A Noninvasive Platform for Imaging and Quantifying Oil Storage in Submillimeter Tobacco Seed

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While often thought of as a smoking drug, tobacco (Nicotiana spp.) is now considered as a plant of choice for molecular farming and biofuel production. Here, we describe a noninvasive means of deriving both the distribution of lipid and the microtopology of the submillimeter tobacco seed, founded on nuclear magnetic resonance (NMR) technology. Our platform enables counting of seeds inside the intact tobacco capsule to measure seed sizes, to model the seed interior in three dimensions, to quantify the lipid content, and to visualize lipid gradients. Hundreds of seeds can be simultaneously imaged at an isotropic resolution of 25 μm, sufficient to assess each individual seed. The relative contributions of the embryo and the endosperm to both seed size and total lipid content could be assessed. The extension of the platform to a range of wild and cultivated Nicotiana species demonstrated certain evolutionary trends in both seed topology and pattern of lipid storage. The NMR analysis of transgenic tobacco plants with seed-specific ectopic expression of the plastidial phosphoenolpyruvate/phosphate translocator, displayed a trade off between seed size and oil concentration. The NMR-based assay of seed lipid content and topology has a number of potential applications, in particular providing a means to test and optimize transgenic strategies aimed at the manipulation of seed size, seed number, and lipid content in tobacco and other species with submillimeter seeds.

The use of tobacco (Nicotiana spp.) dates back to the Mayan civilization (Zagorevski and Loughmiller-Newman, 2012) and in modern times still retains a primary place, as measured by monetary value, among nonfood crop species. Given the well-documented health risks associated with smoking tobacco, finding alternative uses for the crop represents an active area of research. The ease of its genetic transformation (the species was the first plant to be successfully transformed some 30 years ago by Zambrsly et al. [1983]) has prompted a suggestion that it might be used as a vehicle for the synthesis of high-value pharmaceutical compounds (Borisjuk et al., 1999; Davoodi-Semiromi et al., 2009; Morandini et al., 2011) and enzymes (Verma et al., 2010; Agrawal et al., 2011). Its potential biomass production can reach 170 tons ha⁻¹ per year in some commercial varieties (Schillberg et al., 2003), a level that is attractive in the context of biofuel production (Usta, 2005; Andrianov et al., 2010).

Tobacco seed is conventionally considered to have no commercial use, but a production level of up to approximately 1.2 tons ha⁻¹ has been documented in the literature (Patel et al., 1998). The seed contains as much as 40% lipid (Frega et al., 1991), consisting mainly of triacylglycerols with a predominant contribution, about 75%, of linoleic acid. Tobacco seed lipid can be blended with mineral diesel fuel without any need for engine modification or fuel preheating (Usta, 2005). The conversion of tobacco into an energy crop can take advantage of a well-developed biotechnology infrastructure (Gadani et al., 2003). Molecular markers have been applied to characterize the genetic diversity present in breeding germplasm (Ren and Timko, 2001; Pricano et al., 2012). Furthermore, a dense marker-based genetic map has been derived (Bindler et al., 2011), and substantial volumes of DNA sequences have been deposited in the public domain (www.pmgg.org/tgi/).

Studying the biology of lipid deposition in tobacco seed (tissue specificity, seed size, and architecture) is hampered by the small size of the seed (typically <1 mm in diameter; Fig. 1A) and the large number of

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Figure 1. Variability in seed lipid content as estimated using TD-NMR in a range of *Nicotiana* species. A, Typical seed size of (from left to right) oat, soybean, oilseed rape, and tobacco. B, The geographical provenance of wild (red) and cultivated (green) *Nicotiana* spp. accessions. C, Seed lipid content expressed as a percentage of seed dry weight (DW).
seeds formed within each capsule. Preexisting approaches
to characterize the internal structure of seeds and to
quantify the lipid fraction within them (Borisjuk et al.,
2011, 2012; Horn et al., 2011a, 2012) are not readily
applicable to small seeds. To the best of our know-
ledge, there is no method available for the noninvasive
lipid imaging in submillimeter oilseeds as those of the
genus \textit{Nicotiana}.

