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(54) **METHOD AND COMPOSITIONS FOR IMPROVED LIGNOCELLULOSIC MATERIAL HYDROLYSIS**

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(65) **Prior Publication Data**

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Related U.S. Application Data

(60) Provisional application No. 61/579,897, filed on Dec. 23, 2011, provisional application No. 61/579,301, filed on Dec. 22, 2011.

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(51) **Int. Cl.**

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C12N 9/42 (2006.01)
C12N 1/20 (2006.01)
C12N 9/24 (2006.01)
C12P 19/02 (2006.01)
A23K 10/12 (2016.01)

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(52) **U.S. Cl.**

CPC **C12P 19/14** (2013.01); **A23K 10/12** (2016.05); **C12N 1/20** (2013.01); **C12N 9/2434** (2013.01); **C12N 9/2437** (2013.01); **C12N 9/2491** (2013.01); **C12P 19/02** (2013.01); **C12Y 302/01004** (2013.01); **C12Y 302/01025** (2013.01); **C12Y 302/01091** (2013.01)

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(58) **Field of Classification Search**

CPC . C12P 19/14; C12P 19/02; C12N 1/20; C12N 9/3491; C12N 9/92434; C12N 9/2437; A23K 10/12; C12Y 302/01091; C12Y 302/01004; C12Y 302/01025

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(57) **ABSTRACT**

A method of digesting a lignocellulosic material is disclosed. In one embodiment, the method comprises the step of exposing the material to an effective amount of *Streptomyces* sp. ActE secretome such that at least partial lignocellulosic digestion occurs.

7 Claims, 70 Drawing Sheets
(22 of 70 Drawing Sheet(s) Filed in Color)
Specification includes a Sequence Listing.

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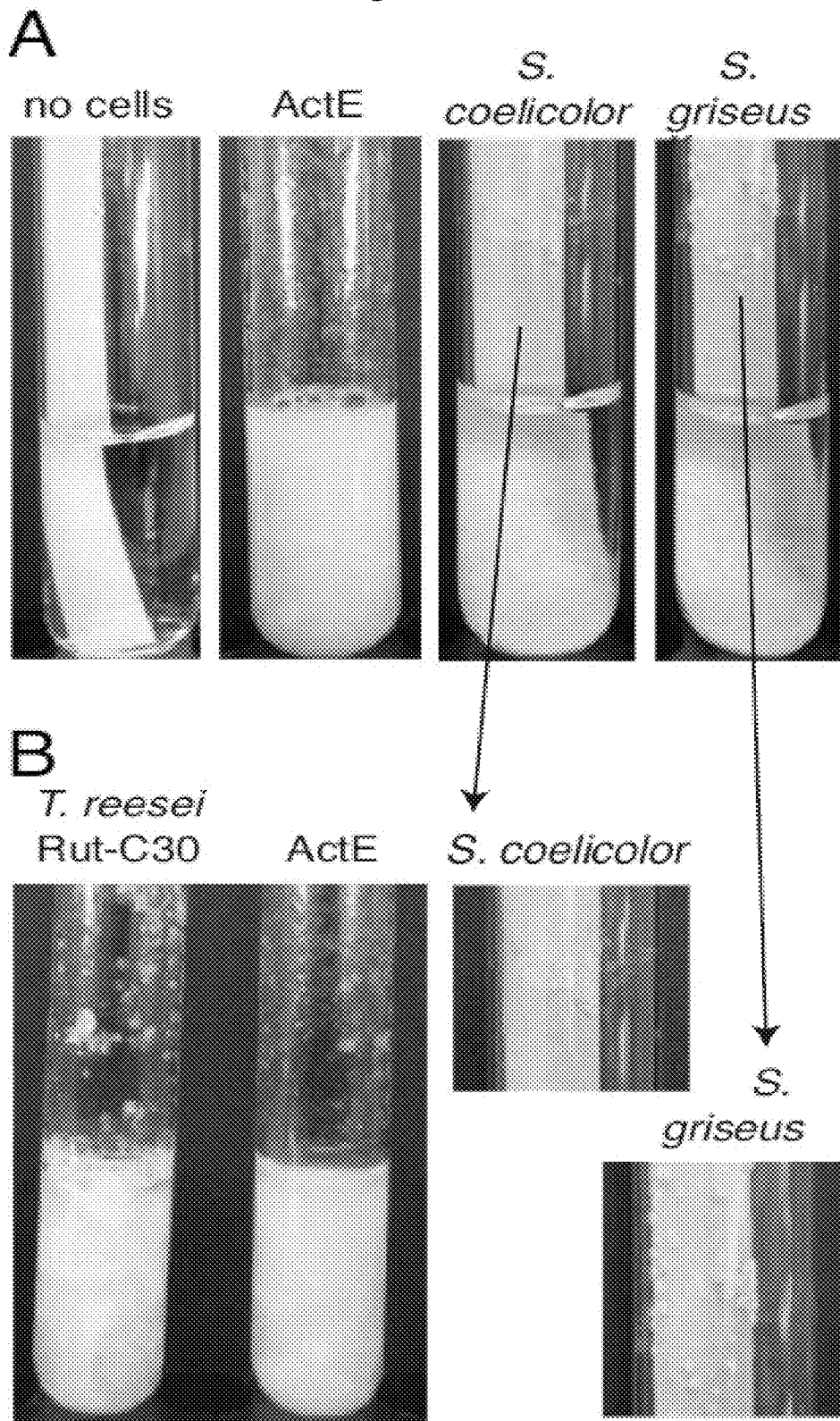
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Figure 1



