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Chapter 17

Isolation of plant cell wall proteins

Running head: Plant cell wall proteins

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Abstract

The quality of a proteomic analysis of a cell compartment strongly depends on the reliability of the isolation procedure for the cell compartment of interest. Plant cell walls possess specific drawbacks: (i) the lack of a surrounding membrane may result in the loss of cell wall proteins (CWP) during the isolation procedure; (ii) polysaccharide networks of cellulose, hemicelluloses and pectins form potential traps for contaminants such as intracellular proteins; (iii) the presence of proteins interacting in many different ways with the polysaccharide matrix require different procedures to elute them from the cell wall. Three categories of CWP are distinguished: labile proteins that have little or no interactions with cell wall components, weakly bound proteins extractable with salts, and strongly bound proteins. Two alternative protocols are described for cell wall proteomics: (i) non-destructive techniques allowing the extraction of labile or weakly bound CWP without damaging the plasma membrane; (ii) destructive techniques to isolate cell walls from which weakly or strongly bound CWP can be extracted. These protocols give very low levels of contamination by
intracellular proteins. Their application should lead to a realistic view of the cell wall proteome at least for labile and weakly bound CWP extractable by salts.

Key Words: *Arabidopsis thaliana*, cell wall, proteomics, plant, bioinformatics, cell wall protein, cell fractionation

1. Introduction

Plant cell wall proteins (CWP) present specific complexities in addition to the difficulties usually encountered in proteome analysis, such as protein separation and detection of scarce proteins (1). They are embedded in an insoluble polysaccharide matrix and interact with other cell wall components, making their extraction challenging. Current models of cell wall structure describe the arrangement of their components into two structurally independent and interacting networks, embedded in a pectin matrix (2,3). Cellulose microfibrils and hemicelluloses constitute the first network; the second one is formed by structural proteins. Three types of CWP can be distinguished, according to their interactions with cell wall components (4). CWP can have little or no interactions with cell wall components and thus move freely in the extracellular space. Such proteins can be found in liquid culture media of cell suspensions and seedlings or can be extracted with low ionic strength buffers. We call this fraction “labile proteins”, most of them have acidic pI ranging from 2 to 6 (Fig. 1A). Alternatively, CWP might be weakly bound to the matrix by Van der Waals interactions, hydrogen bonds, hydrophobic or ionic interactions. Such proteins may be extracted by salts and most of them have basic pI ranging from 8 to 11 (Fig. 1B) so that they are positively charged at the acidic pH of cell walls. Even though most of the cell wall polysaccharides are neutral, negatively charged pectins contain polygalacturonic acid that provides negative charges for interactions with basic proteins.

![Fig. 1](image-url). pIs of labile and weakly-bound CWP. pIs of CWP identified in several proteomic studies (4) were calculated (www.iut-arles.up.univ-mrs.fr/w3bb/d_abim/compo-p.html) after removal of their predicted signal peptides (http://psort.nibb.ac.jp/form.html). Three groups of proteins were considered: (A) labile proteins; (B) salt-extracted proteins, *i.e.* proteins extracted with salt solutions or chelating agent; (C) all proteins. Reprinted from (4), Copyright (2005), with permission from Elsevier.
Such interactions would be modulated by pH, degree of pectin esterification, Ca\textsuperscript{2+} concentration, and by the mobility and diffusion coefficients of these macromolecules (3,5). Finally, CWP can be strongly bound to cell wall components so that they are still resistant to salt-extraction. As examples, extensins are cross-linked by covalent links (6,7) and peroxidases can have a high affinity for Ca\textsuperscript{2+}-pectate (8).

The available techniques described in this chapter allow the extraction of labile and weakly bound CWP. Since labile proteins can be lost during the preparation of cell walls, they must be extracted from tissues by non-destructive techniques such as vacuum infiltration (9), or recovered from liquid culture media from cell suspension cultures or seedlings (10,11). Weakly bound CWP can be extracted with salts or chelating agents from living cells with non-destructive techniques (9,10) or from purified cell walls with destructive techniques. At present, there is no efficient procedure to release CWP strongly bound to the extracellular matrix. Structural proteins, for instance extensins or PRP, can be cross-linked via di-isodityrosine bonds (6,12). Purified cell walls appear as the most suitable material to isolate such proteins. However, until now, extensins have only been eluted with salts prior to their insolubilization from cell suspension cultures (13).

