Sensitive whole mount in situ localization of small RNAs in plants 1 2 Mouli Ghosh Dastidar¹, Magdalena Mosiolek², Michael D. Nodine² and Alexis Maizel^{1,*} 3 4 5 1: Center for Organismal Studies (COS) 6 University of Heidelberg 7 Im Neuenheimer Feld 230 8 69120 Heidelberg 9 Germany 10 11 2: Gregor Mendel Institute (GMI) 12 Austrian Academy of Sciences 13 Vienna Biocenter (VBC) 14 Dr. Bohr-Gasse 3 15 1030 Vienna 16 Austria 17 *: to whom correspondence should be addressed: 18 19 Email: alexis.maizel@cos.uni-heidelberg.de 20 Phone: +49 6221 54 64 56 21 Fax: +49 6221 54 64 24 22 23 **Abstract** 24 Small regulatory RNAs are pivotal regulators of gene expression and play important roles in many 25 plant processes. Although our knowledge of their biogenesis and mode of action has significantly 26 progressed, we comparatively still know little about their biological functions. In particular, knowledge 27 about their spatiotemporal patterns of expression rely on either indirect detection by use of reporter constructs or labor-intensive direct detection by in situ hybridization on sectioned material. None of 28 29 the current approaches allows for a systematic investigation of small RNAs expression 30 patterns. Here, we present a method for the sensitive in situ detection of micro- and siRNAs in intact 31 plant tissues that utilizes both double-labelled probes and a specific cross linker. We determined the 32 expression patterns of several small RNAs in plant roots and embryos. 33 34 **Key words**

Arabidopsis thaliana; micro-RNA; short interfering RNA, in situ hybridization, expression pattern

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Introduction

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Small (20-25nt) RNAs (smRNAs) regulate gene expression in eukaryotes by guiding the transcriptional and post-transcriptional gene-silencing machinery by base pairing to their targets. In plants, smRNAs are involved in development, response to phytohormones and nutrients and ensure genome integrity by mediating the epigenetic regulation of mobile repetitive elements. (Jones-Rhoades et al., 2006; Rubio-Somoza et al., 2009; Sunkar et al., 2007). Our understanding of smRNA biogenesis and action has made significant progresses in the recent years and modern seguencing technologies have tremendously expanded the repertoire of smRNAs. Yet in comparison, the study of their function is lagging behind. A key aspect of the functional characterization of smRNAs is a precise description of their spatial and temporal patterns of expression at the cellular scale. Published examples have revealed that in plants smRNAs are expressed in discrete, tissue or cell type-specific pattern (Juarez et al., 2004; Noqueira et al., 2009; Chitwood et al., 2009; Wollmann et al., 2010; Ori et al., 2007; Cartolano et al., 2007; Douglas et al., 2010; Marin et al., 2010; Carlsbecker et al., 2010; Miyashima et al., 2011; Yu et al., 2015; Husbands et al., 2015). The most common method to infer the expression patterns of smRNAs at the cellular level consists the use of transgenic reporter genes encoding detectable products like Beta-Glucuronidase (uid A), Green Fluorescent Protein (GFP) and Luciferase (LUC), that are placed under the control of putative small RNA precursor promoters (Carlsbecker et al., 2010; Yu et al., 2015; Marin et al., 2010; Noqueira et al., 2009; Nogueira et al., 2007; Chitwood et al., 2009). Alternatively, ubiquitously expressed reporters containing small RNA binding sequence have been used to reveal the small RNA expression pattern as area of absence of reporter expression (Parizotto et al., 2004; Nodine and Bartel, 2010; Marin et al., 2010). However, because this method produces an absence of signal and is influenced by the efficacy of the small RNA, it can be difficult to interpret, and requires the time-consuming, and not always possible, generation of transgenic plants. Direct localization by in situ hybridization with specific anti-sense probes provides high resolution and does not rely on the establishment of such transgenic reporters. Original protocols employed digoxigenin labelled anti-sense RNA probes (Kidner and Martienssen, 2004; Ori et al., 2007; Juarez et al., 2004; Douglas et al., 2010; Cartolano et al., 2007; Noqueira et al., 2009; Chitwood et al., 2009) but were supplanted by use of locked nucleic acid (LNA) oligonucleotide probes that provide increased sensitivity and specificity (Kloosterman et al., 2006; Wheeler et al., 2007; Válóczi et al., 2006). However these protocols suffer from two main disadvantages that preclude their widespread adoption. These methods rely on the use of embedded and sectioned material as a template for hybridization. The generation of these sections is a tedious and lengthy process and some tissues such as the root are particularly challenging to sectioning. Second, the current protocols have a limited sensitivity. Small RNAs are often not be very abundant and current protocols do not take into account the high volatility of small RNAs during the numerous washing steps of the protocols (Pena et al., 2009). The conventional formaldehyde fixation of tissue used in such protocols has been shown to result in substantial loss of smRNAs which can be prevented by the usage of EDC (N-(3-Dimethylaminopropyl)-N'-ethylcarbodiimide hydrochloride) that covalently cross-links smRNA 5' ends to the protein matrix (Pall *et al.*, 2007; Pena *et al.*, 2009). Here, we present an *in situ* hybridization protocol for smRNAs that enables their detection in non-sectioned plant tissues. Our protocol relies on the use of double-labelled LNA probes and a smRNA-specific post-fixation step using EDC for superior sensitivity. We have employed this method to document the expression pattern of seven micro- and short interfering RNAs in Arabidopsis roots and embryos. This approach is highly specific and allows for a semi-quantitative assessment of small RNA abundance. The whole procedure can be preformed using liquid-handling robotic systems and is therefore amenable to high- throughput applications.

