

Conserved sequence motifs in levansucrases and bifunctional β -xylosidases and α -L-arabinases

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Abstract Comparison of the amino acid sequences of two families of glycosyl hydrolases reveals that they are related in a region in the central part of the sequences. One of these families (GH family 68) includes levansucrases and the other one (glycosyl hydrolase family 43) includes bifunctional β -xylosidases and α -L-arabinofuranosidases. The similarity of the primary structure of proteins from these families allows us to consider the invariant glutamate residue as a component of their active center. It is shown for the first time that glycosyl hydrolases recognizing different glycofuranoside residues can have a common sequence motif.

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Key words: Glycosyl hydrolase; β -Fructosidase superfamily; Levansucrase; β -Xylosidase; Multiple sequence alignment

1. Introduction

Levansucrases are a family of bacterial enzymes with a β -fructofuranosyl fructotransferase activity (EC 2.4.1.10). They can produce different types of D-fructose poly- and oligomers (levans etc.) from sucrose. The production process often takes place on the early steps of the plant colonization by various symbiotic and phytopathogenic bacteria [1,2]. As well as polyglucans, fructose polymers are probably implicated in the carries process in humans [3,4]. In addition to the fructosyl transferase activity, levansucrases can hydrolyze sucrose and various fructose polymers (levans, inulin) into monosaccharides. So they also have β -fructofuranosyl fructohydrolase activities [5]. This allows us to consider levansucrases as a family of glycosyl hydrolases (GHs) (family 68) [6,7]. By sequence similarity, this family also includes an extracellular invertase of *Zymomonas mobilis*, which has no fructosyl transferase activity [5–7].

Recently, we have shown that levansucrases have a significant sequence similarity with enzymes of the GH family 32 (sucrases and fructanases) [5]. Enzymes of the two protein families have the same molecular mechanism of hydrolyzing reaction: a double displacement with an overall retention of the anomeric configuration of the fructosyl residue [6,7]. Due to these facts, the two families of GHs have been grouped into the β -fructosidase superfamily [5]. Comparison of the most conserved sites of sequences of the two families of β -fructosidases allowed us to propose several possible components of the active center.

In this work, we compare sequences of levansucrases and GHs from family 43. Enzymes of the latter family usually have β -xylosidase (EC 3.2.1.37) and α -L-arabinofuranosidase (EC 3.2.1.55) activities [6–9]. These enzymes participate in the degradation of plant cell wall material in the rumina of mammals by anaerobic microorganisms [9].

2. Materials and methods

Protein sequences were retrieved from the current sequence databases. GHs compared in this work are listed in Table 1. The alignment of proteins was generated by using the PSI-BLAST program [10]. The program MACAW [11] was used to estimate the probabilities of the independent appearance of the regions of local similarity in the two protein families.

3. Results and discussion

An alignment by using the PSI-BLAST and MACAW programs shows that GHs from families 43 and 68 are related in a region in the central part of the sequences. In this region, there are two highly conserved blocks (Fig. 1) with a probability of obtaining the observed level of similarity by chance (P value) below 10^{-19} (search space $N = 2.7 \times 10^{86}$) for each of them. This low P value results in part from the high similarity between the members of each family. However, for the pairwise comparison of AF005383 of *Caldicellulosiruptor saccharolyticus* and Q43998 of *Acetobacter diazotrophicus* in the first block (sequence E-g/r-s/p-h/q-a/v-Y-k/l-i/h-N-G-K-Y-Y-I-t/f-T-I), the P value is 1.7×10^{-5} ($N = 289$), showing that the relationship is authentic.

