# PRODUCT INFORMATION **TRITC Labeled Lectin**

	Catalog Number:	R-5101-1				e. The success of yo
	Description:	Pure Limax flavus lectin (LFA) from garden	slug, TRITC o	conjugated.	to high leve	block tissue section. els of non specific ba prescent Labeled Lo
	Lot Number:				3. Incubate ti	ssue section with Flu e section with Buffe
	Protein Concentration: (Based on OD280)	1 mg purified LFA TRITC / 1 ml Buffer.				ssue section with Flumbar et. al., (1973).
	TRITC / Protein Ratio: (OD 550 / OD 280)					with Buffer (See rev s by centrifugation.
	Purification Procedure:	Gel filtration performed after conjugation to	remove free	TRITC.	4. Incubate ap	prescent Labeled La pproximately 1x10 <sup>6</sup> e or in a 37°C water
	Carbohydrate Specificity:	Sialic acid (generic term for deriva and N-Glycolylneuraminic acid.	atives of N	I-Acetylneuraminic acid	6. Examine co	with Buffer three tin ells, with or without t ss. (1977). Experime
	Inhibitory Carbohydrate:	N-Acetylneuraminic acid and N-Glycolylneu	raminic acid.			must be protected
	Activity:	10-15 µg/ml will agglutinate type O human e	rythrocytes.			Abs
	Buffer:	0.05M Tris -0.3M NaCl, pH 7.5. Contains 0.0 Dilute with 0.05M Tris -0.1 NaCl, pH 7.5.	)5% sodium az	zide as a preservative.	Т	ITC RITC exas Red™
	Chemical Used for Conjugation:	Tetramethylrhodamine Isothiocyanate, TRITC	2.		Inhibition of lecti	n binding may be ac
	Storage:	Store liquid material frozen in aliquots in a freeze thaw cycles. Clarify by centrifugation.	mber vials or	covered with foil. Avoid	<ul><li>A. Before inc carbohydra</li><li>B. Preincubat</li></ul>	cubating with <b>Fluo</b> the for 30-60 minutes e diluted <b>Fluoresce</b> erature before apply
	Stability:	The liquid material is stable for at least 1 y 0.05% sodium azide added as a preservative.		ed frozen in aliquots with		erature before appry
	Caution:	Refer to the enclosed MSDS for information			Problem	1. Low concent
	Remarks:	seals have sharp edges and the vial itself m tions. Use caution when opening the vial. Fluorescent Conjugates are extremely lights	-	s which can cause lacera-	Weak or no Staining	oligosacchar 2. Low concent 3. Insufficient i
		high protein concentrations or the absence	of salts will	tend to cause aggregation		4. Photobleachi     1. Lectin conjug     2. Insufficient
	References	<ol> <li>Penberton, R.T.,(1970) Voxsang. 1874</li> <li>Miller, R.L., (1981) Fed. Proc. Abst. 40</li> </ol>	:1715.		High Background	3. Autofluoresc
	NO T				Unexpected Staining Pattern	Multiple causes
- All	<b>EY</b> LABORA 107 North Amphlet San Mateo, CA 94	<b>TORIES, INC.</b> tt Blvd. 401	Tel: Fax: Orders:	650-342-3296 650-342-2648 1-800-821-0044 (Outside CA only)		<b>BORATORII</b> mphlett Blvd. CA 94401

# **General Procedure Fluorescent Labeled Lectin**

The following is a general Procedure and Trouble-Shooting Guide. The information is provided only for our experiments are not guaranteed by EY Laboratories, Inc.