Results & Discussion

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We devised a protocol for the detection of small RNAs in whole mount Arabidopsis samples (Figure 1). After initial steps of fixation, dehydration, proteinase treatment and post-fixation, samples are optionally treated with EDC (N-(3-Dimethylaminopropyl)-N'-ethylcarbodiimide hydrochloride), a chemical reacting with free 5' ends present in small RNAs and condensing them with amino groups in the protein matrix of the tissue (Pall et al., 2007; Pena et al., 2009). This step ensures that the small RNAs are not washed away during subsequent steps (Pena et al., 2009). We used LNAmodified probes for hybridization with fixed samples. LNA oligos are high-affinity RNA analogs in which the furanose ring of selected base is locked by an O2',C4'-methylene bridge resulting in increased hybridization properties and specificity towards their targets. The probes were chemically modified on their 5' and 3' ends with a digoxigenin residue. The presence of two digoxigenin group enhances the sensitivity of the probe. Hybridizations were performed overnight at temperatures of 55°C or 65°C depending on the probe (Table 1). After washing and blocking, the hybridized probe was detected by incubating with an antibody against digoxigenin coupled to an alkaline phosphatase followed by its colorimetric detection with chromogenic substrate under the microscope. As a proof of principle, we applied the protocol to the detection of miR390 in the root system of 7-day old Arabidopsis wild type (Col-0). Mir390 is an abundant miRNA in the root produced by a single active locus (MIR390a) (Breakfield et al., 2011; Marin et al., 2010), and expressed in the meristem region of the primary and lateral root primordia (Marin et al., 2010). The specificity of the detection was assessed by performing the procedure either with a probe antisense to the miR390 sequence, a sense probe or no probe at all. We observed strong hybridization signals in the primary root, as well as in the lateral root primordia with the antisense miR390 probe (Figure 2A). We observed very weak to no signal in samples incubated with either the sense or no probe at all (Figure 2B, C). In the samples hybridized with the miR390 antisense probe, signal was more prominent in the cells of the meristem regions in particular in the stele, whereas signal in the endodermis, cortex and epidermis were also observed. No signal was observed in the columella cells. The miR390 pattern observed by *in situ* hybridization was identical to that produced by the transcriptional reporter pMIR390a::GUS:GFP (Marin *et al.*, 2010). MiR390 signal was only detected in the cytoplasm of the cells and more easily visible in cells without a fully developed vacuole (Figure 2D). Together, these results indicate that miR390 can be specifically detected in whole mount Arabidopsis root samples.

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We then tested other less abundant root miRNAs (Breakfield et al., 2011) such as miR160, miR166 and miR167. Whereas miR390 could be easily detected in absence of EDC-crosslinking, in the same conditions, we did not detect any signal for samples incubated with miR160, 166 or 167 antisense probes (Figure 3A, C, E). In samples treated by EDC, we observed signal in the primary root meristem region with all three probes (Figure 3B, D, F). Similar to miR390, these additional miRNA signals were cytoplasmic and easier to visualize in non-fully vacuolated cells. The signal obtained with the miR166 antisense probe was more intense in the epidermis and cortex layers than in the inner layers (Figure 3D), which is in agreement with previous reports (Carlsbecker et al., 2010). Whereas miR160 signal was also more intense in the outer layers (Figure 3B), miR167 was more intense in the stele (Figure 3F). miR172, did not yield any strong signal even in presence of EDC (Figure 3G, H), although it appears to be expressed in roots (Breakfield et al., 2011). We verified that the EDC treatment preserved the specificity of the detection by probing Arabidopsis roots treated with EDC or not with probes against the animal-specific miR124. We did not detect any signal even when the samples had been cross-linked by EDC (Figure 3I, J). In their ensemble, these results show that EDC-crosslinking of small RNAs improves the sensitivity of detection while preserving the specificity.

