

Sub-Cell[®] GT Agarose Gel Electrophoresis Systems

Instruction Manual

Catalog Numbers 170-4401 to 170-4406 170-4481 to 170-4486



Warranty

Bio-Rad Laboratories warrants the Sub-Cell GT, Wide Mini-Sub® Cell GT, and Mini-Sub Cell GT electrophoresis systems against defects in materials and workmanship for 1 year. If any defects occur in the instrument during this warranty period, Bio-Rad Laboratories will repair or replace the defective parts free. The following defects, however, are specifically excluded:

- 1. Defects caused by improper operation.
- 2. Repair or modification done by anyone other than Bio-Rad Laboratories or an authorized agent.
- 3. Use of fittings or other spare parts supplied by anyone other than Bio-Rad Laboratories.
- 4. Damage caused by accident or misuse.
- 5. Damage caused by disaster.
- 6. Corrosion due to use of improper solvent or sample.

This warranty does not apply to parts listed below:

1. Platinum Electrode Wires

To insure the best performance from the Sub-Cell GT electrophoresis systems, become fully acquainted with these operating instructions before use. Bio-Rad recommends that you first read these instructions carefully. Assemble and disassemble the unit completely without casting a gel. After these preliminary steps, you should be ready to cast and run a gel.

Bio-Rad also recommends that all Sub-Cell GT system components and accessories be inspected for damage, cleaned as recommended in this manual, and rinsed thoroughly with distilled water before use.

Record the following for your records:

Model
Catalog No.
Date of Delivery
Warranty Period
Serial No
Invoice No.
Purchase Order No.

For any inquiry or request for repair service, contact Bio-Rad Laboratories after confirming the model and serial number of your instrument.

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Section 1 General Information

1.1 Introduction

The Sub-Cell® GT instruments (basic Sub-Cell® GT cell, Wide Mini-Sub® Cell GT, and Mini-Sub Cell® GT) comprise a comprehensive and flexible gel electrophoresis system that effectively separates nucleic acids using submerged agarose gels. Submarine agarose gels are easy to cast and readily dissipate heat. These gels allow sample underlaying and prevent electrical field discontinuities caused by wicks or sample well dehydration. Agarose gels are ideal for the separation of DNA restriction digestions, polymerase chain reaction (PCR*)-amplified fragments, and genomic DNA and RNA prior to Southern or northern blotting. If operated correctly, agarose gel submarine electrophoresis can effectively separate nucleic acids from 20 base pairs to 20 kilobase pairs in length.

The Sub-Cell GT systems are designed for years of reproducible and rigorous use. These rugged systems incorporate many features that make casting and running agarose gels simple and efficient. The gel caster provides tape-free gel casting in trays. Gels can also be cast in the GT bases using specially designed casting gates. Replaceable electrode cassettes provide a simple way to replace electrode wires. A comprehensive assortment of base and tray sizes, including a variety of preparative, analytical, and multichannel pipet compatible combs, makes these systems ideal for any agarose gel application.

Note: This manual contains instructions for the Sub-Cell GT electrophoresis systems only. Prior to the release of the Sub-Cell GT systems, Bio-Rad supplied similar agarose gel electrophoresis cells: the original Sub-Cell DNA electrophoresis cell, Wide Mini-Sub cell, and Mini-Sub cell systems. This manual does not provide information on these earlier versions. Contact your local Bio-Rad representative for information concerning the original Sub-Cell systems.

Definition of Symbols



Caution, risk of electrical shock



Caution (refer to accompanying documents)

1.2 Safety





The Sub-Cell GT electrophoresis systems are designed for maximum user safety. The buffer chambers are made of 3/16 inch (.476 cm) thick injection-molded acrylic to create a leak-free electrophoresis environment. The safety lids surround the buffer chamber to protect the user from exposure to electrical currents. All Sub-Cell GT systems were designed for indoor use only.

Before use, inspect the GT base for cracks or chips, which may allow the buffer to leak from the base and cause a potential electrical hazard. Additionally, inspect all electrical cables, banana jacks, and plugs for loose connections, cracks, breaks, or corrosion. Do not use any part that is cracked, charred, or corroded. These parts may also cause a potential electrical hazard. Contact your local Bio-Rad representative before using a part that may be considered hazardous.

During electrophoresis, inspect the base and workbench for any signs of buffer leakage. If leaking buffer is detected, disconnect the power to the cell immediately and contact your local Bio-Rad representative.

Power to Sub-Cell GT units is supplied by an external DC voltage power supply. This power supply must be ground isolated in such a way that the DC voltage output floats with respect to ground. All of Bio-Rad's power supplies meet this important safety requirement. The recommended power supply for these units is the PowerPac™ Basic power supply. The PowerPac 300 power supply contains safety features such as no load, overload, rapid resistance change, and ground leak detection capabilities. The maximum specified operating parameters for the Sub-Cell GT systems are given in Table 1.1.

Table 1.1 Sub-Cell GT systems operating parameters

	Sub-Cell	Wide Mini-Sub	Mini-Sub	
	GT Cell	Cell GT	Cell GT	
Maximum voltage limit	200 VDC	150 VDC	150 VDC	
Maximum power limit	40 Watts	45 Watts	10 Watts	
Maximum Buffer temperature	40°C	40°C	40°C	

Electric current to the cell, from the external power supply, enters the unit through the lid assembly, which provides a safety interlock. Current to the cell is broken when the lid is removed. Do not attempt to circumvent this safety interlock, and always turn the power supply off before removing the lid or when working with the cell.

Important: These Bio-Rad instruments are certified to meet IEC 1010-1" safety standards. IEC-certified products are safe to use when operated in accordance with the instruction manual. This instrument should not be modified in any way. Alteration of this instrument will:

- Void the manufacturer's warranty
- Void the IEC 1010-1 safety certification
- Create a potential safety hazard

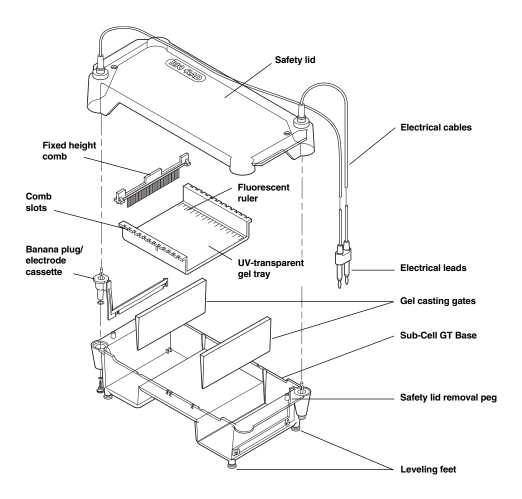
The Sub-Cell GT systems conform to IEC-1010-1 standards: safe between the operating temperatures of 4°C and 40°C and altitudes up to 2,000 meters; safe at a maximum relative humidity of 80% for temperatures up to 31°C, and for temperatures higher than 31°C, safe at a maximum humidity that decreases linearly to 50% at 40°C. Bio-Rad is not responsible for any injury or damage caused by the use of this instrument for purposes other than those for which it is intended, or by modifications of the instrument not performed by Bio-Rad or an authorized agent. There are no user-serviceable parts in this apparatus. To insure electrical safety, do not attempt to service this apparatus.

