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Phloem small RNAs, nutrient stress responses, and systemic mobility

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Abstract

Background: Nutrient availabilities and needs have to be tightly coordinated between organs to ensure a balance between uptake and consumption for metabolism, growth, and defense reactions. Since plants often have to grow in environments with sub-optimal nutrient availability, a fine tuning is vital. To achieve this, information has to flow cell-to-cell and over long-distance via xylem and phloem. Recently, specific miRNAs emerged as a new type of regulating molecules during stress and nutrient deficiency responses, and miR399 was suggested to be a phloem-mobile long-distance signal involved in the phosphate starvation response.

Results: We used miRNA microarrays containing all known plant miRNAs and a set of unknown small (s) RNAs earlier cloned from *Brassica* phloem sap [1], to comprehensively analyze the phloem response to nutrient deficiency by removing sulfate, copper or iron, respectively, from the growth medium. We show that phloem sap contains a specific set of sRNAs that is distinct from leaves and roots, and that the phloem also responds specifically to stress. Upon S and Cu deficiencies phloem sap reacts with an increase of the same miRNAs that were earlier characterized in other tissues, while no clear positive response to -Fe was observed. However, -Fe led to a reduction of Cu- and P-responsive miRNAs. We further demonstrate that under nutrient starvation miR399 and miR395 can be translocated through graft unions from wild type scions to rootstocks of the miRNA processing *hen1-1* mutant. In contrast, miR171 was not transported. Translocation of miR395 led to a down-regulation of one of its targets in rootstocks, suggesting that this transport is of functional relevance, and that miR395, in addition to the well characterized miR399, could potentially act as a long-distance information transmitter.

Conclusions: Phloem sap contains a specific set of sRNAs, of which some specifically accumulate in response to nutrient deprivation. From the observation that miR395 and miR399 are phloem-mobile in grafting experiments we conclude that translocatable miRNAs might be candidates for information-transmitting molecules, but that grafting experiments alone are not sufficient to convincingly assign a signaling function.

Background

The levels of essential inorganic nutrients have to be tightly controlled inside individual cells and organs, but information about nutrient uptake and needs also have to be transferred between organs to optimize nutrient allocation, especially in plants growing under sub-optimal conditions. If an organ experiences nutrient starvation, it needs to communicate its requirements to the other organs in order to increase nutrient uptake or reallocate resources. This type of communication is probably mediated via the phloem. Recent work showed that microRNA (miRNA) 399 is potentially involved in long-distance communication via the phloem following phosphate deprivation [1-3]. miRNAs are short (21-24 nt), nontranslated RNAs that are processed by Dicer-like proteins from large, characteristically folded precursor molecules. The majority of plant miRNAs target transcription factors and is therefore thought to mainly regulate developmental processes. However, recent studies have also identified miRNAs that are involved in responses to nutrient deficiencies. As mentioned earlier, miR399 is strongly induced during phosphate deprivation [4-7], while miR395 drastically increases under growth on low sulfur [8]. In addition to macronutrients like sulfur and phosphate, also a lack of the micronutrient copper leads



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to an accumulation of miR397, 398, 408, and 857 [9-11]. miRNAs 395, 398 and 399 were recently shown to accumulate not only on the whole plant level, but also strongly within the phloem [1]. Since sRNAs accumulating in phloem sap under stress could represent potential longdistance signaling molecules, we used sRNA microarrays from LC Sciences to comprehensively analyze phloem sRNAs. The customized arrays contained, in addition to all known plant miRNAs, a subset of small RNAs (sRNAs) of unknown function that was earlier sequenced from phloem sap of Brassica napus [1]. First we established the miRNA patterns of phloem, leaves and roots of fully nutrient supplied, hydroponically grown oilseed rape plants to subsequently identify candidates that respond to growth under S, Cu or Fe deficiency, respectively. In addition, we used the highly -S induced miR395 as an example to examine whether this specific miRNA can be transported over graft unions when combining WT Arabidopsis with the miRNA biosynthesis mutant hen1-1. The specific aims were 1) to find phloem- and organ-enriched miRNAs, 2) to identify additional miR-NAs that respond to S and Cu deficiencies, 3) to examine whether any miRNAs respond to Fe starvation, and 4) to demonstrate whether miR395 is phloem mobile or not.

Results and Discussion

Phloem sap shows a specific sRNA pattern that is distinct from that of inflorescence stem, leaves and roots

To ensure that the sRNAs observed in phloem sap were not resulting from contamination during sampling, and in order to identify phloem-enriched sRNAs, we performed a microarray hybridization experiment comparing phloem sap to the surrounding inflorescence stem tissue. This resulted in the identification of phloemenriched sRNAs, while others were less abundant in phloem sap than in stem tissue (including phloem) collected after phloem sampling from the sampling site. Signal values for one miRNA per family are depicted in additional file 1. The distribution of ten miRNAs was reevaluated by RNA gel blots from an independent set of plants, what confirmed the microarray results. miRNAs 162, 167, 168, 169, and 399 strongly accumulated in phloem samples as compared to inflorescence stem samples, while miR158, 396 and 397 were stem-enriched. This indicates that phloem samples are not significantly contaminated by the contents of the surrounding inflorescence stem cells, what had already previously been demonstrated [1,12]. The observation that miR167 accumulates in phloem sap confirms an earlier study in pumpkin that found miR167 20-fold enriched in phloem sap as compared to the surrounding vascular tissue [13]. Also the failure to detect miR171 in phloem sap and its low expression in stem samples is in accordance with earlier findings [13,14].

We further used the microarrays to identify sRNAs that preferentially accumulated in phloem sap as compared to leaf and root samples. To this end we grew plants under full nutrition (FN) conditions in three successive, completely independent experiments and compared the sRNA amounts in phloem samples with that of leaves and roots. For inter-array comparisons, signal intensities were normalized to the median signal of each sample. This approach allowed the detection (signal >100) of 161 miR-NAs belonging to 37 families in phloem sap, covering all 17 miRNA families earlier detected in samples from soilgrown Brassica plants by high-throughput pyrosequencing [1] (indicated by the numbers of sequences obtained in additional file 1). In addition, we found several miR-NAs on the arrays that were not identified by the sequencing approach, suggesting that these miRNAs were either not present in soil-grown plants or not identified, possibly due to their low abundance or absence in the steadily growing databases at the earlier time-point of data analysis. A reasonable reproducibility between the experiments was achieved, given that they were completely independent and that miRNAs are known to be strongly influenced by developmental stage and growth conditions [15]. Signal intensities and standard deviations for one representative of each family are depicted in additional file 2. Statistical evaluation using the Students ttest revealed miRNAs that were significantly (p < 0.05)enriched in phloem, leaves or roots (figure 1). miRNAs from four families were more abundant in phloem sap than in leaves and roots under FN, namely miR169 (not statistically significant), 390, 829, 894, and 1132 (not significant) (figure 1). miR1132, together with miR1134 (misnamed miR518), was cloned from wheat [16] and recently from Brachypodium [17]. Both miRNAs are not well characterized, thought to be species-specific, and their possible functions are unknown. However, signal values were well above the microarray noise. Nevertheless this result does not allow a conclusion on whether these miRNAs really occur in Brassica or if the signals represent an artifact (e.g. unspecific cross-hybridization) caused by the microarray technique.