To extend our observations to other tissues, we applied our smRNA *in situ* method to Arabidopsis embryos with a few modifications as outlined in the experimental procedures. Signals corresponding to miR160, miR166, miR167 and miR390, but not miR172, were reproducibly detected above the background signal observed when using the animal-specific miR124 as a negative control (Figure 4). Consistent with the results from post-embryonic roots, we also observed the localization of miR160, miR166, miR167 and miR390 in embryonic radicles. In addition, miR160, miR166 and miR167 were also present in the shoot meristem region and the hypocotyl of embryos. Similar to the results from post-embryonic roots, the use of EDC-crosslinking in the WISH procedure improved the sensitivity of miR160, miR166 and miR167 although this was not required to detect miR390. Moreover, because miR160 probes give strong signal in the innermost cell layers (i.e. vascular tissue precursors) in either the presence or absence of EDC, the use of EDC-crosslinking preserved the cell-specificity of the *in situ* hybridizations (Figure 4A-B).

Given the sensitivity of the method, we tested whether it would be amenable to the qualitative assessment of the abundance of other small RNAs. For this purpose, we designed a probe against

the trans acting (ta)-siRNAs produced by the *TAS3* pathway (tasi-ARFs) and observed tasi-ARF accumulation in either wild typeor in lines in whichthe expression of the *TAS3a* precursor iseither reduced (*tas3a-1*, (Adenot *et al.*, 2006)) or it is over-expressed (*TAS3 OX*, (Marin *et al.*, 2010)). *TAS3 OX* plants have a 2-3 fold increased in tasi-ARFs levels compared to wild type (Marin *et al.*, 2010) whereas *tas3a-1* mutant has only 40% of wild-type levels (Adenot *et al.*, 2006). All samples were treated in parallel and the incubation with the chromogenic substrate was identical for all three genotypes. The hybridization signal intensities observed were lower in the primary of *tas3a-1* mutants compared to wild type (Figure 5A, C) and stronger in the *TAS3 OX* overexpression lines (Figure 5B). These results demonstrate that the method is able to detect siRNAs and can pick up relative differences in their abundance.

To further validate the applicability and specificity of the method, we probed for abundance of small RNAs in plants impaired in the biogenesis of these small RNAs. We first tested the abundance of tasi-ARFs in *rdr6*, *dcl-4* or *drb4*mutant plants that are deficient in tasi-ARF production (Elmayan *et al.*, 1998; Gasciolli *et al.*, 2005; Adenot *et al.*, 2006; Xie *et al.*, 2005). Whereas signal was detected in wild type, no signal could be seen in the *rdr6*, *dcl-4* or *drb4*mutants (Figure 5E, F, G). We then compared the levels of miR390 in wild type and inthe *miR390a-1*, *miR390a-2* and *hen1-5* mutants. Whereas *miR390a-1* is a mutant defective in the processing of the MIR390a precursor (Cuperus *et al.*, 2009), *miR390a-2*has reduced levels of the MIR390A precursor (Marin *et al.*, 2010). The *hen1-5* is a mutant impaired in miRNA biogenesis(Park *et al.*, 2002). The signal intensities were lower in *miR390a-1* mutants compared to wild type and even lower in *miR390a-2* (Figure 5H, I, J) and almost no signal could be detected in *hen1-5* mutant (Figure 5K). The reduced levels of miR390 in the *miR390a-1* and *miR390a-2* mutants was validated by northern blot analysis, which showed an 80% and 50% reduction of mature miR390 levels in respective *miR390a-1* and *miR390a-2* mutants compared to wild type (Figure 5L).

