NMR of $^{31}$P nuclear spin singlet states in organic diphosphates

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Article info
Article history:
Received 16 September 2021
Revised 27 October 2021
Accepted 28 October 2021
Available online 30 October 2021

Keywords:
Phosphorus NMR
Nuclear spin singlet state
Quantum filter

Abstract
$^{31}$P NMR and MRI are commonly used to study organophosphates that are central to cellular energy metabolism. In some molecules of interest, such as adenosine diphosphate (ADP) and nicotinamide adenine dinucleotide (NAD), pairs of coupled $^{31}$P nuclei in the diphosphate moiety should enable the creation of nuclear spin singlet states, which may be long-lived and can be selectively detected via quantum filters. Here, we show that $^{31}$P singlet states can be created on ADP and NAD, but their lifetimes are shorter than $T_1$ and are strongly sensitive to pH. Nevertheless, the singlet states were used with a quantum filter to successfully isolate the $^{31}$P NMR spectra of those molecules from the adenosine triphosphate (ATP) background signal.

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1. Introduction

Organophosphates play a critical role in biology as energy carriers. Adenosine triphosphate (ATP) is the main currency of energy for the cell and is central to cellular metabolism [1,2]. Energy is released and used to drive metabolic processes by breaking the phosphate-phosphate bond, producing adenosine diphosphate (ADP) and inorganic phosphate. At the same time, new ATP is created from ADP via glycolysis, the citric acid cycle, and the electron transport chain, all driven by the breakdown of sugars, fatty acids, and proteins. ATP concentration in vivo is typically several times that of ADP, and their ratio provides information about cellular energy status and mitochondrial function [3,4]. Another organophosphate, nicotinamide adenine dinucleotide (NAD), is a cofactor for many metabolic pathways in which it is converted between its oxidized form NAD$^+$ and its reduced form NADH. The ratio between NAD$^+$ and NADH reflects the oxidative state of the cell [5], and the total NAD concentration can change as a result of aging and neurodegenerative disease [6].

In vivo $^{31}$P NMR spectroscopy and imaging are commonly used to quantify the relative concentrations of ATP, ADP, and NAD non-invasively. Applications include the assessment of traumatic brain injury, investigation of aging, and diagnosis of musculoskeletal diseases [7–11]. However, detection and quantitation of these molecules can be difficult due to spectral overlap, particularly in the −11 to −12 ppm spectral region where lines from ATP, ADP, and NAD all occur. While some lines can easily be resolved with high-field, high-resolution NMR spectroscopy [10,12], lower field strengths used for human MRI make this challenging; thus ADP concentration is often calculated indirectly based on measurements of phosphocreatine [13,14].

To help isolate such overlapping spectral lines, quantum filters and spectral editing pulse sequences can be used to eliminate background signals. While widely used in $^1$H NMR spectroscopy, there are few examples for $^{31}$P. Jayasunder et al. used a multi-quantum filter to remove interfering 2,3-diphosphoglycerate signal from measurements of inorganic phosphate [15], and Tsai et al. used double-quantum filtered HETCOR to study dentin with solid-state NMR [16]. Brindley et al. used spectral editing to suppress phosphomono- and phosphodiester signals [17], Berkowitz and Ben-Bashat et al. explored $^{31}$P-$^1$H quantum coherences for the measurement of ADP in the presence of ATP in vitro [18,19]. A drawback is that ADP appears as an anti-phase signal, which

https://doi.org/10.1016/j.jmr.2021.107101
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can lose intensity if T2* is short. More recently, Ren et al. used spectral editing to improve the NAD⁺ spectrum near an interfering ATP peak. This involved subtracting an inversion spectrum from a reference spectrum, with the inversion time selected to null either the ATP or NAD⁺ signal [20]. This decreased the ATP signal but did not fully remove it or other underlying lines such as uridine diphosphoglucone.

Recently, a new class of quantum filters and spectral editing techniques have been developed based on the long-lived nuclear spin singlet state [21–26]. The singlet state can be created in pairs of coupled nuclear spins when the spins are in the magnetically equivalent or near-equivalent conditions. This condition can be satisfied when the chemical shift is small relative to scalar coupling, such that \( \Delta \nu \ll J \), or through the application of decoupling via CW spin-locking or a pulse train. The resulting singlet states can potentially have long lifetimes far beyond T₁ and can also be isolated from other signals via appropriate pulse sequences. The structure of ADP and NAD, each with a pair of phosphate groups on which \(^{31}\)P singlet order can be prepared, appear to lend themselves to such a detection strategy (Fig. 1).