The first conserved block partly overlaps with known conserved regions of both GH family 43 (the second conserved region according to Sakamoto et al. [8]) and levansucrases (conserved region VII according to Arrieta et al. [12]). This block includes a glutamate residue invariant in both families and a conserved cluster of tyrosine residues (Fig. 1). The sequence between these two elements is highly conserved and corresponds to the pattern: E-x-(SPA)-x(1,2)-(MILAVFY)-x(3)-(NDGA)-(DGNTTE)-x-(YWT)-(YV)-(LFYI). It should be noted that the corresponding site of GH family 32 is also conserved (region VI according to Bezzate et al. [1]) and includes the invariant glutamate residue. In the case of *Saccharomyces cerevisiae* invertase, this Glu was shown to serve as a proton donor in the hydrolysis of the glycosidic bond [13]. However, only one (invertase of *Tritrichomonas foetus*) out of more than 100 sequences in GH family 32 follows the pattern described above. This is a consequence of missing the tyrosine cluster in enzymes of the family. However, the similarity of structures of the first conserved block and the corresponding site of sucrases and fructanases allows us to suppose that the invariant Glu residue is

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block I										block II																			
*****										*****																			
P49943	210	KPLTAGDTERRE	FEASW-MHKYNGKYYE	SYSTGDTHL	-----	-----	-----	-----	-----	LCYATGDN	BYGPFT	----	YQGVILTPV	---	56														
P48791	269	KPLPA-DDPHRR	FEASW-MHKYNGKYYE	SYSTGDTHL	-----	-----	-----	-----	-----	LCYAVGDN	BYGPFT	----	YQGVILEPV	---	52														
AF095243	212	KVMQAGST-RR	FEAGPS-LNKVGNLY	VLOYSTGTSHT	-----	-----	-----	-----	-----	VEIATATK	PDGPYY	----	WKSTLLQPV	---	58														
P48790	195	YSRGSFEGEH	FEAPS-IRKKGDT	YYFTYSSVMHE	-----	-----	-----	-----	-----	LCYATSKH	PTKGFK	----	YGGVIVSNCD	---	218														
JQ1936	195	YSRGSFEGEH	FEAPS-IRKKGDT	YYFTYSSVMHE	-----	-----	-----	-----	-----	LCYATSKH	PTKGFK	----	YGGVIVSNCD	---	218														
