***In situ*/Subcellular localization of arabinogalactan protein expression by fluorescent *in situ* hybridization, FISH.**

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**Summary**

The Arabinogalactan proteins are highly glycosylated and ubiquitous in plants. They are involved in several aspects of plant development and reproduction, however the mechanics behind their function remains for the most part unclear, as the carbohydrate moiety, covering the most part of the protein core, is poorly characterized at the individual protein level. Traditional immunolocalization using antibodies that recognize the glycosidic moiety of the protein cannot be used to elucidate individual proteins distribution, function or interactors. Indirect approaches are typically used to study these proteins, relying on reverse genetic analysis of null mutants or using a reporter fusion system. In the method presented here, we propose the use of RNA probes to assist in the localization of individual AGPs expression/mRNAs in tissues of Arabidopsis by fluorescent in situ hybridization, FISH. An extensive description of all aspects of this technique is provided, from RNA probe synthesis to the hybridization, trying to overcome the lack of specific antibodies for the protein core of AGPs.

**Keywords**
Whole-mount, immunolocalization, RNA-Probes, Arabidopsis, FISH, Arabinogalactan proteins

**Introduction**

Arabinogalactan proteins (AGPs) are cell wall glycoproteins, ubiquitous in the plant kingdom, that have been involved in several biological processes, namely in sexual plant reproduction. The structure of these proteins has an N-terminal sequence that targets the protein to the endoplasmic reticulum, a protein core rich in Pro/Hyp, Ala, Ser, Thr and a C-terminal sequence for the addiction of a glycosylphosphatidyl inositol (GPI) anchor. The protein core defines the different classes of AGPs as the classical, having this typical Pro/Hyp core between the N and the C-terminal sequence, the lysine-rich AGPs that have a lysine rich domain, the fasciclin-AGPs (FLAs) with fasciclin-like domains and finally the AG peptides containing small protein cores with no more than 30 aminoacids residues **(1)**.

For the past twenty years, several approaches have been developed to try to learn more about the biological way of action of these important class of proteins, but exactly due to their structure, with more than 90% of the molecule being sugars, this has been a hard task to perform. AGP glycans are polysaccharide chains O-glycosidically linked to the Hyp residues. The initial studies used the synthetic Yariv reagent that blocks AGP function, and clearly demonstrated the importance of these proteins in several biological processes. The use of monoclonal antibodies available for the glycosidic part of the AGPs have soon shown that AGPs could be used as molecular markers for different stages of plant development and for different cell/tissues involved in sexual plant reproduction and embryogenesis **(2–6)**.

The presence of the GPI anchor that anchors AGPs to the plasma membrane, facing the cell wall, the possibility of cleavage of the anchor by specific phospholipase C, associated with the complex sugar architecture that surrounds the protein core and the specific cell and tissue localization, makes these molecules important candidates for signaling events **(7)**.

This chapter describes in detail, one of the different procedures for study AGPs involved in sexual plant reproduction. The immunocytochemistry technique using the available collection of antibodies that recognize the different sugar epitopes present in AGPs **(8)** is very useful to impart important roles for these proteins in sexual plant reproduction **(2–6,9)**, but the use of other important molecular techniques became essential for unraveling the involvement of a particular AGP in a particular biological process. Very significant to learn about the function of a particular protein is to use reverse genetic analysis every time that null mutants are available **(10,11)**. Also crucial for giving important results are the techniques of promoter analysis, using different type of reporter genes **(12)**. In this chapter we will go in detail over the technique for fluorescent *in situ* hybridization of AGP transcripts, method that allows detection of specific mRNA sequences of AGPs with high sensitivity and accuracy, at individual cell level.

The study of AGPs involvement in *Arabidopsis* *thaliana* ovule development is a challenging process, where timing and optical resolution are key to success. In the complex structure of the ovule, the gametophyte tissues are inaccessible and surrounded by a double layer of sporophyte integuments, a situation/fact that adds complexity to this study. Tools have been developed to ease this ordeal, like the use of reporter fused promoters or cellular terminator together with confocal and phase contrast microscopy. But these techniques are complex and time consuming, requiring besides the preparation of constructs, transformation and selection of transformed offspring, and yet a few more weeks for plantlets to reach maturity and start producing flowers. A swifter way to determine a gene expression pattern is by *in situ*-hybridization **(4,13,14)**.

*In situ* hybridization is traditionally a technique that allows the detection and localization of genes and there transcripts **(13)**. Originally this technique relied on paraffin embedding and sectioning, a time consuming technique that required optimization for the tissue or organ under study, due to variation on cell wall thickness and permeability; moreover, the structural preservation of cells and tissues after processing for paraffin embedding is very poor due to high temperature and exhaustive dehydration with potent solvents used during the procedure. Additionally the probe was labelled with harmful radio isotopes for detection **(15,16)**, the later substitution of this radioactive materials by Digoxigenin **(17–19)** or alkaline phosphatase labeled oligonucleotides **(20)** made this technique safer and more user friendly, however the use of alkaline phosphatase or peroxidase needed optimization of the time of reaction, as the overreaction can create artefacts. Additionally it was very difficult to evaluate the state of preservation of the tissues. With further development of highly stable and sensible fluorochromes that can be conjugated to a wide set of reporter molecules, as antibodies, nucleotides, biotin or digoxigenin, cell biology localization techniques highly increased their sensitivity and field of application; this fact, together with the development of confocal laser scanning microscopy opened the way for tracking specific protein and gene expression in tissues and organs at single cell level, also in plant biology research **(21)**.

In the case of plants, one of the challenge of these in situ methodologies has been to overcome the limitations associated with specific features of most plant cells such as the presence of cell wall which hinders, at least partially, penetration of antibodies and probes to subcellular targets. Regarding *in situ* localization methods for plant ovaries, the situation is even more complex, as stated before, due to small size and inaccessibility of the gametophyte inside tissues.

