



Account/Revue

Designing and building a low-cost portable FT-NMR spectrometer in 2019: A modern challenge



Concevoir et construire un spectromètre FT-RMN portable à bas coût en 2019 : un défi moderne

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ABSTRACT

High-field Fourier-Transform Nuclear Magnetic Resonance (FT-NMR) spectroscopy is a high performance spectroscopic technique that is essential in many analytical fields. The non-destructive nature of NMR makes it a preferred means of analyzing chemical and biological environments. Compact benchtop NMR spectrometers are low-cost alternatives to conventional high-field and high-resolution spectrometers. A research laboratory may want to develop its own compact FT-NMR spectrometer ("home-built benchtop NMR") with a very low financial cost (~10 k€). But why? First of all, to use it punctually as an additional channel (nucleus X) to a high-resolution spectrometer, and also to be able to couple it with complementary physics instruments such as an optical microscope to study spin diffusion in semiconductors, for instance. In addition, a "home-built" NMR spectrometer can be used with a low-field permanent magnet for the quantification of species that does not necessarily require high-resolution, avoiding the need for weekly and expensive cryogenic services. Outside the research laboratory, this portable NMR can be used for the *in situ* analysis of outdoor natural environments. Finally, this compact spectrometer is naturally dedicated to the teaching of NMR technique and is open to the study of the basic electronic functions that constitute an NMR spectrometer. The main question then arises: how to build a robust "Home-Built" NMR? In this article, we describe the realization of an NMR instrument based on electronic components and boards (LNA, ADC, FPGA, ARM, DDS...) easily commercially available, allowing one to obtain a benchtop NMR instrument presenting both a high acquisition dynamics and a good signal-to-noise ratio. © 2019 Académie des sciences. Published by Elsevier Masson SAS. This is an open access article under the CC BY-NC-ND license (<http://creativecommons.org/licenses/by-nc-nd/4.0/>).

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R É S U M É

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La RMN à transformée de Fourier (RMN FT) et à hauts champs est une technique spectroscopique très performante et incontournable dans de nombreux domaines analytiques. Le caractère non destructif de la RMN en fait un moyen privilégié pour analyser les milieux en chimie et en biologie. Les spectromètres RMN à TF compacts, dits aussi de paillasse, sont des alternatives, à bas coût, aux spectromètres classiques hauts champs et haute résolution. Un laboratoire de recherche peut vouloir développer son propre spectromètre RMN compact (*home-built benchtop NMR*) à un coût très réduit (~10 k€). Mais pourquoi ? Tout d'abord, pour l'utiliser de façon ponctuelle comme canal supplémentaire (noyau X) à un spectromètre haute résolution, mais aussi pour pouvoir le coupler à d'autres instruments de physique, comme par exemple à un microscope optique, pour l'étude de la diffusion de spin dans les semi-conducteurs. Par ailleurs, une RMN *home-built* peut être utilisée avec un aimant permanent bas champ pour la quantification d'espèces qui ne nécessitent pas nécessairement une haute résolution, évitant le recours à des services cryogéniques hebdomadaires et coûteux. Hors laboratoires de recherche, cette RMN portable peut être utilisée pour l'analyse *in situ* de milieux naturels en extérieur. Enfin, ce spectromètre compact est naturellement dédié à l'enseignement de la spectroscopie RMN, en étant ouvert à l'étude des fonctions électroniques de base constituant l'instrument. La question principale qui se pose alors est : comment construire aujourd'hui un spectromètre « maison » ? Nous décrivons dans cet article la réalisation d'un spectromètre RMN basé sur des composants et des cartes électroniques (LNA, ADC, FPGA, ARM, DDS...) accessibles commercialement et permettant d'obtenir une RMN de paillasse présentant à la fois une grande dynamique d'acquisition et de bons rapports signal/bruit.

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1. Introduction

Conceived and explored in 1945 in physics laboratories [1–5], Nuclear Magnetic Resonance (NMR) spectroscopy is arguably nowadays one of the most essential tools for analyzing the structure of mineral, organic, or biological molecular systems. This fantastic achievement was possible with the development of Fourier-Transform NMR (FT-NMR) proposed by R.R. Ernst [6], that in turn opened the door to first NMR experiments based on sophisticated multipulse sequences [7,8]. Concomitantly, the application of NMR principles/concepts has also offered fundamental opportunities for medical imaging for robust diagnostic purposes. Behind the vast practical aspects of analysis of matter, NMR basics underlie varied and sophisticated theoretical concepts in physics (data sampling), mathematics (Fourier Transform (FT) principle), electronics (resonating circuit), or quantum mechanics (Hamiltonian, operator density...), as nicely described in many theoretical/practical books that are references for any NMR spectroscopists or experienced users [9–16].

As a brief reminder, Table 1 provides a chronological summary of the main technological/methodological advances from the 1950s to the present days, in conjunction with the magnetic field strength for a continuous enhanced spectral resolution and sensitivity.

Unfortunately, the increasing "race of Tesla" has a significant financial cost (purchase) and periodic maintenance (cryofluids), and hence the need for cryofluid-free, FT-NMR spectrometers is becoming acute again. In addition, nowadays, the availability of rather small permanent magnets (soft iron) with a good magnetic homogeneity (up

to 10^{-4}) and stability and the significant improvement in electronic performance make it possible to offer very compact, versatile benchtop FT-NMR spectrometers at a reduced cost, making again this new generation of instrument very attractive for various applications in chemistry [17].

In this article, we propose to revisit the development and assembly of a very low-cost (~10 Keuros), portable (benchtop) FT-NMR spectrometer by integrating magnetic and electronic classical elements. For this, the essential units required will be described, detailing their respective role in the excitation/detection process of signal, for instance. In a memorial approach, in particular, toward chemists who have generally forgotten the NMR principles, we will start from some (important) basic physical principles of NMR to organize and choose the key block elements necessary for the realization of this "artisanal" or home-made (as anglicists would say) spectrometer. Interestingly, such a project is possible with laboratory instrumentation (function generator, oscilloscope...) accessible at a lower cost. All requested, specific electronic components/devices with their current nomenclature are listed in Appendix and are commercially available.

Proposing an inexpensive, home-made FT-NMR instrument is above all an exciting task of engineering integrated into an educational project, for instance. It obviously cannot be considered as a competition with the new generation of benchtop FT-NMR spectrometers available in the market today from different manufacturers (Magritek, Oxford, ThermoFisher, Nanalysis...), which integrates many optional modules (locking, sample spinning...) [17]. Moreover, although the overall cost (material/electronic

Table 1

Historical overview of main instrumental/methodological achievements.

NMR methodology	CW-NMR			FT-NMR									
				1D		2D homonuclear		2D heteronuclear gradients		3D, 4D, TROSY ...		FAST NMR	
				Supraconductor magnet			Ultra shielded magnet, cryogenic probe						
Size of analyzable molecules (kDa)	< 10						25	35		50	100		
B ₀ (T)	0.70	1.41	2.11	4.22	6.34	9.39	11.74	14.09	17.61	18.78	21.13	22.32	23.48
¹ H Larmor (MHz)	30	60	90	180	270	400	500	600	750	800	900	950	1000
Microprocessors Intel family name				4004		8080			80490		Pentium 4		Intel Core i7
Number of transistors in microprocessors (Intel family)				2.3 × 10 ³		6 × 10 ³			1.2 × 10 ⁶	9.5 × 10 ⁶	42 × 10 ⁶	291 × 10 ⁶	1.1 × 10 ⁹
Years				1965		1975		1985		1995		2005	
	1950	1960			1970		1980		1990		2000		2010

supplies) of the instrument seems attractive at first sight, it does not include the actual human costs (total cost), in particular in view of its pedagogical dimensions in Academic Education. This home-built NMR spectrometer can be seen as an “alive” support instrument for teaching NMR. Finally, this type of mobile NMR apparatus is well suited for scientific research by coupling NMR and another instrument (such as a microscope).

To have an idea, the time needed by a single engineer to build a one-channel mobile NMR may be estimated as two months, including the tests (and assuming that all electronic hardware components have been already purchased).

2. Contribution of modern electronics

Electronics has developed considerably in recent decades, making it possible to build high-performance systems with high functional integration. To illustrate the evolution of the power of electronic component, we have included the number of transistors integrated on the same chip over the years in Table 1. For example in 1965, a microprocessor integrated nearly 2000 transistors on the semiconductor chip. The density of the components has increased by a factor of 10⁶ in 45 years (1965–2010), thereby reducing the size of the electronic circuits. Currently, a simple integrated circuit is capable of performing signal acquisition, processing, and control functions. The modern smartphone is a blatant illustration of the high-tech concentrate contained in such a small volume. As the density of the components increases, the

operating frequency of the hardware circuits has also increased considerably. Hardware can now operate at frequencies of several gigahertz compared to a few hundred kilohertz in the 1970s. These high working frequencies allow complex processes to be carried out with extremely short times.

Another huge leap in the field of electronic equipment is the memory capacity of components. It is now possible to store and manipulate memory spaces of more than 64 gigabytes. Data storage is no longer a real limitation and it is possible not only to acquire large amounts of data, but also to store the ever more sophisticated programs to control and process these data.

The frequency synthesis is now digitally based on DDS (Direct Digital Synthesis) cards over a very wide range of frequencies with very high accuracy. In Table A1 (nomenclature of components) presented in Appendix, we list the references of a synthesis unit capable of generating frequencies from a few kHz to more than 300 MHz. Analogically, the sizes and costs of low-noise preamplifier components, mixers, and other hardware have been reduced. For example, the low-noise preamplifiers in the acquisition chain can be made with a few circuits for a hundred euros.

Finally, an important point to keep in mind is the reduction of the energy consumption of modern components, allowing complete battery-powered systems to be powered, making them autonomous and therefore portable. This aspect was unthinkable in the 1980s. In summary, the performance of electronic components has increased while sizes and costs have been reduced, so the

realization of complex units such as a small NMR spectrometer is possible in research laboratories.