Our work describes how NMR can be exploited to
provide a noninvasive platform for the in vivo analysis
of tobacco seed. The method allows for both the
characterization of seed architecture and the assess-
ment of tissue-specific lipid storage capacity without
any seed destruction. The simultaneous imaging of
some hundreds of seeds can be achieved at a level of
resolution sufficient to assess each seed individually.
The method has been applied to investigate lipid de-
position in the endosperm and embryo of the seed of
various \textit{Nicotiana} species as well as transgenic tobacco
plants, thereby uncovering a previously unrecognized
trade off between seed growth and lipid storage.

\section*{RESULTS}

\subsection*{NMR Analysis Suggests That the Seed of Cultivated
Tobacco Is More Lipid Rich Than That of Its
Wild Relatives}

To assess the diversity of lipid content within the
seed of \textit{Nicotiana} species, a set of 64 accessions held at
the German Federal ex situ GenBank at Leibniz-Institut
für Pflanzengenetik und Kulturpflanzenforschung was
surveyed. The germplasm sample comprised at least
three accessions of each of the 11 species and in 10
cultivars of \textit{Nicotiana tabacum}, which originate from
Europe, Asia, Africa, Australia, and the Americas (Fig.
1B; Supplemental Table S1). Seed lipid content was
determined by applying low-field time domain NMR
(TD-NMR), a high-throughput platform that avoids the
need for prior extraction procedures. All of the seed
samples were heavily pigmented, but they varied in
shape from nearly globular to angular, in their surface
patterning and in their weight (by up to 3-fold). The
range in lipid concentration varied by around 6% dry
weight, with the lowest level found in the seed of \textit{Nicotiana plumbaginifolia} and the highest in \textit{N. tabacum}.
Overall, the seed lipid content of the cultivated species
was higher than that of the wild species (Fig. 1C).

\subsection*{Magnetic Resonance Imaging Allows Structural and
Chemical Imaging of the Intact Capsule}

The conventional means of characterizing the inter-
 internal structure of the capsule (Fig. 2A) usually requires
its prior dissection. By using magnetic resonance imaging
(MRI), the destruction of the capsule can be avoided.
Furthermore, due to the different frequencies of the
proton signals, images from water only and lipid only
only can be acquired. Suppressing one of the signals prior to

\begin{figure}[h]
\centering
\includegraphics[width=0.8\textwidth]{figure2.png}
\caption{The internal structure of the tobacco capsule. A, Transverse
section of the central portion, showing the arrangement of vascular
bundles (hand-dissected capsule). B and C, MRI images showing the
distribution of water (B) and lipids (C). cb, Central vascular bundle; pb,
placental bundle; pls, placental surface; ps, placental stalk; S, seed; ch,
chamber; vb, vascular bundle. See also Supplemental Movie S1.}
\end{figure}
isotropic resolution (Fig. 2C). The signal was mostly concentrated on the internal surface of the capsule, corresponding to the positions of the individual seeds (seen as rainbow-colored dots). An animated scan (Supplemental Movie S1) of an intact capsule illustrates its internal structure, in particular demonstrating the distribution of water (on the right) and the sites of lipid deposition (on the left). As the same capsule provided both sets of data, a virtual reconstruction of the capsule was straightforward to generate. The two-dimensional images shown in Figure 3 illustrate the architecture of the capsule during the main storage phase. The images integrating the water (colored cyan) and lipid (red) status (Fig. 3, B and C) indicate that the most heavily hydrated tissues were the placental vasculature and the regions in contact with the seeds, rather than the placenta itself. The thin water-rich layer inside the seed is the testa (Fig. 3D).

Three-Dimensional Modeling of the Capsule and the Estimation of the Seed Number

Our NMR platform also allows to image the intact capsule three dimensionally and to obtain a count of the seed. This was made possible by employing spin-echo imaging based solely on global chemical shift selective pulses. The details of this method are described in “Materials and Methods.” A resulting three-dimensional reconstruction of the capsule is represented in Figure 4 (see also Supplemental Movie S2). The hydration level of the maternal tissues of the capsule was rather low, especially in the pericarp; the most hydrated portions of the capsule were the vascular bundles, the placental stalk, and the placental bundles. As individual seeds could be readily identified, the number of seeds within the imaged capsule was estimated without a destructive assay. Specifically, the seed count was achieved by dividing the combined volume of lipid filled pixels (derived from the lipid image) by the mean volume of each seed (determined from five manually segmented seeds). The resulting seed number per capsule was 413 ± 37.