Except for miR390, these miRNAs were also phloemenriched as compared to inflorescence stem tissue (additional file 1). miRNAs from the families 156, 159, 160, 162, 164, 165, 166, 167, 393, 394, 396 and 403 were less abundant in the phloem as compared to both, leaves and roots. However, some of these miRNAs (159, 162, and 167) were more abundant in the phloem than in the surrounding stem.

miRNAs from the complete 156, 160, 166, 393, 396, and 528 families were found to be significantly enriched in roots as compared to leaves and phloem. In rice, miRNAs 156 and 166 have earlier been shown occur at higher levels in roots than in leaves [18]. In addition, miR166 has

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miR	P/L	P/R	L/R	miRNA
156	-0.7	-4.3 ***	-3.6 ***	ath-miR156a
	-0.7	-4.3 ***	-3.6 ***	ath-miR156g
	-1.4			ath-miR156h
		-3.3	-2.0	
	-0.9 *	-4.5	-0.0	bna-miR156a
	0.8	-1.5	-2.3 *	gma-miR156b
	-0.9	-4.5 *	-3.6 *	osa-miR156l
	-1.8	-1.3	0.5	pta-miR156a
	-1.6	-1.2	0.4	pta-miR156b
	0.7	-3.0	-3.7	ptc-miR156k
		-4.3 **	-3.4 **	
	-0.9			sbi-miR156e
	-1.4	-1.5	-0.1	smo-miR156b
	-1.5	-3.2	-1.6	smo-miR156c
	1.3	0.0	-1.3	smo-miR156d
159	-1.3 **	-0.6	0.7	ath-miR159a
	-1.3 **	-0.6	0.7	ath-miR159b
	-1.6 ***	-0.8	0.8	ath-miR159c
	-1.4 *	-0.6	0.7	osa-miR159a
	1.4		-	
	-1.0	-0.8	0.9	osa-miR159c
	-1.6 **	-0.8	0.8	osa-miR159d
	-1.7 **	-0.8	0.9	osa-miR159e
	-1.4 *	-0.6	0.8	osa-miR159f
	-2.0 ***	-1.2	0.9	pta-miR159a
	-2.4	-1.8	0.6	pta-miR159b
	-3.8 *	-5.9	-2.0	ptc-miR159e
	-2.0 *			•
	2.0	-1.3	0.7	ptc-miR159f
	-1.0	-0.7	0.9	ptc-miR159d
	-1.9 ***	-1.0	1.0	sof-miR159e
160	-2.4	-4.4	-2.0	ath-miR160a
	-2.5	-4.4 *	-1.9	osa-miR160e
	-2.3	-4.4 *	-2.1 *	ppt-miR160b
	-2.4	-4.6 *	-2.2	ppt-miR160c
	-2.4	-4.6	-2.2	ppt-miR160d
			-3.4	
	-3.0	-6.4		ppt-miR160g
	-3.0	-6.5	-3.5	ppt-miR160h
	-2.3	-4.4	-2.1	ptc-miR160g
	-2.3	-4.3	-2.0	ptc-miR160h
162	-1.7 *	-1.1	0.6	ath-miR162a
	-2.4	-1.7	0.7	osa-miR162b
	-2.1 *	-1.6	0.5	zma-miR162
164	-1.8 **	-2.3 **	-0.6	ath-miR164a
104	-1.0	-2.0		
	-1.7 **	-2.0	-0.6	ath-miR164c
	-1.6 ***	-2.1 **	-0.5	osa-miR164c
	-1.8 **	-2.3 **	-0.5	osa-miR164d
	-1.6 ***	-2.1 **	-0.4	osa-miR164e
	-1.7 ***	-2.1 **	-0.4	ptc-miR164f
	-1.7 ***	-2.2 **	-0.5	sbi-miR164c
165	-2.8 *	-4.8 *	-2.0	ath-miR165a
166	-2.6 **	-4.1 *		
	-2.0		-1.6	ath-miR166a
100	-2.7 *	-4.5	-1.8	osa-miR166e
100			-2.0	osa-miR166i
100	-4.5	-6.5		osa-miR166k
100		-5.0	-1.9	05a-1111100K
100	-4.5 -3.1	-5.0		
100	-4.5 -3.1 -2.6 **	-5.0 -4.0	-1.4	osa-miR166m
100	-4.5 -3.1 -2.6 ** -3.0 *	-5.0 -4.0 -5.0	-1.4 -2.1	osa-miR166m ppt-miR166j
100	-4.5 -3.1 -2.6 ** -3.0 * -4.5	-5.0 -4.0 -5.0 -6.6	-1.4 -2.1 -2.1	osa-miR166m ppt-miR166j ppt-miR166m
100	-4.5 -3.1 -2.6 ** -3.0 * -4.5 -4.2	-5.0 -4.0 -5.0 -6.6 -7.9	-1.4 -2.1 -2.1 -3.7	osa-miR166m ppt-miR166j ppt-miR166m pta-miR166c
100	-4.5 -3.1 -2.6 ** -3.0 * -4.5 -4.2 -2.7 *	-5.0 -4.0 -5.0 -6.6 -7.9 -4.3	-1.4 -2.1 -2.1 -3.7 -1.7	osa-miR166m ppt-miR166j ppt-miR166m
100	-4.5 -3.1 -2.6 ** -3.0 * -4.5 -4.2	-5.0 -4.0 -5.0 -6.6 -7.9	-1.4 -2.1 -2.1 -3.7	osa-miR166m ppt-miR166j ppt-miR166m pta-miR166c