In summary, this protocol is sensitive, robust and efficient as it does not involve the long steps of tissues embedding and sectioning before hybridization. The use of double labelled LNA probe and the stringent hybridation conditions ensure both sensitive and specificity to the method as exemplified by the very low of absence of signal when using either sense probes, unrelatedmiRNA (miR124) or performing hybridization in mutant backgroundswith reduced miRNA abundance. Moreover, the protocol is applicable to both embryonic and mature tissues and amenable to the detection of miRNAs and siRNA (ta-siRNA). The protocol should be applicable to other tissues or plant species. The step of small RNA cross-linking with EDC improves the detection of several small RNAs while preserving specificity. Here we have used an additional step of cross-linking of small RNAs by EDC (N-(3-Dimethylaminopropyl)-N'-ethylcarbodiimide hydrochloride) which improved the sensitivity of detection in particular for low abundant small RNAs. Signals for three miRNAs in particular-miR160, miR166, miR167 could be enhanced specifically in the primary roots in contrast

to non-EDC treated roots where levels of miRNAs were low or below detection level. The method allows the detection of relative differences in the abundance of small RNAs or to profile small RNAs in mutant backgrounds. The expression patterns obtained for the different miRNAs corroborated the ones inferred using indirect methods (miR390, miR166, miR167, miR160 (Marin *et al.*, 2010; Breakfield *et al.*, 2011; Carlsbecker *et al.*, 2010)). In addition, the comparison of the expression patterns between the embryo and the seedling revealed the dynamics of miR172 expression, detected in embryo but less abundant in young plants. Importantly, the protocol is amenable to high-throughput applications since the whole procedure can be performed using liquid-handling robotic systems. This improved method allows the systematic determination of a wide range of small RNAs localization patterns and subsequent inference of their functions.

Experimental procedures

195 Plant material.

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- 196 All lines used in this study are in the *Arabidopsis thaliana* Col-0 ecotype background. All mutant used
- 197 have been previously described: mir390a-1 (Cuperus et al., 2009), mir390a-2 (Marin et al.,
- 198 2010),rdr6 (sgs2-1)(Elmayan et al., 1998; Mourrain et al., 2000), dcl4-1(Xie et al., 2005), tas3a-1,
- 199 OXTAS3 (Marin et al., 2010), drb4 (Adenot et al., 2006), hen1-5 (Park et al., 2002). Embryos were
- dissected in a drop of PBS and immediately transferred to 4% paraformaldehyde on ice.
- 202 Plant growth conditions.
- 203 Plants were grown on 0.5X-MS/0.8% agar (MS-agar) plates in controlled-environment chambers
- 204 under the following conditions: 150 μ mol photon.m⁻².s⁻¹ luminance, 16 hr light, 23°C temperature.
- 206 Probe design.
- 207 LNA-modified oligonucleotide probes were synthesized by Exigon. The probes corresponds to the
- full antisense sequence of the miRNA, are DIG-labeled at both the 5' and 3' ends and contain 4
- 209 LNA-modified base at positions 7, 9, 11, 15 (Table 1).
- 211 Whole mount in situ hybridization
- 212 Only the main steps of the method are described here, a detailed protocol is available as
- 213 supplemental material. Tissues were cross-linked with para-formaldehyde (4% solution) by vacuum
- infiltration until the samples sunk and became translucent in color. Length of fixation varied according
- 215 to the tissue. Root and embryo samples are typically fixed for 45 minutes and 2 hours at room
- 216 temperature, respectively. Once fixed, samples are permeabilized and chlorophyll was removed by
- 217 a 1:1 mixture of 100% ethanol and Histo-Clear®. Cleared root tissues and embryos were then
- 218 digested by either 125 μ g/mL proteinase K for 30 minutes or 75 μ g/mL proteinase K for 15 minutes,
- 219 respectively. For the optional EDC (N-(3-Dimethylaminopropyl)-N'-ethylcarbodiimide hydrochloride)
- 220 cross-linking step, incubation was done at 60°C for 2hrs at 0.16M concentration. The optimal
- 221 hybridization conditions (temperature and probe concentration) were optimized for each probe. For
- seedlings, probes were typically hybridized at 55°C or 65°C at 10nM for 16h. For embryos, all probes
- were hybridized at 65°C with 20nM of probe for 16h.After hybridization, samples were washed and
- incubated with Digoxigenin antibodies followed by NBT/BCIP incubation in the dark for colorimetric
- detection. The time of revelation depends on the probe. For root and embryo samples, revelation
- 226 was done for 1 hour (miR390), 2-4 hours (miR166, miR167 and miR172), 3 hrs for ta-siRNA and 6
- 227 hours (miR160). As negative controls, *in situ* hybridizations using animal-specific miR124 probes
- 228 were performed on root and embryos alongside and under the exact same conditions as the miRNAs
- 229 tested.

