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Spatially resolved spectroscopy using tapered stripline NMR

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Magnetic field B₀ gradients are essential in modern Nuclear Magnetic Resonance spectroscopy and imaging. Although RF/B¹ gradients can be used to fulfill a similar role, this is not used in common practice because of practical limitations in the design of B¹ gradient coils. Here we present a new method to create B¹ gradients using stripline RF coils. The conductor-width of a stripline NMR chip and the strength of its radiofrequency field are correlated, so a stripline chip can be tapered to produce any arbitrary shaped B¹ field gradient. Here we show the characterization of this tapered stripline configuration and demonstrate three applications: magnetic resonance imaging on samples with nL–µL volumes, reaction monitoring of fast chemical reactions (10⁻²–10⁻¹ s) and the compensation of B₀ field gradients to obtain high-resolution spectra in inhomogeneous magnetic fields.

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1. Introduction

In general microcoils for the detection of nL–µL samples are often compromised in spectral resolution. This is due to manufacturing imperfections and very local distortions, i.e. magnetic susceptibility differences of materials [1]. In recent years, we have introduced an alternative stripline-based NMR probe as a route to high-resolution NMR on nanoliter samples [2,3].

An interesting property of striplines, which makes them especially attractive for magnetic resonance imaging (MRI) purposes, is that the local RF field strength above the stripline is directly related to the width of the stripline. This means that, by introducing a tapered modification of the stripline, well-defined arbitrary B¹ gradients are attainable, which can be used to impose a spatial coding of spins along this taper. Using a simple 2D nutation experiment, a 1D-image can be recorded with uncompromised spectral (chemical shift) information. Although MRI is traditionally based on the use of B₀ gradients instead of B¹ gradients, the use of these RF gradients has been explored before [4,5], but its implementation was difficult because of limitations in RF gradient design.

One of the proposed applications for stripline NMR is the analysis of fast chemical reaction kinetics. Using uniform B¹-field stripline NMR, we already demonstrated [6] that chemical reactions can be monitored in a time range of seconds to tens of minutes.

By adjusting the flow rates of the reactants, the delay between mixing and NMR detection can be varied, which translates into a time axis for the ongoing reaction. The main limitation in this system was the finite pathway from the microreactor to the NMR chip, which limited the minimum observable reaction time to about 4.5 s. Many chemical reactions, however, take place at sub-second time-scales, which are beyond the capability of this system.

In this manuscript, we will introduce the design and characterization of tapered striplines that aim at the generation of strong B¹ fields with arbitrary (including linear) gradients along a single axis. As a first application we use these gradients for imaging of small volume (nL–µL) samples. In a second application we use this alternative way of 1D-imaging for fast in-situ reaction screening. We demonstrate that the tapered stripline provides combined chemical structure information and spatial mapping, which allows for reaction profiling on a sub-second time-scale. As a third application we propose to use the perfect control over the shape of the tapered stripline’s B¹ profile to perform NMR experiments in inhomogeneous magnetic fields without loss of spectral information.

2. Experimental section

2.1. Chemicals and materials

Unless stated otherwise, all samples were prepared in fused silica capillaries (Polymicro Technologies) with an inner diameter (ID) of 250 µm and an outer diameter (OD) of 360 µm. FC-40
(Fluorinert), imidazole, deuterium oxide (D₂O), dimethylformamide (DMF) and phosphoryl chloride (POCl₃) were obtained from Sigma–Aldrich. Ethanol was obtained from Merck. 1-Propanol was obtained from Riedel-de Haën. All chemicals were used as received. For the reaction monitoring experiments 1 ml gas tight syringes (VWR 549-0536) were used and connected to the capillaries via a PEEK luer lock connector coupled to a one-piece finger-tight PEEK nut with a FEP sleeve between the nut and the capillary that matches either the 360 µm or 200 µm OD (IDEX Health & Science, P-659, F-120, F-242 and F-239 respectively). Connections between two or three capillaries were made using PEEK unions or T-connectors (IDEX Health & Science, P-779 and P-727 respectively). For pumping two syringe pumps (New Era, NE-1000) were used.

2.2. Tapered stripline chips

In this paper several different tapered stripline chips were used, numbered 1 to 3. Pictures (Fig. S6) and schematic cross sections (Fig. S5) of the tapered stripline chips and probes can be found in the Supplementary information. More detailed information about the fabrication of stripline chips can be found in work of Oosthoek-de Vries et al. [7].

Chip 1: For simple feasibility purposes, a 10 mm long tapered stripline (tapered from 3 to 1 mm) was fabricated using low-loss polytetrafluoroethylene (PTFE) based Printed Circuit Board (PCB, Rogers RT/duroid 5870). The PCB thickness was 0.254 mm, with a copper clad thickness of 17 µm. The stripline was structured by CNC milling with a 0.25 mm mill. Samples were positioned on top of the stripline in a second PCB from which the copper was removed at one side. For MRI testing purposes, four silicone rubber disks sample chambers were fabricated by drilling four holes with a diameter of 0.25 mm and a depth of 0.25 mm at a distance of 2 mm between each hole. The stripline was implemented as a short in a 3/4 λ resonator, resonant at 144 MHz.