1.3 System Components

Each of the Sub-Cell GT systems comes with the components listed in Table 1.2 (see Figure 1.1, page 3, for part description). Check your instrument to be sure all items are present. Note any damage to the unit which may have occurred during shipping. Notify Bio-Rad Laboratories if any items are missing or damaged.

Table 1.2 Sub-Cell GT system components

	Sub-Cell GT System	Wide Mini-Sub Cell GT System	Mini-Sub Cell GT System
Item	Quantity	Quantity	Quantity
GT Base (buffer chamber)	1	1	1
Gel Casting Gates (optional)	2	2	2
Safety Lid and Cables	1	1	1
UVTP Gel Tray	1	1	1
Fixed Position Comb	2	2	2
	(15 well, 1.5 mm thick)	(15 well, 1.5 mm thick)	(8 well, 1.5 mm thick)
	(20 well, 1.5 mm thick)	(20 well, 1.5 mm thick)	(15 well, 1.5 mm thick)
Leveling Bubble	1	1	1
Gel Caster (optional)	1	1	1
Instruction Manual	1	1	1



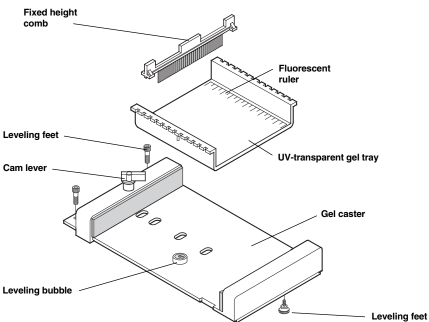


Fig.1.1. Sub-Cell GT components.

1.4 Specifications

Sub-Cell GT System	Wide Mini-Sub Cell GT System	Mini-Sub Cell GT System
42 x 20 x 10 cm	26 x 20 x 7.5 cm	26 x 12 x 6.5 cm
1,500-2,000 ml	650-900 ml	265-320 ml
15 x 15 cm	15 x 7 cm	7 x 7 cm
15 x 10 cm	15 x 7 cm	7 x 7 cm
15 x 15 cm	15 x 10 cm	7 x 10 cm
15 x 20 cm		
15 x 25 cm		
Molded	clear plastic	
Aluminu	ım	
Molded	clear plastic	
Molded	polycarbonate	
Gold-pla	ated brass, 4.4 cm leng	th
Dual, 20	AWG, tinned copper v	vire cable
	• •	,
UV-tran	sparent acrylic plastic (UVTP)
	•	20. ,
•		
	## System ## 42 x 20 x 10 cm ## 1,500–2,000 ml ## 15 x 15 cm ## 15 x 10 cm ## 15 x 20 cm ## 15 x 25 cm ## Molded ## Aluminu ## Molded ## Molded ## Gold-pl: ## Platinun Dual, 20 ## Flame-r Nickel s UV-tran ## Molded Polycari	42 x 20 x 10 cm 1,500–2,000 ml 15 x 15 cm 15 x 10 cm 15 x 15 cm 15 x 10 cm

[†] GT base buffer volumes will vary depending on the size and thickness of the gel used.

Section 2 Operating Instructions

Note: See Section 3, Gel and Electrophoresis Reagent Preparation, for information on the preparation of RNA gels. See References 1 and 2 for more information on DNA and RNA electrophoresis.

2.1 DNA Gel Preparation

DNA agarose gels can be used to separate and visualize DNA of various sizes. Before casting an agarose gel, consult Table 2.1, page 5, to determine the appropriate percent agarose gel to use, based on the size of DNA to be separated.

Procedure

 Determine the amount of agarose (grams) required to make the desired agarose gel concentration and volume. Use Tables 2.1 and 2.2, page 5, as a guide for agarose concentration and gel volume requirements.

Example: For a 1% agarose gel, add 1 gram of agarose to 100 ml of 1x electrophoresis buffer.

Table 2.1 Gel concentration required for DNA separation 1-2

Gel Concentration (%)	DNA Size (Kb)	
0.50	1–30	
0.75	0.8–12	
1.00	0.5–10	
1.25	0.4–7	
1.50	0.2–3	
2–5*	0.01–0.5	

^{*} Sieving agarose such as AmpliSize® agarose

Table 2.2 Gel volume requirements

Gel Size(thickness)	0.25 cm	0.5 cm	0.75 cm	1.0 cm
Base				
7 x 7 cm	10 ml	20 ml	30 ml	40 ml
15 x 7 cm	20 ml	40 ml	60 ml	80 ml
15 x 15 cm	50 ml	100 ml	150 ml	200 ml
Tray				
7 x 7 cm	10 ml	20 ml	30 ml	40 ml
7 x 10 cm	15 ml	30 ml	45 ml	60 ml
15 x 7 cm	20 ml	40 ml	60 ml	80 ml
15 x 10 cm	30 ml	60 ml	90 ml	120 ml
15 x 15 cm	50 ml	100 ml	150 ml	200 ml
15 x 20 cm	70 ml	140 ml	210 ml	280 ml
15 x 25 cm	90 ml	180 ml	270 ml	360 ml

2. Add the agarose to a suitable container (e.g., 250 ml Erlenmeyer flask, Wheaton bottle, etc.). Add the appropriate amount of 1x electrophoresis buffer (see Section 3, Gel and Electrophoresis Reagent Preparation, for electrophoresis buffer preparation) and swirl to suspend the agarose powder in the buffer. If using an Erlenmeyer flask, invert a 25 ml Erlenmeyer flask into the open end of the 250 ml Erlenmeyer flask containing the agarose. The small flask acts as a reflux chamber, allowing long or vigorous boiling without much evaporation.

