

Techniques in Molecular Genetics

Polyacrylamide Gel
Electrophoresis
(PAGE) of Proteins

H.E. Schellhorn

PAGE Electrophoresis-Principle

PAGE can be used to separate proteins and nucleic acids.

Polyacrylamide is a crosslinked polymer of acrylamide and bis-acrylamide

Properties of the gel, especially limiting pore size, are determined by the total concentration of acrylamide-bisacrylamide (%T) and conc. of bis-acrylamide to total acrylamide (%C)

PAGE Electrophoresis-Polymerization

Two additional chemicals play important roles in the polymerization process.

Ammonium Persulfate---produces initiating free radicals when dissolved in water

TEMED- also produces free radicals

PAGE Electrophoresis-Other Chemicals



Sodium Dodecyl Sulfate

β -mercaptoethanol

PAGE Electrophoresis-Discontinuous

Most denaturing gels are composed of a *stacking gel* and a *separating gel*

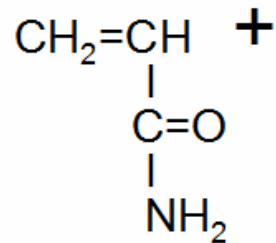
Stacking gel- low porosity, 2 pH units below running buffer

-allows proteins to form a compressed band after a few min.

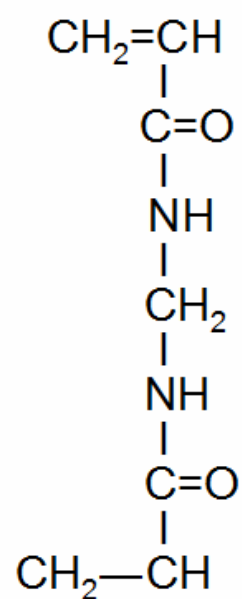
Separating gel- sieves protein according to size

PAGE Electrophoresis-Monomers

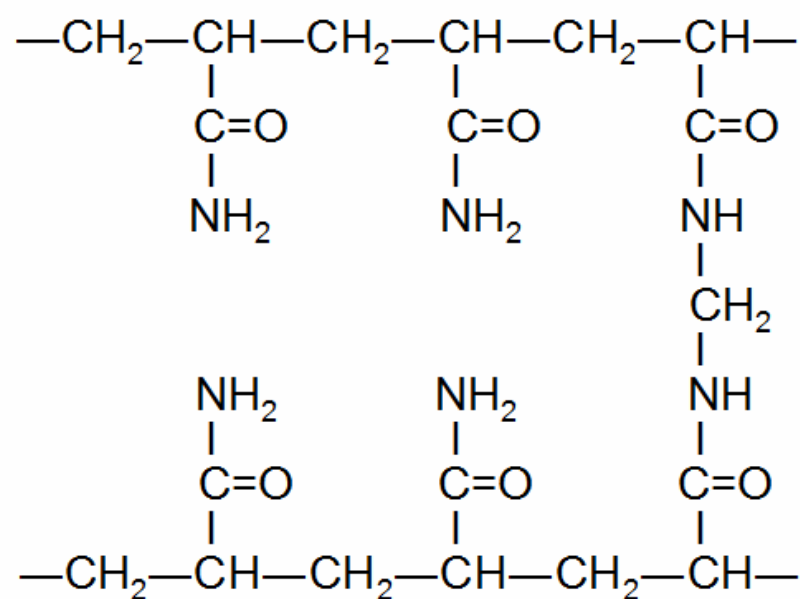
Acrylamide



Bis-acrylamide



Polyacrylamide



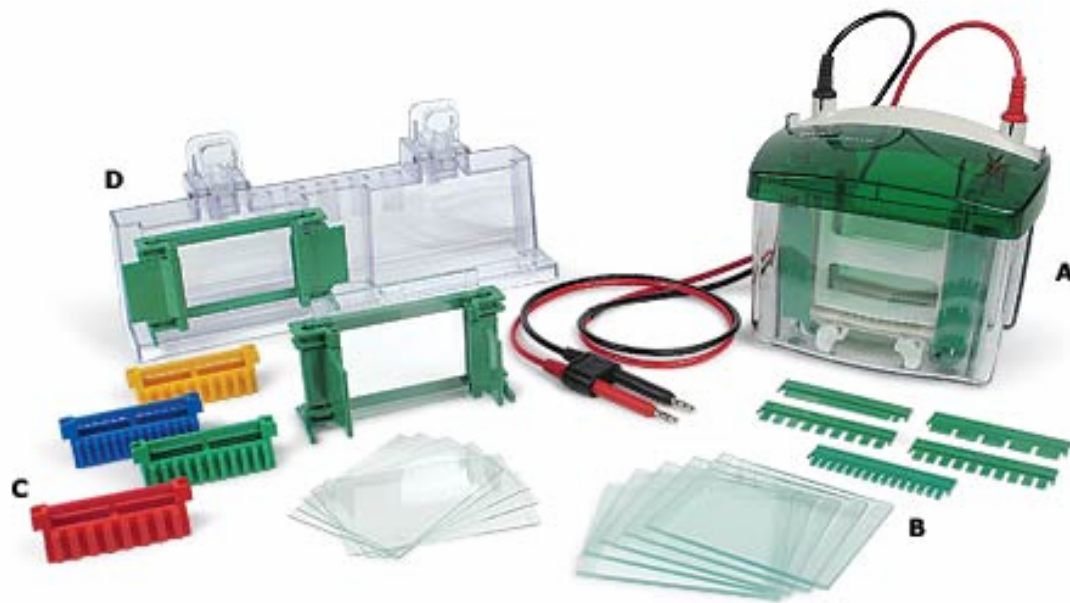
PAGE Electrophoresis-Types

- Native (non-denaturing)
- Two dimensional
- Gradient
- Isoelectric focussing

Common Problems in Using PAGE

Problem	Common Cause	Solution
Unequal lane width	Varying salt conc	Wash samples before
Vertical streaking	overload	Reduce sample
No bands	Not enough protein	Increase protein
Skewed bands	various	Use equal volumes, equal salt conc. in samples
Skewed bands at sides	Standard volume different from sample volume	Make standards up in sample buffer and use same volume as test samples

PAGE Electrophoresis Equipment



- A-Electrophoresis Cell
- B-Glass Plates
- C-Combs
- D-Casting Stand

Electrophoresis-Principle

Electrophoresis i

Fig. 2. Assembling the Mini-PROTEAN 3 cell.

