Immunogold Labeling Analysis of Cell Wall Polysaccharides with Special Reference to
(1;3,1;4)-β-D-glucan in Rice Cell Walls

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[Abstract] Various types of cell wall compositions have evolved to fulfill a wide range of
biological roles during the diversification of land plants. (1;3,1;4)-β-D-glucan (MLG) is a
defining feature of the cell walls in the order Poales (Yokoyama and Nishitani, 2004), which
has multiple functions associated with metabolic, growth, and defense systems. MLG is also a
characteristic component of the matrix polysaccharides that undergo turnover and metabolism,
depending on the tissue and the stage of development (Kido et al., 2015). Determining the
extracellular localization of MLG is essential for elucidating its functions. Electron microscopy
immunogold labeling analysis is a useful technique, which provides an accurate representation
of the extracellular distribution of MLG. This strategy is also applicable to various kinds of cell
wall polysaccharides, which have key roles in regulating growth and differentiation in each
plant species.

Materials and Reagents

1. Gelatin capsule (HF capsule) (Matsuya Co.)
2. Rice plants (see Note)
3. Paraformaldehyde (Electron Microscopy Sciences, catalog number: 12300)
4. Sodium cacodylate buffer solution (TAAB Laboratories BATCH, catalog number:
   90760)
5. Glutaraldehyde (Electron Microscopy Sciences, catalog number: 16220)
6. Tannic acid (Sigma-Aldrich, catalog number: 403040)
7. Phosphate buffer powder (Siyaku, Wako Pure Chemical Industries, catalog number:
   167-14491)
8. Ethanol (Siyaku, Wako Pure Chemical Industries, catalog number: 057-00456)
9. LR White resin (Electron Microscopy Sciences, London Resin Company, catalog
   number: 14381-CA)
10. Monoclonal antibody against (1;3,1;4)-β-D-glucan (mouse IgG, Kappa Light)
    (Biosupplies, catalog number: 400-3)
Note: It is named as (1-3;1-4)-β-D-glucan on Biosupplies' website.

11. Goat Anti mouse IgG pAb, Gold 15 nm, EM (BBI Solutions, catalog number: EM.GMHL15)
12. Phosphate-buffered salts (Takara Bio Company, catalog number: T900)
13. Bovine serum albumin (BSA) (Sigma-Aldrich, catalog number: A2058)
14. Uranyl acetate (Siyaku, Wako Pure Chemical Industries, catalog number: 554-85072)
15. Lead stain solution (Sigma-Aldrich, catalog number: 18-0875-2-25 ml-J)
   Note: This reagent may not be available outside Japan, and can be replaced with alternative lead stains. For alternative, readers are referred to a more general experimental manual for electron microscopy of plant cells (Hall and Hawes, 1991).
16. Molecular sieves (Sigma-Aldrich, catalog number: M6141)
17. Fixative solution (see Recipes)
18. 2% glutaraldehyde solution (see Recipes)
19. 100% ethanol solution (see Recipes)
20. 1x Phosphate-buffered saline (PBS) (see Recipes)

**Equipment**

1. Growth chamber (Nippon Medical & Chemical Instruments, model: LH220S)
2. Ultracut UCT Ultramicrotome (Leica Microsystems)
3. Diaphragm vacuum pump (Leybold-Heraus, model: Divac 2.2 L)
4. Desiccator (Sanplatec, model: PC-250K)
5. Grid mesh (Okenshoji, catalog number: 09-1079)
6. Transmission electron microscope (JEOL USA, model: JEM-1400plus)
7. CCD camera (Olympus soft imaging solutions GmbH, model: Veleta)

**Procedure**

1. Cut the rice leaf blade into pieces measuring ≤ 2 mm in length and immediately immerse them in a fixative solution. The fixative solution should be handled under a fume hood.
Figure 1. Embedding and trimming. a. Equipment and reagent. b. The tissues in the bottom of gelatin capsules filled with fresh 100% LR White resin. c. Polymerization. d. Trimming.

2. The tissue/fixative is placed in a desiccator, which is connected to a diaphragm vacuum pump. The tissue/fixative is vacuum degasified for 1 min at room temperature using the vacuum pump before releasing the vacuum very slowly.

