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# NMR-based metabolomics at work in phytochemistry

R. Verpoorte · Y. H. Choi · H. K. Kim



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**Abstract** Metabolomics is defined as both the qualitative and quantitative analysis of all metabolites in an organism unraveling correlation with other OMICs data. Many of the technologies used in metabolomics have method-specific advantages and drawbacks in terms of diversity of metabolites detected, sensitivity, or resolution. In this paper, the potential of NMR spectrometry applied to metabolomics is reviewed using examples of *Nicotiana tabacum* and *Catharanthus roseus* 

**Keywords** Metabolomics · Metabolic analysis · NMR · Nicotiana tabacum · Catharanthus roseus

## Introduction

In the past years, several genomes have been completely sequenced and the genomes of more organisms are being sequenced. The number of fully sequenced organisms will grow exponentially. This will open novel possibilities of studying the functions of the genes in these organisms. Functional genomics requires a thorough definition of the phenotype, to be able to determine what changes result from up- or down-regulation of genes. A phenotype can be defined by morphological characteristics, by physiological measurements, as well as by biochemical (transcriptomics or proteomics) and chemical (metabolomics) analysis. In recent years, more and more attention is being paid to the chemical characterization of the phenotype. Chemical charac-

R. Verpoorte (⊠) · Y. H. Choi · H. K. Kim Department of Pharmacognosy, Section Metabolomics, Institute of Biology, Leiden University, P.O. Box 9502, 2300RA, Leiden, The Netherlands e-mail: verpoort@chem.leidenuniv.nl terization can be done on the level of macromolecules (e.g., proteomics and characterization of polysaccharides and lignins) or low molecular weight compounds (the metabolome). The metabolome consists of two types of compounds, the primary metabolites and the secondary metabolites. The primary metabolites are compounds involved in the basic functions of the living cell, such as respiration and biosynthesis of the amino acids and other compounds needed for a living cell. Basically all organisms share the same type of primary metabolites, though per class of organisms there may be large differences in the biosynthetic pathways present, e.g., mammalians lack amino acid biosynthesis, as they rely on external sources for their needs. The secondary metabolites are species specific, they play a role in the interaction of a cell with its environment, which can be other cells in the organism or external organisms, e.g., in case of plants to attract pollinators, or to defense itself against pests and diseases. So primary metabolites in plants are important for the growth and agricultural yields, whereas the secondary metabolites are, e.g., concerned with flavor and color of our food and with the resistance of plants against pests and diseases.

Obviously in studying functions of genes metabolomics will be an important tool. Metabolomics is generally defined as both the qualitative and quantitative analysis of all metabolites in an organism. In analyzing a metabolome both the primary and secondary metabolites will be detected. Plants share the primary metabolites, therefore methods that can easily identify and quantify the primary metabolites with high reproducibility are required. One or more methods are required that in all labs and at any time will give the same results, which thus enables a public database which can be used in the same way as the databases for protein and gene sequences. Also for the secondary metabolites such a database is needed, though in that case the occurrence of most compounds is restricted to one or a few species only. The reproducibility is also required for studying the effect of any external condition on the plant metabolome, as one needs to know the biological variation. That requires large numbers of analyses to determine the metabolomic changes as function of daily, seasonal, and developmental variation. The results should function as the control samples for all future analyses.

Thus, reproducibility is the most important criteria for developing a metabolomics technology platform. The other criteria concern the ease of quantitation and identification, the number of metabolites that can be measured, and the time needed for an analysis, including the sample preparation. Obviously a highthroughput methodology is preferred, for which robotization is important. Based on the underlying principle, the methods which one may consider for a metabolomic platform can be grouped as follows:

- chromatographic methods (liquid chromatography, gas chromatography (GC), capillary electrophoresis, thin layer chromatography);
- mass spectrometry (MS);
- nuclear magnetic resonance spectrometry.

A combination of chromatography and one or both of the other two methods also is an option. In Table 1, the various weak and strong points of these methods are summarized. Concerning the mentioned criteria, chromatographic methods have a major problem in reproducibility. Although the intra- and inter-day variability may be reasonable, there is always a problem of getting the same separation in another lab due to small changes in external conditions, chemicals, or solvents. Moreover, due to aging, column performance changes through time. Last but not least, commercial companies always come with improved columns, which usually also means an altered selectivity of the system. A simple search in the literature for the separation of, e.g., tropane alkaloids shows that every year a number of new "improved" separation systems is described for these alkaloids. This is a poor basis for a long-term reproducible database of results of such an analysis. MS also has a problem of reproducibility, mainly because in MS analysis the compounds have to be ionized before measuring. There exist many different types of mass spectrometers with different ways to ionize the molecules, making comparison difficult. Besides the variation of the hardware and the conditions chosen, also a considerable matrix effect exists for the efficiency of the ionization, which means that a compound in different extracts may give different quantitative results.