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231 Imaging: 232 For imaging roots, samples were mounted in a 1:1 mixture of glycerol and 1X TE buffer and detected 233 by light microscopy using Nomarski (Differential Interference Contrast, DIC) from Nikon with a 20X objective. After stopping the colorimetric detection, embryos were transferred to three-welled glass 234 235 slides (Electron Microscopy Science Cat. No. 63418-11), mounted in 70% glycerol/TE buffer and sealed with cover-slips. Slides were subsequently imaged on an automated Pannoramic SCAN 150 236 237 slide scanner (3DHISTECH) with transmitted light and a 20X plan-apochromat objective. Images of 238 embryos were collected using the Panoramic Viewer software (3DHISTECH). 239 240 Northern blotting: 241 Northern blotting was performed as described (Marin et al., 2010) 242 **ACKNOWLEDGEMENTS** 243 244 We thank A. Bleckmann for her comments on the manuscript. This work was supported by the 245 Land Baden-Württemberg, the Chica und Heinz Schaller Stiftung, the CellNetworks cluster of 246 excellence and the Boehringer Ingelheim (to M.G.D, A.M.), and the framework of the first call ERA-NET for Coordinating Plant Sciences (ERA-CAPS) funded by the European Union in the 247 248 framework of the FP7 under the Grant Agreement n°291864, with funding from the Austrian 249 Science Fund (FWF grant number 1476-B16) (M.M and M.D.N). 250 251 **CONFLICT OF INTEREST STATEMENT** 252 The authors declare no conflict of interest. 253 **SUPPORTING INFORMATION** 254 255 Additional Supporting Information may be found in the online version of this article. 256 **REFERENCES** 257 258 Adenot, X., Elmayan, T., Lauressergues, D., Boutet, S., Bouche, N., Gasciolli, V. and 259 Vaucheret, H. (2006) DRB4-dependent TAS3 trans-acting siRNAs control leaf morphology 260 through AGO7. Curr Biol, 16, 927-932. Breakfield, N.W., Corcoran, D.L., Petricka, J.J., Shen, J., Sae-Seaw, J., Rubio-Somoza, I., 261 262 Weigel, D., Ohler, U. and Benfey, P.N. (2011) High-resolution experimental and 263 computational profiling of tissue-specific known and novel miRNAs in Arabidopsis., 22, 163-176. Available at: 264 265 http://eutils.ncbi.nlm.nih.gov/entrez/eutils/elink.fcgi?dbfrom=pubmed&id=21940835&retmode=r ef&cmd=prlinks. 266

- 267 Carlsbecker, A., Lee, J.-Y., Roberts, C.J., et al. (2010) Cell signalling by microRNA165/6 directs
- gene dose-dependent root cell fate. *Nature*, **465**, 316–321.
- 269 Cartolano, M., Castillo, R., Efremova, N., Kuckenberg, M., Zethof, J., Gerats, T., Schwarz-
- Sommer, Z. and Vandenbussche, M. (2007) A conserved microRNA module exerts homeotic
- control over Petunia hybrida and Antirrhinum majus floral organ identity. *Nat Genet*, **39**, 901–
- 272 905.
- 273 Chitwood, D.H., Nogueira, F.T.S., Howell, M.D., Montgomery, T.A., Carrington, J.C. and
- Timmermans, M.C.P. (2009) Pattern formation via small RNA mobility. *Genes Dev.*, 23, 549–
- 275 554.
- 276 Cuperus, J.T., Montgomery, T.A., Fahlgren, N., Burke, R.T., Townsend, T., Sullivan, C.M. and
- **Carrington, J.C.** (2009) Identification of MIR390a precursor processing-defective mutants in
- 278 Arabidopsis by direct genome sequencing. *PNAS*.
- 279 Douglas, R.N., Wiley, D., Sarkar, A., Springer, N., Timmermans, M.C.P. and Scanlon, M.J.
- 280 (2010) ragged seedling2 Encodes an ARGONAUTE7-like protein required for mediolateral
- expansion, but not dorsiventrality, of maize leaves., **22**, 1441–1451.
- 282 **Elmayan, T., Balzergue, S., Béon, F., et al.** (1998) Arabidopsis mutants impaired in
- 283 cosuppression., **10**, 1747–1758.
- 284 Gasciolli, V., Mallory, A.C., Bartel, D.P. and Vaucheret, H. (2005) Partially redundant functions
- of Arabidopsis DICER-like enzymes and a role for DCL4 in producing trans-acting siRNAs.
- 286 *Curr Biol*, **15**, 1494–1500.
- 287 Husbands, A.Y., Benkovics, A.H., Nogueira, F.T.S., Lodha, M. and Timmermans, M.C.P.
- 288 (2015) The ASYMMETRIC LEAVES Complex Employs Multiple Modes of Regulation to Affect
- Adaxial-Abaxial Patterning and Leaf Complexity [OPEN]. THE PLANT CELL ONLINE,
- 290 tpc.15.00454–16.