Chip 2: High-resolution tapered stripline chips were fabricated on fused silica. The chip substrate was 35 mm long and 14 mm wide, the stripline constriction is 10 mm long and tapered from 5 to 1 mm in width. The structured copper stripline was defined on 500 µm thick fused silica wafers using photolithography, magnetron sputtering and electroplating [2]. To minimize susceptibility distortions, fluorinated ethylene propylene (FEP) was hot-pressed into copper-free areas on the chip. A channel for capillary that matches either the 360 µm or 200 µm OD (IDEX Health & Science, P-659, F-120, F-242 and F-239 respectively). Connections between two or three capillaries were made using PEEK unions or T-connectors (IDEX Health & Science, P-779 and P-727 respectively). For pumping two syringe pumps (New Era, NE-1000) were used.

Chip 3: a 15 mm long stripline on fused silica, tapered from 5 to 1 mm, with the same substrate dimensions as chip 2. The chip was produced with the same techniques on the same fused silica wafers as chip 2. Only the second fused silica layer was omitted, instead two Teflon spacers, located at the ends of the chip, were used to space a copper ground plate from the structured layer in the center. The stripline was implemented as a short in a 3/4 λ resonator, resonant at 600 MHz.

In all probes the tapered stripline chip is connected to a coaxial copper tube, which is part of the resonating system. The 1/4 or 3/4 λ resonator has thus roughly the same length as the probe (Fig. S6) and the length of the taper is only a small fraction of the length of the full resonator, therefore the current can be assumed to remain constant along the z-axis of the taper.

2.3. NMR experiments and data treatment

The RF field strength profile and sensitivity profile of the tapered stripline were measured using ¹H NMR at 400 MHz on a 9.4 T solid-state NMR (Agilent-Varian) system with a tapered stripline probe containing chip 3. We characterized these profiles by moving a long sample capillary (ID: 250 µm, OD: 360 µm) in steps over the taper and performing a nutation experiment after every displacement. The top part of the capillary was filled with water. Copper sulfate was added to the water to reduce the $T₁$ (to 16.4 ms) and speed up the experiment. The other part of the capillary was filled with FC-40, a proton-free liquid which is a mixture of perfluorocarbons and is immiscible with water. The capillary was closed off with wax and UV-curable glue and, it was positioned inside the tapered stripline probe, with the water/FC-40 interface on top of the upper contact pad of the stripline chip. The capillary was connected at the bottom of the probe to a translation stage of the optical head of a DVD drive. This stage is moved by a stepper motor with a resolution of 160 µm per step (in our experiment we did two steps, so 320 µm, between each nutation experiment). The stepper motor and translation stage are magnetic and were therefore mounted to a plastic frame fixed to the bottom of the probe, separating motor and stripline by 120 cm. The stepper motor was controlled with an Arduino Uno, and a Python script was used to run the stepper motor and operate the software of the NMR machine, VnmrJ 3.2A, simultaneously. In total 83 steps were made and a total distance of 26.5 mm was analyzed. Processing of the nutation experiments and all other NMR experiments in this article were done using the Matlab based processing package matNMR [8]. For the nutation experiments the following parameters were used: 8 kHz spectral width, 160 ms acquisition time, 30 ms recycle delay, number of transients equal to 2 and a pulse width of 2.2 µs or multiples thereof, with a total of 320 increments.

The static imaging measurements were performed using a nutation experiment. Four silicone disks were measured using ¹H NMR at 144 MHz with tapered stripline chip 1 on a 3.4 T solid-state NMR (Oxford ultra-wide bore magnet, Varian Infinity Plus) system operated by Spinsight software. For the nutation experiment the following parameters were used: 40 kHz spectral width, 25 ms acquisition time, 3 s recycle delay, 4 scans per increment and a pulse width of 1.5 µs or multiples thereof, with a total of 300 increments.

The static measurement of the three liquid plugs was conducted by ¹H NMR at 600 MHz using tapered stripline chip 2 on a 14.1 T solid-state NMR (Oxford 600/89 magnet, Varian VnmrJ) system. For the nutation experiments the following parameters were used: 16 kHz spectral width, 1 s acquisition time, 9.5 s recycle delay, 16 scans per increment and a pulse width of 2.5 µs or multiples thereof, with a total of 300 increments.

