

RESEARCH PAPER

Absolute quantification of *Medicago truncatula* sucrose synthase isoforms and N-metabolism enzymes in symbiotic root nodules and the detection of novel nodule phosphoproteins by mass spectrometry

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Abstract

Mass spectrometry (MS) has become increasingly important for tissue specific protein quantification at the isoform level, as well as for the analysis of protein post-translational regulation mechanisms and turnover rates. Thanks to the development of high accuracy mass spectrometers, peptide sequencing without prior knowledge of the amino acid sequence—*de novo* sequencing—can be performed. In this work, absolute quantification of a set of key enzymes involved in carbon and nitrogen metabolism in *Medicago truncatula* ‘Jemalong A17’ root nodules is presented. Among them, sucrose synthase (SuSy; EC 2.4.1.13), one of the central enzymes in sucrose cleavage in root nodules, has been further characterized and the relative phosphorylation state of the three most abundant isoforms has been quantified. *De novo* sequencing provided sequence information of a so far unidentified peptide, most probably belonging to SuSy2, the second most abundant isoform in *M. truncatula* root nodules. TiO₂-phosphopeptide enrichment led to the identification of not only a phosphorylation site at Ser11 in SuSy1, but also of several novel phosphorylation sites present in other root nodule proteins such as alkaline invertase (AI; EC 3.2.1.26) and an RNA-binding protein.

Key words: Absolute quantification, alkaline invertase, asparagine synthetase, *de novo* sequencing, mass western, *Medicago truncatula*, phosphoproteomics, plant proteomics, root nodules, stable isotope labelling, sucrose synthase.

Introduction

Legumes are able to establish nitrogen-fixing symbiotic relations with soil bacteria generally termed rhizobia. This symbiosis is beneficial for both partners: plants provide a carbon source and nutrients for the microsymbiont which, in return, produces a reduced form of nitrogen directly assimilable by the plant. Sucrose is the main carbohydrate transported via phloem to sink tissues like nodules, where it is metabolized by either SuSy or AI (Morell and Copeland, 1984; Flemetakis *et al.*, 2006). SuSy is a cytosolic enzyme, which catalyses the cleavage of sucrose to fructose and UDP-glucose (Okamoto and Akazawa, 1980). This enzyme is considered primarily responsible for sucrose metabolism in mature nitrogen-fixing nodules, as it has been shown to be essential for symbiotic nitrogen fixation in a variety of legume plants (Gordon *et al.*, 1999; Horst *et al.*, 2007; Kuster *et al.*, 2007). In addition, it has been demonstrated that SuSy is

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Abbreviations: AAT, aspartate aminotransferase; AI, alkaline invertase; AS, asparagine synthetase; GDH, glutamate dehydrogenase; GOGAT, glutamate synthase; GS, glutamine synthetase; LC-MS/MS, liquid chromatography-tandem mass spectrometry; Q-TOF, quadrupole time of flight; SuSy, sucrose synthase.

involved in the regulation of nitrogen fixation under different abiotic stresses, both at the transcriptional and post-transcriptional level (Gonzalez *et al.*, 1995; Gordon *et al.*, 1997; Zhang *et al.*, 1999).

SuSy has been well-characterized at the molecular level in the model legume *Medicago truncatula* (Hohnjec *et al.*, 1999, 2003). Based on transcriptional profiles, five isoforms have been described that are expressed in root nodules. However, a recent multi-dimensional LC-MS/MS-based proteomic profiling study which identified around 400 proteins from *M. truncatula* root nodules was only able to identify SuSy1, the most abundant isoform (Larrainzar *et al.*, 2007). In contrast to shotgun proteomic techniques, targeted MS approaches allow for the detection and quantification of low abundance proteins present in complex samples (Wienkoop and Weckwerth, 2006; Mallick *et al.*, 2007). While western blots are very sensitive and have been widely used for semi-quantitative protein analysis, absolute quantification can only be achieved when used in conjunction isoform specific antibodies and is subsequently scarcely used (Duncan *et al.*, 2006). Thus, the fact that protein quantification by western blots is only relative, the requirement of specific antibodies, together with the limitation in the number of proteins detectable per analysis, has led to the development of alternative LC/MS-based techniques such as the so-called 'mass western' (Lehmann *et al.*, 2008).

In this current work, a selective LC-MS/MS-based method using synthetic isotope-labelled peptides (Wienkoop and Weckwerth, 2006) has been applied for accurate quantification of SuSy proteins and other nodule metabolic enzymes (Fig. 1). Within a single analysis, several proteins involved in root nodule N metabolism as well as SuSy were detected and quantified with high sensitivity in absolute terms. Specific identification of different SuSy isoforms sharing high sequence similarity is presented. Furthermore, phosphopeptide-enrichment techniques have

been applied to analyse the state of SuSy1 phosphorylation as well as for the identification of new phosphorylation sites in several nodule proteins. A schematic overview of the workflow can be found in Fig. 1.