Note: A mark can be put on the lower flask at the same level as the liquid. If evaporation occurs, water can be added to bring the liquid back to the original starting level.

3. The agarose can be melted by boiling on a magnetic hot plate (Step 4a) or in a microwave oven (Step 4b).

Caution: Always wear protective gloves, goggles, and a lab coat while preparing and casting agarose gels. The vessels containing hot agarose can cause severe burns if allowed to contact skin. Additionally, molten agarose can boil over when swirled.

Magnetic hot plate method

4a. Add a stir bar to the undissolved agarose solution. Heat the solution to boiling while stirring on a magnetic hot plate. Bubbles or foam should disrupt before rising to the neck of the flask.

Microwave oven method

- 4b. Place the gel solution into the microwave. Using a low to medium setting, set the timer for a minimum of 5 minutes, stopping the microwave oven every 30 seconds and swirling the flask gently to suspend the undissolved agarose. This technique is the fastest and safest way to dissolve agarose.
- 5. Boil and swirl the solution until all of the small translucent agarose particles are dissolved. With the small flask still in place, set aside to cool to 60°C before pouring.

2.2 Casting Agarose Gel Slabs

There are several ways to cast agarose submarine gels using the Sub-Cell GT systems. Gels may be cast with or without a UV-transparent plastic (UVTP) tray directly on the stage of the Sub-Cell GT bases using the gel casting gates. Gels may also be cast on the removable UVTP trays with the aid of the gel caster or with standard laboratory tape.

Casting gels on the Sub-Cell GT base

- 1. Level the Sub-Cell GT base using the leveling bubble provided.
- 2. Slide the gel casting gates into the slots at opposite ends of the GT base.
- 3. Place the comb(s) into the appropriate slot(s) of the base so that the sample wells are near the cathode (black). DNA samples will migrate toward the anode (red) during electrophoresis.
- 4. Prepare the desired concentration and amount of agarose in 1x electrophoresis buffer (see Section 2.1). When the agarose solution has cooled to 50–60°C pour the molten agarose between the gates.

Warning: Hot agarose (>60°C) may cause the plastic in the cell to warp or craze and will decrease the lifetime of the Sub-Cell base. Warping may also result in sample wells of uneven depth.

- 5. Allow 20–40 minutes for the gel to solidify at room temperature.
- 6. Carefully remove the comb from the solidified gel. Remove the gel casting gates.
- 7. Submerge the gel beneath 2 to 6 mm of 1x electrophoresis buffer (see Section 3, Gel and Electrophoresis Reagent Preparation). Use greater depth overlay (more buffer) with increasing voltages to avoid pH and heat effects.

Casting gels on the base stage with the UVTP tray

- 1. Level the cell using the leveling bubble provided.
- 2. Place the UVTP tray on the GT base.

Note: The Mini-Sub Cell GT requires the 7 x 7 cm UVTP tray for casting in the GT base. The Wide-Mini-Sub Cell GT requires the 15 x 7 cm UVTP tray and the Sub-Cell GT system requires the 15 x 15 cm UVTP tray for casting in the GT base.

- 3. Slide the gel casting gates into the slots at opposite ends of the GT base stage. Insure that the gates are evenly seated in the slots and the gates uniformly contact all edges of the UVTP tray. The weight of the gates provides a tight seal to prevent any leakage problems during gel casting.
- 4. Place the comb(s) into the appropriate slot(s) of the trays so that the sample wells are near the cathode (black). DNA samples will migrate toward the anode (red) during electrophoresis.
- 5. Prepare the desired concentration and amount of agarose in 1x electrophoresis buffer (see Section 2.1). When the agarose solution has cooled to 50–60°C, pour the molten agarose between the gates.

Warning: Hot agarose (>60°C) may cause the tray to warp or craze and will decrease the lifetime of the tray. Warping may also result in sample wells of uneven depth.

- 6. Allow 20–40 minutes for the gel to solidify at room temperature.
- 7. Carefully remove the comb from the solidified gel. Remove the gel casting gates.
- 8. Submerge the gel beneath 2 to 6 mm of 1x electrophoresis buffer (see Section 3, Gel and Electrophoresis Reagent Preparation). Use greater depth overlay (more buffer) with increasing voltages to prevent pH and heat effects.

Removable tray (UVTP) gel casting using a Gel Caster or Mini-Gel Caster

- 1. Level the Gel Caster or Mini-Gel Caster using the leveling feet in the gel caster and the leveling bubble provided.
- 2. Disengage and slide the movable wall to the open end of the Gel Caster or Mini-Gel Caster by turning and lifting the cam peg upward.

Note: If casting more than one gel with the Gel Caster, add the removable gel casting wall to the gel caster. The removable wall will allow casting using two 15×10 cm trays, four 7×10 cm trays or one 15×10 cm and one 15×15 cm trays.

- Place the open edge of the UVTP tray against the fixed wall of the Gel Caster or Mini-Gel Caster.
- 4. Slide the movable wall against the edge of the UVTP tray (Figure 2.1).
- 5. To seal the open tray ends, engage the cam peg by turning and pressing downward simultaneously.
- 6. When the cam peg has dropped into the appropriate slot, turn the peg in either direction until resistance is felt. This action seals the edges of the tray for casting.
- 7. Place the comb(s) into the appropriate slot(s) of the tray.

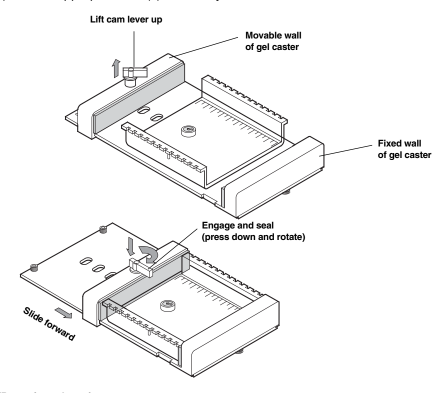


Fig. 2.1. Sealing the UVTP tray for gel casting.

8. Prepare the desired concentration and amount of agarose in 1x electrophoresis buffer (see Section 2.1). When the agarose solution has cooled to 50–60°C pour the molten agarose between the gates.

Warning: Hot agarose (>60°C) may cause the tray to warp or craze and will decrease the lifetime of the tray. Warping may also result in sample wells of uneven depth.