The 1D nutation measurements on the three silicone rubber blocks were done with the same chip and NMR system. For the standard 2D arrayed nutation experiment the following parameters were used: 100 kHz spectral width, 50 ms acquisition time, 4 s recycle delay, 4 scans per increments and a pulse width of 2 µs or multiples thereof, with a total of 360 increments. For the 1D nutation experiment the following parameters were used: 25 kHz spectral width, 10 s recycle delay, 16 scans per increments, 40 µs per cycle, a pulse width of 2 µs and a total number of 360 acquired points.

For the experimental determination of the spatial resolution chip 3 was used on the 400 MHz Varian-Agilent system. A 10 cm long capillary was filled with H₂O and closed off on both sides with wax. The following parameters were used for the nutation experiment: 5 kHz spectral width, 1 s acquisition time, 2 s recycle delay, 1 scan per increment and a pulse width of 2.5 µs or multiples thereof, with a total of 1600 increments. During the entire experiment a B₀ gradient of 12.5 G/m was applied using the Z1 shim coil. The spatial resolution was determined by averaging the FWHM of the peak in the indirect dimension at 33 positions of the taper.
The reaction monitoring measurements were performed using $^1$H NMR at 600 MHz with tapered stripline chip 2 on the 14.1 T solid-state NMR system. The liquids were pumped through the microfluidic system using gas-tight syringes and syringe pumps. The pumps were inside a fume hood and connected with 6 m long capillaries (250 μm ID) to union connectors on the bottom of the probe. From there on one liquid was directed through a 250 μm ID capillary and the other liquid through a 100 μm ID capillary. Both capillaries were connected to a T-connector on top of the probe. The 100 μm ID capillary was led all the way through the connector and into the outlet capillary (250 μm ID). Hereby we created a system in which the inner capillary contains liquid A and between the inner and the outer capillary there is liquid B. The end of the inner capillary was modified by Reith Laser (Wijchen, the Netherlands). The idea was to create a kind of micronozzle (Fig. S4), which should improve mixing, however we were only able to reduce the mixing time to a third of a second and besides that the mixer also caused some flow instability, for instance because gas bubbles built up inside the nozzle. The outlet capillary was positioned over the tapered stripline and exited the probe at the lower end (total length 1 m), were the reaction mixture was collected in a glass flask and the reaction was quenched with a sodium acetate solution in water. The micronozzle was positioned exactly on top of the narrowest side of the taper (Fig. S3) to be able to measure the mixture really from the start of the reaction.

The flow rates for DMF and POCl$_3$ were 9.0 and 3.0 μL/min respectively. Before the measurement started the system was allowed to stabilize for half an hour. A back pressure regulator was also tested to see if it stabilized the system, however, no positive effect was observed, instead the syringe pumps stalled more often and therefore the regulator was removed. The following parameters were used for the nutation experiment: 8 kHz spectral width, 1.5 s acquisition time, 3 s recycle delay, 1 scan per increment and a pulse width of 2 μs or multiples thereof, with a total of 256 increments.

The $B_0$ gradient compensation experiments were conducted on the 400 MHz NMR system, running $^1$H NMR experiments in tapered stripline chip 3. The sample was a capillary filled with 1-propanol and a $B_0$ gradient was created by lowering the probe into the stray field of the NMR magnet. A pulse sequence based on the sequence with consecutive adiabatic full passages, described by Meriles et al. [9], was used. The adiabatic passages had a linear sequence with consecutive adiabatic full passages, described by the stray field of the NMR magnet. A pulse sequence based on the probe. From there on one liquid was directed through a 250 μm ID capillary and into the outlet capillary (250 μm ID). Hereby we created a system in which the inner capillary contains liquid A and between the inner and the outer capillary there is liquid B. The end of the inner capillary was modified by Reith Laser (Wijchen, the Netherlands). The idea was to create a kind of micronozzle (Fig. S4), which should improve mixing, however we were only able to reduce the mixing time to a third of a second and besides that the mixer also caused some flow instability, for instance because gas bubbles built up inside the nozzle. The outlet capillary was positioned over the tapered stripline and exited the probe at the lower end (total length 1 m), were the reaction mixture was collected in a glass flask and the reaction was quenched with a sodium acetate solution in water. The micronozzle was positioned exactly on top of the narrowest side of the taper (Fig. S3) to be able to measure the mixture really from the start of the reaction.

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### 3. Tapered stripline design

A typical configuration for a tapered stripline consists of five layers (Fig. 1): a central copper layer is electrolytically deposited on a fused silica substrate, which acts as an electrical insulator, these two layers together are the stripline chip. The chip is sandwiched between two copper ground planes. The bottom one is in direct contact with the fused silica, however, the top ground plane is separated from the central copper layer with Teflon spacers to ensure electrical insulation and to create space for the sample. The central copper layer is divided in three parts. The middle part carries the current and generates the RF field, its width is varied over the length of the taper. The outer copper parts are not part of the electric circuit, but are in place for susceptibility matching.