3. Some specificities of a benchtop FT-NMR spectrometer

Classical or benchtop/mobile FT-NMR spectrometer is a complex instrument, but it can be easily divided into distinct and clearly identified functional blocks [18]. In practice, this instrumentation involves many domains of electronics such as radiofrequency (RF), audio chain, power electronics, low-noise preamplification, and transmission lines. In this first decade of the 21st century, FT-NMR spectrometers are fully digital and multi-nucleus. Gradually, they are connected by internet, and therefore they can be remotely controlled and (unfortunately for experimenters) fully automated. The two operating modes of FT-NMR spectrometers (excitation and detection steps) are performed by fast microprocessors and executed by suitable softwares.

As a matter of principle, the channels (^1H , ^{13}C , X...) of spectrometers are similar but independent of each other, all of them are synchronized by a master clock; a dedicated pulse sequencer controlling the complex multi-channel pulse sequences of the spectrometer. This sequencer takes into account the different transmission delays and hardware dead times (such as the pre-scan delay) to produce a reliable and precise pulse timing controlled by the software. The duration of the (hard or soft) electromagnetic pulses (μs to ms) and the delays between events (μs to s) are those actually achieved by the electronics, without the need to add small compensation delays in the pulse programs. These specificities are made possible, thanks to the very high speed and stability of modern electronics.

The assembly of mobile NMR consists of three distinct parts: the magnet, probe, and data acquisition and processing spectrometer. The magnet and probe depend on the chosen application: low field or high field. We discuss these aspects later in this article.

The electronics of our mobile NMR spectrometer is designed to acquire very low value signals (less than μV) over a high dynamic range (the noise factor of the detection chain is less than 2 dB), and the sampling is adapted to the required dynamics (8, 12, 16, or 24 bits). The same applies to frequency synthesis and quadratic demodulation, which can be adapted to a very wide range of frequencies. Note that all the electronic hardware components (RF power amplifier, mixer, preamplifier...) have been chosen to work in a wide frequency range (from 1 to 400 MHz). This allows the portable FT-NMR spectrometer to work at low field (with a static permanent magnet) or at high frequency (with a superconducting magnet).

4. Sensitivity and resolution

The homogeneity and strength of the magnetic field of a permanent magnet define the sensitivity and spectral resolution of any NMR experiment. The temporal magnetic stability of a magnet obviously limits the duration of the measurements. Homogeneity and magnetic stability must allow a coherent accumulation of signals in phase.

NMR is often defined as rather insensitive and high-field magnets are needed to achieve robust sensitivity and resolution to analyze complex molecules. In high-resolution NMR, chemical shifts must be measured with a resolution of about 0.01 ppm to solve the J couplings of a few Hertz. Magnetic fields are of the order of several Tesla and are generated by superconducting magnets cooled by helium and liquid nitrogen. A system of compensation for inhomogeneity (shims) is essential to obtain the required resolution.

Permanent magnets with low field ($<0.94\text{ T}$ or 40 MHz for ^1H) are called low resolutions due to their low sensitivity and homogeneity. In an inhomogeneous field, the magnetic strength varies in relation of the spatial position in the magnet. Such effect leads to significant broadening or/and overlapping of resonance peaks on spectra, thus preventing exploitable spectral resolution. Nevertheless, the measurement of distribution of relaxation times or diffusion coefficients of molecules is accessible at low fields. Quantitative analyses are also possible. The characterization of materials (liquid, powder) can be carried out at low resolution with a home-made spectrometer that is much less cumbersome, less expensive in cryogenic maintenance, and accessible because it is close to the synthetic bench of molecules. The latter reduces the cost and workflow of any research activity. The mobile NMR spectrometer we are describing is part of this approach: it is compact and can be used with a low-field magnet at Larmor frequencies ranging from 1 to 15 MHz in the set presented.

Although intended for low-field, low-frequency applications, as mentioned above, the design and electronic components of our mobile NMR spectrometer can operate at frequencies up to 300 MHz (see the [Table in the Appendix](#)). As a result, this mobile NMR spectrometer can also be used, without major modifications, at high field using the magnet and the probe part of the industrial spectrometer. In this type of application, the interest is methodological: development of a specific auxiliary channel, magnetization manipulation to control radiation damping, and development of new strategies (multiple acquisitions).

5. Back to the principles of NMR and practical consequences

The nucleus of atoms (^1H , ^{13}C , ^{15}N , ^{19}F , ^{31}P ...) is characterized by a quantum quantity, the spin, which is specific to each nucleus and determines its magnetic properties. The magnetic moment, μ , of a nucleus is directly related to I by its gyromagnetic ratio, γ , which is a constant depending on the nature of the nucleus. The magnetic moment being quantified, the energy states are degenerated.

When this nucleus is immersed inside the static magnetic field, \mathbf{B}_0 , the degeneration of energy level is lifted, leading to several energy levels according to the spin value (Zeeman effect). If the nucleus is then subjected to a radiofrequency wave at a specific frequency (Larmor frequency), the atomic nucleus will absorb the energy of this radiation and release it during relaxation mechanisms. This process at the origin of the NMR phenomenon depends on

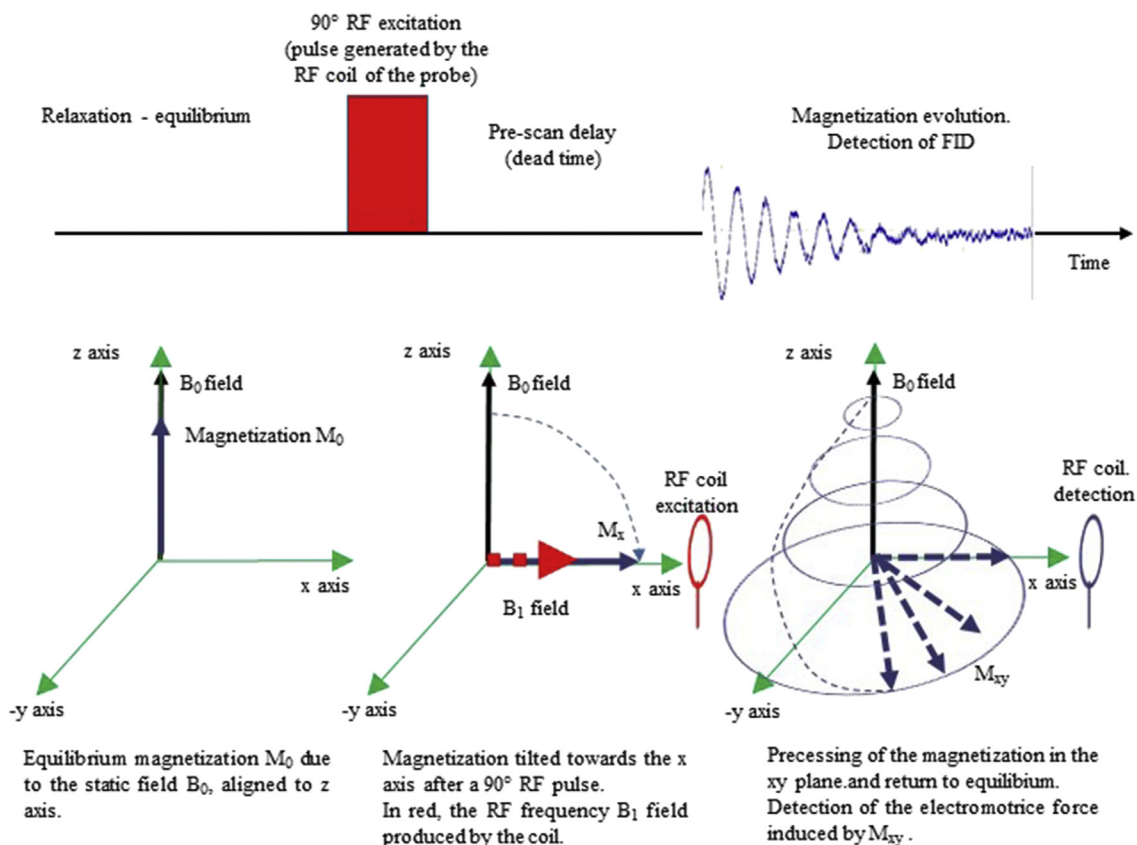


Fig. 1. Pulse-acquisition 1D NMR experiment. The macroscopic magnetization is aligned along the B_0 axis (Oz) at equilibrium state, ideally at the end of the relaxation delay. After an RF excitation pulse, M_0 is flipped in the Oxy plane. The precession of this magnetization gives the impulse response (or FID) which is detected.

the magnetic field and the properties of the molecules. The study and measurement of an isolated atom is not possible due to a lack of sensitivity of NMR; the result is that we are still dealing with a very large set of nuclear spins, the macroscopic result of which we will measure: magnetization. In the presence of a magnetic field B_0 , the resultant of macroscopic magnetization is collinear to B_0 at equilibrium and proportional to the spin populations (number of nuclei).

Under the action of B_0 , the magnetic moment, μ , of any magnetically active nucleus ($I \neq 0$) is subjected to a force,

$$\mathbf{C} = \vec{\mu} \wedge \vec{B}_0, \quad (1)$$

which causes a spontaneous precessional movement around the z axis (\parallel to the B_0 axis) at the Larmor frequency (resonance condition) (see Fig. 1),

$$\nu_{\text{Larmor}} = \nu_0 = -\gamma B_0 / 2\pi \quad (2)$$

where γ is the gyromagnetic ratio of the considered nucleus.

At equilibrium, all nuclei of a sample are subjected to B_0 . The resulting macroscopic magnetization (M_0) is oriented along the Oz axis. The first step of any FT-NMR experiment

is to excite the spin system with a (generally short) RF pulse to remove it from its equilibrium state by tilting the M_0 magnetization in the xy plane (see Fig. 1). The return of M_0 to equilibrium (denoted Free Induction Decay (FID)) is then detected and processed by FT to obtain a frequency spectrum in Hertz or expressed in parts per million (ppm). The magnetization, M_0 , observable for a set of nuclei of spin I , is given by the following law:

$$M_0 = \frac{N\gamma^2 \hbar I(I+1)}{3kT} B_0 \quad (3)$$

where T is the sample temperature, N is the number of nuclei per unit volume, I is the spin of the nucleus, k is the Boltzmann constant, and B_0 is the static magnetic field.