#### **Tissue Sections**

		Inssue Section	5	
1.	Wash and block tissue section. Do not use serum products, they contain glycoproteins which may lead to high levels of non specific background. After blocking, rinse briefly with Buffer (See reverse side).			
2.	Dilute Fluorescent Labeled Lectin to desired concentration 20-100 µg/ml using Buffer.			
3.	Incubate tissue section with Fluorescent Labeled Lectin for 30 minutes in a moist chamber.			
4.	Wash tissue	section with Buffer three times.		
5.	Examine tiss	ue section with Fluorescent microscope. Use	appropriate filter.	
	Ref. M. Imm	bar et. al., (1973). Intnl. Journal of Cancer, 12	2, 93-99	
		Cell Suspension	n	
1.	Wash cells with Buffer (See reverse side.)			
2.	Collect cells by centrifugation.			
3.	Dilute <b>Fluorescent Labeled Lectin</b> to $100 \mu$ g/ml using Buffer.			
4.	Incubate approximately $1 \times 10^6$ cells with 1 ml diluted Fluorescent labeled Lectin for 15 minutes at room temperature or in a 37°C water bath.			
5.	Wash cells with Buffer three times using centrifugation.			
6.	Examine cells, with or without fixation with Fluorescent microscope. Use appropriate filter.			
	Ref. K. Phiss. (1977). Experimental Pathology, 14, S15			
Flue	prochromes m	ust be protected from light. Perform incu	ibation, when practical, in a dark room or	
	ered in foil.		, , , , , , , , , , , , , , , , , , ,	
		Absorption and Emission	on	
		Absorption/Excitation Ra		
	FIT		517 nm	
	TRITC 554 nm		570 nm	
	Tex	xas Red <sup>™</sup> 596 nm	615 nm	
		Carbohydrate Inhib	bition	
Inhi	bition of lectin	binding may be accomplished by using one of	two procedures:	
A. B.	carbohydrate Preincubate	for 30-60 minutes at room temperature. NO	, incubate section or cells with inhibitory TE: Complete inhibition may NOT occur. nhibitory carbohydrate for 30-60 minutes at	
		TROUBLE SHOOTING	GUIDE	
	Problem	Cause	Solution	
		1. Low concentration of specific	Causes #1 - #3	
v	Weak or no	oligosaccharide on sample.	a. Increase incubation time.	
	Staining	<ol> <li>Low concentration of lectin conjugate.</li> <li>Insufficient incubation time.</li> </ol>	b. Increase concentration conjugate.	
		4. Photobleaching	a. Avoid exposure to light.	
		1. Lectin conjugate is too concentrated.	<ul> <li>a. Decrease concentration of Lectin conjugate.</li> </ul>	
		,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,	b. Shorten incubation times.	
		2. Insufficient washing.	a. Perform multiple washings and prolong	
1	High		washing time.	
Background		2 Autofluoroscont comple	a Use fluoreshrome with different excitation	

	<li>b. Shorten incubation times.</li>
ficient washing.	a. Perform multiple washings and prolong
	washing time.
fluorescent sample.	<ol> <li>Use fluorochrome with different excitation</li> </ol>
	and emission spectrum.
	<li>b. Use a different lectin conjugate (enzyme or</li>
	colloidal gold).
	a. Perform control reactions.
causes	b. Use other cytochemical technique to prove
	or disprove the findings.
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### MSDS for Fluorescent labeled Purified Proteins Continued - page 2 of 2.

## **MATERIAL SAFETY DATA SHEET**

Effective Date: March 31, 2006 Revision 4 Page 1 of 2

#### **PRODUCT IDENTIFICATION**

Purified proteins labeled with fluorescein isothiocyanate (FITC),
tetramethylrhodamine isothiocyanate (TRITC), or Texas Red a trademark of
Molecular Probes for the sulfonyl chloride derivative of sulforhodamine 101
FP-01, RP-01, TP-01, F-1102 to F-9000, R-1102 to R-9000, T-1102 to T-9000, FA-
2100 to FA-2701, RA-2100 to RA-2701, TA-2100 to TA-2701, FAF-001 to FAF-
2354, RAF-001 to RAF-2354, TAF-001 to TAF-2354, FAL-1104 to FAL-4701,
RAL-1104 to RAL-4701, TAL-1104 to TAL-4701, FA-01 to FA-013, TA-01 to
TA-013, DM1011F to DM1064F, FNP-01 to FNP-05, BA-101, BA-102, BA-612.
Protein A, Avidin (egg white), Glycosylated Bovine Serum Albumin, Lectins,
Secondary and Monoclonal Antibodies labeled with FITC, TRITC, or Texas Red®

#### **EMERGENCY INFORMATION**

EY Laboratories, Inc. 107 North Amphlett Blvd. San Mateo, CA 94401 EMERGENCY PHONE: 650-342-3296

#### HAZARDOUS COMPONENTS

Specific protein(s) as listed on the vial label. Solutions are at a concentration generally greater than 0.5mg protein/ml. Biological activity of these labeled proteins will vary. FITC, TRITC, and Texas Red® are possible carcinogens in their pure form. Compounds with similar chemical structures are known to be reactive with proteins and other biomolecules. The complete properties of the dyes after labeling have not been evaluated. These compounds should be treated as potentially hazardous. All solutions contain less than 0.05% sodium azide as a preservative.