The spectral resolution performance of striplines of constant width has been studied before [2,3]. The arguments presented there also apply to the present case. Because the plane in which the copper stripline is defined is extended with the copper outer parts one can approximate the $B_0$ field distortion induced by this plane using simple arguments based on Maxwell’s equations. If the plane is oriented parallel to the field and if the contact pads are sufficiently long then the field will be nearly homogeneous along the sample volume probed by the stripline. There will be a small $B_0$ field offset, but in essence the gradient can be negligibly small and the resolution in the spatially resolved spectra remains uncompromised. In practice the main field distortion will be due to finite sample lengths and liquid or solid susceptibility plugs are needed.

The $B_0$ field strength on top of a stripline carrying a current $I$ can be approximated using Ampère’s circuital law:

$$\oint_C Bds = \mu_0 I - B_0 \equiv \frac{\mu_0 I}{2W + h}$$

where $C$ is a closed rectangular path (dimensions: $w \times h/2$) around the central strip of the stripline, $s$ is the infinitesimal in $C$, $\mu_0$ is the magnetic permeability and $w$ and $h$ are the width of the confined part and height of the stripline chip, respectively. In this approximation we assumed that the $B_0$ field is parallel to the surface of the strip and nearly constant.

For imaging purposes, $w$ has to be a function of the spatial coordinate, $z$, along the axis of the stripline. We can simplify...
Eq. (1) further by eliminating the constants $a_0$ and $I$. Also, there is a minimum $B_1$ field where the taper is at its widest:

$$b(z) = \frac{1}{2(w(z) + d)} \quad \text{(2)}$$

$$b_{\text{min}} = \frac{1}{2(w_{\text{max}} + d)} \quad \text{(3)}$$

where $d$ is equal to $h/2$. In order to obtain a linear RF gradient over the stripline, $b/b_{\text{min}}$ must be a linear function of $z$:

$$b(z) = \frac{w_{\text{max}} + d}{w(z) + d} = zG + 1 \quad \text{(4)}$$

where $G$ is the strength of the $B_1$ gradient. So it follows that for a stripline width profile:

$$w(z) = \frac{w_{\text{max}} + d}{zG + 1} - d \quad \text{(5)}$$

a normalized linear gradient in $B_1$ of strength $G$ is generated. Eq. 2 shows plots of $B_1$ and $w$ as functions of $z$. All tapered striplines in this manuscript were fabricated with the taper shape described by Eq. (5) (tapered from 5 to 1 mm). Finite Element Method Magnetics (FEMM, a Matlab toolbox) was used to model a cross section of this tapered stripline and to calculate an approximate 2D RF field profile between the ground planes. This was done for 50 positions along the taper and these 2D RF field profiles were joined to produce a 3D profile. A current of 20 A was used for the central strip and the ground planes were set to $-10$ A.

The cross-sectional profile, Fig. 3a, shows large field distortions at the edges of the central strip, however, the $B_1$ field between the center of the central strip and the center of the ground planes is nearly homogeneous. For narrower sections of the tapered stripline an RF gradient will be present perpendicular to the strip. For those sections the plane of constant $B_1$ field will be curved, since there is both a gradient parallel and perpendicular to the strip. However, for wider sections of the tapered stripline this perpendicular gradient is negligible in size.

In Fig. 3b a cross section is shown parallel to the strip at half the height between the central strip and the ground plane. The figure shows that the tapered stripline generates a $B_1$ field that is homogeneous perpendicular to the axis of the taper and only deviates at the edges. For this slice the gradient field was calculated, Fig. 3c, and shown to be constant over the entire length of the taper. Again only the edges show a deviation from the tapers RF gradient strength.

5. Experimental verification of the RF profile

We characterized the RF field profile experimentally by displacing a water/FC-40 interface stepwise over the taper and performing a nutation experiment every 320 $\mu$m. At zero displacement only FC-40, which is a proton-free perfluorocarbon mixture, was on top of the taper and the water was on top of one of the broad connection pads on the chip. As the water/FC-40 interface was moved over the chip, a larger section of the taper was covered by water. Each nutation experiment then gave an RF profile, which tells us what RF field strengths were felt by the water sample. By subtracting each RF profile from the preceding one we can determine the RF field for a 320 $\mu$m long section of the tapered stripline. In total 83 displacements were executed, giving the full RF field profile of the tapered stripline (Fig. 4). As expected the tapered stripline gives a nearly perfect linear RF field profile. The field has a steep transition (at 7 mm displacement) going from the broad contact pads of the chip to the narrowest side of the taper and the field at the contact pads is, with 18 kHz, relatively low.

Fig. 2. Plots of the stripline width $w$ (top) and the corresponding linear $B_1$ field (bottom) as a function of the position. $G$ is chosen such that the $w_{\text{max}}/w_{\text{min}} = 5$ for a 10 mm long taper, $h = 800 \mu$m, $I = 1$ A.