AF005383	219	-----	KIDGLEBGS	HAYKINGKYYIT	IRWEKGGIREE	-----	-----	-----	-----	CVYRADK	IDGPYE	----	GRVVLSDTL	---	1076														
AF005383	1081	-----	PAPFMFDSG	-INKIGNTYYY	SYCTNFYSGAR	PQG-----	-----	-----	-----	SPPAGVIA	YMTSKS	----	EMGPWE	----	YKGVILKNPG	---	202												
O52374	1515	-----	PAPYMFDSG	-INKIGNTYYY	SYCTNFAQ	-RPQG-----	-----	-----	-----	SPPAGAI	YMTSKS	----	EMGPWE	----	YKGVILKNPG	---	202												
P45796	241	-----	DAPYLFDSG	-IHKYNGKYYY	SYCINFAGT	-HPQQ-----	-----	-----	-----	YP-AGEI	GYMVS	DN-	EMGPFT	----	YKGVILKNPY	---	332												
Q45071	244	-----	DAPFMFDSG	-LHKYNGTYYY	SYCINFGGT	-HPAD-----	-----	-----	-----	KPP-GEIG	YMTSSS	----	PMGPFT	----	YRGHFLKNPG	---	207												
P95470	207	VLMDDSQAGS	QAQIEAPF-IL	RKGDYYYLF	ASWGLCCR	KGDST-----	-----	-----	-----	YHLVVGR	SKQ	----	VTGPYLDK	----	TGRDNQGGG	---	71												
P94522	210	-----	GGALBAPT	-LTYONGY	YYLMSV	FDKCCD	GVNST-----	-----	-----	YKIA	YGRSKS	----	ITGPYLDK	----	SGKSMLEGGG	---	43												
O07078	211	-----	NGALBAPT	-LTYONGY	YYLMSV	FDKCCD	GVNST-----	-----	-----	YKIA	YGRSKS	----	ITGPYLDK	----	SGKSMLEGGG	---	53												
P42256	195	-----	THAEGSY	-MFCYGDY	YYLFS	SAGICCGY	DTSM-----	-----	-----	PASGEE	YHIK	VCR	STSS	----	PTGDFVDS	----	DGTACTDGGG	---	60										
P42293	219	-----	HSRIEG	GVLYNP	DTQYYLY	LYSYGGL	DATGGY-----	-----	-----	NIRVAR	SKK	----	PDGPYYDA	----	EGNPMLDVRG	---	190												
P94489	173	KVIFEG-TDR	KLTAAPH-LY	HIGNY	YLLTA	AEAGGTRY	EHAAT-----	-----	-----	IARSA	N	----	IEGPYEVH	----	PDNPILTSWHP	---	294												
P07129	173	KVIFKG-TPI	KLTAAPH-LY	IINDY	YLLTA	AEAGGTRY	EHAAT-----	-----	-----	IARSS	R	----	IDGPYEVH	----	PDNPILTSWHP	---	296												
AF107211	173	KVIFKG-TPI	KLTAAPH-LY	HIGNY	YLLTA	AEAGGTRY	EHAAT-----	-----	-----	IARSS	H	----	IEGPYEVH	----	PDNPILTSWHP	---	296												
O52729	173	KIIFEG-TDI	KLTAAPH-LY	IINDY	YLLTA	AEAGGTRY	EHAAT-----	-----	-----	IARSK	H	----	IEGPYEVH	----	PDNPILTSWHP	---	296												
AF040720	173	EIIYKG-TD	IATBGP-LY	IINDY	YLLTA	AEAGGTRY	QHSET-----	-----	-----	IARSK	T	----	IEGPYEVH	----	PDNPILTSWHP	---	299												
P77713	173	KTIFTG-TPL	CYTBGAH-LY	RHAGY	YLLMA	AEAGGTRY	EHAAT-----	-----	-----	VLRSK	N	----	IDGPYEVH	----	PDNPILTSWHP	---	297												
