

# INSTRUCTION MANUAL

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## **SERVA Ge/™ Native PAGE Starter Kit** Precast Vertical Gels for Electrophoresis

(Kat.-Nr. 43201.01)



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# 1. SERVAGel™ Native PAGE Starter Kit

## 1.1. General information

SERVAGel™ Native PAGE Starter Kit contains in addition to ready-to-use Tris/Glycine gels electrophoresis running and sample buffer for performing native PAGE. The included SERVAGel™ TG gels are designed for vertical slab gel electrophoresis and suited for discontinuous separation according to Laemmli (Nature 277, 680 [1970]). They do not contain SDS and thus, can be operated with native buffers. SERVA offers homogenous or gradient precast gels featuring various acrylamide concentrations (T).

Benefits of the product for the user:

- simple, fast handling
- high resolution, sharp bands, best reproducibility
- made from top-quality chemicals
- gels prepared in unbreakable plastic cassette, leakage-free
- long separation distance, cm-scale at front of cassette allows reproducible runs
- marking of anode and cathode for error-free assignment
- extra tool provided for easy and safe opening of cassette at the end of run
- compatible with many commercially available electrophoresis tanks (e.g. SERVA BlueVertical 102, Hoefer Mighty Small™ SE 260, Hoefer miniVE™, NOVEX XCell II®, etc.)

The precast gels are manufactured according to proprietary methods developed by SERVA Electrophoresis GmbH and are subject to strict quality control. Each production batch has assigned a unique lot number. In the event of queries, please quote this lot number along with the catalogue number.

## 1.2. Kit components

SERVAGel™ Tris/Glycine gels (Cat. No. selectable)	4 pieces
Tool for opening of cassette	1 piece
10x Towbin running buffer (Cat. No. 42558)	400 ml
2x Tris/Glycine native sample buffer (Cat. No. 42528)	1 ml

Each gel is packed individually sealed in an aluthene bag. It is protected from desiccation by a layer of filter paper moistened with gel buffer.

**Cassette :**

Outer dimensions	10 cm x 10 cm
Number of sample wells	12
Volume of well	35 µl

**Gel:**

Material	Acrylamide/N, N'-methylene bisacrylamide
Dimensions separation gel	Length 7 cm x width 8 cm
Thickness of gel layer	1 mm

**Note:**

Reagents needed for the subsequently staining of the gels are not included in the kit and must be ordered separately. A detailed Western Blotting protocol for the SERVAGe™ TG gels is available on request:

Phone: +49-6221-1384044, E-mail: tech.service@serva.de

**1.3. Composition of gels**

SERVAGe™ TG gels are offered at various acrylamide concentrations (T). The gels contain **no SDS**. Depending on the electrophoresis buffer used, you determine whether native or denaturing conditions prevail. The separation ranges of gels for **denatured** proteins are shown in table 3.1.(p. 20)

**Acrylamide concentration (T):** 8 %, 10 %, 12 %, 14 %, 16 %, 4 – 12 %, 8 – 16 %, 4 – 20 %

**Cross linker concentration (C):** 2.6 %

**Stacking gel:** 4 % T, 2.6 % C

**Gel buffer:**

Stacking gel 125 mM Tris/HCl, pH 6.8

Separation gel 375 mM Tris/HCl, pH 8.8

**1.4. Storage conditions**

Kit components	Storage temperature
10x Towbin running buffer	+15 °C - +30 °C
SERVAGe™ TG gels	2 – 8 °C
2x Tris/Glycine native sample buffer	2 – 8 °C

Do **not freeze** the gels **or leave them at room temperature for longer periods** as this may impair their separation properties.

If stored at the recommended temperature at least usable until: see expiry date on package.

## 2. Handling of gel cassettes/electrophoresis procedure

### **Safety information:**

*For safety reasons always wear suitable protective gloves and clothing, when you work with gels and appending solutions.*

1. Remove gels from cardboard box. If only one gel is required, immediately place the remaining gels again to storage at 2 – 8 °C. Cut open aluthene bag along the upper edge using scissors. Remove gel.
2. Place the gel into the electrophoresis chamber so that the opened (“u-shaped”) side of the cassette is facing towards the cathode buffer tank. Follow the manual of your electrophoresis chamber supplier for detailed instructions.
3. Add the electrophoresis buffer. Pull the comb steadily out of the gel; remove eventually remaining gel rests above the sample wells. Rinse the sample wells thoroughly, avoiding and/or removing any air bubbles.
4. Apply samples. Load those sample wells without samples with sample buffer (1x).
5. Close the electrophoresis chamber and connect to power supply. Switch on power supply and begin electrophoresis.  
Conditions: see paragraph 3.
6. On completion of electrophoresis, switch off power supply, disconnect the electrophoresis chamber, remove electrophoresis buffer and remove cassettes.
7. To open cassette hold cassette upright with its bottom end supported by a table or bench. Place the corner of the key marked by an arrow at the upper right-hand end of the grooved edge of the cassette (also marked by an arrow) and break open the cassette with a swift blow from above on the key. Turn around the cassette and open the other side in the same way.
8. To remove the gel, carefully detach the plates so that the gel remains on one. Gels can now be stained or used for blotting.