With the revised method presented here, localization of transcripts in Arabidopsis ovules is made easier, more sensitive and less time consuming; by making a whole-mount fixation, probe hybridization and detection, we eliminate the need for embedding and sectioning, reducing time and difficulty of the procedure. RNA probes have the advantage that RNA-RNA hybrids are very thermostable and are resistant to digestion by RNases. This allows the possibility of post-hybridization digestion with RNase to remove non-hybridized RNA and therefore reduces the possibility of background staining.  Appropriate fixation and permeabilization are essential steps to optimally preserve cells and RNAs for efficient hybridization, messenger RNA detection and high cellular preservation and visualization. Also the use of fluorescent conjugated antibodies to localize digoxigenin-labelled hybrids enables higher sensitivity, clearer images and higher resolution of labeling at subcellular level **(22,23)**, its combination with DAPI staining **(24)** provides additional evaluation of cell structure and integrity.

**Materials**

1. **Buffers**
	1. DNA Extraction Buffer: Tris-HCl pH 7.5 200 mM, NaCl 250 mM, EDTA 25 mM, SDS 0.5% (w/v), in ddH2O. Always prepare fresh.
	2. RNA dilution buffer: 6× SSC (see buffers 1.6), 3.2% (w/v) Paraformaldehyde in RNase free H2O. Always prepare fresh.
	3. PBS (Phosphate-buffered saline): KCl 2.7mM, NaCl 137mM, Na2HPO4 10 mM, KH2PO4 1.8 mM. Adjust pH to 7.4 with HCl. Filter sterilize or autoclave.
	4. Hybridization buffer (*see* **Table 1**).
	5. 50× Denhardt: BSA 1% (w/v), Ficoll 400 1% (w/v), Polyvinylpyrrolidone 1% (w/v), in RNase free water. Filter sterilize with a 0.2 µM Filter Aliquot and store at -20oC.
	6. 20× SSC: Trisodium citrate 300 mM, Sodium chloride 3 M in RNase free H2O. Adjust pH to 7.0 with 1 M HCl. Aliquot and store at -20oC.
2. **Culture media**
	1. LB/ampicilline/IPTG/X-Gal culture plates : Bacto Tryptone 1 % (m/v), Bacto Yeast Extract 0.5 % (w/v), NaCl 1 % (w/v), Bacto Agar 1.5 % (m/v), in ddH20.Adjust pH to 7.5 with NaOH.Sterilize by autoclaving,let cool to 45oC and add,filter sterilized, ampicillin to 100 µg/mL, IPTG to 0.1 mM and X-Gal (in DMF) 30 µg/mL. Plate immediately and store at 4oC wrapped in aluminium foil.
	2. SOC: Bacto Tryptone 2 % (w/v), Bacto Yeast Extract 0.5 % (w/v), NaCl 10 mM, KCl 2.5 mM, MgCl2 10 mM, MgSO4 10 mM, Glucose 10 mM. Sterilize by autoclaving**.** SOC culture medium is used for the recovery of transformed bacterial cells.
3. **Solutions**
	1. TE: Tris-HCl pH 7.5 10 mM, EDTA 1 mM, in ddH2O. This lightly saline solution helps to devolve DNA.
		1. Tris/EDTA: Tris-HCl pH 7.5 100 mM, EDTA 10 mM, in RNase free ddH2O. Aliquot and store at -20oC. This variation of the TE solution is used in the hybridization buffer.
	2. Miniprep solution 1: TRIS pH 8.0 25 mM, EDTA pH8.0 10 mM, Glucose 50 mM, in ddH2O. Always prepare fresh.

After removing the growing medium in the first step of the plasmid purification protocol, the cells must be suspended in a uniform manner for the alkaline lysis to take place. This solution will allow just that.

* 1. Miniprep Solution 2: SDS 1% (w/v), NaOH 0.2 N, in ddH2O. This solution will burst bacterial cell by attacking the cell membrane with a detergent and high pH. In this way the plasmids will be released into the solution. Always prepare fresh.
	2. Miniprep Solution 3: KOAc 3 M, Glacial acetic acid 2 M, in ddH2O. Store at 4oC, check for contaminants before use. After bursting the cells, proteins and all other cellular molecules are released to the alkaline solution. This solution will neutralise the solution and precipitate SDS, proteins and other impurities.
	3. Miniprep Solution 4: RNase (Thermo, EN0531) to 20 µg/mL in TE (see solutions 3.1). If possible prepare fresh. RNA co-purification is common with this miniprep protocol and can be a problem to the restriction of the purified plasmids. A simple and easy way to get rid of this contamination is a simple treatment with RNase.
	4. BSA 5%: Bovine serum albumin 5% (w/v) in PBS (see buffers 1.3). Aliquot and store at -20oC. Dilute at a ratio of 1:5 in PBS to prepare 1% BSA. Bovine serum albumin is used as a blocking agent in the immunodetection reactions.
	5. Fixative Solution: Paraformaldehyde 4% (w/v), Tween20 0.001% (v/v), in PBS (see buffers 1.3). Always prepare fresh.To prepare 1% paraformaldehyde in PBS make a dilution 1:4 of this solution in PBS. This fixative solution is used to prevent degradation of cellular content, prior to the dehydration step. It also acts on mRNA unfolding, easing RNA probe ligation.
	6. Methanol series in PBS: Make methanol dilutions of 70% (v/v), 50% (v/v), and 30% (v/v), in PBS (see buffers 1.3). Store at -20oC. During the dehydration/rehydration process a methanol series is used to dehydrate and rehydrate the dissected pistils. This step serves as a first permeabilization to help further accessibility of probes to intracellular targets. The final dehydration prior hybridization eliminates water excess in tissues that would hinder hybridization reaction, it also helps eliminating any remaining active enzyme used during permeabilization.
	7. Macerozyme R-10: Macerozyme R-10\* (cat.no. 28302, SERVA Germany) 2% (w/v), in RNase free H2O. Filter sterilize with a 0.45 µM Filter,Aliquot and store at -20oC. Macerozyme R-10 has various enzymatic activities to partially degrade cell wall components The cell walls are one principal barriers for probe penetration, by treating the tissue with a pectinase and cellulase cocktail the cell walls will become more permeable without loosening there integrity.