Microimaging of Lipids within the Embryos and Endosperms of Intact Seeds

The testa of the mature tobacco seed normally renders the endosperm and embryo inaccessible except by destructive assay (Fig. 5A). High-resolution three-dimensional imaging and modeling of the mature tobacco seed was achieved by using high-field MRI (17.6 Tesla) in combination with a custom-built 2-mm-i.d. solenoid coil as a detector. This procedure compromised neither the subsequent germination performance nor the plant growth (data not shown). A standard three-dimensional spin-echo sequence was used to acquire a three-dimensional image of the lipid distribution (see “Materials and Methods”) within the seed, followed by image segmentation in order to relate the distribution of the NMR signal to individual organs (Melkus et al., 2009). The model shown in Figure 5B, which features both the endosperm and the embryo, allowed their respective volumes to be estimated. This allows accurate calculation of the endosperm/embryo ratios as well as estimation of their contribution to seed biomass. The distribution of lipid could be assessed at 25-µm isotropic resolution as is illustrated in Figure 5, C to E (see also Supplemental Movies S3–S5). Lipid levels appeared to be highest in the endosperm, a
structure composed of larger cells than are present in the embryo (Fig. 5A). Lipid deposition in the embryo was associated with the radicle rather than with the cotyledon (Fig. 5E), and no lipid was deposited in the testa.

Cross-Species Comparison of Seed Architecture and Lipid Distribution

The NMR method in principle enabled a cross-species comparison to be made of seed architecture and lipid distribution. For this purpose, we performed a calibration of the lipid signal (see “Materials and Methods”) and applied this improved measurement scheme to individual seeds of cultivated *N. tabacum* and wild species *Nicotiana sylvestris* and *Nicotiana tomentosiformis* (considered as ancestors; Ren and Timko, 2001). Comparisons between *N. sylvestris* or *N. tomentosiformis* and *N. tabacum* showed that the former formed smaller seeds and distinct morphology (Fig. 6, A–C). The indication from the NMR-based, three-dimensional reconstructions was that for all three species the volume of the embryo was less than that of the endosperm (Fig. 6, D–F). The ratio between these two volumes was 6.2 in *N. tabacum* but much higher in *N. sylvestris* (14.4) and *N. tomentosiformis* (22.7). However, the distribution of lipids in the embryo was similar among all three species, in each case showing a degree of preferential accumulation in the radicle (Fig. 6, G–I), although most of the lipid was deposited in the endosperm (Fig. 6,
J–L). The lipid concentration in both the embryo and endosperm was invariant across the three species.

**Heterologous Expression of BoPPT1 in Tobacco Seed Alters the Pattern of Lipid Accumulation**

The *Brassica oleracea* gene *BoPPT1* encodes the plastidial phosphoenolpyruvate/phosphate translocator PPT1, a protein that plays a role in lipid storage (Schwender and Ohlrogge, 2002; Andre and Benning, 2007). In tobacco plants where *BoPPT1* was driven by the seed-specific promoter *LegB4*, the transgene was activated most strongly during the main storage phase (Supplemental Fig. S1A). The wild type and the transgenic lines were not differentiated with respect to their pattern of lipid accumulation during seed development (Supplemental Fig. S1B). However, by seed maturity, the lipid content of the latter was statistically greater (up to 13%) than that of the wild type, while at the same time, there was no significant alteration in fatty acid composition (Supplemental Fig. S1C).

