

Protocol: DNA Staining in Gels with dsGreen

dsGreen is a fluorescent dye that binds specifically to double-stranded DNA. There are three variants of the staining protocol: gel soaking, gel pre-staining, and sample pre-staining.

Gel soaking

Classical method for agarose and polyacrylamide gels.

1. Run sample(s) in an agarose or polyacrylamide gel.
2. In a beaker, add 10 μ L of the [10,000 \$\times\$ dsGreen solution in DMSO](#) to 100 mL of 1 \times TE, TBE, or TAE buffer (for mini gels), or 50 μ L of the [10,000 \$\times\$ dsGreen solution in DMSO](#) to 500 mL of 1 \times TE, TBE, or TAE buffer (for mid-sized gels). Mix thoroughly with a spatula, rod, or magnetic stirrer.
3. Pour the diluted dsGreen solution into an appropriate tray or pan and submerge the gel.
4. Soak the gel for 5–10 min.
5. View or document the gel using available light source and an green/yellow filter. Transilluminators with blue light, or with UV low pressure mercury lamp (254 nm) can be used to visualize gels stained with dsGreen. High pressure mercury lamp (365 nm) can be used too, but this light source gives somewhat less efficient excitation.

Gel pre-staining

This method is suitable for **agarose** gels only, but not for PAAG. Note — this staining method can sometimes cause bands to warp or form smears. Use gel soaking in this case.

1. Boil the agarose in buffer to dissolution using a microwave or heating appliance.
2. While still fluid, add 1 μ L of the [10,000 \$\times\$ dsGreen solution in DMSO](#) per each 10 mL of gel solution. Mix thoroughly.
3. Pour the gel and let it solidify.
4. For best results, add 1 μ L of the [10,000 \$\times\$ dsGreen solution in DMSO](#) per each 10 mL of buffer near the anode («+», red wire).
5. Run the samples. Real-time monitoring of migrating bands under a 254 nm low-pressure mercury lamp is possible.
6. View or document the gel using available light source and an green/yellow filter. Transilluminators with blue light, or with UV low pressure mercury lamp (254 nm) can be used to visualize gels stained with dsGreen. High pressure mercury lamp (365 nm) can be used too, but this light source gives somewhat less efficient excitation.

Sample pre-staining

Least sensitive, most economical method.

1. Mix 25 μ L of DMSO and 1 μ L of the [10,000 \$\times\$ dsGreen solution in DMSO](#).
2. Add 1 μ L of the solution to each sample to be separated on an agarose or polyacrylamide gel.
3. Run the samples. Real-time monitoring of migrating bands under a 254 nm low-pressure mercury lamp is possible.
4. View or document the gel using available light source and an green/yellow filter. Transilluminators with blue light, or with UV low pressure mercury lamp (254 nm) can be used to visualize gels stained with dsGreen. High pressure mercury lamp (365 nm) can be used too, but this light source gives somewhat less efficient excitation.

Lumiprobe Corporation

201 International Circle, Suite 135
Hunt Valley, Maryland 21030
USA
Phone: +1 888 973 6353
Fax: +1 888 973 6354
Email: order@lumiprobe.com

Lumiprobe GmbH

Feodor-Lynen-Strasse 23
30625 Hannover
Germany
Phone: +49 511 16596811
Fax: +49 511 16596815
Email: de@lumiprobe.com

Lumiprobe RUS Ltd

Kotsyubinsky street, 4
121351 Moscow
Russian Federation
Phone: +7 800 775 3271
Email: ru@lumiprobe.com