niR	P/L	P/R	L/R	miRNA
167	-2.5 **	* -2.4 **	0.1	ath-miR167a
	-3.0 **		0.0	ath-miR167c
	-2.5 **		0.1	ath-miR167d
	-2.6 **		0.2	ppt-miR167
	-2.5 **		0.2	ptc-miR167f
	-2.3	-2.4	-	'
	-2.7 **		0.4	ptc-miR167h
168	0.5	0.5	0.1	ath-miR168a
	0.6	0.6	0.0	osa-miR168a
	0.8	0.7	-0.1	osa-miR168b
	0.9 *	0.9	0.1	sof-miR168b
169	5.5	5.9	0.4	ath-miR169a
	5.1	5.4	0.3	ath-miR169b
	5.7	5.8	0.1	ath-miR169d
	4.7	5.6	0.9	ath-miR169h
	7.3	2.7	-4.6	osa-miR169d
	5.2	5.5	0.4	osa-miR169e
	7.8	3.1	-4.6	osa-miR169n
	4.6	-2.6 *	-7.2 *	osa-miR169q
	3.6	4.3	0.7	ptc-miR169ab
	4.7	5.4	0.7	ptc-miR1690
	4.4	4.5	0.1	ptc-miR169q
	5.3	5.9	0.6	ptc-miR169s
	6.3	5.0	-1.2	ptc-miR169t
	4.0	4.8	0.8	ptc-miR169u
	4.4	4.8	0.4	ptc-miR169v
	4.4	4.3	-0.1	ptc-miR169x
	3.3	-3.0	-6.3	zma-miR169d
	-0.9	-3.3	-2.4	zma-miR169e
319	-1.8			
519		-1.0	0.8	ath-miR319a
	-1.7	-0.8	0.9	ath-miR319c
	-1.8	-0.8	1.0	gma-miR319a
	-1.3	-1.2	0.1	gma-miR319c
	0.7	-0.5	-1.2	osa-miR319a
	-1.6 **	-0.6	1.0	ppt-miR319a
	-1.7	-1.1	0.6	pta-miR319
	-1.7 **	-0.7	1.0	ptc-miR319e
	1.8	0.1	-1.6	ptc-miR319i
390	1.5 *	1.8 *	0.4	ath-miR390a
	0.6	0.1	-0.5	pta-miR390
	1.2	1.7 *	0.5	ppt-miR390c
391	-3.3	1.6 *	4.9 ***	ath-miR391
393	-2.2	-4.1	-1.9	ath-miR393a
	-2.0	-3.8	-1.7	bna-miR393
	-1.7	-4.0 *	-2.3	osa-miR393b
394	-1.2	-1.0	0.2	ath-miR394a
396	-8.2	-9.3	-1.1	ath-miR396a
	-11.7	-12.9	-1.2	ath-miR396b
	-5.0 *	-9.3	-4.3	osa-miR396d
	-2.3	-6.5	-4.2	ptc-miR396f
	0.0	-5.5	-5.5	ptc-miR396g
103	-3.0 **		0.6	ath-miR403
528	3.1	-1.1	-4.2 **	osa-miR528
	3.1 3.5 *			
329		2.9	-0.6	ath-miR829.1
394	<u> </u>	1.4	-1.2	ppt-miR894
1132	3.0	0.1	-3.0 *	tae-miR1132
				npared organ
				pared organ

Figure 1 List of miRNAs that were enriched in phloem, leaves or roots, respectively, in plants grown under full nutrition. Only families where at least one member showed a statistically significant differential accumulation in one organ are shown (p < 0.05, n = 3). Values are log2s between P/L: phloem vs. leaves, P/R: phloem vs. roots and L/R: leaves vs. roots. Markedly (log2 values >1 or <-1, indicating a two-fold difference) phloem-enriched miRNAs are marked in blue, leaf-enriched in green, and root-enriched in red. The statistical significance is indicated as: * p < 0.05; ** p < 0.01; *** p < 0.001.

been described to be expressed in roots of *Medicago truncatula*, where it functions in root and nodule development [19]. In Arabidopsis, miRNAs 156 and 160 occur root-enriched [20], and miR160 has been implicated with root development [21,22].

miR391 was the only miRNA that accumulated in leaves as compared to roots and phloem sap (figure 1). In an earlier study, miR391 was found to appear preferentially in rosette leaves of Arabidopsis, as compared to seedlings, flowers and siliques [23]. According to the same publication, miR391 targets a beta-fructofuranosidase, but its function is currently not well understood. Although miR391 is regarded as being related to miR390, differing in only 5 nt [24], both miRNAs showed a quite distinct organ distribution: while miR391 was clearly leafenriched, miR390 was slightly, but significantly phloemenriched, indicating that both miRNAs might still have distinct localizations and functions.

Interestingly, the unknown sRNAs represented on the chip were, except for Bn_PsRNA_24, significantly more abundant in phloem sap as compared to leaves and roots (figure 2). All Bn_PsRNAs were additionally more abundant in roots than in leaves. Most of these differential unknown sRNAs had a length of 24 nt, and only five had a length of 21 nt characteristic for miRNAs (figure 2). Precursor and target predictions using mfold and psRNA-Target, respectively (data not shown), provided no conclusive evidence that any of these sRNAs could represent a novel miRNA following recently published criteria [25]. On the one hand, the inability to successfully predict targets and precursors of the Brassica sRNAs could be due to the limited EST genome sequence of Brassica napus publicly available. On the other hand, it could indicate that they are no miRNAs, but rather siRNAs, as yet unclassified sRNAs, or breakdown products of larger RNAs. However, the observation that they accumulate in phloem sap makes them interesting candidates for future studies.

Phloem small RNA patterns change under nutrient deficiency

Since three miRNAs, miR395, 398 and 399, had been previously shown to accumulate in the phloem under the corresponding nutrient stress conditions [1], we intended to identify additional nutrient-responsive phloem sRNAs. They could represent novel information transmitters during nutrient deprivation, as has been suggested for miR399 under phosphate deficiency [2]. To induce nutrient deprivation, we raised *Brassica napus* plants in hydroponic cultures under FN and omitted the respective nutrient from the medium for two (-S, -Cu experiments) or three weeks (-Fe experiment), respectively, before samples were collected. Under -S and -Cu conditions the plants did not show any obvious stress symptoms at the

sRNA	sRNA P/L P/B L/B							
Bn PsRNA 03	7/∟ 3.8 *	- ,	-1.7 *					
		<mark>2.1</mark> 0.2						
Bn_PsRNA_04	3.4 3.6 *	0.2 3.5 *	-5.1					
Bn_PsRNA_05			-0.1					
Bn_PsRNA_07	5.7 *	-0.7	-6.4					
Bn_PsRNA_10	6.9 **	0.1	-3.8					
Bn_PsRNA_20	7.0 ***	1.1	-5.9					
Bn_PsRNA_24	-1.4 *	-4.0	-2.6					
Bn_PsRNA_26	3.0 *	1.3	-1.6					
Bn_PsRNA_27	3.2 *	1.1	-2.1					
Bn_PsRNA_29	2.4 *	1.3	-1.1					
Bn_PsRNA_31	5.3 *	3.3 *	-2.0					
Bn_PsRNA_35	5.6 *	3.6 *	-2.0					
Bn_PsRNA_41	4.4 *	3.4 *	-0.9					
Bn_PsRNA_47	4.9 *	3.8 *	-1.2					
Bn_PsRNA_56	5.1 *	<mark>3.5</mark> *	-1.6					
Bn_PsRNA_57	3.1 *	1.8	-1.3					
Bn_PsRNA_65	7.9 **	2.2 *	-5.7					
Bn_PsRNA_67	4.0 *	3.4 *	-0.6					
Bn_PsRNA_69	5.4 *	3.7 *	-1.7					
Bn_PsRNA_72	6.9 *	3.1 *	-3.7					
Bn_PsRNA_83	6.2 *	<mark>5.8</mark> *	-0.4					
higher in phloem than in the compared organ higher in leaves than in the compared organ higher in roots than in the compared organ								
Figure 2 List of unknown sRNAs that were organ-enriched grown under full nutrition. List of unknown sRNAs, sequenced from <i>Brassica</i> obloem sap [1]. that showed statistically significant differences be-								

under full nutrition. List of unknown sRNAs, sequenced from *Brassica* phloem sap [1], that showed statistically significant differences between phloem sap, leaves and roots, respectively (p < 0.05, n = 3). Values are log2s between P/L: phloem vs. leaves, P/R: phloem vs. roots and L/R: leaves vs. roots. Markedly (log2 values >1 or <-1, indicating a two-fold difference) phloem-enriched miRNAs are marked in blue, leaf-enriched in green, and root-enriched in red. The statistical significance is indicated as: * p < 0.05; ** p < 0.01; *** p < 0.001.

time of sampling. However, omitting Fe led to chlorosis symptoms in very young upper leaves after 4-5 days of stress (data not shown).