Jones-Rhoades, M.W., Bartel, D.P. and Bartel, B. (2006) MicroRNAS and their regulatory roles

- 292 in plants. *Annu Rev Plant Biol*, **57**, 19–53.
- 293 Juarez, M.T., Kui, J.S., Thomas, J., Heller, B.A. and Timmermans, M.C.P. (2004) microRNA-
- mediated repression of rolled leaf1 specifies maize leaf polarity. *Nature*, **428**, 84–88.

295 Kidner, C.A. and Martienssen, R.A. (2004) Spatially restricted microRNA directs leaf polarity

- 296 through ARGONAUTE1. *Nature*, **428**, 81–84.
- 297 Kloosterman, W.P., Steiner, F.A., Berezikov, E., de Bruijn, E., van de Belt, J., Verheul, M.,
- 298 Cuppen, E. and Plasterk, R.H. (2006) Cloning and expression of new microRNAs from
- zebrafish. Nucleic Acids Research, 34, 2558–2569.
- 300 Marin, E., Jouannet, V., Herz, A., Lokerse, A.S., Weijers, D., Vaucheret, H., Nussaume, L.,
- 301 Crespi, M.D. and Maizel, A. (2010) miR390, Arabidopsis TAS3 tasiRNAs, and their AUXIN
- 302 RESPONSE FACTOR targets define an autoregulatory network quantitatively regulating lateral
- 303 root growth., **22**, 1104–1117.

- 304 Miyashima, S., Koi, S., Hashimoto, T. and Nakajima, K. (2011) Non-cell-autonomous
- 305 microRNA165 acts in a dose-dependent manner to regulate multiple differentiation status in
- 306 the Arabidopsis root., **138**, 2303–2313.
- 307 Mourrain, P., Béclin, C., Elmayan, T., et al. (2000) Arabidopsis SGS2 and SGS3 genes are
- required for posttranscriptional gene silencing and natural virus resistance. *Cell*, **101**, 533–542.
- Nodine, M.D. and Bartel, D.P. (2010) MicroRNAs prevent precocious gene expression and
- enable pattern formation during plant embryogenesis. *Genes Dev.*, **24**, 2678–2692.
- Nogueira, F.T.S., Chitwood, D.H., Madi, S., Ohtsu, K., Schnable, P.S., Scanlon, M.J. and
- Timmermans, M.C.P. (2009) Regulation of small RNA accumulation in the maize shoot apex.
- 313 *PLoS Genet.*, **5**, e1000320.
- Nogueira, F.T.S., Madi, S., Chitwood, D.H., Juarez, M.T. and Timmermans, M.C.P. (2007) Two
- small regulatory RNAs establish opposing fates of a developmental axis. *Genes Dev.*, **21**,
- 316 750–755.
- 317 Ori, N., Cohen, A.R., Etzioni, A., et al. (2007) Regulation of LANCEOLATE by miR319 is required
- for compound-leaf development in tomato. *Nat Genet*, **39**, 787–791.
- Pall, G.S., Codony-Servat, C., Byrne, J., Ritchie, L. and Hamilton, A. (2007) Carbodiimide-
- mediated cross-linking of RNA to nylon membranes improves the detection of siRNA, miRNA
- and piRNA by northern blot. *Nucleic Acids Research*, **35**, e60–e60.
- Parizotto, E.A., Dunoyer, P., Rahm, N., Himber, C. and Voinnet, O. (2004) In vivo investigation
- of the transcription, processing, endonucleolytic activity, and functional relevance of the spatial
- distribution of a plant miRNA. *Genes Dev.*, **18**, 2237–2242.
- 325 Park, W., Li, J., Song, R., Messing, J. and Chen, X. (2002) CARPEL FACTORY, a Dicer
- Homolog, and HEN1, a Novel Protein, Act in microRNA Metabolism in Arabidopsis thaliana.
- 327 *Curr Biol*, **12**, 1484–1495.
- Pena, J.T.G., Sohn-Lee, C., Rouhanifard, S.H., et al. (2009) miRNA in situ hybridization in
- formaldehyde and EDC-fixed tissues. *Nat. Methods*, **6**, 139–141.
- 330 Rubio-Somoza, I., Cuperus, J.T., Weigel, D. and Carrington, J.C. (2009) Regulation and
- functional specialization of small RNA-target nodes during plant development. Curr. Opin.
- 332 *Plant Biol.*, **12**, 622–627.
- 333 Sunkar, R., Chinnusamy, V., Zhu, J. and Zhu, J.-K. (2007) Small RNAs as big players in plant
- abiotic stress responses and nutrient deprivation. *Trends Plant Sci.*, **12**, 301–309.
- 335 Válóczi, A., Várallyay, E., Kauppinen, S., Burgyán, J. and Havelda, Z. (2006) Spatio-temporal
- accumulation of microRNAs is highly coordinated in developing plant tissues., 47, 140–151.
- Wheeler, G., Válóczi, A., Havelda, Z. and Dalmay, T. (2007) In situ detection of animal and plant
- 338 microRNAs. *DNA Cell Biol*, **26**, 251–255.
- 339 Wollmann, H., Mica, E., Todesco, M., Long, J.A. and Weigel, D. (2010) On reconciling the
- interactions between APETALA2, miR172 and AGAMOUS with the ABC model of flower

