

Identification and Characterization of Dextran α -1,2-Debranching Enzyme from *Microbacterium dextranolyticum*

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Supplementary Materials

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Table S1. Putative GHs detected by LC-MS/MS.

Accession	-10logP	Coverage (%)	No. of peptides	Signal peptide	Average mass	GH family	Description
MBM7461785.1	338.32	29	43	Yes	113904	GH9	endoglucanase
MBM7463898.1	206.35	13	15	Yes	131812	GH65	trehalose/maltose hydrolase-like predicted phosphorylase
MBM7461588.1	180.35	11	13	No	123490	GH97	hypothetical protein
MBM7463772.1	54.35	2	2	Yes	130172	GH92	putative alpha-1,2-mannosidase

1 MKPVSPTAGE PEMRRAPRRT RAAMGALLTA PLLVTGVLGA GAAAESGSA PTGANTSNGW EMATTDPFSA DYHPTFTGNG ■ Carbamidomethylation (+57.02)

81 NFAARVPAQG QGYSAADVAT QFQVAGLYAA HLPNEARASA PAWTGLTVSD GSGSFDQAFS APCVVGSICQ LEDAQLSGGV

161 SVASDHGGYQ GSGFTQGWGN ASAGALLYAN GAQPDAAYDL IVR**YAAGNPG DNSANVR**SLH VSVGDTSAI ALPPSPQGDW

241 DTWMQAR**LPL GALTDRALT LSCSSTPGAD CRVNVDSIAI VPRDGALPTT PPDLTMDAR**G LADYRQSLDV STGAIITSAR

321 WTSPSGNVSD VAYTVLTD RG HDDRALVRVD VTPQWSGDLE VTDVLDTRPA TFVAGTVTHR DDAAGTIGVD AAITGSGITA

401 SIASTLAGPG QR**AEARPVGL PSGSIAQK**LT QSVESGKTY S FVKYVGLTT HDGADTAGLA **KAASTAAAQ GYDAVRASND**

481 **GAWADIWK**GD IQVTGDDALQ QQIRASRFYL LASVNADRPW SPSPAGLSSD GYGGHVFDW ETWMWPSLVA QDPEIAAAVL

561 KYRSDRIDDA RANAAAGHD GIRFPWEGAL DGTEQTATL FGETEQHITA DVALAFWQYY LSTGDRTWLQ NEGWPVISGA

641 ADFWASRVEL GDDGQYHNG VTPPDEWAGK HDSAYTNVA AAQTMRMATR **AAGVLGTSAK PAWQTIADK**M FMPHDDALGI

721 TPEYAGYNHG VIKQADVMT TYPWGYEQSD ELTAR**NLAYY ASR**VTTSGGP SMTDAIHSIV SAELGNVCDA WYYTQQSGTP

801 **FMRAPFDQFS EER**EGGAFTT TTGAGGLQE FYYGYTGLRM QEDGITLAPI LPPSLTDMTV AGLHYQGRTF DVKIGTTSTD

881 VTLTSGPALT VHTADGDKTV STDAALRIPT RTANDCEATS GYGTLLGSL S APAGGDNGAG TLQYPGSSNF PTGTFDMTDF

961 DVYRDGDTLR VYTVSGEIT NPWGGNGMSV QLLHTYLRLP SNDSAPARSG AVPALPGTNA DLESPWDLVL VGNGRATGSG

1041 PGTGLYDASG ALVAVVELSV TPRRHQIVLS FPESALKGAD FVKVGYVAAM LVNAESNEGL SNIRPALDCE APGSPDWVSQ

1121 WRICGGLGTI SSASPYD TDT RDPNIIK**VFV PEGR**TQQGDV LK**TEGPSILP FVALK**AAPGP SPSPSDSGSA SPNPSDSGSA

1201 SPSPSDSGPG SPSASASAP TTSGAAGGP SADRLGATGG ESPLPLAVIG GVLLLLGAAG VLMRRRSLRR DGA

Supporting Peptides:

Peptide	Uniq	-10lgP	Mass	ppm	m/z	z	RT	Scan	#Spec	Start	End	PTM
K.AASTAAAQGYDAVR.A	Y	82.23	1421.6898	31.9	711.8749	2	-	532	2	462	476	
R.AALTLSG(+57.02)SSTPGADC(+57.02)R.V	Y	71.23	1665.7450	27.9	833.9030	2	-	569	1	257	272	Carbamidomethylation
R.VNVDSIAIVPR.D	Y	63.76	1181.6768	27.5	591.8619	2	-	655	1	273	283	
R.YAAGNPGDNSANVR.S	Y	57.69	1404.6382	28.8	703.3466	2	-	466	1	204	217	
R.ASNDGAWADIWK.G	Y	55.54	1332.6099	27.6	667.3306	2	-	710	1	477	488	
R.LPLGALTDR.A	Y	55.52	954.5498	27.3	478.2952	2	-	615	1	248	256	
R.AAGVLGTSAKPAWQTIADK.M	Y	54.21	1884.0105	30.2	629.0297	3	-	657	1	691	709	
K.TEGPSILPFVALK.A	Y	49.94	1370.7809	26.7	686.4160	2	-	749	1	1163	1175	
R.APFDFSEER.E	Y	40.92	1224.5410	31.2	613.2969	2	-	598	1	804	813	
L.PSGSIAQK.L	Y	38.90	786.4235	30.4	394.2310	2	-	558	1	421	428	
R.DGALPTTTPDLTMDAR.G	Y	37.46	1669.7981	27.3	835.9291	2	-	671	1	284	299	
R.AEARPVGLPSGSIAQK.L	Y	32.12	1579.8682	34.2	527.6480	3	-	557	1	413	428	
R.NLAYYASR.V	Y	19.27	956.4716	30.6	479.2577	2	-	531	1	756	763	
K.PAWQTIADK.M	Y	19.14	1028.5291	29.9	515.2872	2	-	579	1	701	709	
K.VFVPEGR.T	Y	16.38	802.4337	32.1	402.2370	2	-	501	1	1148	1154	
total 15 peptides												

Figure S1. Peptides of *M. dextranolyticum* GH65 protein (MddDE) identified by LC-MS/MS. Coverage map (*top*) and properties (*bottom*) of MddDE-derived peptides identified by LC-MS/MS are shown.