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## Microrna Regulation of Nodule Zone-Specific Gene Expression in Soybean

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#### Article Info

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#### Abstract:

Nitrogen is a paramount important essential element for all living organisms. It has been found to be a crucial structural component of proteins, nucleic acids, enzymes and other cellular constituents which are inevitable for all forms of life. In the atmosphere, the percentage of nitrogen is very high (N<sub>2</sub>, 78%) compared to other inorganic gases. However, most organisms have practically no direct access to this nitrogen. While plants can not directly uptake nitrogen from atmosphere, they are capable of assimilating other forms of nitrogen, for example ammonium (NH<sub>4</sub><sup>+</sup>) and nitrate (NO<sub>3</sub><sup>-</sup> ). For agricultural crop production, artificial fixation of nitrogen is heavily utilized, and it is an expensive process that requires high temperatures (at least 400 °C) and pressures (around 200 atm). It has been conspicuously demonstrated that indiscriminate use of fertilizer hampers soil physical, chemical and micro biological properties and also a potential risk to environment e.g. water quality. Besides, chemically manufactured fertilizers are depleted from soils in various ways, for instance; denitrifying bacteria, volatilization, and leaching. Consequently, it results relatively poor availability of nitrogen to get into plants. On the flipside, only 1-2% of the nitrogen fixation in the world occurs through the natural process of lightening. Notably, microbial fixation is well characterized in diazotrophs for example; Rhizobia and Frankia, and blue-green algae. Against the backdrop, we are accentuated on an environmentally friendly and the most sustainable approach to increase productivity for legume and non-legume crops. Till today, the term biological nitrogen fixation (BNF) has received much attention as a sustainable alternative; this process facilitates atmospheric nitrogen to convert into ammonia by rhizobia in specialized plan organs termed "root nodules". This review article seeks to better understand plant mechanisms involved in the development of root nodules in soybean.

#### Introduction:

Soybean (Glycine max) is one of the most important oil crops and a source of animal feed protein in the world. It has a salient feature to fix atmospheric nitrogen through symbioses with compatible rhizobia that yields to determinate type nodule (Oldroyd, Murray et al. 2011). Biological nitrogen fixation in soybean nodules reduces the use of chemical nitrogen fertilizers resulting in cost-savings to producers and minimizes environmental damage due to nitrogen run-off. A better understanding of how nodules form and function is important for selection or generation of soybean genotypes with better nitrogen fixation capacity. Soybean nodules originate from root cortex via de novo cell differentiation (Oldroyd 2013). Consequently, two major nodule development zones are formed for instance; the nodule primordium (Npr) in the middle and it is encircled by nodule parenchyma (Npa). At later time point, the Npr gives rise to N-fixation zone and the Npa holds vascular bundles. It is not clear what early signaling pathways driving the conspicuous development of the nodule zones. My research is aimed at filling this knowledge gap by illustrating the molecular signatures that paves the way to cellular differentiation in root nodule development in soybean. Based on initial evidence obtained by the Subramanian lab, we hypothesize that microRNAs (miRNAs) play important regulatory roles in spatio-temporal expression of their target genes during nodule developmental in soybean. For instance, the regulation of auxin sensitivity by miR160 has been found to be crucial for formation of nodule primordia and vasculature in the parenchyma (Marie Turner 2013). Against this backdrop, this review article focused on nuclear and cytoplasmic transcriptome as well

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determine the relative abundance and differentially expressed bacteria multiply within the host cells and differentiate into the mRNAs and regulatory role of miRNAs in cell differentiation and nitrogen fixing bacteroids (Udvardi and Day 1997) (Oldroyd nodule development.