Fig. 3. (a) $B_1$ field of a cross section of the tapered stripline, taken at the middle of the taper. (b) $B_1$ field halfway between the center strip and the top ground plane. (c) $B_1$ gradient strength at the same height as figure (b).
we prefer the longer method, in which no assumptions have to be made.

The same tapered stripline chip was used to determine the maximum gradient strength. The proton amplifier was set to maximum power (450 W) and a nutation experiment was performed. A maximum RF field of 630 kHz was observed with a field difference of roughly 350 kHz over the 15 mm length of the taper, which is equivalent to a gradient strength of 55 G/cm. This is on the same order as the \( B_0 \) gradients in standard liquid-state probes, which range from 50 to 100 G/cm. However, nowadays specialized diffusion probes are commercially available with pulsed gradients that are an order of magnitude stronger (1000–3000 G/cm). Nevertheless, 55 G/cm is a decent gradient strength and some small modifications to the chip, for instance a reduction in the length of the taper, should allow us to reach \( B_1 \) field gradients of more than 100 G/cm.

6. Imaging of static samples

The RF gradient of the tapered stripline can be used for static imaging experiments. The spatial information is encoded during the excitation pulse as an RF-dependent phase. The RF frequencies can be determined from a nutation experiment. After a Fourier transform in both the direct and indirect dimension the data obtained are a 1D-image in the \( F_1 \) dimension, whereas the spectral information is contained in the \( F_2 \) dimension. A more extensive description of this \( B_1 \) imaging method and the processing is given in the Supplementary information.

The imaging capability of the tapered stripline chip was first tested on a set of disks. Four silicone rubber disks with a diameter of 250 \( \mu \)m were placed 2 mm apart in a straight line along the axis of the tapered stripline and a nutation experiment was performed. The image (Fig. 5) clearly visualizes the capability of our method to map the distinct positions of the silicone disks. The peaks in the skyline plot (\( F_1 \) dimension) are 3.6 kHz wide, corresponding to a spatial width of 273 \( \mu \)m, which is in good agreement with the diameter (250 \( \mu \)m) of the silicone rubber disks. In the \( F_2 \) dimension we observe one narrow peak from the rapidly rotating methyl group in the silicone rubber.

From the same data we could extract the sensitivity profile of the tapered stripline (Fig. 4c) by integrating the peak in the direction of the RF field strength. This first resulted in a modulated profile, because the steps of the stepper motor were not exactly equal in size. However, they were periodic, so averaging of the neighboring data points resulted in a smooth profile. As expected from the reciprocity theorem, the sensitivity profile of the tapered stripline is equal to the RF field profile. This means that for quantitative measurements the results have to be corrected with the experimental profile of Fig. 4c, or one can also simply make use of an internal standard. The sensitivity profile deviates from the RF profile at very low and high displacements. The deviation at low displacements can be explained by the fact that after connecting the sample capillary to the stepper motor, the capillary is not yet under tension and the water/FC-40 interface does not move. The RF profile measured in the consecutive nutation experiments is thus the same. Subtracting the profiles results therefore in zero intensity.

The RF field profile can also be determined with the help of a static linear \( B_0 \) gradient, where the resonance frequency encodes for position [11]. Then in a single nutation experiment the full RF field profile can be obtained. However, in that case the initial shimming, the linearity of the applied \( B_0 \) field and uniformity of the sample all have to be perfect. Because the measurement of the RF profile only has to be done once to verify the design concepts,
In general a single nutation experiment in combination with the tapered stripline gives a 1D-image of the sample, but it also contains all the chemical shift information, thus it gives both spatial and spectral information. To demonstrate that the chemical shift information is preserved, another measurement was performed. A capillary was filled with three plugs of different compounds, each separated by FC-40, as displayed in Fig. 6. From these data, the position as well as the chemical composition of the sample plugs could be deduced straightforwardly. The spectra that were taken from the image show that the spectral resolution is good.

If one is only interested in the spatial information and not in the spectral information, then the nutation experiment can be performed in a single 1D experiment [12,13] instead of a 2D array. A train of equal pulses is given and the acquisition is done by sampling a single data point between each pulse pair. The resulting FID gives a nutation spectrum, and thus a 1D image, after a single Fourier transformation. With this method one does not have to run an array and can reduce the experimental time by a factor of several hundred (depending on the number of points and the desired signal to noise ratio). We tested the method on three silicone rubber blocks (Fig. 7) and found minimal differences between the results of the standard 2D arrayed nutation and the 1D nutation.