Initial analysis of the expression of selected genes that are known to be altered by the respective nutrient stress clearly confirmed that the plants were nutrient deficient in all three kinds of stress experiments performed (additional file 3). As expected, S starvation led to an increase in the expression of the two high-affinity sulfate transporters st1 (AJ416460) and st2 (AJ311388), especially in roots. Copper deprivation was confirmed by a slight decrease in the amount of Cu-Zn SOD transcripts, while the amount of the high-affinity copper transporter COPT1 increased markedly. Fe deprived plants showed only a slight reduction in the expression of the iron storage protein ferritin LSC30 in leaves and roots, accompanied by an increase in the transcript of the root-specific iron transporter IRT1 in roots (additional file 3).

Subsequently, material from the same batch of plants was used for dual-color microarray hybridizations of stressed and FN samples. Since only one array per stress experiment was hybridized, we applied specific criteria to only identify the most drastic positive changes (>fourfold increases, log2 >2) upon stress treatments and furthermore restricted the analyses to abundant sRNAs with signal intensities of >100 in one of the two (FN or stressed) samples.

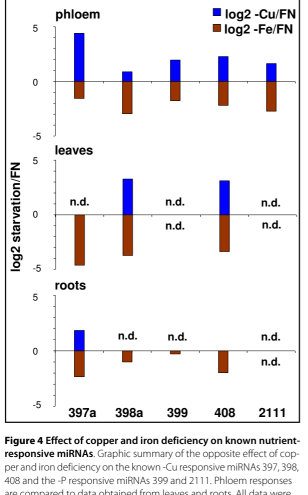
The response to S deficiency was characterized by a dramatic increase of the known -S-responsive miR395 (the at-miR395a signal increased from 280 to 76369).

While the amount of no additional miRNA increased, the amount of miR397 decreased (figure 3).

Growth under copper deficiency is known to induce a number of physiological responses, including the expression of specific miRNAs. Recently, the transcription factor SPL7 (SQUAMOSA promoter binding protein-like7) has been found to be a central regulator of the copper-deficiency response. It is able to induce the expression of miRNAs 397, 398, 408, 857, different copper transporters, and a copper chaperone [26]. Accordingly, our miRNA microarrays showed that copper deficiency led to a more than four-fold increase of the known copper-responsive miRNAs 397 and 408 that target laccases [1,11] in phloem sap. miR397 also accumulated in roots, but remained undetectable in leaves, while 408 responded positively in leaves and not in roots (figure 4).

sRNA No	-S/FN	-Cu/FN	-Fe/FN	sRNA name	sRNA No	-S/FN	-Cu/FN		-Fe/FN	sRNA name
158	n.d.	n.d.	2.4	ath-miR158b	397	-2.0	4.4	Ť	-1.6	ath-miR397a
			FN -Fe			n.d.	4.0	Ť	n.d.	osa-miR397b
				ath-miR158b		-6.7	4.1	Ť	-1.7	ptc-miR397b
395	8.1	-0.4	-0.5	ath-miR395a	398	0.2	0.9	Ť	-3.0	ath-miR398a
	6.7	-0.4	-0.4	ath-miR395b		0.3	0.9	Ť	-2.8	ath-miR398b
	8.1	-0.3	-0.5	mtr-miR395a					FN -Fe	ath-miR398a
	9.0	-0.3	-1.5	mtr-miR395b						amininosoa
	8.1	-0.3	-0.6	mtr-miR395g	399	n.d.	3.3	Ť	-1.8	ath-miR399d
	8.6	-0.3	-0.7	mtr-miR395h		n.d.	3.6	Ť	-1.7	ath-miR399e
	8.6	-0.3	-0.7	mtr-miR395p		-0.2	2.9	Ť	-1.7	ath-miR399f
	8.2	-0.4	-0.6	osa-miR395a		-0.1	2.5	Ť	-1.7	mtr-miR399a
	8.1	-0.4	-0.6	osa-miR395b		-0.2	2.2	Ť	-1.8	osa-miR399h
	9.1	-0.5	-1.0	osa-miR395c		0.2	2.0	Ť	-1.7	ptc-miR399h
	10.0	-0.3	n.d.	osa-miR395f		n.d.	3.1	1	-0.6	ptc-miR399j
	9.0	-0.4	-1.3	osa-miR395o		-0.2	2.2	X	-1.7	tae-miR399
	9.2	-0.5	-1.6	osa-miR395t					FN -Fe	bna-miR399
	9.3	n.d.	n.d.	osa-miR395u					-	bha-minto99
	11.1	n.d.	n.d.	osa-miR395v	408	n.d.	2.3	Ť	-2.2	ath-miR408
	9.8	-0.3	n.d.	osa-miR395w		n.d.	2.2	Ť	-7.4	osa-miR408
	9.6	-0.2	-0.5	ppt-miR395		n.d.	2.3	Х	-7.5	sof-miR408e
	10.8	-0.1	0.1	ptc-miR395a					FN -Fe	ath-miR408
>4-fold	increas	ed			857	n.d.	n.d.	-	n.d.	ath-miR857
>4-fold	decrea	sed			2111	n.d.	1.7	Ť	-2.7	bna-miR2111
									FN -Fe	bna-miR2111

Figure 3 List of nutrient-responsive sRNAs. List of sRNAs that showed a strong positive reaction to S, Cu or Fe deprivation, respectively, shown as log2 values of stressed vs. FN samples. Only sRNAs that fulfilled the criteria described in the Methods section (positive response, log2 >2 in one of the stress treatments, signal value >100 in FN or deprived sample) in at least one of the comparisons are listed. The insets show results obtained by miRNA sqRT-PCR (after 25 cycles) from an independent experiment. To allow a better overview, values for known nutrient starvation-responsive miRNAs (398 and 857 for -Cu and 2111 for -P) were included, although they only showed a negative response or were not detectable. Arrows indicate directions of changes obtained in a second, independent -Cu experiment. nd.: not detectable (both, FN and stress, signal values <100). X: not on chip.



are compared to data obtained from leaves and roots. All data were obtained from miRNA array hybridization experiments. Differences between stress and control plants are shown as log2 values, only Arabidopsis miRNAs are depicted. n.d.: not detectable.