# nitrogen:

Atmospheric nitrogen percentage is very high (N2, 78%) uncovered thousands of genes to be induced during Nod factor compared to other inorganic gases (Mary Elvira 1932). However, signaling and its resulting ethylene (ET) biosynthesis throughout most of the organisms have practically no direct access to this the multiple development stages of indeterminate nodule nitrogen. Nevertheless, plants can not directly uptake nitrogen (Larrainzar, Riely et al. 2015). Albeit the molecular mechanism of from atmosphere, but they are capable of assimilating only very such regulation is not well understood. There has been a large-scale specific forms of nitrogen, for example ammonium (NH4+) and transcriptome analysis of B. japonicum-inoculated and mocknitrate (NO3-) (Bytnerowicz and Fenn 1996, Peter M. Vitousek inoculated soybean root hairs. It has showed that a total of 1,973 1997) (Sponseller, Gundale et al. 2016). Virtually, nitrogen has soybean genes differentially expressed during root hair infection, been found to be a crucial structural component of proteins, nucleic particularly NFR5 and NIN genes (Libault, Farmer et al. 2010). acids, enzymes, and other cellular constituents which are inevitable Nevertheless, the signaling mechanisms directing the cellular for all forms of life (O'Brien, Vega et al. 2016). For agricultural differentiation of nodule are not known. crop production, artificial fixation of nitrogen is heavily utilized. It is an expensive process that requires high temperatures (approx. Sovbean root nodule organogenesis: 400 °C) and pressures (approx. 200 atm) (Witschi 2000). It has been conspicuously demonstrated that indiscriminate use of N Soybean (Glycine max) has a genome size of 1.1 to1.5 Gb, it is fertilizer hampers the diversity of the bacterial community and partially diploidized tetraploid. It decreases soil C and N concentrations (Verzeaux, Alahmad et al. is one of the most important oil crops and a source of animal feed 2016). Notably, it has been demonstrated as a potential risk to protein in the world (soybase.org/sb\_about.php). It has a salient environment e.g. water quality (Zhao, Sha et al. 2016) (Sponseller, feature to fix atmospheric nitrogen through symbioses with Gundale et al. 2016). Besides, chemically manufactured fertilizers compatible rhizobia that yields to determinate type nodule are depleted from soils in various ways, for instance; denitrifying (Udvardi and Day 1997) (Oldroyd, Murray et al. 2011). bacteria, volatilization, and leaching (Johnson 1996, Peter M. Notwithstanding of the economic and environmental importance, Vitousek 1997). Consequently, it results relatively poor there has been very few studies about quantitative trait loci (QTL) availability of nitrogen to get into plants. On the flipside, over 90 that controlling BNF traits, for instance nodule number, ration of % of the nitrogen fixation in the world occurs through the natural nodule dry weight with nodule number, and shoot dry weight process of lightening and microorganisms. Furthermore, microbial (SDW). It has been reported via composite interval mapping that fixation is well characterized in diazotrophs for example; Rhizobia approximately six QTLs bears very small effect on BNF traits and Frankia, and blue-green algae (Cheng 2008). It has been (Santos, Geraldi et al. 2013). Besides, it has been demonstrated in demonstrated that Bradyrhizobium strains substantially escalated earlier studies that nodules originate from root cortex via de novo soybean grain yield, and protein content up to 57% and 26%, cell differentiation into two different cell types, parenchymal and respectively (Zimmer, Messmer et al. 2016). Against the backdrop, primordium (Celine Charon 1997) (Oldroyd & Downie 2008; we are accentuated on an environmentally friendly and a Oldroyd 2013). In addition, early nodulin genes in legume for sustainable approach to increase the productivity for legume and instance; Enod 40 gene reported to be expressed in root pericycle non-legume crops. Literature mining depicted that biological during the rhizobia infection and later it occupied in the dividing nitrogen fixation in soybean nodules reduces the use of chemical cortical cells (H. Kouchi and S. Hata 1993). Among the two major nitrogen fertilizers resulting in cost-savings to producers and nodule development zones, the nodule primordium (Npr) in the minimizes environmental damage due to nitrogen run-off.

#### Rhizobia infection leads to the root nodule development:

factors. This signal is perceived by the LysM receptor like kinase yielding to biological N2 fixation (Li, Zhao et al. of host plant, it induces the root hair curling, and bacteria get 2015). Even though, it is not clear what early signaling pathways interaction, bacteria are released from the infection threads into the primordia (Marie Turner 2013).

