

Q-PAGETM Precast Gel

Quick & Quality

Gel Type: TGN Gel	Cassette size: Mi	4-15%	
% of Gel Well No.	10%		
12 wells	QP4210	QP4510	
15 wells	QP4220	QP4520	

Storage and stability- Store Q-PAGE™ Precast Gels at 4°C for periods up to 12 months. Do not freeze Q-PAGE™ Precast Gels. Remove tape and comb before electrophoresis.

Description

Q-PAGE™ TGN (Tris-Glycine Novel) Precast Gels are ready-to-use acrylamide gels for SDS-PAGE running in Tris-Glycine buffer system. unique formula, Q-PAGE™ TGN Precast perform enhanced speed, better separation, and longer shelf life as compared with conventional Laemmli Tris-HCl gels. The protein migration patterns in Q-PAGE™ TGN series, however, are similar with typical Laemmli Tris-HCl gels, and thus Q-PAGE™ TGN Precast Gels are compatible to traditional SDS-PAGE and subsequent analyses.

Q-PAGE™ TGN Precast Gels are available in gradient (4 to 15%) and fixed (10%) concentrations of polyacrylamide in 12- and 15-well formats. Two available cassette sizes, Mini (10 x 8.3 cm) and Midi (10 x 10 cm), are compatible with most popular protein electrophoresis systems. Q-PAGE™ Mini (QP4XXX) Gels are suitable for Bio-Rad® and other systems. Q-PAGE™ Midi (QP5XXX) Gels are suitable for Invitrogen® XCell SureLock® Mini-Cell, Invitrogen® Mini Gel Tank, Hoefer SE260, and other systems.

Key Features:

User-friendly gel cassette:

Numbered and framed wells for sample loading Labeled warning sign and green tape as reminder

Enhanced gel performance:

Enhanced gel electrophoresis speed

Better band separation

Stable for shipping at ambient temperature

Easy compatibility:

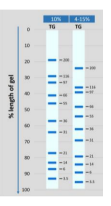
Available as homogeneous and adjusted gradient gels for a wide range of protein separation.

Compatible with most popular protein electrophoresis systems

Recommendations/Tips for Gel Running

- 1. Remove comb and tape before adaption.
- 2. Use fresh 1X running buffer for the inner cathode chamber.
- 3. Rinse the wells before sample loading.
- 4. Try 200 V first, and optimize the voltage and running time if needed. Do not set voltage lower than 100 V.

O-PAGE™ TGN Gel Migration Charts





ROTH					
		(QP4210) (QP4210)			
		(QP4220) (QP4220)			
		(QP4510) (QP4510)			
	_	(QP4520)			

Bands correspond to the migration of Mark12 Unstained Standard.

Procedures for Using Q-PAGE™ TGN Precast Gel

Sample Preparation for SDS-PAGE

- Mix protein sample with 2X sample buffer.
- Heat the diluted samples at 95°C for 5 min or at 70°C for 10
- Cool the diluted samples to 4°C and spin down the water condensed on tube surface. (If there is high viscosity part at bottom of tube, transfer supernatant to a new tube.)

Prepare Q-PAGE™ for Sample Loadina

- Open the blister tray of Q-PAGE™ Precast Gel.
- Briefly rinse the gel cassette with ddH₂O.
- Remove tape and comb; avoid squeezing the gel.
- Adapt Q-PAGE™ to electrophoresis system; instruction is provided below. (BioRad Mini-PROTEAN® Core Electrophoresis System is recommended.)
- Use a pipette to gently wash the wells with running buffer to remove residual storage buffer.
- 6. Fill the wells with running buffer prior to sample loading.
- 7. Load samples and pre-stained protein marker into numbered
- 8. Fill both inner and outer chambers with running buffer to the highest level. Ensure gel wells are completely covered.

Power Settina for Runnina Q-PAGE™

Optimize the voltage and running time if needed.

Voltage*1	150 V	200 V*2	250 V*3	300 V*3
Running Time*4	40-60 mins	30-40 mins	25-35 mins	15-25 mins
Expected Current				
Initial (per gel)	40-50 mA	50-60 mA	80-90 mA	90-100 mA
Final (per gel)	10-20 mA	25-30 mA	35-40 mA	40-50 mA
Expected temperature	25-30°C	25-40 °C	25-40°C	25-40°C

^{*1} Set voltage higher than 100 V is recommended.

^{*2} Try 200 V first, and optimize the voltage and running time if needed.

