11]). After collecting reference images and adjusting the magnetic field shims, the 3D-CSI $^{31}$P dataset was acquired and processed as described in the caption of Fig. 3. On the left of Fig. 3, a transverse MR image of the head has been overlaid with the corresponding localization grid of the CSI. In addition, spectra localized entirely in brain volumes (voxels) were subsampled and overlaid on the image. On the right of Fig. 3, the rightmost spectrum of the grid has been magnified. The quality of the localized spectra is indicated by the flat baseline, separation of the signals in the phosphomonoesters (Etn-P and Cho-P) and phosphodiesters (Gro-P-Etn and Gro-P-Cho) regions, and a signal-to-noise ratio (SNR) larger than 2 for the Pi signal.

3.2. Double-tuned coils

The low sensitivity of $^{31}$P-MRS makes it important to obtain the highest SNR for its in vivo applications. This can be achieved, at least in part, by obtaining a highly homogeneous static magnetic field ($B_0$-field), which usually is adjusted using the $^1$H water signal as a reference. Further, we have proved previously that some heteronuclear schemes enhance the in vivo sensitivity of $^{31}$P signals by $^1$H excitation (i.e., $^1$H decoupling, nuclear Overhauser enhancement [NOE], and polarization transfer) [12,13]. We also have proved that, to obtain the most benefit from these heteronuclear schemes, the adjustment of the $B_0$-field needs to be thorough. We have developed a fast and reliable automated method for this based on $^1$H CSI localization [14]. As stated before and shown in Figs 3 and 4, it also is necessary to collect $^1$H MR images to anatomically place the in vivo localized $^{31}$P signals. To facil-
Fig. 1. $^{31}$P MR spectroscopy of perchloric acid extracts of human lymphocytes. Examples of $^{31}$P spectra from normal human lymphocytes (bottom spectrum; corrected pellet weight = 0.16 g) and chronic lymphocytic leukemia (CLL) lymphocytes (top spectrum; corrected pellet weight = 0.65 g) acquired and processed as described in [5]. $^1$H-decoupled $^{31}$P MR spectra were obtained at 25°C and 162 MHz (9.4 T). A 16 K free-induction decay (FID) was acquired in 1.02 s using a 45° pulse (8.6 µs), broadband composite-pulse $^1$H-decoupling, and a total repetition time of 2.7 s. One to three 225-minute acquisitions with 5,000 transients were accumulated for each sample as needed to achieve an adequate signal-to-noise ratio. Chemical shifts ($\delta$) were referenced to glycerophosphocholine (Gro-P-Cho; 0.494 ppm at pH 8.0). Considering that the concentration of nucleotide triphosphates (NTP) is constant in CLL and normal lymphocytes, the triplet of the $\beta$ phosphate of NTP was used to scale the spectra (inset D) to ease comparisons. Due to the lower pellet weight and the scaling of the spectra, the noise in the normal lymphocyte spectrum appears larger. In the main panel, the signals of phosphoethanolamine (Etn-P) and phosphocholine (Cho-P) are labeled. The insets show expanded regions of the spectra. Inset A shows the region where the phosphodiester signals resonate (1.2–0.3 ppm). Glycerophospho-ethanolamine (Gro-P-Etn) and glycerophosphocholine (Gro-P-Cho) are labeled in the inset. Inset B shows the spectral region where the doublets of the terminal phosphates of the NTP ($\gamma$-NTP) and nucleotide diphosphates (NDP) moieties ($\beta$-NDP) resonate ($-5.3$ and $-6.0$ ppm), while inset C shows the region where the doublets of the $\alpha$-phosphates from the NTP and NDP moieties resonate ($-9.8$ and $-12.8$ ppm). Several derivatives of diphosphonucleosides and diphospho-dinucleotides also resonate in the area shown in inset C (X and Y regions). Inset D shows the spectral region where the triplets of the $\beta$-phosphate of the NTP moieties resonate ($-20.8$ and $-21.4$ ppm).