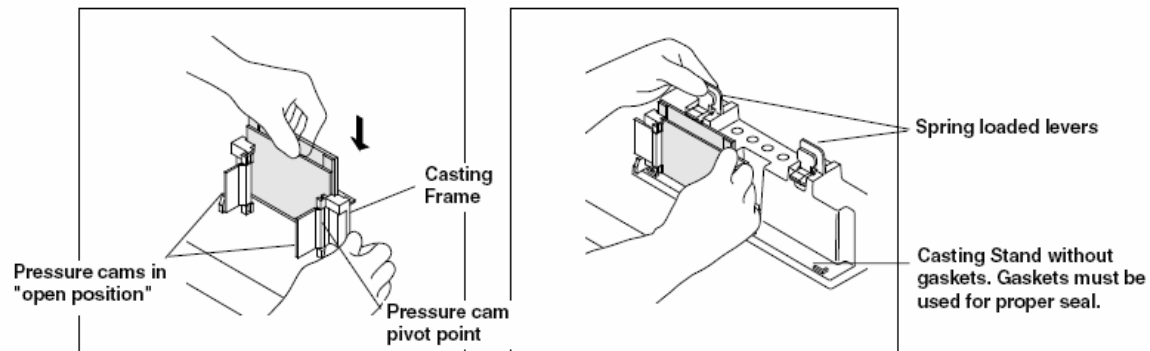
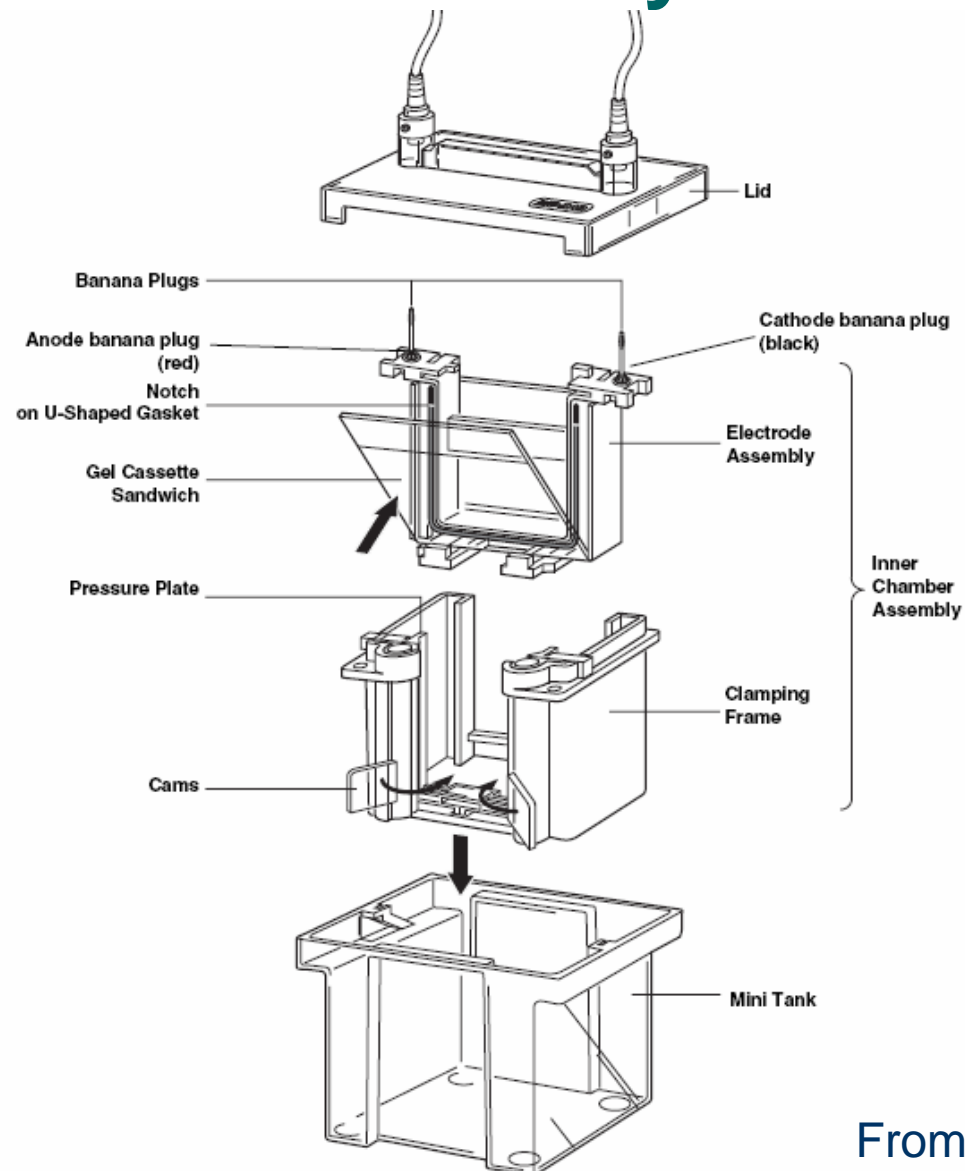


Fig. 3. Assembling the Mini-PROTEAN 3 Casting Frame and Casting Stand.