The known -Cu-responsive miR398 that targets Cu/Zn superoxide dismutases also increased, but only nearly two-fold. A similar accumulation was also detected in leaves, but not roots (figure 4). miR857 that was found to be copper-responsive in Arabidopsis [11] was undetectable in the phloem, leaves and roots of rapeseed in the present study (figure 3), probably caused by the different species, compartment, developmental stage and milder stress treatment analyzed. Surprisingly, also the phosphate-deficiency-responsive miR399 increased more than four-fold (figure 3). This indicates a slight phosphate limitation in the -Cu plants, although the plants were supplied with the same amount of P as in all other experiments. The same was also observed in an independent repetition of the experiment (indicated by arrows in figure 3). Interestingly, miR2111 that was recently found to also respond to phosphate starvation [14] was also accumulating under -Cu, confirming the noticeable phosphate deficiency already evidenced by the increase of miR399 (figure 3). Our results thus confirm that copper deficiency up-regulates miRNAs that mainly target mRNAs of enzymes that use copper as cofactors, namely the multicopper proteins laccases and copper zinc superoxide dismutases (Cu/Zn SOD). As already discussed by Abdel-Ghany and Pilon [11], this mechanism is thought to save Cu for the most important copper-containing proteins like plastocyanin that is a key protein of photosynthesis [11].

Under iron deficiency only miR158 increased in the phloem more than four-fold (ath-miR158a increased from 231 to 1201), what was verified by sqRT-PCR in an independent experiment (inset in figure 3). miR158 was described as a non-conserved miRNA from Arabidopsis that could, for example, not be detected in citrus [27]. miR158 is predicted to target a pentatricopeptide repeatcontaining protein of unknown function, a lipase, and xyloglucan-fucosyl transferases [28]. None of these potential targets has an obvious connection to iron uptake or metabolism, and thus the increase of miR158 might be a secondary effect on plant development. Moreover, the accumulation of miR158 seemed to be phloem sap-specific, as it could not be observed in leaf or root samples (see data submitted to GEO, series accession number GSE20263). Comparative high-throughput sequencing of FN and -Fe samples would help to clarify if an as yet unknown (and therefore not represented on the chip) sRNA increases under -Fe, or if there is really no small RNA accumulating during this deprivation response.

Interestingly, however, miRNAs 397, 398, 399, 408 and 2111 notably decreased during iron starvation, showing an opposite response to their increases observed under -Cu (figure 3, figure 4). This response was verified for miR398, 399, 408 and 2111 by sqRT-PCR from a set of independently grown plants (inset in figure 3). Decreases in the levels of -Cu-responsive miRNAs were visible not only in the phloem, but also in leaves and comparably weak in roots (figure 4). A decrease of these Cu starvation-responsive miRNAs suggests that copper uptake is stimulated by iron deficiency, as has already been observed in Brassica and other plant species [29,30]. The need for higher Cu uptake under -Fe could be explained by the fact that many iron and copper-containing enzymes can substitute for each other when one of the two elements is present at suboptimal levels, e.g. SODs, cytochrome oxidase, or diiron oxidase [31,32].

Interestingly, a phloem response opposite to the -Cu reaction under -Fe was also observed for the -P-responsive miRNAs 399 and 2111, which were more than two-(399), respectively more than four-fold (2111) decreased. The responses of miR399 and miR2111 were undetectable in leaves and roots (figure 4). This confirms the

observation from a previous study that demonstrated that miR399 responds stronger to -P in phloem sap than in leaves and roots [2]. The decrease of -P-responsive miR-NAs in phloem sap suggests that Fe deficiency positively influences P uptake and metabolism, what has already been demonstrated in earlier studies e.g. [33,34]. The other way around, high Fe can lead to lower P concentrations in the plant [34]. If more Fe is taken up during growth under -Cu in order to replace Fe in Cu-containing enzymes, this could explain the observed increase of the -P-responsive miRNAs in phloem sap under Cu deprivation.

Taken together, the data from the -Cu and -Fe experiments indicate a tight link between iron and phosphate metabolism that has earlier been described. Moreover, they suggest a close linkage between iron and copper uptake, although it is known that in higher plants this link is at least not as close as, for example, in yeast or *Chlamydomonas*, where iron uptake is directly Cu-dependent [35,36]. It is interesting to note that the tissues/compartments analyzed react differentially to specific stress triggers, but the physiological meaning of this observation needs to be evaluated in future experiments.

Specific miRNAs that accumulate in phloem sap under stress are also mobile in grafting experiments

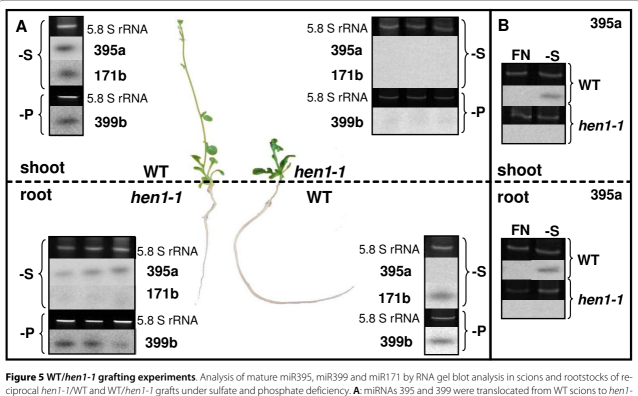
Whether miRNAs are mobile between cells and over long distance is still a matter of debate and evidence for transport only exists for one single miRNA, miR399, that was able to move from shoots to roots in a miR399 overexpressor as scion/WT as rootstock graft situation [2,3]. Because miR395 is comparably well studied, its targets have been validated in Arabidopsis, and it strongly accumulates under sulfur starvation, also within the phloem, we chose this miRNA to examine whether additional miRNAs are mobile in vivo. To this end, we performed grafting experiments using hen1-1 mutants and WT plants. hen1-1 mutants are inhibited in sRNA methylation and, as a consequence, the levels of several miRNAs are markedly decreased [37]. RNA gel blot analysis of the different miRNAs further analyzed in our study confirmed that hen1-1 mutants did not contain any of these mature miRNAs at detectable levels (data not shown). In all grafting experiments, hen1-1 mutants retained their typical phenotype, mainly characterized by growth retardation (figure 5A), what indicates that not all necessary miRNAs can be translocated between the grafting partners. After the establishment of graft unions, successful grafts were transferred to media lacking a specific nutrient for two weeks, and miRNA abundance was analyzed in the different parts of the graft by RNA gel blots. We first examined the abundance of the phosphate-dependent miR399 in scions and rootstocks under phosphate starvation as a positive control. As expected, miR399 was not only clearly detectable in WT rootstocks and scions, but also in *hen1-1* rootstocks of independent grafts with similar signal strength as in phosphate starved WT rootstocks (figure 5A). Our data thus confirmed the translocatability of miR399 from shoots to roots in a graft situation. We further chose miR171 as a negative control, since this miRNA has neither been detected in phloem sap by sRNA sequencing [1,14,38] nor by our sRNA array experiments (additional file 1). As assumed, we detected a signal in the WT rootstocks and scions, but not in the mutant parts of the grafts, making a phloem translocation of miR171 highly unlikely (figure 5A).

