

# Neopro4 Mini Vertical electrophoresis system full set

NB-12-8001



# Neopro Mini Vertical Electrophoresis System full set #Cat: NB-12-8001

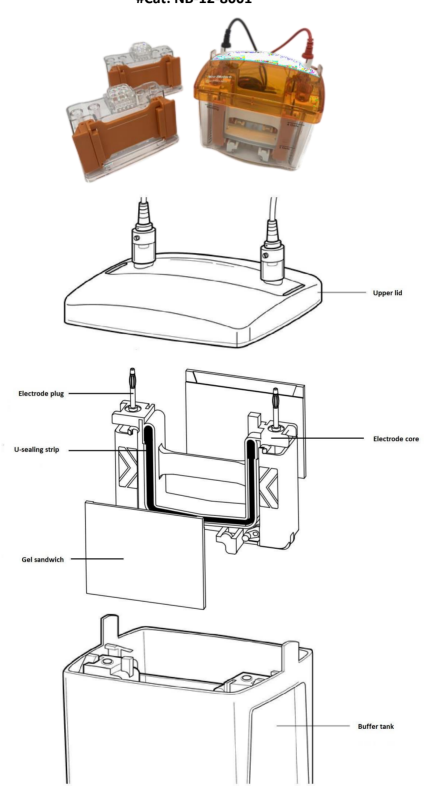


Figure 1: Neo-PRO-Mini electrophoresis installation



# **Chapter 1 Generalization**

#### 1.1 Brief introduction

NeoPRO Mini Vertical Electrophoresis System is used to run pre-casting gel and hand- casting gel, at most 4 pieces of gels are available simultaneously. It is compatible with 1-D and 2-D electrophoresis application. The gel-casting frame and spacer glass plate make the activity of hand-casting gel simple and avoid the leakage.

### 1.2 Component

In order to achieve best performance, please read the instruction manual carefully and operate the instrument strictly according to the instruction manual.

Spacer glass plate: It is the relatively high glass plate with fastening side strip. There are 3kinds of glass plate with the thickness of 0.75mm, 1.0mm, 1.5mm respectively.

Short glass plate: It is the relatively short glass plate and spacer glass plate to form the gelsandwich clamp.

Gel-casting frame: It is put on the desk and align the short glass plate with spacer glass plate, and make sure to form the gel sandwich clamp.

Components of gel clamp: 1 piece of gel casting frame, a piece of spacer glass plate and apiece of short glass plate.

Gel base: Pressure leverage grip the components of gel to sealing gasket closely to avoid the leakage of gel.

Gel sandwich: It consists of short glass plate and spacer glass plate as well as polymer gel.

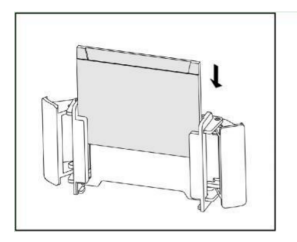
Replacement plate for single gel: Transparent mound pressing baffle, it is used to cast 1 pieceof gel or 3 pieces of gels.

Electrophoresis core: It is used to hold the gel sandwich and supply the U-sealing strip and upper and lower electrode as well as connecting plug. Anode is marked by red while the cathode is marked by black.

#### 1.3 Technical specification

Buffer tank and upper lid: The buffer tank and the upper lid are closed to make sure the electrophoresis is conducted smoothly. The power is off when the upper lid is opened. The tank and upper lid are compatible with other electrophoresis module, such as the electrophoresis transfer, 2-D single direction electrophoresis and electrophoresis elution etc.





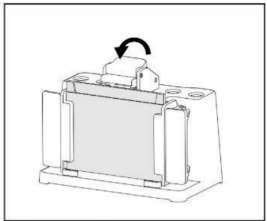


Figure 2: Installation of gel casting frame and gel base

# Maximum volume of sample

	Width			
Hole	ofhole	0.75 mm	1.0 mm	1.5 mm
quantity				
5	12.7 mm	70 μΙ	105 μΙ	160 µl
9	5.08 mm	33 μΙ	44 µl	66 µl
10	5.08 mm	33 μΙ	44 µl	66 µl
15	3.35 mm	20 μΙ	26 μΙ	40 µl
IPG	6.2 mm		420 µl	730 µl
Prep/2-D				
Reference we	ell 3.1 mm	13 μΙ	17 μl	30 µl
Sample well	71.7 mm	310 µl	400 μl	680 µl

# Compatibility of chemical reagent:

All the components of NeoPRO Mini Vertical Electrophoresis System should be kept clear of acetone and ethyl alcohol. The damage caused by using the organic reagent is not covered by the guarantee policy.