development., 137, 3633–3642.
Xie, Z., Allen, E., Wilken, A. and Carrington, J.C. (2005) DICER-LIKE 4 functions in trans-acting small interfering RNA biogenesis and vegetative phase change in Arabidopsis thaliana. *PNAS*, 102, 12984–12989.
Yu, N., Niu, Q.-W., Ng, K.-H. and Chua, N.-H. (2015) The role of miR156/SPLs modules in Arabidopsis lateral root development. *Plant J.*, 83, 673–685.

TABLES

Table 1. Different microRNAs and their probe sequences, hybridization temperatures and probe concentrations

miRNA	Probe sequence	Hybridiza	Probe
		tion	conc.
		temp.	
miR160	/5DigN/TGGCATA+CA+GG+GAG+CCAGGCA/3Dig_N/	55°	10nM
miR124	/5DigN/ACTGATA+TC+AG+CTC+AGTAGGCAC/3Dig_N/	65°	10nM
miR166	/5DigN/GGGGAAT+GA+AG+CCT+GGTCCGA/3Dig_N/	65°	10nM
miR167	/5DigN/TAGATCA+TG+CT+GGC+AGCTTCA/3Dig_N/	65°	10nM
miR172	/5DigN/ATGCAGC+AT+CA+TCA+AGATTCT/3Dig_N/	55°	10nM
miR390	/5DigN/GGCGCTA+TC+CC+TCC+TGAGCTT/3Dig_N/	65°	10nM
tasiARF	/5DigN/AAGGCCT+TA+CA+AGG+TCAAGAA/3Dig_N/	65°	10nM

Figures and legends

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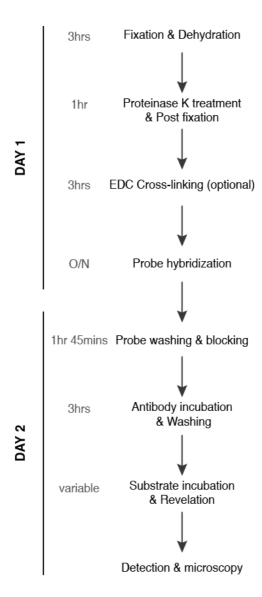


Figure 1. Overview of the protocol

A detailed step-by-step description of the procedure can be found in the supplemental material.

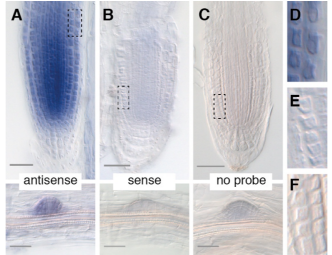


Figure 2. Visualization of miRNAs in whole mount Arabidopsis roots

Whole mount *in situ* hybridization of double digoxigenin labelled locked nucleic acid miRNAs probes carried out on *Arabidopsis thaliana* (wild type, Col-0) roots. miR390 signals were specifically detected both in primary root (upper panels) and lateral root primordia (lower panel) only with the antisense probe (A). Little to no signal were detected with the sense (B) or in absence of probe (C). (D-F) Higher magnification views of the areas delineated in (A-C). Scale bars are 50μ m.