NMR spectrometry is a physical measurement of the resonances of magnetic nuclei, such as <sup>1</sup>H, <sup>13</sup>C, or <sup>15</sup>N in a strong magnetic field. Each compound has a highly specific spectrum. The only variables are the solvent used and the magnetic field strength. The solvent is easy to standardize. The field strength is not a problem with <sup>13</sup>C NMR and even in case of <sup>1</sup>H NMR it can be overcome (see below). The high reproducibility makes that NMR among the three methods mentioned is the most suited for a public domain metabolomics database that may serve for many years to come.

Concerning the criteria of quantitation, the chromatographic methods have shown through the years to be very suitable for quantitative analysis. However, they always require calibration curves for the compounds that are quantified, as each compound will give a different detector response. Only in GC in combination with a flame ionization detector (FID) one may compare peak intensities with each other, and thus make an estimation of the absolute quantities of all compounds detected. UV-detection in HPLC, and MS in HPLC and GC will show totally different sensitivities for each compound, some even not being detected at all. For complex samples with a number of compounds, it is not feasible to make calibration curves for all compounds. That means only a relative comparison for each compound in different analyses is possible, but no absolute quantitation can be done, i.e., one cannot compare the amounts present of different compounds. This is a major disadvantage, e.g., one cannot make any conclusion about the conversion of one compound into another in case

Table 1 Description of chromatographic methods used in metabolomics

Methods	Target metabolites	Sample preparation	Reproducibility	Resolution	Detector
TLC	General	Simple	Low	Low	UV, MS, color reagents
GC	Non-polar (low molecular)	Elaborate derivatization	Medium	High	FID, TCD, NPD, MS
HPLC CE	Polar (chromophore) Ionic	Elaborate –	Medium Medium	Medium High	UV, RI, MS, ELSD, fluorescence, NMR UV, MS

of a decrease of one compound and an increase in another related compound, i.e., conclusions about stoichiometry cannot be drawn from the data. For qualitative analysis the hyphenated chromatographic methods (HPLC with diode array detector, -MS, and/ or NMR, GC-MS) are the most powerful as they offer both retention behavior and physical characteristics as a tool for identification. MS allows the determination of the molecular weight, and in case of high resolution also of the elemental composition, but this is not always sufficient to determine the structure. Tandem MS might be of help to identify in such case the compound through its fragmentation pattern, but in case of novel compounds further spectral data are required. The performance for NMR can be improved by using two-dimensional (2D) NMR methods (see below), which even may enable structure elucidation of novel compounds in a mixture. In fact <sup>13</sup>C NMR spectra were already used in the early 1980s for fingerprinting essential oils (Formáček and Kubeczka 1982). We described the use of <sup>1</sup>H NMR spectra for the fingerprinting of plant cell cultures (Schripsema and Verpoorte 1991). It has been used ever since for the analysis of sugars in cell cultures and media (e.g., Kraemer et al. 1999, 2002; Schripsema and Verpoorte 1991). Also other compounds have been analyzed in plant extracts by means of NMR, e.g., pyrrolizidine alkaloids (Pieters et al. 1989) and hop bitter agents (Hoek et al. 2001). With the increasing field strength of the NMR-magnets and the improved resolution the application of NMR for metabolomics became feasible.

Here we will first discuss the general approach to metabolomics and the sample preparation methods and show some examples of the use of NMR-metabolomics in phytochemical studies. The weak points and possible solutions will be discussed.

## Sample preparation

In metabolomics studies there are three phases. The first one is a qualitative phase in which as many as possible of the signals observed in the NMR are assigned to compounds. After this phase it is first necessary to determine the biological variation for the system studied, that means measuring a large number of samples, e.g., young and old leaves, different times of the day, and different stages of development of the plant. The last step is to compare the model system under different experimental conditions or compare, e.g., diseased or transgenic plants with the normal wild type plant. An important aspect is of course the extraction method. Plants contain a wide variety of compounds with totally different polarities. No single method is able to detect all metabolites in a single operation. To extract as many as possible we developed some years ago a two-phase extraction method in which chloroform-methanolwater (2:1:1) was used. This procedure extracts both polar and polar compounds in the same operation. This method gives good results, but is quite elaborate. Therefore, a series of single solvents were tested for the extraction of the metabolites from Arabidopsis (unpublished data, Fig. 1). By multivariate analysis, the spectra were compared in order to see which solvent covered the widest range of compounds. Methanolwater (1:1) came out as the best one in terms of the diversity (Choi et al. 2006; Hendrawati et al. 2006;