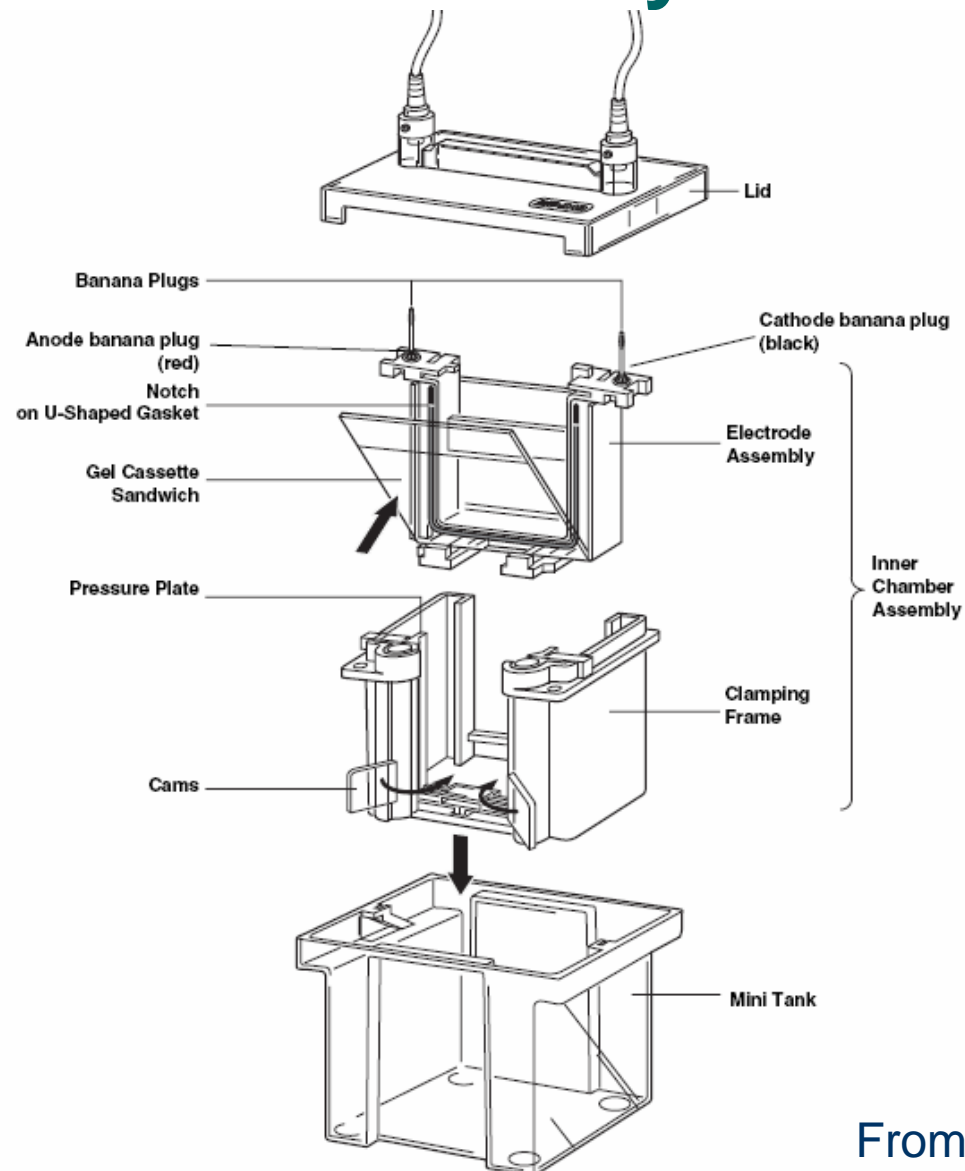
From Biorad Manual

Using the Biorad PAGE System...