In practice, the magnetization, and therefore the sensitivity of a NMR experiment, is directly dependent on the value of B_0 . The exact magnetic-field dependence of the NMR sensitivity (considering the detection coil) can be written as the energy (E in Joule) detected in a coil of volume, V , namely:

$$E = M_0 \cdot B_0 \cdot V \quad (4)$$

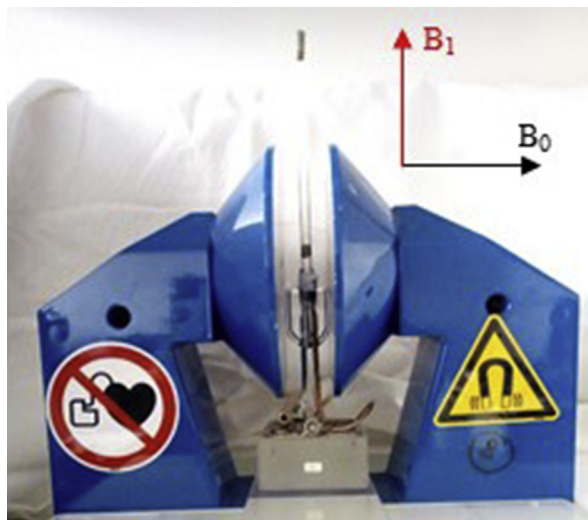


Fig. 2. Permanent horizontal magnet and its probe placed at the center of the magnet gap with a sample tube of 5 mm, inside.

where \mathbf{B}_0 and \mathbf{M}_0 are the static magnetic field and the magnetization per unit of volume, respectively.

A priori, it is therefore preferable to work with high magnetic fields, which are also at the origin of high-resolution NMR in liquids. For instance, at room temperature (300 K) and for a \mathbf{B}_0 field of 11.7 T (500 MHz), the detectable magnetization is about 3×10^{-6} (in SI unit), or an energy of about $10^{-4} \mu\text{J}$ for a sample volume V of 1 cm^3 ($E = \mathbf{M}_0 \cdot \mathbf{B}_0 \cdot V$). This corresponds to a reception power of about $1 \mu\text{W}$ for an acquisition time of 0.1 ms. As seen, the strength of detected signal is therefore very low. *De facto*, using a so-called low-field NMR, where the \mathbf{B}_0 strength

varies between 0.15 and 1.5 T, makes the experimental NMR detection much more critical.

From an electronic viewpoint, the voltage of detected signal (before its amplification) is of low amplitude, with an order of microvolt (μV). In practice, the experimental conditions as well as the instrumentation electronics must therefore be optimized to reach acceptable signal-to-noise ratios (SNR). This challenge can be met, thanks to the performance of modern electronic components. In addition, the transmission power required to energize a frequency band of about 10 kHz (for the ^1H nucleus) for a 1 cm^3 volume sample is about 11 W (@500 MHz or 11.7 T). For a low-field spectrometer, an excitation power of about 5 W for a pulse duration of about 20 μs is sufficient to generate an instantaneous tilting of 90° of \mathbf{M}_0 , i.e. in the xy plane.

6. NMR instrumentation

6.1. The magnet

In the applications of this study, we use a permanent magnet that is sufficiently homogeneous and stable to provide consistent NMR measurements. Considering the analytical purposes of low-field NMR and its transportability, the weight of the permanent magnet to be used must be reasonable ($< 80 \text{ kg}$), and generate at least a field of 0.15 T (1500 Gauss), leading to a resonant frequency for protons around 6.4 MHz. The axis of the magnet is horizontal and perpendicular to the probe (coil), which consists of a vertical solenoid, and in which the sample is positioned (5-mm tube) (see Fig. 2).

The NMR probe (the antenna) is carefully placed at the magnetic center of the magnet to both excite (with a perpendicular field, \mathbf{B}_1) and detect the signal along one of the four axes perpendicular to \mathbf{B}_0 ($x, -x, y, -y$). The probe includes its frequency and impedance system (tuning and

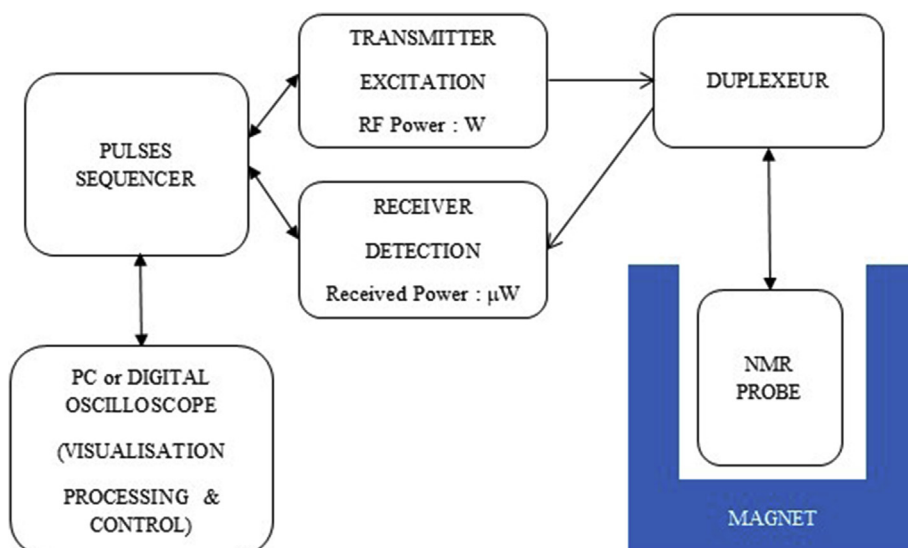


Fig. 3. Simplified "block" diagram of an NMR spectrometer.

matching) that is connected to the electronic chain of the equipment. Note that it is easy to add two external coils on either side of the magnet to produce a field gradient that can potentially be used for possible monodimensional imaging applications. As the static magnetic field is not homogeneous over all the surface of the magnet, it is therefore necessary to accurately measure the value of B_0 for determining the Larmor frequency (ν_0). To this aim, a field map can be made within the permanent magnet to determine the exact value using a gaussmeter (Teslameter) that is slowly moved into the air gap of the magnet. In our case, the value measured at the center of the air gap is 1492 Gauss, which corresponds to a resonance frequency, ν_0 , of 6.28 MHz for ^1H .

6.2. Excitation vs detection

To perform FT-NMR experiments, three steps are necessary: i) create a homogeneous magnetic field, B_0 ; ii) excite the spin system to tilt it from its equilibrium state by means of an excitation coil generating a B_1 field perpendicular to B_0 ; and iii) detect the FID signal by means of a receiver coil (which is also the same generating the B_1 field excitation), the whole satisfying the Larmor's resonance condition, $\nu_{\text{exc}} = \nu_{\text{dec}} \approx \nu_0$.

The "block" diagram setting up the main parts of an NMR spectrometer is shown in Fig. 3, which shows the permanent magnet and the NMR probe inside the magnet, an electronic "transmitter" block responsible for exciting the spin system with RF powers of several dozen watts, and a second "receiver/detection" block that amplifies the FID signal of very low power (μW) and demodulates the signal

in two components of low frequency (separate operations). These two steps can be controlled by a "programmable sequencer" that generates the multiple RF pulses trains and controls the delays during the experiment. For very simple experiments (pulse-acquisition), a laboratory pulse generator is sufficient. Finally, data processing and visualization can be performed using a simple oscilloscope, but in a more modern way using a computer (PC) with its software interface. It should be noted that the probe is connected to the excitation and reception blocks *via* a unit called duplexer. This element can be seen as a signal "switcher" that aims to protect the preamplifiers of the detection channel from high RF excitation power. The duplexer directs all RF power to the probe during the excitation step (by closing access to the receiving channel) first, and then redirects the detected NMR signal to the preamplification/demodulation chain.

For a spectrometer operating at frequencies of the order of 15 MHz and with relatively low powers, a simple structure based on a π quarter-wave filter can be used to perform this function. It is a design strategy for a very low-cost mobile NMR. First, the π filter requires only a few electronic components at a very low cost (self and capacitivities), capable of withstanding high powers (several Watts). In addition, it works automatically and there is no need to control it during acquisition, simplifying any pulse programs. The realization of the circuit is simple and fast. We will describe how to make a quarter-wave duplexer in Section 8.4 for a frequency around 6.4 MHz. Because the π filter (or quarter-wave filter) is a circuit tuned to the Larmor frequency of the nucleus under consideration, its use is limited to a small frequency band.

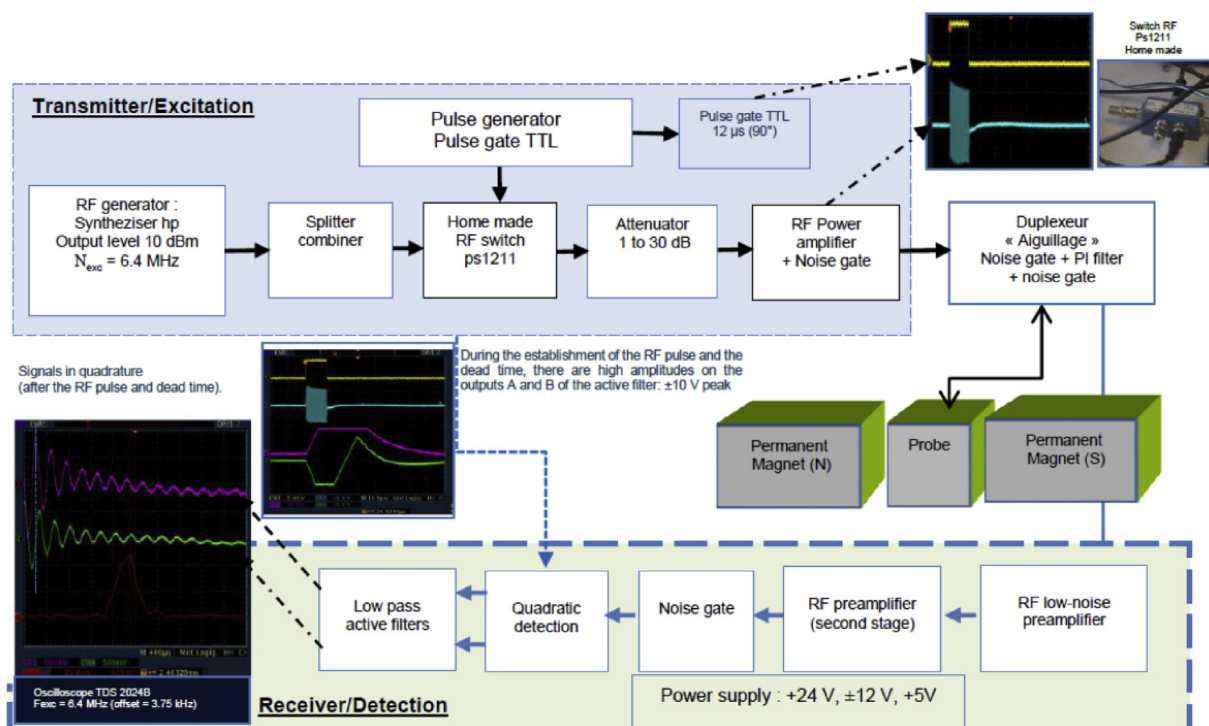


Fig. 4. Schematic description of the NMR spectrometer blocks along with the visualization of some of key signals obtained. Note the presence of the duplexer.