**Fig. 1** Score plot of PCA (PC1 vs PC3) based on <sup>1</sup>H NMR spectra of the *A*. *thaliana* extracts. *I* Acetone, 2 acetone–MeOH (1:1), 3 MeOH, 4 AcCN, 5 MeOH– water (1:1), 6 water. The *ellipse* represents the Hotelling T2 with 95% confidence in score plots



Liang et al. 2006a, b; Widarto et al. 2006). By using these solvents in deuterated form, extracts can be measured directly after extraction, without any need of evaporation and reconstitution of the extract. To avoid shifts due to differences of pH (Schripsema et al. 1986) a buffered solvent ( $KH_2PO_4$ ) is used (pH 6.0).

# Transgenic tobacco plants overexpression microbial genes for salicylate biosynthesis

We have overexpressed the microbial genes encoding the isochorismate pathway leading to salicylate (SA). The genes, isochorismate synthase from *Escherichia*  *coli*, and isochorismate lyase from *Pseudomonas fluorescens*, result in the constitutive production of SA in tobacco plants. These plants have an increased resistance against viral and fungal infection (Verberne et al. 2000). This raised the question if any secondary metabolites pathways were altered, either due to the SA that is involved as signal compound in the systemic acquired resistance (SAR), or due to channeling of chorismate away from the other important chorismate pathways branches, leading among others to phenylalanine (and from there to all phenylpropanoids) and tryptophan. In a series of papers, various targeted approaches have been described, which led to the conclusion that terpenoids (by GC-analysis, Nugroho et al. 2002a), and alkaloids

Table 2 Characteristic <sup>1</sup>H chemical shifts of common metabolites in plants [CD<sub>3</sub>OD-KH<sub>2</sub>PO<sub>4</sub> in D<sub>2</sub>O (pH 6.0)]