Reagents needed for the subsequently staining of the gels are not included in the kit and must be ordered separately. A detailed Western Blotting protocol for the SERVAGE™ TG gels is available on request:

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### 3. Electrophoresis protocols

**Note:** The specified separation range of gels is determined by SDS PAGE.

#### 3.1. Separation range of gels

Acrylamide concentration (%)	Separation range (Mr 10 <sup>3</sup> )
8	40 - 250
10	30 - 200
12	20 - 200
14	10 - 100
16	5 - 70
4 - 12	30 - 300
8 - 16	20 - 250
4 - 20	6 - 200

#### 3.2. Running buffer preparation

Dilute 10x Towbin running buffer 1:10 (composition see appendix, page 23).

#### 3.3. Sample preparation

- Mix your sample with the same volume **2x Tris/Glycine native sample buffer** (composition see appendix, page 23). The maximal well volume is 35 µl. **Do not heat sample !**
- Rinse wells with running buffer.
- Load samples and start electrophoresis.

##### 3.3.1. Recommended sample quantity

Amount/band	Staining method	SERVA product
0.1 – 0.5 µg protein	SERVA Blue, Coomassie <sup>®</sup> Brilliant Blue	DensiStain BlueG Soln., SERVA Blue R Staining Kit
10 - 50 ng protein	Silver staining	Silver Staining Kit Native PAGE

### 3.4. Electrophoresis conditions

Electrophoresis is carried out under the following conditions:

**Limiting voltage:** 130 V

Amperage will decrease during run from initial ca. 15 mA/gel to ca. 5 mA/gel.

**Duration:** depending on sample, one hour to several hours. No standard protocol available, has to be defined by the user.

## 4. Staining

### **Safety information:**

*For safety reasons, always wear protective gloves and clothing, when working with fixing and staining solutions.*

For best results use user-friendly staining kits from SERVA like SERVA DensiStain Blue G Staining Solution (Cat. No. 35078.01), SERVA Blue R Staining Kit (Cat. No. 42531.01) or SERVA Silver Staining Kit Native PAGE (Cat. No. 35077.01).

You can also use other common staining protocols as e.g. the protocol described in paragraph 4.1:

### 4.1. Staining with SERVA Blue R

#### 4.1.1. Reagents and solutions

<b>Fixative</b>	20 % (w/v) trichloroacetic acid (Cat. No. 36913)
<b>Stock solution 1</b>	0.2 % SERVA Blue R in 90 % (v/v) ethanol (Cat. No. 11093) (Solve 100 mg SERVA Blue R (Cat. No. 35051) in 50 ml ethanol)
<b>Stock solution 2</b>	20 % (v/v) acetic acid
<b>Destainer</b>	20 % (v/v) ethanol, 5 % (v/v) acetic acid, 1 % (w/v) glycerol (Cat. No. 23176)
<b>Preservation solution</b>	30 % (v/v) ethanol, 5 % (w/v) glycerol

#### 4.1.2 Protocol

Carry out all fixing and staining work on a shaker at moderate speed (50 rev/min). The specified times apply to incubation at room temperature. Shorter staining and destaining times can be achieved by increasing the temperature.

<b>Fixation</b>	Fix gel in 20 % (w/v) trichloroacetic acid for 30 min., wash gel for 1 min. in distilled water before staining.
<b>Staining</b>	<b>Stock solution 1</b> and <b>2</b> are mixed in equal parts and the gel is incubated for 30 min. in the solution. (Staining solution can be re-used for 2 - 3 xs.)
<b>Destainer</b>	Rinse gel after staining for <b>1 minute with dest. water</b> and incubate for <b>2 x 60 minutes</b> in destainer. If background is not clear enough, destain gel for 20 – 30 minutes in 40 % ethanol/10 % acetic acid/2 % glycerol.
<b>Preservation</b>	Incubate gel over night in preservation solution. The gel can then be dried in a drying frame.



## 4. Trouble shooting

<b>Problem</b>	<b>Possible cause</b>	<b>Countermeasure</b>
<b>No current</b>	Unclosed circuit	Check contacts/leads at source of current and separation chamber; check buffer level
<b>Low current</b>	Wrong adjustment of parameters at power source	For limiting amperage select the maximum voltage recommended for the chamber; for limiting voltage select maximum amperage
<b>'Smile effect' at buffer front</b>	Overheating	Pre-cool buffer; cooling via cooling circulator or a reduction in amperage
<b>Slow migration of buffer front</b>	Running buffer fully consumed	Always use fresh running buffer
<b>Blurred bands</b>	Diffusion after application of samples	Apply samples quickly; begin electrophoresis straight away
	Diffusion after separation	Transfer gel to fixing or staining solution immediately after electrophoresis
<b>Irregular bands</b>	Sample volumes too low or too different	Apply at least 5 µl sample; use approx. the same amounts of sample
	Differing saline content of samples	Desalinate samples as required (dialysis, gel filtration)
<b>Formation of stripes</b>	Precipitation of sample	Centrifuge or filter sample
<b>Wide, partially smeared bands</b>	Lipophilic substances in the sample	Remove substances prior to electrophoresis
<b>More bands than expected</b>	Protease activity	Add protease inhibitor; minimise time between sample preparation and run

## 5. Appendix

### Composition of buffers:

#### 10x Towbin running buffer

Components	Concentration
Tris	0.25 M
Glycine	1.92 M

#### 2x Tris/Glycine native sample buffer

Components	Concentration
1 M Tris-HCl pH 6.8	0.126 M
Glycerol	20 %
0.1 % (w/v) Bromophenol blue	0.02 %
Water, deion.	