For wide band applications (*i.e.* frequencies from MHz to a few hundred MHz), a duplexer based on fast RF electronic switches with fast diodes is mandatory. This type of duplexer is much more complex to implement because a pulse program within a microprocessor must control it. The description of such a circuit is out of scope for the version of the mobile NMR spectrometer we present in this article. We are targeting here a low-cost mobile NMR, with laboratory instrumentation, a duplexer π filter, and a low-field magnet operating at 0.15 T (or 6.4 MHz for ^1H).

6.3. The lock system

An electronic locking system may be easily added to make the mobile NMR device capable to continuously control the stability of the magnet, to avoid a slow shift of the magnetic field over time [see for instance, pp. 44 to 50 of ref. 19]. The lack of lock control can be particularly troublesome when working at high magnetic fields, and/or when recording long-term acquisition NMR experiments. Considering the aims (pedagogical aspects), application domains (1D NMR, detection of abundant nuclei), and the cost of this home-made instrument, no locking system has been integrated into the instrument.

7. NMR device synoptic

For clarity, the general electronic diagram of a NMR spectrometer is schematically proposed in Fig. 4. As seen, the attenuators and tuning/matching circuits are present at the input and output of the power amplifier to avoid excess power that can be harmful to the detection chain. The excitation step generates one or more RF pulses at the Larmor frequency. These pulses are directed *via* the duplexer to the NMR probe to tilt \mathbf{M}_0 perpendicularly to the z axis. The high-frequency resonance free induction decay signal (HFFID) is then routed *via* the duplexer to the detection circuit. The detection circuit includes at least two very low-noise RF preamplifiers to amplify this HFFID signal. The signal is then demodulated with an intermediate frequency, in quadrature (quadratic detection leading to a 90° phase shift of the signal), to generate a real (R) and an imaginary (I) component accordingly to Eq. 5,

$$S(t) = A \times \{\cos(2\pi\nu_1 t + \phi) + i \sin(2\pi\nu_1 t + \phi)\} \quad (5)$$

where A is the amplitude and ϕ is the phase of the detected signal. This signal is then filtered to eliminate unwanted frequency components. At the output of the active "low-pass" filter, the signal is lowered to the 10-kHz range (often wrongly called « audio filter »). The R and I components of the signal can then be observed and checked on an oscilloscope or digitized *via* an analog-to-digital converter (ADC) for further processing.

Although the spectrometer is optimized to work around the 6.4 MHz frequency, it can easily operate efficiently over a wider band. All the electronic components required for the construction of a low-cost NMR spectrometer are listed in Table A1, with the adequate associated nomenclature. The choice of the bandwidths of the various elements

present in the nomenclature makes it possible to work up to frequencies around 500 MHz. The assembly of the various blocks can be made entirely on a printed circuit board or more simply by connexion components (BNC or SMA outputs).

8. Key electronic elements of the spectrometer

Let us now review the key elements of the equipment (probe, switch, RF preamplifier...) in the order of the synoptic block diagram presented in Fig. 4.

8.1. RF pulse generator

The first important function of an NMR spectrometer is the generation of the basic frequencies (BF) necessary for excitation/reception of signal. The RF excitation pulse is realized by means of a switch connected to a frequency synthesizer and a pulse generator. However, to perform a synchronous detection (resonance), part of this frequency must be directed to the detection circuit.

The generation of RF pulses can be performed in different ways. The first one consists of using a standard frequency and pulse generator present in any physicist laboratory. To ensure sufficient stability and accuracy for NMR, the resolution of the function generator must be at least 10^{-6} . Typically, an arbitrary function generator of type HP33120A or Agilent 33220A operating up to 15 and 20 MHz, respectively, with a power of 10 dBm is perfectly suited for a low-field NMR spectrometer. For pulses, the pulse generator can also be used to produce square-shaped pulses. For pulses of the order of 10 μs , a pulse generator with a resolution of less than 100 ns is sufficient (*e.g.*, Keysight 33220A). The time and frequency parameters are adjusted from the control panels of these laboratory instruments. In this case, only very simple single-pulse sequences are possible: 90° pulse - acquisition; 180° inversion pulse - acquisition, etc. The primary advantage of laboratory equipment is their availability at a lower cost, their speed, and simplicity of implementation (no software development for pulse programs). The second way to generate pulse sequences is to use a board based on an Advanced Reduced instruction set computing Machines (ARM) microcontroller, a Field Programmable Gate Area (FPGA) unit, and a Direct Digital Synthesis AD9959 (DDS), which allow digital frequencies up to 300 MHz to be synthesized. The advantage in this case is to have a very compact and powerful unit to generate the basic pulses and frequencies. It can be developed, for example, from an NXPLPC1768 ARM, an Altera Cyclone II FPGA [20, 21], and a DDS. This board is capable of sequencing four fully independent channels. The FPGA allows the most complex pulse sequences to be performed, and the ARM component then manages both the system and data processing. This option allows stand-alone battery operation for outdoor (off-laboratory) use.

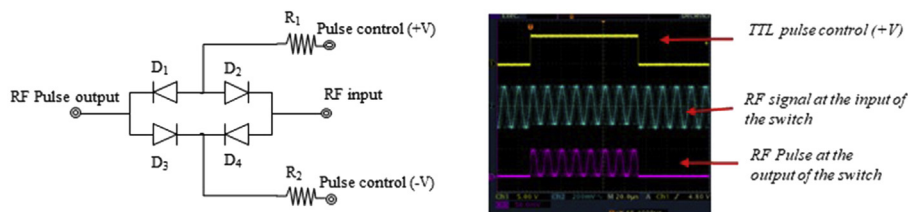


Fig. 5. (On Left) Structure of an RF switch based on fast diodes. (On Right) Example of input and output signals from an RF switch. Here, the RF pulse has a length of 80 μ s.

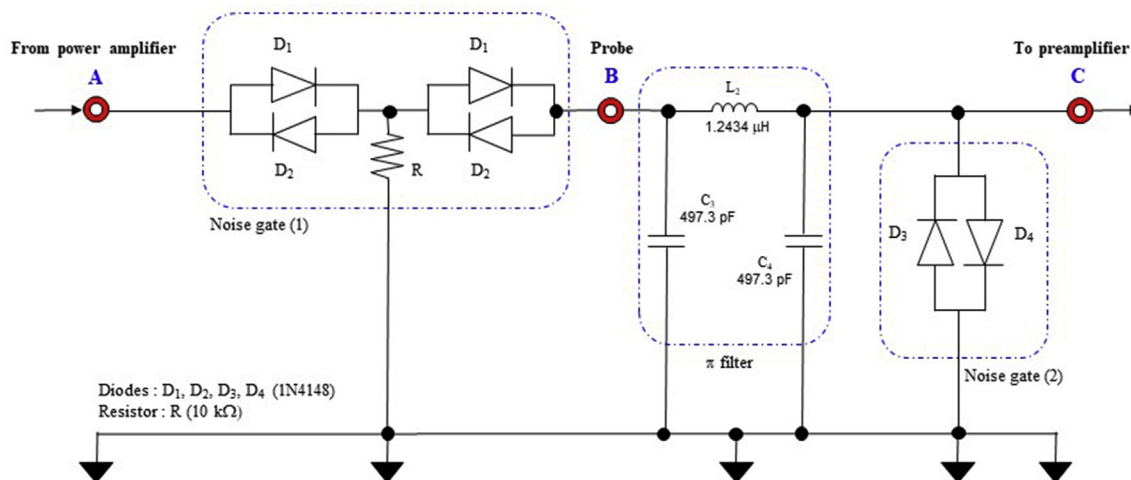


Fig. 6. Electrical diagram of the duplexer, including two diode arrays and the π filter. The π filter has a minimum RF signal at the receiver (port C) when there is a maximum on the probe (port B). The "noise gate 1" isolates any signal feedback from the probe to the power amplifier to minimize losses during detection. The "noise gate 2" protects the detection chain from residual voltages eliminated by the π filter.

8.2. The switch

The role of this circuit is to switch the RF signal for a controlled pulse duration. This element can be realized by means of four fast diodes (type 1N4148) mounted in head to tail or with a PS1211 mini-circuit component (switch), which offers fast switching times (< 500 ns) and very low channel imbalance (see Fig. 5).

8.3. The RF power amplifier

As our mobile NMR is aimed to work from low to high field, it is necessary to have a broadband RF excitation pulse. The excitation provided to the spin system must be sufficiently high, with relatively short RF pulses (10–100 μ s) to excite a bandwidth covering the spectral width of an observed nucleus. The selected power amplifier must be able to operate in pulse mode but also in continuous wave (CW), if nuclear decoupling or any other type of low power excitation needs to be achieved. It is imperative that the

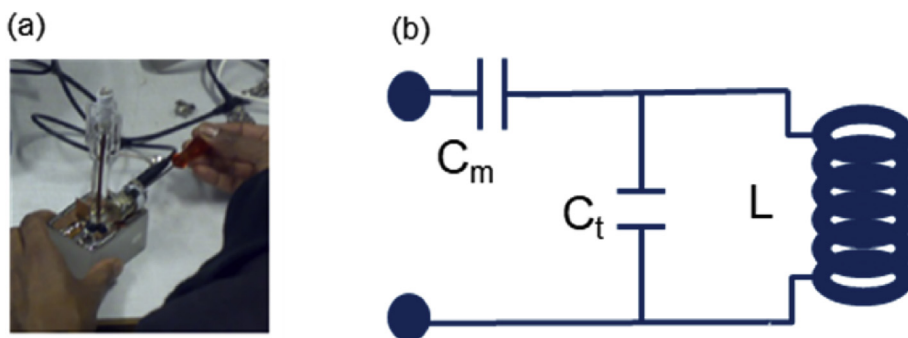


Fig. 7. (a) Example of a probe made of glass with its integrated capacitive devices C_m and C_t . (b) NMR probe wiring diagram: C_m and C_t are adjustable capacitances from 5 pF to 500 pF, inductance coil, $L = 2\text{--}3$ μ H.

amplifier has a good linearity to adjust the RF pulse accurately. A power amplifier generates a high noise level at the output even at rest. It is therefore necessarily not activated. We used for our prototype a ZHL-5W-1 unit operating in a frequency range from 1 to 500 MHz.