Metabolites	Chemical shifts (ppm) and coupling constants (Hz)					
Alanine	$\delta$ 1.48 (H-3, d, $J$ = 7.2 Hz)					
Ascorbic acid	$\delta$ 4.52 (H-4, d, J = 2.0 Hz)					
Caffeic acid	$\delta$ 7.52 (H-7, d, $J = 15.9$ Hz), $\delta$ 7.12 (H-2', d, $J = 2.1$ Hz), $\delta$ 7.03 (H-6', d, $J = 8.4, 2.0$ Hz), $\delta$ 6.88 (H-5', d, $J = 8.4$ Hz), $\delta$ 6.29 (H-8', d, $J = 15.9$ Hz)					
Chlorogenic acid	$\delta$ 7.60 (H-7', d, $J = 15.9$ Hz), $\delta$ 7.15 (H-2', d, $J = 2.1$ Hz), $\delta$ 7.05 (H-6', d, $J = 8.4$ , 2.0 Hz), $\delta$ 6.87 (H-5', d, $J = 8.4$ Hz), $\delta$ 6.36 (H-8', d, $J = 15.9$ Hz), $\delta$ 5.33 (H-3, td, $J = 10.0$ , 4.8 Hz), $\delta$ 4.21 (H-5, br q, $J = 3.1$ Hz), $\delta$ 3.8 (H-4 dd $J = 9.9$ 3.2 Hz), $\delta$ 2.17 (H-2 $\beta$ H-6 $\beta$ m), $\delta$ 2.04 (H-2 $\alpha$ H-6 $\alpha$ m)					
Citramalic acid <sup>a</sup>	$\delta$ 2.80 (H- $\beta$ , d, J = 15.7 Hz), $\delta$ 2.61 (H- $\beta$ , d, J = 15.7 Hz)					
Citric acid <sup>a</sup>	$\delta$ 2.74 (H- $\beta$ , d, J = 17.6 Hz), $\delta$ 2.56 (H- $\beta$ ', d, J = 17.6 Hz)					
Fatty acids	$\delta 0.88 (\text{H-}\omega, \text{t}, J = 7.5 \text{ Hz})$					
Ferulic acid	$\delta$ 7.56 (H-7', d, J = 15.9 Hz), $\delta$ 7.19 (H-2', d, J = 2.1 Hz), $\delta$ 7.10 (H-6', d, J = 8.4, 2.0 Hz), $\delta$ 6.88 (H-5', d, J = 8.4 Hz), $\delta$ 6.33 (H-8', d, J = 15.9 Hz)					
Fumaric acid	$\delta$ 6.56 (s)					
α-Glucose	$\delta$ 5.20 (H-1, d, $J = 3.8$ Hz)					
$\beta$ -Glucose	$\delta$ 4.59 (H-1, d, J = 7.9 Hz)					
GABA	$\delta$ 2.28 (H-2, d, J = 7.5 Hz), $\delta$ 1.88 (H-3, q, J = 7.5 Hz), $\delta$ 1.88 (H-4, t, J = 7.5 Hz)					
Glutamic acid	$\delta$ 2.07 (H-3, m), $\delta$ 2.36 (H-4, m)					
Glutamine	$\delta 2.15$ (H-3, m), $\delta 2.47$ (H-4m)					
<i>p</i> -Hydroxy benzoic acid	$\delta$ 7.91 (H-3, H-5, d, J = 8.8 Hz), $\delta$ 6.92 (H-2, H-6, d, J = 8.8 Hz)					
Inositol	$\delta$ 4.00 (H-2, t, $J = 2.8$ ), $\delta$ 3.61 (H-4 and 6, t, $J = 9.9$ ), $\delta$ 3.44 (H-1 and 3, dd, $J = 9.9$ , 2.9 Hz), $\delta$ 3.24 (H-5, t, $J = 9.3$ Hz)					
Isocitric acid <sup>a</sup>	$\delta$ 4.00 (H-α, d, $J$ = 4.2 Hz), $\delta$ 3.04 (H- $\beta$ , td, $J$ = 11.7, 4.2 Hz), $\delta$ 2.56 (H- $\gamma$ , dd, $J$ = 15.4, 7.3 Hz), $\delta$ 2.56 (H- $\gamma'$ , dd, $J$ = 15.4, 7.7 Hz)					
Kaempferol	$\delta$ 8.04 (H-2' and H-6', d, $J$ = 8.6 Hz), $\delta$ 6.74 (H-3' and H-5', d, $J$ = 8.6 Hz), $\delta$ 6.52 (H-8, d, $J$ = 2.0 Hz), $\delta$ 6.28 (H-6, d, $J$ = 2.0 Hz)					
α-Linolenic acids	$\delta$ 0.96 (H- $\omega$ , t, J = 7.5 Hz), $\delta$ 1.30 (CH <sub>2</sub> brs)					
Malic acid <sup>a</sup>	$\delta$ 4.32 (H- $\alpha$ , dd, J = 6.6, 4.7 Hz), $\delta$ 2.80 (H- $\beta$ , dd, J = 16.6, 4.7 Hz), $\delta$ 2.61 (H- $\beta$ ', dd, J = 16.6, 6.6 Hz)					
Proline	$\delta$ 4.08 (H-2, dd, $J = 8.6, 6.4$ Hz), $\delta$ 2.34 (H-3, m)					
Quercetin	$\delta$ 6.99 (H-5', d, J = 8.6 Hz), $\delta$ 6.52 (H-8, d, J = 1.8 Hz), $\delta$ 6.28 (H-6, d, J = 1.8 Hz)					
Rhamnose in flavonoid	$\delta$ 1.10 (3- <i>O</i> -rhamnosylglucoside, H-6, d, $J = 6.1$ Hz)					
Sinapic acid	δ 7.48 (H-7, 16 Hz), δ 6.93 (H-2, H-6, s), δ 6.37 (H-8, 16 Hz)					
Steroids	$\delta$ 0.78 (H-18, s), $\delta$ 0.85 (H-19, s), $\delta$ 0.89 (H-21, d, $J = 7.0$ Hz)					
Succinic acid <sup>a</sup>	$\delta$ 2.54 (s)					
Sucrose	$\delta$ 5.40 (H-1, d, $J$ = 3.8 Hz), $\delta$ 4.17 (H-1, d, $J$ = 8.5 Hz)					
Threonine	$\delta$ 1.32 (H-5, d, $J = 6.6$ Hz)					
Tryptophan	δ 7.20 (H-5', t, <i>J</i> = 7.4 Hz), δ 7.29 (H-6', t, <i>J</i> = 7.5 Hz), δ 7.32 (H-2', s), δ 7.54 (H-7', d, <i>J</i> = 8.1 Hz), δ 7.73 (H-4', d, <i>J</i> = 7.9 Hz)					
Tyrosine	$\delta$ 7.19 (H-2, H-6, t, J = 8.5 Hz), $\delta$ 6.86 (H-3, H-5, t, J = 7.5 Hz)					
Valine	$\delta$ 1.00 (H-3a, d, J = 6.8 Hz), 1.05 (H-3b, d, J = 6.8 Hz)					