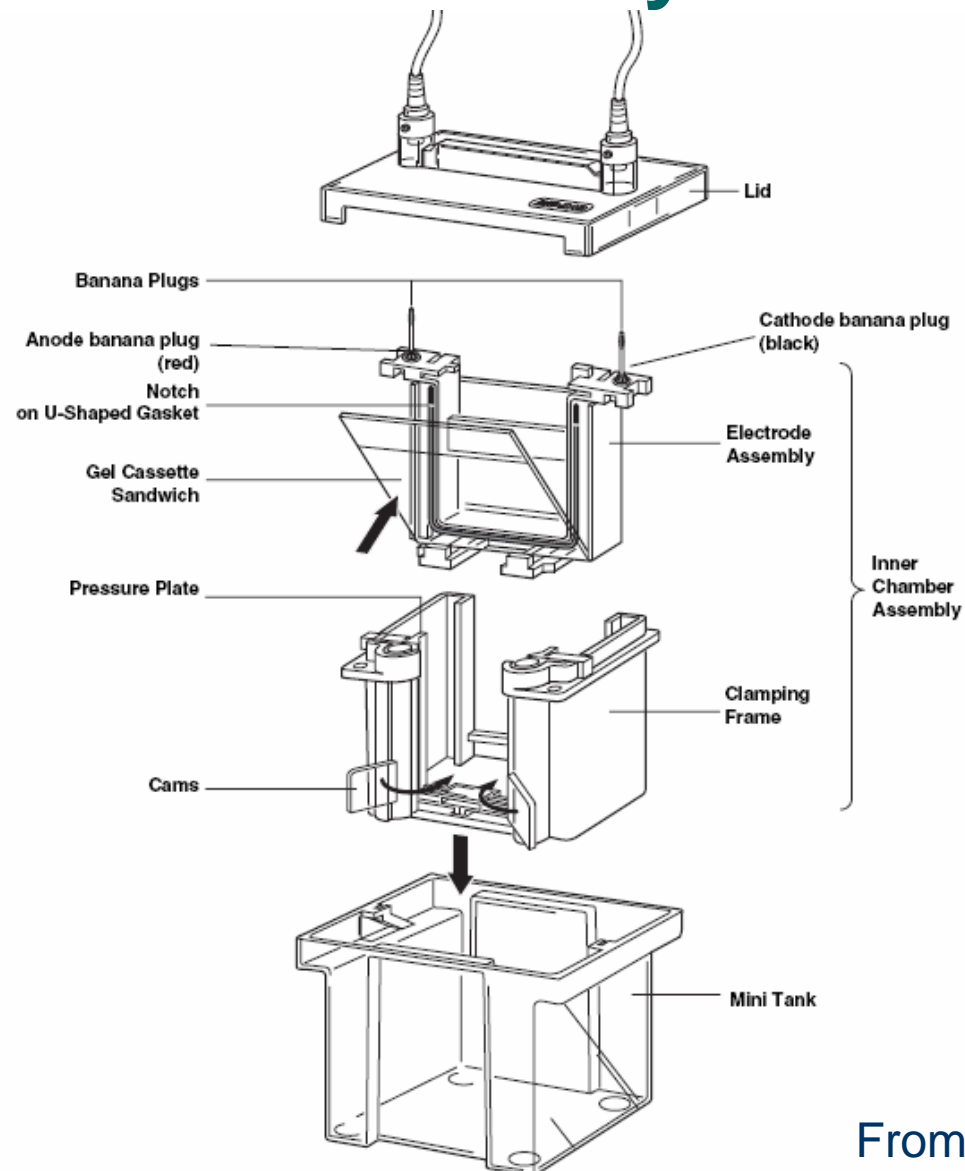
From Biorad Manual

Using the Biorad PAGE System...



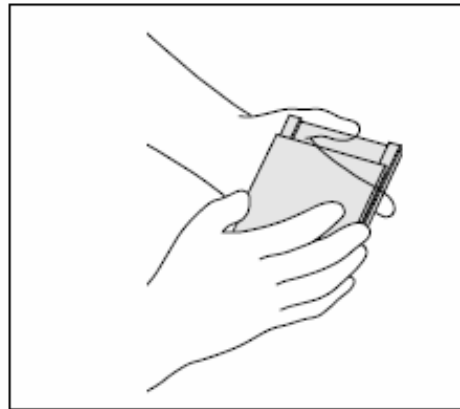
From Biorad Manual

Using the Biorad PAGE System...

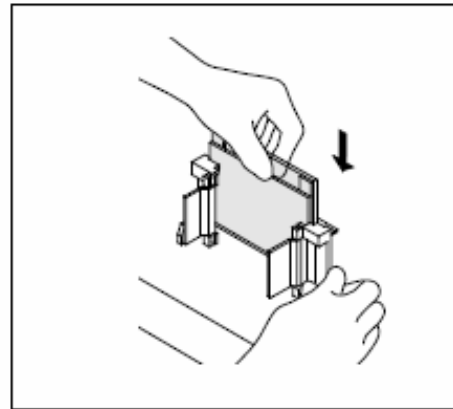


From Biorad Manual

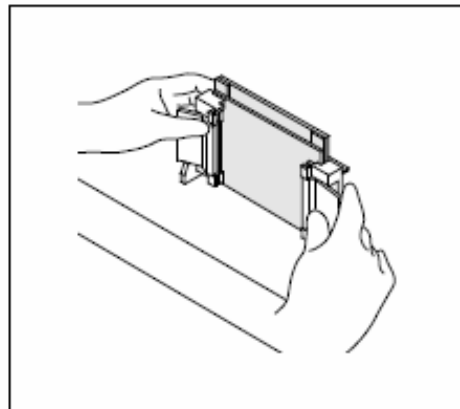
Using the Biorad PAGE System...



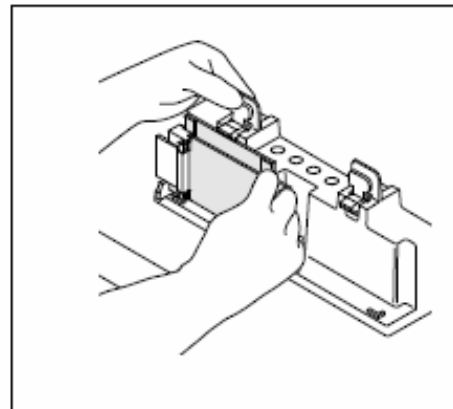
4a. Place a Short Plate on top of the Spacer Plate.



4b. Slide the two plates into the Casting Frame keeping the Short Plate facing front.



4c. Lock the pressure cams to secure the glass plates.

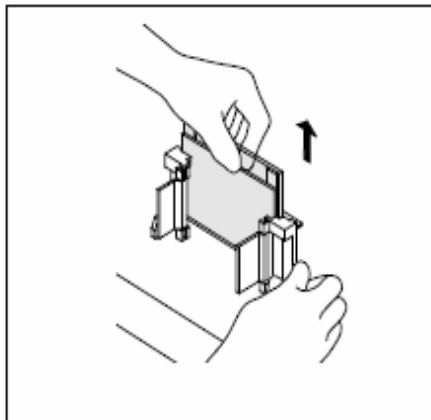


4d. Secure the Casting Frame in the Casting Stand by engaging the spring loaded lever.

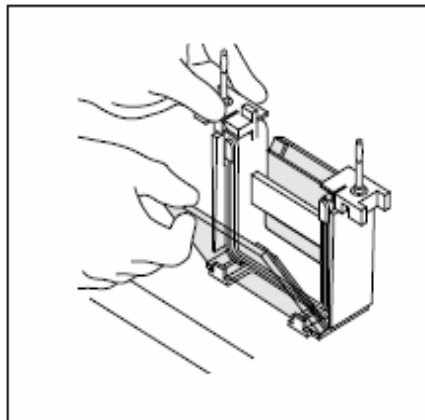
Fig. 4. Assembling the Mini-PROTEAN 3 casting stand and frame.

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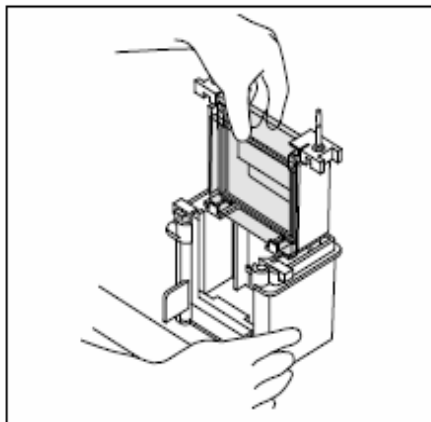
Using the Biorad PAGE System...