8.4. The duplexer

As already mentioned above, the duplexer has a very important role, in particular, to protect, during reception, the low-noise preamplifier chain from the high RF power emitted during excitation. It is a three-port system A, B, and C (see Fig. 6), where the first port (A) is connected to the power amplifier, the port (B) is connected to the probe, and the output (C) is directed to the low-noise detection pre-amplifier chain. For broadband systems, it is preferable to use a switch network electronically controlled by the pulse sequencer. In low-frequency (low-field) NMR applications, a structure based on diode arrays and a quarter-wave filter can be used; the diagram is shown in Fig. 6. A first network of four head-to-tail diodes (noise gate 1) acts as an automatic switch: these diodes are blocked (open circuit) for small amplitudes, thus preventing any return of the signal from the probe to the power amplifier. The quarter-wave system, referred to as the π filter, has an input (B)/output (C) phase shift of 90° . It is called a quarter wave because it has a phase shift optimized at 90° around a given frequency, in our case, around 6.28 MHz. This filter consists of

a choke and two capacitors. The filter input is connected to the probe (port B) and the output (port C) to the low-noise preamplifier chain. For a specific frequency, a quarter-wave filter has a minimum amplitude at its output (C) when its input (B) is maximum. To reinforce the protection of the preamplifiers, a second network of head-to-tail diodes can be incorporated to protect the detection circuit by short-circuiting (at the threshold of the diodes) any residual voltages. The filter setting must combine input and output impedance matching, as well as phase shift and attenuation at the resonance frequency. The characteristics of the quarter-wave filter were adjusted to the network analyzer with the Smith diagrams of the input (S11) and output (S22) impedances as a function of frequency. The input (S11) and output (S22) impedances have been adjusted to have an actual value as close as possible to $50\ \Omega$ (in practice $44.5\ \Omega$ at the resonance frequency of 6.28 MHz), while maintaining a 90° phase shift between the input and output.

8.5. The NMR probe

Different probe body shapes (solenoid, saddle coil) can be produced and used depending on the adaptability to the magnet. The most simple detection antenna consists of a solenoid coil. A probe body made entirely of glass (boro-silicate) can be used (see Fig. 7a). For a 10-mm probe, the solenoid is wound on the glass tube with an inner diameter

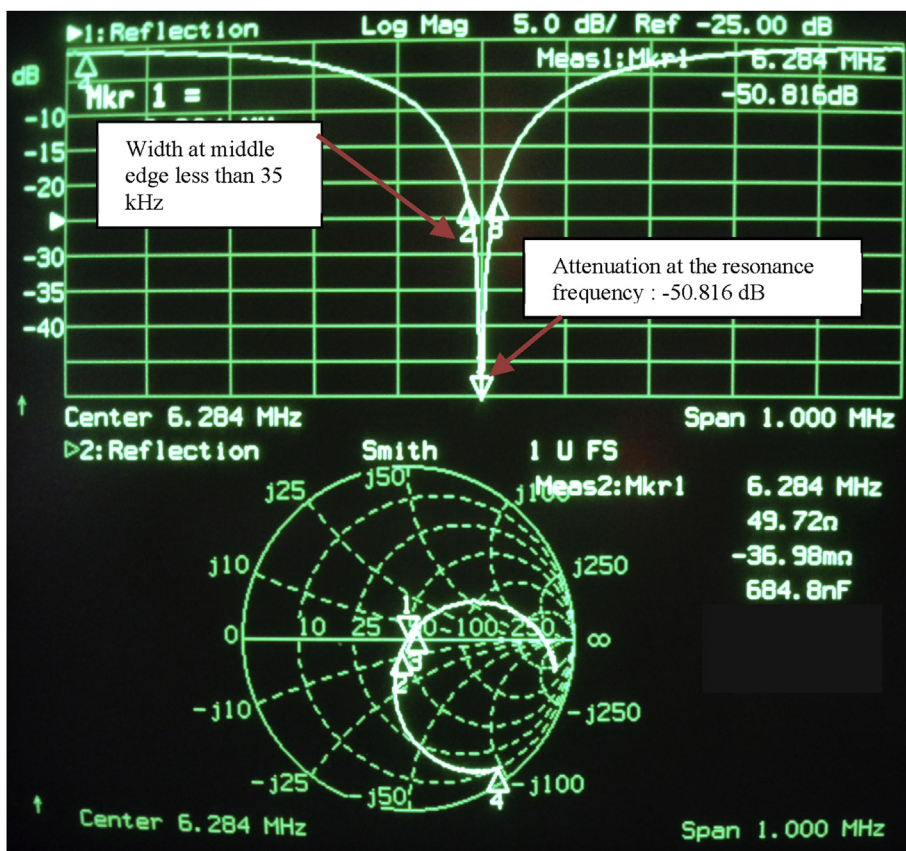


Fig. 8. Picture showing the measurement of the frequency tuning and impedance matching curve (top) and on the Smith chart (bottom) on a network analyzer.

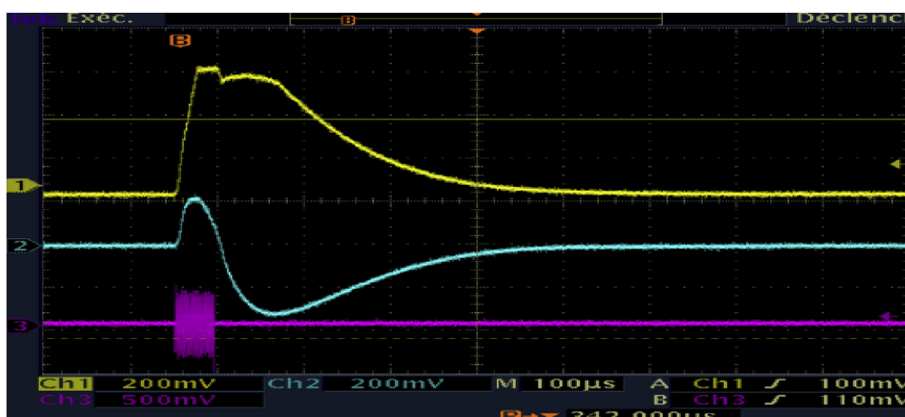


Fig. 9. Observation of the probe's dead time: about 250 μ s. At the top and in the middle are displayed the two quadrature detection channels (yellow and blue). At the bottom is displayed the rectangular (purple) RF excitation pulse (50 μ s of length).

of 10 mm, thus allowing the detection of a large amount of analyte, but small diameter (5 mm) could be used. The capacitive network is integrated into the probe. The solenoid is made of rigid copper wire with a diameter of 1–1.5 mm and includes about 20 turns with a diameter of 10 mm over a length of 20 mm; the better the quality of the winding, the smaller the artefacts will be. The elementary structure of the frequency tuning and impedance matching system is described in Fig. 7b. A network of capacitors adjustable from 5 to 500 pF allows frequency tuning (C_t) and impedance matching (C_m) for a coil of the order of 2.2 μ H. The tuning allows the probe to be adjusted to the resonance frequency ($\nu_0 \approx \nu_1 = 1/2\pi\sqrt{LC}$). Impedance matching is performed at this frequency to adjust the probe circuit to 50 Ω and minimize the energy losses.

Frequency tuning and impedance matching are performed using a network analyzer to accurately position the probe's attenuation peak at the desired resonance frequency while observing the impedance evolution on the Smith chart (see Fig. 8). Impedance matching consists of refining and lowering the resonance peak as low as possible to absorb all the energy in the coil. In our case, the resonance is adjusted with the network analyzer at 6.284 MHz, the attenuation obtained is -50.816 dB with a very good impedance matching (49.72 Ω), and consequently a good detection sensitivity is obtained for a low-field spectrometer. The half-height width obtained at this frequency is 35 kHz, giving a quality coefficient Q of 179, a very acceptable value.

Once the frequency tuning has been achieved, the second important parameter to be estimated is the dead time (or response time) of the probe (see Fig. 9). The dead time is the time required for the probe to dissipate the energy stored during excitation. Some of the excitation energy is sent to the probe via the duplexer, but once the RF pulse has disappeared, significant energy remains in the probe. The higher the quality factor Q of the probe, the higher the dead time is. This is a disadvantage for detection, because it will be necessary to delay the measurement of this time: i) not to destroy the low-noise preamplifiers and ii) not to superimpose the low detection signal on the high amplitude due to the probe signal.

In the model presented in this article, the dead time is about 250 μ s before the two quadrature detection channels return to zero (see Fig. 9). The detection of the useful signal needs to be delayed by this delay and the detection circuit is reinforced, by a network of diodes, to support the peak amplitudes of several volts for a few hundred microseconds.

8.6. The receiving chain: radio frequency preamplifier

The signal from the probe is too weak and cannot be used directly, and so it must be amplified. The amplification chain consists of two stages with radiofrequency preamplifiers in series. The noise factor of a preamplifier cascade is almost equal to that of the first stage. A "BA01500" preamplifier with the lowest factor is placed first, followed by a RF2046 home-made broadband preamp unit. The low-noise amplifier LN BA01500-35 has a frequency range varying from 0.1 to 500 MHz with a gain of 35 dB. The intermodulation product power at the output is $+12$ dBm (output IP3), and the undistorted input power is -35 dBm (1 dB compression point). Thus, the maximum permissible input power is therefore -35 dBm.