7. Spatial resolution

The theoretical spatial resolution of this imaging method is – analogous to the resolution in the spectral dimension – determined by the length of the recorded signal. In the spatial dimension this is equivalent to the longest pulse in the nutation experiment, which is limited by the bandwidth that has to be excited. For example, if protons are observed at 600 MHz in a bandwidth of 10 ppm (6 kHz), then the maximum pulse width allowable is 1/(6 kHz) = 167 μs. This limit on the resolution arises from the need to have a pulse that is uniformly effective over the entire range of Larmor frequencies of the sample. If the maximum pulse width is 167 μs,
then the resolution in the indirect dimension is $1/(167 \mu s) = 6$ kHz. Since the shape of the RF field gradient is linear, the resolution in the indirect dimension ($R_{\text{indirect}}$) can be directly converted by Eq. (6) into the resolution of the spatial dimension ($R_{\text{spatial}}$) if the length of the taper ($L_{\text{taper}}$) and the RF field at the narrowest ($RF_{\text{max}}$) and widest ($RF_{\text{min}}$) part of the taper, or the RF gradient ($G_{RF}$) are known.

$$R_{\text{spatial}} = \frac{L_{\text{taper}} \cdot R_{\text{indirect}}}{RF_{\text{max}} - RF_{\text{min}}} = \frac{R_{\text{indirect}}}{G_{RF}}$$

Eq. (6) shows that the spatial resolution can be optimized by matching the length of the taper to the length of the sample and by maximizing the RF gradient. The latter can be achieved by moving to higher RF pulse powers, which is again limited by the bandwidth. Instead it is limited by the practical limit of the pulse length (to prevent RF heating and arcing) and the sensitivity of the probe. The voxel should be large enough so that it contains at least the number of spins necessary to give a detectable signal in a reasonable amount of time. For example, for a $^{1}H$ image of water in a capillary ($250 \mu m$ ID), recorded with a receiver bandwidth of $5$ kHz, a S/N of 5 in the FID (single scan) and a sensitivity of $10^{14}$ spins/$\sqrt{Hz}$ the spatial resolution is $10$ $\mu m$.

8. Reaction monitoring

Bart et al. [2] showed that the stripline design can be used to monitor reactions in a flow NMR setup. Many other flow NMR systems have been described in literature. Most of them use high magnetic fields and only make minor adjustments to the standard liquid-state NMR probes [14–17]. In recent years several excellent benchtop NMR machines have entered the market, with very homogeneous fields that are stable over longer periods. This has literally brought the NMR machine inside the fume hood and has already given several good examples of low-field NMR reaction monitoring [18–21]. However, there are few examples of flow NMR systems [22] in which the hardware is adapted to very low detection volumes (on the order of a few microliters down to several picoliters) that match those of a microfluidic system. Microfluidic systems are used in microreactor synthesis, the production of microdroplets for biochemistry and several other fields. They offer many advantages over traditional systems [21–26] including perfect control over temperature and reaction time, less solvent and good safety control. So they make it possible to study more violent and fast reactions.

The next challenge in flow NMR is to be able to monitor fast chemical reactions. Reactions of a few seconds or less are difficult to monitor with high spectral resolution because the acquisition length is on the same time scale, although ultrafast 2D NMR techniques [27, 28] have made it possible to obtain more information on this short time scale. Up to now, sub-second reactions can only be monitored with stopped-flow and rapid injection NMR techniques [29–31], however, these methods are laborious. Our tapered stripline setup is optimized for the small sample volumes of microfluidic systems and by applying the nutation sequence to a continuous flowing reaction mixture (with a fixed flow rate) we should be able to produce a kinetic profile of the reaction, in the same way as we would get a one-dimensional image of a static sample.

We tested our method on the first step of the Vilsmeier–Haack reaction, which was discovered in 1927 as a new way to functionalize an electron-rich aromatic ring with an aldehyde [32]. This so-called formylation reaction consists of three steps: the formation of the extremely reactive iminium ion, also called the Vilsmeier reagent; the reaction of an electron-rich arene with this reagent...
and an aqueous work-up step (Fig. 8). In the first step N,N-dimethylformamide (DMF) reacts with phosphoryl chloride (POCl₃). Via several intermediates the oxygen atom of the DMF gets exchanged for a chlorine atom and an iminium ion ($R_2C(=N)R_3^+$) is formed. Note that the $C(=O)$–$N$ bond has a double bond character [33], therefore, the DMF and all intermediates can have two peaks for the methyl groups.

The reaction, in particular the first step, is very fast and exothermic and serious thermal hazards have been reported [34]. The safety risks can be brought to a minimum by using a microfluidic flow setup instead of the standard reaction flask.

9. Experimental results reaction monitoring

The reaction monitoring setup described in the experimental section was used to study the first step in the Vilsmeier–Haack reaction: the formation of the iminium ion. It was already known from literature that this step would be fast (completed within 90 s in a solution of acetonitrile [35]). Using our method we monitored the first 2.5 s of the reaction of the pure compounds.