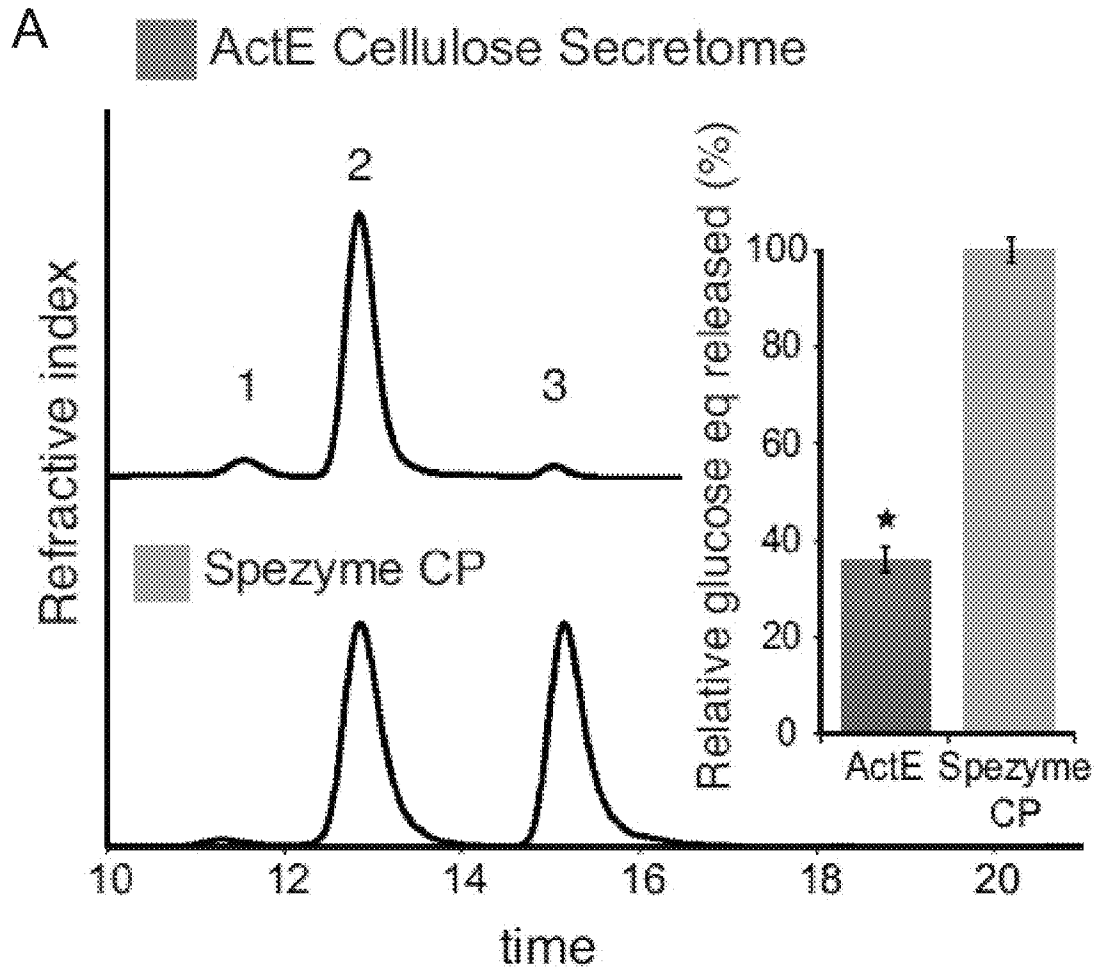


Figure 2

B

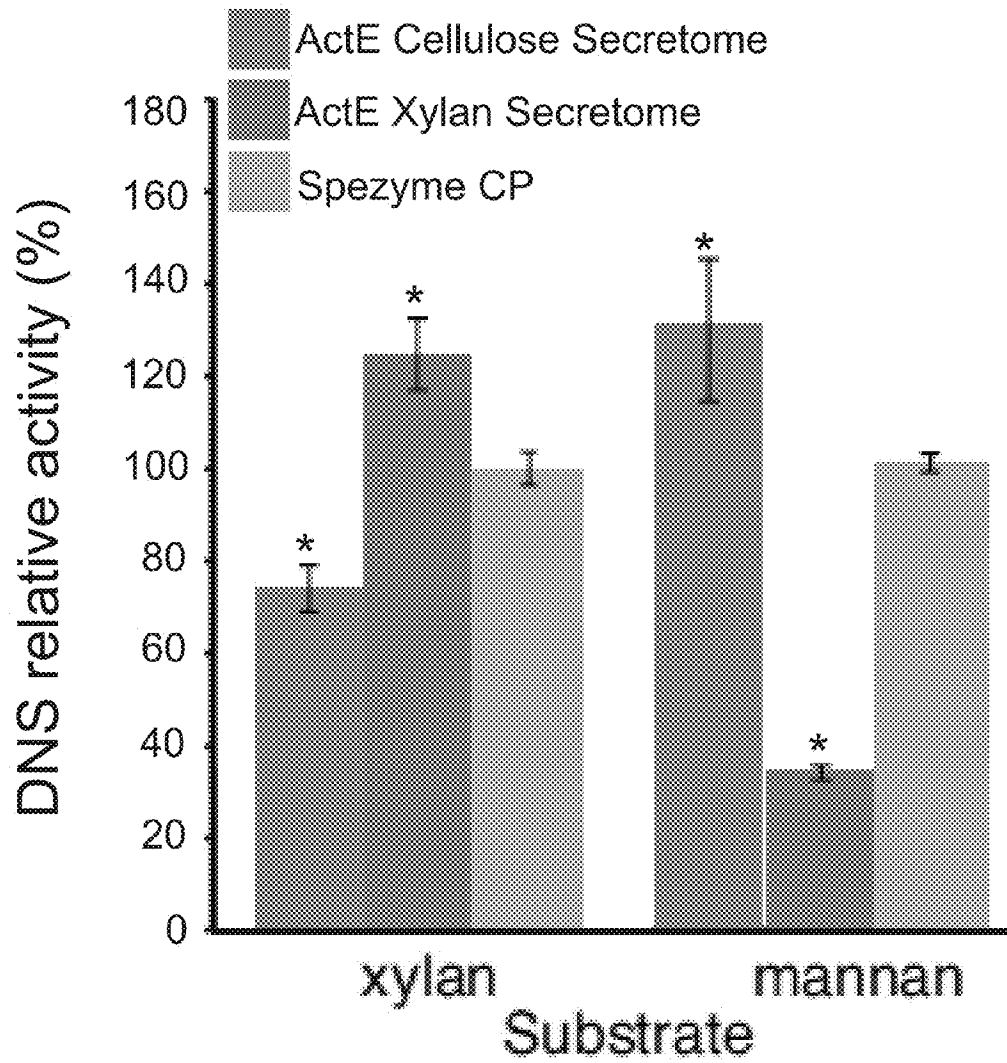


Figure 2

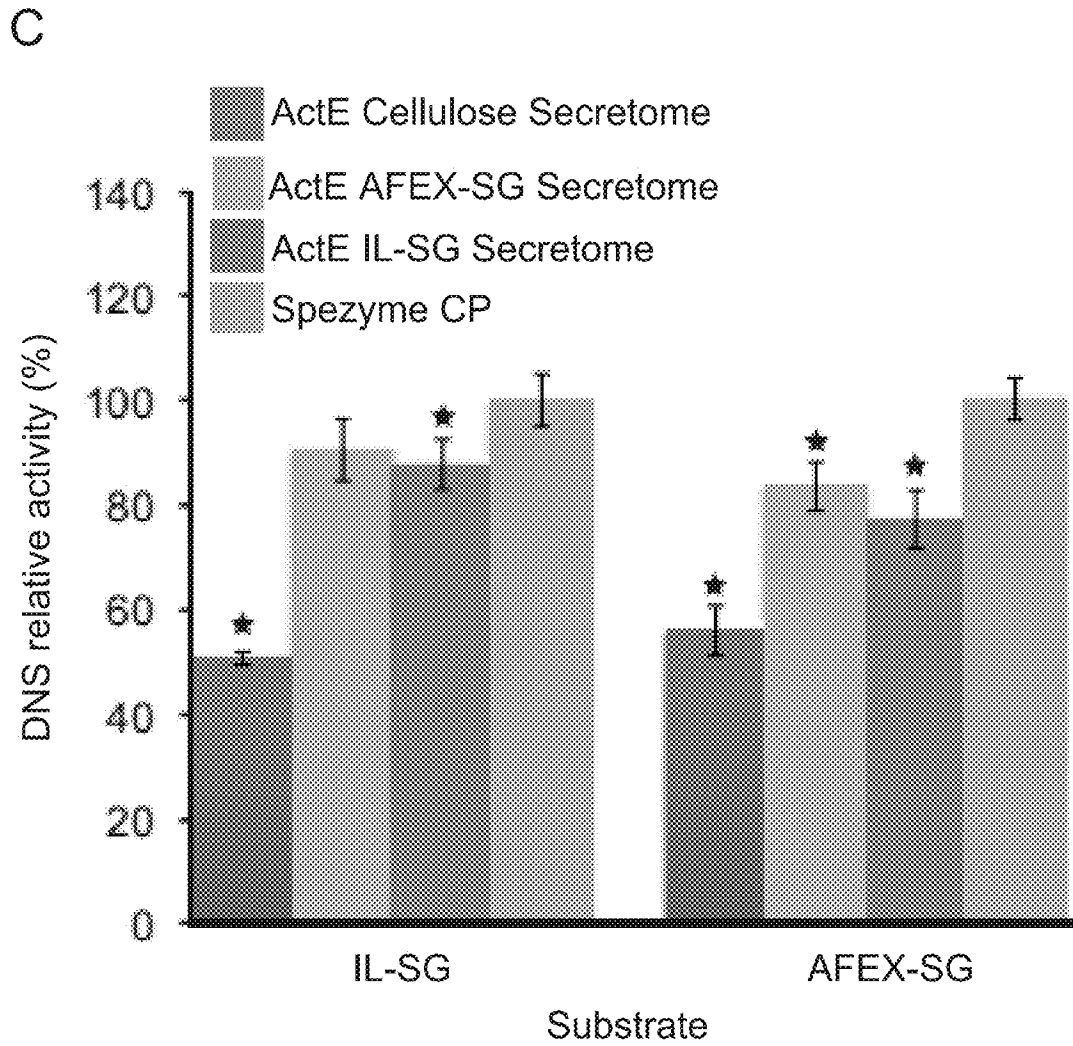
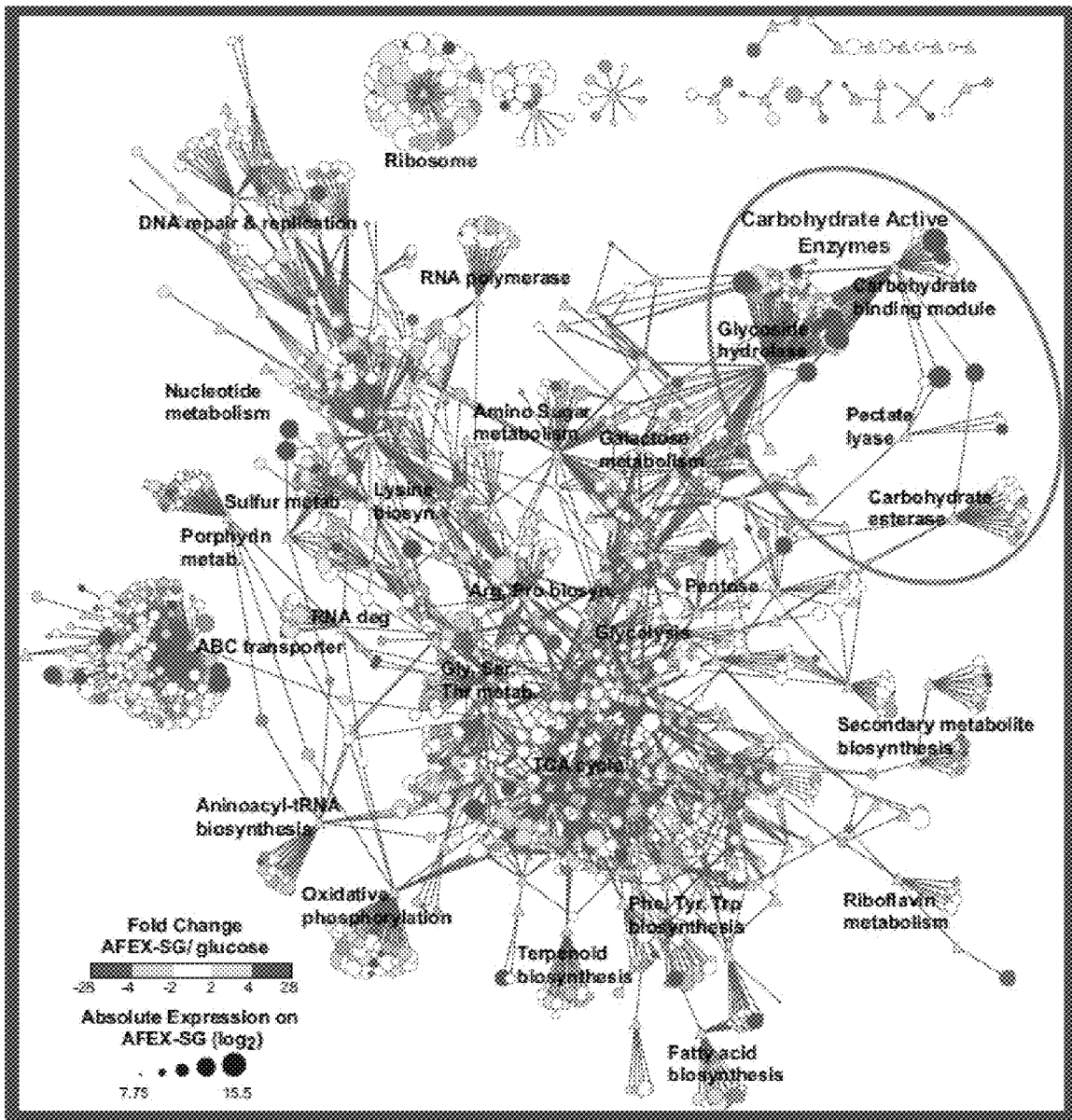