The comb of NeoPRO Mini Vertical Electrophoresis System should be kept clear of 100%TEMED, whose structure will be damaged by long time friction of TEMED.

# 1.4 Safety instruction

The power is off when the upper lid is opened, never try to operate the electrophoresis when there is no upper lid.

Note: The safety standard is applied to the Neo Biotech product from design stage to production stage, and it would be safe in case of proper and right operation. Please do not improve or mend it by any means.



# **Chapter 2 Installation and basic operation**

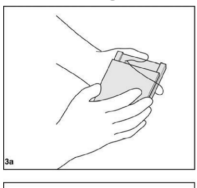
# 2.1 Preparing gel plate

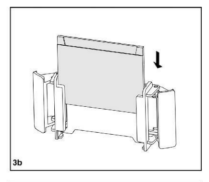
#### Hand casting gel

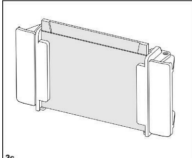
- 1. Note to installation of glass plate clamp and gel casting frame.
- a. Put the gel casting frame vertically on the horizontal desk and leave the hinge of gel castingframe open.
- b. Choose the spacer glass plate according to the thickness of gel and put the short glass plateabove (Please refer to the figure 3a).
- c. Keep the marked tip of spacer glass plate up, slide 2 pieces of glass plates into the gel castingframe, make the face of short glass plate forward (Please refer to the figure 3b).

Note: Keep the 2 pieces of glass plates on the same level and right direction of marker. Theleakage occurs in case of wrong direction of glass plate and non-aligning.

- d. Close the hinge of gel casting frame if the glass plate is in the right position, clamp the glass plate tightly in gel casting frame (Please refer to figure 3c). Check the bottom of glass plate is level.
- e. Keep the hinge of gel casting frame outward and put the gel casting frame on the gel gasket of gel base, then put the spring leverage on the spacer glass plate (Please refer to figure 3d).
- f. Repeat step a-e to make another gel.







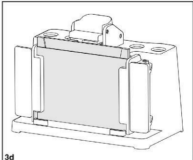


Figure 3: Installation of gel casting frame and gel base



#### **Gel casting**

# a. Non continued polyacrylamide gel

- 1. Put the comb into the gel clamp completely and mark the glass plate 1 cm blow the lower end of tooth. The mark indicates the height of separated gel.
- 2. Mix all the reagent except APS and TEMED to make gel monomer solution, vacuum degassing for more than 15 minutes.
- 3. Put APS and TEMED in monomer solution after being vacuum degassed, and use the pipette or sample adding pipette to inject the solution between glass plate until reach the marker. Keep stable injection in order to avoid mixing with air.
- 4. Cover the surface of solution by water or tert-amyl alcohol (2- methyl, 2- butanol). Note: Pleaseinject the water slowly and stably in order to avoid mixing with solution.
- 5. Leave the gel from 45 minutes to 60 minutes until it is polymerization. Cleanse the surface of gel completely by double distilled water. Do not let the matters of alcohol kind stay on the gel more than 1 hour in order to avoid the dehydration of upper part of gel.

Note: Such separated gel can be store in room temperature overnight. Add 5 ml of 1.5MTri-HClwith ratio of 1 to 4 and buffer with pH8.8 (Laemmli system) to the separated gel to avoid being dried. In case of other buffer, please add 5 ml of 1 fold diluted solution.

- 6. Prepare spacer gel monomer solution, mix all the reagents except APS and TEMED, vacuum degassing for more than 15 minutes.
- 7. Use the filter paper to dry the surface of separated gel before injecting spacer gel.
- 8. Add APS and TEMED in the degassed spacer gel monomer solution and inject the solutionuntil it reaches the level the glass plate.
- 9. Insert comb and make sure that the back of comb and the glass plate are on the same level.
- 10. Leave the gel from 45 minutes to 60 minutes until it is polymerization.
- 11. Take out the comb slightly and cleanse the surface of gel by distilled water and buffer.
- 12. Cleanse the used gel clamp frame and gel casting frame by distilled water and ion-removingwater.