miRNA profiles of parenchyma and primordial tissues and plant cells, surrounded by membrane of plant origin. These 2013). Till now, integration of genetic and genomic approaches has revealed twenty-six genes to be involved in nodule Root nodule a sustainable alternative to fix atmospheric development of Medicago truncatuala and Lotus japonicum (Kouchi, Imaizumi-Anraku et al. 2010). In addition, deep sequencing of the Medicago truncatula root transcriptome has

middle which is encircled by nodule parenchyma (Npa). At later time point, the Npr gives rise to N-fixation zone and the Npa holds vascular bundles. Lately, a β- expansin gene, GmEXPB2 fused with GUS reporter gene which was observed to be preferentially In the natural environment, plants are continuously confronted expressed in nodule vascular trace and nodule vascular bundles. It with pathogenic and symbiotic microbes. Symbioses involves indicated that GmEXPB2 might be crucial for nodule mutual exchange of diffusible signal molecules, first endophytic organogenesis. Over expression of GmEXPB2 contrast to bacteria (rhizobia) are attracted by the plant root exudates suppressed GmEXPB2 transgenic lines found to be escalated flavonoids which are perceived and triggered the bacterial nodule number, nodule mass and nitrogenase activity. It further nodulation (nod) genes. Consequently, the bacteria synthesize suggested that GmEXPB2 might have influenced over root specific lipochito-oligosaccharides, called nodulation (Nod) architecture, nodule formation and development, and profoundly

access into the host epidermis through infection threads (ITs) and driving the conspicuous development of the nodule zones. Against initiate cell division within the root cortex, leading to the the backdrop, to understand the regulation of auxin sensitivity by progression of the root nodule meristem. In later stages of the miR160 which is believed to be crucial for the formation of nodule



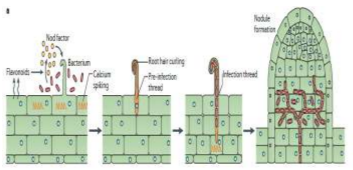


Figure 1a: Illustrating the progression of root nodule development through Rhizobial bacterial infection in the plant root leading to the determinate nodule (Oldroyd 2013).

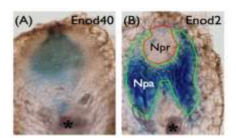


Figure 1b: Nodule development zones A. Nodule primordial zone (Enod 40 gene) in the middle B. surrounding parenchyma (Enod 2 gene), differentiated from cortex (collected from Sen Subramanian

#### Regulatory small RNAs biogenesis and its molecular functions:

Regulatory small RNAs are ranged between 20 to 24 nucleotides which are ubiquitous elements of endogenous transcriptomics, a common response to exogenous viral infections and introduced double-stranded RNA (Axtell 2013). Three core enzymes families, for instance; RNAdependent RNA polymerase (RDR), Dicer like (DCL), and Argonaute (AGO) proteins paves the way of small RNA biogenesis and function in plants. Firstly, ribonuclease type III or DICERLIKE1 involves in the yield of a Figure 3: Gene expression events occurring in typical plant cell fold-back precursor RNA or primary miRNA (primiRNA) transcripts using an RNA templates in the nuclei. Later, the It has been found in several studies that most plant miRNAs are resulting miRNA-miRNA duplex which is originated in nucleus non-coding RNA, and small 21-24 nucleotide long (Cuperus, then translocated into cytoplasm. The guided miRNAmolecule is Fahlgren et al. 2011). It requires DCL1-clade DCL for their incorporated into ARGONAUTE (AGO) to form an active RISC biogenesis and AGO1-clade AGO for their function (Wu, Zhou et complex to specific target RNAs that are complementary to the al. 2010, Manavella, Koenig et al. 2012). miRNA, and this process eventually follows up mRNA cleavage, In rice (Oryza sativa), DCL3 has been reported in the biogenesis represses the translation of the mRNAs or Chromatin modification. of 24nt long miRNA that This phenomenon accentuated as an inhibition or silencing of the incorporated in AGO4 to regulate the target gene expression gene expression, which play a crucial role in the developmental primarily through mRNA cleavage process in plant and animal (Chapman and Carrington 2007) (Wu, Zhou et al. 2010). Argonaute proteins (AGO) form RNA (Axtell 2013).