To mitigate all of these $^1$H requirements during a $^{31}$P study, our team designed several double-tuned RF antennas with highly sensitive $^{31}$P and $^1$H channels. Moreover, with these double-tuned antennas, $^{31}$P and $^1$H spectroscopy can be carried simultaneously, thereby minimizing adjustment and image requirements [15–17]. A comparison of the double-tuned probes with conventional, single-tuned ones showed that, despite double tuning, there is no significant loss in $^{31}$P sensitivity when the $^1$H channel provides the needed performance. For studies at 1.5 T, volume probes using a four-ring birdcage design were developed with the resonators operating in quadrature mode to provide improved sensitivity, excellent B1 homogeneity, and reduced pow-
Fig. 2. $^{31}$P MR spectroscopy from a normal human head. Proton-decoupled, NOE-enhanced $^{31}$P spectra from a normal human head were collected at 1.5 T (TR = 1,000 ms, spectral width = ± 1,000 Hz, 256 acquisitions, and a 45° rectangular pulse). The chemical shift ($\delta$) is expressed in parts-per-million (ppm), using phosphoric acid as the reference at 0 ppm (internal reference $P_\alpha$ of NTP at $-10.01$ ppm). Assignments: PCr, phosphocreatine, the rest as in Fig. 1. In spectrum A, the broad signal under the sharp resonance corresponds mainly to phospholipids. This broad signal was cancelled out during acquisition in spectrum B by applying an off-center Gaussian pulse ($-300$ Hz) that saturated this signal as described elsewhere [6].

Simultaneous application of $^1$H decoupling and NOE of in vivo $^{31}$P signals significantly improves SNR and enhances spectral resolution [12]. We have acquired $^1$H-decoupled, NOE-enhanced $^{31}$P spectra localized to defined regions of different human tissues in vivo [12, 15,18,22]. The usual scheme to obtain $^1$H-decoupled and fully NOE-enhanced $^{31}$P spectra from human tissues in vivo includes the use of Waltz-4 modulation for proton decoupling and continuous wave bi-level excitation for NOE [12]. An example of the application of $^1$H-decoupling and NOE to $^{31}$P signals is shown in Fig. 4. Localized spectra of a non-Hodgkin’s lymphoma tumor in vivo are shown before (lower spectrum) and after (upper spectrum) $^1$H-decoupling and NOE were applied. Higher SNR and better resolved peaks in the phosphomonoester (Etn-P and Cho-P) and NTP regions were obtained in the $^1$H-decoupled, NOE-
enhanced spectrum without processing for resolution enhancement. This demonstrates that \(^1\)H-decoupling and NOE of \(^{31}\)P signals permits obtaining more information about the *in vivo* metabolism of human tissues than was possible previously and should enhance the utility of this technique for studying human disorders.

4. **In vivo markers of disease using \(^{31}\)P MRS**

Since the advent of larger bore spectrometers, human studies in healthy individuals have been carried out showing the usefulness of *in vivo* \(^{31}\)P MRS to study normal muscle [23–33,4,35–53], liver [54,55], kidney [56], brain [4,7,57–62], testes [63], spleen [64], skin [65,66], heart [67,68], breast [69], uterus [70], and blood [71] and references therein. These studies have attempted to give actual molar concentration to phosphorus metabolites and magnesium, find metabolic fluxes of phosphorus compounds under particular conditions, and determine the normal intracellular pH values on the tissues studied. Examples of markers of disease using \(^{31}\)P MRS can be found throughout the literature in which the levels of \(^{31}\)P metabolites and/or their fluxes are modified due to specific illnesses. In *vivo* \(^{31}\)P spectral alterations in acquired and genetic pathological human muscle conditions such as dermatomyositis and polymyositis [72], muscle dystrophies [73–76], ischemia [77], sporadic inclusion body myositis [78], progressive supranuclear palsy [79], malignant hyperthermia [80], fibromyalgia [81], putaminal psoriasis [82], retinitis pigmentosa [83], Parkinson disease [84], McArdle’s syndrome [85], adenyl-succinate lyase deficiency [86], hypo-ß-lipoproteinemia [87], and some mitochondrial myopathies (cytochrome bc1 deficiency [88], carriers of 11778 mtDNA mutation [89], NARP syndrome [90], phosphofructokinase deficiency [91,92], Leber disease [93–95], and pyruvate dehydrogenase complex deficiency [96] have been reported. The usual change found using \(^{31}\)P MRS in these diseases is a reduction in the bioenergetics of the tissue. In addition, the effects of lipoic acid [97] and coenzyme Q10 [98,99] as treatment for mitochondrial cytopathies (MC) have been reported. Q10 has been shown to be especially promising for the treatment of mitochondrial diseases and perhaps for other diseases that involve compromised bioenergetics. The effects of thyroid hormones [100,101], insulin [102], exertional heatstroke [103], chronic fatigue syndrome [104], uremia [105], and phosphocreatine treatment [106] on muscle metabolism demonstrated by \(^{31}\)P MR spectroscopy also have been reported.
In the brain, markers of disease have been demonstrated by $^{31}$P MR spectroscopy in a number of organic diseases (i.e., seizures [107], neonatal intraventricular hemorrhage [108], migraine [109,110], head injury [111], lupus erythematosus [112], infarction [113], multiple sclerosis [114], hepatic encephalopathy [115], hydrocephalus [116], progressive supranuclear palsy [80], retinitis pigmentosa [83], epilepsy partialis continua [117], hypoxia-lipoproteinemia [87], cluster headache [118], and MC [89,95,119,120]) as well as the brain effect of the treatment of MC with coenzyme Q$_{10}$ [99]. Further, a group of diseases that have not been considered with a formal organic origin also have shown alteration on the in vivo $^{31}$P MR spectroscopy of the brain usually characterized by changes in bioenergetics or phospholipid content (i.e., schizophrenia [121–130], bipolar disease [131–133], affective disorders [134], Alzheimer’s disease [135–137], autism [138], depression [139], and dislexia [140]). Other human tissues with pathological conditions studied by in vivo $^{31}$P MR spectroscopy have included liver (cirrhosis [141–145], diffuse hepatic disease [146], alcoholic disease [147–149], acute alcohol abstinence [150], hepatitis A infection [151], chronic hepatitis and its response to treatment [152], and the acute effect of ethanol and fructose [153]); bone (osteomalacia [154] and osteoporosis [155,156]); and heart (cardiomyopathy [157,158], coronary disease [158–160], mitral stenosis [161], and Friedreich ataxia [162] and its treatment with antioxidants [163]). Transplanted kidney [164,165] and psoriasis of the skin [166] also have been studied.