Figure 2

Figure 4



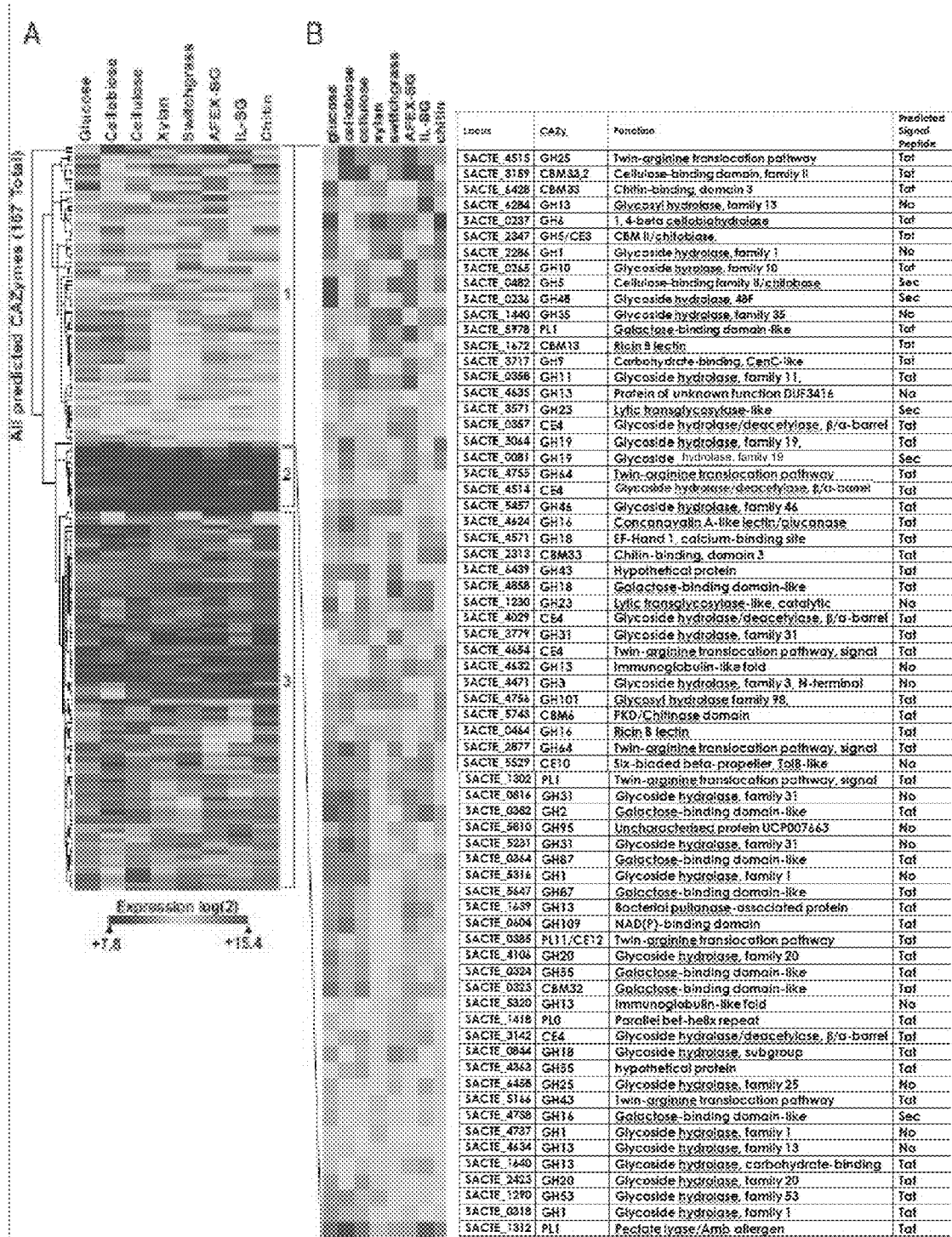


Figure 5 A-B

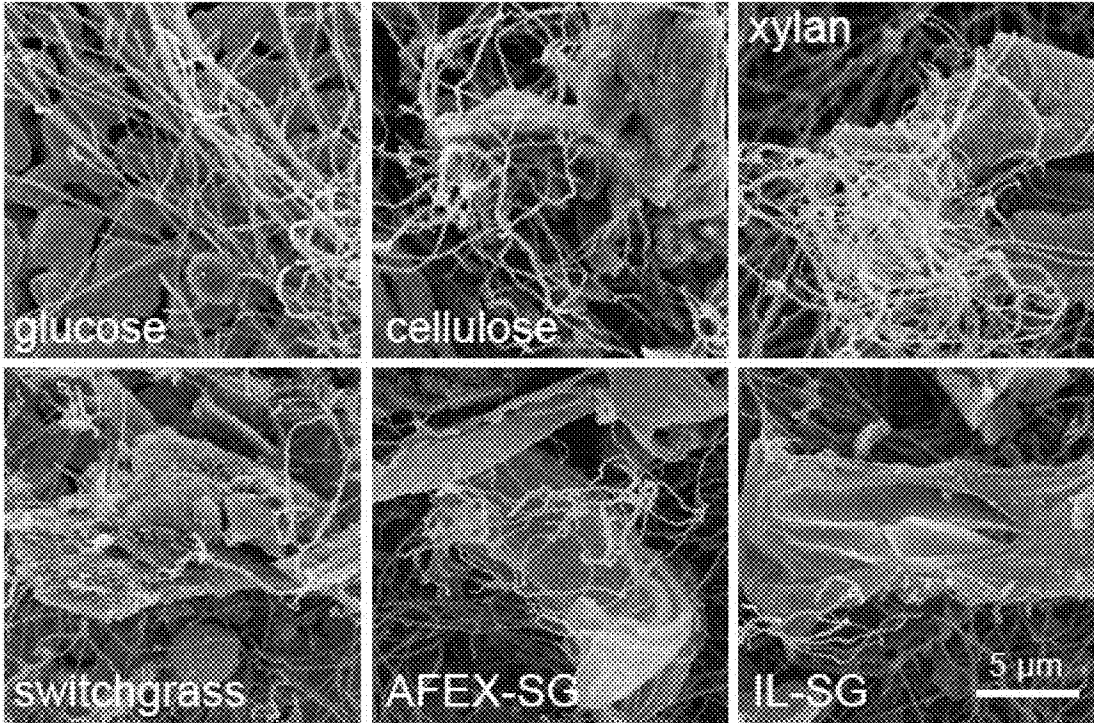


Figure 6

Figure 7A

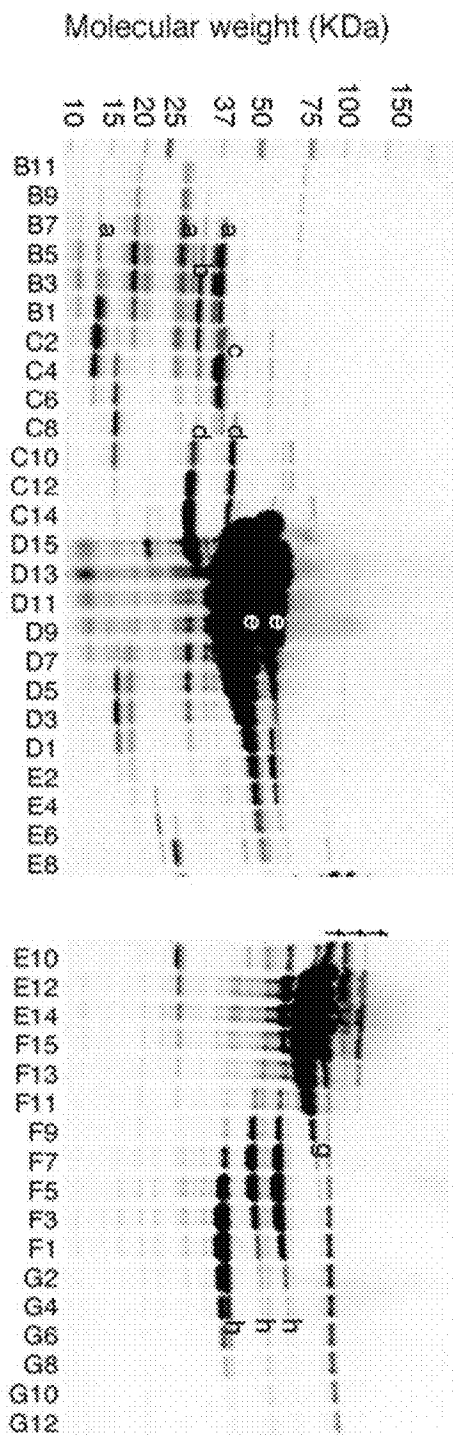
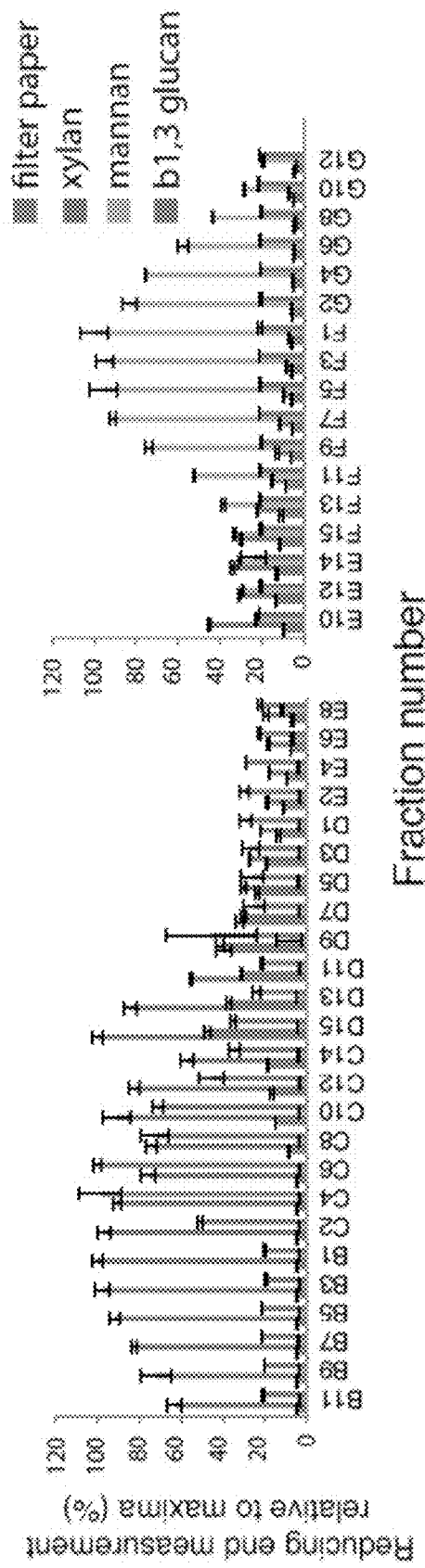


