

ELISABETH PHARMACON Ltd. Rokycanova 4437/5

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EliDNATM PS Green Instructions for Use

Package:

Ref. No. Package ED01 1 ml ED01s 50 μl

Storage:

The dye should be stored at 4 °C and protected from light. When stored under these conditions, the dye will retain full activity until the expiration date indicated on the label.

Product description

EliDNATM PS Green is a new generation of green fluorescent nucleic acid dye intended for gel staining. The dye is non-toxic and non-mutagenic. Therefore, making it a safer alternative to ethidium bromide. EliDNATM PS Green can be used with a standard UV transilluminator (300 nm) or with instruments using visible light (~500 nm). Gels stained with EliDNATM PS Green are compatible with various downstream applications such as gel extraction, cloning and many others.

Protocol

Prestaining:

Prepare the desired volume of agarose gel solution according to your protocol. For a 100 ml agarose solution add 4-6 μ l of **EliDNA**TM **PS Green** and mix it properly. Allow the gel solution with the dye to cool down to ~60 °C before casting it into the gel tray. After the gel has solidified, load the samples and perform electrophoresis. Image the gel with a UV transilluminator or with a visible light with ~500 nm using yellow or green filter.

Poststaining:

Prepare and run the agarose gel according to your protocol. For 100 ml of buffer or distilled water use 20-30 μ l of **EliDNA**TM **PS Green**. Store the staining solution at room temperature in the dark. Use a suitable container and place the gel into the solution. Recommended staining time is 10-20 min. Depending on the thickness of the gel and its concentration, staining time and the volume of the dye may be adjusted.



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Troubleshooting

- If you experience low fluorescence signal of DNA bands, try to add more dye. Signal intensity depends on the number of bands and their concentration.
- We do not recommend repeated melting of gel containing the dye. This may lead to reduced signal intensity.
- Smeared DNA bands may be caused by excessive DNA concentration. Reduce the amount of loaded DNA or perform poststaining.
- If there are any discrepancies in bands migration, try to reduce the amount of loaded DNA or use less dye.

Related products

| REF | Name of product | UV light | Blue LED | Green/blue LED | In gel | Post-st. | Loading | dsDNA | ssDNA | RNA |
|------|-------------------------------------|-------------|-------------|-------------------|--------|----------|---------|-------|------------|------------|
| ED01 | EliDNA [™] PS Green | ✓ | ✓ | ✓ | ✓ | ✓ | Х | 1 | √ ¹ | √ ¹ |
| ED02 | EliDNA [™] PS Green Plus | ✓ | ✓ | ✓ | ✓ | ✓ | Х | ✓ | ✓ | √ |
| ED03 | EliDNA [™] PS Green Ultra² | ✓ | ✓ | ✓ | ✓ | ✓ | Х | ✓ | ✓ | √ |
| ED04 | EliDNA [™] PS Red | ✓ | Х | Х | ✓ | ✓ | Х | ✓ | ✓ | √ |
| ED05 | EliDNA [™] LD Green | ✓ | ✓ | ✓ | Х | Х | ✓ | ✓ | ✓ | √ |
| ED06 | FliDNA™LD Red | | Х | Х | Х | Х | 1 | 1 | 1 | |

¹ Since ssDNA and RNA are single stranded, you may experience a lower signal intensity.

Manufacturer:

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Catalog number



Batch code



Use by (last day of month)



Upper limit of temperature



Manufacturer



Contains sufficient "N" tests

²The dye with the highest sensitivity.