In vivo $^{31}$P MR spectroscopy of human tumors deserves special consideration. The characteristic $^{31}$P spectral pattern of tumors seen in vitro, which, unlike the majority of diseases studied is not related to bioenergetics but is instead related to modifications of intermediates of phospholipid metabolism, al-
so has been found \textit{in vivo}. The pattern consists of relatively high concentrations of Etn-P and Cho-P. Several general reviews have been published on the subject [167–172], as well as papers on specific tumor histopathologies (lymphomas [173–176]; sarcomas [177–180]; and breast [69,181–185], lung [186, 187], brain [188], prostate [189], and liver carcinomas [190]) as well as follow up of anticancer treatment [191–195].

In the following sections, specific examples of the study of markers of disease using $^{31}$P MR spectroscopy in MC, schizophrenia, non-Hodgkin’s lymphoma, and chronic lymphocytic leukemia are described.

4.1. $^{31}$P MRS studies in mitochondrial cytopathies

Due to their initial non-specific symptoms, differential diagnosis of MC must be done against the rest of the ailments grouped as neuromuscular diseases (ND). The ND group includes a variety of pathologies of acquired and genetic origin, common among children, with complicating identification and characterization due to overlapping clinical manifestations that delay instituting specific treatment. In the case of MC, the usual underlying alteration is a genetic mutation that produces malfunctions of the oxidative phosphorylation, with concomitant reductions in the cellular energy state. This energy deficit is demonstrated by decreased synthesis capacity for adenosine triphosphate (ATP, the most abundant intracellular NTP) by the respiratory chain, which in turn affects all tissues; especially those that are highly active (i.e., muscle, heart, nervous system). Although measurement of respiratory and phosphorylating activities in intact mitochondria isolated from the muscle of patients with ND is a very powerful \textit{in vitro} method of identifying MC, the heterogeneity of these diseases can prevent their diagnosis using this methodology. It has been recognized that the non-invasive nature of $^{31}$P MRS and its ability to visualize NTP, NDP, Pi, and PCR and their changes during exercise make it a promising technique to aid MC diagnosis and to monitor therapy [24,27,31,32,34,36,40,42,50,94,99,196,198].

