

DNA-PAINT KIT MASSIVE-sdAB-FAST 1-PLEX

Expiration after 6 months

(For research use only)

CONTENT

SECONDARY SINGLE DOMAIN ANTIBODIES

- FluoTag®-XM-QC Anti-Mouse IgG kappa light chain (Clone: 1A23) + FAST docking site F1 (To be measured with Imager F1) or FluoTag®-XM-QC Anti-Rabbit IgG (Clone: 10E10) + FAST dock ing site F2 (To be measured with Imager F2)
- Concentration: 5 μ M Protein, 5 μ M DNA (1 DNA strand per protein)
- Volume: 100 µL
- Storage: -20 °C
- Storage buffer: PBS, 50 % glycerol + 0.05 % NaN₃
- Recommened dilution: 1:100 1:500 (for optimal results the dilution needs to be optimized depending on the target accessiblity and expression level)

IMAGERS

- Imager F1 or F2 Cy3B or ATTO 655
- Concentration: 1 μM in TE buffer (10 mM Tris, 1 mM EDTA, pH 8)

	 Volume: 300 μL
	 Storage: 20 °C (1 µM imager solutions are stable for multiple freeze-and-thaw cycles)
	Optional: Prepare 50 μL aliquots and store at -20 °C. Working aliquots can be stored at 4 °C for short-term or -20 °C for long term
	stable in plastic tubes.
	BUFFERS
	 Antibody incubation buffer, 30 mL, store at 2-8 °C
	Note: For longer-term storage we recommend to store aliquots at -20 °C.
	 Washing buffer (10×), 20 mL, store at room temperature (to be diluted 1:10 in water before use) Imaging buffer, 50 mL, store at room temperature
SAMPLE PREP.	1. Prepare sample using a protocol optimized for your target and primary antibody staining.
PROTOCOL	2. Wash with washing buffer (1×).
	3. Dilute secondary DNA-PAINT single domain antibodies in Antibody incubation buffer.
	4. Incubate for 30 - 60 min at room temperature.
	5. Wash three times with washing buffer (1×).
	6. Optional: Incubate fiducial markers.
	7. Wash with imaging buffer before adding the final imaging solution with imager strands.
	8. Before imaging: Add imager strands diluted in imaging buffer. We recommend a starting concentra- tion of 1 nM. However, the optimum imager concentration strongly depends on the target and la- beling density. Thus, the imager concentration should be adjusted such that distinct single molecule blinking events can be observed.
	9. After imaging, exchange buffer to washing buffer (1×) for storage at 4 °C.
IMAGING	Exposure time: 30-50 ms
	 Laser-Intensity: ~300 W/cm² (561 nm) and ~500 W/cm² (640 nm). This intensity might vary due to
PARAMETERS	different illumination depths/modes (TIRF/HILO). For dense targets we recommend increasing the laser power to enhance blinking.
	• Total imaging time/target: 5-15 min (Depends on target density and applied imager concentration)
	• Temperature: The kit is optimized for image acquisition at 21-25 °C. At higher temperatures shorter
	exposure times and higher laser powers are required.

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