<sup>a</sup> Changeable by pH and concentration

(by HPLC-analysis, Nugroho et al. 2002b) were not effected, whereas flavonoids (by HPLC-analysis, Nugroho et al. 2002c) showed lower levels, similar conclusions were drawn from a non-targeted approach using centrifugal partitioning chromatography to prefractionate the extract followed by LC–MS to analyze the fractions (Halim et al. 2003). This was a major effort as each of the different groups of compounds required the validation of a suited method.

The next step was to use this model system for an NMR-based metabolomics approach, to learn how this approach compared with the elaborate targeted approach.

The first step was the qualitative analysis. A number of compounds could easily be identified. To facilitate the identification we started to build a database with NMR spectra of common plant metabolites. This database contains now more than 300 compounds, and more than 600 spectra (both 1D and 2D spectra). Some of the very common diagnostics signals are summarized in Table 2. They represent, e.g., phenolics, the anomeric protons of sugars, amino acids, and organic acids. In each area of the NMR, one may look for specific groups of compounds (Fig. 2).

Using this method extensive studies have been made of the metabolomes of the tobacco plants (Choi et al. 2004a, 2006). We used the two-phase or deuterated solvent extraction method, and measured the NMR spectra. For analyzing the large data set of NMR spectra, multivariate analysis (using SIMCA-P, Umetrics, Umeå, Sweden) was used. From the results we were able to clearly distinguish these in wild type and transgenic plants. To introduce the NMR spectra into the biostatistics program, they have to be processed into digital quantifiable data. This is done by dividing the spectra into buckets (or bins) of 8 or 16 Hz, peak height in each bucket is integrated and introduced into a text file used for further statistical analysis. This processing can be done by the NMR-software (e.g., AMIX, Bruker Biospin GmbH, Rheinstetten, Germany). For robustness of the bucketing high reproducibility of chemical shift is required. By using a buffered NMR-solvent, the reproducible chemical shift



**Fig. 2** Typical <sup>1</sup>H spectra of *Arabidopsis* [400 MHz, CD<sub>3</sub>OD–KH<sub>2</sub>PO<sub>4</sub> in D<sub>2</sub>O (pH 6.0)] (adapted from Hendrawati et al. 2006)

can be obtained. However, there are still problem with organic acids such as citric, citramalic, isocitric, malic, and succinic acid. The chemical shift of these signals may vary considerably depending on pH of the solution. In order to solve the problem the region in which they occur can be bucketed by larger size (e.g., 40 Hz). However, one should always be aware of the risk that a compound may show changes in its NMR spectra, e.g., due to complexity of the mixture, or due to concentration effects (Rhee et al. 2004). Table 3 shows the example of the variation of chemical shifts of citric acid and malic acid in different concentration.

The signals responsible for the separation found in the score plot of principle component analysis (PCA) were identified from the loading plot (Fig. 3). Assigning these signals to various known tobacco constituents resulted in the conclusion that the transgenic plants had lower levels of flavonoids, chlorogenic acid, and sugars, and higher levels of malic acid and alanine (Choi et al. 2004a). The alkaloid and terpenoid levels were not changed. Similar conclusions as for the targeted approach (see above), but as extra information the increased levels of sugars and alanine were observed in this approach. This nicely showed the great potential of studying metabolic changes in a single NMR-analysis.

#### Phytoplasma infected Catharanthus roseus

The same approach as for tobacco was applied to *Catharanthus roseus* plants infected with different

**Table 3** <sup>1</sup>H chemical shifts of H- $\beta$  of citric and malic acids in different concentrations [CD<sub>3</sub>OD–KH<sub>2</sub>PO<sub>4</sub> in D<sub>2</sub>O (pH 6.0)]

Concentration (mg/mL)	Citric acid		Malic acid		
	Η-β	Η-β'	Η-β	Η-β'	
1	$\delta$ 2.73 (d, J = 15.0 Hz)	$\delta$ 2.62 (d, $J = 15.0$ Hz)	$\delta$ 2.68 (dd, $J = 14.0, 3.3$ Hz)	$\delta$ 2.37 (dd, $J = 14.0, 10.0$ Hz)	
5	$\delta$ 2.89 (d, $J = 15.0$ Hz)	$\delta$ 2.77 (d, J = 15.0 Hz)	$\delta$ 2.84 (dd, $J = 15.0, 5.0$ Hz)	$\delta$ 2.67 (dd, $J = 15.0, 7.0$ Hz)	
10	$\delta$ 2.87 (d, $J = 13.3$ Hz)	$\delta$ 2.74 (d, $J = 13.3$ Hz)	$\delta$ 2.85 (dd, $J$ = 15.0, 5.0 Hz)	$\delta$ 2.72 (dd, $J = 15.0, 7.0$ Hz)	