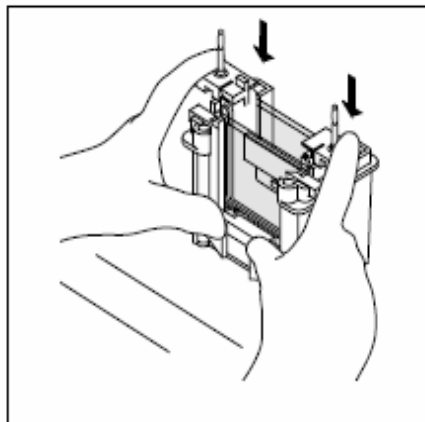
5a. Remove the Gel Cassette Sandwich from the Casting Frame.



5b. Place Gel Cassette Sandwich into the Electrode Assembly with the Short Plate facing inward.



5c. Slide Gel Cassette Sandwiches and Electrode Assembly into the clamping frame.



5d. Press down on the Electrode Assembly while closing the two cam levers of the Clamping Frame.

Using the Biorad PAGE System...

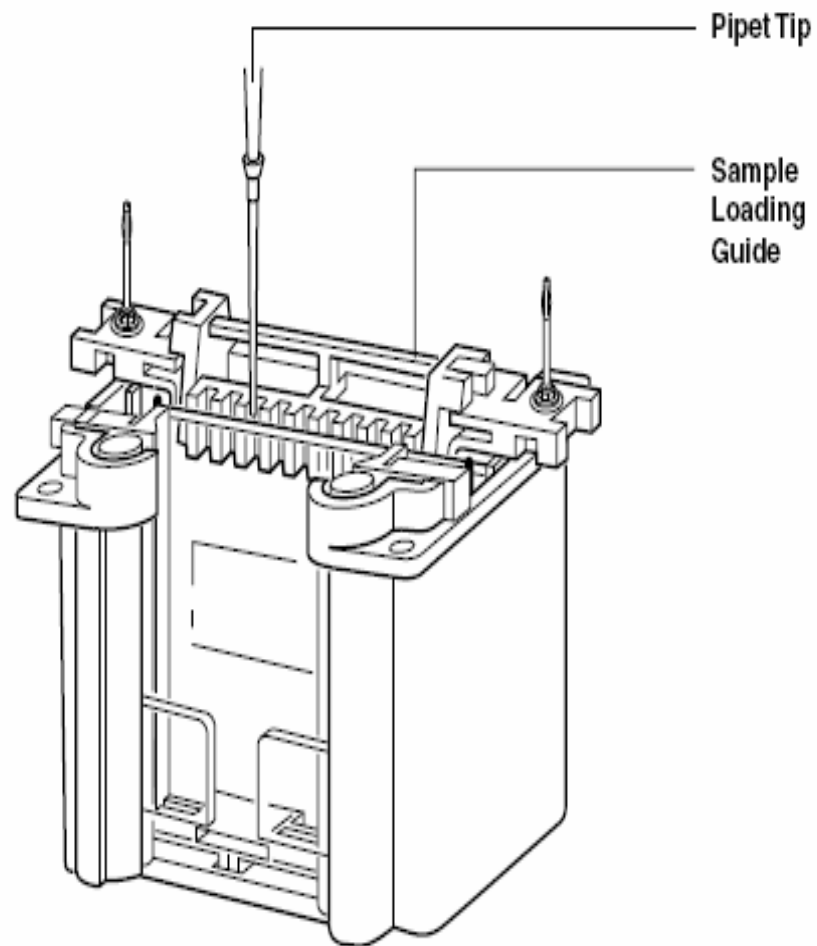
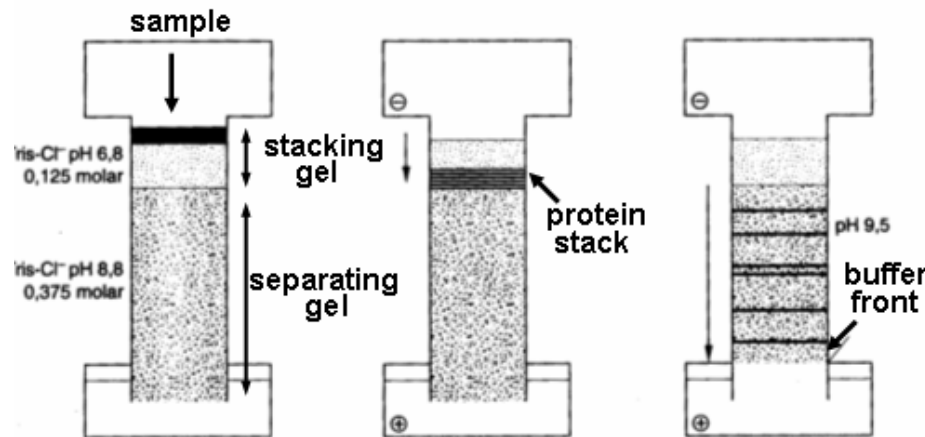


Fig. 6. Using the Sample Loading Guide (patent #5,656,145).