- 9. Allow 20-40 min for the gel to solidify at room temperature.
- 10. Carefully remove the comb from the solidified gel.
- 11. Disengage the cam peg by turning and lifting upward. Slide the movable wall away from the tray. Remove the tray from the Gel Caster or Mini-Gel Caster.
- **Note:** While the gel is solidifying, a light seal is formed between the gasket and the gel (especially for low percentage agarose gels [<0.8%]). Before moving the wall away from the tray, carefully lift the tray on one side to release the seal or use a spatula to break the seal between the agarose and gasket.
- 12. Place the tray onto the leveled Sub-Cell base so that the sample wells are near the cathode (black). DNA samples will migrate toward the anode (red) during electrophoresis.
- 13. Submerge the gel beneath 2 to 6 mm of 1x electrophoresis buffer (see Section 3, Gel and Electrophoresis Reagent Preparation). Use greater depth overlay (more buffer) with increasing voltages to avoid pH and heat effects.

Removable tray (UVTP) gel casting using tape

- 1. Seal the ends of the UVTP gel tray securely with strips of standard laboratory tape. Press the tape firmly to the edges of the gel tray to form a fluid-tight seal.
- Level the gel tray on a leveling table or workbench using the leveling bubble provided with the instrument.
- 3. Prepare the desired concentration and amount of agarose in 1x electrophoresis buffer (see Section 2.1). When the agarose solution has cooled to 50–60°C pour the molten agarose into the gel tray.
 - **Warning:** Hot agarose (>60°C) may cause the tray to warp or craze and will decrease the lifetime of the tray. Warping may also result in sample wells of uneven depth.
- 4. Allow 20-40 min for the gel to solidify at room temperature.
- 5. Carefully remove the comb from the solidified gel.
- 6. Remove the tape from the edges of the gel tray.
- 7. Place the tray onto the leveled Sub-Cell base so that the sample wells are near the cathode (black). DNA samples will migrate toward the anode (red) during electrophoresis.
- 8. Submerge the gel beneath 2 to 6 mm of 1x electrophoresis buffer (see Section 3, Gel and Electrophoresis Reagent Preparation). Use greater depth overlay (more buffer) with increasing voltages to avoid pH and heat effects.

2.3 Electrophoresis

After the agarose gel has solidified, sample loading and electrophoresis can begin. Agarose gels can be run in many different types of electrophoresis buffers. Nucleic acid agarose gel electrophoresis is usually conducted with either Tris-Acetate-EDTA (TAE) buffer or Tris-Borate-EDTA (TBE) buffer. While TAE buffers provide faster electrophoretic migration of linear DNA and better resolution of supercoiled DNA, TBE buffers have a stronger buffering capacity for longer or higher voltage electrophoresis runs. Bio-Rad offers premixed 50x TAE and 10x TBE buffers, as well as individual buffer reagents for use with the Sub-Cell GT systems.

1. Prepare samples for gel loading. The maximum sample loading volumes for Bio-Rad's combs are listed in Section 6.2. Loading volume is dependent upon the type of comb used (*i.e.*, well thickness and length) and thickness of the gel.

- 2. When loading volume is determined, add standard nucleic acid sample loading dye to a final 1x concentration to make samples dense for underlaying into sample wells (see Section 3, Gel and Electrophoresis Reagent Preparation, for sample loading dye preparation).
- 3. Load the samples into the wells using standard pipets. Multichannel pipets can only be used for loading samples with the Bio-Rad MP combs (see Section 6.2).
 - **Note:** Sample wells are often difficult to see. Well visualization can be enhanced by placing black paper or tape under the base or trays in the are of comb placement and well formation.
- 4. Place the lid on the DNA cell carefully. Do not disturb the samples. The Sub-Cell GT system lids attach to the base in only one orientation. To attach the lid correctly, match the red and black banana jacks on the lid with the red and black banana plugs of the base.
- 5. Power requirements vary depending on gel thickness, length, agarose, concentration, and type of electrophoresis buffer used. Refer to Tables 2.3 and 2.4 below for relative sample migration rates for the different Sub-Cell GT systems and for DNA size migration with sample loading dyes.

Note: Buffer recirculation is not required for most standard DNA and RNA agarose gel electrophoresis. If buffer recirculation is required, simply turn off the power supply, remove the safety lid, and mix the running buffer as desired. After the buffer has been mixed, reconnect the safety lid and continue with electrophoresis.

Table 2.3 Relative sample migration rates *

Cell Type	Voltage	Bromophenol Blue Migration Rate
Sub-Cell GT cell, 15 x 15 cm gel	75 V	3.0 cm/hr
Wide Mini-Sub cell GT, 15 x 10 cm gel	75 V	4.5 cm/hr
Mini-Sub cell GT, 7 x 10 cm gel	75 V	4.5 cm/hr

^{*} These sample migration rates were determined based on a 0.5 cm thick 1.0% agarose gel using Bio-Rad's Molecular Biology Certified Agarose in 1x TAE electrophoresis buffer (diluted from Bio-Rad's Premixed 50x TAE Buffer). Migration rates will vary depending on the voltage, current, and type of agarose or buffer used.

Table 2.4 DNA size migration with sample loading dyes

Agarose** Concentration (%)	Xylene Cyanol	Bromophenol Blue
0.5–1.5	4–5 Kb	400–500 bp
2.0-3.0**	750 bp	100 bp
4.0-5.0**	125 bp	25 bp

^{**} Sieving agarose such as AmpliSize agarose.

2.4 Nucleic Acid Staining and Visualization

Gels can be removed from the Sub-Cell GT base or gel tray for nucleic acid staining. The gel can also remain on the UVTP gel tray for staining.

Ethidium bromide staining procedure

1. Place the gel into the appropriate volume of 0.5 μ g/ml ethidium bromide (EtBr) stain for 15–30 min. Use enough staining solution to cover the entire gel.

Caution: Ethidium bromide is a suspected carcinogen and should be handled with extreme care. Always wear gloves, eye glasses, and a laboratory coat. Dispose of used EtBr solutions and gels appropriately (Review EtBr Material Safety Data Sheet [MSDS] for proper disposal methods).

2. Destain the gel for 10–30 min in dH₂O with the same volume used for staining.

Note: Ethidium Bromide can be removed from the DNA with extended destaining. This will cause lower sensitivity of detection. However, insufficient destaining will create higher background fluorescence.