When analyzing grafts grown under sulfate starvation, we observed the translocation of miR395 from WT scions to *hen1-1* rootstocks in different independently grafted plants. We also observed signals for miR395 in WT scions, but not in WT rootstocks (figure 5A). However, miR395 has been previously shown to be expressed in roots under sulfur starvation [39], and we could also detect signals in roots of intact WT plants (figure 5B). This result could be reproduced in several independent experiments. This could indicate that miR395 translocation from shoot to root is required for root miR395 expression in the WT, but further experiments will be needed to substantiate this assumption. The earlier studies of miR399 translocation do not allow any conclusions about the (non) existence of such a crosstalk, since a comparable graft situation of a stressed WT rootstock with an "unstressed" (not miRNA-producing) scion cannot be achieved when grafting overexpressors with WT plants [2,3].

For both, miR399 and miR395, we only found signals in hen1-1 rootstocks and never in hen1-1 scions, indicating that mobility was restricted to the direction from shootto-root in Arabidopsis seedlings (figure 5A). The reason for this unidirectional translocation might lie in the early developmental stage analyzed, where roots constitute the only real sink organ that needs nutrient supply from the phloem translocation stream. However, the results do not rule out that mobile miRNAs can reach other organs than roots at different developmental stages with different source-sink relationships. Our experiments also did not allow concluding whether mature miR395 or its PT is the translocated species. In the case of miR399, however, it has been previously shown that exclusively mature miRNA and not PTs is transported through graft unions [2]. In addition, no miRNA precursors were detectable in B. napus phloem sap [1], suggesting that mature miRNAs are the translocated molecules.

The graft translocation of miR395 coincides with a down-regulation of the target APS4

To examine whether the translocation of miR395 from WT shoots into *hen1-1* roots might have physiological



ciprocal hen1-1/W1 and W1/hen1-1 grafts under sulfate and phosphate deficiency. **A**: miRNAs 395 and 399 were translocated from W1 scions to hen1-1 rootstocks but not in the opposite direction, miR171 was immobile. One representative result is shown for WT, and three replications for hen1-1 roots and shoots. The hen1-1 graft parts kept their growth retardation phenotype, indicating that not all necessary miRNAs could be transferred. The 5.8 ribosomal RNA band served as a loading control. **B**: Control of miR395 expression in WT and hen1-1 mutant plants. In WT plants miR395 was induced by sulfate deficiency in shoots and roots, while no signal was detected in hen1-1 mutants under both conditions.

functions, we analyzed the levels of three experimentally validated mRNA targets of miR395, the ATP sulfurylases *APS1* and *APS4* and the low affinity sulfate transporter *AtSULTR2;1* [8,39]. As a general observation, the transcript levels of all three targets seemed to be higher in shoots of *hen1-1* as compared to WT plants (additional file 4). In addition, the experiments showed that only the level of ATP sulfurylase *APS4* mRNA, but not of *APS1* or the low affinity sulfate transporter *SULTR2;1*, was notably

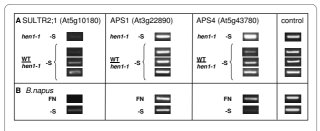


Figure 6 Analysis of the targets of miR395 in roots. Analysis of the mRNA levels of the miR395 targets SULTR2;1, APS1 and APS4 by semiquantitative RT-PCR. **A**: PCR results from root tissue of hydroponically grown Arabidopsis *hen1-1* mutants and WT/*hen1-1* rootstocks (35 cycles, UBC10, At5g53300 served as a control). **B**: Changes of target mR-NAs in *B. napus* roots under -S compared to full nutrition (FN) (35 cycles, UBP1B, At1g17370 served as a control). decreased in grafted hen1-1 rootstocks as compared to non-grafted -S starved roots of hen1-1, while housekeeping genes remained constant (figure 6A). A similar reduction of levels of APS4, but not the other two targets, could be observed in B. napus WT roots grown under sulfur starvation (figure 6B). These results indicate that APS4 mRNA might be a target of miR395 in roots, and interestingly, this mRNA has previously been shown to exhibit root-specific expression [40]. The observation that the other miR395 target SULTR2;1 was up- and not downregulated under -S conditions (figure 6A and 6B, [39]) was earlier explained by the spatially differential expression of SULTR2;1 and miR395 in xylem parenchyma and companion cells, respectively [39]. It was suggested that one of the major functions of miR395 was the down-regulation of SULTR2;1 expression in the phloem to restrict SULTR2;1 expression exclusively to the xylem [39].

Is the transport of specific miRNAs of biological relevance in intact plants?

Most miRNAs are believed to act in a locally restricted manner, in contrast to the mobile class of siRNAs [41]. Their limited mobility is suggested by the closely correlating patterns of miRNA transcription and activity [42],

the spatial restriction of miRNA gene expression [43,44], and the limited area of mature miRNA localization [45]. However, phloem mobility of miR399 across graft unions has been demonstrated in earlier studies by grafting miR399 overexpressor with WT plants [2,3]. In this study, we observed the transport of miR395 and 399 from WT scions to *hen1-1* mutant rootstocks. Moreover, one of the miR395 targets, APS4, was down-regulated in grafted mutant roots. This indicates that miR395, like miR399, is transported from shoot to root to down-regulate its target(s). However, the question whether such a miRNA transport is physiologically relevant remains, since members of the miR395 and 399 families can indeed be synthesized in roots of wild type plants under the respective stress [7,39] (figure 5B). Interestingly, expression of miR-NAs 395 and 399 was shown to be highly overlapping, being predominant in vascular tissue, especially in root phloem companion cells (CC) [7,39].

Different scenarios could explain the observation that specific miRNAs are present in phloem sap and mobile in grafting experiments: 1) None of the phloem miRNAs is specifically targeted for translocation, but instead a portion of all miRNAs highly expressed in CC leaks into sieve elements. No miRNA would represent a signaling molecule. 2) A portion of all miRNAs highly expressed in CC reaches phloem sap, but some of these miRNAs can act as long-distance regulators under certain physiological conditions. 3) Selected miRNAs synthesized in CC are specifically targeted for transport and only these are released into the phloem stream. In this case, all miRNAs present in the phloem would be translocatable information transmitters.

No matter how miRNAs reach phloem sap, they would then be swept away from source to sink organs (in our system from shoots to roots). The translocated miRNAs would probably exit the translocation stream into sink CC in an unspecific manner, as rather unselective unloading of macromolecules into sink tissues has been suggested [46]. Here, they would down-regulate their target mRNAs, no matter whether they are intended to function as signaling molecules or not.

If certain miRNAs should indeed be translocated to transmit information, one possible rationale could be that roots are unable to synthesize sufficient amounts of these miRNAs under stress, or that they need a trigger from the shoot to initialize miRNA synthesis. This might be suggested by the absence of mature miR395 in WT rootstocks of grafted plants that was, however, well detectable in roots of complete WT plants (figure 5). Another explanation might be that some organs experience nutrient deprivation earlier than others, and that the translocated miRNAs serve to coordinate physiological responses with plant parts that are not yet stressed and therefore do not yet synthesize stress-responsive miRNAs themselves. This would resemble the situation in grafted plants, where only scions of the graft produced the stressinduced miRNAs (stressed WT in this study, overexpressors in [2]), while rootstocks did not (*hen1-1* mutants in this study, non-stressed WT in [2]).