Figure 7B



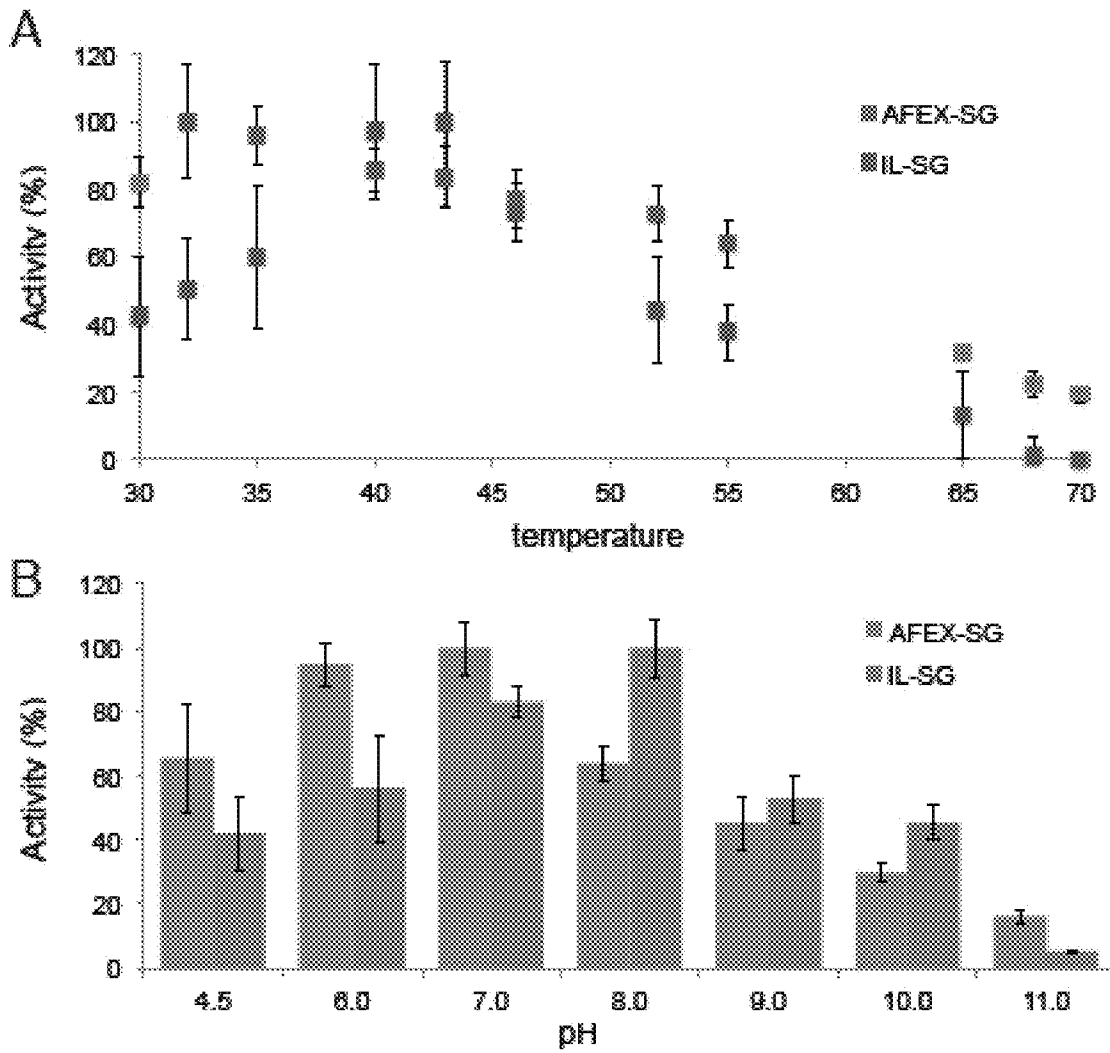


Figure 8 A-B

Figure 9

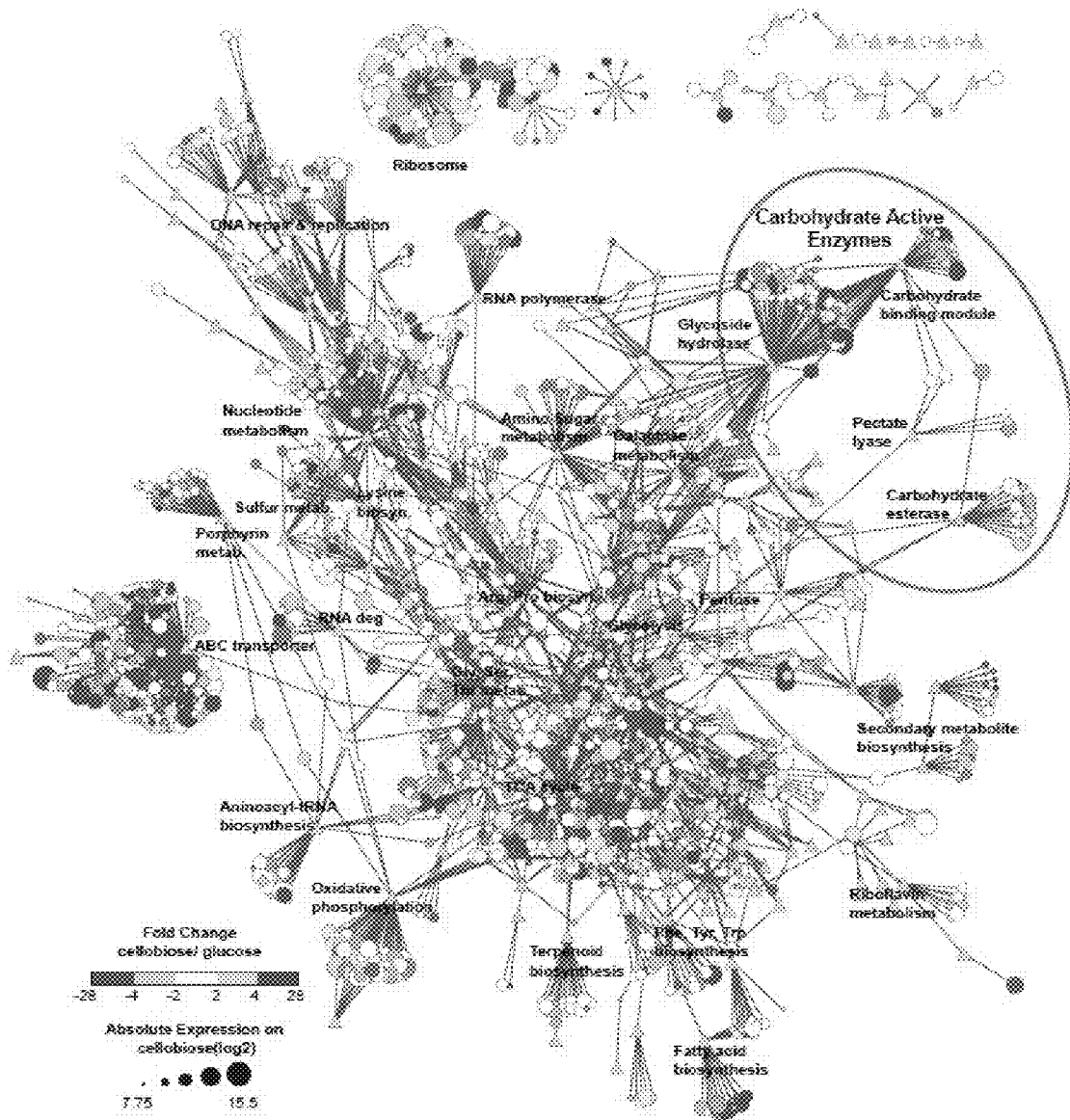