## b. Continued polyacrylamide gel

- 1. Mix all the reagent except APS and TEMED to make gel monomer solution (Please refer to gellist of chapter 4), vacuum degassing for more than 15 minutes.
- 2. Put APS and TEMED in monomer solution after being vacuum degassed, and inject the solution until it reaches the level the glass plate.
- 3. Insert comb and make sure that the back of comb and the glass plate are on the same level.
- 4. Leave the gel from 45 minutes to 60 minutes until it is polymerization.
- 5. Take out the comb slightly and cleanse the surface of gel by distilled water and buffer. And cleanse the used gel casting frame and gel base by distilled water and ion-removing water.

#### 2.2 Installation of electrophoresis module and adding sample

Clean and dry Neo-PRO-Mini electrophoresis buffer tank.

Electrophoresis module (Electrophoresis core can only be used for 1 piece gel or 2 pieces of gels, the shared module should be used in case of 3 pieces of gels or 4 pieces of gels.



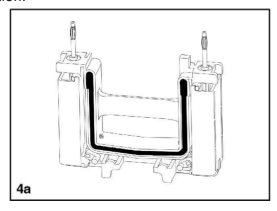
## 1. Installation

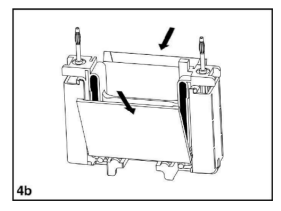
Note: If only run 2 pieces of gels, please use electrode head electrophoresis core. If only run 4pieces of gels, please use the electrode head electrophoresis core and mushroom head electrophoresis core, each module for 2 pieces of gels.

- a. Put the opened gel-casting frame on the clean and horizontal surface of desk (Please refer to the figure 4a).
- b. Adjust the short glass plate inwardly, put the first gel sandwich on the gel supporter, there are 2gel supporters on both two sides of bottom of module, now there is a 30 degree angle between gelplate and module. Be careful to put the first gel and keep balance, then put the second gel on the other side of gel supporter. There are totally 2 pieces of gels slant from the center (Please refer to the figure 4b). Note: Be sure to adjust the short glass plate inwardly and then put the gels on the two sides of gel clamping frame. The gel clamping frame requires 2 pieces of gels to form the functional module. In case of running odd number of gel (1 piece of gel or 3 pieces of gels), please use single gel plate (Please refer to figure 4b).
- c. Push 2 pieces of gel plates slightly to the center, close to U-sealing Strip, to make sure that the short glass plate is under the concave tank of U-sealing Strip.
- d. Compress the gel plate tightly by one hand, while close the red gel clamp frame by another hand and make it in the best position. Or hold the whole unit and stabilize the gel plate and close gel clamp frame on both two sides in order to make it be locked in the right position (Please refer to the figure 4c). The gel clamp frame pushes the gel plate to make it and concave tank of red gelgasket matched together tightly in order to avoid the leakage of liquid (Be sure the short glassplate is under the bottom of concave tank of U-sealing Strip). Thus, use the buffer to cleanse the sample and start to add the sample (Please refer to the figure 4b).

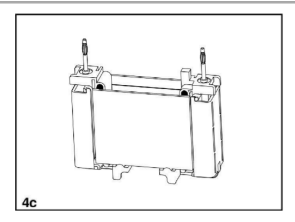
Important note: Never try to close the gel clamp frame when the gel plate is not under the bottom of concave tank of U-sealing Strip. In order to avoid the movement of gel plate in the process of being locked, please use single hand to compress the gel plate on the two sides of gel clamp frame stably and evenly.

**Note:** Please do not put mushroom head electrophoresis core in the electrophoresis tank when 1-2 gels are running, otherwise the surplus heat is generated which affects the result of electrophoresis separation.