\*Activity: Pectinase 0.5 Umg-1, Hemicellulase 0.25 Umg-1, Cellulase 0.1 Umg-1

* 1. DAPI staining solution: To make a 5 mg/mL DAPI stock solution (14.3 mM for the dihydrochloride DAPI or 10.9 mM for the dilactate DAPI), dissolve 10 mg in 2 mL of deionized water (dH2O) or dimethylformamide (DMF). You may have to sonicate the dihydrochloride DAPI as it is harder to dissolve in water than the dilactate DAPI. Dilute the DAPI stock solution to 300 nM in PBS, and use about 20 µL directly over the pistil containing wells. DAPI (4', 6-diamidino-2-phenylindole) is a very useful stain for the nucleus. In this version of fluorescent *in situ* localization it serves a dual purpose, first it will help to locate the cell nucleus. The labelling of DAPI and the probe should not overlap since in the conditions used the RNA probe should have no affinity to the nucleic DNA. Observation of the integrity of the nucleus, as revealed by DAPI staining, permits proper evaluation of the hybridization signal in the cytoplasm and good cell structural preservation.
	2. Mowiol mounting medium: Weigh 6 g of analytical-grade glycerol in a 50 mL disposable plastic conical centrifuge tube. Add 2.4 g of Mowiol 4-88 (sigma, 81381) and stir thoroughly to mix the Mowiol with the glycerol. Add 6 mL of distilled water and let the solution rest at room temperature for 2 h. Mix in 12 mL of 0.2 M Tris buffer (2.42 g Tris/l00 ml water, adjust pH to 8.5 with HCl) and incubate the solution in a water bath at 50°C for 10 min with occasional stirring to dissolve the Mowiol. Clarify the mixture by centrifugation at 5000 *g* for 15 min. Aliquot and store at -20oC.

The solution is stable, and the pH is retained for at least 12 months at -20°C, and for about a month at room temperature. Mowiol is a water soluble hydrocolloid muco-adhesive based on poly (vinyl alcohol), that carries both the advantage of antifade properties, and creating a solid seal of the final preparation. Stained slides mounted with Mowiol retain fluorescence when stored at 4oC in the dark.

**Equipment:**

1. 0.2 and 1.5 µL centrifuge tubes.
2. 15 and 50 mL conical tubes.
3. Centrifuge.
4. Fine tipped Forceps.
5. G26 (or higher)-hypodermic Needles.
6. Humidity chamber (see **Figure 3**).
7. Hybridization oven.
8. Kitazato.
9. Micropipettes.
10. Nanodrop®.
11. Parafilm®.
12. Teflon coated reaction slides.
13. Termocycler.
14. Vaccum pump.
15. Vacuum chamber.
16. Vortex.

**Enzymes:**

1. Taq based PCR system.
2. PstI and NcoI.

**Commercial Kits:**

1. NBT-BCIP® solution (sigma, 72091).
2. pGEM®-T Easy cloning kit(Promega, A1360).
3. JM109 Competent Cells (Promega, L2005).
4. T7/SP6 DIG-RNA labelling kit (Roche, Cat. No. 11 175 025 910).

**Method**

This extended protocol for Fluorescent *in situ* hybridization is a broad guide through all the steps from the extraction of the template DNA, to the synthesis of the probe. Finally the hybridization itself will be divided in preparation of the hybridization slides, probe hybridization and probe detection.

1. **Probe preparation**

Probe design and synthesis is one of the most important steps for a successful *in situ* hybridization. Careful planning must be made to choose the correct probe sequence and length. Has it will affect the probe specificity and penetration ability.

AGPs are a diverse group of proteins **(1)** where splicing is generally absent, making the selection of an mRNA region to use has target for a RNA probe fairly simple. The design process must include however a careful verification of the selected sequence specificity, the most preserved sequences being the N-terminal and GPI anchor signals. AG-Peptides due to their very short sequence are most challenging for this task, but short specific probe sequences can be obtained. The optimal length for a RNA probe is between 200 and 1500 bases.

Another aspect of the use of RNA probes is there sensitivity to degradation, an extra effort must be made to keep all materials and solutions RNase free. A minute amount of RNase may destroy a complete batch of precious RNA probe.

* 1. **DNA-Extraction**

Genomic DNA will serve has a template to create the probe sequence that will be inserted in a plasmid, that in turn will be used to synthesise the RNA probe.

The first step will be to extract DNA. In the following method we propose to use a rapid and simple method to isolate genomic DNA suitable for PCR amplification, based on the method of Edwards *et al*. (1991) **(25)**.

All steps are performed at room temperature and centrifugations are at 13.000 *g*.

* + - 1. Prepare the DNA Extraction Buffer (see buffer 1.1).
			2. Use the lid of a 1.5 mL centrifuge tube to punch out a disc from a young leaf into the tube (*see* **Note 1.1**), and grind it thoroughly in a small mortar and pestle. Immediately add 400 μL of extraction buffer, and vortex for 5 seconds (*see* **Note 1.2**).
			3. Spin at max RPM for 2 min to pellet the tissues debris, and transfer 400 μL of the supernatant into a fresh 1.5 mL centrifuge tube (avoid taking debris from the pellet). Add 350 μL isopropanol, mix by gently inverting the tubes a few times and leave at room temperature for 2 minutes. Then spin 5 minutes at full speed to pellet the DNA.
			4. Remove all the supernatant and let air dry for a few minutes. (Do not let the pellet get too dry, otherwise it will be very difficult to suspend the genomic DNA).
			5. Dissolve the pellet in 50 μL TE (solution 3.1), or PCR grade H20 by gentle shaking. DO NOT VORTEX. Check the integrity of the gDNA by agarose gel electrophoresis (*see* **Note 1.3**).
	1. **pGEM Cloning**

After extracting the gDNA can be used as a template to amplify the target gene sequences by PCR using an enzyme with 3´ adenylation activity (see notes pGEM cloning 2.1).

Perform a PCR cleanup and quantify your PCR product.