Figure 10

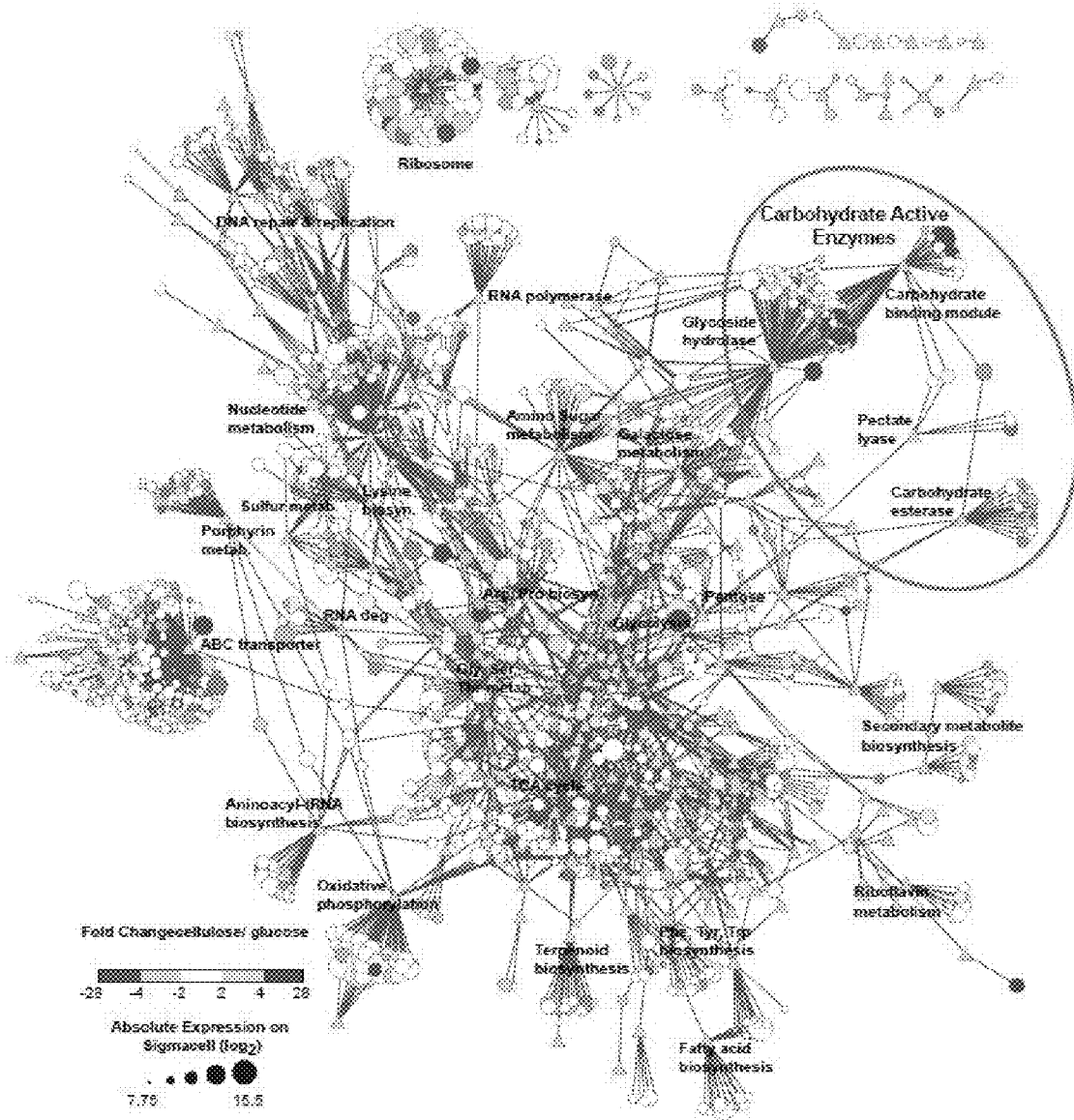


Figure 11

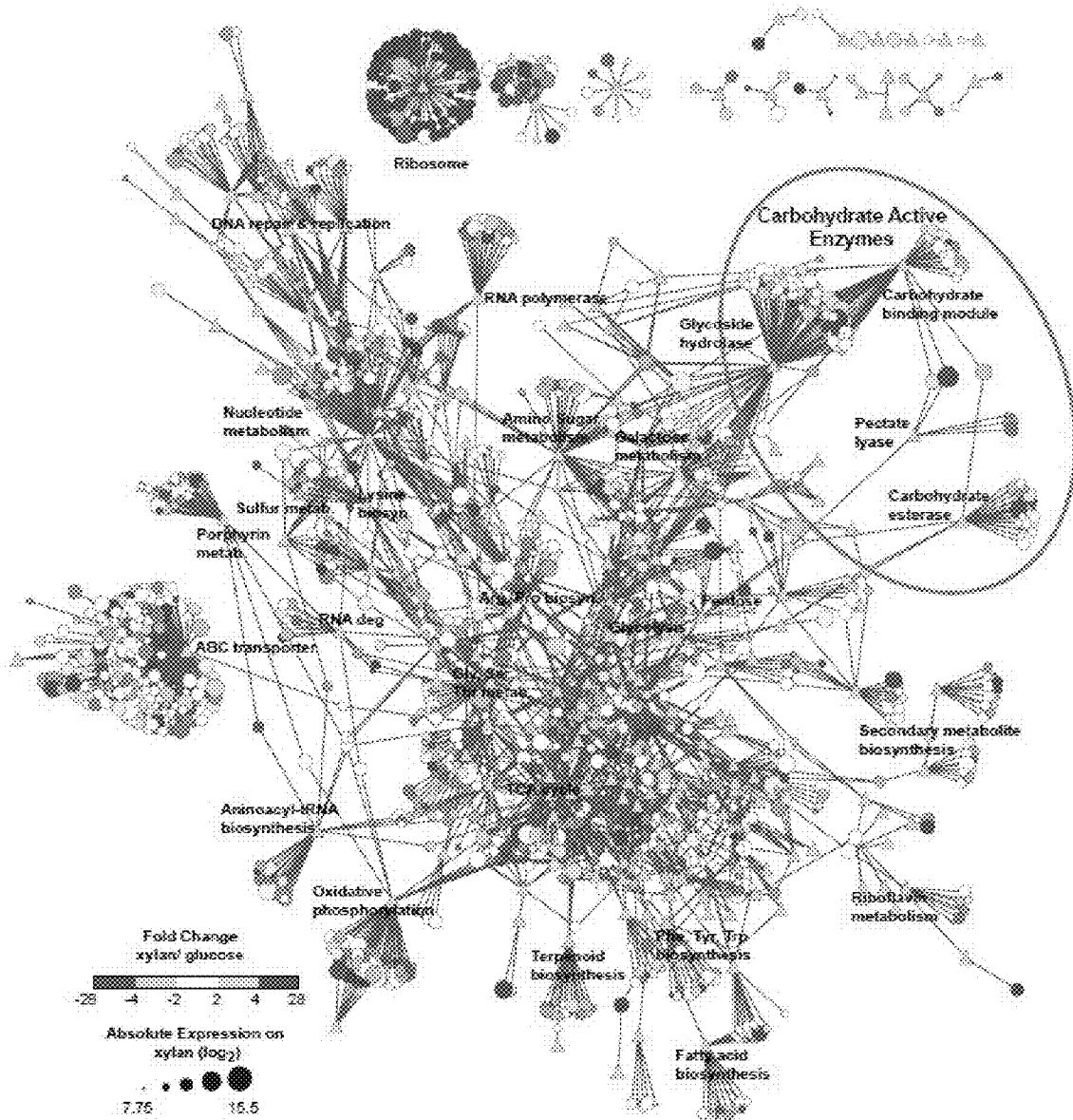


Figure 12

