

GEL STAINING OF NUCLEIC ACIDS

GEL STAINING SOLUTIONS

Nucleic acid dyes for agarose and acrylamide gels, for laboratories and practical applications. We offer you high-quality stains with high sensitivity (silver stains), standard stains (ethidium bromide) in user-friendly dropper bottles, and non-toxic green-fluorescent dyes for blue or UV light. The following table provides an overview of the various stains and their respective sensitivity levels.

Summary: Staining of Nucleic Acid in Gels

Staining	Art. No.	Sensitivity	Description
Roti®-Black N	N769	<0,1 ng/mm ²	Silver staining of DNA polyacrylic amide gels, visible under white light.
Roti®-Black NSeq	P081	<0,1 ng/mm ²	Silver staining of DNA sequencing gels, visible under white light.
Roti®-Load DNastain	5783, 5784, 6472	1,5 ng/mm ²	Sensitive, non-toxic fluorescent staining of DNA (excitation 320 nm and 490 nm/blue light). Combined with gel loading buffer for fragments >500 bp (DNastain 1), 100-2000 bp (DNastain 2), or <500 bp (DNastain 3).
Roti®-GelStain	3865	1,5 ng/mm ²	Sensitive fluorescent staining of DNA (excitation: 290-320 nm wave length). Non-toxic, noncarcinogenic.
Ethidium bromidesolution	2218 (1 %) HP46 (0.5 %) HP47 (0.025 %)	1,5 ng/mm ²	Quick standard staining of nucleic acid, fluorescent (excitation: 254-360 nm wave length).
Roti®-Methylen blue staining concentrate	0648	10 ng/mm ²	Reversible, non-toxic blue staining of DNA, visible under white light.

ROTI[®]-GELSTAIN

Mechanismus

- Ready-to-use staining mixture prepared from a variety of reagents, dye component: orange-red to brown powder.
- Benzimidazoles typically bind to the minor groove of helical nucleic acid. No intercalation takes place.
- Fluorescence staining, requires UV-excitation.
- Excitation maximum (DNA bound): 290-320 nm
Emission maximum (DNA bound): 515 nm

Hazar

None known. Not membrane permeable.

Applications

Suitable for all agarose gels.

Not suitable for staining of PAA gels.

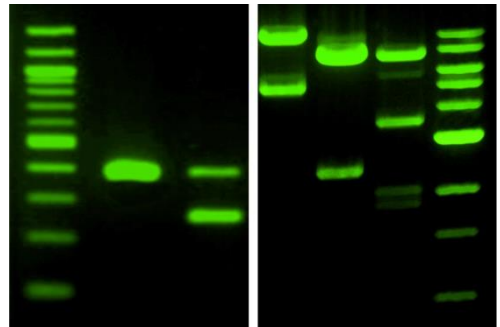
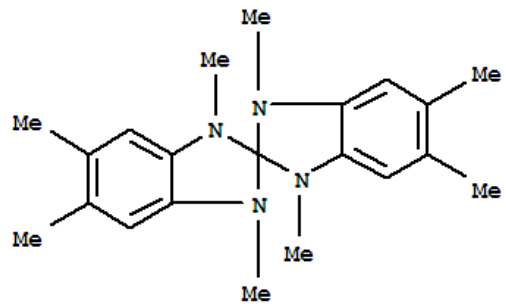
Sensitivity: 1.5 ng/mm² (0.2 ng/band).

Staining

- Addition to gel (liquid agarose) [ca. 5 µl/100 ml]
- Addition to running buffer [ca. 25 µl/100 ml]
- Staining after gel run [ca. 25 µl/100 ml]
- Addition to loading buffer /nucleic acid not recommended
- Staining is reversible

Please note

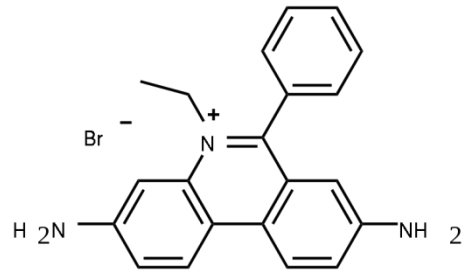
- When applied during the gel run: Nucleic acid that has been loaded with Roti[®]-GelStain shows slightly altered running behaviour when compared to free nucleic acid.
- Staining depends on secondary and tertiary structure of the nucleic acids. Due to the optimal formation of the minor groove, longer fragments of linear dsDNA are stained more efficiently than short fragments (below 500 bp), ssDNA, short RNA or cyclic dsDNA.
- DNA quantitation in the gel is possible, as long as the both DNAs compared are having the same secondary and tertiary structure. For instance, cyclic DNA should only be quantified by comparison with a cyclic DNA standard.