- 3. Rinse the gel briefly with dH₂O to remove any residual staining solution.
- 4. Place the gel on a UV transilluminator for nucleic acid visualization and analysis. DNA-ethidium bromide complexes may be illuminated with UV light of 254, 302, or 366 nm. Sensitivity decreases with illumination at higher wavelengths. However, nicking of DNA will increase below 302 nm. Table 2.5 gives the percentage of transmittance of UV light through 1/4" (.64 cm) UV-transparent plastic.

Note: Nucleic acids in the gel can be visualized through the UVTP trays. If a UVTP tray is not used, place household plastic wrap between the UV transilluminator and the gel to avoid contaminating the transilluminator with nucleic acids or EtBr.

Table 2.5 Percent UV transmittance through 1/4" (.64 cm) UV-transparent plastic (UVTP)

	Approximate		
Wavelength (nm)	% Transmittance		
254	0		
302	80		
366	90		

5. Photograph the gel using standard cameras and film (e.g., Bio-Rad's Standard Polaroid Gel Documentation System) or with CCD-based digitized image analysis systems (e.g., Gel Doc™ 1000 UV fluorescent gel documentation system). Gels are generally photographed with a yellow, orange, or red interference filter. Red filters generally give the cleanest background. Bio-Rad offers a full-line of standard photography and CCD-based imaging systems for nucleic acid gel analysis.

2.5 Note on Blotting

Nucleic acids within the gel can be transferred to membranes using the techniques of Southern and northern blotting. It is beyond the scope of this instruction manual to include blotting procedures. Consult references #1 and #2 for blotting techniques. Bio-Rad offers a full line of nitrocellulose and positively charged nylon membranes, as well as vacuum and electrophoretic blotting apparatus for Southern and northern blotting.

Section 3 Gel and Electrophoresis Reagent Preparation

RNA agarose formaldehyde gels

For 100 ml of a 1% agarose formaldehyde gel prepare as follows:

62 ml of 1.6% melted agarose

20 ml 5x MOPS electrophoresis buffer (1x final concentration)

18 ml 12.3 M (37.5%) formaldehyde (2.2 M final concentration)

Caution: Formaldehyde solutions and formaldehyde vapors are toxic. When handling solutions or gels that contain formaldehyde use a chemical hood. Always wear gloves, eye glasses, and a laboratory coat when using formaldehyde. See the MSDS for safety information.

Nucleic acid electrophoresis buffers¹⁻²

DNA agarose gel electrophoresis is usually performed using either Tris-Acetate-EDTA (TAE) or Tris-Borate-EDTA (TBE). While TAE buffers provide faster electrophoretic migration of linear DNA and better resolution of supercoiled DNA, TBE buffers have a stronger buffering capacity for longer or higher voltage electrophoresis runs. Bio-Rad offers premixed 50x TAE and 10x TBE buffers for use with the Sub-Cell GT systems. RNA formaldehyde gels require a MOPS [3-(N-morpholino)-propanesulfonic acid] electrophoresis buffer.

1x Tris-Acetate-EDTA (TAE)—40 mM tris (pH 7.6), 20 mM acetic acid, and 1 mM EDTA.

50x Stock (1 liter)—dissolve in 600 ml distilled water:

242 g Tris base (FW = 121)

57.1 ml glacial acetic acid

100 ml 0.5 M EDTA (pH 8.0).

Fill to a final volume of 1 liter with distilled water.

1x Tris-Borate-EDTA (TBE)—89 mM tris (pH 7.6), 89 mM boric acid, 2 mM EDTA

10x Stock (1 liter)—dissolve in 600 ml distilled water:

108 g Tris base (FW = 121)

55 g boric acid (FW = 61.8)

40 ml 0.5 M EDTA (pH 8.0)

Fill to a final volume of 1 liter with distilled water.

1x MOPS Buffer (RNA Gels)—0.02 M MOPS [3-(N-morpholino)-propanesulfonic acid] (pH 7.0), 8 mM sodium acetate, 1 mM EDTA (pH 8.0)

5x Stock (1 liter)—dissolve in 600 ml DEPC-treated distilled water:

20.6 g MOPS

13.3 ml 3 M sodium acetate (DEPC treated), pH 7.4

10 ml 0.5 M EDTA (DEPC-treated), pH 8.0

Fill to a final volume of 1 liter with DEPC-treated distilled water.

Caution: DEPC is a suspected carcinogen. Always wear gloves, eye glasses, and a laboratory coat. Use caution when handling DEPC containing solutions. Consult the DEPC MSDS (Material Safety Data Sheet) for more information.

DNA and RNA sample loading dye¹⁻²

A convenient 10x sample buffer stock consists of 50% glycerol, 0.25% bromophenol blue, and 0.25% xylene cyanole FF in 1x TAE buffer. Only 1–10 ml of the 10x loading dye should be prepared.

RNA sample preparation¹⁻²

Prior to loading RNA onto an agarose formaldehyde gel prepare each RNA sample as follows:

6 μl RNA in DEPC-treated water

10 µl 5x MOPS buffer (final concentration 1.67x)

9 µl 12.3 M formaldehyde (final concentration 3.7 M)

25 µl formamide (final concentration 50% v/v)

Caution: Formamide is a teratogen. Always wear gloves, eye glasses, and a laboratory coat. Use caution when handling formamide. Consult the formamide MSDS for more information.

Ethidium bromide solution

Add 10 mg of EtBr to 1 ml distilled water. Bio-Rad offers EtBr solutions (10 mg/ml).

Section 4 Care and Maintenance

4.1 Cleaning Sub-Cell GT Components

 All Sub-Cell GT system parts should be washed with a mild detergent solution in warm water.

Note: Be careful not to snag or break the electrode wire in the GT base while cleaning.

2. Rinse all parts thoroughly with warm water or distilled water and air dry, if possible.

4.2 Compatible Cleaning Agents

Chemically compatible cleaners must be used to insure long life of parts. These include:

Aqueous solutions of soaps and mild detergents:

Bio-Rad Cleaning Concentrate (catalog number 161-0722) Dishwashing liquid

• Organic solvents:

Hexane

Aliphatic hydrocarbons

Do not leave plastic parts to soak in detergents more than 30 minutes. A short detergent rinse typically is all that is required.

Caution: Do not use the following chemicals to clean Sub-Cell GT parts. Exposure to these chemicals may cause the plastic parts to crack, craze, etch, or warp.