2. Materials

A major problem in proteomics is the occurrence of keratins that can contaminate materials and working solutions. The presence of keratins can prevent the identification of proteins of interest by mass spectrometry. It is necessary to pay attention to all possible sources of contamination at all steps of the following protocols. Powder-free gloves should be permanently worn and washed with soap before their first use. Chemicals should be reserved for proteomic studies and should not be manipulated with spatula. Buffers should be filtered on 0.22 µm pore size filters. Glass plates for electrophoresis should be cleaned with alcohol before use.

2.1. Extraction of labile or weakly bound CWP by non-destructive techniques

2.1.1. CWP extraction and analysis from liquid culture medium of seedlings

1. Murashige and Skoog (MS) culture medium: Murashige and Skoog (14) liquid medium (Sigma Chemical, St Louis, MO, USA) is supplemented with 10 g/L sucrose and adjusted to pH 5.8 with KOH.
2. PVPP (Sigma, St. Louis, MO, USA) is treated with acid to increase polymerization and to remove metal ions and contaminants. One g PVPP in 10 mL 10% HCl is boiled for 10 min, filtered through a G4 filter, and rinsed until neutral pH is reached. The residue is dehydrated with acetone and grinded in a mortar to obtain a fine powder (15).
3. Low binding 12 kDa cutoff Spectra/Por\textsuperscript{®} cellulose ester (CE) dialysis bags (Merck Eurolab Poly Labo, Strasbourg, France).
4. Centriprep\textsuperscript{®} device (MWCO: 10 kDa) (Millipore Corporation, Bedford, MA, USA).
5. Bradford protein assay (Coomassie\textsuperscript{®} Protein assay Reagent Kit, Pierce, Rockford, IL, USA) (16).
6. Immobilized pH gradient (IPG) buffers and 13 cm-strips pH 4-7 or 6-11 (GE Healthcare Europe GmbH, Orsay, France).
7. 2-DE (2-dimensional electrophoresis) sample buffer: 7 M urea, 2 M thiourea, 4% (w/v) CHAPS, 65 mM DTE, 0.5% (v/v) IPG buffer (pH 4-7 or 6-11), bromophenol blue trace.
2.1.2. CWP extraction and analysis from cell suspension cultures

1. Gamborg liquid medium: Gamborg B5 medium supplemented with 20 g/L sucrose, 2.5 μM naphthalene acetic acid (Sigma Chemical, St Louis, MO, USA) and adjusted to pH 5.7 with KOH (17).
2. Cell washing buffer: 50 mM sodium acetate buffer pH 6.5, 10 mM DTT, 1 mM PMSF, 1% ethanol, 50% glycerol.
3. C1 protein extraction buffers: 0.15 M NaCl in cell washing buffer.
5. C3 protein extraction buffer: 0.2 M CaCl₂ dihydrate in cell washing buffer.
6. C4 protein extraction buffer: 2 M LiCl in cell washing buffer.
7. C5 protein extraction buffer: 50 mM 1,2-cyclohexanediamine tetracetic acid (CDTA) in cell washing buffer.
8. Low binding 2 kDa cutoff Spectra/Por® CE dialysis bags (Merck Eurolab Poly Labo, Strasbourg, France).
9. One mL Hi-Trap SP Sepharose cation exchanger (GE Healthcare Europe GmbH, Orsay, France).
10. Hi-Trap SP equilibration buffer: 10 mM MES-KOH, pH 5.2.
11. Hi-Trap SP elution buffer: 10 mM MES-KOH, pH 5.2, 2 M NaCl.
12. Econo-Pac® 10DG desalting column (Bio-Rad, Hercules, CA, USA).
13. Desalting column equilibration buffer: 50 mM ammonium formate.
14. Bradford protein assay (Coomassie® Protein assay Reagent Kit, Pierce, Rockford, IL, USA) (16).
15. 1-DE (1-dimensional electrophoresis) sample buffer: 62 mM Tris-HCl, pH 6.8, 2% SDS, 10% glycerol, 5% mercapto-ethanol.
16. Resuspending solution: 1 M thiourea, 10 mM DTT, 1% (v/v) protease inhibitor cocktail for plant (Sigma, St. Louis, MO, USA) in UHQ water (see Note 1). Prepare as required.
17. Immobilized pH gradient (IPG) buffers and 13 cm-strips pH 4-7 or 6-11 (GE Healthcare Europe GmbH, Orsay, France).
18. 2-DE sample buffer: 7 M urea, 2 M thiourea, 4% (w/v) CHAPS, 65 mM DTE, 0.5% (v/v) IPG buffer (pH 4-7 or 6-11), bromophenol blue trace.

