

Datasheet for 600-901-379S**RFP Antibody****Overview**

Description:	Anti-RFP (CHICKEN) Antibody - 600-901-379S
Item No.:	600-901-379S
Size:	25 µL
Applications:	ELISA, WB, FC, IF, IHC, Multiplex
Reactivity:	RFP, rRFP, tdTomato
Host Species:	Chicken

Product Details

Background:	Fluorescent proteins such as Discosoma Red Fluorescent Protein (DsRed) from sea anemone Discosoma sp. mushroom or green fluorescent protein (GFP) from Aequorea victoria jellyfish are widely used in research practice. Fusion RFP and GFP commonly serve as marker for gene expression and protein localization. As DsRed and GFP share only 19% identity, therefore, in general, anti-GFP antibodies do not recognize DsRed protein and vice versa. Structurally, Discosoma red fluorescent protein is similar to Aequorea green fluorescent protein in terms of its overall fold (a β -can) and chromophore-formation chemistry. However, Discosoma red fluorescent protein undergoes an additional step in the chromophore maturation and obligates tetrameric structure. Rockland offers many controls, monoclonal, and polyclonal antibodies for RFP.
Synonyms:	chicken anti-RFP antibody, DsRed, rDsRed, Discosoma sp. Red Fluorescent Protein, Red fluorescent protein drFP583
Host Species:	Chicken
Clonality:	Polyclonal
Format:	IgY

Target Details

Gene Name:	DsRed
Reactivity:	RFP, rRFP, tdTomato
Immunogen Type:	Recombinant Protein

Immunogen:	Red Fluorescent Protein (RFP) fusion protein corresponding to the full length amino acid sequence (234aa) derived from the mushroom polyp coral <i>Discosoma</i> .
Purity/Specificity:	RFP Antibody was prepared from egg yolks by a multi-step process which includes filtration, delipidation, salt fractionation and extensive dialysis against the buffer stated above. RFP Antibody was tested by western blot.
Relevant Links:	• UniProtKB - Q9U6Y8 • 600-901-379 SDS

Application Details

Tested Applications:	ELISA, WB
Suggested Applications:	FC, IF, IHC, Multiplex (Based on references)
Application Note:	Anti-RFP is designed to detect recombinant RFP. Anti-RFP antibody has been tested by ELISA and western blot to detect RFP. Use either alkaline phosphatase or peroxidase conjugated polyclonal anti-RFP to detect RFP or RFP containing proteins on western blots. Optimal titers for applications should be determined by the researcher. This product shows optimal performance by western blot.
Assay Dilutions:	All assays should be optimized by the user. Recommended dilutions (if any) may be listed below.
ELISA:	1:10,000
WB:	1:2,000 - 1:3,000

Formulation

Physical State:	Liquid (sterile filtered)
Concentration:	1.05 mg/mL by UV absorbance at 280 nm
Buffer:	0.02 M Potassium Phosphate, 0.15 M Sodium Chloride, pH 7.2
Preservative:	0.01% (w/v) Sodium Azide
Stabilizer:	None

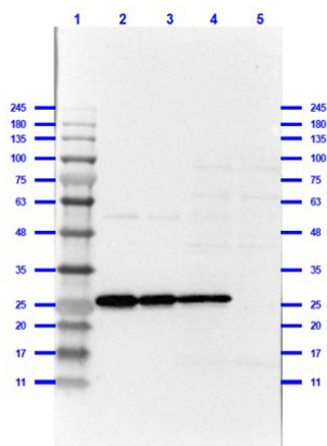
Shipping & Handling

Shipping Condition:	Dry Ice
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Storage Condition: Store vial at -20° C or below prior to opening. This vial contains a relatively low volume of reagent (25 µL). To minimize loss of volume dilute 1:10 by adding 225 µL of the buffer stated above directly to the vial. Recap, mix thoroughly and briefly centrifuge to collect the volume at the bottom of the vial. Use this intermediate dilution when calculating final dilutions as recommended below. Store the vial at -20°C or below after dilution. Avoid cycles of freezing and thawing.

Expiration: Expiration date is three (3) months from date of receipt.

Images



Western Blot

Western Blot of Chicken Anti-RFP Antibody.

Lane 1: Prestained MW marker (p/n MB-210-0500).

Lane 2: RFP (p/n 000-001-379) 0.1µg reduced [+].

Lane 3: RFP (p/n 000-001-379) 0.05µg reduced [+].

Lane 4: RFP/HEK293T (p/n 000-001-379/W09-001-GX5) 0.5µg/10µg reduced [+].

Lane 5: HEK293T (p/n W09-001-GX5) 10µg reduced [-].