Fig. 3 Score (a) and loading plot (b) of methanol-water fraction for wild type and CSA-line #16 leaves and veins following PCA (PC1 vs PC2). WNL wild type noninoculated leaf, WIL wild type inoculated leaf, WSL wild type systemic leaf, CNL CSA non-inoculated leaf, CIL CSA inoculated leaf, CSL CSA systemic leaf, WNV wild type non-inoculated vein, WIV wild type inoculated vein, WSV wild type systemic vein, CNV CSA noninoculated vein. CIV CSA inoculated vein, CSV CSA systemic vein (adapted from Choi et al. 2004a)



types of phytoplasmas (Choi et al. 2004b). Again a clear separation could be obtained for the plants, analysis of the markers for the differences led to the conclusions as shown in Fig. 4. Also, in this case, the macroscopic view on the metabolome via the NMR-analysis clearly showed major changes in the infected plants. In this case, a considerable effort was made to identify a number of compounds using 2D NMR spectrometry, this among others allowed the identification of the iridoids (see Fig. 5).

## **Brassicaceae NMR-based metabolomics**

From the results with the tobacco and *C. roseus* it became clear that biostatistical software (e.g., SIMCA-P) especially developed for metabolomics-like applications might be an advantage. Also, we learned that in general the two-phase extraction system was quite elaborate and thus not the preferred one for a highthroughput systems. Moreover, the chloroform phase mostly only showed strong signals due to fatty acids, which can be quantified as group, but individual acids cannot be observed in the 1D NMR spectra. On the other hand, the methanol-water phase gave an excellent overview of the plant's metabolome, and particularly on a broad range of secondary metabolites. A direct extraction with deuterated methanol-water solvent thus seems to be most suited (unpublished data). For the identification of compounds further 2D NMR methods were explored to identify compounds in the crude extracts. Also, the problem of comparing spectra Fig. 4 Schematic pathway for the biosynthesis of terpenoid indole alkaloid, phenylpropanoid, and phenolic acid. The increased metabolites in *C. roseus* leaves infected by phytoplasma are shown in *bold font* (adapted from Choi et al. 2004b)



8.0 7.9 7.8 7.7 7.6 7.5 7.4 7.3 7.2 7.1 7.0 6.9 6.8 6.7 6.6 6.5 6.4 6.3 6.2 6.1 ppm

**Fig. 5** HMBC spectra of water fraction of phytoplasma (UDI-NESE) infected *C. roseus* leaves. In HMBC spectra (C), *1* correlation of H-3 and C-5 of secologanin, *2* correlation of H-3 and C-4 of secologanin, *4* correlation of H-3 and carbonyl group of secologanin, *5* correlation of H-7 and carbonyl group of 6 correlation of H-7' and C-6' of chlorogenic acid, *7* correlation of H-7' and carbonyl group of chlorogenic acid, *8* correlation of H-3 and C-5 of loganic acid, *9* correlation of H-3 and C-1 of loganic acid, 10 correlation of H-3 and C-4 of loganic acid, 11 correlation of correlation of H-3 and carbonyl group of loganic acid, 12 correlation of H-2' and C-1' of chlorogenic acid, 13 correlation of H-2' and C-3', 14 correlation of H-2 and C-1 of gallic acid derivatives, 15 correlation of H-2 and C-3 of gallic acid derivatives, 16 correlation of H-2 and C-1' of chlorogenic acid, 18 correlation of H-8' and C-1' of chlorogenic acid, 18 correlation of H-8' and C-1' of chlorogenic acid, 18 (adapted from Choi et al. 2004b)



**Fig. 6** Two-dimensional J-resolved NMR spectra of phenylpropanoids of *B. rapa* leaves in the range of  $\delta$  6.25–6.50 (**a**);  $\delta$  5.32– 5.42 (**b**). The spectra were measured in methanol-d<sub>4</sub> (adapted from Liang et al. 2006a)

obtained at different field strength needed to be solved. These matters were addressed in a series of experiments with tobacco, *Brassica rapa* and *Arabidopsis thaliana*.