2.1.3. Extraction and analysis of CWP from rosette leaves

2.1.3.1. Extraction of proteins

1. Recovering solution: 0.3 M mannitol, 66 mM DTT, 330 mM thiourea, 3.3% (v/v) protease inhibitor cocktail for plant (Sigma, St. Louis, MO, USA) (see Note 1). Prepare as required.
2. R1 protein extraction buffer: 1 M NaCl, in recovering solution. Adjust pH to 6.9 with 0.5 N NaOH. Prepare as required
3. R2 protein extraction buffer: 0.2 M CaCl₂ dihydrate, in recovering solution. Adjust pH to 6.9 with 0.5 N NaOH. Prepare as required.
4. R3 protein extraction buffer: 2 M LiCl, 0.3 in recovering solution. Adjust pH to 6.9 with 1 N NaOH. Prepare as required.
5. R4 protein extraction buffer: 50 mM CDTA, in recovering solution. Adjust pH to 6.9 with 5 N NaOH. Prepare as required.
6. Malate dehydrogenase (MDH) assay mixture: 50 mM Tris-HCl, pH 7.8 (2.15 mL), 50 mM MgCl₂ hexa-hydrate (300 μL), 150 mM DTT (100 μL), 10 mM NADP (150 μL), 30 mM malic acid (300 μL). Store DTT and NADP solutions in single use aliquots at -20°C.
Store malic acid solution at -20°C no longer than one month. Prepare the MDH assay mixture as required

2.1.3.2. Analysis of labile CWP
1. Low binding 2 kDa cutoff Spectra/Por® CE dialysis bags (Merck Eurolab Poly Labo, Strasbourg, France). Store at -20°C.
2. Resuspending solution: 1 M thiourea, 10 mM DTT, protease inhibitor cocktail (1% v/v) in UHQ water. Prepare as required.
3. Immobilized pH gradient (IPG) buffers and 7 cm-strips pH 4-7 (GE Healthcare Europe GmbH, Orsay, France).
4. 2 DE-sample buffer: 7 M urea, 2 M thiourea, 4% (w/v) CHAPS, 65 mM DTE, 0.5% (v/v) IPG buffer (pH 4-7 or 6-11), bromophenol blue trace.

2.1.3.3. Analysis of weakly-bound CWP
1. Low binding 2 kDa cutoff Spectra/Por® CE dialysis bags (Merck Eurolab Poly Labo, Strasbourg, France). Store at -20°C.
2. One mL Hi-Trap SP Sepharose cation exchanger (GE Healthcare Europe GmbH, Orsay, France).
3. Hi-Trap SP equilibration buffer: 10 mM MES-KOH, pH 5.2.
4. Hi-trap SP elution buffer: 10 mM MES-KOH, pH 5.2, NaCl 2 M.
5. Econo-Pac® 10DG desalting column (Bio-Rad, Hercules, CA, USA).
6. Desalting column equilibration buffer: 50 mM ammonium formate.
7. 1-DE sample buffer: 62 mM Tris-HCl, pH 6.8, 2% SDS, 10% glycerol, 5% mercaptoethanol.
8. Resuspending solution: 1 M thiourea, 10 mM DTT, 1% (v/v) protease inhibitor cocktail in UHQ water. Prepare as required.
9. Immobilized pH gradient (IPG) buffers and 7 cm-strips pH 4-7 (GE Healthcare Europe GmbH, Orsay, France).
10. 2 DE-sample buffer: 7 M urea, 2 M thiourea, 4% (w/v) CHAPS, 65 mM DTE, 0.5% (v/v) IPG buffer (pH 4-7 or 6-11), bromophenol blue trace.