Conclusions

This study demonstrates that the phloem sap sRNA complement is distinct from that of stems, leaves and roots, and that a set of phloem-enriched sRNAs exists. It also shows that the abundance of several phloem sap sRNAs changes under nutrient deficiency conditions. While the results confirmed that the known miRNAs reacting to -S or -Cu, respectively, also respond in phloem sap, they provided no clear indications that the response to -Fe involves miRNA regulation, despite of influencing copper uptake/metabolism.

Grafting studies between WT plants and *hen1-1* mutants demonstrated that two phloem stress-reactive miRNAs, 395 and 399, can indeed be transported from shoot to root in Arabidopsis seedlings, and that this translocation leads to a reduction of the amount of their target mRNAs in roots. The grafting experiments also revealed that not all miRNAs are phloem translocatable, since miR171 did not move.

Therefore, this study demonstrates that identifying phloem-enriched macromolecules and analyzing their translocation in grafting studies is a very useful approach to distinguish between phloem translocatable and nonmobile molecules. It is tempting to classify miR395 and 399 as systemic signaling molecules, because they not only move from source to sink, but also induce a measurable effect on their target mRNAs in sink tissue in grafting experiments. However, we conclude that profiling phloem components combined to grafting studies is still not sufficient to doubtless decide whether a phloemtranslocatable macromolecule is really a long-distance signal or not.

Methods

Plant material and growth conditions

For hydroponic growth, *Brassica napus* (cv. Drakkar, Serasem GIE, la Chapelle d'Armentiers, France) seeds were germinated on wet filter paper for 1 week. Germ buds were transferred to plastic boxes containing nutrient medium for 10 weeks. Nutrient medium: 0.6 mM NH₄NO₃, 1 mM Ca(NO₃)₂*4H₂O, 0.04 mM Fe-EDTA, 0.5 mM K₂HPO₄, 0.5 mM K₂SO₄, 0.4 mM Mg(NO₃)₂*6H₂O. Micro nutrients added: 0.8 μ M ZnSO₄*7H₂O, 9 μ M MnCl₂*4H₂O, 0.1 μ M Na₂MoO₄*2H₂O, 23 μ M H₃BO₃, 0.3 μ M CuSO₄*5H₂O. The pH was adjusted to 4.7 with 37% HCl. Nutrient solutions were changed after 4 weeks, and then renewed once a week. After 5 to 6 weeks, media were constantly aerated by an aquarium air pump (Sera, Heinsberg). Sulfur and copper starvation were applied for two, and iron starvation for three weeks before flowering started by changing to medium without sulfur, copper, or iron, respectively. Here, 0.5 mM K₂SO₄ were substituted by 0.5 mM K₂HPO₄ and instead of ZnSO₄*7H₂O and CuSO₄*5H₂O as micro nutrients, 1 μ M ZnCl₂ and 1 μ M CuCl₂*2H₂O were added for low sulfate experiments. For copper deprivation, the 0.3 μ M CuSO₄*5H₂O were omitted from the full nutrient solution. For low iron experiments Fe-EDTA was omitted from the medium.

For the growth of Arabidopsis thaliana WT (ecotype Ler-0) and hen1-1 [47] mutant plant seeds (NASC code N6583) were surface-sterilized in 70% (v/v) ethanol for 3 min and further incubated in 20% sodium hypochlorite solution containing 0.1% (v/v) surfactant (Triton X-100) for 10 min. After exhaustive washing with sterile water, seeds were placed on plates on half-concentrated MS medium [48] supplemented with 1% (w/v) sucrose and solidified with 0.7% (w/v) agar. After keeping them in the dark for three days at 4°C, seeds were germinated by transferring the plates in a growth chamber under controlled long day conditions (16 h day, 8 h night) at 25°C for 13 days. For hydroponic cultivation these plantlets were transferred into plastic boxes containing the nutrient solution previously described in [49] with minor modifications in the content of magnesium sulfate, boric acid and potassium dihydrogen phosphate (4 mM MgSO₄*7H₂O and 0.1 mM H₃BO₃, 2.5 mM KH₂PO₄). The hydroponic growth was carried out under short day conditions (8 h day at 20°C, 16 h night at 16°C). For sulfur deprivation experiments starvation was applied directly after the transfer of plantlets to hydroponic culture with nutrient solution omitting all sulfate-containing components for two weeks. Instead of MgSO4*7H2O 0.8 mM MgCl₂*6H₂O were added to the medium. Phosphate starvation was performed analogously in nutrient solution that contained potassium nitrate instead of potassium dihydrogen phosphate.

Micrografting experiments

For micrografting experiments four-day-old *Arabidopsis thaliana* wild type and *hen1-1* mutant seedlings were cut transversely using a sterile small razor blade part and combined within silicon tubing (0.3 mm internal diameter) as previously described [50]. The grafts were grown on 1.5% (w/v) agar plates with half-strength MS medium for nine days under controlled short day conditions. Successfully grafted plantlets were subsequently grown hydroponically for two weeks before plant material from stock and scion was harvested. To avoid contaminations, the area close to the graft union was omitted from sampling and grafts were microscopically inspected for

adventitious root formation, what led to exclusion from analysis.

Sampling and RNA isolation

Phloem sampling from *Brassica napus* plants was performed as described earlier [1,12] from 4 - 8 small punctures into the inflorescence stems. After discarding the first droplets to avoid contaminations, 500 μ l to 1.5 ml phloem sap from three independent sets of plants were obtained, yielding about 10-50 μ g of total RNA. Total RNA from phloem sap was isolated by Trizol LS reagent (Invitrogen) according to manufacturer's instructions.

RNA from 100 mg frozen material of stem, leaf and root tissue of *Brassica napus* and *Arabidopsis thaliana*, respectively, was extracted using the normal Trizol reagent. Total RNA from all samples was dissolved in 25 μ l DEPC-treated water and RNA concentrations were determined photometrically with a Biophotometer (Eppendorf).

Microarray hybridization

Microarray assays were performed by LC Sciences (Houston, Texas). The assays started from 2 to 5 µg total RNA samples that were size fractionated using a YM-100 Microcon centrifugal filter (Millipore) and the sRNAs (< 300 nt) isolated were 3'-extended with a poly(A) tail using poly(A) polymerase. An oligonucleotide tag was then ligated to the poly(A) tail for later fluorescent dye staining. Two different tags were used for the two RNA samples in dual-sample experiments. Hybridization was performed overnight on µParaflo microfluidic chips using a micro-circulation pump (Atactic Technologies). On the commercial microfluidic chip, each detection probe consisted of a chemically modified nucleotide coding segment complementary to a known target plant miRNA (from miRBase, http://microrna.sanger.ac.uk/ <u>sequences/</u>, releases 10.0 (-S), 10.1(-Fe) or 11.0 (-Cu)). The known plant miRNAs were mainly from Arabidopsis thaliana, Oryza sativa, Populus trichocarpa and Physcomitrella patens. Among the total number of unique miRNA sequences (release 10.0, 623 miRNAs, 10.1, 653 miRNAs and 11.0, 714 miRNAs) all arrays contained a constant number of 154 miRNAs from Arabidopsis thaliana. Additionally to these known miRNAs, the customized array contained a set of 85 sRNAs of unknown function that were derived from an earlier high-throughput sequencing experiment of phloem sap [1] (sequences and accession numbers in additional file 5). Coding segments were coupled to a spacer segment of polyethylene glycol to place the coding segment away from the substrate. The detection probes were prepared by in situ synthesis using PGR (photogenerated reagent) chemistry. The hybridization melting temperatures were balanced by chemical modifications of the detection probes. For hybridization 100 μ L 6 × SSPE buffer (0.90 M NaCl, 60 mM Na₂HPO₄, 6 mM EDTA, pH 6.8) containing 25% formamide at 34°C were used. After hybridization, signals were detected after fluorescence labeling using tag-specific Cy3 and Cy5 dyes. Hybridization images were collected using a laser scanner (GenePix 4000B, Molecular Devices) and digitized using Array-Pro image analysis software (Media Cybernetics). Data were analyzed by first subtracting the background and then normalizing the signals using a LOWESS (locally-weighted regression) filter.