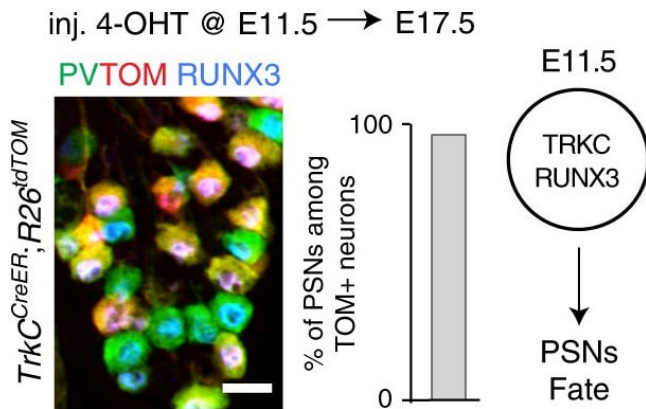
Blocking Buffer: BlockOut® Universal Blocking Buffer (p/n MB-073) for 1hr at RT.

Primary Antibody: Chicken Anti-RFP at 1µg/mL overnight at 2-8°C.

Secondary Antibody: Rabbit anti-Chicken IgG HRP (p/n 603-4302) at 1:40,000 for 30 mins at RT.

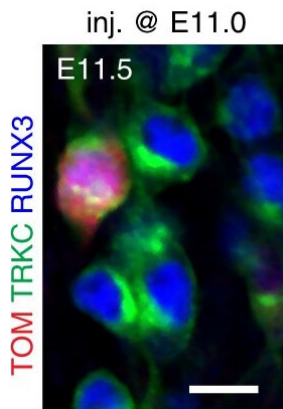
Predicted/Observed: 27kda.

c



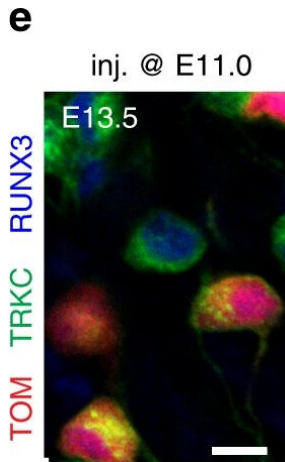
Immunocytochemistry

Differential expression of TRKC in PSNs prior to the cell death period. a Scheme of our working hypothesis. b, c Temporal fate mapping of TRKC PSNs by 4-OHT induction. TrkCCreER mice allow temporary activation of CreER in the TRKC+ cells 2 h after 4-OHT injection^{21,22}. Immunostaining for PV, RFP and RUNX3 on E17.5 DRG sections (c) and graph showing distribution of PV+/RUNX3+ PSNs among the TOM+ cells (n = 4). Scale bar: 20 μ m. d Quantification of PSNs at C5 and C7. ***P < 0.001, one-way analysis of variance (ANOVA) with Sidak's multiple comparisons test (n = 2–3). The window of PSNs cell death is shown. e TRKC expression in E11.5 ISL1+ (and RUNX3+, whose staining is not shown for more visibility) DRG neurons. Scale bar: 50 μ m. f TRKC levels in PSNs of e illustrated by color coding; dark blue indicates the lower and red the higher TRKC levels. From here, all observations are done at brachial levels (C5–8). g Distribution of TRKC levels in PSNs from e. h Distribution of TRKC levels in PSNs in E11.5 DRG neurons (from g). The data exhibit a Poisson-like distribution (one representative animal), with the mean used to define the two different categories of TRKC intensity (TRKCHigh and TRKCLow). i Projection of seven images of RUNX3+/TRKC+ PSNs from one brachial DRG; dots indicate TRKC-labeled neurons and color codes reveal TRKC intensity as shown in h. j Projection image of smFISH for pan Ntrk3 and Ntrk3 full length (FL) transcripts in E11.5 DRG, visualized at high magnification in (1) and (2) (images show full projection); right panel shows color coding of Ntrk3 FL levels in red; the brighter, the higher levels. k Distribution of the number of Ntrk3 FL molecules in E11.5 DRG neurons by smFISH, normalized to pan Ntrk3 (Ntrk3 FL represent 68% of all Ntrk3 transcripts). l TrkCCreER;R26tdTOM mice were injected at E9.75 with 4-OHT and analyzed at E11.5 (n = 3). m, n Frequency distribution (m) and pie chart (n) of TOM+/TRKC+ neurons from l according to their level of TRKC intensity. Source data are available as a Source Data file Figure provided by CiteAb. Source: Nat Commun, PMID: 31515492.