2.2. Extraction of weakly-bound CWP by destructive techniques

2.2.1 Cell wall preparation
1. MS solid medium: Murashige and Skoog liquid medium (Sigma Chemical, St Louis, MO, USA) is supplemented with 20 g/L sucrose and 12 g/L agar, and adjusted to pH 5.8 with KOH.
2. PVPP (Sigma, St. Louis, MO, USA) is treated with acid to increase polymerization and to remove metal ions and contaminants. One g PVPP in 10 mL 10 % HCl is boiled for 10 min, filtered through a G4 filter, and rinsed until neutral pH is reached. The residue is dehydrated with acetone and grinded in a mortar to obtain a fine powder.
3. Nylon nets (1.5 mm pore size and 25 µm pore size).
4. Waring blender with a 2 L flask (SEB Moulinex, Ecully, France).
5. Grinding buffer: 10 mM acetate buffer, pH 4.6, 0.4 M sucrose, 0.2 % (v/v) protease inhibitor cocktail for plant (Sigma, St. Louis, MO, USA) (see Note 1).
6. Cell wall purification buffers: 5 mM acetate buffer, pH 4.6, 0.6 M or 1 M sucrose, 0.2 % (v/v) protease inhibitor cocktail.
2.2.2 Extraction and separation of proteins
1. H1 protein extraction buffer: 5 mM acetate buffer, pH 4.6, 0.2 M CaCl₂, 0.1% protease inhibitor cocktail for plant (Sigma, St. Louis, MO, USA).
2. H2 protein extraction buffer: 5 mM acetate buffer, pH 4.6, 2 M LiCl, 0.1% protease inhibitor cocktail.
3. Econo-Pac® 10DG desalting column (Bio-Rad, Hercules, CA, USA).
4. Desalting column equilibration buffer: 50 mM ammonium formate.
5. Bradford protein assay (Coomassie® Protein assay Reagent Kit, Pierce, Rockford, IL, USA) (16).
6. 1-DE sample buffer: 62 mM Tris pH 6.8 (HCl), 2% SDS, 10% glycerol, 5% mercapto-ethanol.

3. Methods

The choice of a protocol to extract CWP for proteomic analysis is dependent on the plant material and of the type of proteins to be released from cell walls. Working on living cells is probably the best solution to avoid intracellular contamination. This is possible for cell suspension cultures or seedlings grown in liquid medium as well as for any plant organ that can be infiltrated under vacuum with extraction buffers. Both labile and weakly-bound CWP can be released. When this is not possible, it is necessary to purify cell walls. The main problem is to avoid intracellular contaminants that will stick non-specifically to cell walls. Only weakly-bound CWP can be extracted from purified cell walls since labile CWP are lost during cell wall preparation.

Another important point is the choice of the extraction solution. For example, a solution of 0.3 M mannitol infiltrated in living tissues such as leaves can solubilize a few CWP expected to be located only in intercellular spaces. Indeed, identified proteins are acidic, suggesting no ionic interactions with negatively charged cell wall components (9). NaCl is usually used for extraction of proteins retained by ionic interactions in the cell wall. LiCl can extract hydroxyproline-rich glycoproteins from intact cells in Chlamydomonas reinhardii (18). Calcium chloride is probably the most efficient salt to extract CWP (9,19). The ability of acidic and neutral carbohydrates to strongly chelate calcium (20,21) might explain, through a competition mechanism, that proteins or glycoproteins weakly bound to cell wall polysaccharides can be selectively solubilized by CaCl₂. CDTA, a chelating agent, solubilizes Ca²⁺-pectate. It releases a small number of proteins having domains of interaction with polysaccharides, notably proteins showing homology to lectins. This suggests an interaction of these proteins with polysaccharides associated to pectins (9).