P45982	171	AIWKALKD	VILWPGPH-LY	KKDG	YLLMA	AEAGGTRY	EHAAT-----	-----	-----	VARSKE	L	----	IEGPYEVH	----	PDNPILTSWHP	---	279												
consensus										E.Pz-&j&bG.YY&u										o.z-z&GzPz...zG&uu									
P05655	328	KPLIASNTVT	DELEBRAN-VFKM	NGKYYL	ETDSR	SGSKMTI	-----	-----	-----	DGITS	NDI	YML	GYV	SNS	----	LTGPYKPLN	----	KTGLVLKMDLD	---	69									
P94468	328	KPLIASNTVT	DELEBRAN-VFKM	NGKYYL	ETDSR	SGSKMTI	-----	-----	-----	DGITS	NDI	YML	GYV	SNS	----	LTGPYKPLN	----	KTGLVLKMDLD	---	69									
P21130	328	KPLIASNTVT	DELEBRAN-VFKM	NGKYYL	ETDSR	SGSKMTI	-----	-----	-----	DGINS	NDI	YML	GYV	SNS	----	LTGPYKPLN	----	KTGLVLKMDLD	---	68									
AB010272	346	KPLVASNTVA	DEBRAN-IFKM	MNKYYL	ETDSR	SGSKMTS	-----	-----	-----	DGIND	KDV	YML	GP	GDS	----	LNGP	PNP	IN	ETGL	VNM	NLD	---	65						
P11701	487	TPLLSS	TMVSDLEBRN-VV	KLGD	KYYL	ETASRL	NHGS	NNDAWN	KANEV	VGD	NV	ML	GYV	SDQ	----	LTNGYKPLN	----	NSGV	VLT	ASV	-P	228							
Q55242	533	TPLVTS	HMTVDEBRPS-VV	KMGK	KYYL	ETASR	INKST	DAEGT	VAA	ARE	AVG	DD	V	ML	G	FV	S	----	LRGK	YR	PLN	-G	SGV	VLT	ASV	-P	354		
Q43998	387	SPLISAN	CVNDQTERPO-VY	LHNG	KYYL	ETISH	-RTTF	-----	-----	AAGV	D	GP	D	GV	-Y	CF	V	GD	----	IRSD	EP	MNY	GS	GL	TM	---	125		
Q46654	273	PPLITAV	GVNDQTERPH-FV	QD	KYYL	ETISH	-KFTY	-----	-----	ADNLT	G	P	D	GV	-Y	CF	V	SD	----	LTGP	Y	PMN	-S	GL	V	---	71		
O54435	273	PPLVTA	GVNDQTERPH-FV	QD	KYYL	ETISH	-KFTY	-----	-----	ADGLT	G	P	D	GV	-Y	CF	V	SD	----	LTGP	Y	PMN	-S	GL	V	---	71		
O52408	273	PPLVTA	GVNDQTERPH-FV	QD	KYYL	ETISH	-KFTY	-----	-----	ADGLT	G	P	D	GV	-Y	CF	V	SD	----	LTGP	Y	PMN	-S	GL	V	---	71		
O68609	289	PPLVTA	GVNDQTERPH-FV	QD	KYYL	ETISH	-KFTY	-----	-----	ADGLT	G	P	D	GV	-Y	CF	V	SD	----	LTGP	Y	PMN	-S	GL	V	---	71		
Q60114	264	PPLVTA	GVNDQTERPH-FV	QD	KYYL	ETISH	-HSTY	-----	-----	ADGLS	G	P	D	GV	-Y	CF	V	SD	----	LTGP	Y	PMN	-S	GL	V	---	87		
Q60115	262	PPLLTAF	GVNDQTERPH-VI	FQ	NG	LYL	ETISH	-DSTY	-----	-----	ADGLT	G	S	D	L	-Y	CF	V	SD	----	LTGP	Y	PMN	-S	GL	V	---	79	
O02490	284	GCHDGV	WECPA-VLK	V	D	G	KWV	ILN	242																				
P00724	215	GFLGYQ	YECFG-LI	E	V	P	T	E	Q	D	S	K	S	293															