From Biorad Manual

SDS-PAGE: Separation process



SDS (Na-dodecylsulfate) -> charge

stacking of proteins between Cl⁻ and Gly

Solutions for 2 Laemmli gels

		7 %	10 %	12.5 %
Separating gel	4x Lower Tris	3.7 ml	3.7 ml	3.7 ml
	40% Acrylamide Solution	2.6 ml	3.8 ml	4.7 ml
	H ₂ O	8.7 ml	7.5 ml	6.6 ml
	APS	75 µl	75 µl	75 µl
	TEMED	7.5 µl	7.5 µl	7.5 µl
Stacking gel	4x Upper Tris	1.25 ml	1.25 ml	1.25 ml
	40% Acrylamide Solution	0.5 ml	0.5 ml	0.5 ml
	H ₂ O	3.25 ml	3.25 ml	3.25 ml
	APS	75 µl	75 µl	75 µl
	TEMED	7.5 µl	7.5 µl	7.5 µl

Solutions for SDS-PAGE:

Ammoniumperoxodisulfate (= APS): 10%

4x Lower Tris (buffer for separating gel): 1.5 M Tris/Cl pH 8.8 + 0.4% SDS

4x Upper Tris (buffer for upper gel): 0.5 M Tris/Cl pH 6.8 + 0.4% SDS

10x Laemmli running buffer (1 l):

Tris 30.25 g

Glycine 144 g

SDS 10 g

2x concentrated sample buffer (=SB) (40 ml):

1 M Tris/Cl pH 6.8 5 ml

β-Mercaptoethanol 4 ml

SDS 1.84 g

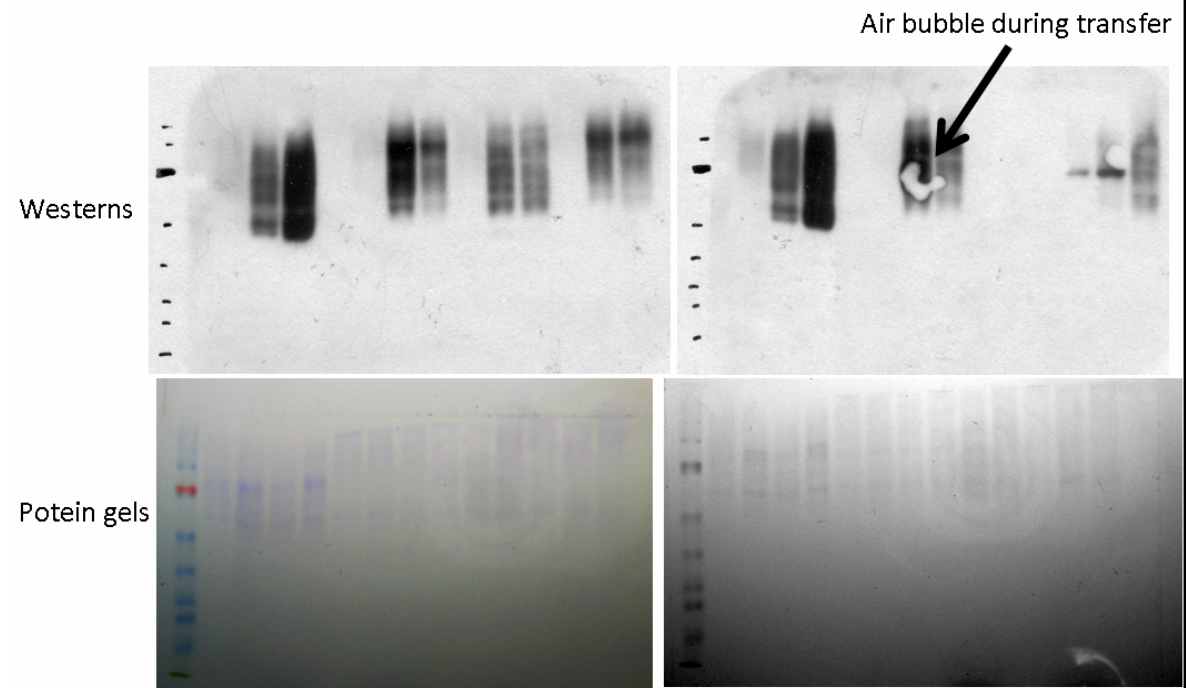
Glycerol 8 ml (heat in Microwave)

Bromphenol blue (a few grains until colored)