A simple protocol of \textit{in vivo} $^{31}$P MRS of active muscle can be used in the differential diagnosis of MC in pediatric patients with ND symptomatology. Under this protocol, we studied 23 ND-diagnosed patients (4–17 years-old) and 12 age-matched healthy volunteers with 100% compliance. Twelve of the ND patients already were confirmed as having MC based on isolated mitochondria studies and/or formal genetic determina-

4.2. Brain studied by $^{31}$P MRS in schizophrenia [11]

Changes in schizophrenia have suggested that the physiopathology of the disease probably is due to a low energy state in several areas of the brain. Accordingly, coenzyme Q$_{10}$ (CoQ$_{10}$) has been hypothesized as improving brain energy states and cognitive functions in schizophrenia. We conducted a double-blind, placebo-controlled crossover study to examine the correlation...
Fig. 5. Non-localized $^{31}$P MR spectroscopy of calf muscle during activation. A series of spectra sampled from a muscle activation study carried out as described in the text is shown at the top of the figure. Each spectrum is a 15-second $^1$H-decoupled, NOE-enhanced $^{31}$P spectra of the right calf muscle (TR = 1s, 512 points spectral width of $\pm 1,000$ Hz, 15 acquisitions, and a 45$^\circ$ rectangular excitation pulse) collected serially before, during, and after calf activation. Spectral assignments are as described in Figs 1 and 2. The spectra on this figure show representative temporal changes in PCr and NTP during the activation protocol. The graph at the bottom of the figure shows the temporal changes in integration values of PCr (squares), Pi (diamonds), and $\beta$-NTP (triangles) during the activation study. The dotted line corresponds to the PCr recovery after exercise which, under the conditions studied, is proportional to the ATP re-synthesis in the respiratory chain.

between cognitive results and the in vivo signals in brain-localized $^{31}$P MRS, testing the effect of CoQ$_{10}$ on impaired cognitive functions in schizophrenia patients under neuroleptic treatment. Ten schizophrenia patients were selected randomly to receive either placebo or CoQ$_{10}$ during a period of 8 weeks, with the counterpart in the following 8 weeks. At the end of each arm, both cognitive studies and localized brain $^{31}$P MRS were performed. However, only 7 of the 10 patients had the $^{31}$P MRS studies done in both arms. Six matched normal volunteers also were used as controls for the brain $^{31}$P MRS study.

The schizophrenia patients ($n = 10$) studied showed a slightly significant improvement in attention ($p \leq 0.05$) when treated with CoQ$_{10}$ compared to placebo, but other cognitive tests failed to show differences for the group as a whole. When the patients were divided by below-normal (BN) and near-normal (NN) memory scores, a regular practice in schizophrenia research, the patients with BN scores ($n = 4$) showed significant improvement in verbal learning ($p \leq 0.01$), graphmotor speed ($p \leq 0.04$), and attention ($p \leq 0.004$) during CoQ$_{10}$ treatment. Patients with NN scores ($n = 6$) did not show significant changes in any of the cognitive studies during CoQ$_{10}$ treatment.

Table 1 shows the mean brain PCr values obtained by $^{31}$P MRS in the volunteer group ($n = 6$) and in the schizophrenia group ($n = 7$). As shown in the table, while on placebo, the schizophrenia group showed significantly lower brain PCr values than did normal volunteers ($p \leq 0.001$). The mean brain PCr value for schizophrenia patients increased significantly during the CoQ$_{10}$ period ($p \leq 0.03$ against placebo), but this value still was significantly below that of the normal volunteers ($p \leq 0.03$). When the schizophrenia group was divided into BN ($n = 3$) and NN ($n = 4$) memo-
ry score subgroups, there was a significant increase in mean brain PCR value during CoQ_{10} treatment for the BN group (\( p \leq 0.03 \)) and a non-significant increase for the NN group. We also analyzed these data on an individual basis, comparing the levels of PCR in the different voxels in each patient’s brain during placebo or CoQ_{10} treatment. These results are shown in Fig. 6. In this analysis, two of the three patients in the BN subgroup showed statistically significant increases in PCR during CoQ_{10} treatment (\( p \leq 0.001 \) in both) with no changes in the third patient. The NN scores subgroup showed non-significant variations in PCR values.

The correlation of the cognitive studies with our 31P MRS results supports the link between bioenergetic impairment found in schizophrenia may be explained at least in part by a problem in bioenergetics. In addition, the correlation between improvements in cognitive parameters and increased brain PCR levels in schizophrenia patients after CoQ_{10} treatment seen in this work also supports the link between a bioenergetic impairment and the low cognitive scores found in schizophrenia, a possible therapeutic use of CoQ_{10} for patients afflicted with schizophrenia, and the value of 31P-MRS to follow up on these therapeutic changes.