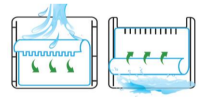
^{*3} For higher voltage conditions, please use fresh running buffer for inner and outer chambers

⁴ Running time varies depending on gel percentage, running buffer, temperature, and power supply.

Remove Q-PAGE™ Gel from Cassette

Open cassette immediately after electrophoresis. Avoid ael drvina.

- 1. Insert the cassette opener into corners of cassette.
- 2. Sequentially pry the opener to separate the two plates.
- 3. Gently pull two plates apart from the top of cassette.
- 4. Carefully detach the gel either from the bottom or the top side of the cassette.
 - -Avoid diagonally peeling the gel from the corner.
 - -Use water to help ael detachment if needed.



5. Gently remove the gel for further staining or Western blotting.

Gel Stainina

Proteins separated using Q-PAGE™ Precast Gels can be further stained with most popular staining reagents, such as Coomassie dyes (R-250 or G-250), Silver-stain solution, and FluoroStain™ Protein Fluorescent Staining Dye. (Cat. No. PS1000)

Transferring Protein from Q-PAGE™ to Blotting Membrane

- After protein separation using Q-PAGE™, gently detach Q-PAGE™ from cassette and then equilibrate the gel in transfer buffer.
- Pre-soak blotting membrane and filter papers in transfer buffer.
 *Activate PVDF membrane in methanol before soaking in transfer buffer.
 - **Prepare 6 filter papers for one gel/membrane sandwich.
- Assemble transfer sandwich by orientating cathode, sponge, filter papers, gel, membrane, filter papers, sponge, and anode. The protein goes to the direction of cathode to anode.
- Carefully move roller over the gel/membrane to remove air bubbles and excess buffer until complete contact is established.
- 5. Insert transfer cassette into transfer module. Notice that black side of cassette should be next to black side of module.
- Fill transfer tank with pre-cooled transfer buffer to the highest water level.
- Set constant voltage at 100 V. Transfer for 90 minutes at low temperature condition. Pre-stained protein marker should be visible on the membrane after transfer is completed. Transfer of proteins to the membrane can be checked using Ponceau S staining before blocking step.

Supplemental Information for Using Q-PAGE™ Precast Gel Adapting Q-PAGE™ Mini Precast Gel to BioRad Mini-PROTEAN® Core

 After removing comb and tape, place the Q-PAGE™ Mini Precast Gel with notched plate facing toward inner chamber.

- Align the notched plate to ensure the edge sits just below the notch at the top of green gasket.
- Gently press gel cassette toward green gasket and then lock gel cassette with two green arms. Avoid squeezing the cassette and gel.
- Fill inner chamber with running buffer to check tightness of seal.
 If necessary, reassemble and check the seal again.
- Fill inner chamber with running buffer to ensure gel wells are completely covered.
- 6. Fill outer chamber with running buffer to the highest level.

Adapting Q-PAGE™ Mini Precast Gels to other electrophoresis system, please follow the manufacturer's instruction.

Buffer recipes

2X sample buffer with reducing agent

62.5 mM Tris-HCl pH 6.8, 2% SDS, 25% (v/v) glycerol, 0.01% bromophenol blue, 5% β -mercaptoethanol or 100 mM DTT (added fresh)

10X Tris-Glycine running buffer

30.0 g Tris base, 144.0 g Glycine, 10.0 g SDS. Bring up the volume to 1 L with ddH₂O.

1X running buffer

Dilute 100 ml 10X running buffer with 900 ml ddH₂O.

10X transfer buffer

 $30.0\,g$ Tris base, 144.0 g Glycine. Bring up the volume to 1 L with ddH2O.

1X transfer buffer

*Cool 1X transfer buffer to 4°C before using.

Dilute 100 ml 10X transfer buffer with 200 ml methanol and 700 ml ddH₂O.

**Add SDS to 0.1% to promote transfer of high molecular weight proteins.

Related Products: Q-PAGE™ Precast Gel

Туре	TGN			
Cassette	Mini		Midi	
Well No.	12 well	15 well	12 well	15 well
10%	QP4210	QP4220	QP5210	QP5220
4-15%	QP4510	QP4520	QP5510	QP5520

Туре	Bis-Tris			
Cassette	Mini		Midi	
Well No.	12 well	15 well	12 well	15 well
8%	QP2110	QP2120	QP3110	QP3120
12%	QP2310	QP2320	QP3310	QP3320
4-12%	QP2510	QP2520	QP3510	QP3520

More information can be found on website: (Procedures and Troubleshooting)

