

TRBC1/TRBC2 double-labeling: accuracy in the identification of T neoplasms by flow cytometry

Characterization of the $\alpha\beta$ TCR repertoire at the isoform level by flow cytometry



Different dyes available

ADVANTAGES

The combined use of anti-TRBC1 and TRBC2 allows the detection of clonal T populations with high accuracy, similar to kappa/lambda analysis in B cells.

1

Fast and reliable detection of T clonalities.

2

Increases diagnostic sensitivity, reduces artifacts of single labeling.

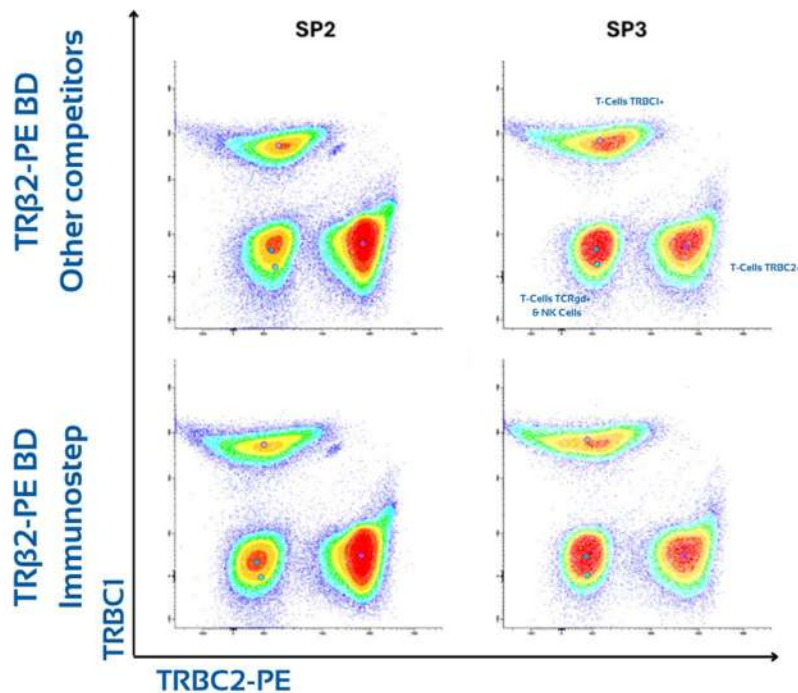
3

Improved quality of analysis, ideal for clinical lab.

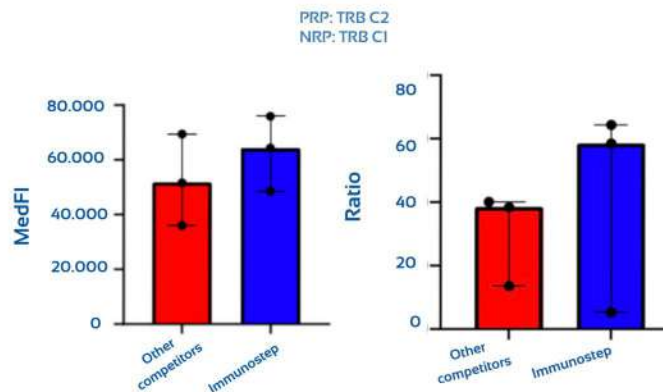
4



Do you want more information?
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all the details.

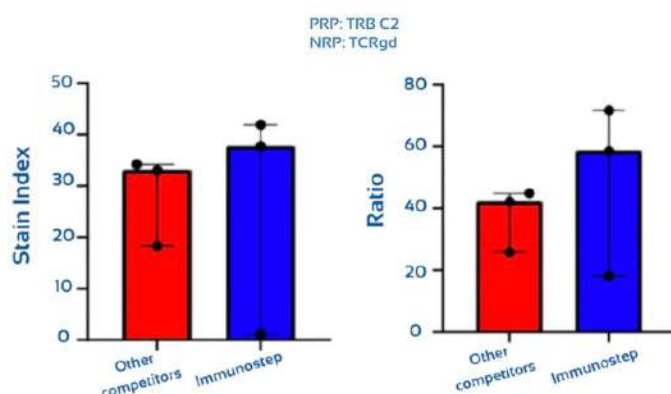


TCRCB1 and TCRCB2 are mutually exclusive allow clear and non-overlapping gating.



	Median MedFI (min-max)	Ratio (min-max)
Other competitors	50,451 (36,054 - 69,363)	38,49 (13,60 - 40,17)
Immunostep	64,275 (48,566 - 75,822)	58,59 (5,33 - 64,45)

Better performance than other competitors.



Specificity of the TRBC2-PE antibody, distinguishing its background reactivity on non-target cells (TCRγδ or TRBC1-positive target cells (TCRγδ or TRBC1 positive).

- The exclusive use of TRBC1 can induce artifacts ("dim TRBC1"), present in 27% of the samples. Double labeling completely resolves these artifacts and allows a more precise identification of the expressed TRBC isoform.
- Tested in different protocols with optimal performance in all of them: EuroFlow™ FACS Lysis, staining-lysis (NH 4 Cl)-wash, lysis (NH 4 Cl)-staining-wash.
- Evaluated in the basic and extended module T.

Anti-Human TCR Cβ 1 (JOVI-1)

	REF	Σ		[A]	
PURE	JOVIPU	1 mg	1 mg/ml		
FITC	JOVIF	100 test	2 µL/test	0,02 mg/ml	
Dy-634	JOVIDY634	100 test	2 µL/test	0,02 mg/ml	
PE-Cyanine7	JOVIPC7	50 test	3 µL/test	0,03 mg/ml	
					RUO

Anti- Human TCR Cβ 2 (SAM.2.rMAb)

	REF	Σ		[A]	
PE	TCRCB2PE	100 test	2 µL/ test	0,5 mg/ml	
					RUO