* + 1. **Cloning PCR Products with pGEM®-T Easy Vectors** (Promega, A1360).
			- 1. Set up ligation reactions, with 5 µL of 2× Rapid Ligation Buffer, 1 µL of pGEM®-T Easy Vector (50 ng), 37.5 µg of purified PCR product, 1 µL of T4 DNA Ligase and PCR grade H2O to a final volume of10 µL. Vortex the 2× Rapid Ligation Buffer vigorously before use (*see* **Note 2.2**).
				2. Mix the reactions by pipetting (*see* **Note 2.3**). Incubate overnight at 4°C to maximize ligations.
		2. **Transformation of JM109 High Efficiency Competent Cells.**
1. Prepare LB/ampicillin/IPTG/X-Gal plates (see materials 2.1).
2. Centrifuge the ligation reactions briefly. Add 2μl of each ligation reaction to a sterile 1.5 mL tube on ice. Prepare a control tube with 0.1 ng of uncut plasmid.
3. Place the JM109 Competent Cells (Promega, L2005) in an ice bath until just thawed (~5 minutes). Mix cells by gently flicking the tube. Carefully transfer 50 μL of cells to the ligation reaction tubes. Use 100 μL of cells for the uncut DNA control tube. Gently flick the tubes and incubate on ice for 20 minutes. Then Heat-shock the cells for 45–50 seconds in water bath at exactly 42°C. DO NOT SHAKE. Immediately return the tubes to the ice. After 2 min add 950 μL room temperature SOC medium (materials 2.2) to the ligation reaction transformations and 900 μL to the uncut DNA control tube. Incubate for 1.5 hours at 37°C with shaking (~150 r.p.m.).
4. Plate 100 μL of each transformation culture onto duplicate LB/ampicillin/IPTG/ X-Gal plates. For the uncut DNA control, a 1:10 dilution with SOC is recommended. Incubate all plates overnight at 37°C. Select white colonies.
5. Use M13 primers in conjugation with your reverse primer to confirm to orientation of your insert. This will determine which reaction T7 RNA Polymerase or SP6 RNA Polymerase will produce your antisense (or detection) probe **(26, 27)** (*see* **Note 2.4**).
6. After this initial screening select one colony and proceed with multiplication of the probe containing plasmid.
	* 1. **Miniprep protocol (28).**

The plasmid containing the probe sequence will be used as a template for SP6 and T7 RNApolymerase **(26, 27)**. We will now proceed with the multiplication and purification of the probe containing plasmid.

1. Inoculate 5 mL of LB/amp (75 µg/mL) (*see* **Note 3.1**) overnight at 37°C with vigorous agitation.
2. Fill a 1.5 mL Eppendorf tube and centrifuge for 1 min at 10 000 *g* (*see* **Note 3.2**). Discard the supernatant and repeat until you have spun down 5 mL.
3. Then suspend the pellet in 100 µL of Miniprep solution 1 (materials Solution 3.2) and mix in 20 µL of 10 mg/mL of lysozyme. Let stand at room temperature for 2 min (for 1 mL: 0.01g Lysozyme in 1 mL 0.250 mM tris pH 8.0).
4. Add 200 µL of Miniprep solution 2 (materials Solution 3.3) mix well and place on ice for 5 min, then add 150 µL of Miniprep solution 3 (material solution 3.4) vortex gently and place back on ice for 5 min.
5. Centrifuge at 4°C, for 5 min at 12000 *g*. Transfer the supernatant to a new tube (*see* **Note** **3.3**).
6. Add 400 µL of phenol: chloroform. Vortex and centrifuge at 4°C, for 2 min at 12000 *g*.
7. Transfer the aqueous upper phase to a new tube (*see* **Note 3.4**). Add 1 mL of 100% ethanol at room temperature.
8. Leave to stand for 2 min at room temperature.
9. Centrifuge at 12000 *g* for 5 min, pour off the ethanol and let the pellet dry.
10. Dissolve the pellet in 50 µL of TE/RNase (Materials Solution 3.4), quantify your purified Plasmid by Nanodrop and send one sample for sequencing.
	1. **RNA Probe Synthesis and labelling**

In the following steps the purified plasmid will be prepared for the synthesis and labeling of the probe with the T7/SP6 DIG-RNA labelling kit (Roche, Cat. No. 11 175 025 910) (*see* **Note 4.1**).

* + 1. **Linearize the plasmid**

For the proper synthesis of the RNA probe by T7 or SP6 RNA polymerase the plasmid most first be linearized, or it will produce uneven sized and possibly unspecific sequences.

1. Carefully label two 1.5 mL tubes, with Sense (Tube 1) and Anti-Sense (Tube 2) and add the volume equivalent to 2.5 µg purified plasmid to each.
2. Digest tube 1 with PstI and tube 2 with NcoI (*see* **Note 4.2**).
3. Spin down and incubate both tubes at 37°C for 15 min then stop the reaction by heating the tubes at 95°C for 5 min.
4. Clean up the linearized plasmids (*see* **Note 4.3**). Quantify both the purified linearized plasmids by Nanodrop.
	* 1. **Digoxigenin-labelled RNA Probe Synthesis.**

In the next step, the Roche Dig Labeling Kit will be used to synthesize and label the RNA probe in one single step. In the described procedure both control and detection probe will be synthesized. It is of the outmost importance to provide high a quality template for the reaction to occur properly.

1. Add the equivalent of 1 μg of purified linearized plasmid to a 200µL microtube. Place the microtube on ice and add the following reagents according to the mixing order (*see* **Table 2**).
2. Mix gently and incubate at 37°C for 2 h. To stop the reaction add 2 µL of 0.2 M EDTA (pH 8.0) (*see* **Note 4.4**). Probes may be stored at -20°C for up to one year (*see* **Note 4.5**).
	* 1. **Probe labelling quantification** (please *see* **Note 4.6**).

Now determine the labelling efficiency, as it may be inconsistent. This step will ensure the control of the reaction and allows normalizing the probe for the future hybridization essays. The simplest way to determine the labelling efficiency is to compare the probe that we synthesized to standard solution, a standard solution is provided as a control in the previously used labelling kit. An efficiency test is proposed in the kit instructions, but we find it to be overcomplicated and more expensive. The following approach consists of simply comparing the diluted standard to a dilution of our probe to find out with some degree of precision the concentration of our probe.