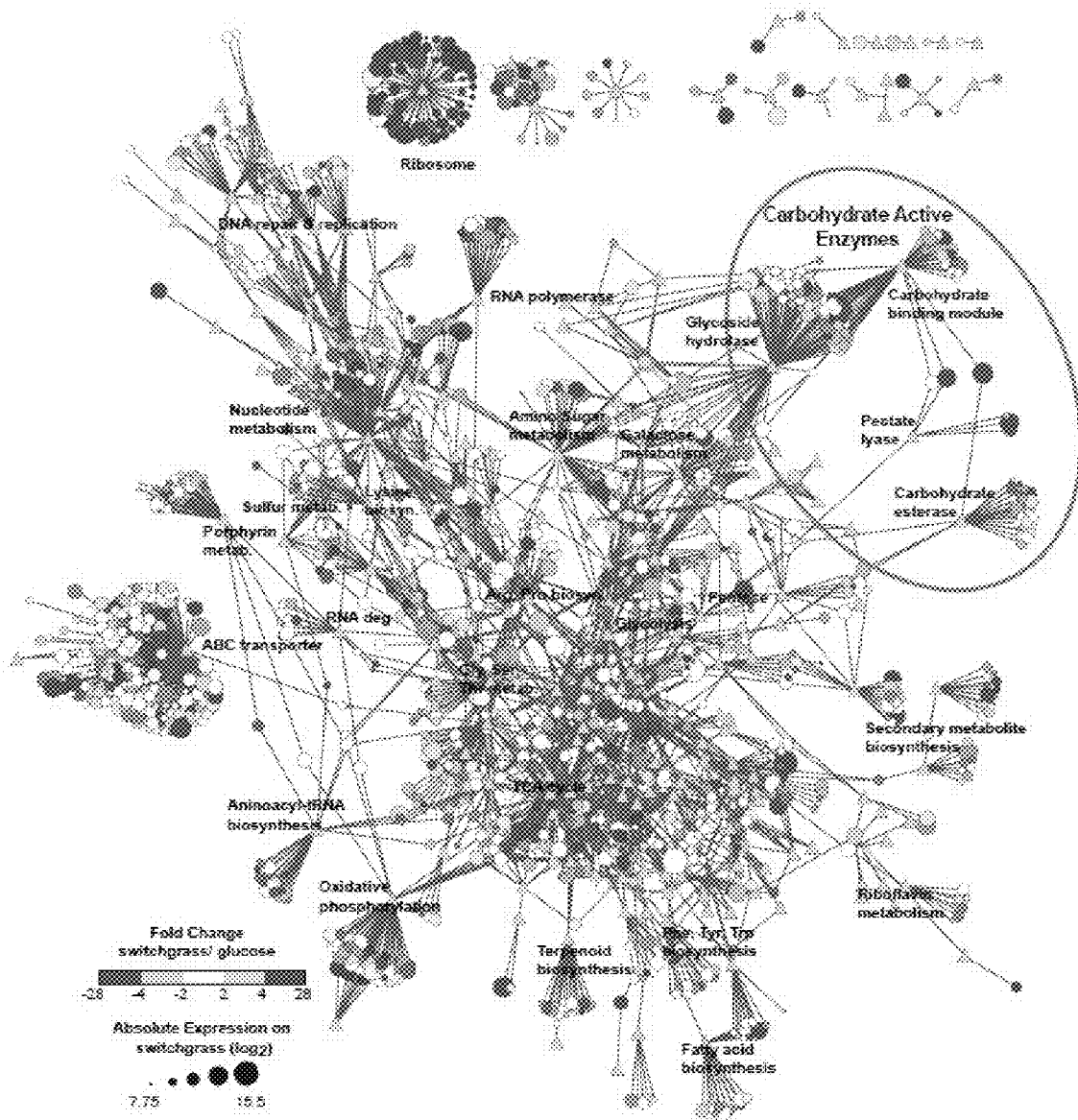


Figure 13

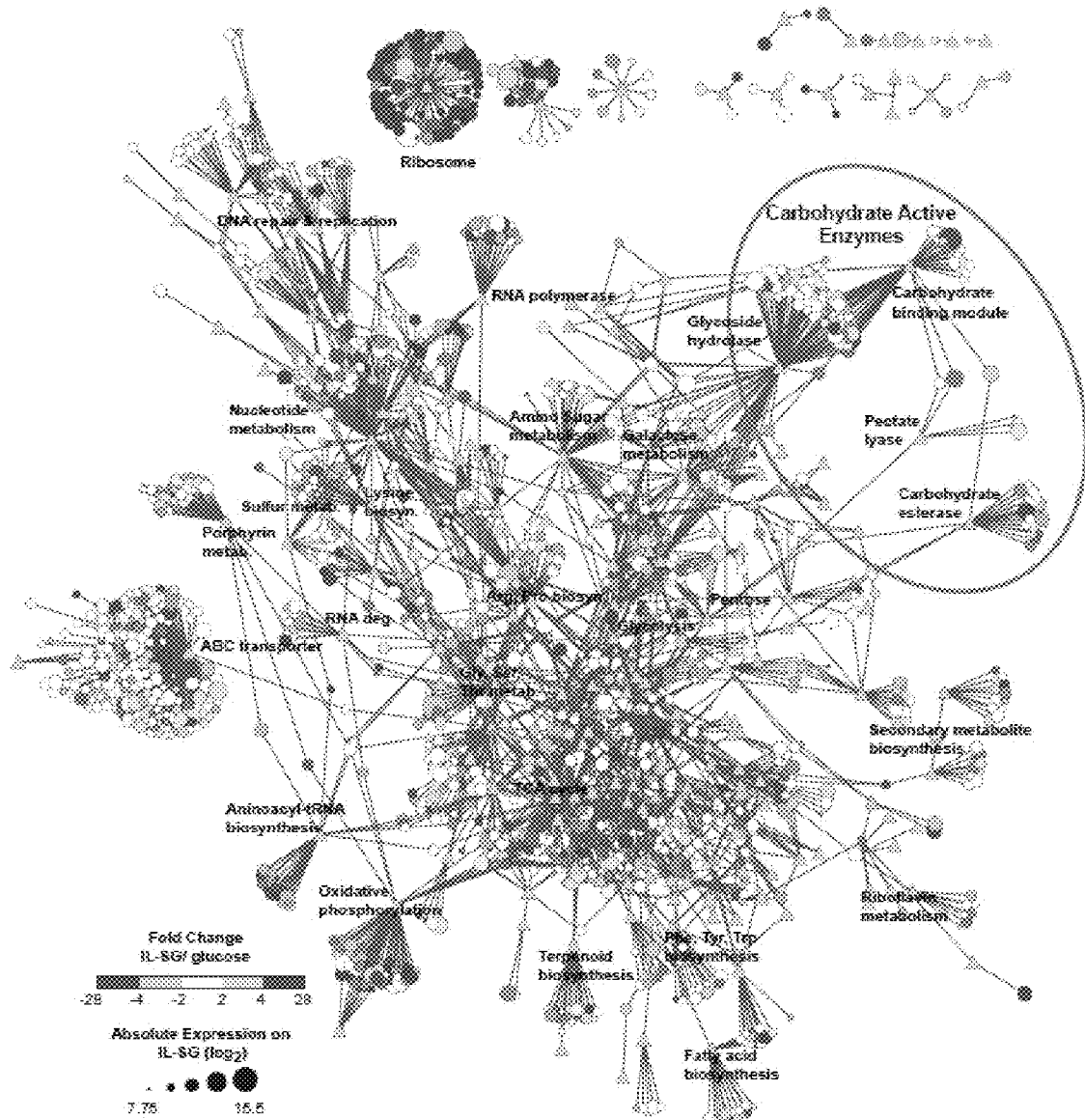
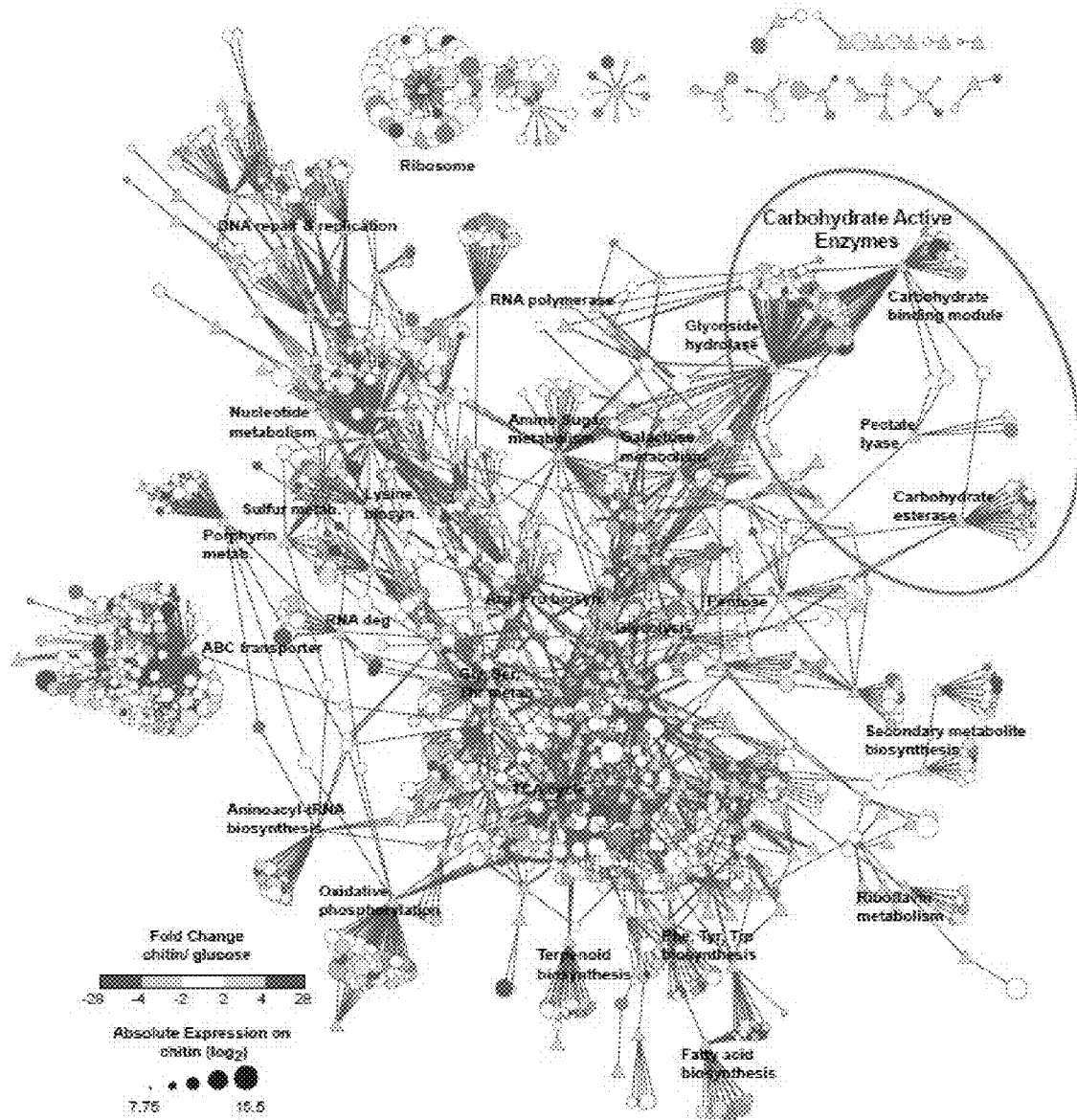