3.1. Extraction of labile or weakly bound CWP by non-destructive techniques

3.1.1. Liquid culture medium of seedlings
1. Soak A. thaliana ecotype Columbia seeds (see Note 2) in tap water for 2 h, then sterilize in diluted bleach (2.4% w/v) for 45 min, and rinse several times with deionized water.
2. Sterilized seeds (100 mg) are germinated and grown in MS liquid culture medium in ten 1 L-flasks on a rotary shaker (90 rpm) at 26 °C in the dark (22). Each flask contains 130 mL of culture medium. After 14 days, etiolated seedlings are harvested and the culture medium is filtered through nylon net (60 µm) to remove cell debris.
3. Collect 900 mL of culture medium (see Note 3). Mix with 9 g PVPP. Shake the mixture at 4 °C for at least 30 min, filter and centrifuge to pellet the insoluble residue. Dialyze against 10 L distilled water during 10-12 h at 4 °C using a dialysis bag (MWCO: 12 kDa) with 3 changes. Reduce the volume of the sample by repeated centrifugations (3500 × g for 15 min at 4 °C) through a Centriprep® device (MWCO: 10 kDa) to about 1 mL.


5. Dilute 250 µg proteins extracted from culture medium with 250 µL 2-DE sample buffer. Proteins are separated by 2-DE using 13 cm- IPG gel strips pH 4-7 or 6-11 for the first dimension.

3.1.2. Cell suspension cultures

1. A cell suspension culture of *A. thaliana* ecotype Columbia is grown on Gamborg liquid medium. From this culture, 50 mL (25 g) is transferred every 2 weeks to 250 mL fresh medium in 1 L Erlenmeyer flask and shaken at 70 rpm in an orbital shaker, under continuous light (30 μE.m⁻².s⁻¹) at 22°C.

2. Wash cells of 7-day old *A. thaliana* suspension culture with water and pellet them by centrifugation at 200 × g. Plasmolyze by successive immersion in 25 % glycerol, 50 % glycerol for 10 min each, and finally wash in 50 % cold glycerol. All subsequent extractions are performed at 0°C except otherwise stated.

3. Prior to protein extraction, wash cells with cell washing buffer in order to remove contaminant proteins coming from broken cells that non-specifically stick to cell walls.

4. Extract proteins by washings of the plasmolyzed cells under gentle stirring (30 min) in the proportion of 25 mL of pelleted cells per 50 mL of solution. First extraction is performed with C1 buffer.

5. Wash cells with the same extraction buffer, then with 50 % glycerol before centrifugation at 200 × g for 5 min.

6. Extract weakly-bound CWP with C2, C3, C4 or C5 buffer in the same way (see Note 4).

7. Exhaustively dialyze protein extracts at 4°C against 20 L H₂O using a dialysis bag (MWCO: 2 kDa). Measure the protein content of each extract using the Bradford protein assay.

8. Dialyze against Hi-Trap SP equilibration buffer and apply to a Hi-Trap SP Sepharose column equilibrated with Hi-Trap SP equilibration buffer at 1 mL.min⁻¹. Elute the retained basic proteins with Hi-Trap SP elution buffer at 1 mL.min⁻¹. Desalt the basic proteins on an Econo-Pac® 10DG desalting column equilibrated with desalting column equilibration buffer. Freeze-dry the eluate. Resuspend the dry residue in 40 µL 1-DE sample buffer and separate proteins by 1-DE on a 10%-17% gradient polyacrylamide gel (16.5 x 13.5 x 0.15 cm).