1. Make 3–10 fold dilutions of the probe in RNA dilution buffer, 4% Paraformaldehyde 6× SSC in H2O (materials buffers 1.2), starting with 1 µL of probe. And make 4 10-fold dilutions with the labelled RNA control standard, supplied with the kit (vial 5). Pipette 1 µL of the original solutions followed by 1 µL of all dilutions on a nylon membrane (*see* **Note 4.7**).
2. Fix the nucleic acid to the membrane by cross-linking with UV-light or baking for 30 min at 120°C.
3. Place the membrane strip in a 50 mL conical tube. Block for 5 min with 10 mL 5% BSA (Materials 3.6), in an orbital shaker with mild agitation. Discard the blocking solution and incubate for 90 min at room temperature with a solution of 1/1000 of anti-Digoxigenin antibody conjugated to alkaline phosphatase (Sigma, A1054), in 1% (w/v) BSA.
4. Wash 3 times for 10 min with PBS (Materials 1.3), in an orbital shaker 80 r.p.m. Drain thoroughly and transfer to a Petri dish. Add 5 mL of NBT-BCIP® solution (sigma, 72091), cover with aluminium foil and place in an orbital shaker at 80 r.p.m. Let react until the last dilution of the control starts to appear. Stop the reaction by washing out the NBT-BCIP® solution with ddH2O.
5. Compare the labeling of the probe with the scale made using the RNA labelled control (*see* **Note 4.8**).
6. **In Situ Hybridization.**

The pistils will be prepared to undergo hybridization with the probe previously set, in two steps: First the pistils will be fixated and then the cells permeabilized to permit the probe penetration, and finally the hybridization and detection reaction will be performed.

* 1. **Fixation.**

Dissecting the ovaries prior to the fixation will help with the fixation, probe penetration and visualization of the final results. The formaldehyde fixation will allow to preserve the tissues, reduce the activity of proteins and help to unfold the target mRNA.

* + 1. Select flowers and extract pistils. Place the pistil on a piece of double-face duct tape on a microscopy slide. Under a stereoscope remove the valves of the pistil using fine gauge needles (G26 or higher) and fine point tweezers, by cutting along the valve suture ridges. Be careful not to sever the ovules from the septum and maintain the transmitting tissue intact (**Figure 2**).
		2. Immediately immerse the samples in ice cold fixative solution, 4% Paraformaldehyde 0.001% Tween 20 in PBS (solution 3.7). Apply sustained vacuum of -70 KPa for 15 min to help the fixative to penetrate into tissues. Maintain in the fixative solution for 18 h or overnight at 4°C.
		3. Wash the fixed pistils by immersing them for 10 min in cold PBS, with mild agitation. Repeat 3 times. If necessary, preserve the dissected fixed pistils at 4°C in 0.1% Paraformaldehyde in PBS until use, to avoid reversion of paraformaldehyde fixation.
	1. **Slide preparation**

To address the preparation of the slides for the hybridization, some protocols call for the use of extremely expensive slides or hybridization chambers. We propose the use of simple and affordable multi-well Teflon printed diagnostic slides, which restrict reagents to tissue samples, with some good cleaning and polylysine coating.

* + 1. **Slides wash (**Please *see* **Note 5.1**).
1. Place the slides in a staining rack and cover with a cleaning solution, 70% Ethanol 0.1% Triton X100, with mild agitation for 20 min.
2. Wash the slides by dipping the staining rack in ddH2O with mild agitation for 10 min. repeat this process 4 times. Carefully dry the rack before dipping briefly in 100% ethanol and let the slides dry out in a dust free environment.
	* 1. **Polylysine coating**
3. Place the clean slides in a clean 12×12cm square Petri dish.
4. Make a solution of 0.001% poly-l-Lysine (Sigma P4707) in ddH2O.
5. Cover each well of the slides with 50 µL of the poly-l-lysine solution.
6. Place the closed Petri dishes in a 40°C oven overnight to dry the solution.
7. Keep the coated slides at room temperature in a dust free environment (*see* **Note 5.2**).
	* 1. **Permeabilization**

Permeabilize the pistils to ease penetration of the probe; by first fixing the pistils to the poly-l-lysine coated slides and then dehydrate, rehydrate, digest briefly with a cell wall-degrading enzyme cocktail and finally dehydrate for storage or immediate processing.

* + 1. **Pistil adhesion to the coated slides.**
1. Transfer up to two fixed dissected pistils to each polylysine coated well (*see* **Note 5.3**).
2. Let air dry for 20 min at room temperature, and immediately proceed to the dehydration step.
	1. **Dehydration-rehydration and mild cell wall enzymatic digestion.**
3. Dehydrate by incubating for 5 min with an increasing concentration series of methanol in PBS, 30%, 50%, 70% 100%. Then rehydrate by incubating for 5 min with a decreasing concentration series of methanol in PBS, 70%, 50%, 30%. And finally for 5 min in PBS (*see* **Note 6.1**).
4. Transfer to a humidity box and immediately cover the pistils with 30 µL 2% macerozyme (cat.no. 28302, SERVA Germany) in PBS. Seal the humidity box with Parafilm and incubate for 60 min at 45°C.
5. Remove the macerozyme solution and wash with PBS three times, 5 min each. Then, incubate each well with 30 µL proteinase K (20 µg/mL) for 20 min at 37°C. Then wash 3 times with PBS for 5 min.
6. Wash with ddH2O for 5 min (*see* **Note 6.2**).
7. Dehydrate again in an increasing series of methanol in PBS. Make an additional incubation with 100% methanol, and leave to air dry.
8. Slides can be immediately used or stored in a dry sealed box at -20°C for up to 2 years (*see* **Note 6.2**).
	1. **Hybridization:**

To proceed to the final stage of the protocol, the hybridization and detection, first the previously dehydrated tissues will be incubate with the labeled probe, and finally the hybridized probe will be detected by immunolocalization of the Digoxigenin.