ETHIDIUM BROMIDE

Mechanism

- Intercalation between base pairs of nucleic acids
- Very strong binding to dsDNA
- Fluorescence staining, requires UV-excitation
- Excitation maxima
DNA bound: 330 nm, 500 nm (Excitation possible at 260 - 350 nm)
Unbound: 210 nm, 285 nm, 470 nm
Emission maximum: 605 nm



Hazar

Dangerous up to toxic, mutagenicity may not completely be excluded

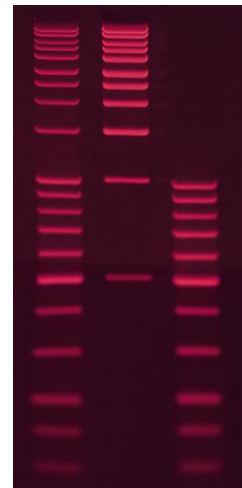
Applications

Suitable for all agarose gels and PAA gels.

Sensitivity: 1.5 ng/mm² (0.2 ng/band)

Staining

- Addition to loading buffer /nucleic acid [ca. 2 µg/ml]
- Addition to gel (liquid agarose) [ca. 0.2 µg/ml]
- Staining after gel run [ca. 2 µg/ml, 10 mins, optional: destaining for 15 mins.]
- Staining in principle reversible, but hard to remove.



Please note

- When applied during the gel run: Nucleic acid that has been loaded with ethidium bromide shows slightly altered running behaviour when compared to free nucleic acid.
- Linear dsDNA will be stained more intensely than ssDNA, RNA, or cyclic dsDNA.
- Quantitation of DNA in the gel is possible if compared DNA has identical secondary and tertiary structure; e.g.: compare cyclic DNA only with equally cyclic DNA.

METHYLENE BLUE

Mechanism

- Ionic binding to phosphoric acid of nucleic acids results in weak binding
- Blue staining, visible in white light

Hazard

None known.

Applications

Only applicable as staining solution after the gel run.

Only suitable for thin agarose gels of up to approx. 1.5 % agarose and PAA gels.

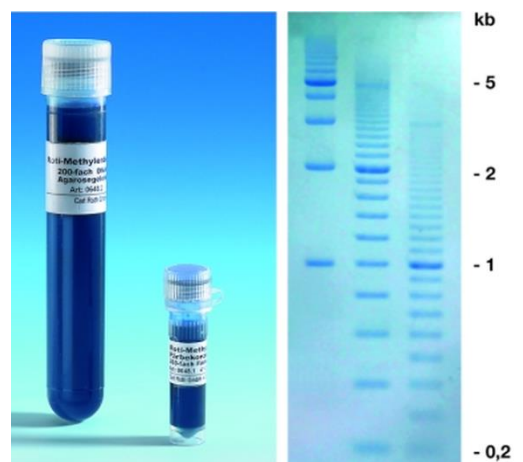
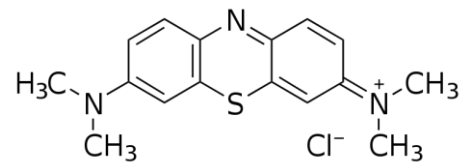
Sensitivity: 10 ng/mm² (1.5 ng/band)

Staining

- Staining for 10 - 30 min.
- Destaining necessary for 15 min. to over night.
- Reversible staining, easy to remove.

Please note

- Methylene Blue Staining Concentrate contains a solvent, causing DNA to precipitate after addition to the loading buffer.
- dsDNA/RNA will be stained more intensely than ssDNA/RNA



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