9. Freeze-dry the acidic and neutral proteins from the Hi-Trap SP Sepharose column effluent. Solubilize the dry residue with a minimal volume of resuspending solution and desalt on an Econo-Pac® 10DG desalting column. Freeze-dry the proteins, dissolve in 2-DE sample buffer and perform a 2-DE using a 7 cm-IPG gel strip pH 4-7 for the first dimension.

3.1.3. Rosette leaves

3.1.3.1. Extraction of proteins

1. Sterilize *A. thaliana* ecotype Columbia seeds by soaking in diluted bleach (2.6% w/v) for 5 min and rinse several times with deionized water. Sow the seeds on humid compost in 10 x 10 cm pots and cover the pots with a plastic film. Remove the plastic
film after 48 h and transfer the cultures in a growth chamber at 70% humidity, with a photoperiod of 9 h light at 110 µE.m⁻².s⁻¹ at 22°C, and 15 h dark at 20°C. Plants should be moderately watered with a nutrient solution once a week.

2. Remove carefully 4- to 5-week-old plants (Fig. 2A) from the pots and wash compost off with deionized water. Cotyledons and yellowish leaves should be systematically removed from plants. Process whole plants for vacuum-infiltration as follows. Make a small noose with a piece of string and pass the root through the noose. Tighten the noose around the collar then twist the root around the string and wrap in parafilm. In a large beaker, immerse completely the rosettes first in distilled water for a few seconds in recovering solution. Put the beaker with the immersed rosettes in a dessicator connected to a vacuum pump (Fig. 2B). Vacuum-infiltrate the rosettes for 2 min after starting the pump. Reintroduce carefully air in the dessicator after vacuum breakage (Fig. 2C). Transfer the infiltrated plants to a centrifuge tube, with the collar at about 1 cm at the edge of the tube (Fig. 2D). Paste the lower part of the root outside of the tube with adhesive tape. Introduce at the bottom of the centrifuge tube 300 µL of recovering solution. Centrifuge infiltrated plants in swinging buckets at 200 × g for 17 min at 20°C (see Note 5). Collect the apoplastic washing fluids with a micropipette and estimate the volume. Vacuum-infiltration and centrifugation should be repeated twice.

Fig. 2. Vacuum-infiltration of rosette leaves. Four steps of the procedure are illustrated: (A) 4-5 week-old plants; (B) vacuum-infiltration of immersed rosettes in a dessicator connected to a vacuum pump; (C) rosette leaves after vacuum-infiltration, note the darker part of a leaf after successful infiltration (black arrow); (D) infiltrated plant transferred to a centrifuge tube, note the drop of solution containing the protease inhibitor cocktail at the bottom of the tube (black arrow).
3. Assay the apoplastic fluids for malate dehydrogenase (MDH) activity in order to detect cytoplasmic contaminations. Measure MDH activity at room temperature in 3 mL MDH assay mixture and one-twentieth of the volume of the recovered apoplastic fluids. Reduction of NADP is followed at $\lambda=340$ nm. Pool only those apoplastic washing fluids with no detectable MDH activity.

4. Vacuum-infiltrate rosettes with R1, R2, R3 or R4 buffer. Check for MDH activity on the recovered apoplastic fluids as described above. Discard any apoplastic washing fluids with MDH activity. Pool the remaining apoplastic washing fluids free of MDH activity.

3.1.3.2. Analysis of labile CWP by 2-DE

1. Exhaustively dialyze the apoplastic washing fluids from rosettes infiltrated with the recovering solution at 4°C against deionized water in low binding 2 kDa cutoff Spectra/Por® CE dialysis bags. Freeze-dry the dialysates. Resuspend the dry residues in 3 mL of resuspending solution and desalt on an Econo-Pac® 10DG desalting column equilibrated with desalting column equilibration buffer for the complete removal of mannitol. Freeze-dry the eluate.

2. Solubilize the dry residue in 2-DE sample buffer and separate proteins by 2-DE using a 7 cm-IPG gel strip pH 4-7 for the first dimension.