Figure 14



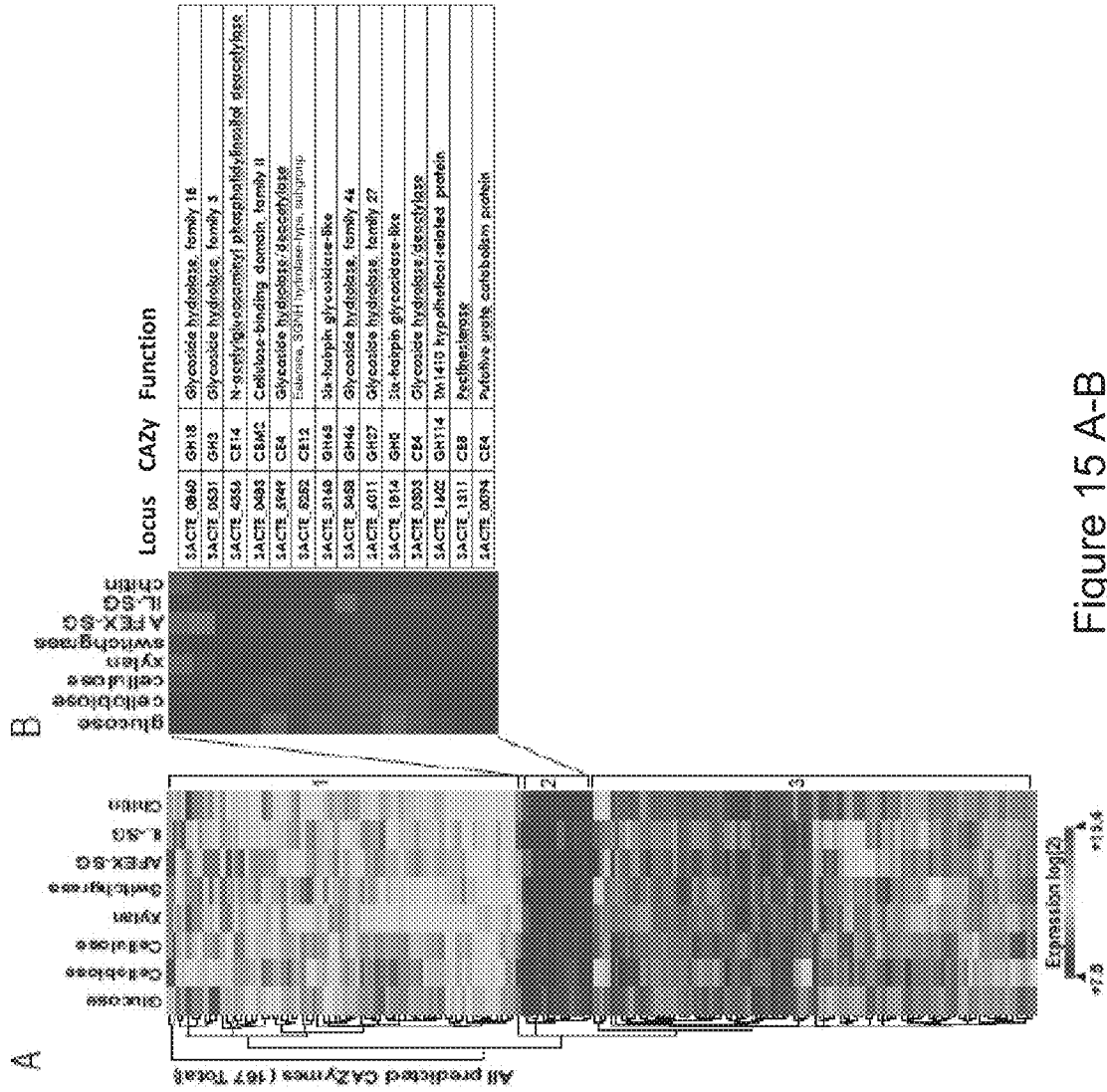


Figure 15 A-B

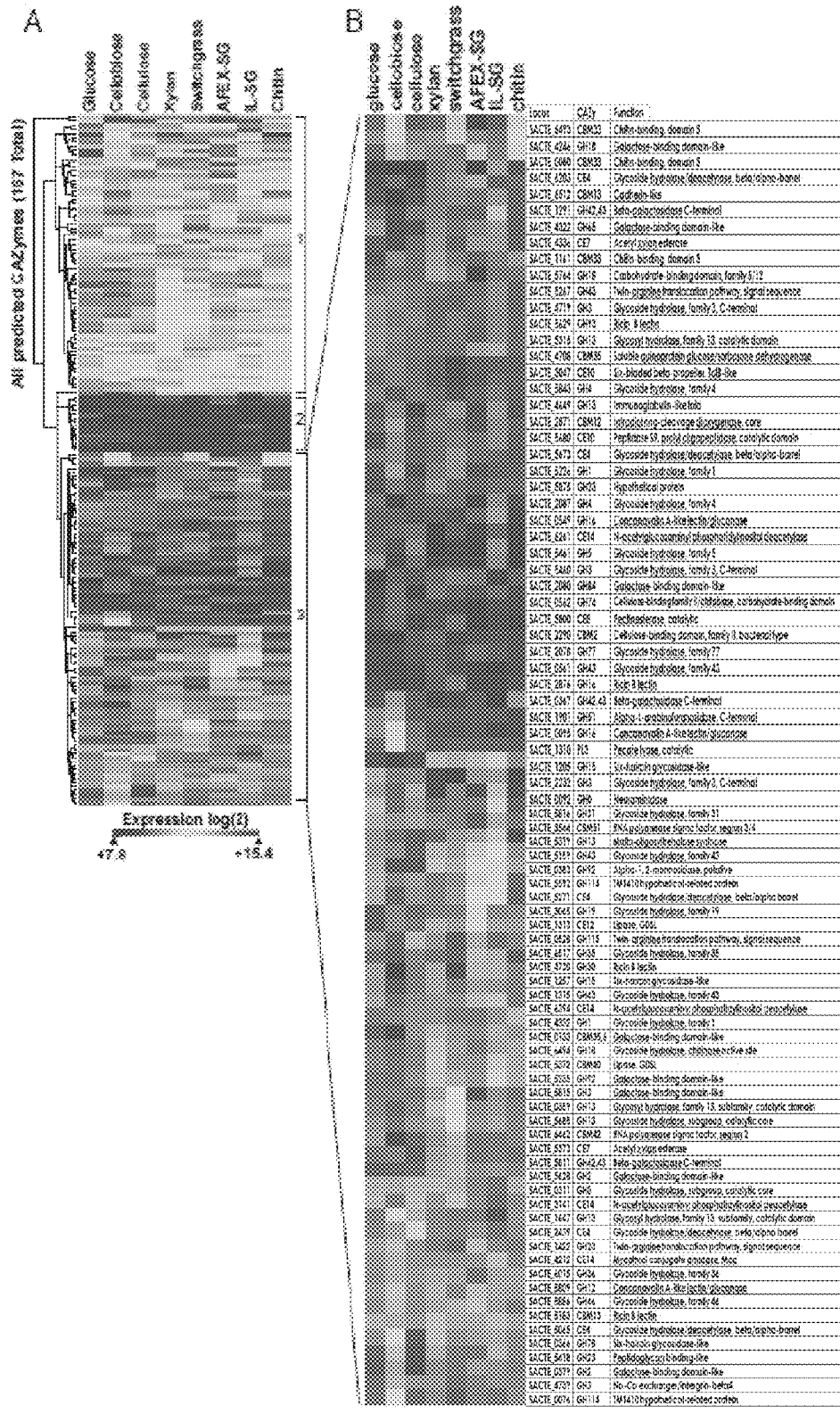


Figure 16 A-B

Figure 17

Protein band ^a	Locus	Catalytic domain	CBM	Functional class ^b
a	SACTE_3159	CBM33	CBM2	cellulase
b	SACTE_0265	GH10	CBM2	xylanase
c	SACTE_4755	GH64		beta-1,3-glucanase
d	SACTE_0482	GH5	CBM2	cellulase
e	SACTE_0237	GH6	CBM2	cellulase
f	SACTE_0236	GH48	CBM2	cellulase
g	SACTE_3717	GH9	CBM2	cellulase
h	SACTE_2347	GH5	CBM2	mannanase

^a Protein bands labeled in Figure 3A were identified by MALDI-TOF mass spectrometry. ^b Function identified by assays of individual fractions from ion exchange chromatography.

Figure 18

Figure 18: Spectra count of proteins identified on each substrate, where top 95 % spectra covered were bolded on glucose, cellobiose, cellulose, xylan switchgrass, AFEX-SG, IL-SG and chitin, respectively.

Locus	CAZy	Identified Proteins (414)	glucose	cellobiose	cellulose	xylan	switchgrass	AFEX-SG	IL-SG	chitin
SACTE_0237	GH6	ACTE_1_4-beta cellobiohydrolase	0	462	3965	8	3330	3300	3520	0
SACTE_0236	GH48	ACTE_Glycoside hydrolase, 48F	0	44	1296	0	577	905	1045	0
SACTE_2347	GH5,CE3	ACTE_Cellulose-binding family II/chitinase, carbohydrate-binding domain	0	28	624	0	185	240	205	0
SACTE_3159	CBM33,2	ACTE_Cellulose-binding domain, family II, bacterial type	0	4	564	29	493	646	847	8
SACTE_0482	GH5	ACTE_Cellulose-binding family II/chitinase, carbohydrate-binding domain	0	11	156	0	92	113	153	0
SACTE_0265	GH10	ACTE_Glycoside hydrolase, family 10	0	13	90	778	118	220	101	8
SACTE_0357	CE4	ACTE_Glycoside hydrolase/deacetylase, beta/alpha-barrel	0	12	87	938	94	195	190	3
SACTE_4439		ACTE_Catalase, N-terminal	0	65	83	30	50	20	30	271
SACTE_0562	GH74	ACTE_Cellulose-binding family II/chitinase, carbohydrate-binding domain	0	0	83	0	29	48	30	0
SACTE_0358	GH11	ACTE_Glycoside hydrolase, family 11, active site	0	0	67	649	165	324	119	7
SACTE_4343		ACTE_Bacterial extracellular solute-binding protein, family 5	67	298	53	60	55	43	30	19
SACTE_1546		ACTE_Bacterioferritin	0	117	32	42	32	22	24	35
SACTE_1310	PL3	ACTE_Pectate lyase, catalytic	0	57	26	14	226	96	48	25
SACTE_4638		ACTE_Chondroitin AC/alginate lyase	0	41	23	135	34	59	59	14
SACTE_5668		ACTE_Alpha/beta hydrolase fold-1	29	30	21	23	0	0	9	39
SACTE_3717	GH9	ACTE_Carbohydrate-binding, CenC-like	0	49	21	0	342	432	143	0
SACTE_3590		ACTE_Phospholipase C, phosphatidylinositol-specific, X domain	0	5	21	0	0	4	7	0
SACTE_4571	GH18	ACTE_Ef-Hand 1, calcium-binding site	0	9	20	5	18	14	10	1758
SACTE_2172		ACTE_Citrate synthase-like, core	19	95	20	51	47	12	15	46
SACTE_5978	PL1	ACTE_Galactose-binding domain-like	0	5	20	525	287	245	10	0
SACTE_6428	CBM33	ACTE_Chitin-binding, domain 3	0	3	20	0	6	5	8	0
SACTE_2313	CBM33	ACTE_Chitin-binding, domain 3	0	11	19	0	12	0	11	308
SACTE_0366	GH78	ACTE_Six-hairpin glycosidase-like	0	4	17	30	0	0	5	20
SACTE_1604		ACTE_Pyridine nucleotide-disulphide oxidoreductase, class I, active site	72	473	16	96	25	18	26	57
SACTE_4702		ACTE_Protein of unknown function DUF756	0	26	16	18	0	2	26	26
SACTE_2059		ACTE_Manganese/iron superoxide dismutase, C-terminal	0	83	13	6	13	16	0	62
SACTE_4755	GH64	ACTE_Twin-arginine translocation pathway, signal sequence	2	92	12	23	33	30	30	24
SACTE_4945		ACTE_Extracellular solute-binding protein, family 3	38	26	12	18	16	4	22	9
SACTE_4730	GH30	ACTE_Ricin B lectin	0	40	12	41	22	25	51	7
SACTE_2556		ACTE_Enolase, N-terminal	57	84	12	47	10	4	19	0
SACTE_1041		ACTE_ABC-type glycine betaine transport system, substrate-binding domain	25	71	11	8	6	6	10	17
SACTE_4246	GH18	ACTE_Galactose-binding domain-like	0	20	10	15	7	9	11	1115
SACTE_0880		ACTE_S-adenosylmethionine synthetase, central domain	0	15	10	25	6	3	12	33
SACTE_5330		ACTE_Peptidase S1/S6, chymotrypsin/Hap	0	13	10	6	9	11	11	14
SACTE_0464	GH16	ACTE_Ricin B lectin	0	7	10	14	6	4	8	9
SACTE_5335		ACTE_ABC transporter, substrate-binding protein, aliphatic sulphonates	4	58	9	24	4	11	9	35
SACTE_2232	GH3	ACTE_Glycoside hydrolase, family 3, C-terminal	8	32	9	31	0	0	0	9
SACTE_0132		ACTE_hypothetical protein	0	0	9	0	6	0	16	0
SACTE_2289		ACTE_Bacterial extracellular solute-binding, family 1	0	13	9	0	0	0	7	0
SACTE_1130		ACTE_Galactose-binding domain-like	0	43	9	55	22	11	6	0
SACTE_5457	GH46	ACTE_Glycoside hydrolase, family 46	0	461	8	148	21	23	12	57
SACTE_3803		ACTE_Pyridoxal phosphate-dependent transferase, major domain	6	165	8	14	11	0	13	23
SACTE_1422	GH23	ACTE_Twin-arginine translocation pathway, signal sequence	27	19	8	34	0	0	2	19
SACTE_3306		ACTE_6-phosphogluconate dehydrogenase related protein	11	86	8	18	8	0	9	18
SACTE_5230		ACTE_Xylose isomerase-like, TIM barrel domain	0	15	8	1019	150	183	245	6
SACTE_0383	GH92	ACTE_Alpha-1,2-mannosidase, putative	0	0	8	8	0	0	0	6
SACTE_5647	GH87	ACTE_Galactose-binding domain-like	0	46	8	96	75	63	36	0
SACTE_4676		ACTE_Bacterial extracellular solute-binding protein, family 5	9	25	8	12	11	3	9	0
SACTE_0642		ACTE_Peptidase S33, tripeptidyl-peptidase C-terminal	2	74	8	0	15	0	0	0
SACTE_3779	GH31	ACTE_Glycoside hydrolase, family 31	0	5	8	0	0	0	0	0