Materials and methods

Plant growth conditions and nodule harvest

Medicago truncatula 'Jemalong A17' plants were inoculated with *Sinorhizobium meliloti* strain 2011. Plants were grown in 1.0 l pots with a mixture of vermiculite:perlite (5:2 v:v) as substrate under controlled environment conditions (14/10 h day/night, 22/16 °C temperature, and 70/60% relative humidity). The photosynthetic photon flux was 600 $\mu\text{mol m}^{-2} \text{s}^{-1}$. When plants were 10 weeks old, root nodules were collected, frozen in liquid nitrogen and stored at -80 °C for further analysis.

Protein extraction

Nodules (0.1 g fresh weight) were homogenized in a mortar and pestle with 0.5 ml ice-cold homogenization buffer (50 mM HEPES, pH 7.8). Homogenates were centrifuged at 2000 g at 4 °C for 15 min and supernatants were collected and kept as nodule plant fractions.

Generation of stable isotope-labelled peptides

For absolute quantification of SuSy and proteins involved in nodule N assimilation, isoform-specific $^{13}\text{C}/^{15}\text{N}$ leucine-labelled peptides were synthesized (Thermo Electron, Ulm, Germany) and employed as internal standard peptides. Two approaches were used for the selection of peptides, empirical (i) and theoretical (ii).

(i) Many of the proteins that were to be targeted for quantitation had already been previously identified by MS (Larrainzar *et al.*, 2007). Consequently, peptides derived from these proteins were already known and can be found in the ProMEX spectral library (Hummel *et al.*, 2007). From this list of peptides, isoform-specific peptides were selected according to the recommendations of Thermo Electron in the instructions for the design of heavy peptide standards. The guidelines advise on sequence length and on the avoidance of certain amino acids that are common targets of uncontrolled modifications and in some cases were limiting factors for the successful selection of candidate peptides.

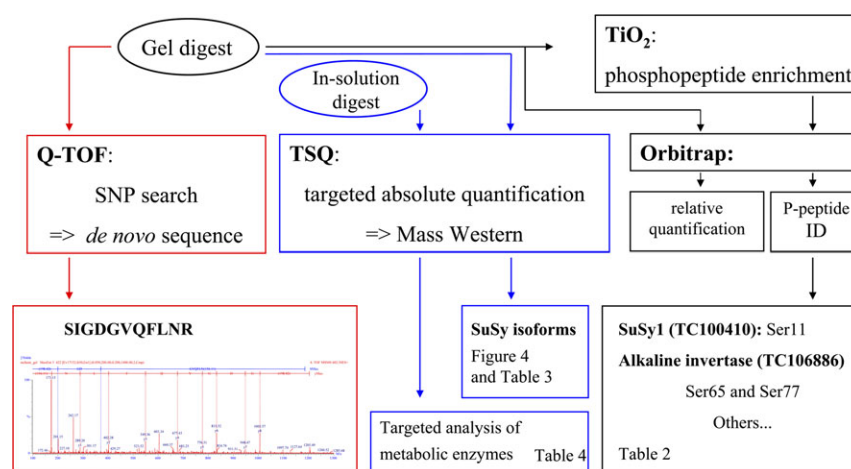


Fig. 1. Schematic workflow of the different MS analysis carried out. ID, identification.

(ii) Peptides selected for quantitation that had not been identified by MS in previous studies were designed based on proteotypic peptide prediction. In the case of the SuSy isoforms where peptides from SuSy1 have previously been identified, the peptide sequences were aligned and proteotypic peptides from the other isoforms selected based on similarity to a known peptide from SuSy1. These peptides displayed changes in amino acid composition that resulted in significant mass differences and thus a specific proteotypic peptide for each isoform (Fig. 2, framed segment).

Detailed information about peptide sequences and specific single reaction monitoring transition data can be found in Table 1. Peptide mass shifts caused by methionine oxidation reactions were below 1% and, therefore, did not significantly influence the quantification.

SDS-PAGE and in-gel protein digestion

Prior to mass western analyses, plant nodule protein extracts (80 µg of protein per sample) were separated by SDS-PAGE on 6% (w/v) polyacrylamide gels. After electrophoresis, proteins were visualized using Gel-Code Blue Stain reagent (Pierce Biotechnology, Rockford, USA). For the specific analysis of SuSy isoforms and determination of phosphorylation levels, gel bands corresponding to a molecular weight of approximately 90 kDa were cut out and standard synthetic peptides were added. For the identification of novel phosphoproteins, bands in the range between 25–60 kDa were also excised from the gels. A total number of four gel band replicates were analysed. In-gel digestion was carried out as previously described by Shevchenko *et al.* (1996).

In-solution protein digestion

Aliquots containing 50 µg of plant nodule proteins were digested overnight at 37 °C with Poroszyme immobilized trypsin beads

(1:10, v/v, Applied Biosystems, Darmstadt, Germany). After centrifugation to remove beads the peptide mixtures were desalted using SPEC C18 columns according to the manufacturer's instructions (Varian, Darmstadt, Germany). Desalted digest solutions were dried and pellets stored at –20 °C until use. Four biological replicates were analysed.