4.3. Non-Hodgkin’s lymphoma studies by in vivo 31P MRS

Identification of metabolic or genetic variables from a tumor that correlate with success or failure of an individual treatment is of great clinical interest. Early knowledge of the likelihood of treatment failure would be of great use, because alternative therapies could then be considered. Currently, the commonly used indicator of treatment prognosis in non-Hodgkin’s lymphomas (NHL) is the international prognostic index (IPI) [199, 200]. This index is based on clinical parameters related to tumor extent and host status and not to specific genetic or metabolic information about individual patients or their tumors.

Several groups have attempted to find tumor-specific genetic markers of disease in NHL to correlate with outcome in an attempt to determine whether such information improves prognostic accuracy beyond the IPI [201–207]. Non-genetic markers of disease in cancer also may be important to correlate with treatment response and outcome. In vivo 31P MR spectroscopy of human tumors has shown a characteristic 31P spectral pattern: the tumor levels of phospholipid-related phosphomonoesters, particularly Etn-P and Cho-P, have been known to be elevated in human tumors compared to most normal tissues since the first in vivo 31P MRS observation of a rhabdomyosarcoma in the hand [208]. Several general reviews have been published on the subject [167–172], on specific tumor histopathologies [69, 173,190], and on specific treatments [191–195].

We have studied patients with NHL using localized 31P MRS before treatment to demonstrate whether pretreatment levels of Etn-P plus Cho-P correlate with clinical parameters related to treatment response and outcome [19]. Three-dimensional, H-decoupled, NOE-enhanced CSI of 31P spectra was acquired from each patient’s major tumor area in a 1.5 T clinical imager [10] using custom-built, dual-tuned (31P/H) RF antennas, as described elsewhere [18,209]. The tumor-containing voxels in the CSI dataset were selected using the IR images as reference. Voxel-shifting [10] was used to reduce the amount of contamination in the tumor. An example of the procedure is shown in Fig. 4. Tumor voxels were extracted from the CSI datasets and summed when necessary to produce one spectrum per patient. The sum of the phosphate signals of the phospholipid-related phosphomonoesters, Etn-P and Cho-P (Etn-P + Cho-P), and the P_{i} signal of NTP in the resulting spectra were integrated manually to obtain the tumor [Etn-P + Cho-P]/NTP ratio. Each patient was studied in the 30 days prior to instituting a new treatment regimen. Treatments were at the discretion of the clinician. The response to treatment was determined following the World Health Organization criteria using bi-dimensional radiological measurements in serial CT scans.

We evaluated the pretreatment [Etn-P + Cho-P]/NTP levels in tumors for their ability to predict the patient’s long-term (6-month) treatment response. Clinically, patients were classified into complete (CR) and not-complete responders (NCR). The difference between CR and NCR was significant (CR: 1.45 ± 0.15, mean ± standard error, \( n = 10 \) vs. NCR: 2.28 ± 0.15, \( n = 17 \), \( p \leq 0.001 \)). A Fisher test for [Etn-P + Cho-P]/NTP also was significant (sensitivity = 70%, specificity = 71%, positive predictive value [PPV] of 58%, and overall accuracy = 70%, \( p \leq 0.04 \)). When the patients were divided into risk groups depending on their IPI, the [Etn-P + Cho-P]/NTP ratio was significant for the low-risk (CR: 1.60 ± 0.21, mean ± standard error, \( n = 5 \) vs. NCR: 2.46 ± 0.25, \( n = 6 \), \( p \leq 0.03 \)) and low-intermediate risk populations (CR: 1.30 ± 0.20, \( n = 5 \) vs. NCR: 2.20 ± 0.18, \( n = 6 \), \( p \leq 0.01 \)). Using the average medians for CR and NCR of the low-risk group (1.99) and of the low-intermediate group (1.67), variable IPI-dependent cutoffs for [Etn-P + Cho-
Table 1: Brain Phosphocreatine Measured by $^{31}$P MR Spectroscopy in Schizophrenia

| Group                        | Brain PCr (arbitrary units) | Significance                  | Placebo vs. CoQ$_{10}$ |
|------------------------------|----------------------------|-------------------------------|-------------------------|
| Normal group ($n=6$)         | 4.20 ± 0.16                | Normal vs. placebo vs. CoQ$_{10}$ | $p \leq 0.001$          |
| Schizophrenia whole group ($n=7$) | 3.19 ± 0.18                | CoQ$_{10}$ vs. placebo       | $p \leq 0.03$           |
| Below normal (BN) score group ($n=3$) | 2.99 ± 0.26                | CoQ$_{10}$ vs. placebo       | $p \leq 0.03$           |
| Near normal (NN) score group ($n=4$) | 3.39 ± 0.22                | N.S. b] $p \leq 0.03$         |

aMean ± standard error.
bN.S., not significant.