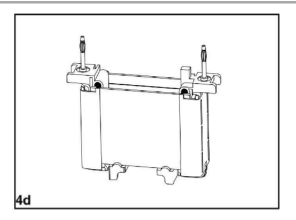


Figure 4: Installation of Neo-PRO-Mini electrophoresis module

# 2. Sample adding

- a. Inject the buffer into the tank, from the outside of outer tank and follow lower part of glass plate, just flood the short glass plate slightly.
- b. Add the sample before or after the electrophoresis core putting in the electrophoresis tank, bothof two methods are available to obtain satisfied result.
- c. Add the sample into the hole by syringe and sample adding pipette.

Note: Make the sample fall onto the bottom of hole slowly and evenly. Be sure not to puncture thebottom. Note: Anode and cathode should be flooded by the buffer and on the same level.

3. Put the electrophoresis module in the buffer tank.

Note: The volume requires, 2 pieces of gels with 700ml, 4 pieces of gels with 1000ml.

There are two places of buffer tank for two modules: Electrode head electrophoresis core is put infront of the mushroom head electrophoresis core.

- a. Put the Neo-PRO-Mini buffer tank on the flat table, make the front side (The side indicating thewords of 2 gels and 4 gels) forward. If the direction is correct, the red mark of edge of tank shouldbe on the right side and the black left.
- b. If running 2 pieces of gels, please use the electrophoresis core with tall leads. Put it on the backside of the tank and make the red (+) corresponding to the red mark on the right side of tank.
- c. If running 4 pieces of gels, please use not only the electrophoresis core with tall leads but also the mushroom head electrophoresis core that should put in the front side of the tank. Make the red(+) of the two corresponding to the red mark on the right side of tank. Note: The wrong direction and position will make the upper lid impossible to be closed.
- d. Pour the buffer into the tank until reach the marker.
- 4. Neo-PRO-Mini installation of buffer tank

Put the upper lid on the buffer tank, and make sure the plug into socket, the protruded partof upper lid can help avoid the error. Note: The two protruded parts of buffer tank should go through the slot of upper lid in order to make it open and close smoothly. At this time, please compress the upper lid continuously with thumb until is done.



#### 5. Power condition

- a. Insert the tall leads into the socket of power supply correctly.
- b. Power on Neo-PRO-Mini, the constant voltage is 200V is recommended for SDS-PAGE and most of native PAGE. 200V voltage is used to run 2 pieces of gels and 4 pieces of gels. The customer choose the voltage according to the real situation. SDS-PAGE needs around 35 minutesunder the voltage of 200V.
- 6. Extract gel
- a. Power off power supply and disconnect tall leads when the elextrophoresis is done.
- b. Lift the upper lid, take out the electrophoresis core carefully, and pour out the buffer. In orderto prevent buffer leakage, please pour out the buffer before opening the clamp.
- c. Take out the gel plate.
- d. Separate the two glass plates carefully and take out the gel.
- e. Put the gel downward and immerse the gel and glass plate in the buffer to make them separated.
- f. Cleanse the Neo-PRO-Mini electrophoresis core and buffer tank by the ion-removing distilledwater.



# **Chapter 3 Trouble shooting**

Problem	Cause	Solution
Smile effect – band pattern curvesupward at both sides of the gel	<ul> <li>Center of the gel running hotter than either end</li> <li>Power conditions excessive</li> </ul>	<ul> <li>Buffer not mixed well or buffer in upper chamber too concentrated. Remake buffer, ensuring thorough mixing, especially when diluting 5x or 10x stock.</li> <li>Decrease the power setting from 200 V to 150 V or fill lower chamber to within 1 cm of top of short plate.</li> </ul>
Vertical streaking of protein	<ul><li>Sample overloaded</li><li>Sample precipitation</li></ul>	<ul> <li>Dilute sample, selectively remove predominant protein in sample, or reduce the voltage about 25% to minimize streaking.</li> <li>Centrifuge sample before addition of SDS sample buffer, or decrease %T of the gel*</li> <li>The ratio of SDS to protein should be enough to coat each protein molecule with SDS, generally 1.4:1.</li> <li>It may require more SDS for some membrane protein samples.</li> </ul>
Lateral band spreading	<ul> <li>Diffusion of the wells prior to turning on the current.</li> <li>lonic strength of the sample lower than that of the gel.</li> </ul>	<ul> <li>Minimize the time between sample application and turning on the power start-up.</li> <li>Use same buffer in sample as in the gel or the stacking gel.</li> </ul>
Skewed or distorted band	<ul> <li>◆ Poor polymerization around wells</li> <li>◆ Salts in sample</li> <li>◆ Uneven gel interface</li> </ul>	<ul> <li>Degas stacking gel solution completely prior to casting; +C2:C5 increase ammonium persulfate and TEMED concentrations by 25%, for stacking gel or low %T, leave APS the same and double the TEMED concentration.</li> <li>Remove the salts by dialysis, desalting, column, etc.</li> <li>Decrease the polymerization rate.</li> <li>Overlay gels very carefully.</li> </ul>