Phosphopeptide enrichment

For the identification of novel nodule phosphopeptides, gel digests corresponding to protein bands within the range of 25–60 kDa were loaded onto titanium dioxide columns (TiO₂, TopTips) purchased from SunChrom (Friedrichsdorf, Germany). Phosphopeptide enrichment was carried out as previously described by Mazanek *et al.* (2007).

Determination of phosphorylation stoichiometry

To determine the relative phosphorylation states of proteins, corresponding bands were excised from SDS-PAGE gels, digested, and analysed by MS. Relative abundance was assessed by comparing peak areas for the phosphorylated and non-phosphorylated tryptic peptides (see also 'Orbitrap settings for the identification of phosphopeptides and relative phosphorylation state analysis', and the workflow diagram in Fig. 1).

Q-TOF settings for de novo sequencing analysis

For *de novo* sequencing, gel digests corresponding to SuSy were loaded onto a HPLC system (Waters 2695 Alliance, Waters GmbH, Eschborn, Germany) directly coupled to a Q-TOF mass spectrometer (Waters GmbH, Eschborn, Germany). Peptides were eluted using a 15-min gradient from 0–80% (v/v) acetonitrile (ACN)/0.1% (v/v) formic acid (FA) using a C18 column (150×1 mm, Jupiter 4µ

TC100410 SuSy1	MATERLTVHSLKERLDETITANRNEILALLSRLEAKGKILQHHQVIAEFEEIPEDSRQ	60
TC94447 SuSy3	MAS--LTRSTSLRERFDESLTAHRIEILALLSRLEAKGKILQHHQVIAEFEEIPEEKQ	58
TC98648 SuSy4	MSTQAKYVRVPSIRDRVQDTLSAHRNELISLLSRYVAQGGKILQPHNLIDELENIHQE-D	
TC99016 SuSy5	-----	
TC95820 SuSy2	-----	
TC100410 SuSy1	KLTDGAFGEVLRSTQEAIVLPVVALAVRPRPGIWEYLRVNVHALVVENLQPAEFLKFK	120
TC94447 SuSy3	KLVNGAFGEVLRSTQEAIVLPVVALAVRPRPGVWEYLRVNVHSLVVDLRAEYLFKFK	118
TC98648 SuSy4	HLKDGPFGEIISQAQEAIVLPVFAVAVRPRPGVWEYVRVNVFELNVEQLSISEYLSFK	
TC99016 SuSy5	-----	
TC95820 SuSy2	-----	
TC100410 SuSy1	ELVDGSANGNFVLELDFEPFTASFPPTLNKSSIGNGVQFLNRHLSAKLFHDKESLHPLLE	180
TC94447 SuSy3	ELVEGSSNGNFVLELDFEPFNASFPPTLNKSSIGNGVQFLNRHLSAKLFHDKESLHPLLE	178
TC98648 SuSy4	ELVDGKINENFVLELDFEPFNASFPPTSSSIGNGVQ-----	
TC99016 SuSy5	-----	
TC95820 SuSy2	-----SIGDGVQFLNR-----	
TC100410 SuSy1	FLRLHSYKGTLMNDRIQNPDSLQHVLRKAEYLSITIDPETPYSEFHRFQEIIGLGRW	240
TC94447 SuSy3	FLRLHMYNGKTMVNDRIQNLDSLQHVLRKAEYLSITIDPETPYSEFHRFQDSGLGRW	238
TC98648 SuSy4	-----	
TC99016 SuSy5	-----	
TC95820 SuSy2	-----VEDHLSKLAPDTLYSEFETVVLQGMGFERGW	30
TC100410 SuSy1	GDTAERVLESIQLLLDLLEAPDPTLETFLDRIPMVFNVVILSPHGYFAQDDVLGYPTDG	300
TC94447 SuSy3	GDTAERVLEMIQLLLDLLEAPDPTLETFLGRIPMVFNVVILSPHGYFAQDNLGYPTDG	298
TC98648 SuSy4	-----	
TC99016 SuSy5	-----GLPDTG	6
TC95820 SuSy2	GDTAARVLETMHLILDILQAPDPTLETFLGRVPMVFNVVILSPHGFQANVLGLPDTG	
TC100410 SuSy1	GQVVYILDQVRALESEMLSRIRKQGLDIIIPRILITRLLPDAVGTTCGQRLEKVVYDTEHC	360
TC94447 SuSy3	GQVVYILDQVRALENEMLRIRKQGLDIKPRILITRLLPDAVGTTCGERLEKVVYDTEHC	358
TC98648 SuSy4	-----	
TC99016 SuSy5	GQVVYILDQVRALEFELLQKTELQGLNVKPKILVWTRILIPNAKGTTCNQELEPIIKTKHS	66
TC95820 SuSy2	GQVVYILDQVRALENEMLRIRKQGLDFTPRILIVTRILIPDAKGTTCNQRLERVSQDGYT	90

Fig. 2. Alignment of SuSy isoforms sequences of *M. truncatula*. The *de novo* peptide sequence is included in bold as part of SuSy2 isoform. Framed segment indicates the proteotypic peptides from the SuSy isoforms selected for absolute quantification.