To allow inter-array comparisons of FN samples, signal intensities were normalized to the median signal intensity of each sample and p-values of the t-test were calculated for the three replicates of each organ (phloem, leaves, and roots). Signals with p-values lower than 0.05 were regarded as being differential.

For the stress experiments (two color hybridizations), the ratio of the two sets of detected signals (log2 transformed, balanced) and p-values of the t-test were calculated and signals with p-values lower than 0.01 were regarded as being differential. Since only one array per stress was hybridized, we further restricted the data evaluation to sRNAs that showed a signal intensity of >100 in the FN or the stressed sample, an accumulation upon stress, and a more than four-fold difference (log2s of >2 or <-2) between stress and FN. All microarray data have been submitted to GEO, series accession No. GSE20263.

Semi-quantitative RT-PCR

For semi-quantitative RT-PCR (sqRT-PCR), Trizol isolated RNA was cleaned with the RNeasy Plant Mini Kit (Qiagen) and a DNase I digest following the manufacturers instructions was performed. For nutrient stressresponsive marker gene and miRNA target transcript analysis, 500 - 1000 ng RNA were used for cDNA synthesis in the presence of 2.5 μ M oligo(dT)₂₀ primer (Qiagen), 0.5 mM dNTPs, 5 mM DTT (Invitrogen), 40 U RNase-OUT RNase Inhibitor (Invitrogen) and 200 U M-MLV reverse trancriptase (Promega) in 1× M-MLV reverse transcriptase reaction buffer (Promega) in a final volume of 20 μ l. The reverse transcription reactions were carried out in a Primus Thermocycler (Peqlab) at 50°C for 45 min followed by 70°C for 15 min to denature the reverse transcriptase enzyme. 2 µl of the reverse transcription reaction were used for each PCR amplification with gene specific oligonucleotide primer pairs (additional file 6). The reaction mixtures containing 1.5 mM MgCl₂ (Invitrogen), 0.2 mM dNTPs (Promega), 0.2 µM of both forward and backward primer and 2 U of Paq5000 DNA Polymerase in a 50 µl volume of 1× Paq5000 DNA polymerase buffer (Agilent Technologies) were divided into three equal volumes in reaction tubes and semi-quantitative RT-PCR was performed with different cycle numbers under the following conditions: 30 s at 94°C, 30 s at 55°C, 1 min at 72°C and a 10 min end-elongation step at 72°C. The PCR reaction was stopped after a certain number of cycles and PCR products were separated electrophoretically in 2% (w/v) agarose gels for size estimation and semi-quantitative analysis.

PCR of mature miRNAs was performed by following the method of Shi and Chiang [51]. Total RNA $(1 \mu g)$ was first polyadenylated by a poly(A) polymerase [E-PAP, Poly(A) Tailing Kit (Ambion)] at 37°C for 1 h in a 50-µL reaction mixture containing 1× E-PAP buffer, 2.5 mM MnCl₂, 1 mM ATP and 1 U E-PAP. Samples were purified from E-PAP by a further RNA extraction using TriFast FL reagent (Peqlab) and resolved in 50 µl DEPC-treated water. 10 µl of the polyadenylated RNA samples were used as a template for reverse transcription performed as described above using 0.5 µg poly(T) adapter instead of the oligo(dT)₂₀ primer. miRNAs were subsequently amplified using 1 μ l of the reverse transcribed sample, miRNA-specific forward and poly(T) adapter-specific reverse primers (additional file 6) under the same PCRcycler conditions used in sqRT-PCR described above.

RNA gel blot analysis

Gel blot analyzes were performed on 15% denaturing urea gels as described earlier [1,52].

Additional material

Additional file 1 Comparison of miRNA abundance in phloem sap vs. inflorescence stem. Comparison of sRNA microarray analysis of stem tissue (green) and phloem sap (blue) of *Brassica napus*. Only known miRNAs present on the commercial array, only one member per family are depicted. The upper graphs show the signal intensities on the array while the lower depict the log2 differences between phloem and inflorescence stem. Insets show RNA gel blot analyses of selected miRNAs from an independent experiment. Numbers indicate the number of sequences that were previously obtained by phloem sap sequencing [1], asterisks (*) indicate sequences from miRNA stars.

Additional file 2 Comparison of sRNA abundances in phloem, leaves and roots. sRNA microarray comparison of phloem (blue), leaf (green) and root (red) tissue of *Brassica napus* plants from biologically independent replications (n = 3). To allow inter-array comparison, signal intensities were normalized to the median signal of each sample. Only known miRNAs present on the commercial array and only one member per family are depicted.

Additional file 3 Transcript analysis of known nutrient stress-specific genes. Transcript analysis of known nutrient stress-specific genes in leaf and root tissue of hydroponically grown *Brassica napus* plants by semiquantitative RT-PCR after 25, 30 and 35 cycles under -S, -Cu and -Fe compared to full nutrition (FN).

Additional file 4 Accumulation of three miR395 targets in WT and hen1-1 shoots grown under full nutrition. Levels of the targets SULTR2;1, APS1 and APS4 in shoots as detected by sqRT-PCR (35 cycles, UBC10, At5q53300 served as a control). FN: full nutrition.

Additional file 5 Sequences of the unknown phloem sap sRNAs represented on the microarrays. Phloem sap small RNA sequences of *Brassica napus* (Bn_PsRNAs) that were contained on the sRNA microarray (sequences were derived from high-throughput sequencing of *B. napus* phloem sap published in [1]). Additional file 6 List of oligonucleotides used. Oligonucleotide sequences used for the detection of nutrient stress-specific miRNAs by RNA gel blots or by semi-quantitative RT-PCR, for the analysis of miR395 target genes, and for transcript detection of nutrient-responsive genes by semi-quantitative RT-PCR.

Authors' contributions

AB and FS carried out the plant growth, stress and microarray experiments. AB was also involved in microarray data analysis and evaluation. JP carried out the micrografting experiments, miRNA and target analyses. AB and JP drafted the manuscript. JK conceived of the study, participated in its design, coordination, data analysis, and drafted the manuscript. All authors read and approved the final manuscript.

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