To test this method, we created singlet states in the \(^{31}\)P pairs of ADP and NAD⁺. We measured the longitudinal relaxation time (T₁) and singlet relaxation time (T₂) under both neutral and basic conditions. We then used a singlet quantum filter via the SUCCESS (pKa = 0.9 and 2.9–4.4) and the NAD⁺ phosphate groups (pKa = 6.8–7.2) remains partially protonated, likely driving \(^{31}\)P T₁ relaxation [28–30]. Given pH = 7 and pKₐ = 6.8, 39% of \(^{31}\)P groups are protonated at a given time. To measure singlet lifetime with these protons fully removed, a basic 50 mM ADP solution was prepared in 0.5 M sodium hydroxide solution, giving a pH > 13 and dissociating over 99.99% of the protons from the \( \alpha \)-ADP phosphate group. This increased T₁ significantly and T₂ by a factor of eight (Table 1). However, T₂/T₁ was still on the order of unity, indicating that other relaxation processes are also significant, likely from anti-correlated chemical shift anisotropy [31].

Next, we tested the \(^{31}\)P SUCCESS sequence on a mixture containing 30 mM ATP, 3 mM ADP, and 3 mM NAD⁺ in a pH 7.0 phosphate buffer. Fig. 2a shows a spectrum of the mixture measured with a 90-FID sequence. Both ATP and ADP exhibit splitting patterns indicating weakly coupled \(^{31}\)P groups (additional ATP peaks at \( \pm 20 \) ppm are not shown), and the ADP peaks are on the shoulders of the ATP peaks. One of the NAD⁺ peaks lies directly beneath an ATP peak while the other is resolved. Fig. 2b shows the results after using the SUCCESS sequence to target ADP. Using line fitting for quantitation, we find that 35% of the ADP signal is retained, compared with only 1% of the ATP signal, resulting in a factor of 35-fold enhancement for ADP signal contrast. Fig. 2c shows the results after targeting NAD⁺ instead. For NAD⁺, 43% of the signal is retained, compared with only 1.7% of the ATP signal, resulting in a factor of 25-fold enhancement for NAD⁺ signal contrast.

3. Discussion and conclusion

Our results show that it is possible to create, and detect with NMR, nuclear spin singlet states on pairs of \(^{31}\)P nuclei in common biomolecules. However, these singlet states relax significantly faster than T₁ under typical conditions, similar to the results reported by Korenchan et al. in tetrabenzyl pyrophosphate [31]. Measurements under basic conditions show that nearby protons are partly to blame, but their removal does not necessarily lead to a long-lived state. The strongest relaxation mechanism is likely chemical-shift anisotropy (CSA), which is absent in \(^{1}\)H systems but is often strong in \(^{13}\)C, \(^{15}\)N, and \(^{31}\)P systems [27,31,32]. Long-lived singlets can still be created in \(^{13}\)C and \(^{15}\)N pairs when molecules are small and highly symmetric. While it is possible that \(^{31}\)P systems with the appropriate criteria also exist, rotation about the single bonds connecting phosphate groups makes it unlikely for this class of molecules.

Despite the short singlet lifetime, the SUCCESS sequence performed well at removing ATP background NMR signal and isolating the spectra of ADP and NAD⁺. This method is potentially useful for in vivo MR spectroscopy at lower 1.5 T and 3 T magnetic field strengths typical of human and animal imaging, particularly when the broader lines in vivo lead to stronger spectral overlap. As a special case of zero-quantum filters, singlet filtration techniques such as SUCCESS and others are not significantly affected by B₀ inhomogeneity. While we based our technique on the Sarkar three-pulse sequence for singlet preparation and readout [33], other sequences such as M2S, SLIC, and APSOC might be more appropriate at lower

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**Fig. 1.** Structures of ATP, ADP, and NAD⁺. Singlet order can be prepared on the \(^{31}\)P pairs of ADP and NAD⁺ (circled). The triphosphate group of ATP does not support a singlet, because as a spin-0 state, the singlet must be composed of an even number of coupled spin-1/2 nuclei.
magnetic field strengths, especially for NAD+, whose spins enter the nearly-equivalent regime at 1.5 T [34–36]. In that regime, it might be possible to also eliminate spin-locking for singlet preservation, so that specific absorption rate (SAR) can be minimized. The SUCCESS sequence can also be improved by using gradients as part of the quantum filter for isolating singlet state, but these were not available on our spectrometer. The use of phase cycling alone can lead to imperfect signal cancellation if there are small frequency shifts during acquisition, which might be why some ATP signal remained in both cases.