Table 1. Triple quadrupole tune settings for the specific standard peptides used for absolute quantification

Peptide sequences employed in this work, as well as their corresponding precursor masses and product ions are given. Stable isotope-labelled leucine residues are marked in bold.

TC code	Name	Peptide sequence	Standard peptide			Native peptide		
			Precursor ion (<i>m/z</i>)	Product ions (<i>m/z</i>)	Collision energy	Precursor ion (<i>m/z</i>)	Productions (<i>m/z</i>)	Collision energy
TC100391	AS	STYAWGLEAR	580.7	738.3	19	577.2	731.3	19
TC100393	AS	STFAWGLEAR	572.8	552.2	18	569.3	545.2	18
TC106729	GS1a	ITEIAGVVVSFDPK	741.4	243.9	36	737.9	243.9	36
TC106808	GS1b	HETADINTFLWGVANR	617.6	702.3	19	615.3	702.3	19
TC106913	GS2	IHIEAYGEGNER	465.5	475.1	14	463.2	475.1	14
TC94780	GOGAT	EVLVDFDNLPLK	704.9	1166	17	701.4	1159	17
TC94631	AAT2	ATAELLGADNPAIK	752.4	785.4	22	748.9	785.4	22
TC106918	AAT1	TEEGKPLVLDVVR	487.9	917.6	19	485.6	910.6	19
TC94623	AAT	IPSGHGYDDFEVVR	534.2	744.3	18	530.9	739.3	18
TC94704	GDH	GLDIPSLLK	481.8	564.3	15	478.3	557.3	15
TC100410	SuSy1	VHSLKER	438.2	639.3	18	434.7	632.3	18
TC100410	SuSy1	ALESEMLSR	521.7	851.4	18	518.2	851.4	18
TC95820	SuSy2	ALENEMLAR	527.2	862.4	18	523.7	862.4	18
TC94447	SuSy3	ALENEMLR	491.7	791.3	17	488.2	791.3	17
TC99016	SuSy5	ALEEELLQK	540.2	888.5	18	536.7	888.5	18

Proteo 90A, Phenomenex, Aschaffenburg, Germany). *De novo* sequencing was performed using Mass Lynx software (Waters GmbH, Eschborn, Germany). The software was programmed to screen for the doubly charged precursor masses 521.7 and 602.8 of the known SuSy peptides (ALESEMLSR and SIGNGVQFLNR, corresponding to SuSy1; TC100410), a mass 527.2 for ALENEMLAR (corresponding to isoform SuSy2; TC95820) and a mass of 602.8 plus a mass shift of 0.5, in an attempt to identify a possible sequence variation due to amino acid differences encoded by single nucleotide polymorphisms (SNPs), for example, glutamine (Q)/glutamic acid (E) or asparagine (N)/aspartic acid (D) exchange.

Triple quadrupole settings for absolute quantification

For the identification and absolute quantification of different SuSy isoforms, a one-dimensional (1D) nano-flow LC-MS/MS system equipped with a pre-column (Agilent, Zeesen, Germany) was employed. Peptides were eluted using a monolithic column (Merck, Darmstadt, Germany) of 15 cm length and 0.1 mm internal diameter during a 30 min gradient from 5% to 100% (v/v) methanol/0.1% (v/v) FA with a controlled flow rate of 0.7 µl min⁻¹. 1 pmol of each internal standard peptide was added to the protein sample prior to analysis, as peptides at this concentration have been shown to lie within the linear range of detection (Wienkoop and Weckwerth, 2006). MS analysis was performed on a TSQ Quantum Discovery MAX mass spectrometer (Thermo Electron, Ulm, Germany) operated in the positive mode and tuned to its optimum sensitivity for each standard peptide as previously described (Glinski and Weckwerth, 2005). Scan width for all MRMs (multiple reaction monitoring) was 0.7 mass units, and resolution was set to 0.25 and 0.7 mass units for Q1 and Q3, respectively. The dwell time per transition was 50 ms, spray voltage was set to 1.8 kV and temperature of heated transfer capillary was set to 150 °C. Collision energies used for recorded transitions are summarized in Table 1. Absolute quantification was performed based on peak integration.

Orbitrap settings for the identification of phosphopeptides and relative phosphorylation state analysis

After phosphopeptide enrichment, TiO₂-peptide eluates were loaded on a 1D nano-flow LC-MS/MS system equipped with pre-column

(Agilent, Zeesen, Germany). A monolithic column (Merck, Darmstadt, Germany) of 15 cm length and 0.1 mm internal diameter was coupled to an Orbitrap LTQ XL mass spectrometer (Thermo Electron, Ulm, Germany). Peptides were eluted during a 50-min gradient from 5–100% (v/v) methanol/0.1% (v/v) FA with a controlled flow rate of 0.5 µl min⁻¹. For precursor masses determinations, resolution and accuracy of the Orbitrap were set to 3000 ppm and 2 ppm, respectively. MS/MS measurements were performed on wideband mode. Spray voltage was set to 1.8 kV and temperature of the heated transfer capillary was set to 150 °C. Potential phosphopeptide candidates were evaluated using MSQuant 1.4 software. A post-translational modification (PTM) score for localization, which estimates the probability to find an actual phosphorylation site based on fragment ions (Olsen *et al.*, 2006), is shown in Table 2. For the relative quantification of phosphorylation levels, gel digests were directly analysed using the same settings as described for phosphopeptide identification.