1. Prepare a hybridization box, by placing a few conical tube caps with formamide on the bottom of a humidity box (*see* **Note 7.1**).
2. After selecting the correct amount of probe to use (*see* **Note 7.2**).
3. Calculate the volume of hybridization solution that you will require, about 25 µL per well and prepare your hybridization buffer (300 mM NaCl, 10 mM Tris/EDTA, 1× Denharts solution, 50% Formamide, 10% Dextran sulphate, 200 µg/mL tRNA) accordingly (see Materials Buffer 1.4).
4. Place all aliquots required on ice and allow thawing.
5. Place the Dextran sulfate aliquot in a 75°C water bath for 15 min, as it is highly viscose, it will also help to pipette with a clipped tip. Place the unfrozen tRNA stock aliquot in boiling water for 15 s before adding to the hybridization buffer.
6. Mix all volumes from the stocks according to the mixing order in **Table 2** (Materials Buffer 1.4).
7. Mix the hybridization buffer with the appropriate probe volume.
8. Add 25 µL of the hybridization solution to each well; take care to not put different probes in adjacent wells.
9. Cover the wells with a small piece of Parafilm®, place the slide in the modified humidity chamber and seal with Parafilm®. Incubate at 50°C overnight. The Parafilm® pieces will ensure appropriate contact between tissue samples and probe, making a small hybridization chamber over tissues and avoiding the reagents evaporation during the incubation.
	* 1. **Post hybridization washing**

Extract the excess probe which has not hybridized with samples (*see* **Figure 4d**).

1. Remove the conical tube caps containing formamide from the humidity chamber and carefully remove the Parafilm® covering the wells with fine tips forceps (*see* **Note 7.3**).
2. Wash 4 times for 2 min with 4× SSC at room temperature.
3. Wash 4 times for 2 min with 2× SSC at room temperature.
4. Wash 2 times for 15 min with 0.1× SSC at 50°C. These stringency conditions are standard but they can be adjusted depending on the probe sequence to detect; lower stringency with lower temperature and higher SSC concentration allows hybridization between sequences that are less similar.
5. Finally wash with PBS for 5 min.
	1. **Immunolocalization of the hybridized probe**

The final step is the immunolocalization.

1. Place your slides in a humidity box.
2. Incubate samples in wells for 5 min with 5% (w/v) BSA at room temperature to block non-specific reactive groups of tissues.
3. Incubate for 90 min at room temperature with the primary antibody, mouse anti-Digoxigenin (Sigma, 11333062910), in a solution of 1/5000 in 1% (w/v) BSA.
4. Wash 3 times for 5 min with 1% (w/v) BSA.
5. Incubate for 45 min at room temperature, with the secondary antibody, anti-mouse conjugated with FITC (Sigma, F0257), in a solution of 1/25 in PBS (*see* **Note 7.4**).
6. Wash 6 times for 3 min with PBS.
7. Stain for 5 min with about 20 µL of a 0.3 mM DAPI solution in PBS (*see* **Note 7.5**).
8. Then wash 6 times for 3 min each time with PBS and mount the slides with Mowiol **(30,31)**.
9. Observe under fluorescence or confocal microscope (*see* **Note 7.6**).
10. **Controls:**

To all assays the following tests must be considered to verify and validate the specificity of the hybridization probe and also of the antibodies used.

* 1. *In situ* Hybridization using the sense probe. This control evaluates the unspecific binding of the probe. The results should be negative, with no detectable hybridization.
	2. Immunolocalization replacing the primary anti-dig antibody by buffer. This control evaluates the unspecific binding of the secondary antibody. No signal should be detected.
	3. Immunolocalization replacing the secondary fluorescent-labelled antibody by buffer. This control evaluates if there is any source of unspecific autofluorescence, which may contaminate the results. No autofluorescence results should be detected in the range of the fluorescent-labelled antibody.

**Notes**

1. **DNA Extraction Notes:**

This crude DNA extraction method is a cheap and fast way to purify DNA.

It is not as efficient as other commercially available solutions. However, this protocol employs readily available reagents, and provides DNA suitable for PCR.

* 1. It is important to avoid using too much plant material.

If too much material is used, it will result in an increase of gDNA degradation by interfering with the buffer denaturing capacity. It will also make it difficult to separate the DNA containing supernatant from the plant material debris.

In contrast this same method can be used to extract PCR suitable DNA from a smaller sample, like a single cotyledon, in which case all volumes should be reduced by 50% and DNA precipitation (Step 1.1.1.7.) should be performed at -20oC with final centrifugation at 4oC.

* 1. At this stage, the samples can be kept at room temperature for up to one hour with no major damage to the extraction results, while you finish other samples that you may be preparing.
	2. You may find that some RNA was co-purified, it should not be much of a problem for normal PCR. However, if necessary treat by adding 1 µL of RNase solution and incubate at 37oC for 20 min.
1. **pGEM Cloning Notes:**

The pGEM cloning system lets you easily clone regular PCR products without any additional complicated steps.

* 1. Most common PCR enzymes will leave a 3´ Adenine overhang, which is necessary for pGEM cloning. Error rate for a small amplicon are low. However, if you prefer to use a proofreading enzyme verify that they have 3´ adenylation activity or perform this as an additional step.
	2. Assuming a probe size of 750 bp and a 3:1 ratio molar of insert to linearized plasmid.

The pGEM cloning system should perform perfectly with a wide range of insert to linearized plasmid ratios. However if you experience difficulties with the ligation, try to increase the amount of insert.

* 1. Use 0.5 ml tubes with low DNA binding capacity. In alternative regular PCR grade 200 µL may also be used.
	2. The pGEM vector features a polylinker flanked by M13 sites. You may use this feature to select plasmids that were recombined with the insert in a desirable manner. To do so select plasmids by PCR using primers M13 forward and the revers primer used to generate your insert.

Always select T7 to SP6 RNA polymerase ligation site insert oriented plasmids, this avoids confusion during synthesis of the RNA probe. Because of our selection T7 RNA polymerase always synthesizes Sense (or control) probe and SP6 RNA Polymerase always produce the anti-sense (detection) Probe.