A problem we encountered was the mixing of the liquids. At the small length scales in the capillary and the slow flow rates the flow is no longer turbulent, instead there is a laminar flow. So mixing occurs only on the basis of diffusion, which is too slow to study the fast reaction kinetics. We tried to overcome this problem by using a micro nozzle, as described in the experimental section. This did improve mixing, but not enough to obtain fully quantitative kinetic information.

Nevertheless, this method can still give extremely valuable information about the reaction mechanism. The spectra of the Vilsmeier–Haack reaction, Fig. 9, give detailed information of the first 1.5 s of the reaction. At $t = 0$ s three peaks are observed, corresponding to the methyl groups (2.88 and 3.05 ppm) and $N$–$H$ group (8.13 ppm) of DMF. These lines quickly broaden in an asymmetric way in the upfield direction and keep this shape for 20 s (as was determined with the method of Oosthoek-de Vries et al. [71]). This can be explained partly by incomplete mixing, but most likely it indicates the dynamic formation of one of the intermediates. Additional broadening may be caused by intramolecular chemical exchange, namely the hindered rotation around the $C(=O)$–$N$ or $C(=Cl)$–$N$ bonds, and by intermolecular chemical exchange (for instance the addition and reduction of a chlorine atom to the positively charged intermediate).

After a few hundred milliseconds peaks emerge (3.38, 3.54 and 9.46 ppm) of what we assume to be the Vilsmeier reagent. Remarkably these peaks also have an asymmetric shape, only with a broadening in the downfield direction.

From the 2D matrix we integrated the DMF peaks and the peaks of the Vilsmeier reagent and the possible intermediates. The signal intensity along the taper is scaled according to the sensitivity profile of the tapered stripline (Fig. 4), therefore the conversion (Fig. 10) was calculated by dividing the signal of the Vilsmeier reagent and the intermediates by the integral of the full spectrum. The reaction does not reach 100% conversion because an excess of DMF (3:1 v/v) was used to keep the viscosity of the reaction mixture low, so the curve levels off to the theoretical maximum conversion at 27.7%.

Our reaction monitoring method shows some very interesting changes in the reaction mixture of the Vilsmeier–Haack reaction at short reaction times on the order of several hundreds of milliseconds. However, slower processes are also present and after a long reaction time the NMR spectrum shows more changes, i.e. narrowing of the methyl peaks and formation of several side products. The Vilsmeier–Haack reaction is clearly a very complex reaction and more research has to be done to fully understand the mechanism and to assign all the peaks. In the present context it is not possible to give a full, detailed description of this reaction. However, these preliminary results underline the potential of our method to study chemical processes at a time scale of tens of milliseconds up to several seconds.

10. B₀ gradient compensation

As a third application example for the tapered stripline approach we will discuss the possibility for arbitrary $B₀$ gradient compensation.

Between 2000 and 2005 the Pines group published a series of papers [9,36–39] in which they describe how $B₁$ field gradients can be used to retrieve spectral resolution in inhomogeneous magnetic ($B₀$) fields. Their methods can be used in portable single-sided NMR systems or in other systems where perfect $B₀$ homogeneity is impossible to realize using standard shimming techniques.

Fig. 9. Spectra of the Vilsmeier–Haack reaction, showing the methyl region at different reaction times. The most upfield peak at 0 s was set to 2.88 ppm (the literature value for DMF in CDCl₃). The spectra are corrected for the sensitivity profile of the tapered stripline.

Fig. 10. Conversion of dimethylformamide in the first step of the Vilsmeier–Haack reaction. The flow rates are 9 μL/min for DMF and 3 μL/min for POCl₃. The conversion was calculated from a nutation measurement with 256 increments (1 scan each), acquired in a total time of 20 min.
In an inhomogeneous field a phase shift of the local magnetization is needed to compensate for the difference in Larmor precession due to the $B_0$ variations over the sample. The Pines group first used composite z-rotation pulses to create this position dependent phase shift [36,37], but later they switched to consecutive adiabatic passages [9,38,39], because these are more effective over a large bandwidth. Since our tapered stripline produces a linear $B_0$ field gradient, any $B_0$ field profile can be compensated for.