3.1.3.3. Analysis of weakly-bound CWP

1. Exhaustively dialyze the apoplastic washing fluids from rosettes infiltrated with R1, R2, R3 or R4 buffer against Hi-Trap SP equilibration buffer as described above. Apply to a Hi-Trap SP Sepharose column equilibrated with Hi-Trap SP equilibration buffer at 1 mL.min$^{-1}$. Elute the retained basic proteins with Hi-Trap SP elution buffer at 1 mL.min$^{-1}$. Desalt the basic proteins on an Econo-Pac® 10DG desalting column equilibrated with desalting column equilibration buffer. Freeze-dry the eluate. Resuspend the dry residue in 40 $\mu$L 1-DE sample buffer and separate proteins by 1-DE on a 10%-17% gradient SDS-polyacrylamide gel (16.5 x 13.5 x 0.15 cm).

2. Freeze-dry the acidic and neutral proteins in the Hi-Trap SP Sepharose column effluent. Solubilize the dry residue with a minimal volume of resuspending solution and desalt on an Econo-Pac® 10DG desalting column equilibrated with desalting column equilibration buffer. Freeze-dry the proteins and perform a 2-DE.

3.2. Extraction of weakly bound CWP by destructive techniques

3.2.1. Cell wall preparation

1. Soak *A. thaliana* ecotype Columbia seeds (*see Note 2*) in tap water for 2 h, then sterilize in diluted bleach (2.4 %) for 45 min, and rinse several times with deionized water. Sow the seeds (150 mg) in a Magenta box (6 cm x 6 cm) containing 50 mL of solid MS medium. Grow seedlings at 23°C in the dark for 11 days (*see Note 6*).

2. Harvest hypocotyls (around 2 cm high) of an average of 20 Magenta boxes as follows. First, remove carefully the solid MS medium carrying the seedlings from each box. Then, cut hypocotyls below cotyledons and above root with a pair of scissors. Wash the 1 cm-long hypocotyls with distilled water onto a nylon net (1.5 mm pore size) to remove all the cut cotyledons and seed coats that stick to hypocotyls (*see Note 7*). Transfer the hypocotyls into 500 mL of grinding buffer and add PVPP (1g/10g fresh weight of hypocotyls) to complex phenolic compounds.

3. Grind the mixture in cold room using a Waring blender at full speed for 15 min.
4. Separate cell walls from soluble cytoplasmic fluid by centrifugation of the homogenate for 15 min at $1000 \times g$ and $4^\circ C$. Further purify the pellet by two successive centrifugations in 500 mL of cell wall purification buffers, 0.6 M and 1 M sucrose respectively.

5. Wash the residue with 3 L of cell wall washing buffer on a nylon net (25 µm pore size) in order to eliminate all soluble compounds. Grind the resulting cell wall fraction in liquid nitrogen in a mortar with a pestle prior to lyophilization. Starting with 16 g fresh weight of hypocotyls, this procedure usually results in 1.3 g dry powder.

3.2.2. Extraction of proteins

1. Typically, 0.65 g of lyophilized cell walls is used for one experiment. Extract proteins by successive salt solutions in this order: two extractions with 6 mL H1 buffer, followed by two extractions with 6 mL H2 buffer. Resuspend cell walls by vortexing for 5-10 min at room temperature, and then centrifuge for 15 min at $4000 \times g$ and $4^\circ C$. Supernatants from the same extracting buffer are pooled.

2. Desalt supernatants using Econo-Pac® 10DG desalting columns equilibrated with desalting column equilibration buffer. Lyophilize the extracts and resuspend in H2O2.


4. Add 1-DE sample buffer. Separate proteins by 1-DE on a 12.5% SDS-polyacrylamide gel.

3.3. Analysis of CWP

3.3.1. Specific constraints for separation by electrophoresis and protein identification by mass spectrometry

The separation of CWP by classical two-dimensional gel electrophoresis (2-DE) is difficult. Since most CWP are basic glycoproteins (Fig. 1C), they are poorly resolved by this technique (22). They are better separated by 1-DE. However, protocols including chromatographic steps to separate proteins prior to 1-DE are available (24,25). In this chapter, a method able to separate acidic and basic CWP is proposed for a better resolution of these two types of CWP in 2-DE or 1-DE respectively.