## 7. Order information

Precast gels	Cat No.
SERVAGel™ TG 8 %Tris-Glycine (10 gels)	43208.01
SERVAGel™ TG 8 %Tris-Glycine ( 6 gels)	43208.02
SERVAGel™ TG 8 %Tris-Glycine ( 2 gels)	43208.03
SERVAGel™ TG 10 % Tris-Glycine (10 gels)	43210.01
SERVAGel™ TG 10 % Tris-Glycine ( 6 gels)	43210.02
SERVAGel™ TG 10 % Tris-Glycine ( 2 gels)	43210.03
SERVAGel™ TG 12 % Tris-Glycine (10 gels)	43212.01
SERVAGel™ TG 12 % Tris-Glycine ( 6 gels)	43212.02
SERVAGel™ TG 12 % Tris-Glycine ( 2 gels)	43212.03
SERVAGel™ TG 14 % Tris-Glycine (10 gels)	43214.01
SERVAGel™ TG 14 % Tris-Glycine ( 6 gels)	43214.02
SERVAGel™ TG 14 % Tris-Glycine ( 2 gels)	43214.03
SERVAGel™ TG 16 % Tris-Glycine (10 gels)	43216.01
SERVAGel™ TG 16 % Tris-Glycine ( 6 gels)	43216.02
SERVAGel™ TG 16 % Tris-Glycine ( 2 gels)	43216.03
SERVAGel™ TG 4 - 20 % Tris-Glycine (10 Fertiggele)	43230.01
SERVAGel™ TG 4 - 20 % Tris-Glycine ( 6 Fertiggele)	43230.02
SERVAGel™ TG 4 - 20 % Tris-Glycine ( 2 Fertiggele)	43230.03
SERVAGel™ TG 8 - 16 % Tris-Glycine (10 Fertiggele)	43231.01
SERVAGel™ TG 8 - 16 % Tris-Glycine ( 6 Fertiggele)	43231.02
SERVAGel™ TG 8 - 16 % Tris-Glycine ( 2 Fertiggele)	43231.03
SERVAGel™ TG 4 - 12 % Tris-Glycine (10 Fertiggele)	43232.01
SERVAGel™ TG 4 - 12 % Tris-Glycine ( 6 Fertiggele)	43232.02
SERVAGel™ TG 4 - 12 % Tris-Glycine ( 2 Fertiggele)	43232.03

<b>Equipment</b>	
BlueVertical mini slab gel system BV 102	BV 102
Blue Power 500 Plus power supply	BP-500Plus
BlueBlot Wet 100 tank blotter (10 x 10 cm)	BB 100
BlueFlash Semi-Dry blotter medium (15 x 15 cm)	BF-M
<b>Protein marker</b>	
Protein MW Standards for Native PAGE (12 – 450 kDa)	39064.01
<b>Staining reagents and kits:</b>	
SERVA <i>Dens</i> Stain Blue G Staining Solution (2x concentrated, 500 ml)	35078.01
SERVA Blue R Staining Kit (2 x 500 ml)	42531.01
SERVA Silver Staining Kit Native PAGE (25 mini gels)	35077.01
SERVA Blue G	35050
SERVA Blue R	35051
Amido black10 B (50 g)	12310.01
Ponceau S solution (0.2 %, 500 ml)	33427.01
Silver nitrate	35110
<b>Buffer etc.</b>	
SERVA Tris-Glycine native electrophoresis buffer (10x)	42530
SERVA Tris-Glycine native sample buffer (2x)	42528
<b>Buffer etc.</b>	
Towbin buffer 10x, for native PAGE and for Western Blotting	42558
Semi-Dry blotting buffer kit (3 x 500 ml)	42559
Glycine	23390
Tris(hydroxymethyl)aminomethane	37186
Bromophenol blue, sodium salt	15375
Ethanol, undenatured, absolute	11093
Glycerol	23176
Trichloroacetic acid, 20 % solution	36913
<b>Membranes</b>	
Immobilin (PVDF), 9 x 12 cm, pore size: 0.2 µm (10 sheets)	42579.01
Immobilin (PVDF), 26.5 cm x 3.75 m, pore size: 0.2 µm (1 roll)	42574.01
Fluorobind (PVDF), 10 x 10 cm, pore size: 0.2 µm (20 sheets)	42573.01
Fluorobind (PVDF), 25 cm x 3 m, pore size: 0.2 µm (1 roll)	42571.01

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