Problem	Cause	Solution
Lanes constricted at the bottom of the gel	<ul> <li>Ionic strength of samplehigher than the surroundinggel</li> </ul>	Desalt sample and neighboringsamples.
Run taking unusually long	<ul><li>Running buffer too concentrated</li><li>Excessive salt in sample</li></ul>	<ul><li>Check buffer protocol, dilute if necessary</li><li>Desalt sample</li></ul>
Run too fast	<ul><li>Running or reservoir buffertoo dilute</li><li>Voltage too high</li></ul>	<ul> <li>◆ Check buffer protocol, dilute if necessary</li> <li>◆ Decrease voltage by 25–58%</li> </ul>
Doublets observed where single proteinspecies is expected (SDS-PAGE)	◆ A portion of the protein may have been re-oxidized during the run or may not have been fully reduced prior to the run	<ul> <li>Prepare fresh sample buffer solution if over 30 days old; Increase concentration in the sample buffer;Substitute DTT for BME</li> </ul>
Fewer bands than expected and oneheavy band at the dry front	<ul><li>Protein(s) migrating at the dye front</li><li>Protein degradation</li></ul>	<ul> <li>Increase the %T of the resolving gel</li> <li>Use protease inhibitors, e.g.,</li> <li>PMSF,etc</li> </ul>
Fewer bands than expected and oneheavy band at the dry front	<ul> <li>Protein(s) migrating at thedye front</li> <li>Protein degradation</li> </ul>	<ul> <li>Increase the %T of the resolving gel*</li> <li>Use protease inhibitors, e.g., PMSF,etc.</li> </ul>
Upper buffer chamber leaks	<ul><li>Upper buffer chamber overfilled</li><li>Improper assembly</li></ul>	<ul> <li>Keep buffer level below the top of the spacer plate</li> <li>Be sure U-shaped electrode core gasket is clean, free of cuts, and lubricated with buffer</li> <li>Be sure short plate is under the notchon the gasket, not on top of it</li> </ul>
Leaking during hand casting	<ul> <li>Chipped glass plates</li> <li>Spacer plate and short plate not level</li> <li>Casting stand gasket is dirty, flawed, or worn out.</li> </ul>	<ul> <li>Ensure glass plates are free of flaws</li> <li>Ensure plates are aligned correctly</li> <li>Wash the gasket if it is dirty,</li> <li>replace casting stand gaskets if</li> <li>flawed orworn out</li> </ul>



Problem	Cause	Solution
Poor end well formation	<ul> <li>Incorrect catalyst formation</li> <li>Monomer solution not degassed. Oxygen inhibits polymerization</li> </ul>	<ul> <li>Prepare fresh catalyst solution, or increase the catalyst concentration of the stacking gel to 0.06% APS and 0.12% TEMED</li> <li>Degas monomer solution immediately prior to casting the stacking gel</li> </ul>
Webbing/excess acrylamide behind thecomb	◆ Incorrect catalyst concentration	<ul> <li>Prepare fresh catalyst solution, or increase the catalyst concentration of the stacking gel to 0.06% APS and 0.12% TEMED</li> </ul>
The pressure cams on the casting frame are difficult to close or make noise when closed	Powder residue has built up at the pivot of the pressure cams	Rinse or wipe off the powder residuebefore each use

# **Chapter 4 Quality guarantee**

- (1) The warranty is 2 years since the date of sales.
- (2) The warranty excludes the following situations otherwise it is charged.
  - a. No presentation of warranty card and invoice.
  - b. The invoice is revised.
  - c. Improper operation or accident factors.
  - d. The damage is caused by the user's repair.
  - e. Out of the warranty, the instrument is still in usage after repair.