H₂O 23.2 ml

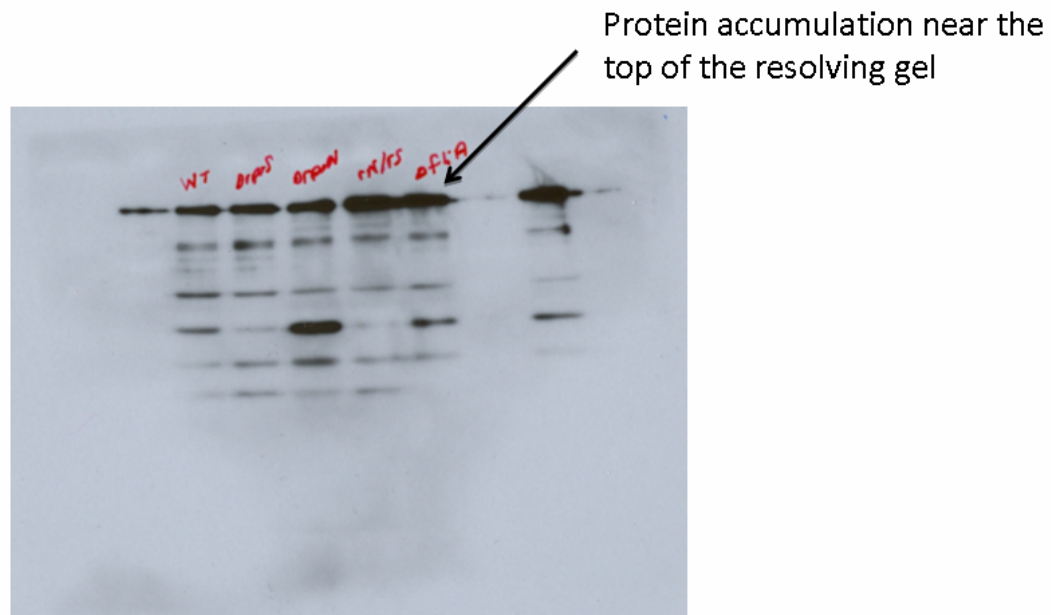
Bad SDS-PAGE gel examples

SDS-PAGE gels run without SDS in running buffer



Bad SDS-PAGE gel examples

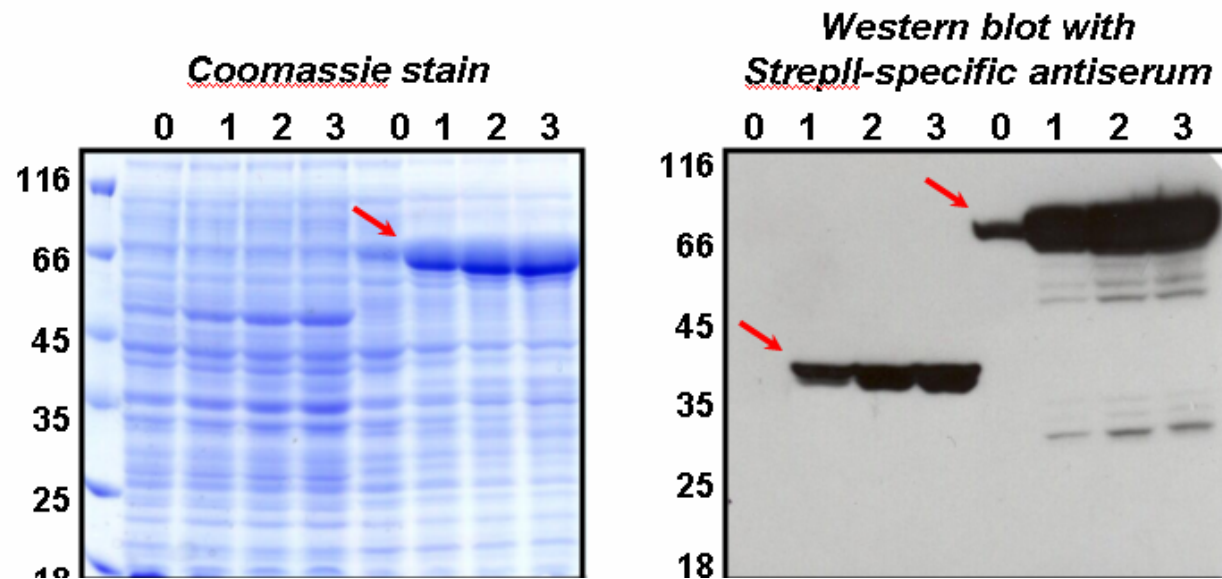
SDS-PAGE GEL: WESTERN



SDS-PAGE: Staining and de-staining

Staining solution: 30% MeOH, 10% Acetic Acid, 0.1% Coomassie Blue R250

Destaining solution: 20% MeOH, 8% Acetic Acid

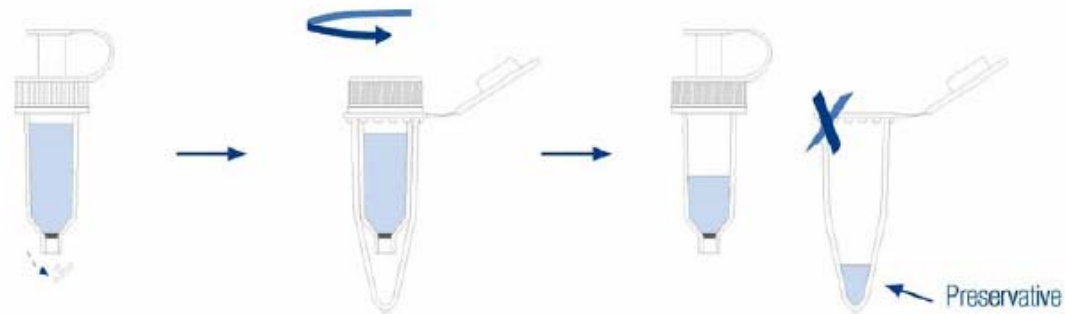


His-Tag purification-Steps

1. Elimination of the Preservative

Remove the lower cap of the Spin-Column, place in a microtube and centrifuge, then discard the preservative residue collected in the tube.

Note: In all centrifugation processes carried out in the procedure, normally a mild centrifugation (1,000 – 1,500 rpm) is sufficient.

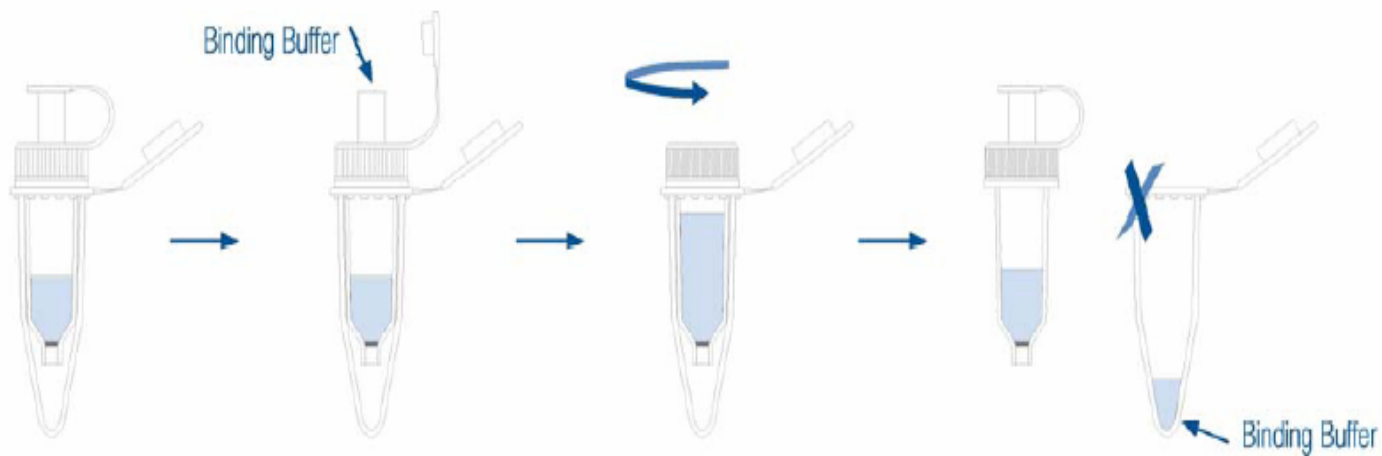


His-Tag purification-Steps

2. Equilibration of the Spin-Column

Introduce the Spin-Column in a microcentrifuge tube and add binding buffer through the top. Centrifuge and discard the residue obtained.