Frequently, in 1-DE, proteins are not well separated from one another, and a protein sample can contain a mixture of proteins. However, the peptide mass mapping technology using high resolution (< 20 ppm) MALDI-TOF mass spectrometry (MS) permits the identification of several proteins from a mixture. Search engines such as MS-FIT from Protein Prospector (http://prospector.ucsf.edu/ucsfhtml4.0/msfit.htm) or MASCOT (http://www.matrixscience.com/search_form_select.html) allow multiple searches. In case of difficulties, proteins can also be identified by peptide sequencing using LC (liquid chromatography)-MS/MS systems (9).

3.3.2. Use of bioinformatics for the evaluation of the efficiency of an extraction protocol

The reliability of protein profiling for a compartment like the cell wall, strongly depends on the quality of the preparation. Unfortunately, the classical methods to check the purity of a particular fraction are not conclusive for proteomic studies, since the sensibility of the analysis by mass spectrometry is 10 to 1000 times more sensitive than enzymatic or immunological tests using specific markers. Our experience in the field has shown that the most efficient way to evaluate the quality of a cell wall preparation is (i) to identify all the proteins extracted from the cell wall by mass spectrometry, and (ii) to perform extensive bioinformatic analysis to determine if the identified proteins contain a signal peptide, and no
retention signals for other cell compartments. Comparison of the results obtained with different programmes is necessary to ensure a reliable prediction: PSORT allows predicting any sub-cellular localization (http://psort.ims.u-tokyo.ac.jp/form.html); TargetP looks for the presence of signal peptides for protein secretion or of transit peptides for mitochondrion or chloroplast targeting (http://www.cbs.dtu.dk/services/TargetP/); Aramemnon compares the results of several programmes predicting the presence of signal peptides and trans-membrane domains (http://aramemnon.botanik.uni-koeln.de/). It is then possible to conclude about the quality of the cell wall preparation by calculating the ratio of predicted secreted proteins to intracellular ones.

4. Notes
1. Protease inhibitor cocktail for plant is required to prevent proteolysis during the extraction procedure. Proteolysis induces the production of smaller broken proteins that can be spread over 1-D or 2-D polyacrylamide gels. Thus, proteolysis can prevent the identification of both broken proteins and other proteins of interest by mass spectrometry. Moreover, the occurrence of these polypeptides is a great problem for quantitative and comparative proteomics.
2. Seeds germinate in culture media that are favourable to development of bacteria or fungi. Due to the high amount of seeds (150 mg) introduced in a culture flask or in a Magenta box, the possible contamination events are multiplied. So, seeds should be carefully sterilized, and the healthy state of plants should be very good during their production in greenhouses.
3. Culture media should be processed immediately after recovery. Otherwise, proteolysis may occur even if they are stored frozen.
4. No more than two successive extractions with salt solutions should be performed. Otherwise, cells are damaged and intracellular contaminants are released in the culture medium and can stick non-specifically to cell walls (10).
5. Be careful setting minimal acceleration in order to avoid seriously damaging the vacuum-infiltrated plants during the centrifugation step. Imperatively centrifuge in swinging buckets to get undamaged plants during spinning.
6. All seedlings should grow at about the same rate to reach the same size after 11 days. If germination is not homogeneous, place the boxes at 6°C during 2 days to allow all seeds to start germination without growth. Then, all boxes can be put at 23°C for 11 days.
7. Cotyledons should be carefully removed. They contain few protein species but each of them in a huge amount. Due to their density, cotyledons co-sediment with cell walls. As a consequence, few cotyledons induce a significant contamination during extraction of cell wall proteins, especially by storage proteins. This contamination prevents the identification of proteins of interest by mass spectrometry.

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References