#### HEALTH HAZARD INFORMATION

EXPOSURE LIMITS:	None established. The toxicological properties of these products have not
	been thoroughly investigated. Care should be taken when handling any of these materials.
	these materials.
EFFECTS OF	Causes localized eye, skin, or mucous membrane irritation. Some sensitive
OVEREXPOSURE:	individuals may develop a chronic allergic reaction with exposure. The
	known effects are due to the protein. No specific effects of the bound dye are known at this time.
ROUTES OF EXPOSURE:	Inhalation of powders and skin contact with liquids are the primary routes of exposure. Care should be taken to avoid the formation of aerosols when
	handling any of the solutions.

#### PHYSICAL CHARACTERISTICS

APPEARANCE: SOLUBILITY:

Powders are a light orange. Solutions will be yellow to dark purple. Powders are completely soluble in many biological buffers and water. I liquids are completely miscible in water and biological buffers.

## FIRE AND EXPLOSION HAZARDS

Not considered to be a vire hazard. At high concentrations the chemicals may emit toxic fumes. Such high concentrations are not normally found in a research laboratory.

EXTINGUISHING MEDIA: SPECIAL FIRE FIGHTING CRECAUTIONS:

Dry chemical powder or CO<sub>2</sub>. Wear self-contained breathing apparatus and protective clothing to prevent contact with skin and eyes.

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NOTE: Most solutions contain less than 0.05% sodium azide as a preservative. Azide may react with lead and copper plumbing to form explosive metal azides. Flush with copious amounts of water when disposing material in the sink.

# **REACTIVITY DATA**

STABILITY: HAZARDOUS POLYMER INCOMPATIBILITY:	ATION: hazardous. Vill NOT of Alcohols, si agents, and	Stable. Decomposition products are not known to be hazardous. Will NOT occur. Alcohols, strong bases and acids, strong oxidizing agents, and heat. (Lead and copper may react with sodium azide).	
<b>SPILL / LEAK PROCEDU</b> MATERIAL RELEASE / SPILL:	SE / Avoid contact with powder or liquid. Clean up spill with a paper towel soaked in household bleach. Do not allow solutions to dry on environmental surfaces. Wash affected area with detergent after the area		
WASTE DISPOSAL:	Local, State, and Federal reg	ose of paper waste in accordance with all gulations. Due to the small quantities of ucts are generally not considered to be	

#### **EMERGENCY FIRST AID PROCEDURES**

May be harmful if swallowed, inhaled, or allowed to absorb through the skin. Wash contacted area with water for 15 minutes. If inhaled remove to fresh air. Report exposure to the appropriate safety official. Consult a physician if irritation occurs or if there is any indication of an allergic response, such as watering eyes, sneezing, or difficulty breathing.

environmental hazards. All of these proteins are fully biodegradable.

#### SPECIAL HANDLING PRECAUTIONS

VENTILATION:	No special ventilation is required but it is recommended to handle these reagents in a fume hood when possible.
EYE PROTECTION:	Required. Goggles or safety glasses with a side shield are recommended.
RESPIRATORY	Recommended as a safety precaution, specifically when working with
PROTECTION:	powders. An approved respirator may be required for those individuals
	already known to be sensitive to these materials.
PROTECTIVE GLOVES:	Required when handling any of these materials.

#### SPECIAL PRECAUTIONS

This material is for research and experimental application only. It is not intended for food, drug, household, agricultural, or cosmetic use. All materials should be handled only by technically qualified individuals experienced with working with potentially hazardous chemicals. The above information is correct to the best of our knowledge. The user should make independent decisions regarding completeness of the information, based on all sources available. EY Laboratories, Inc. shall not be held liable for any damage resulting from handling or contact with the above product.



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