Note: As binding buffer, generally 20 mM disodium phosphate, 500 mM NaCl, 10 mM imidazole pH 7.5 is used.



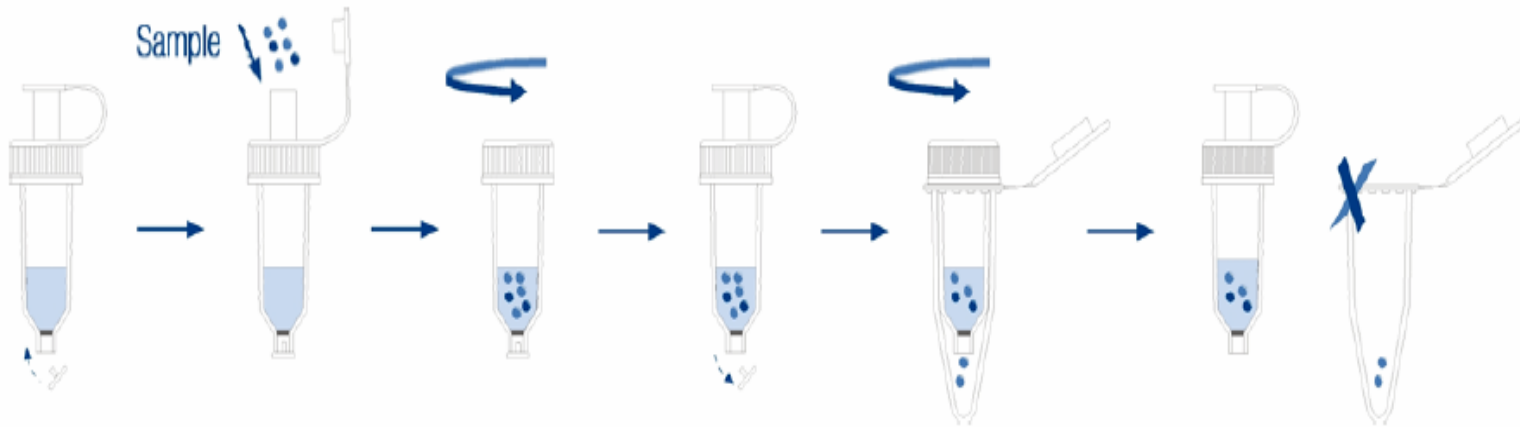
His-Tag purification-Steps

3. Application of the Sample

Add the sample containing the histidine-tagged protein keeping the lower cap in its place.

Manually shake the Spin-Column to maximize contact between the resin and the target-protein.

Remove lower cap, introduce the Spin-Column in a microcentrifuge tube and centrifuge (thus eliminating the proteins not retained in the column).

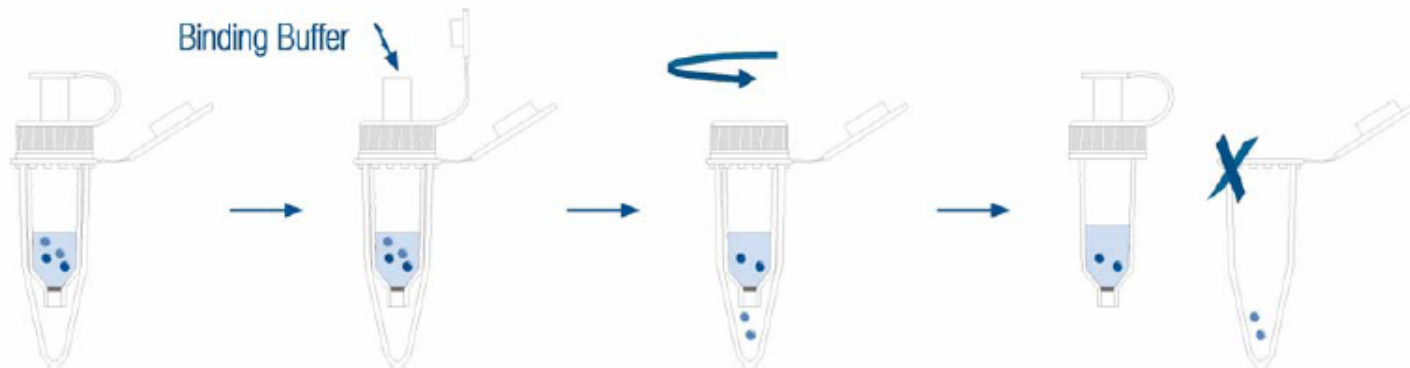


ABT Manual

His-Tag purification-Steps

4. Washing of the Spin-Column

Introduce the Spin-Column in a microcentrifuge tube and add the binding buffer through the top. Centrifuge and discard the residue gathered in the tube.



His-Tag purification-Steps

5. Elution of the pure protein

Add the elution buffer with the lower cap of the Spin-Column in place. Manually shake to drive the elution of the target-protein. Remove the lower cap, introduce the Spin-Column in a microcentrifuge tube and centrifuge, finally collecting the pure protein in the tube.

Note: As elution buffer, 20 mM disodium phosphate, 500 mM NaCl, 500 mM imidazole pH 7.5 is generally used. This concentration of imidazole is usually enough to provoke the elution of the target-protein. However if the desired result is not achieved then the concentration may be increased up to 2.0 M.

Note: Other reagents that may be used to elute the protein are histidines and ammonium chloride.