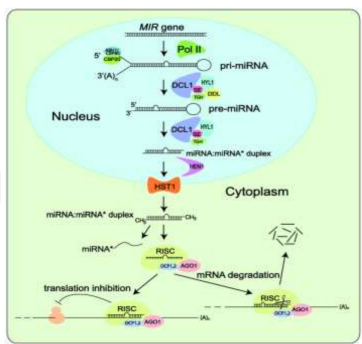
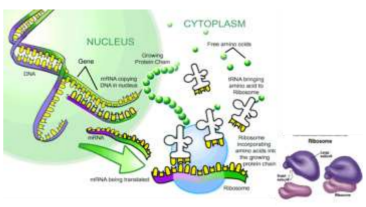


Figure 2: Regulation of gene expression events via RISC complex



inducing silencing complexes (RISC) with small RNAs which is known as post-transcriptional gene silencing. It has typically four domains, for instance: N-terminal, PAZ, MID and PIWI domains. The MID-PIWI lobes are belongs to the C-terminus. It has been studied that MID-domains contains the specificity loop to recognize and bind to the 5'-phosphate of smRNAs. The PIWI domains contained the catalytic active site D-E-D-H/D. PAZ domain anchored the 2-nt overhang at the 3' end of miRNAs. The N-terminal domain involved in the separation of miRNA-miRNA duplex and the slicer activity of the mRNA (Song, Smith et al. 2004). There has been an expansion and duplications of AGO



family members during plant evolution (Singh, Gase et al. 2015). soybean mosaic virus and among them

The functional diversification of AGOs is indicating 71 genes projected to induce in defense response (Hui Chen 2016). sRNAdirected regulatory pathways. The binding preference of These findings suggested the

Arabidopsis, 10 AGO have been extensively studied (Liu et al. during viral infection.

recognized distinct structural features in miR165/miR166 duplex Soybean- Bradyrhizobium

meristem by decoying miR165/miR166 and subsequent repression Subramanian et al. 2008. They sequenced

Hu et al. 2011). Notably, 22 AGO proteins have been reported in were inoculated with B.

Soybean (Glycine max). It has been found that genome duplication japonicum. It helps to detect 20 conserved miRNAs loci based on in Soybean resulted such a proliferation of AGOs. For example: its the similarity to miRNAs in

genome encodes two copies of AGO1, AGO2, AGO5, AGO4/9, another plant species. In addition, 35 novel miRNAs were AGO6 and AGO7 (Xiang Liu 2014). However, the molecular identified based on potential hairpin

function of the plant AGO genes yet not very clear.

There are several miRNA families that are conserved across the sequences (Subramanian, Fu et vast evolutionary distances from

flowering plants to mosses (Cuperus, Fahlgren et al. 2011). It has in the regulation of legume rhizobium symbiosis.

been observed in another study

that miRNA, and its target pairing found to be stable for a identified in soybean by Turner prolonged periods of plant evolution.

On the flip side, another group demonstrated that conserved plant miR160, miR164 and miR393 miRNAs and their targets are to somehow flexible. For instance; found to be involved in auxin signaling (Turner, Yu et al. 2012).

miR159 is a highly conserved miRNA that targets not only a

mRNA, SGN-U567133 (Buxdorf, Hendelman et al. 2010). A determinate and indeterminate

higher level of the SGN-U567133 transcript and exhibited defects family members that governs in leaf and flower development. This result suggests that miR159 the formation of root nodule.

involves in a post-transcriptional regulation. Additionally, it is It has been illustrated that auxin receptor gene family hushed by found to be crucial for the normal tomato development.

Recently, the identification of miRNAs in the regulation of These plant roots found to be hypersensitive to auxin and yielded photoperiodic pathways in soybean

have been reported through high throughput sequencing and qRT- advocated that only minimal/reduced auxin signaling is required PCR. Six libraries were

under short day treatment, similar

of 163 miRNAs families were

novel predicted miRNAs. As

expected, significant differences in abundance between short day Turner and long day treatment was

observed (Wenbin Li 2015). These findings provided evidence of Organ specific expression of profile of miRNA and the potential miRNA in the regulation of

flowering time that ultimately affects the seed yield and quality of genes (Glyma10g10240 and Glyma17g05920) which were the

The complex regulatory network of miRNA-mRNA interactions highly expressed in soybean nodule. Likewise, three potential during viral infection has been

revealed via small RNA seq (sRNA), degradome seq, and genome- demonstrated to be highly expressed in the nodules than in the wide transcriptome analysis.

folds abundance compared with

mock-inoculated control demonstrated through sRNA seq and nodules. analysis. Among them 105 miRNAs

were identified as potential targets of 125 transcripts that has been namely, miR482, miR1512, and validated by degradome seq

analyses. In addition, 2679 genes were detected via genome wide numbers while root length and transcriptomic analysis. These

genes have been differentially expressed during infection of expected, there were differential