Fig. 1. Multiple alignment of the region of similarity in GH families 43 and 68. The distance from the protein termini is indicated. At the top of the figure, the two conserved blocks described in the paper are indicated by asterisks. The consensus shows residues, which are conserved in the majority of sequences of both families: u indicates a bulky hydrophobic residue (I, L, V, F or M); @ indicates an aromatic residue (F, Y or W); & indicates a hydrophobic residue (I, L, V, F, M, Y, W, A, C or P); j indicates a positively charged residue (K, R or H); b indicates N or D; o indicates a small residue (G, S or A); z indicates a polar residue (D, E, N, Q, H, K, R, S or T). The residues that are conform to the consensus are highlighted. In the consensus, the sequence corresponding to the pattern of GH families 43 and 68 (see text) is underlined. The regions known to be conserved in GH families 43 [8], 68 [12] and 32 [1] are boxed. The sequences are from the current sequence databases with the accession number for each sequence indicated in the most left column (for origin of the sequences see Table 1).

Table 1
Glycosyl hydrolases analyzed in this work

Family number ^a	Enzyme	Organism	Length	GenBank accession number	SWISS-PROT accession number
43	β-Xylosidase and arabinofuranosidase	<i>Bacteroides ovatus</i>	325	U04957	P49943
43	β-Xylosidase	<i>Prevotella ruminicola</i>	319	Z49241	P48791
43	β-Xylosidase	<i>Cochliobolus carbonum</i>	328	AF095243	
43	β-Xylosidase and arabinofuranosidase	<i>Clostridium stercorarium</i>	473	D13268	P48790
43	β-Xylosidase and arabinofuranosidase	<i>Clostridium stercorarium</i>	473	JQ1936	
43 ^b	β-xylosidase and arabinofuranosidase	<i>Caldicellulosiruptor saccharolyticus</i>	1347	AF005383	
43/10 ^c	β-Xylanase	<i>Caldicellulosiruptor sp. Rt69B.1</i>	1779	AF036924	O52374
43	β-Xylanase	<i>Paenibacillus polymyxa</i>	635	X57094	P45796
43	β-Xylanase	<i>Bacillus subtilis</i>	513	Z99113	Q45071
43	α-L-Arabinase	<i>Pseudomonas fluorescens</i>	347	Y10458	P95470
43	α-L-Arabinase	<i>Bacillus subtilis</i>	313	Z75208	P94522
43	α-L-Arabinase	<i>Bacillus subtilis</i>	324	D85132	O07078
43	α-L-Arabinase	<i>Aspergillus niger</i>	321	L23430	P42256
43	ORF	<i>Bacillus subtilis</i>	469	D31856	P42293
43	β-Xylosidase	<i>Bacillus subtilis</i>	533	U66480	P94489
43	β-Xylosidase	<i>Bacillus pumilus</i> IPO	535	X05793	P07129
43	β-Xylosidase	<i>Bacillus pumilus</i> PLS	535	AF107211	
43	β-Xylosidase	<i>Bacillus sp. KK-1</i>	533	AF045479	O52729
43	β-Xylosidase and arabinofuranosidase	<i>Selenomonas ruminantium</i> GA192	583	AF040720	O52575

Table 1 (continued)
Glycosyl hydrolases analyzed in this work

Family number ^a	Enzyme	Organism	Length	GenBank accession number	SWISS-PROT accession number
43	β -Xylosidase	<i>Escherichia coli</i>	536	U70214	P77713
43	β -Xylosidase and arabinofuranosidase	<i>Butyrivibrio fibrisolvens</i>	517	M55537	P45982
68	Levansucrase	<i>Bacillus subtilis</i>	473	Z99121	P05655
68	Levansucrase	<i>Bacillus stearothermophilus</i>	473	U34874	P94468
68	Levansucrase	<i>Bacillus amyloliquefaciens</i>	472	X52988	P21130
68	Levansucrase	<i>Bacillus sp. V230</i>	487	AB010272	
68	Levansucrase	<i>Streptococcus mutans</i>	797	M18954	P11701
68	Levansucrase	<i>Streptococcus salivarius</i>	969	L08445	Q55242
68	Levansucrase	<i>Acetobacter diazotrophicus</i>	584	L41732	Q43998
68	Levansucrase	<i>Erwinia amylovora</i>	415	X75079	Q46654
68	Levansucrase	<i>Rahnella aquatilis</i>	415	U91484	O54435
68	Levansucrase	<i>Pseudomonas syringae pv. glycinea</i>	415	AF037443	O52408
68	Levansucrase	<i>Pseudomonas syringae pv. phaseolicola</i>	431	AF052289	O68609
68	Levansucrase	<i>Zymomonas mobilis</i>	423	L33402	Q60114
68	Invertase	<i>Zymomonas mobilis</i>	413	L33403	Q60115
32	Invertase	<i>Tritrichomonas foetus</i>	550	U66071	O02490
32	Invertase	<i>Saccharomyces cerevisiae</i>	532	Z46921	P00724

^aAccording to the classification of GHs [6,7].

^bThe protein consists of two homologous domains (see Fig. 1).

^cThe C-terminal domain belongs to family 43 and the N-terminal domain belongs to family 10 [6,7].

a component of the active center in enzymes of GH families 43 and 68. The conserved tyrosine cluster and preceding glycine residue (Fig. 1) also can be essential for the enzymatic activities.

The similarity found between the sequences of enzymes with β -D-fructofuranosidase and α -L-arabinofuranosidase activities is the first case showing that GHs recognizing different glycofuranoside residues can have a common sequence motif.

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