Fig. 6. Treatment response to coenzyme Q$_{10}$ in schizophrenia. The brain PCr obtained using in vivo $^{31}$P MR spectroscopy in each patient during placebo and Coenzyme Q$_{10}$ (CoQ$_{10}$) as the mean ± standard error of the brain voxels selected are shown. Patients were segregated according to those with below-normal memory scores and those with near-normal memory scores. The two asterisks denote those brain PCr values during CoQ$_{10}$ treatment that are significantly higher than those during placebo ($p \leq 0.001$ in both). As shown in the figure, the two significant PCr increases during CoQ$_{10}$ treatment happened in patients with below-normal memory scores who were the patients with significant responses on the cognitive studies (see text).

PJ/NTP were recognized that proved to be better predictors of long-term treatment response in a Fisher test. These variable cutoffs are such that, for patients at higher risk, with larger IPI values, a lower [Etn-P + Cho-P]/NTP is needed to predict positive outcome. Using these cutoffs, the Fisher test significance of the whole group increases to $p \leq 0.0002$, and its sensitivity and specificity improve to 80% and 94%, respectively.

These results suggest that we have identified an independent metabolic marker for disease. As far as we are aware, this is the first observation of an association between in vivo metabolite levels in tumors and clinical outcome, although genetic variations have been documented [207]. In any case, it is likely that, the more closely a variable can predict treatment outcome, the more likely the variable is causally connected to the underlying mechanism(s) responsible for the response to the treatment. This is true particularly when the variable is a specific intra-tumor metabolite or group of metabolites, as is the case in the present work. Thus, our results suggest that part of the metabolic pathway involved in a tumor being triggered into apoptosis is associated with the phospholipid precursors Etn-P and Cho-P. Although a detailed analysis of the causal possibilities is not feasible at present, the use of [Etn-P + Cho-P]/NTP levels in a clinical setting to guide therapy in individual patients appears to be possible once they have been confirmed in a larger patient cohort. In this work, we concluded that the pretreatment levels of Etn-P and Cho-P in NHL in vivo correlate with individual responses to treatment and time to treatment failure. This suggests that an independent factor in the variability of NHL tumor responses to therapy was identified and that it can be used as a predictor of treatment response.
4.4. Chronic lymphocytic leukemia studies by $^{31}$P MRS [5]

Chronic lymphocytic leukemia (CLL) is a unique disease because it offers a link between in vivo and in vitro studies. CLL cells can be isolated easily from fresh blood so that cellular conditions are determined by the patient’s clinical and treatment status, and cell extracts suitable for $^{31}$P MRS analysis at high fields can be prepared immediately after isolation. Thus, the advantages associated with $^{31}$P MRS studies of cultured cell lines can be obtained in a human tumor through ex vivo studies of CLL cell extracts. Furthermore, CLL and NHL both are B-lymphoid malignancies. The similar cellular origin of both tumors offers the possibility of comparing the ex vivo CLL results with our ongoing studies of NHL in vivo. In this preliminary CLL study, levels of phospholipid-related metabolites (Cho-P, Etn-P, Gro-P-Chol, Gro-P-Etn) of extracts from CLL leukemia lymphocytes and normal human lymphocytes were quantified using phosphorus MRS. Figure 1 shows an example of the comparison between normal (lower spectrum) and CLL lymphocytes (upper spectrum) in this study. The CLL cells vs. normal lymphocytes showed significant increases of Etn-P ($8.11 \pm 2.10 \text{ mean } \pm \text{ standard error, } \mu \text{mol/g wet weight, } n = 12 \text{ vs. } 3.63 \pm 1.10, n = 3; \mu \text{mol/g wet weight, } n = 12 \text{ vs. } 0.36 \pm 0.09, n = 3; \mu \text{ mol/g wet weight, } n = 10 \text{ vs. } 0.17 \pm 0.05, n = 3; p \leq 0.003$), Cho-P ($2.10 \pm 0.37, n = 12 \text{ vs. } 0.11 \pm 0.05, n = 3; p \leq 0.004$), and Gro-P-Chol ($0.26 \pm 0.03, n = 10 \text{ vs. } 0.11 \pm 0.05, n = 3; p \leq 0.004$), and Gro-P-Etn ($0.33 \pm 0.03, n = 10 \text{ vs. } 0.17 \pm 0.05, n = 3; p \leq 0.003$). Further, the phospholipid precursor ethanolamine was studied in blood and was found to be reduced significantly in CLL patients ($4.6 \pm 1.6 \mu \text{M, } n = 25$) compared to normal volunteers ($7.7 \pm 2.5, n = 12; p \leq 0.001$). Increased intermediates with depletion of precursors suggest the presence of sustained phospholipid metabolism activation in CLL.