3. Incubate the tissue/fixative for 2 h at 4 °C.

4. Rinse the tissues gently three times with 50 mM sodium cacodylate buffer (pH 7.4) for 15 min each time.

5. The tissues are dehydrated using a graded ethanol series (70%, 80%, 90%, and three times at 100%) at 4 °C for 30 min in each solution. Swirl occasionally.

6. Immerse the tissues three times with a 50:50 mixture of 100% (v/v) ethanol and 100% (v/v) LR White resin for 30 min each time at room temperature.

7. Transfer the tissues to 100% LR White resin and immerse the tissues three times for 1 h each time at room temperature. This step ensures that no residual ethanol will be carried over since other chemical like ethanol or else prevents polymerization of LR resin.

8. The tissues are settled to the bottom of gelatin capsules filled with fresh 100% LR White resin (Figure 1b). Stand the gelatin capsules in the test-tube rack for polymerization overnight at 50 °C (Figure 1c).

9. Turn the tissue block upside down and secure it in a UCT ultramicrotome. Chuck and trim the block to expose the tissue surface area up to 1.0 sq mm using a razor blade (Figure 1d).
10. Cut the block to 80 nm with a diamond knife on a UCT ultramicrotome (Figure 2a). Dip up the serial sections floating on water in the boat of the diamond knife with a nickel grid mesh, and mount the ultrathin sections on the grid (Figure 2b).

11. Incubate the grid with a droplet of the mouse anti-(1;3,1;4)-β-D-glucan antibody (diluted 1:1,000 in a 1% BSA solution, depending on the abundance of the target primary antibody) at room temperature for 90 min (Figure 3a).

12. Rinse the grid three times with 1% BSA/1x PBS for 1 min each time.

13. Float the grid on a droplet of goat anti-mouse IgG coupled to 15-nm gold particles 1:20 in 1% BSA/1x PBS at room temperature for 1 h. When detecting other primary antibody, prepare goat anti-mouse IgG coupled to gold particles to match the primary antibody.

14. Rinse the grid and the sections three times with 1x PBS.

15. Immerse the grid and the sections in 2% glutaraldehyde solution and are air-dried under a fume hood.

16. Immerse the sections in a droplet of 2% uranyl acetate, which had been prepared by diluting with distilled water, for 15 min (Figure 3b).
17. Rinse the sections with distilled water.

18. Immerse the sections with Lead stain solution at room temperature for 3 min. Rinse the sections with distilled water.

19. Observe the sections using a JEOL JEM 1200EX transmission electron microscope at an accelerating voltage of 80 kV.

20. Digital images (2,048 x 2,048 pixels) are obtained with a CCD camera.

Representative data

![Representative data](image)

**Figure 4. Immunogold-based localization of MLG is shown in the leaf blades.** a. The images in (a) are of the cell walls of the epidermal cell with with a wart-like protuberance (WP). Scale bar = 5 μm. b. A higher magnification of the rectangle in (a). Scale bar = 1 μm. c. The images in (c) are of the cell walls of the cortical sclerenchyma. Scale bar= 5 μm. d. A higher magnification of the rectangle in (c). Scale bar = 1 μm.

**Notes**

Rice (*Oryza sativa* L. cv Nipponbare) plants are grown in a growth chamber at 28 °C under a 15/9 h light/dark cycle (light at 150 μmol m⁻² s⁻¹). This protocol is performed using the fifth leaf blades of the rice plants.

**Recipes**

1. Fixative solution
   
   Prepare fixative solution by dissolving 0.4 g of paraformaldehyde, 0.05 g of tannic acid, and 0.1 ml of 10% glutaraldehyde in 9.9 ml of 50 mM sodium cacodylate buffer (pH
7.4). Heat this solution in the fume hood on the hotplate/stirrer to approx. 70 °C until the solution clears.

2. 2% glutaraldehyde solution

Dissolve an appropriate volume of Wako Pure Chemical Industries ready-made powder in distilled water according to manufacture’s protocol, to make a 0.1 M phosphate buffer of pH 7.4 at room temperature. Prepare 2% glutaraldehyde solution by diluting 10% glutaraldehyde in the phosphate buffer.

3. 100% ethanol solution

To prepare the 100% ethanol solution, use 100% bulk ethanol with molecular sieves in the bottom of the bottle

4. 1x PBS

Dissolve 10 PBS tablets in distilled water to make a total volume of 1,000 ml

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References