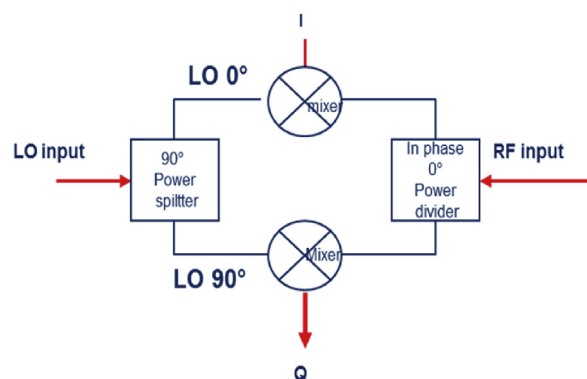


Fig. 10. Structure of a Quadrature Intermediate Frequency Mixer demodulator (QIFM). Mixers: X2L-06–414 from Pulsar Microwave.

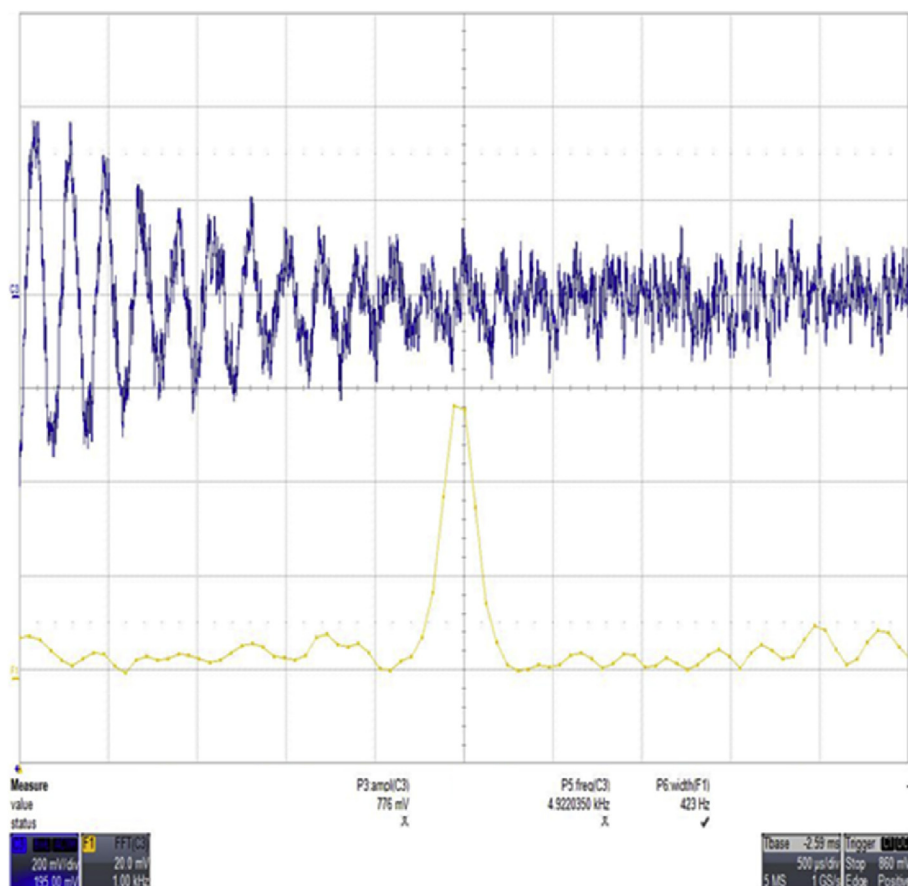


Fig. 11. ^1H signal (one component) of a water sample (5 mm) obtained (one SCAN: $\text{NS} = 1$) after the demodulation stage and low pass filtering, observed on a digital oscilloscope with FFT. Top: FID component. Bottom: spectrum: presence of the peak at 4.92 KHz. Note the linewidth of 423 Hz. The Larmor frequency was $\nu_0 = 6.4$ MHz. The RF excitation pulse length is 50 μs . See Appendix for the details of the experimental setup.

Our home-made preamplifier is made with three RF2046 components in series: the overall gain of the channel measured at the network analyzer (without distortion) is 35 dB at 6 MHz and 31 dB at 1 MHz and the operating frequency range is from DC continuous to 1 GHz. The output intermodulation product (IP_3) is +23 dBm and the noise factor (NF) is equal to 3.8 dB. This configuration of the elements as well as their performances allows us to ensure the detection of signals on a large dynamic range without distortion: typically from some 100 μV to 14 mV.

8.7. Quadrature demodulation

Quadratic demodulation is based on the principle of synchronous detection to obtain real and imaginary components of the NMR signal. Its structure is based on a Quadrature Intermediate Frequency Mixer (**QIFM**) demodulator whose schema is given (see Fig. 10). It consists of a 90° hybrid splitter, two mixers, and a 0° combiner (summing). Each of these elements is listed in the table of components (see Table A1).

At the input of the 90° hybrid divider, we have the base frequency of the Local Oscillator (**LO**) input spectrometer; this is actually the Larmor frequency of the spectrometer. At the input of the 0° divider, we have the FID: this is the high-frequency response RF. At the output of the quadratic demodulator on channels I (imaginary) and Q (quadratic), the signals are of the form: i) $\cos [2\pi(\nu_{\text{rf}} + \nu_0)t] + \cos [2\pi(\nu_{\text{rf}} - \nu_0)t]$; ii) $\sin [2\pi(\nu_{\text{rf}} + \nu_0)t] - \sin [2\pi(\nu_{\text{rf}} - \nu_0)t]$. We therefore obtain two quadrature components with all the "sum" and "difference" frequencies of the demodulated signals: $2\pi(\nu_{\text{rf}} + \nu_0)$ and $2\pi(\nu_{\text{rf}} - \nu_0)$, plus harmonics. Only low-frequency signals in the 10-kHz range are interesting, and the signal must be filtered to retain only the lowest frequencies using a low-pass filter.

8.8. Double active audio low-pass filter

An active double low-pass filter is required to keep only the low frequencies of the demodulated signals (see Fig. 11). Each filter is designed to charge the demodulators on 50 Ω and consists of two stages allowing an overall gain of about 30 dB in voltage. As around the resonance, the

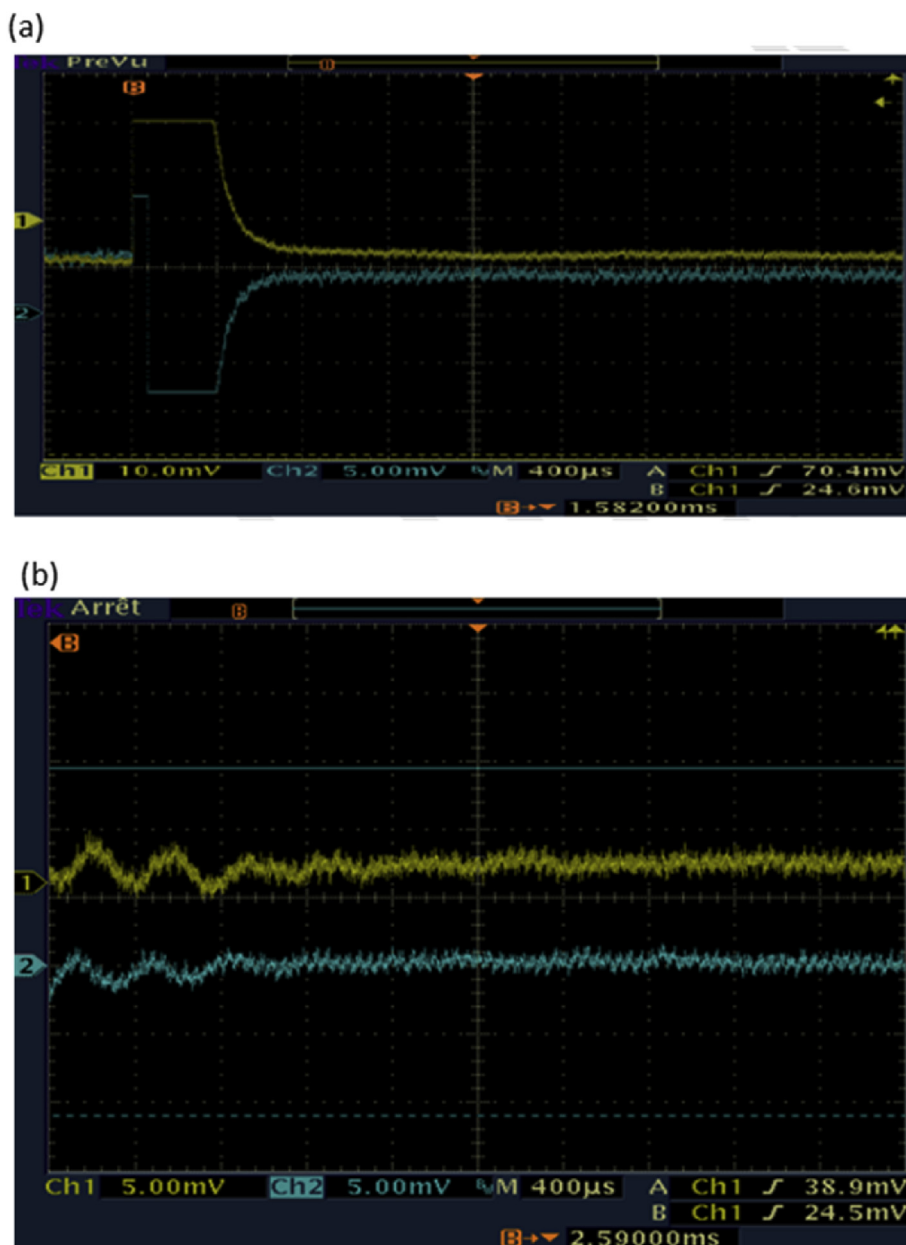


Fig. 12. (a) No NMR tube: real (high, channel 1) and imaginary (low, channel 2) components at the output of the audio frequency amplifier after an RF excitation pulse length of 50 μ s. (b) Example of one-scan quadrature detection of a low-magnitude ^1H signal (5-mm water tube). See Appendix for the experimental setup.

measured frequency differences are about ten kilohertz, the filter bandwidths are adjustable over a range from 100 Hz to 500 KHz.