AGO and sRNA is mainly assigned by the sequence of sRNA. In regulatory role miRNA that governed the target gene expression

2014). It has been demonstrated that AtAGO10 like AtAGO1, it Furthermore, the regulatory role of microRNAs (miRNAs) during

than involved by AtGO1. AtAGO10 found to regulate shoot apical japonicum mutualistic association was studied first by

of homeodomain-leucinezipper (HD-ZIP) gene expression (Zhu, approximately 350000 small RNAs of soybean root sample which

forming precursors in Soybean EST as well as shotgun genomic

al. 2008). These findings advocated the potential role of miRNAs

In another study, 120 hairpin-forming precursor genes have been

et al. In addition, they reported three novel miRNAs for instance;

Moreover, the plant hormone subset of MYB mRNAs but also observed to target a non MYB auxin is thought to have a pivotal role in nodule organogenesis in

mutant tomato transgenic line (miR159-resistant line) showed type of nodule. It indicates a redundancy and diversity of miRNAs

overexpressed microRNA393.

normal nodule. This observation

for determinate nodule

constructed using Illumina Solexa, for instance; 0, 8, and 16 h development. Likewise, overexpressed microRNA160 hushed a set of repressor auxin response

time points considered for the long the long day treatment. A total transcription factor. These plant roots were hypersensitive to auxin and observed not to be reluctant

reported which covered 318 plant miRNAs, and unclassified 81 in epidermal responses to rhizobia. Notably, it yielded to lower sized nodule primordium (Marie

> 2013). This observation indicated that auxin hypersensitivity inhibits nodule organogenesis

targets were also studied. Two

target of miR169 but detected to be

targets of gma-new-miR13587

roots. As expected, gma-newmiR13587 found to be poorly There has been a total of 253 soybean miRNAs found to be two-expressed in the nodules than in the roots (Turner, Yu et al. 2012). There was an inverse expression pattern observed in between roots

Li et al., studied the transgene expression of three novel miRNAs

miR1515 in Soybean. They noticed a significant increase of nodule

later root density were normal in all tested miRNA lines. As



of these miRNAs in supernodulating expression nonnodulating soybean mutants. They

reported that 6 novel miRNAs decoyed 22 predicted target genes. ENOD40p:HF-GFP-RPL18 for primordial

And it was estimated via real

time polymerase chain reaction and qRT-PCR (Li, Deng et al. Glycine max root nodules that 2010). It advocates that miRNAs

have the signatory roles in soybean nodule development.

Sequencing of small RNAs and Parallel analysis of RNA ends studies which characterized (PARE) libraries revealed to

including 178 novel soybean

expressed in nodule tissue not in

and RNAi silencing approach to

formation (Zhe Yan 2015).

Therefore, this study showed a list of miRNAs and their potential nodule by Bradyrhizobium japonicum. target of nodulation genes.

In the model legume (Medicago truncatula), 25 conserved miRNA zones are formed for instance, the families and 100 novel miRNA

reads were detected by high-throughput sequencing. The nodule parenchyma (Npa). At later expression of MtHAP2-1 (encodes a

restricted by miR169a which is

nodule (Combier, Frugier et al.

overexpression of miR166, it

(Boualem, Laporte et al. 2008). To get

insights into key genes of nodule zones, transcript profiles of dai, 7 dai, 10 dai & 14 dai). specific cells/tissues were

M. truncatula using laser

capture micro dissection. It has been demonstrated from the of microRNA localization between nodule primordium and comprehensive gene expression map

Moling et al. 2013). These

controlled by the presence or

absence of miRNAs.

transformation has been applied as tool

type or tissue specific promoter

cloning technology to develop binary

vectors. INTACT method used to capture biotin tagged nuceli from obtain cytoplasmic transcriptomes data. specific cell types and TRAP

method used for profiling of mRNAs or foot printing of individual **Techniques to determine cell type specific expression profiles:** ribosomes (Ron 2014). TRAP

methodology is not required tissue fixation or single cell suspension. It has been successfully used

human cultured cells. Multiple

translation on a single mRNA. To

ribosomes, and polysomes can be

and al. 2014). In this study, we would

perform polysome isolation deploying cassettes

tissues, and ENOD2p:HF-GFP-RPL18 for parenchymal tissues in

express an epitope tagged version of ribosomal protein L18.