Results and discussion

Detection of different SuSy isoforms in *M. truncatula* root nodules

Although detection of three SuSy isoforms using isoform-specific antibodies has been reported in maize (Duncan *et al.*, 2006), multi-dimensional LC-MS/MS and western blot analysis did not allow detection of SuSy isoforms other than SuSy1 in *M. truncatula* root nodules (Larrainzar *et al.*, 2007). This is probably due to the relatively low abundance of the other isoforms, together with the similarity they share at the amino acid sequence level, which limits an MS/MS-based discrimination. Figure 2 represents an alignment of the N-terminal region of the five described SuSy isoform sequences. Sequences for SuSy1 and SuSy3 (TC94447) appear to be complete, whereas the known sequences for SuSy2 (TC95820),

SuSy5 (TC99016), and SuSy4 (TC98648) are estimated to be 72, 40, and 19%, respectively. To improve the detection of different SuSy isoforms in root nodules further, two approaches were carried out: (i) targeted screening for peptide sequence variation using a Q-TOF mass spectrometer with high mass accuracy; and (ii) multiple reaction monitoring on a triple quadrupole based on predicted proteotypic peptides.

Identification of a new SuSy peptide sequence

When comparing sequences of protein isoforms, it is common to find certain amino acid differences encoded by SNPs [i.e. glutamine (Q) for glutamic acid (E) or asparagine (N) for aspartic acid (D)]. These sequence variations are translated into mass differences in peptides enabling isoform discrimination using MS. Looking for this type of polymorphism, the predicted amino acid sequences of the SuSy isoforms were aligned (Fig. 2) and examined for potential candidates with this type of polymorphism. A SuSy1 tryptic peptide containing some of the above-mentioned residues was subsequently chosen (SIGNGVQFLNR). After screening the gel sample for masses corresponding to the doubly charged peptide, two ions were selected: one corresponding to a mass-to-charge ratio (m/z) of 602.8 derived from a SuSy1 tryptic peptide, and the other peptide of unknown sequence with an m/z

of 603.3. *De novo* sequencing was performed on the unknown ion and was identified as the peptide (SI)GDGVQFLNR (Fig. 3), which did not match any of the previously reported sequences (Fig. 2). The observed mass difference corresponds to a shift starting from ion y_7 , explained by a replacement of D instead of the original N residue present in SuSy1. The only other possible combination producing a 603.3 m/z peptide would be the replacement of isoleucine (I) for leucine (L) in the N-terminal region. However, as this region appears to be well-conserved in the amino acid sequence of SuSy in different legume plants (data not shown), this alternative appears to be less likely. Furthermore, peptides derived from both SuSy1 and SuSy2 were detected during the analysis of 1D-PAGE samples (see Materials and methods and Fig. 2), further supporting the hypothesis that the new peptide is driven from SuSy2 (Fig. 2). Interestingly, SuSy2 is the second most abundant isoform in *M. truncatula* root nodules at the transcript level (Hohnjec *et al.*, 2003).

Mass western of different SuSy isoforms in root nodules

For the specific absolute quantification of protein isoforms, a technique based on high resolution multiple reaction monitoring in a triple quadrupole mass spectrometer, called mass western, has recently been established (Lehmann *et al.*, 2008). Instead of antibodies, isoform-specific stable isotope-labelled peptides are synthesized. These are then employed as internal standards and added to protein samples, which are subsequently digested either in gel or directly in solution (Wienkoop and Weckwerth, 2006). As the concentration of the standard peptides is known, absolute quantification of native proteins is achievable in $\text{fmol } \mu\text{g}^{-1}$ of total protein.

In the present study, four standard peptides specific for each of the isoforms SuSy1, 2, 3, and 5 were selected (Table 1). As only 19% of the SuSy4 sequence is

Table 2. Phosphopeptides found in *M. truncatula* root nodules after TiO_2 -enrichment and MSQuant validation (PTM: post-translational modification)

Protein	Peptide sequence	Phospho site	Mascot score	MSQuant PTM score
TC106886	AGLDNYDNYS(p)PGGR	Ser65	69	114
	SGFNTPAS(p)SAR	Ser77	48	73
TC101080	KHASS(p)PPDSPSQ	Ser21	53	60
	DSVEKPTYVR			
TC97295	HQSAAT(p)PTPTAGAR	Thr33	40	70