Chlorinated hydrocarbons

Carbon tetrachloride

Chloroform

· Aromatic hydrocarbons

Benzene

Phenol

Toluene

Methyl ethyl ketone

Acetone

Alcohols

Methanol

Ethanol

Isopropyl alcohol

Do not use abrasive or highly alkaline cleaners on Sub-Cell GT parts.

Do not expose Sub-Cell GT parts to temperatures >60°C. Do not sterilize Sub-Cell GT parts by autoclaving or dry heat.

4.3 Maintenance Schedule

Item	Look For	Frequency	Action
All parts	Dried salts, agarose, grease, and dirt	Each use	Clean parts as described in Section 4.1
Electrical cables	Breaks or fraying	Each use	Replace cables
Trays	Chips or cracks	Each use	Replace tray
Electrode wires	Breaks	Each use	See Section 4.4 (Electrode cassette replacement)
Cable connections or (banana jacks and holders plugs)	Looseness	Weekly	Replace banana jacks banana plug
GT base	Crazing, cracks, or leaks	Monthly	Replace GT base or banana plug holder o-ring

4.4 Electrode Replacement

The Sub-Cell GT systems allow easy replacement of broken electrode wires by removing the banana plug/electrode wire assembly and ordering a new assembly from Bio-Rad (Figure 4.1). Order the new assembly using the part description and catalog numbers listed in Section 6, Product Information.

- 1. Remove the thumb screw and rubber gasket from the banana plug chamber of the Sub-Cell GT base to release the banana plug/electrode wire assembly. Do not discard this thumb screw or rubber gasket (keep these parts with the GT base).
- 2. Remove the broken wire assembly by lifting upward on the banana plug. Discard the broken assembly.
- 3. Insert the new assembly into the banana plug chamber of the GT base. Make sure the electrode wire guard guides are properly seated into the electrode wire guard slots in the bottom of the GT base.
- 4. Replace and tighten the thumb screw and rubber gasket to secure the assembly in the base and to form a leak-free seal in the banana plug holder chamber.

Note: Test for buffer leakage by filling the base with water and checking for leakage of water through the banana plug chamber of the base. If leakage occurs, tighten the thumb screw.

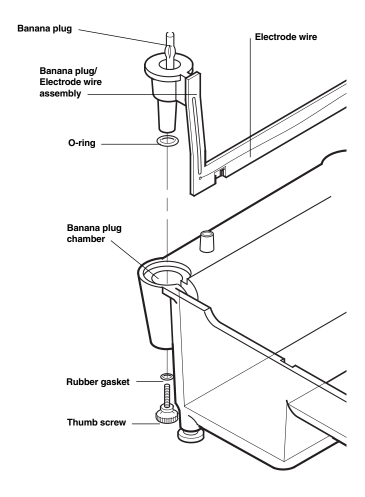


Fig. 4.1. Removal of banana plug/electrode wire assembly.

4.5 RNase Decontamination

Sub-Cell GT parts can be cleaned with a mild detergent and treated for 10 minutes with 3% hydrogen peroxide (H_2O_2), and then rinsed with 0.1% DEPC- (diethyl pyrocarbonate) treated distilled water, to eliminate RNases prior to using the Sub-Cell GT systems for RNA gels.¹⁻² Consult references #'s1–2 for other suggestions regarding the use of DEPC in RNase decontamination.

Caution: DEPC is a suspected carcinogen. Always wear gloves, eye glasses, and a laboratory coat. Use caution when handling DEPC-containing solutions. Consult the DEPC MSDS for more information.

Do not attempt to decontaminate RNase from Sub-Cell GT parts using extreme dry heat.

Note: Several commercial products are available for eliminating RNase contamination. RNase *ZAP*® (Ambion) is a safe, simple, and effective method that if used properly does not craze or fog the Sub-Cell GT parts. See manufacturer's instructions for proper use.

Section 5 Troubleshooting

Symptoms	Cause	Solutions
Slanted lanes (bands)	Gel not fully solidified Comb warped or at an angle	Let gel solidify for at least 30–45 min. Check alignment of comb.
Curved line or distortion of lanes (bands)	Bubbles in sample wells	Remove bubbles prior to electrophoresis.
Differential relative mobilities	Sample spilled out of wells	Samples should have proper density. Apply carefully.
	Unit not leveled	Level unit. Place on steady work bench.
Curved bands, smiles	Sample overload	Reduce load.
Ragged bands	Sample density incorrect	See sample application instructions.
	Sample well deformed	Carefully remove comb, especially from soft gels. Be sure gel has solidified. Cooling soft gels aids in comb removal.
	Excessive power or heating	Reduce voltage. See electrophoresis instructions.
Band smearing and streaking	Agarose has improper endosmosis (m _r)	Consult Bio-Rad about agarose.
	Salt concentration in sample too high	Reduce salt concentration to ≥ 0.1 M.
	Excessive power and heating	Reduce voltage. See electrophoresis instructions.
	Sample spilled out of well	Apply sample carefully. Increase gel thickness for large sample volumes. Adjust comb height.
	Incomplete digest, nuclease contamination, bad enzyme	Heat sample. Check enzyme activity. Digest sample further.

Symptoms	Cause	Solutions
	Sample wells cast through the gel. Sample leaks along bottom of running surface.	Comb should be placed 1 to 2 mm above the base of the running surface.
	Sample overload	Dilute sample.
Bands sharp but too few bands seen	Gel agarose percentage too high	Lower agarose percentage.
	Incomplete digest	Check enzyme activity, digest further.
High MW bands sharp; Low MW bands smeared	Gel agarose percentage too low	Increase agarose percentage. Switch to polyacrylamide.
Gels crack	Too high voltage gradient, especially with low melting temperature agarose or low gel strength gels	Reduce voltage. Run gel at lower temperature.