Besides SAR, another drawback for human imaging is the two to three-fold signal loss that results from relaxation and from quantum mechanical limitations of using a singlet filter. To attain the same signal-to-noise ratio as a conventional scan would require either four to nine times more averages or the use of voxels with two to three times the volume. However, current spectral editing techniques for NAD+ also have a time penalty, as they require two scans, one reference and one with an inversion. While relaxation times are shorter in vivo for NAD+ we do not expect longitudinal or singlet relaxation to be significantly worse. Given that T1 of up to 2 s has been measured in the brain at 7 T, T2 ~ 1 s is expected, which is still sufficiently long compared to the sequence time. The 31P T2 of NAD+ has not been determined in vivo, but literature values for T2 of other metabolites are in the 10 to 250 ms range [37,38]. This would lead to losses during preparation and readout for all forms of quantum filters. However, some of the short T2 measurements are actually a reflection of the homonuclear couplings rather than the true T2, so further investigation is needed [39]. Application to ADP may be more challenging, as a T1 in vivo of only 870 ms has been measured at 9.4 T, but the relaxation times should be longer at lower field where there is less CSA relaxation.

In conclusion, we found that 31P nuclear spin singlet states in ADP and NAD+ have relatively short lifetimes, but such states can nevertheless be used to isolate the NMR spectral signature of these molecules. This result suggests there is utility in exploring singlet states even in systems where short lifetimes might be expected due to strong chemical shift anisotropy or out-of-pair couplings. In particular, when spin relaxation properties are sensitive to parameters such as pH, the T2/T1 ratio might also provide useful information about the chemical environment.

4. Methods

NMR spectroscopy was performed on a Bruker DMX 200 MHz spectrometer with a 1H/X dual channel probe (81 MHz for 31P). Reagents were purchased from Sigma Aldrich (St. Louis, MO, USA). ADP, ATP, and NAD+ were in the form of sodium salts. For T1 and T2 measurements, solutions were 50 mM of either ADP or NAD+ in 69 mM pH 7.0 phosphate buffer (neutral conditions) or
in 500 mM NaOH giving a pH > 13 (basic conditions). For SUCCESS measurements, a solution of 30 mM ATP, 3 mM ADP, and 3 mM NAD$^+$ was prepared in 69 mM pH 7.0 phosphate buffer. Conventional $^{31}$P spectra were acquired with a 90- FID sequence, and T1 was measured with a standard inversion recovery sequence. Chemical shifts were referenced to inorganic phosphate. TS was measured with a standard inversion recovery sequence. NAD$^+$ was prepared in 69 mM pH 7.0 phosphate buffer. Conventional $^{31}$P spectra were acquired with a 90- FID sequence, and T1 was measured with a standard inversion recovery sequence. Chemical shifts were referenced to inorganic phosphate. T2 was measured with the SUCCESS sequence (below) using a series of spin-lock nutation times ($\tau_C$). The $^{31}$P spin-lock nutation frequency was 615 Hz. In all cases 1 H decoupling was applied during acquisition (CW, 150 Hz nutation frequency).

The SUCCESS technique has been described previously and is shown in Fig. 3 [21,26]. It is a Sarkar sequence combined with a filter for the singlet state (a type of zero-rank filter) [33]. In this pulse sequence, a singlet precursor state is prepared on a selected spin throughout. The 24-phase cycle is then returned to transverse magnetization for readout. Pulse phases for $\phi_1$, $\phi_2$, $\phi_3$, and $\phi_\text{acq}$ of the polyhedral phase cycle are given in Table 2. $\phi_2$ was 0 throughout. The 24-phase cycle was chosen to ensure filtering of coherences up to rank $k = 3$, the maximum possible for three spin-1/2 nuclei. Delay times for ADP were $\tau_1 = 12.02 \text{ ms}$, $\tau_2 = 17.16 \text{ ms}$, and $\tau_3 = 2.57 \text{ ms}$. The center frequency was $-10.74 \text{ ppm}$. For NAD$,^+$ delays were $\tau_1 = 15.5 \text{ ms}$, $\tau_2 = 16 \text{ ms}$, and $\tau_3 = 12.5 \text{ ms}$, and the center frequency was $-11.675 \text{ ppm}$. For both molecules, the spin-locking time was $\tau_4 = 100 \text{ ms}$ and the spin-lock nutation frequency was $\nu_\text{acq} = 615 \text{ Hz}$. 8192 averages were taken with a delay of 14 s between measurements. A large number of averages were used for the SUCCESS demonstration so that residual ATP signal could be quantified. Metabolites were quantified in the reference and SUCCESS spectra by fitting each phase spectrum with Lorentzian lines. SUCCESS was also performed on the individual 50 mM ADP and NAD$^+$ solutions to quantify signal losses without interference from ATP.

### Declaration of Competing Interest

The authors share royalty interest in U.S. Patent 20150042331 covering the SUCCESS method.

### Acknowledgments

We acknowledge support from the NSF, Army, DARPA, and ARO MURI.

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