The pulse sequence (Fig. 11) we used is very similar to the one used by the Pines group. An adiabatic half passage is used to create transverse magnetization. Then a windowed acquisition is started in which between every two acquisition points a set of two adiabatic full passages are given. The amplitude of the second adiabatic passage is optimized such that phase produced by the double passage matches the phase introduced by the $B_0$ field gradient. (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)

Under a small $B_0$ gradient of 13 G/m the spectrum looks very crowded, however, the compensation pulse sequence is able to fully recover the spectral resolution. At a larger gradient of 172 G/m the lines in the standard 1D spectrum are broadened so much that they fully overlap and the resulting spectrum consists of only a single broad peak. With the compensation pulse sequence it is still possible to recover most of the spectral information and to distinguish the five peaks. However, the peaks are broadened and

11. Experimental results $B_0$ gradient compensation

The $B_0$ gradient compensation pulse sequence was tested at three positions in the magnet: the homogeneous center, a slightly inhomogeneous part (13 G/m) and deeper in the stray field of the magnet (172 G/m). Fig. 12 displays the spectra obtained at these three positions with both a single 90° pulse and with the $B_0$ gradient compensation pulse sequence. Clear differences are visible in the spectra acquired in the homogeneous center of the magnet. The apparent J-coupling is larger in the spectrum obtained with the compensation pulse sequence. In fact the J-coupling has remained the same, but the chemical shift is scaled. In this pulse sequence the chemical shift only evolves during the time between the sets of adiabatic full passages. The J-coupling, on the other hand, is unaffected by the adiabatic sweeps and is active during the entire windowed acquisition. For the measurements in Fig. 12 the compensation sequence was run with 100 μs long adiabatic full passages and a point-by-point acquisition of 1 point per 250 μs. So the chemical shift evolves only 50 μs between each point and the chemical shift is thus scaled by a factor 5.

Under a small $B_0$ gradient of 13 G/m the spectrum looks very crowded, however, the compensation pulse sequence is able to fully recover the spectral resolution. At a larger gradient of 172 G/m the lines in the standard 1D spectrum are broadened so much that they fully overlap and the resulting spectrum consists of only a single broad peak. With the compensation pulse sequence it is still possible to recover most of the spectral information and to distinguish the five peaks. However, the peaks are broadened and

![Fig. 11. Pulse sequence for obtaining $B_0$ gradient compensated spectra: after an adiabatic half passage (gray) a windowed acquisition is started. Between the acquisition points (black dots) sets of two adiabatic full passages (yellow) are given. The amplitude of the second adiabatic passage is optimized such that phase produced by the double passage matches the phase introduced by the $B_0$ field gradient. (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)](image)

![Fig. 12. Spectra of 1-propanol. The top spectra were acquired in the homogeneous part of the magnet. The middle and bottom spectra were acquired in the stray field of the magnet. The spectra on the left are simple 1D spectra, acquired after a single 90° pulse. The spectra on the right are acquired with the $B_0$ compensation sequence.](image)
the triplets and multiplet are no longer visible. This can be explained by the shape of the stray field of the magnet, which is used to create the magnetic field gradient. For a small section of the stray field the gradient is approximately linear, but as the gradient becomes larger this approximation no longer holds. In that case our tapered stripline, with its linear gradient, does not perfectly match the $B_0$ field gradient anymore.

12. Conclusions and outlook

The constriction in a stripline NMR detector can be shaped such that a linear $B_1$ gradient is obtained. This tapered stripline can be used for 1D imaging and may offer several advantages over imaging with $B_0$ gradients. Furthermore in this approach the chemical shift information is completely preserved. We have shown the applicability of the tapered stripline by monitoring the first 2.5 s of the Vilsmeier–Haack reaction in a microfluidic tapered stripline chip. Hereby we demonstrated that it is possible to study reactions by NMR on a time scale on the order of hundreds of milliseconds up to several seconds. The combination of flow chemistry and tapered stripline NMR is in principle a very promising one. Flow chemistry allows one to perform reactions in a very controlled manner and with perfect control over timing and temperature, bringing the exploration of more exotic reactions, which were previously regarded as too dangerous or difficult to perform, within reach. Stripline NMR offers all the advantages of NMR and sensitivity-wise the stripline design is the preferred choice to match the nL–µL sample volumes in microfluidics. Future research should be aimed at a better integration of the microfluidic system into the probe, and at improved mixing to obtain quantitative data to allow analysis of the pure reaction kinetics.

As a final application we showed that the tapered stripline can be used to compensate $B_0$ gradients and retrieve the spectral resolution obtained in homogeneous fields. In principle the shape of the tapered stripline can be modified to create any desired RF field profile. So one could use a tapered stripline with a linear gradient to map the $B_0$ field profile, and from that profile design a stripline chip with a $B_1$ field profile that matches the $B_0$ field profile. This stripline can then be used to obtain high resolution spectra in inhomogeneous $B_0$ fields.

Another obvious extension of the tapered stripline is the measurement of spatial diffusion in microfluidic systems. Work in this direction is ongoing.

Because stripline chips are flat and easily scalable, it is possible to apply them in benchtop spectrometers or even smaller miniaturized NMR systems [40,41]. These systems are relatively sensitive and low in cost, which makes them ideal for medical and industrial applications.

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Appendix A. Supplementary data

Supplementary data associated with this article can be found, in the online version, at http://dx.doi.org/10.1016/j.jmr.2015.12.021.

References