**Method Design for Simultaneous Monitoring of a Large Number of Seeds**

Having demonstrated that the heterologous expression of BoPPT1 in the seed had a positive impact on lipid accumulation, it was of interest to elucidate which organ was responsible for this effect. This required the simultaneous high-resolution imaging of many seeds. NMR-based imaging is a priori taken as less reasonable for screening tasks due to its low sample throughput. To overcome this limitation here, we developed a novel approach by three-dimensional imaging of multiple seeds and computation of the virtual model seeds (see “Materials and Methods”). Again high-field-strength MRI was used. A small tube containing 99 seeds of a BoPPT1 transgenic line (PPT7) was inserted into a larger tube containing 138 wild-type seeds. The position and size of the wild-type (dark gray) and transgenic (light gray) seeds is illustrated in Figure 7A and the schematic tissue pattern of the seed in Figure 7B. To derive the probable pattern of lipid distribution within each genotype, one virtual model was generated from the signal derived from all the 99 transgenic seeds (Fig. 7C) and a second from all the 138 wild-type seeds (Fig. 7D). When the tissue scheme was merged with the virtual lipid models to highlight the effect of the transgene, it was apparent that the increased quantity of lipid accumulated by the transgenic seed was accompanied by an altered lipid concentration in both the embryo and the endosperm. In the former, most of the alteration was associated with the hypocotyl/radicle, while in the latter it was the region surrounding the embryo that was the most affected. In addition, it was observed that the transgenic seeds were approximately 19% smaller than the wild-type ones and that their lipid content was approximately 13% higher. The transgenic seeds varied with respect to size by a factor of about two, and their size was inversely proportional to their lipid concentration (Fig. 7E). An equivalent correlation was not obvious among the wild-type seeds (Fig. 7F).

**DISCUSSION**

In this work, we propose to use NMR as a noninvasive platform for the multicomponent analysis of submillimeter seeds in vivo. We demonstrate how to count seeds inside the intact tobacco capsule, measure seed sizes, model the seed interior, quantify lipid content, and visualize lipid distribution. Using our approach, hundreds of seeds can be imaged simultaneously.

**Figure 7.** High-throughput MRI visualization and quantification of lipid content in intact tobacco seeds. A, The wild type (dark gray) and transgenic (light gray) “virtual seed” inside the NMR tubes. B, Arrangement of tissues within the tobacco seed. C and D, A representative transgenic (C) and wild-type (D) virtual model seed. E and F, Relationship between seed size and lipid content (given in percentage of volume) along an individual transgenic (E) and wild-type (F) seed. The size of the ellipses is determined by $a$ of the principal components (dark gray 1 $a$; light gray 2 $a$), and the obliqueness of the ellipse indicates the correlation between the size and the lipid concentration of the seed specimen.
These high-resolution NMR methods are a significant step toward the most wanted applications for the model plant Arabidopsis (Arabidopsis thaliana), where numerous oil storage mutants are available (Baud and Lepiniec, 2009; Wallis and Browse, 2010).

NMR is a Versatile Tool for in Vivo Analysis of Tobacco Seeds

While the application to date of NMR to study seed biology has been productive (for review, see Borisjuk et al., 2012), the focus so far has been on large-seeded species, such as Cocos nucifera, Arachis hypogaea, Pinus monticola, and others (Table I). Because the sample size is of matter in NMR hardware design, submillimeter seeds remained challenging for being imaged using MRI. Thus, the seeds of tobacco and Arabidopsis (both of which are well established as model species) have so far remained inaccessible. To the best of our knowledge, only few NMR studies were devoted to small seeds (Manz et al., 2005; Seongjin, 2008), but despite their technical advance, they were not quite appropriate either for lipid detection or for detailed topological studies. In contrast, our protocols have been able to image individual tobacco seeds and their major internal organs. The combination of a high-throughput quantitative measurement of lipid facilitated by low-field TD-NMR and imaging based on high-field MRI has succeeded in a simultaneous imaging of hundreds of seeds in a way in which seed size and lipid content could be assessed on an individual seed basis. The level of resolution achieved (25 μm isotropic) was as much as 100-fold above that obtained by MRI as practiced in the clinical field (Table I). The data captured from hundreds of seeds could be integrated to produce a model of an individual seed reflecting an averaged lipid distribution (Fig. 7). The way is now open to applying MRI to small seeds to derive data with respect to seed composition and internal structure.