The two filter channels must be adjusted to have the same frequency and phase responses throughout the useful bandwidth. This is to avoid unbalancing between the real and imaginary components of the signal, which would degrade the spectrum in frequency (quadrature artifact, zero frequency...). Precise gain and phase adjustments in the bandwidth are made with a network analyzer, the only true and effective way to visualize

the response of both channels as a function of frequency.

9. Setting of spectrometer

9.1. Probe control

The tuning of the probe to the resonance frequency has been described above.

9.2. Cancellation of the DC offsets

Any DC voltage offset in the detection circuit generates a zero frequency component that will affect the quality of the spectra. To reduce this unwanted effect, it is important to carefully cancel the offset voltages of the active double low-pass audio filter to have a signal close to zero when the input signal is nul.

9.3. Signals at the output of the audio amplifier without sample

At first, we work without putting a sample in the magnet to minimize the imperfections of the assembly: adjustment of the LO level (10 dBm) and cancellation of continuous offsets. In the absence of a sample, the voltage offsets are less than 1 mV. In addition, note that the deadtime of the probe after the RF pulse is 400 μ s (see Fig. 12a).

9.4. Signals at the output of an audio amplifier with sample

The imperfections of the assembly having been minimized, we introduce our sample (a 5-mm tube of water), and we visualize the precession signals in quadrature mode (see Fig. 12b). Note the very low magnitude of the signals.

9.5. Optimization of the 90° flip angle

The NMR signal obtained at this step, although observable, is relatively small in amplitude. To maximize the amplitude of the FID, it is now necessary to adjust the strength excitation pulse for a tilting of \mathbf{M}_0 equal to 90°.

For this reason, the duration of the pulse is varied until a magnetization is obtained whose intensity in the xy plane is maximum (see Fig. 13). The amplitude variation of the magnetization vector is observed as a function of the excitation pulse duration. In our case, the pulse duration is between 13 and 14 μ s. Another approach is to determine the pulse duration corresponding to an angle of 180°, for which the macroscopic magnetization is then null (tilted to the $-Oz$ axis).

9.6. Increase in the S/N ratio

The signal-to-noise ratio of an NMR experiment involves numerous factors such as: i) the magnetization \mathbf{M}_0 , ii) the filling factor and the quality factor of the probe, iii) the noise factor of the preamplifier, and so on [22]. After optimization of all the electronic hardware and parameters, we can also increase the SNR during the acquisition. The SNR can be improved by averaging the FID signals over a large number of acquisitions (NS scans). Indeed, on an average of NS acquisitions, the noise will be averaged toward zero due to its random origin, whereas the useful signal will increase on average. Thus, the SNR is improved at the root of NS as displayed in Fig. 14.

10. Possible applications

Benchtop FT-NMR allows many applications in chemistry, such as the analysis or characterization of small molecules, or the monitoring of chemical reactions. The teaching of NMR methodologies (effects of excitation/inversion pulses, pulse calibration, optimization of acquisition parameters, as well as the description of some experiments as the measurement of relaxation

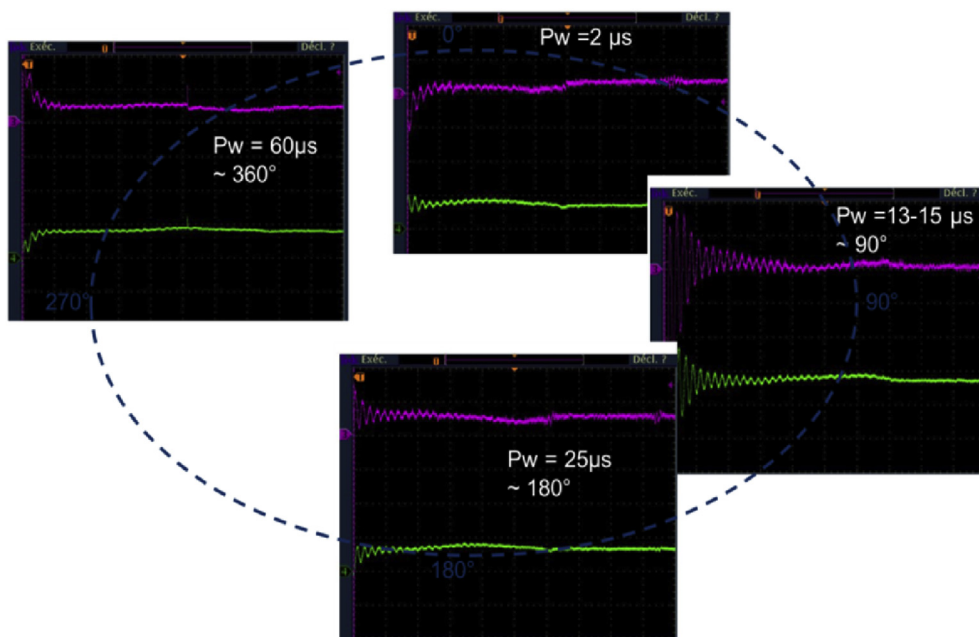


Fig. 13. Adjustment of the RF excitation pulse duration to obtain a maximized signal corresponding to a 90° pulse.

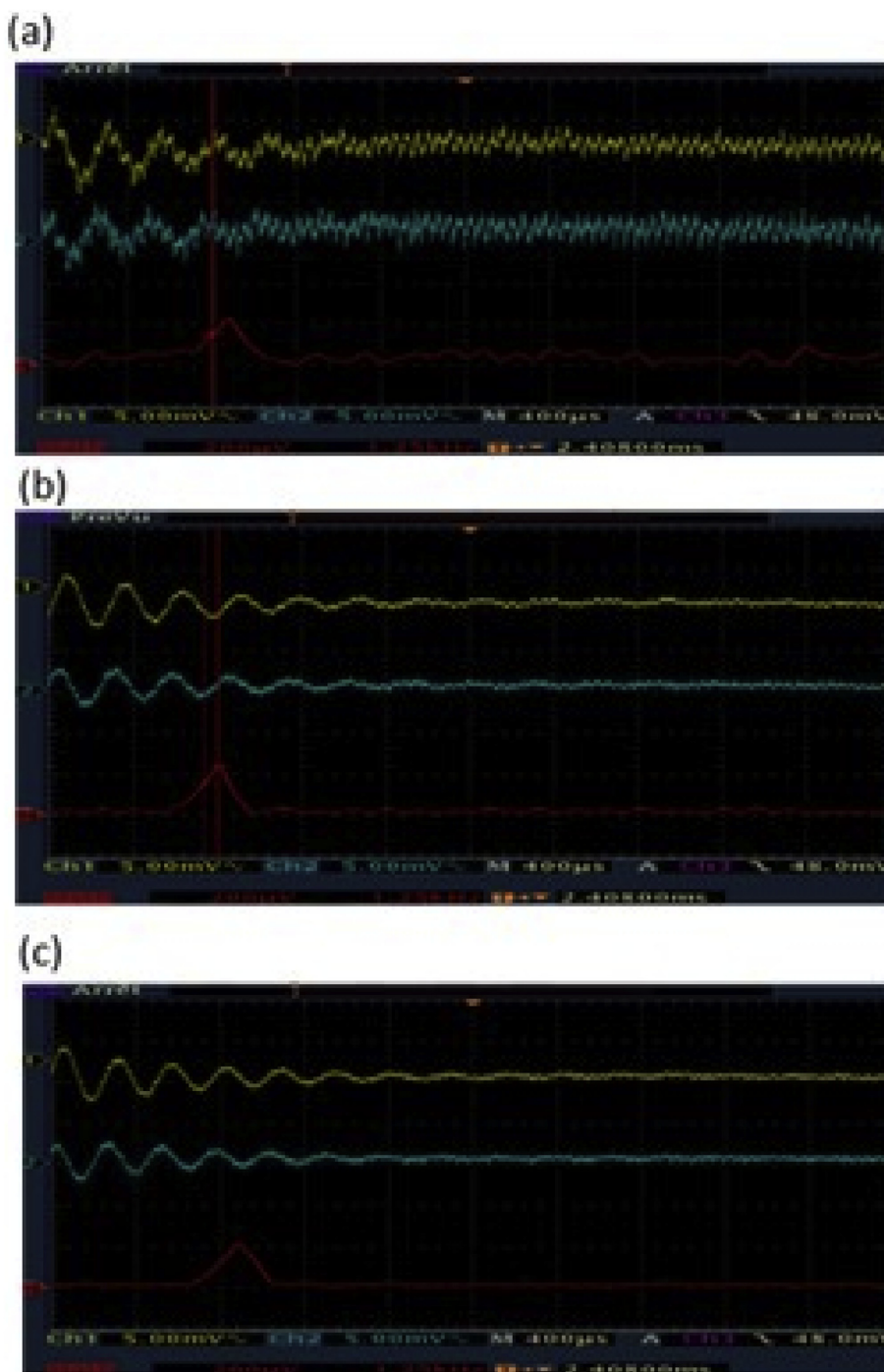


Fig. 14. Effect of averaging on the SNR. Comparison of the FIDs obtained by adding successively: (a) 2, (b) 32, and (c) 64 scans (5-mm water tube). On (c), note the vertical line setting at 3.45 kHz. See Appendix for the experimental set up.

parameters, spin echoes...) is also another aspect of their use. The portable low-field NMR is particularly suitable for nuclei with high natural sensitivity, such as ^1H , ^{19}F , ^{31}P , and ^{29}Si . However, other « exotic » nuclei with high gyromagnetic ratios such as ^7Li , ^{11}B , and ^{23}Na can also be detected [27,28].

A (home-built) mobile NMR spectrometer can also be used to develop projects that aim to couple NMR and optical microscopy in a semiconductor [23] to locally polarize nuclear spins (dynamic polarization) and measure the spatial profile of nuclear magnetization using an optically detected NMR, with good spatial resolution. This

fundamental study of nuclear spin diffusion does not require a high magnetic field (less than 0.15 T). For this kind of experiment, NMR functionalities are reduced to signal excitation and detection for parameter optimization, without the need for frequency resolution.

