

GFP-Booster for Immunofluorescence of GFP-Fusion Proteins

For the immunofluorescence of GFP-fusion-proteins in fixed cells.

Only for research applications, not for diagnostic or therapeutic use

1. Introduction

Green fluorescent proteins (GFP) and variants thereof are widely used to study protein localization and dynamics in living cells. However, photo stability and quantum efficiency of GFP are not sufficient for Super-Resolution Microscopy (e.g. 3D-SIM or STED) of fixed samples. In addition, many cell biological methods such as BrdU-staining, EdU-Click-iT™ treatment or Fluorescent *In Situ* Hybridization result in disruption of the GFP signal. The GFP-Booster_Atto488, a specific GFP-binding protein coupled to the fluorescent dye ATTO 488 (from ATTO-TEC), reactivates, boosts and stabilizes your GFP signal (for a complete list of recognized GFP variants, please visit the FAQ section at www.chromotek.com).

2. Content

Reagent	Code	Quantity
GFP-Booster_Atto488	gba488	100 µg

Storage buffer: 1x PBS, 0.09% sodium azide

3. Optical properties

ATTO 488: Excitation range 480 - 510 nm (λ_{abs} = 501 nm)
Emission range 520 - 560 nm (λ_{fl} = 523 nm)

For further information please refer to <http://www.atto-tec.com>

4. Stability and Storage

Shipped at ambient temperature. Upon receipt store at +4°C.
Stable for 6 month. Do not freeze. Protect from light.

5. Protocol

- Fixation:** Fix cells seeded on coverslips in 3.7% formaldehyde in PBS for 10 min at room temperature.
- Wash samples three times with PBS containing 0.1% Tween 20 (PBST).
- Permeabilization:** Add PBS containing 0.5% Triton X-100 to samples and incubate for 5 min at room temperature.
Alternatively permeabilize cells by incubating samples in 100% methanol for 5 min at -20°C.
- Wash samples twice with PBST.
- Blocking:** Add 4% BSA in PBST to samples and incubate for 10 min at room temperature.
- GFP-Booster incubation:** Dilute GFP-Booster 1:200 in blocking buffer and incubate for 1 h at room temperature.
Note: For multiplexing protocols you can combine GFP-Booster with any other antibody.
- Wash samples three times for 5-10 min in PBST.
- If required counter stain with DNA fluorescent dyes, e.g. DAPI.
- Mounting:** Rinse sample shortly in water to prevent salt crystal formation. Mount in VectaShield (Vector Labs) or other mounting media with anti-fading agents and seal mounted coverslips with clear nail polish.

Suggested buffer composition

Buffer	Composition
Fixation buffer	3.7% formaldehyd in PBS
Permeabilization buffer	PBS; 0.5% Triton X-100
Wash buffer	PBS; 0.1% Tween 20
Blocking buffer	4% BSA (w/v); PBS; 0.1% Tween 20

**Support/
Troubleshooting**

Please refer to our FAQ section at www.chromotek.com or contact support@chromotek.com

Related products

GFP Toolbox	code
GFP-Trap [®] _M	gtm-20; gtm-100; gtm-200; gtm-400
GFP-Trap [®] _M Kit	gtmk-20
GFP-Trap [®] _A	gta-20; gta-100; gta-200; gta-400
GFP-Trap [®] _A Kit	gtak-20
GFP-multiTrap	gtp-96; gtp-480
Blocked agarose beads	bab-20
Blocked magnetic beads	bmp-20
GFP antibody	3h9

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