MRI Analysis Provides Insights into Lipid Topology in the Tobacco Seed

The topology of lipid accumulation in the seed is conventionally obtained using destructive sampling, and only few techniques are capable of tissue-specific analysis (Schiebold et al., 2011; Horn et al., 2012). The noninvasive NMR approach, extended here to small seeds, has demonstrated the internal distribution of lipid within the seed and has allowed for an evaluation of the relative contribution of the embryo and the endosperm to the overall quantity of lipid accumulated by the seed. We have shown that as in other crop species, tobacco accumulates lipid in both its endosperm and embryo, a pattern that concurs with the outcome of conventional studies (Tomlinson et al., 2004). At maturity, a nonuniform distribution of lipid in both the endosperm and the embryo (Fig. 5) was observed, providing evidence for a spatial incongruity in lipid storage activity during seed development. While the pattern of lipid deposition varied from seed

<table>
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<tr>
<th>Plant Species Seed Size Method</th>
<th>Field Strength (Proton Frequency)</th>
<th>Resolution (Slice Thickness)</th>
<th>Reference</th>
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<tr>
<td>Pine seed</td>
<td>6</td>
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<td>8.5 Tesla</td>
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<td>(Pinus monticola)</td>
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<tr>
<td>Soybean seed</td>
<td>6</td>
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<td>17.6 Tesla</td>
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<td>(Triticum aestivum)</td>
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<tr>
<td>Olives fruit</td>
<td>6</td>
<td>Spin-echo imaging</td>
<td>17.6 Tesla</td>
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<td>(Olea europaea)</td>
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<td>Barley grain</td>
<td>5</td>
<td>Spin-echo imaging</td>
<td>17.6 Tesla</td>
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<tr>
<td>Cotton seed</td>
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<td>11.7 Tesla</td>
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<td>Tobacco seed</td>
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<td>Spin-echo imaging</td>
<td>17.6 Tesla</td>
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<tr>
<td>(N. tabacum)</td>
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to seed, the integrated data set obtained from a large population of seeds has provided a global picture in tobacco: In the endosperm, the highest level of lipid was associated with the region surrounding the embryonic radicle and in large cells, while in the embryo, most of the lipid present was deposited in the radicle/hypocotyl. The embryo in various species accumulates significant quantities of lipid (barley [Hordeum vulgare], Neuberger et al., 2008; rapeseed [Brassica napus], Neuberger et al., 2009; cotton [Gossypium hirsutum], Horn et al., 2012; soybean [Glycine max], Borrisjuk et al., 2005; and oat [Avena sativa], Hayden et al., 2011) but commonly shows completely different gradients in oil distribution compared with tobacco. In dicotyledonous species, the embryo lipid concentrations are generally highest in the cotyledons, and in the monocotyledonous, the preferred site is its analog scutellum. In tobacco, the major site of accumulation in the embryo was the radicle, but, as in a number of other species, in particular jatropha (Jatropha curcas, Gu et al., 2012), oat (Hayden et al., 2011), and castor bean (Ricinus communis, Brown et al., 2012), most of the lipid in the seed was deposited in the endosperm.

The NMR platform provided a tool to show that the ratio between the volumes of the endosperm and the embryo differs between the cultivated species of tobacco N. tabacum and its ancestral species N. sylvestris (maternal progenitor) and N. tomentosiformis (paternal progenitor). Molecular diversity and contribution of parent genomes have been reported in a number of studies (Ren and Timko, 2001; Fricano et al., 2012), while seed architecture and lipid topology received much less attention. We created seed models of all three tobacco species, which differed by size and lipid content (Fig. 6). Seeds of both wild tobacco progenitors are smaller in size and comprise at least 2-fold higher endosperm/embryo ratios compared with N. tabacum. With respect to seed type, this observation is in agreement with general trends in angiosperm seed evolution, namely, that seeds with small embryo and abundant endosperm are rather primitive and evolution shows a general trend toward mature seeds with little or no endosperm in which the embryo occupies most of the seed (Finch-Savage and Leubner-Metzger, 2006). The imaging approach presented here offers a ready method of relating topological, biochemical, and molecular approaches, creating a comprehensive view of seed biodiversity and evolution.