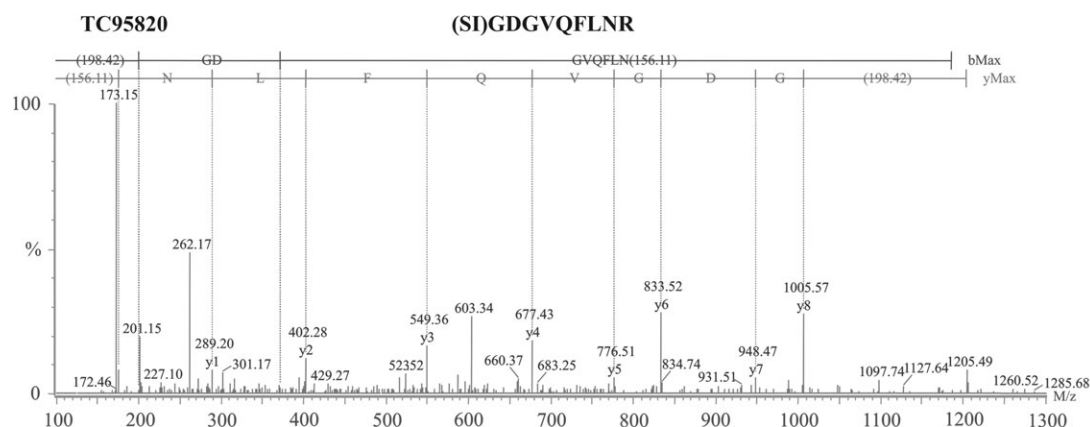


Fig. 3. Q-TOF driven MS/MS spectrum of the *de novo* sequence for a tryptic peptide with m/z 603.3 (SIGNGVQFLNR).

available (Fig. 2), this isoform was excluded from the analysis. Out of the four isoforms analysed, absolute quantification was only possible for three of them (Table 3; Fig. 4). SuSy5 appeared either to be below the detection limit or not present in nodules under the conditions investigated. Results show that SuSy2 is the second most abundant isoform present in *M. truncatula* root nodules. This finding further supports the hypothesis that the *de novo* peptide sequence belongs to this second

Table 3. Absolute quantification of three different SuSy isoforms in *M. truncatula* root nodules after in-gel digestion (n=4)

TC Code	Name	Specific amount ^a		
		fmol µg ⁻¹ protein	pmol per gel slide	ng per gel slide
TC100410	SuSy1	160±2.21	12.8	1000
TC95820	SuSy2	2.1±0.03	0.170	16
TC94447	SuSy3	0.3±0.25	0.023	2
TC99016	SuSy5	n.d.	n.d.	n.d.

^a n.d., Not detected.

isoform. Due to the high sensitivity of this technique, SuSy3 was also successfully identified, despite its much weaker signal (Fig. 4). Absolute quantification of the different isoforms shows that the protein ratios for SuSy1:SuSy2:SuSy3 are 1:0.013:0.0018 (Table 3). These results are in agreement with a recent transcript analysis in pea nodules (Marino *et al.*, 2008), where the ratio for the three isoforms was 1:0.002:1.7 × 10⁻⁵. From a technical point of view, the four orders of magnitude range determined in a single analysis corresponds to the current maximum capacity for a MS-based analysis (Wienkoop *et al.*, 2006; Wienkoop and Weckwerth, 2006). This limitation is mainly due to the limit of detection of each peptide and the loading capacity of the columns. Nevertheless, the sensitivity (2 ng) in the present study has been shown to be comparable to those of western blots (Lehmann *et al.*, 2008). A significant advantage of the mass western is that it allows for absolute quantification and for multiple targets to be analysed in a single run. Furthermore, mass western provides a direct measurement of protein concentration, not available by transcript analysis.