11. Conclusion

The low-field FT-NMR spectrometer presented in this article is fully operational. This revisited instrument can be exploited to carry out simple NMR measurements such as the determination of chemical shifts (δ), spin echoes, measurement of $T_2(^1\text{H})$ or $T_1(^1\text{H})$ relaxation times, etc. On the other hand, it provides a pedagogical approach to the teaching of NMR to students at all levels, from the physical concept to the electronic realization through the many possible analysis applications [24, 25, 26]. It is also an excellent demonstration element at scientific events for a more general public. Finally, it can be inserted in real experiments of physics requiring NMR to be coupled with another measurement technique such as optics, for example.

Noteworthy technical developments of the instrument are possible. Thus if the permanent magnet is equipped with an additional pair of coils with currents in the range of 1–3 A, it can then be injected to create additional field gradients to \mathbf{B}_0 . The spectrometer is then able to perform some simple 1D imaging experiments or to experimentally measure molecular diffusion coefficients.

Although sensitivity is reduced at low-field strength, quantification and measurement of relaxation and diffusion parameters of neat samples or solutions are possible [27, 28]. This kind of portable NMR spectrometer could then appear the ideal tool for outdoor NMR analysis of ecosystems.

Conflicts of interest

The authors declare no conflicts of interest.

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Appendix

Definitions of acronyms and notations used

ARM: Advanced Reduced instruction set computing Machines

ADC: Analog-to-Digital Converter

BNC: Bayonet Neill–Concelman (connector)

\mathbf{B}_0 : Static magnetic field

BF: Basic Frequency

\mathbf{B}_1 : RF coil. The term \mathbf{B}_1 is used to characterize the field, in the plane, orthogonal to the static \mathbf{B}_0 field. In the

terminology frequently encountered in the fields of NMR, the \mathbf{B}_1 coil is a resonant LCR circuit tuned to the Larmor frequency. This coil is used for the detection and the excitation of the nuclei in FT-NMR experiments. L is inductance, C is capacitance, and R is resistance.

CW: Continuous Wave

dB: Decibel

dBm: Power in decibel: 0 dBm = 1 mW/50 Ω

DDS: Direct Digital Synthesis

FFT: Fast Fourier Transformation

FID: Free Induction Decay

FT: Fourier Transformation

FPGA: Field Programmable Gate Area

HFFID: High-Frequency Free Induction Decay

LO: Local Oscillator

M_0 : Magnetization at equilibrium

NF: Noise Factor

NMR: Nuclear Magnetic Resonance

Q: Quality factor

QIFM: Quadrature Intermediate Frequency Mixer

RF: RadioFrequency

SMA: SubMiniature version A (connector)

SNR: Signal-to-Noise Ratio

TROSY: Transverse Relaxation Optimal Spectroscopy

TTL: Transistors Transistors Logics

T_2 : Spin–spin relaxation time

T_1 : Spin–lattice relaxation time

ν_0 : Larmor frequency (MHz)

Table A1

Characteristics and nomenclature of electronic components.

Characteristics	Designation	Number of devices	Frequency range (MHz)
1500 G permanent magnet, 40-mm air gap, NMR homogeneity on a useful diameter of 20 mm, divergent polar pieces on a diameter of 200 mm, height 400 mm	TE2M C4-1367 1500 G	1	6.2–6.8
Directional coupler 30 dB, 3W, SMA (Pulsar)	C5-08-411	2	5–1000
SMA connector (pulsar)		20	0.1–1000
Mixer level 7 dBm SMA (Pulsar)	X2L-06-414	2	(LO/RF) 1–1000; (IF) DC-1000
Power divider/combiner, 2 way (0°), phase balance: 2°, amplitude balance: 0.3 dB	P2-09-411	3	5–1000
Insertion loss 0.8 dB, SMA (Pulsar)			
90° hybrid power divider Phase balance 2°, amplitude balance 0.4 dB, insertion loss 0.5 dB, SMA (pulsar)	QE-01-412	1	2–10
Low-noise RF preamplifier NF = 2, gain: 35 dB, RF output power: 5 dBm Absolute maximum rating input	BA01500-35	1	0.1–500
RF power: –35 dBm (Elhyte)			

Table A1 (continued)

Characteristics	Designation	Number of devices	Frequency range (MHz)
Low-noise RF preamplifier NF = 2.9, gain: 25 dB, output RF power: 5 dBm (mini circuit)	ZFL1000LN+	1	0.1–1000
Low-noise RF preamplifier RF; NF = 3.8, gain: 23 dB, RF output power: 5 dBm (home made)	RF2046	3	DC -1000
Power amplifier	ZHL 5W–1	1	
Operational amplifier	OP27	1	DC - 8
Operational amplifier	OP37	1	DC - 60
Resistor	1 k Ω , 10 k Ω	10	
Variable capacitor	5 pF–500 pF	4	
Probe (home made)		1	6.1–6.8
Sequencer board with ARM + DDS (home made)	NXPLPC1767+ DDS 9959	1 1	0.01–301
Or frequency generator and laboratory pulse generator (see text).			
Analog or digital oscilloscope with FFT		1	
AC/DC adapter 24 V/6.4 A	AHM150PS24	1	
Total estimated cost			~10 k€

Experimental setup (except otherwise in the text)

All NMR and test experiments have been made with the electronic setup outlined in Fig. 4. A permanent magnet with a field of 1500 G (6.4 MHz) is used for the static magnetic B_0 field (see Fig. 2). A home-made solenoid-type probe device and a 5-mm water sample tube were used for the tests. The frequency and the RF pulse duration have been set with the frequency generator and the pulse generator mentioned in the text. Data have been acquired and processed using some digital oscilloscopes with sensitivities of 1 or 2 mV/div.

Characteristics and nomenclature of the electronic components

Table A1 provides the characteristics and nomenclature of the electronic components used.

References

- [1] F. Bloch, *Phys. Rev.* 70 (1946) 460–474.
- [2] P.E. Bloembergen, *Phys. Rev.* 73 (1948) 679–746.
- [3] F. Bloch, W.W. Hansen, M. Packard, *Phys. Rev.* 70 (1946) 474–485.
- [4] J. Keeler, *Understanding NMR Spectroscopy*, John Wiley & Sons, 2005, 978-0-470-01786-9.
- [5] M. Goldman, *Actualité Chimique* 273 (2004) 57–60.
- [6] R.R. Ernst, W.A. Anderson, *Rev. Sci. Instrum.* 37 (1966) 93–102.
- [7] E.L. Hahn, *Phys. Rev.* 80 (1950) 580–594.
- [8] J. Jeener, Lecture notes from the Ampère Summer School in Basko Polje, Yugoslavia, September 1971, later reprinted, in: M. Goldman, M. Porneuf (Eds.), *NMR and more, in honor of Anatole Abragam*, Éditions de Physique, Les Ulis, France, 1994.
- [9] A. Abragam, *Principle of Nuclear Magnetism*, Oxford University Press, 1961.
- [10] I. Solomon, *Phys. Rev.* 99 (1953) 559–565.
- [11] D. Canet, *La RMN: concepts et méthodes*, InterÉditions, Paris, 1991.
- [12] A. Abragam, M. Goldman, *Nuclear Magnetism Order and Disorder*, International Series of Monographs in Physics, Oxford University Press, New York, 1982.
- [13] R.R. Ernst, G. Bodenhausen, A. Wokaun, *Principles of Nuclear Magnetic Resonance in One and Two Dimensions*, Clarendon Press, Oxford, UK, 1987.
- [14] M. Goldman, *Introduction aux méthodes quantiques de calcul en RMN de haute résolution*. Centre d'études nucléaires de Saclay, Institut de recherche fondamentale, Département de physique générale, Service de physique du solide et de résonance magnétique, CEA, Palaiseau: GERM, École thématique RMN, Figeac, France, 1996.
- [15] J.-Y. Lallemand, *Spectroscopie, résonance magnétique nucléaire: Cours de majeure de chimie*, Palaiseau (France), Département de chimie, École polytechnique, 2003.
- [16] D. Neuhaus, M.P. Williamson, *The Nuclear Overhauser Effect in Structural and Conformational Analysis*, VCH Publishers Inc., New York, 1989.
- [17] E. Auguste, E. Lucas, L. Poigniez, D. Ramamurthy, H. Vaneekhouste, P. Lesot, *Spectra Anal.* 315 (2017) 32–43.
- [18] A. Louis-Joseph, *RMN: Théorie pratique, cours*, Enseignement d'approfondissement, Département de physique, École polytechnique, Palaiseau, France, 2015, pp. 1–197.
- [19] A.E. Derome, *Modern NMR Technique for Chemistry Research*, Organic Chemistry Series, vol. 6, Pergamon Press, New York, 1987, pp. 38–50.
- [20] J. Gallegos Pablo Ignacio, A. Louis-Joseph, *Pilotage programmable de spectromètre RMN. Conception expérimentale micro et nanoélectronique*, PHY573A, Rapport d'enseignement d'approfondissement, Département de physique, École polytechnique, Palaiseau, France, 2013.
- [21] A. Louis-Joseph, B. Thiébaud, *Introduction aux microprocesseurs, familles 68K/MCF5407. Électronique, composants et systèmes: enseignement expérimental, support de cours*, Département de physique, École polytechnique, Palaiseau, France, 2015, pp. 1–126.
- [22] D.I. Hoult, R.E. Richards, *J. Magn. Reson.* 24 (1976) 71–85.
- [23] D. Paget, *Phys. Rev. B* 24 (1981) 3776–3793.
- [24] A. Bienfait, L. Bonin, R. Antonelli, A. Louis-Joseph, *Instrumentation de résonance magnétique nucléaire, X2009, Rapport d'enseignement d'approfondissement*, Département de physique, École polytechnique, Palaiseau, France, 2011.
- [25] F. Lan, M. Weiss, *Résonance magnétique nucléaire: construction d'un spectromètre et application à un gaussmètre de précision*. X2011. Rapport d'enseignement d'approfondissement, Département de physique, École polytechnique, Palaiseau, France, 2013.
- [26] A. Louis-Joseph, A. Nauton A, D. Coupvent-Desgravier, J.-P. Korb, *Diffusion-fundamentals.org* 29 (2017) 1–6. Online journal.
- [27] B. Blümich, *Trends Anal. Chem.* 83 (2016) 2–11.
- [28] S.D. Riegel, G.M. Leskowitz, *Trends Anal. Chem.* 83 (2016) 27–38.