NMR Analysis Indicates a Trade Off between Seed Size and Lipid Level in the Tobacco Seed

The NMR platform was able to simultaneously monitor seed size along with its lipid content in tobacco. This advantageous feature can be exploited for the analysis of mutant or transgenic individuals in which lipid content, seed production, and seed architecture are potentially altered. An example of such an analysis was provided by monitoring the consequences of heterologously expressing BoPPT1 in the tobacco seed. In these seeds, it was possible to detect not just an altered level of lipid deposition, but also the localized accumulation of lipid in distinct regions of the endosperm and embryo.

In the Brassicaceae species oilseed rape and Arabidopsis, phosphoenolpyruvate is believed to be an important precursor of fatty acid synthesis (Schwender and Ohlrogge, 2002; Andre and Benning, 2007; Lonjen and Schwender, 2009). The supply of this metabolite from the cytosol to the plastid is catalyzed by PPT (Knappe et al., 2003), which, at least in soybean seed, is present at the same site as lipid accumulates (Borrisjuk et al., 2005). In Arabidopsis, the down-regulation of PPT depresses lipid accumulation (Prabhakar et al., 2010). Our work demonstrated that its up-regulation raised lipid accumulation by approximately 13%. However, given PPT’s proposed metabolite translocator function (and, hence, its importance for lipid synthesis; see Linka and Weber, 2010), the observed increase in seed lipid content was rather modest. Thus, it is likely, as suggested by Hayden et al. (2011), that other factors, such as the availability of specific intermediates and cofactors, must act to limit lipid storage in the seed. Metabolic control theory (as well as recent data provided by Lonien and Schwender [2009]) suggests that major perturbations of a single step have a lesser impact than do multiple, smaller ones. Therefore, the conclusion is that the up-regulation of PPT will need to be combined with additional interventions where the aim is to further improve the lipid storage capability of the tobacco seed.

The NMR imaging analysis revealed a strong negative correlation between seed size and lipid concentration (Fig. 7E). Assuming that seed metabolism is source limited and that a number of biosynthetic pathways compete with one another for a common substrate (s), any increase in lipid content would be expected to negatively impact seed size. The up-regulation of PPT, a key component of carbon delivery to the plastid, intensified this competition, resulting in a higher lipid concentration but a smaller seed. While seed number per capsule was unaffected by the presence of the BoPPT1 transgene (data not shown), an alteration in the physiological and developmental regulation of lipid accumulation in the seeds (Baud and Lepiniec, 2010) cannot be excluded. Thus, a trade off between growth and storage during the course of seed development is likely to be one of the determinants of lipid yield.

MATERIALS AND METHODS

Plant Materials

Investigated tobacco (Nicotiana spp.) seeds of 36 accessions belonging to 10 Nicotiana wild species and 28 accessions of cultivated tobacco (Nicotiana tabacum) were obtained from the GenBank collection maintained at the Leibniz-Institut für Pflanzenbiochemie and Kulturpflanzenforschung. Details including the geographical origins are given in Supplemental Table S1. Nine wild species were represented by three accessions each. For Nicotiana rustica,
nine accessions belonging to three different varieties (var asiatica, var brasólica, var rusticā) described by Danert (1963) were considered. For N. tabacum, we followed the nomenclature of Danert (1961), dividing the species into 10 varieties. All 10 varieties were considered. For var alpīs and var segāttā, only two samples were available in the GenBank collection, whereas all others were represented by three accessions.

NMR Techniques

Hardware

For the low-field NMR experiments, a Bruker MQ60 was used. The NMR instrument has a magnetic field strength of 1.5 Tesla. The probe head allows for analyzing samples with a diameter of up to 5 mm and a length of 1 cm. All high-field experiments were conducted on a vertical 17.6-Tesla Bruker Avance II wide-bore system operating at 750 MHz. The Micro 2.5 gradient set (Bruker) had an i.d. of 40 mm and a maximum gradient strength of 1 Tesla m⁻¹.