One of the major problems of NMR-metabolomics is the overlap of many signals, particularly in some crowded regions. The use of 2D NMR spectrometry can improve this considerably. In 2D J-resolved spectra, the second dimension gives for each signal the coupling constant (Fig. 6a, Liang et al. 2006a). In this way, the congestion is to a great extend solved. Moreover, the coupling constant is an excellent diagnostic tool for the classification of signals as belonging to a certain class of compounds (e.g., cinnamic acid derivatives, Fig. 6a) (Liang et al. 2006a). Moreover, by projecting the signals of 2D J-resolved spectra on the F2 axis (the chemical shift), each signal is reduced to a singlet, i.e., each proton is now observed as a singlet (Fig. 7). The spectra are comparable in appearance to <sup>13</sup>C NMR spectra. This simplifies the spectra enormously, as well as the identification of known compounds as the overlap of signals is much less. The most important aspect of these projected 2D J-resolved spectra is that all secondary effects are eliminated, consequently the spectra are independent of the field



**Fig. 7** <sup>1</sup>H NMR (a), 2D J-resolved (b), and projected 1D Jresolved (c) spectrum of healthy *Nicotiana tabacum* leaves in the range of  $\delta$  6.25–6.50 (adapted from Choi et al. 2006)

strength (Choi et al. 2006). This makes them to excellent candidates for storage in a large database for longterm use in metabolomics. The disadvantage of these spectra is that because of the projection the intensity of the signals of the different protons cannot be compared anymore. For absolute quantitation, the intensities of the different peaks in the second dimension need to be added.

For the identification of compounds 2D NMR spectrometry methods are of great value. Signals can be identified as well as their coupling with other protons or carbons. Figure 6 shows part of the spectrum of *B. rapa* in which from the 2D J-resolved spectrum it was clear that the congested signals at  $\delta$  6.3–6.5 were due to cinnamic acid derivatives as they all have protons with a large coupling constant (16 Hz, Liang et al. 2006a). A single separation step of the extract was made to have an enriched phenolic compound fraction, and thus reduce the NMR time needed to run the various 2D NMR spectra. Further 2D NMR spectra were used to elucidate their structures. From the COSY and HMBC spectra, it can be learned that malic acid moiety attaches to carboxylic group of



 $R_1 = OCH_3$ ,  $R_2 = OH$ , trans-5-hydroxyferuloyl malate

 $R_1 = OH, R_2 = H, trans-caffeoyl malate$ 

 $R_1 = H, R_2 = H, trans-coumaroyl malate$ 

 $R_1 = OCH_3$ ,  $R_2 = H$ , trans-5-hydroxyferuloyl malate

R<sub>1</sub> = OCH<sub>3</sub>, R<sub>2</sub> = OCH<sub>3</sub>, trans-5-sinapoyl malate



 $R_1 = OCH_3$ ,  $R_2 = OH$ , *cis*-5-hydroxyferuloyl malate

 $R_1 = OH, R_2 = H, cis$ -caffeoyl malate

 $R_1 = H, R_2 = H, cis$ -coumaroyl malate

 $R_1 = OCH_3$ ,  $R_2 = H$ , *cis*-5-hydroxyferuloyl malate

$$R_1 = OCH_3$$
,  $R_2 = OCH_3$ , *cis*-5-sinapoyl malate

Fig. 8 Chemical structures of phenylpropanoids identified in *B. rapa* leaves

phenylpropanoids. The presence of the attached H-2 of malic acid was confirmed by 2D J-resolved spectra (Fig. 6b). From the various 2D NMR experiments, a whole series of cinnamic acid derivatives was identified as shown in Fig. 8, both *cis*- and *trans*-isomers were found (Liang et al. 2006a). Whether the *cis*-derivatives are artifacts or already formed in the plant remains to be determined.

Using the NMR-based metabolomics approach we studied the effects of a methyl jasmonate treatment of *B. rapa* plants (Liang et al. 2006b). The score plot of the PCA (Fig. 9) shows a clear separation between the treated and non-treated plants. From the loading plot, it can be concluded that the phenylpropanoids discussed above increased in the treated plants, also an

indole glucosinolate such as neoglucobrassicin and indole acetic acid increased in the treated plants.

### Virus infected tobacco

The various tools we developed since our first tobacco experiments were applied on a study of tobacco mosaic virus (TMV) infected tobacco plants. This system has been studied extensively on the level of genes and proteins to clarify the SAR in plants. As already discussed above SA does play an important role in SAR. Interestingly very little work has been done on the role of secondary metabolism in SAR, as still the function of several of the pathogen-related proteins is unknown, they might play a role in secondary metabolism. We thus applied our NMR-metabolomics platform technology on this system (Choi et al. 2006). In Fig. 10, the PCA score plot is shown. As expected the metabolome of the leaf changes through time, similar changes are observed in the infected leaves (mainly in PC1), but on top of that in PC2 or PC3 clear differences between leaves from the infected and control plants can be observed.