1. **Miniprep protocol**
	1. Use 50 mL tubes as they provide better oxygenation, which will promote growth. This process can be scaled up. Most importantly avoid letting your culture growth for too long.
	2. Two tubes may be used simultaneously to pellet down the bacterial cells, reducing the need for a counterweight in centrifugations and speeding the process, the two pellets can be joined at the resuspension step.
	3. It is best to leave some supernatant than to risk transferring of any the white precipitate. If it is too difficult to do so centrifuge again.
	4. It is of primordial importance to not disturb the phases and to only transfer the aqueous upper phase, it is best to lose some than to contaminate your plasmid extract. If however you unfortunately transferred some of the other phases simply repeat step 6 and 7.
2. **RNA Probe Synthesis and labelling**
	1. Assume that you selected a plasmid that features an insertion with the orientation T7 to SP6. In this way SP6 polymerase will produce an anti-sense probe and T7 will produce the sense probe. Therefore henceforth, to avoid confusion, we will use the term sense for the probe synthesized with T7 RNA polymerase and antisense to the probe synthesized with SP6 RNA Polymerase. If you have chosen to do otherwise, it is of little importance just adjust accordingly.
	2. The use of fast digest enzymes is of course optional. But it saves a lot of time to this already long procedure. To further speed and simplify the process a thermocycler may use to act as an incubator, simply perform the restriction on 200µL tubes.
	3. Use a simple PCR purification kit (Like, Thermo K0701 “GeneJET PCR Purification Kit”), to remove enzymes and salts. Make sure to make the final elution with 20 µL of RNase free H2O**.** Run 1 µL in an agarose gel to check for nicked or supercoil plasmid, has both this forms can cause disruptions on the probe synthesis process.
	4. The RNA synthesis quality should be verified by running 1 µL of the reaction next to the labeled control (Vial 5) in an agarose gel electrophoresis with ethidium bromide staining. A good RNA synthesis results in a simple band. If a smear is observed, the reaction was probably contaminated with RNase and/or the RNase Inhibitor was not added to the reaction mix. If multiple bands are visible, then most probably the plasmid linearization and purification step failed, please repeat step 1.2.1 and check for nicked or coiled plasmid.
	5. RNA probes are easily degraded. To preserve the probe avoid thawing and freezing cycles, for a longer preservation consider making aliquots.
	6. It is very important to determine the labelling efficiency, has it may be inconsistent. Several times we observed different probe yields in technical replicates, furthermore it is useful to determine the probe concentration. So that the correct amount is used in the hybridization solution.
	7. Spot immediately to avoid degradation of the Probe, space the spots so they do not overlap. It is best to have the scale and the test probe side by side for easy comparison. **Figure 1** is an example of what the spot distribution should look like.
	8. The labeled control standard dilution will allow you to determine the relative concentration your probe (see **Figure 1**).
3. **Slide preparation**
	1. This step is especially important when reusing slides or if they seem to be dirty.
	High quality coating on the slides is of the outmost importance, as it is what will insure that solutions won’t spread and mix between wells. A good quality of the slides is one of the key issues for this technique. Slides can be cleaned and reused, just let them soak in a solution of ethanol 30% with some detergent for a couple of hours and gently brush them with a soft tooth brush, just prior to using this wash method. There are a lot of solution changes in this protocol. However, time consuming to prepare the polylysine-coated slides will help significantly to keep the pistils in place **(29)**.
	2. Polylysine coated slides can be prepared in advance, and if stored in a sealed rack at -20oC they can be preserved for up to one year without losing their properties. Just let them reach room temperature before opening the accumulation of condensation. Always inspect the slides under the microscope, to check the coating and cleaning, before use.
	3. Some humidity must be present at the surface of the dissected pistils to promote adhesion to the polylysine coated slides, but avoid transferring to much fixative solution it will take a long time to dry. A good way to get read of the excess moisture is to lightly touch with a lint free tissue, before placing the pistils on the slide.
	4. **Dehydration-rehydration and mild enzymatic digestion of cell wall**

Going up the methanol dehydration series it is unfortunately very common for some material to loosen from the polylysine coating, please proceed with special care. If necessary don’t be afraid to reposition the pistils in the reaction wells. In the higher concentrations of methanol take care not to let the solution evaporate. If necessary add more solution during incubation.

Plant cell walls are the principal barrier for the penetration of the Probe. Partial digestion of them will not only facilitate the penetration of the probe but also will help with the removal off unlinked probe thus reducing non-specific labelling.

* 1. Make sure to store slides with pre-treated/dried samples in small batches in clean air tight containers and also let them reach room temperature before opening the container, to avoid water condensation on samples. A cheap and easy way to prepare an air-tight container, is to put up to 4 slides in a 12×12cm square Petri dish and seal it with a double layer of Parafilm.

**Hybridization:**

A humidity box is a versatile tool that can be purchased, or built, to maintain a saturated atmosphere around the slides avoiding the evaporation of solutions. We always build our humidity boxes by placing a few damp paper towels to bottom of a tip box. See **Figure 3** for how to build a simple reusable humidity chamber that also can be used for routine immunofluorescence.

Use 20 to 100 ng of probe for each 1 mL of hybridization solution. Please note that some consideration and assays should be performed to ascertain the perfect amount of probe to use. Consider the expression level of the gene and the concentration of the probe. Assuming that your labeled probe has a concentration of near to 20 µg/µL, 1/20 or 1/50 ratio of probe to hybridization buffer is usually a good starting point to detect an averagely expressed gene. However, most probably you will need to make adjustments on the quantity of probe to use. If you do not get any labelling with this standard dilution, double the amount of probe until you do. Do the inverse if you get to much background with the anti-sense Probe.

It is not unusual that some material may stick to the Parafilm®, make sure to check the Parafilm® before discarding. If you do find that some material did stick to the Parafilm® use your forceps to place it back in the well.

This proportion is for genes with an average expression. When the signal is very weak the quantity of probe used in the hybridization buffer must be adjusted. Genes with low expression are always more difficult to detect.

DAPI should only label a bright blue intact nucleus, and therefore will act as one of the controls for integrity of the tissue has well as control for the specificity of the probe, as they should not overlap. Always check your DAPI solution before staining your slides. It is not uncommon for the DAPI solution to degrade before the expected expiration date, and it is very difficult to wash out (*see* **Figure 5**).

