



CERTIFICATE OF ANALYSIS

PRODUCT NAME:	GLYKO® β (1-4,6)GALACTOSIDASE (Jack Bean)		
PRODUCT CODE:	GKX-5012		
LOT NUMBER:	DG33 022a		
FORMULATION:	Lyophilized from 20 mM sodium citrate phosphate (pH 6.0)		
RECONSTITUTION:	Dissolve the lyophilizate in 201 μ l of high purity water to obtain the described formulation. Since the enzyme will be more concentrated than is required for most applications, dilute further with buffer as needed.		
SUGGESTIONS FOR USE:	Conditions for use vary depending on the application, since the enzyme hydrolyzes β (1-4) and β (1-6) linkages significantly more rapidly than β (1-3) linkages. For example, for non-selective hydrolysis of non-reducing terminal galactose, a final concentration of at least 4 U/ml is recommended whereas a final concentration of <1 U/ml is recommended for selective hydrolysis of β (1-4)- and β (1-6)-linked galactose.		
STORAGE:	-20°C until redissolved. Store redissolved enzyme at 2-8°C or -20°C but avoid repeated freeze-thaw cycles.		
PACK SIZE:	5 Units		
EXPIRATION:	March 2019		
QUALITY CONTROL			
1. Specific activity ¹ :	Passed	(Specification: ≥ 70 U/mg)	
2. Protease assay ² :	Passed	(Specification: "Not Detectable")	
3. Contaminants ³ : (except as noted below)	Passed	(Specification: $\leq 0.001\%$)	
α -Galactosidase	0.0049%		
β -Glucosidase	0.0401%		
β -Xylosidase	0.0038%		

Authorized Signature

- One unit of Jack Bean β -Galactosidase is defined as the amount of enzyme required to catalyze the release of one μ mole of p-nitrophenol per minute from pNP- β -galactopyranoside at pH 3.5 and 37°C.
- No protease activity was detectable after incubation of the enzyme with 0.2 mg resorufin-labeled casein for ~18 hours at 37°C based on Schickaneder E, Hösel W, von der Eltz H, Geuß U. Casein-resorufin, a new substrate for a highly sensitive protease assay. Fresenius Z. Anal Chem. 1988 330:360.
- The absence of exoglycosidase contaminants was confirmed by extended incubations with the corresponding pNP-glycosides: α -fucosidase, α -mannosidase, β -mannosidase, β -N-acetylhexosaminidase, α -N-acetylgalactosaminidase, α -galactosidase, α -glucosidase, β -glucosidase and β -xylosidase. The absence of contaminating sialidase was confirmed by extended incubation with MU-NANA. **Note: this enzyme has activity on pNP- β -D-fucoside (although reduced relative to the standard substrate).**