Over the last one decade, there has been several microarrays-based

transcriptional variations deployed in nodule formation. It has been identify 284 nodule miRNAs, more than 500 target genes, and embedded with couple of short comings, for instance; relative late time points study, incomplete representation of plant genes, miRNAs. It has been reported that ENOD93 only found to be discrimination of close paralogs, and reduced sensitivity. Lately, next generation sequencing technology have widened the horizon other plant parts of Soybean. Ectopic expression of miR393j-3p of transcription analyses in different legume species to detect symbiosis induced changes in late nodule developmental stages. ENOD93 expression showed a significant reduction in nodule Against this backdrop, we are accentuated to reveal early transcriptional changes induced in determinate type of soybean

In determinate type of nodule, two major nodule development

nodule primordium (Npr) in the middle and it is encircled by

time point, the Npr converted to N-fixation zone and the Npa CCAAT binding transcription factor) to meristematic zones was contained vascular bundles. Of these facts, it is not clear what early signaling pathways driving the conspicuous development of the found to be critical for the development of indeterminate type of nodule zones. In this context, mechanisms regulate the distinct gene expression profiles in Npr and Npa cell types has not 2006). In another study, HDZIPIII transcripts were inhibited by understood clearly. The proposed research study is aimed at filling this knowledge gap by illustrating the molecular signatures that dropped the number of symbiotic nodule and lateral root paves the way to cellular differentiation in root nodule development in soybean considering four different time points (5

The hypothesis is microRNAs (miRNAs) play important investigated at different time points from indeterminate nodules of regulatory roles in spatio-temporal expression of their target genes during nodule developmental in soybean. For example, a gradient parenchyma cells could result in distinct differentiation of these that selected genes enriched in different cell/tissue types (Limpens, cell types. To test this hypothesis, one has to obtain both cell typespecific miRNA and transcriptome (miRNA target) profiles. Since, findings indicated that organ specific gene expression could be the majority of miRNA regulation occurs in the cytoplasm, we reasoned that comparison of nuclear and ribosomal transcriptome profiles would reveal genes whose expression is potentially Recently, Agrobacterium rhizogenes mediated hairy root regulated by post transcriptional mechanisms such as miRNA cleavage. Combining this information with cell type-specific for exploring cell type specific gene expression in tomato. Cell miRNA profiles, and to test the above hypothesis and identify key miRNA-target pairs important for nodule cell differentiation. The introduced into INTACT and TRAP constructs via gateway use of translating ribosome affinity purification (TRAP) of nodule zone cells, namely from parenchyma and primordial tissues, to

## TRAP methods:

TRAP is termed translating ribosome affinity purification, to date in organisms ranging from D. melanogaster to mice and combines cell-type-specific transgene expression with affinity purification of translating ribosomes. It supersedes the need for ribosomes or Polyribosomes (polysomes) are engaged in tissue fixation, and facilitates to study the cell type-specific mRNA profiles of any genetically defined cell type. It has been evaluate the translation state of an mRNA, ribosomal subunits, successfully used to date in organisms ranging from D. melanogaster to mice, and human cultured cells. Multiple isolated from detergent-treated cell extracts (Heiman, Kulicke et ribosomes or Polyribosomes (polysomes) are engaged in



translation on a single mRNA. To evaluate the translation state of 15. Physiology 162(2013): 2042–2055.Oldroyd GE, Downie JA. an mRNA, ribosomal subunits, ribosomes, and polysomes can be isolated from detergent-treated cell extracts.

In this study, the polysome isolation using gene cassettes ENOD40p:HF-GFP-RPL18 for primordial tissues, ENOD2p:HF-GFP-RPL18 for parenchymal tissues in Glycine max root nodules that express an epitope tagged version of ribosomal protein L18 RPL18(Heiman, Kulicke et al. 2014, Ron 17. 2014).

Relative abundance and differentially expressed mRNAs profile in two different tissue specific zones would help to understand the 18. Sponseller, R. A., et al. (2016). "Nitrogen dynamics in effect of regulatory role of miRNAs in cell differentiation and nodule development.

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