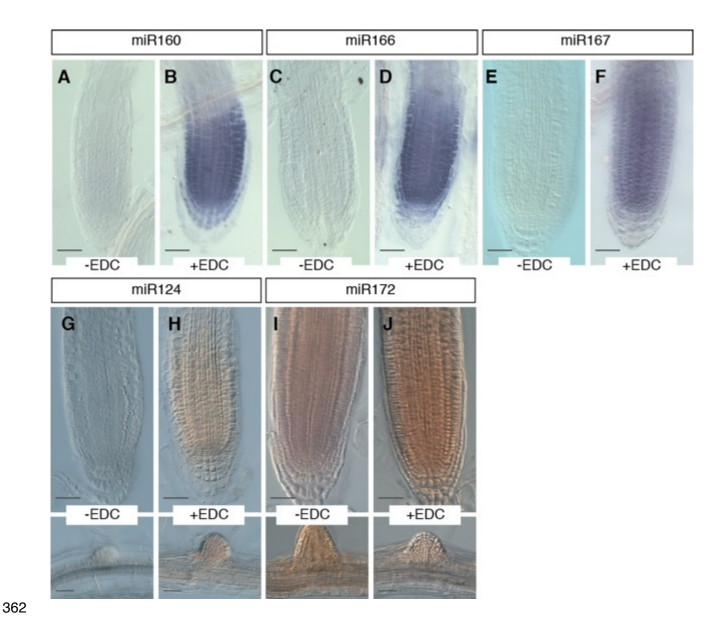


Figure 3. Increased sensitivity by use of EDC fixation

Expression of miR160, miR166, miR167, miR124 and miR172 in absence (A, C, E, G, I) and in presence of EDC crosslinking (B, D, F, H, J). EDC-treatment allows the detection of miR160, 166 and 167 signal. In the same conditions, the mouse miRNA miR124 and the Arabidopsis miR172 do not show any signal. Scale bars are 50 μ m.

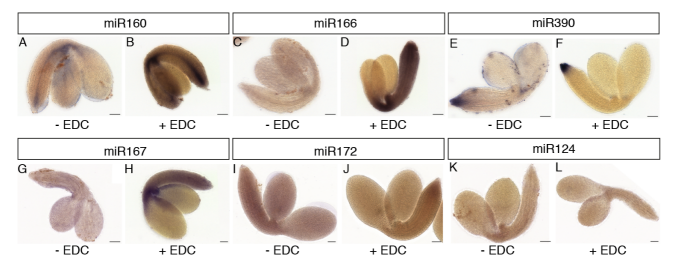


Figure 4. miRNA detection in embryos

Images of embryos hybridized with probes corresponding to miR160, miR166, miR390, miR167 and miR172 *in situ* hybridizations in either the absence (A, C, E, G, I) or presence of EDC-crosslinking (B, D, F, H, J) are shown. Probes against the mouse miRNA miR124 were used as negative controls and were performed either without (K) or with (L) EDC-crosslinking. Scale bars are 50 μ m.

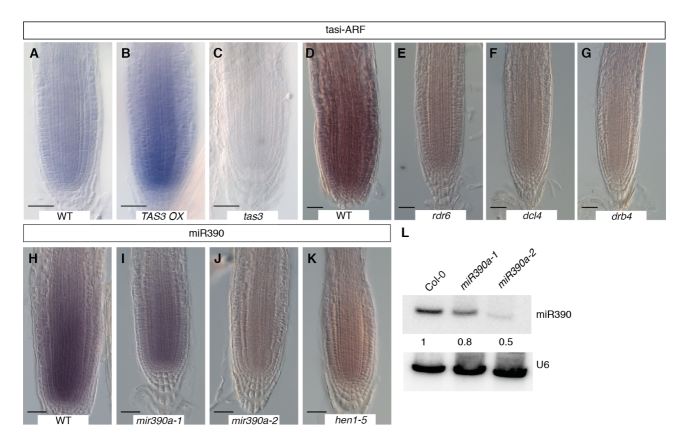


Figure 5. Semi-quantitative detection of small RNA abundance.

(A-G) Detection of *TAS3*-derived ta-siARFs ta-siRNAs in roots of wild type plants (A, D), in plants expressing the precursor *TAS3* (B) or in plants impaired in ta- siRNAs biogenesis such as the *tas3* mutant (C), the *rdr6* (E), *dcl4* (F) or *drb4* (G) mutants. (H-K) Detection of miR390 in wild type (H) plants, a in mutant partially affected in miR390 processing from the *MIR390a* precursor (*mir390a-1*, I), in a mutant with reduced levels of the precursor (*mir390a-2*, J), in a mutant destabilizing mature miR390 by lack on terminal methylation (*hen1-5*, K). Signals are reduced in the *miR390a-1* mutant and absent in *mir390a-2* and *hen5* mutants. Scale bars are 50µm. (L) Northern blot for miR390 in root RNA of 7-day old wild type, *mir390a-1* and *miR390a-2* seedlings confirming the reduction of miR390 abundance in these mutants. U6 serves as loading control. Values indicate relative abundance of miR390 compared to wild type (Col-0).