Section 6 Product Information

6.1 Sub-Cell® GT Systems

Catalog Number	Product Description
170-4401	Sub-Cell GT System, with 15 x 10 cm tray
170-4402	Sub-Cell GT System, with 15 x 15 cm tray
170-4403	Sub-Cell GT System, with 15 x 20 cm tray
170-4404	Sub-Cell GT System, with 15 x 25 cm tray
170-4481	Sub-Cell GT System, with 15 x 10 cm tray and gel caster
170-4482	Sub-Cell GT System, with 15 x 15 cm tray and gel caster
170-4483	Sub-Cell GT System, with 15 x 20 cm tray and gel caster
170-4484	Sub-Cell GT System, with 15 x 25 cm tray and gel caster
170-4405	Wide Mini-Sub Cell GT System, with 15 x 7 cm tray
170-4485	Wide Mini-Sub Cell GT System, with 15 x 7 cm tray and mini-gel caster
170-4406	Mini-Sub Cell GT System, with 7 x 7 cm tray
170-4486	Mini-Sub Cell GT System, with 7×7 cm tray and mini-gel caster
Sub-Cell Sy	stems with PowerPac Basic Power Supply, 100–120/220–240 V
164-0302	Sub-Cell GT Cell and PowerPac Basic Power Supply
164-0301	Wide Mini-Sub Cell GT Cell and PowerPac Basic Power Supply
164-0300	Mini-Sub Cell GT Cell and PowerPac Basic Power Supply

6.2 Sub-Cell GT System Accessories

Catalog Number	Product Description
Sub-Cell GT	Systems
170-4410	Sub-Cell GT Base
170-4411	Sub-Cell GT Safety Lid with Cables
170-4412	Gel Caster
170-4413	Sub-Cell GT Electrode (Anode), red
170-4414	Sub-Cell GT Electrode (Cathode), black
170-4415	Sub-Cell GT Gel Casting Gates
170-4416	GT UVTP Gel Tray, 15 x 10 cm
170-4417	GT UVTP Gel Tray, 15 x 15 cm
170-4418	GT UVTP Gel Tray, 15 x 20 cm
170-4419	GT UVTP Gel Tray, 15 x 25 cm
Wide Mini-S	ub Cell GT Systems
170-4420	Wide Mini-Sub Cell GT Base
170-4421	Wide Mini-Sub Cell GT Safety Lid with Cables
170-4422	Mini-Gel Caster
170-4423	Wide Mini-Sub Cell GT Electrode (Anode), red
170-4424	Wide Mini-Sub Cell GT Electrode (Cathode), black
170-4425	Wide Mini-Sub Cell GT Gel Casting Gates
170-4416	GT UVTP Gel Tray, 15 x 10 cm
170-4426	GT UVTP Gel Tray, 15 x 7 cm
Mini-Sub Ce	II GT Systems
170-4430	Mini-Sub Cell GT Base
170-4431	Mini-Sub Cell GT Safety Lid with Cables
170-4422	Mini-Gel Caster
170-4432	Mini-Sub Cell GT Electrode (Anode), Red
170-4433	Mini-Sub Cell GT Electrode (Cathode), Black
170-4434	Mini-Sub Cell GT Gel Casting Gates
170-4435	GT UVTP Gel Tray, 7 x 10 cm
170-4436	GT UVTP Gel Tray, 7 x 7 cm

Sub-Cell Systems Combs Fixed Height Combs For Sub-Cell GT and Wide Mini-Sub Cell GT Systems

Catalog Number	Well Number	Thickness (mm)	Well Width (mm)	Well Volume Capacity* (μΙ)
170-4443	10	0.75	9.87	37.0
170-4444	10	1.50	9.87	74.0
170-4445	15	0.75	5.52	20.7
170-4446	15	1.50	5.52	41.4
170-4447	20	0.75	4.84	18.2
170-4448	20	1.50	4.84	36.4
170-4449	30	1.50	2.69	20.2

Adjustable Height Combs For Sub-Cell GT and Wide Mini-Sub Cell GT Systems (Adjustable height combs require a comb holder [catalog number 170-4320])

Catalog Number	Well Number	Thickness (mm)	Well Width (mm)	Well Volume Capacity⁺ (μl)
170-4325	10	0.75	9.87	37.0
170-4326	10	1.50	9.87	74.0
170-4323	15	0.75	5.52	20.7
170-4324	15	1.50	5.52	41.4
170-4321	20	0.75	4.84	18.2
170-4322	20	1.50	4.84	36.4
170-4344	30	1.50	2.69	20.2

Preparative Combs For Sub-Cell GT and Wide Mini-Sub GT Systems (Adjustable height combs require a comb holder [catalog number 170-4320])

Catalog Number	Well Number	Thickness (mm)	Well Width (mm)	Well Volume Capacity* (μΙ)
170-4440	1	1.50	106.43	800.0
170-4441*	2	1.50	50.29	377.0
170-4442	4	1.50	26.42	200.0
170-4328*	1	1.50	106.43	800.0

Multi-channel Pipet Compatible (MP) Fixed Height Combs For Sub-Cell GT and Wide Mini-Sub Cell GT Systems

Catalog Number	Well Number	Thickness (mm)	Well Width (mm)	Well Volume Capacity* (μΙ)
170-4450	10	0.75	5.82	21.8
170-4451	10	1.50	5.82	43.6
170-4452	14	0.75	5.82	21.8
170-4453	14	1.50	5.82	43.6
170-4454	18	0.75	2.91	10.9
170-4455	18	1.50	2.91	21.8
170-4456	26	0.75	2.91	10.9
170-4457	26	1.50	2.91	21.8

Fixed Height Combs for Mini-Sub Cell GT

Catalog Number	Well Number	Thickness (mm)	Well Width (mm)	Well Volume Capacity* (µI)
170-4460	1	1.50	43.43	325.7
170-4461	2	1.50	20.32	152.4
170-4462	8	0.75	5.54	20.8
170-4463	8	1.50	5.54	41.6
170-4464	15	0.75	2.59	9.7
170-4465	15	1.50	2.59	19.4

Adjustable Height Combs for Mini-Sub Cell GT (Adjustable height combs require a comb holder [catalog 170-4331])

Catalog Number	Well Number	Thickness (mm)	Well Width (mm)	Well Volume Capacity* (μΙ)
170-4342	1	1.50	43.43	325.7
170-4333	8	1.50	5.54	41.6
170-4332	15	1.50	2.59	19.4

 $^{^{\}star}\,$ Well volume capacity was determined based on a well depth of 0.5 cm.