High-Throughput Quantification of Oil Content Using Low-Field TD-NMR

The determination of lipid content using low-field TD-NMR (using MQ60) is based on detecting the spin echo at 7 ms echo time (TE). Mature intact seeds were loaded into the tubes at 20°C, and the spin-echo signal was averaged over 16 measurements. The signal height of the spin echo is proportional to lipid content in seeds (Borijusk et al., 2011; Horn et al., 2011b), given that seeds are dry. The proportionality constant was estimated from a range of samples with known lipid content. About 20 mg of seed was used for individual measurements (with up to eight replicates each). Additionally, following the NMR analysis, seed and some tobacco leaf samples were powdered to obtain conventional estimates of their total lipid content (using gas chromatography). Both noninvasive and conventional assays were superposed (Supplemental Fig. S2). A high correlation between the independent measurement methods was achieved. Therefore, low-field NMR enables accurate, rapid quantification of lipid content. Notably, the NMR analysis takes approximately 1 min per sample and thus allows high-throughput applications.

Two-Dimensional Multislice Imaging of the Capsule

A spin-echo sequence with a frequency-selective VAPOR suppression preparation module was used to acquire the MRI data. An in-house-built 20-mm-i.d. birdcage resonator was used for data acquisition. In detail, the VAPOR suppression parameters were adjusted using global spectroscopic measurements. The transmitting frequency was set on resonant of water. For water suppression, a bandwidth of 3000 Hz with an offset of 500 Hz (away from the lipid signal) was used, and the pulse powers were manually adjusted to achieve optimal water signal suppression. The same procedure was carried out for lipid suppression (bandwidth, 2000 Hz, offset, ~2.5 Hz). The repetition time (TR) was 3000 ms in both experiments, allowing almost complete T1 relaxation. The TE was kept as short as possible (6.2 ms) to minimize effects of T2 relaxation. During one repetition, 33 slices with a slice thickness of 750 µm were acquired. With a field of view (FOV) of 12 × 12 mm² and a matrix size of 192 × 192, an isotropic in plane resolution of 62 µm could be achieved. Averaging the experiment six times resulted in a total experimental time of 57 min for each scan. This approach was applied to create data sets for Figure 2, B and C, and Figure 3, B to D.

Three-Dimensional Imaging of the Capsule

A chemical shift-selective spin-echo sequence (TE = 6.5 ms and TR = 2000 ms, two averages) was used to visualize the whole fruit (Neuberger et al., 2008). The signals from water and lipid were acquired in an interleaved manner during the experiment time of about 18 h. Chemical shift-selective hermite pulses with a bandwidth of 2000 Hz were used for global excitation and refocusing of the protons. The pulses were given on-resonance for water to acquire water images and with an offset of ~2.800 Hz to generate lipid images. An isotropic resolution of 109 µm (FOV = 28 × 14 × 14 mm², matrix = 256 × 128 × 128) was achieved for both the water and the lipid data sets. By interleaving the water and lipid acquisitions, the scan time could be reduced by a factor of two, compared with a sequential acquisition of water and lipid with the same TR. This method was used to acquire the data sets for Figure 4B. In-house Java-based software was used for data reconstruction, while the visualization of the capsule was performed with AMIRA software (VSG). The estimation of the seed number was derived from image segmentation using AMIRA.

Three-Dimensional Imaging of Single Tobacco Seed

A standard three-dimensional spin-echo sequence was used to acquire images from the lipid distribution of single dried tobacco seeds. A water suppression module was not necessary as the residual bound water did not contribute any detectable signal to the experiment. The data were acquired using a 2-mm-i.d. solenoid coil. With an FOV of 4 × 4 × 1 mm³ and a matrix size of 160 × 160 × 40, an isotropic resolution of 25 µm could be achieved. The shortest possible TE was 6.4 ms. With two averages and a TR of 3000 ms, the data set was acquired in ~11 h. This technique was used to acquire the data sets for Figure 5. In-house software was used for data reconstruction, while the visualization and manual segmentation for modeling was performed with AMIRA. For Figure 6, D to L, a similar sequence was used with the following parameters: TE 6.5 ms, matrix 350 × 140 × 140, FOV 12.5 × 5 × 5 mm, isotropic resolution 36 µm, TR 3500 ms, two averages, and 36-h total measurement time.