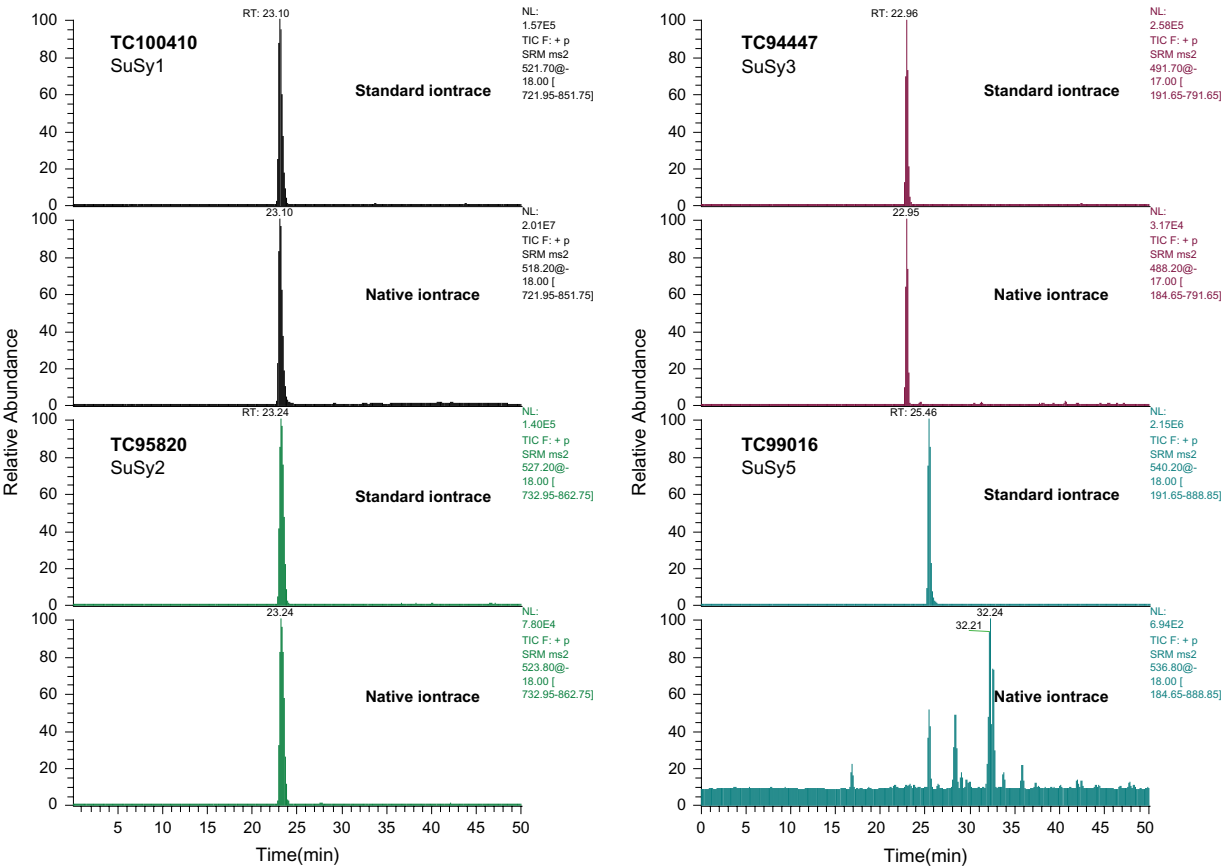


Fig. 4. Mass western of four SuSy isoforms detected in *M. truncatula* root nodule protein extract using a triple quadrupole mass spectrometer and specific stable isotope-labelled standard peptides (see Table 1).

Absolute quantification of proteins involved in N metabolism present in complex nodule protein samples

Absolute quantification enables the analysis of complete metabolic pathways in complex protein samples in a single analysis. In contrast to the previous analysis, where the starting materials employed were in-gel digests, absolute quantification was also carried out using nodule extract protein samples as the starting material. Internal standard peptides for glutamine synthetase (GS), glutamate dehydrogenase (GDH), asparagine synthetase (AS), and aspartate aminotransferase (AAT) were synthesized (peptide sequences shown in Table 1), and these metabolic enzymes, together with SuSy, were subjected to absolute quantification (Table 4). Regarding SuSy, only the most abundant isoform, SuSy1, representing about 1% of the total nodule proteins, was detected. The reason for this might be a higher signal-to-noise ratio of a complex protein sample compared to the pre-fractionated gel sample used in the previous analysis. However, the absolute amount of SuSy1 appears to be slightly higher in the complex nodule protein analysis indicating that the in-gel analysis may be even more sensitive.

Most of these enzymes involved in nodule N metabolism (GS, GDH, AAT) are well characterized and their activities are routinely assessed (Vance, 2008). Although most of the N exported from temperate climate nodules is in the form of asparagine, detection of AS activity in plant tissues is extremely difficult (Lea *et al.*, 2007). The reasons for this are not clear, but they may be related to the presence of inhibitors (Joy *et al.*, 1983), asparaginase activity (Streeter, 1977) or protein instability (Huber and Streeter, 1985). To overcome these limitations, quantification of AS gene expression has been suggested to be a good semi-quantitative indicator of its activity levels, showing a good correlation with asparagine:aspartate ratios transported to the shoot via the xylem (Antunes *et al.*, 2008). The strategy shown in the present paper may

provide a more accurate alternative for the quantification of AS in nodules.

Identification of phosphorylation sites and determination of the phosphorylation state of SuSy1 in nodules under normal growth conditions

SuSy has been shown to be phosphorylated in a number of tissues in different plants. While initial evidence that SuSy was phosphorylated came from work on maize cell cultures (Shaw *et al.*, 1994), it was Huber *et al.* (1996) who first characterized a phosphorylation site at Ser15 in SuSy isolated from maize leaves. Further studies have described other phosphorylation sites in maize (Hardin *et al.*, 2003) and legume plants like soybean (Zhang and Chollet, 1997; Zhang *et al.*, 1999; Komina *et al.*, 2002). In particular, Zhang *et al.* (1999) described a putative seryl-phosphorylation domain near the N-terminus of the protein (LTRVHS¹¹LRER) in soybean. This motif, which was reported to be the primary phosphorylation site of nodule soybean SuSy *in planta*, is present in practically all the SuSy sequences reported to date.

In the present work, MS analysis revealed that SuSy1 from the nodules of *M. truncatula* is also phosphorylated at Ser11, the same residue identified as phosphorylated in soybean and homologous to Ser15 in maize SuSy1 (VHS¹¹LKER; spectra can be found in ProMEX database <http://promex.mpimp-golm.mpg.de/home.shtml>). In addition, the relative phosphorylation state of the protein was analysed and quantified by comparing peak areas of the phosphorylated and non-phosphorylated tryptic peptides derived from the same analysis (see also 'Determination of phosphorylation stoichiometry', and the workflow diagram in Fig. 1).