Use DAPI and FITC filter setting to detect the signals. Plant cell walls have autofluorescence at almost every wave length, for the best results use a confocal microscope if possible. Set the excitation Laser to 355 nm for DAPI and 488 nm for FITC. Emission filters between 450 and 460 nm for DAPI, and for FITC 515–525 nm. If you use an epifluorescence microscope set the filters to Excitation⁄Emission (nm) 358/461 for DAPI and 485/530 for FITC.

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**Tables:**

**Table 1: Hybridization buffer**

|  |  |
| --- | --- |
|  | **Volume of hybridization buffer** |
| **Stocks**  | **Final concentration** | **50 µL** | **100 µL** | **150 µL** | **200 µL** | **250 µL** | **Mixing order** |
| RNase free H2O | n.a. | 5.5 µL | 11 µL | 16.5 µL | 22 µL | 27.5 µL | **1** |
| NaCl 5M | 300 mM | 3 µL | 6 µL | 9 µL | 12 µL | 15 µL | **2** |
| Tris/EDTAsolutions 3.1.1 | 10mM/1mM | 5 µL | 10 µL | 15 µL | 20 µL | 25 µL | **3** |
| 50× Denhardt | 1x | 1 µL | 2 µL | 3 µL | 4 µL | 5 µL | **4** |
| Formamide | 50% | 25 µL | 50 µL | 75 µL | 100 µL | 125 µL | **5** |
| 50% Dextran sulfate | 10% | 10 µL | 20 µL | 30 µL | 40 µL | 50 µL | **6** |
| tRNA (20 mg/mL) | 200 µg/mL | 0.5 µL | 1 µL | 1.5 µL | 2 µL | 2.5 µL | **7** |

**Table 2. Probe synthesis reaction mixture order**

|  |  |  |  |
| --- | --- | --- | --- |
| Vial  | Reagent | Volume | Mixing order |
| n.a. # | RNase free PCR grade H2O | To 20 µL | 1 |
| 8 | 10x Transcription buffer | 2µL | 2 |
| 10 | Protector RNAse inhibitor | 1 µL | 3 |
| 7 | 10x NTP labeling Mixture | 2 µL | 4 |
| FOR **SENSE** PROBE\* |
| 11 | RNA polymerase T7 | 2 µL | 5 |
| FOR **ANTI-SENSE** PROBE\* |
| 12 | RNA polymerase SP6 | 2 µL | 5 |

#(Sigma, W1754); \*According to pGEM insert orientation

**Figure Legends:**

**Figure 1. Dot blot distribution example.**

This is an example of a dot blot comparing the 5 dilutions of the control probe with 3 test probes. The five 10 fold dilutions of the control probe correspond to, 1µg/µL – 0.1 µg/µL – 0.01 µg/µL – 0.001 µg/µL – 0.0001 µg/µL, from left to right. This scale can be used to determine the approximate concentration of your labeled probe. If the fourth spot of your probe matches the color of the 4th spot of the control probe then your probe as a concentration of approximately 1 µg/µL, like for probe A. but if by the time the higher dilution of the control starts to appear, the last spot of your probe is still not visible use the last spot for your probe and multiply by a factor of -10 fold for each missing spot. For Probe B the first spot is similar to the fourth spot of the control then it has the relative concentration of 0.001 µg/µL. However, if like in the case of probe C, the last spot is similar to a lower dilution of the control a multiplication of 10 fold must be applied for each spot to the left, in this case 1 mg/µL.

**Figure 2. Pistil dissection.**

To expose the ovules of the Arabidopsis pistil, place the pistil on a piece of double faced duct tape so that the valve sutures on one of the sides face the top (a). With a G26 gage hypodermic needle make a shallow incision following the suture line exposing the ovules. Proceed in the same manner on the other side (b). Finally using a pair of fine tip tweezers pull the semidetached valves and finish the cutting the opposite side of valve suture detaching it from the pistil (c).

**Figure 3. Setting a hybridization box.**

A humidity box is a fundamental tool in this procedure, it will avoid the evaporation of solutions during the incubation periods. To make one use a tip box, tin foil and some double face duct tape (a). Cover the top of the box with a layer of double face duct tape (b) then cover it with aluminium foil (c). Remove the excess foil (d) and remove the tip holding rack. In the bottom of the box place some damp paper towels (e). To make it suitable for the hybridization incubation place some 50 mL conical tubes caps field of formamide on the bottom of the box (f) This will saturate the air in the sealed chamber and reduce the evaporation of the hybridization solution, as after the hot incubation the leads can be easily removed to minimize the exposure to formamide vapors. Put back the tip holding rack, it will serve as a nice support for the slides.

**Figure 4. Hybridization tips.**

Place the slides in a humidity chamber to avoid evaporation of solutions, additionally different probes can be used on the same slide if some space is left between wells, for example anti-sense probe was applied to wells® 1 and 5 and control probe can be applied to wells 4 and 8 (a). Small squares of Parafilm® can be used to cover the wells to avoid excessive evaporation of the hybridization solution (b). A pipette tip attached to a vacuum pump, under low negative pressure (-20 KPa), can be used to swiftly remove solutions from the wells (d), speeding the wash steps.

**Figure 5. Examples of Confocal Whole-mount FISH (Fluorescent *in situ* hybridization) in Arabidopsis Pistils results.**

Examples of ovules after hybridization with the same AtAGP12 probe and counterstained with DAPI. On the left a perfectly preserved ovule was stained with a fresh solution of DAPI (a), on the right the ovule was not treated with paraformaldehyde and was stained with an old DAPI solution (b).

The probe should only hybridize with the RNA present in the cell cytoplasm, giving a clear bright green image of the cells were the transcript is present (orange arrows), and should not be seen on the nucleus (pink arrow). DAPI stain is more than just a counterstain, it also serves to verify the conservation of the tissues. In perfectly preserved tissues the DAPI stain should be restricted to the cell’s nucleus (yellow arrows), in poorly preserved or degenerated tissues DAPI stain spreads to the cytoplasm (white arrow) and it is very common for the probe to be detected along the cell wall (b). Spoiled DAPI solution will stain the cell walls (b).