Three-Dimensional Imaging of Multiple Seeds and Computation of the Virtual Model Seeds

For sample preparation and data acquisition, two different types of seeds (transgenic PPT7 line and the wild type) of mature seeds were examined in this experiment. While one of them (PPT7) was placed inside a 3-mm NMR tube (2.5-mm i.d.), the other species (the wild type) was poured carefully around the 3-mm NMR tube after it was inserted into a 5-mm NMR tube (4-mm i.d.). Both the NMR tubes were filled with vegetable oil for B1 correction and calibration. The complete sample was placed into an in-house-built 5-mm scroll coil (Hayes et al., 1985; Grant et al., 2001). As no water signal was present, a standard three-dimensional spin-echo sequence (TR = 1750 ms, TE = 5.7 ms, FOV = 9 × 4.5 × 4.5 mm³, matrix size = 320 × 160 × 160) without a suppression module could be used to acquire data sets with a 28-µm isotropic resolution. The total experimental time was 25 h.

Data Reconstruction and Analysis

The acquired data set was zero filled by a factor of three in each direction before it was reconstructed by a three-dimensional inverse fast Fourier transformation using our software. To exclude strong B1 inhomogeneity at the side of the three-dimensional data set along the axis of the scroll coil and artifacts caused by insufficient filtering in the frequency (readout) direction, the data set was cropped to a size of 768 × 480 × 480 voxels. To correct the data for the residual B1 inhomogeneity within the data set, a calibration had to be applied. This correction could be performed using the external vegetable oil, which also served as a known standard (density 0.914 mm³, TR 11 h). This technique was used to acquire the data set. Furthermore, this data set was cropped to a size of 768 × 480 × 480 voxels.

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The resulting plasmid was transformed into (ocs) terminator, resulting in the expression cassette LegB4P/BoPPT1/ocs. To phospho BoPPT1 complementary DNA fragment was cloned between the LegB4 promoter from \textit{Vicia faba} (Bäumlein et al., 1991) and 3′-region of octopine synthase (ocs) terminator, resulting in the expression cassette LegB4P/BoPPT1/ocs. To generate a binary vector, the LegB4P/BoPPT1/ocs was cloned into pGPTV-bar (Becker et al., 1992) plasmid replacing the Gus gene cassette. All plasmid manipulations were carried out according to the methods described by Sambrook et al. (1989). The resulting plasmid was transformed into Agrobacterium \textit{tumefaciens} strain EHA105 and transferred into tobacco 'SNN' according to the method of Horsch et al. (1985).

The presence of PPT expression cassette in transformed tobacco lines was confirmed by PCR analysis of isolated genomic DNA using primers binding at 3′-region of LegB4 promoter and 5′-region of BoPPT1 gene sequences (5′-CACACACGTTCTGTCACAGG-3′ for LeB4 and 5′-ACTGCGCTAGCTCAGTTT-3′ for PPT). More than 90% of tobacco lines selected on kanamycin showed the presence of gene-specific PCR fragments. Tobacco PPT plants were further analyzed for the presence of introduced cassette by Southern DNA analysis, and the expression of BoPPT1-specific mRNA was confirmed by northern analysis (Supplemental Fig. S1A). Analysis of fatty acid composition in seeds of tobacco plants was performed by gas chromatography as previously described (Borisjuk et al., 2005).

**Supplemental Data**

Supplemental Data

The following materials are available in the online version of this article.

**Supplemental Figure S1.** Analysis of transgenic tobacco plants heterologously expressing the \textit{B. oleracea} gene BoPPT1.

**Supplemental Figure S2.** TD-NMR-based measurement of the lipid content of tobacco leaf and seed tissue is consistent with NMR and gas chromatography-based data.

**Supplemental Table S1.** \textit{Nicotiana} spp. accessions sampled from the IPK GenBank collection.

**Supplemental Movie S1.** Visualization of water (right) and lipid (left) distribution in the tobacco capsule.

**Supplemental Movie S2.** Three-dimensional model of the tobacco capsule.

**Supplemental Movie S3.** Three-dimensional model of the tobacco seed, showing the embryo (green) surrounded by endosperm.

**Supplemental Movie S4.** Topology of lipid accumulation in the embryo of a tobacco seed.

**Supplemental Movie S5.** Topology of lipid accumulation in the endosperm of a tobacco seed.

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**LITERATURE CITED**


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