b


Immunocytochemistry

PSNs with high TRKC levels preferentially survive the cell death period. a Temporal fate mapping of TRKCHigh PSNs by 4-OHT (low dose, 0.02 g/kg). b–d Injection of TrkCCreER;R26tdTOM mice with low dose of 4-OHT at E11.0; DRG analyzed at E11.5 with recombination in few (b), preferentially high TRKC PSNs (c, d) ($P < 0.001$). Frequency distribution of TOM+ PSNs according to TRKC intensity (c) and pie charts (d) illustrating the large proportion of TOM+ cells among TRKCHigh PSNs. Scale bar: 50 μ m. e, f Percentage of recombined PSNs at E11.5 and E13.5 in DRGs from TrkCCreER;R26tdTOM animals after 4-OHT injection at E11.0 (* $P < 0.05$, Student's t-test; $n = 2$ litters with 6 embryos, E11.5; 2 litters with 5 embryos, E13.5). g The percentage of labeled PSNs does not change between E11.5 and E13.5 in R26CreERT2;R26tdTOM embryos injected at E11.0 with 0.032 g/kg 4-OHT ($n = 4$). Similarly, the recombination rate in TOM+ PSNs does not change between E14.5 and E16.5 in TrkCCreER;R26tdTOM embryos after injection at E14.0 with 0.02 g/kg 4-OHT ($n = 2$). Unpaired Student's t-test. h Whole-mount immunostaining for TRKC, NF160 and RFP of E13.5 forelimb from TrkCCreER;R26tdTOM embryos injected with low dose 4-OHT at E11.0. Insert shows restricted number of TOM+ fibers dispersed amongst TRKC+ axons. Scale bar: 200 μ m. i Pattern and color-coded depth (in micrometers) of innervation of TRKC+, NF160+ and RFP+ nerve fibers (processed from h) of E13.5 forelimb from TrkCCreER;R26tdTOM embryos injected with a low dose of 4-OHT (0.02 g/kg) at E11. The pattern and depth color code reveal similar territories (in all dimensions, xyz) of innervation of the TOM+ PSNs compared to all axons (NF160). Scale bar: 200 μ m. j Scheme illustrating the preferential selection of TRKCHigh PSNs during the cell death period. Source data are available as a Source Data file Figure provided by CiteAb. Source: Nat Commun, PMID: 31515492.

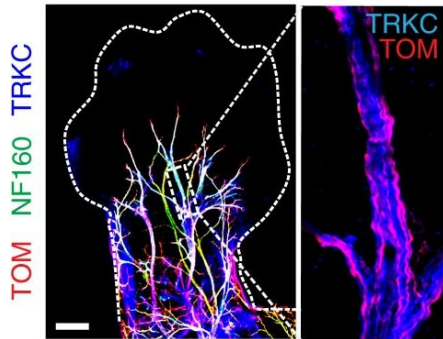


Immunocytochemistry

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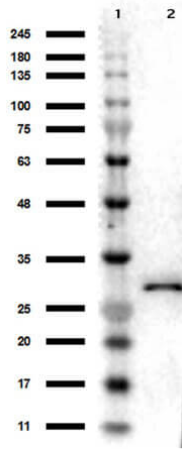
h

E13.5 *TrkC^{CreER}; R26^{tdTOM}*
inj @ E11.0



Immunohistochemistry

PSNs with high TRKC levels preferentially survive the cell death period. a Temporal fate mapping of TRKCHigh PSNs by 4-OHT (low dose, 0.02 g/kg). b–d Injection of *TrkCCreER;R26tdTOM* mice with low dose of 4-OHT at E11.0; DRG analyzed at E11.5 with recombination in few (b), preferentially high TRKC PSNs (c, d) ($P < 0.001$). Frequency distribution of TOM+ PSNs according to TRKC intensity (c) and pie charts (d) illustrating the large proportion of TOM+ cells among TRKCHigh PSNs. Scale bar: 50 μ m. e, f Percentage of recombined PSNs at E11.5 and E13.5 in DRGs from *TrkCCreER;R26tdTOM* animals after 4-OHT injection at E11.0 (* $P < 0.05$, Student's t-test; $n = 2$ litters with 6 embryos, E11.5; 2 litters with 5 embryos, E13.5). g The percentage of labeled PSNs does not change between E11.5 and E13.5 in *R26CreERT2;R26tdTOM* embryos injected at E11.0 with 0.032 g/kg 4-OHT ($n = 4$). Similarly, the recombination rate in TOM+ PSNs does not change between E14.5 and E16.5 in *TrkCCreER;R26tdTOM* embryos after injection at E14.0 with 0.02 g/kg 4-OHT ($n = 2$). Unpaired Student's t-test. h Whole-mount immunostaining for TRKC, NF160 and RFP of E13.5 forelimb from *TrkCCreER;R26tdTOM* embryos injected with low dose 4-OHT at E11.0. Insert shows restricted number of TOM+ fibers dispersed amongst TRKC+ axons. Scale bar: 200 μ m. i Pattern and color-coded depth (in micrometers) of innervation of TRKC+, NF160+ and RFP+ nerve fibers (processed from h) of E13.5 forelimb from *TrkCCreER;R26tdTOM* embryos injected with a low dose of 4-OHT (0.02 g/kg) at E11. The pattern and depth color code reveal similar territories (in all dimensions, xyz) of innervation of the TOM+ PSNs compared to all axons (NF160). Scale bar: 200 μ m. j Scheme illustrating the preferential selection of TRKCHigh PSNs during the cell death period. Source data are available as a Source Data file Figure provided by CiteAb. Source: Nat Commun, PMID: 31515492.



Western Blot

Western Blot Results of Chicken Anti-RFP Antibody. Lane 1: Opal PreStained Molecular Weight Marker (p/n MB-210-0500). Lane 2: RFP (p/n 000-001-379), load 50ng. Primary Antibody: Anti-RFP 1 μ g/mL overnight at 4°C. Secondary Antibody: Goat Anti-Chicken HRP (p/n 603-103-126) at 1:40,000 for 30min at RT. Blocking: BlockOut (p/n MB-073) for 30min at RT. Expect: 27kDa.

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