6.3 Related Bio-Rad Products

Power Supplies

164-5050	PowerPac Basic Power Supply, $100/120-220/240 \ V$
164-5052	PowerPac HC Power Supply, 100/120-220/240 V

Blotting Membranes

162-0153	Zeta-Probe® Positively Charged Nylon Blotting Membrane, sheets, 9 x 12 cm, 15
162-0154	Zeta-Probe Positively Charged Nylon Blotting Membrane, sheets, 10 x 15 cm, 15
162-0155	Zeta-Probe Positively Charged Nylon Blotting Membrane, sheets, 15 x 15 cm, 15
162-0156	Zeta-Probe Positively Charged Nylon Blotting Membrane, sheets, 15 x 20 cm, 15
162-0157	Zeta-Probe Positively Charged Nylon Blotting Membrane, sheets, 20 x 20 cm, 15
162-0158	Zeta-Probe Positively Charged Nylon Blotting Membrane, sheets, 20 x 25 cm, 3
162-0159	Zeta-Probe Positively Charged Nylon Blotting Membrane, roll, $30\ cm\ x\ 3.3\ m,\ 1$
162-0165	Zeta-Probe Positively Charged Nylon Blotting Membrane, roll, $20\ cm\ x\ 3.3\ m,\ 1$
162-0190	Zeta-Probe GT (Genomic Tested) Positively Charged Nylon Blotting Membrane, sheets, 9×12 cm, 15
162-0191	Zeta-Probe GT (Genomic Tested) Positively Charged Nylon Blotting Membrane, sheets, 10×15 cm, 15

Catalog Number	Product Description			
162-0192	Zeta-Probe GT (Genomic Tested) Positively Charged Nylon Blotting Membrane, sheets, 15×15 cm, 15			
162-0193	Zeta-Probe GT (Genomic Tested) Positively Charged Nylon Blotting Membrane, sheets, 15×20 cm, 15			
162-0194	Zeta-Probe GT (Genomic Tested) Positively Charged Nylon Blotting Membrane, sheets, 20×20 cm, 15			
162-0195	Zeta-Probe GT (Genomic Tested) Positively Charged Nylon Blotting Membrane, sheets, 20×25 cm, 3			
162-0196	Zeta-Probe GT (Genomic Tested) Positively Charged Nylon Blotting Membrane, roll, 30 cm \times 3.3 m, 1			
162-0197	Zeta-Probe GT (Genomic Tested) Positively Charged Nylon Blotting Membrane, roll, 20 cm \times 3.3 m, 1			
162-0090	Supported Nitrocellulose Membrane, 0.45 micron, sheets, 7 x 8.4 cm, 10			
162-0091	Supported Nitrocellulose Membrane, 0.45 micron, sheets, 10 x 15 cm, 10			
162-0092	Supported Nitrocellulose Membrane, 0.45 micron, sheets, 15 x 15 cm, 10			
162-0093	Supported Nitrocellulose Membrane, 0.45 micron, sheets, 20 x 20 cm, 10			
162-0094	Supported Nitrocellulose Membrane, 0.45 micron, roll, 30 cm x 3 m, 1			
162-0095	Supported Nitrocellulose Membrane, 0.20 micron, sheets, 7 x 8.4 cm, 10			
162-0096	Supported Nitrocellulose Membrane, 0.20 micron, sheets, 15 x 15 cm, 10			
162-0097	Supported Nitrocellulose Membrane, 0.20 micron, roll, 30 cm x 3 m, 1			
Vacuum Blot	tting Apparatus			
165-5000	Model 785 Vacuum Blotter			
165-5001	Model 785 Vacuum Blotter System, 120 VAC			
165-5002	Model 785 Vacuum Blotter System, 220/240 VAC			
Semi-Dry Tra	ansfer Cells			
170-3940	Trans-Blot® SD Semi-Dry Electrophoresis Transfer Cell			
UV Crosslink	king Chamber			
165-5031	GS Gene Linker® UV Chamber, 120 VAC			
165-5032	GS Gene Linker UV Chamber, 220 VAC			
165-5033	GS Gene Linker UV Chamber, 240 VAC			
165-5034	GS Gene Linker UV Chamber, 100 VAC			

Number	Product Description
Gel Reager	nts and Standards
162-0019	Low Melt Preparative Grade Agarose, 100 g
170-8200	AmpliSize DNA Size Standard, 50-2,000 bp
161-0404	Bromophenol Blue, 10 g
161-0423	Xylene Cyanole FF, 25 g
161-0433	Ethidium Bromide Solution, 10 ml, 10 mg/ml
Electropho	resis Buffers
161-0733	10x Tris/Boric Acid/EDTA (TBE), 1
161-0743	50x Tris/Acetic Acid/EDTA (TAE), 1
161-0719	Tris, 1 kg
DNA Gel Im	nage Analysis and Documentation Systems
170-8170	Molecular Imager Gel Doc XR System-PC
170-8171	Molecular Imager Gel Doc XR System-MAC

Section 7 References

Catalog

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- 2. **Current Protocols in Molecular Biology**, Greene Publishing Associates and Wiley-Interscience, 1989.

Additional Reading

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- 6. Bittner, M., Kupferer, P. and Morris, C.F., Anal. Biochem., 102, 459 (1980).
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- 10. Dretzen, G., Bellard, M., Sassone-Corsi, P. and Chambon, P., *Anal. Biochem.*, **112**, 295 (1981).

^{*} The Polymerase Chain Reaction (PCR) process is covered by patents owned by Hoffmann-LaRoche. Use of the PCR process requires a license.

^{**} IEC 1010-1 is an internationally accepted electrical safety standard for laboratory instruments.



Bio-Rad Laboratories, Inc.

Life Science Group Web site www.bio-rad.com USA 800 4BIORAD Australia 61 02 9914 2800 Austria 01 877 89 01 Belgium 09 385 55 11 Brazil 55 21 3237 9400 Canada 905 364 3435 China 86 21 6426 0808 Czech Republic 420 241 430 532 Denmark 44 52 10 00 Finland 09 804 22 00 France 01 47 95 69 65 Germany 089 318 84 0 Greece 30 210 777 4396 Hong Kong 852 2789 3300 Hungary 36 1 455 8800 India 91 124 4029300 Israel 03 963 6050 Italy 39 02 216091 Japan 03 6361 7000 Korea 82 2 3473 4460 Mexico 52 555 487 670 The Netherlands 0318 540666 New Zealand 0508 805 500 Norway 23 38 41 30 Poland 48 22 331 99 99 Portugal 351 21 472 7700 Russia 7 495 721 14 04 Singapore 65 6415 3188 South Africa 27 861 246 723 Spain 34 91 590 5200 Sweden 08 555 12700 Switzerland 061 717 95 55 Taiwan 886 2 2578 7189 United Kingdom 020 8328 2000