With respect to their $^{31}$P MR spectra, CLL lymphocytes are characterized by elevated phosphomonoesters (Etn-P and Cho-P) and mobile phosphoesters (Gro-P-Chol and Gro-P-Etn) in comparison to normal lymphocytes. This pattern of phospholipid-related metabolites cannot be accounted for by elevated serum ethanolamine. Although steady-state concentrations of Etn-P, Cho-P, Gro-P-Chol, Gro-P-Etn, CDP-ethanolamine, and CDP-choline are determined by a complex interaction of phospholipid synthetic pathways (from ethanolamine and choline), catabolic pathways (phospholipases A2, C, and D), and reentry of intermediates into synthetic pathways, probable causes for observed patterns can be inferred. For example, an increase of Gro-P-Etn and Gro-P-Chol upon mitogen stimulation is compatible with a concurrent activation of phospholipase A2 known to occur upon stimulation [210,211]. Similar to mitogenically stimulated normal human lymphocytes, our study shows that CLL cells have higher Gro-P-Etn, CDP-ethanolamine, and Gro-P-Chol than controls, which suggests an activation of phospholipase A2 in CLL lymphocytes. Alternatively, activation of the kinases of ethanolamine and choline could be responsible for the elevated Etn-P and Cho-P seen in CLL cells if this activation is such that it compensates for lower substrate levels (lower serum ethanolamine in CLL patients than in controls).

A high concentration of Etn-P has been observed with stimulation of phospholipases C or D in a variety of cancer cell lines [212–214]. Gillham and Brindle suggested a tentative connection between the elevated levels of plasma fatty acids seen in cancer patients and the increased phosphomonoester signals observed in $^{31}$P MR spectra of tumors [214]. Dixon and Tian in a murine lymphoma infiltrating the liver support this hypothesis of phospholipase C or D activation [215, 216]. In this tumor model, high Etn-P levels relative to Cho-P, and a lack of elevated Gro-P-Etn or Gro-P-Chol, were found. Etn-P synthesis was increased relative to that of Cho-P, and Etn-P synthesis via the CDP-ethanolamine synthetic pathway was decreased compared to the normal liver [215]. From these results, the authors suggested that their lymphoma model has high levels of Etn-P due to phospholipid breakdown via phospholipase C or D in combination with rate-limiting activity of CTP:EtP cytidylyltransferase [215]. Our results differ from these with regard to the elevation of Gro-P-Etn and Gro-P-Chol and a greater increase of Cho-P compared to Etn-P in CLL than in normal lymphocytes. Hence, we believe that sustained activation of phospholipase C or D is not the mechanism for our findings.

Although our data support phospholipase A2 or kinases activation in CLL, a formal mechanism by which these alterations are associated with malignancy still needs to be elucidated. However, regardless of the mechanism, there is a clear association between the malignant phenotype and elevated levels of Etn-P and Cho-P in both NHL and CLL cells. Moreover, several investigators have observed elevated levels of these compounds in other tumors as well [167,174,176,183, 215–231]. Since there is a reasonable possibility that this elevation is due to pathways that involve the control of fundamental cellular processes (such as apoptosis [232]), further investigation is warranted.
5. Summary

As illustrated above, there are many diseases in which the observation of in vivo metabolism can provide helpful clinical information, both for diagnosis and treatment. As our abilities to measure metabolites improve, these unique markers of disease, which clearly are related to the disease status in an individual patient, will become more useful in diagnosing and treating each unique patient.

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