M. truncatula nodule SuSy1 was found to be hyperphosphorylated, as the phosphopeptide was 1.7-fold more abundant than the non-phosphorylated peptide. The meaning of this hyperphosphorylation state remains the subject of much debate. In soybean, *in vitro* studies suggest that nodule SuSy sucrose-cleavage activity is not dependent on the phosphorylation state of the protein, but rather influences the attachment of this enzyme to the membrane (Zhang *et al.*, 1999). However, studies in maize have shown that leaf SuSy is activated when phosphorylated at its primary site (Ser15), while phosphorylation at Ser170 would trigger degradation of the enzyme (Hardin *et al.*, 2003, 2004). More recently, it has been suggested that sucrose concentration and phosphorylation may regulate SuSy oligomerization and F-actin association in maize kernels and seedlings (Duncan and Huber, 2007). Regarding membrane attachment, Komina *et al.* (2002) investigated the phosphorylation state of membrane and soluble SuSy forms of nodules in soybean. Using several non-MS-based approaches, the microsomal SuSy fraction was found to be hypophosphorylated

Table 4. Absolute quantification of several proteins involved in C and N metabolism in *M. truncatula* root nodules after gel free shotgun analysis

TC Code	Name	fmol mg ⁻¹ protein ^a
TC100410	SuSy1	195.2±23.7
TC95820	SuSy2	n.d.
TC94447	SuSy3	n.d.
TC100391	AS	76.5±2.5
TC100393	AS	90.5±8.5
TC106729	GS1a	98.9±1.0
TC106808	GS1b	29.6±8.6
TC106913	GS2	23.8±5.8
TC94631	AAT2	44.3±0.9
TC106918	AAT1	5.1±0.9
TC94623	AAT	20.7±4.1
TC94704	GDH	51.1±9.3

^a n.d., Not detected.

relative to the soluble form (on an equivalent SuSy protein base). Although abiotic stresses seem to have a negative impact on SuSy abundance, as well as on its phosphorylation levels (data not shown), the physiological meaning of a state of hyper- or hypophosphorylation remains to be elucidated.

Other phosphoproteins found in nodules

SuSy is one of the most characterized nodule proteins in legumes. Due to its relatively high abundance in root nodules and, specifically with regard to *M. truncatula*, its hyperphosphorylation state, it is understandable that the conserved Ser11 phosphopeptide has been described several times. In order to identify novel putative phosphorylated proteins in *M. truncatula* root nodules, a phosphopeptide enrichment strategy based on TiO₂-affinity chromatography was carried out. As a result, several other phosphopeptides were identified (Table 2). Among them, two phosphopeptides (AGLDNYDNYS⁶⁵PGGR and SGFNTPAS⁷⁷SAR) belonging to AI (TC106886), the other enzyme with a sucrose-cleavage activity in nodules. In contrast to SuSy, relative peak area quantification of both peptides indicated that AI is mostly hypophosphorylated in nodules. The non-phosphorylated to the phosphorylated forms of Ser65 and Ser77 were determined to be in a ratio of 1:0.01:0.03, respectively. It has been suggested that AI is post-transcriptionally regulated through an interaction with a phospho-binding 14-3-3 protein in developing barley grains (Alexander and Morris, 2006). However, to our knowledge this is the first report where the actual phosphorylation sites of the protein are described. Several other phosphopeptides were found that belong to a putative RNA-binding protein (TC101080) and a hypothetical protein (TC97295), which is similar to a universal stress response protein described in *Arabidopsis thaliana* (At4g27320) (Table 2). Although, thus far, little is known about the function of these proteins in nodules, they may be considered interesting candidates for further studies given the roles of phosphorylation in metabolic regulation.

Nevertheless, several other nodule proteins have been suggested to be phosphorylated and were not identified in this present study. This is the case for both GS (Lima *et al.*, 2006) and phosphoenolpyruvate decarboxylase (Zhang *et al.*, 1995) from *M. truncatula*. Previously reported phosphopeptides from GS did not appear to be detectable after tryptic digestion, but may be detected using different proteolytic enzymes. In the case of phosphoenolpyruvate decarboxylase, phosphopeptides were not detected using the current approach as its molecular weight was out of range of the analysed bands subjected to TiO₂ enrichment. Therefore, future analysis may include a wider range of molecular weights to identify new nodule protein subject to protein phosphorylation.

Conclusion

This study has demonstrated for the first time the absolute quantification of three distinct *M. truncatula* SuSy isoforms in an independent tissue-specific manner. It is shown that, besides the nodule-enhanced SuSy1, two other protein isoforms are present in root nodules at quantifiable amounts. *De novo* sequencing allowed for the identification of a new peptide sequence, most probably driven from the second most abundant isoform SuSy2, which complements the existing sequence information based on transcript data. Absolute quantification of N metabolism enzymes was carried out, enabling the discrimination of different protein isoforms from a single analysis, even when those isoforms were present at very low amounts. Nodule SuSy1 was shown to be hyperphosphorylated at Ser11 in *M. truncatula* and novel phosphoproteins present in nodules were identified. All peptides found in this study can be viewed in the ProMEX spectral library (Hummel *et al.*, 2007).

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