By using the various 2D NMR methods, a number of signals that are clearly involved in the SAR response and direct defense against the infection could be identified (see above). In Fig. 11, the various changes are summarized. Apparently some of the effects occur in the infected leaves and SAR leaves, whereas others only in the SAR or the infected leaves. The meaning of the various changes need of course further study. But obviously this systems biology type of approach using NMR-based metabolomics, gives many novel leads for further study.

#### NMR-based metabolomics platform

These various examples nicely show the great potential of NMR as a tool for having a macroscopic view on the plant metabolome. It is thus of great interest in functional genomics studies as well as in systems biology studies on all kind of biological processes. But such a platform has also many other fields of application, such as quality control of botanicals and food (Choi et al. 2005; Kim et al. 2005; Yang et al. 2006); phenotyping of plants for breeders rights (Fernie et al. 2004; Fiehn et al. 2000; Fukusaki and Kobayashi 2005; Roessner et al. 2002); studies on the compounds related to pharmacological activity in medicinal plants (Joshi et al. 2002); dereplication of plant extracts in bioprospecting







Fig. 10 Score plot of PCA of healthy and infected *Nicotiana* tabacum leaves by TMV. ● lower leaves of healthy plants, ■ upper leaves of healthy plants, ▲ local-infected leaves of TMV-infected plants, ♦ SAR leaves of TMV-infected plants. a PCA for lower and upper leaves in healthy plants and inoculated and SAR leaves, b PCA for lower leaves in healthy plants and inoculated leaves in the infected plants, c PCA for upper leaves in healthy plants and sAR leaves in the infected plants. The ellipse represents the Hotelling T2 with 95% confidence in score plots. Numberings on the plot are the dates after infection (adapted from Choi et al. 2006)

screening programs (Nord et al. 2001); equivalence of genetically modified plants with non-GM plants (Fernie et al. 2004; Manetti et al. 2004; Noteborn et al. 2000; Roessner et al. 2001, 2002).

The great success of the NMR-based metabolomic studies on urine for, e.g., diagnosis of diseases (Bollard et al. 2005; Kochhar et al. 2006; Lindon et al. 2003) and studying toxicity of compounds in animal models (Gavaghan et al. 2000, 2002; Potts et al. 2001) can serve as a good example how plant metabolomics may further develop. Analyzing the much more complex metabolome of plants if compared with the various bodies fluids such as urine and serum, will be a major challenge for the coming years. Though the number of compounds that are detected in an extract is limited, still the NMR spectra give a good picture of what really is present. Minor compounds might not be seen, but the major trends are clear. By combining the NMRmetabolomic data with a more targeted approach (LC-MS, GC-MS) for those compounds that seem to be of interest one would have the broadest possible information on the metabolome. It can be compared with a microscope zooming in on smaller units. The NMR gives us the general overall view, the macroscopic view. Of course the question could be raised how many compounds can be seen in NMR. The answer is many, but the number of compounds actually present in an extract is much smaller. In fact not all the common compounds we have in our database are detected in all plant extracts. By using both 1D and 2D NMR spectrometry, one may say that about 50-100 compounds can be detected, and the absence (or below threshold) of another 50-100 can also be concluded. For the further development of the NMR platform first of all a database with all common compounds is required. But Fig. 11 Proposed metabolomic alterations in the *Nicotiana tabacum* leaves infected by TMV. • decreased, • increased, • transient increased, • previous results from other *Nicotiana* species (e.g., *N. undulata*, or *N. rustica*, *N. glutinosa*), • based on general plant biosynthesis (adapted from Choi et al. 2006)



that also requires agreement on the solvents used for the NMR-metabolomics. In our experience, buffered methanol-water (1:1) is an excellent solvent for studying a broad range of primary and secondary metabolites. This might be further completed by a nonpolar solvent, though the compounds found in such extracts are usually lipids and terpenoids. Lipids can be identified as total groups, but not as individual compounds by means of 1D NMR. For terpenoids, the possibilities for identification of individual compounds are better.

Our present database of more than 300 common compounds of plants showed to be very useful for identification of compounds in the NMR spectra. A further computerized identification using the projected 2D J-resolved spectra would be an important improvement.

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