Figure 18 (Continued)

SACTE_0746	ACTE_Twin-arginine translocation pathway, signal sequence	0	0	7	0	0	0	2	47
SACTE_4468	ACTE_Bacterial extracellular solute-binding, family 1	0	4	7	0	0	0	0	43
SACTE_4363 GH55	ACTE_hypothetical protein	0	72	7	18	16	14	19	12
SACTE_5166 GH43	ACTE_Twin-arginine translocation pathway, signal sequence	0	62	7	37	23	24	24	4
SACTE_1369	ACTE_Glyceraldehyde 3-phosphate dehydrogenase, NAD(P) binding domain	10	176	7	10	30	29	36	0
SACTE_1364	ACTE_Phosphoglucose isomerase, conserved site	2	147	6	16	14	13	13	20
SACTE_5519	ACTE_YD repeat	9	0	6	0	0	0	0	0
SACTE_0244	ACTE_N-acetylmuramoyl-L-alanine amidase, family 2	6	27	5	22	0	0	7	25
SACTE_4612	ACTE_hypothetical protein	0	8	5	200	67	30	31	7
SACTE_3458	ACTE_Surface protein from Gram-positive cocci	9	21	4	14	5	3	7	82
SACTE_4198	ACTE_NAD(P)-binding domain	6	66	4	19	14	5	8	24
SACTE_2033	ACTE_Nucleoside diphosphate kinase, core	0	7	4	8	12	4	4	16
SACTE_1367	ACTE_Triosephosphate isomerase	0	151	4	13	0	4	10	7
SACTE_0379 GH2	ACTE_Galactose-binding domain-like	0	72	4	10	6	3	10	3
SACTE_5371	ACTE_Neuraminidase	0	4	4	5	2	2	3	3
SACTE_5629 GH93	ACTE_Ricin B lectin	0	80	4	0	0	4	3	2
SACTE_4858 GH18	ACTE_Galactose-binding domain-like	0	17	3	18	5	5	4	384
SACTE_0844 GH18	ACTE_Glycoside hydrolase, subgroup, catalytic core	0	19	3	0	0	0	0	155
SACTE_5589	ACTE_Gamma-glutamyltranspeptidase	9	33	3	31	0	0	0	4
SACTE_5240	ACTE_hypothetical protein	0	30	3	3	0	0	3	0
SACTE_0604 GH109	ACTE_NAD(P)-binding domain	3	2	3	0	0	0	2	0
SACTE_4231	ACTE_Serine/cysteine peptidase, trypsin-like	0	4	3	9	8	0	0	0
SACTE_5267 GH43	ACTE_Twin-arginine translocation pathway, signal sequence	4	2	3	0	0	0	0	0
SACTE_4727	ACTE_Isocitrate/isopropylmalate dehydrogenase	0	17	2	5	0	0	3	36
SACTE_1302 PL1	ACTE_Twin-arginine translocation pathway, signal sequence	0	8	2	4	116	43	7	7
SACTE_6439 GH43	ACTE_Glycoside hydrolase, family 43	0	66	2	9	4	7	3	4
SACTE_6549	ACTE_Peptidase S11, D-alanyl-D-alanine carboxypeptidase A	0	0	2	0	0	0	0	3
SACTE_3711	ACTE_Glycosyl transferase, family 20	0	22	2	3	3	2	3	0
SACTE_3064 GH19	ACTE_Glycoside hydrolase, family 19, catalytic	0	21	0	37	19	12	0	1333
SACTE_5764 GH18	ACTE_Carbohydrate-binding domain, family 5/12	0	3	0	0	0	0	0	365
SACTE_0081 GH19	ACTE_Glycoside hydrolase, family 19	0	40	0	26	0	0	5	322
SACTE_1701	ACTE_Htaa	4	78	0	77	0	9	0	77
SACTE_6494 GH18	ACTE_Glycoside hydrolase, chitinase active site	0	0	0	0	0	0	0	69
SACTE_2768	ACTE_Surface protein from Gram-positive cocci	0	13	0	12	0	0	0	54
SACTE_2062	ACTE_DSBA oxidoreductase	0	22	0	6	5	0	0	51
SACTE_0860 GH18	ACTE_Carbohydrate-binding domain, family 5/12	0	0	0	0	0	0	0	44
SACTE_4908	ACTE_Exoribonuclease, phosphorolytic domain 2	9	13	0	43	26	21	24	33
SACTE_0926	ACTE_Vitamin B6 biosynthesis protein	15	13	0	19	3	2	0	25
SACTE_3097	ACTE_Heat shock protein Hsp70	39	4	0	0	0	0	0	25
SACTE_4178	ACTE_FAD-dependent pyridine nucleotide-disulphide oxidoreductase	5	21	0	17	0	0	3	21
SACTE_3666	ACTE_Histidine phosphatase superfamily, clade-1	30	0	0	17	0	3	0	21
SACTE_1858	ACTE_Bacterial stress protein	0	25	0	17	11	0	0	19
SACTE_4683	ACTE_Malate dehydrogenase, active site	90	19	0	2	0	0	0	19
SACTE_2384	ACTE_Phosphotransferase system, EIC component, type 1	0	0	0	0	0	0	0	19
SACTE_0634	ACTE_Periplasmic binding protein	0	3	0	26	0	0	0	17
SACTE_3078	ACTE_Ketose-bisphosphate aldolase, class-II	0	314	0	161	26	14	12	16
SACTE_4669	ACTE_Pyridoxal phosphate-dependent transferase, major domain	12	24	0	24	10	7	4	16
SACTE_3685	ACTE_Molybdate/tungstate binding	0	7	0	0	4	0	0	14
SACTE_3452	ACTE_Cobaltchelataase, CobN subunit	22	0	0	15	7	7	0	13
SACTE_4039	ACTE_IMP dehydrogenase/GMP reductase	9	3	0	4	0	0	2	12
SACTE_1356	ACTE_Transaldolase, active site	0	9	0	13	3	0	0	12
SACTE_4472	ACTE_Glucosamine-6-phosphate isomerase, subgroup	0	0	0	0	0	0	0	12
SACTE_1702	ACTE_Htaa	0	0	0	0	0	0	0	12
SACTE_0450	ACTE_Dak kinase	0	16	0	0	7	8	8	11

Figure 18 (Continued)

SACTE_3389		ACTE_Peptidase M24B, X-Pro dipeptidase/aminopeptidase P, conserved sit	6	38	0	7	6	2	3	11
SACTE_4607		ACTE_Protein of unknown function DJF1557	0	10	0	18	5	3	0	11
SACTE_6308		ACTE_Periplasmic binding protein	5	25	0	23	0	0	0	11
SACTE_5342		ACTE_Nitrite/sulphite reductase, hemoprotein beta-component, ferredoxin	0	6	0	0	3	0	4	9
SACTE_5751		ACTE_Twin-arginine translocation pathway, signal sequence	0	45	0	16	33	4	3	9
SACTE_5260		ACTE_FAD-dependent pyridine nucleotide-disulphide oxidoreductase	0	11	0	4	0	0	0	9
SACTE_0264		ACTE_Protein of unknown function DJF541	0	7	0	0	0	0	0	9
SACTE_4493		ACTE_Superoxide dismutase, Nickel-type	17	10	0	30	20	11	23	8
SACTE_4145		ACTE_Basic membrane lipoprotein	4	13	0	4	2	0	4	8
SACTE_4515	GH25	ACTE_Twin-arginine translocation pathway, signal sequence	0	45	0	35	6	12	21	7
SACTE_5263		ACTE_Dimeric alpha-beta barrel	3	25	0	9	2	0	2	7
SACTE_3038		ACTE_Bacterial stress protein	174	253	0	140	31	12	0	7
SACTE_3164		ACTE_hypothetical protein	0	0	0	7	2	0	0	7
SACTE_2049		ACTE_Ribose 5-phosphate isomerase, actinobacteria	12	18	0	5	0	0	0	7
SACTE_3319		ACTE_Pyridine nucleotide-disulphide oxidoreductase, class-II, active site	0	0	0	0	0	0	4	6
SACTE_3012		ACTE_AMP-binding, conserved site	18	0	0	0	13	2	3	6
SACTE_2323		ACTE_Fumarylacetoacetase, C-terminal-related	0	0	0	0	0	0	0	6
SACTE_2490		ACTE_S-adenosyl-L-homocysteine hydrolase, conserved site	0	0	0	0	5	0	0	5
SACTE_1949		ACTE_Peptidase M4, thermolysin	0	604	0	86	0	0	0	5
SACTE_0985		ACTE_N-acetyl-gamma-glutamyl-phosphate reductase	0	0	0	0	0	0	0	5
SACTE_3327		ACTE_NAD(P)-binding domain	62	1	0	0	21	15	31	4
SACTE_5418	GH23	ACTE_Peptidoglycan binding-like	0	9	0	34	12	15	10	4
SACTE_2585		ACTE_Phosphoribosyltransferase	6	35	0	10	9	0	7	4
SACTE_5220		ACTE_Aconitase/3-isopropylmalate dehydratase large subunit, alpha/beta	45	26	0	46	12	4	3	4
SACTE_4843		ACTE_Aminotransferase class-III	0	9	0	0	2	0	0	4
SACTE_0639		ACTE_Formyl transferase, N-terminal	0	53	0	5	0	0	0	4
SACTE_4243		ACTE_Endoribonuclease L-PSP/chorismate mutase-like	0	13	0	0	0	0	0	4
SACTE_4738	GH16	ACTE_Galactose-binding domain-like	0	208	0	10	11	14	0	3
SACTE_2260		ACTE_Acyl-CoA dehydrogenase/oxidase, N-terminal	0	0	0	0	5	0	0	3
SACTE_4078		ACTE_MG5-like	12	7	0	20	4	0	0	3
SACTE_5657		ACTE_MaoC-like dehydratase	0	14	0	3	4	0	0	3
SACTE_4436		ACTE_Uncharacterised protein family UPF0182	0	9	0	3	0	0	0	3
SACTE_4946		ACTE_ABC transporter, conserved site	0	3	0	0	0	0	0	3
SACTE_6131		ACTE_Purple acid phosphatase-like, N-terminal	0	0	0	0	0	0	0	3
SACTE_0324	GH55	ACTE_Galactose-binding domain-like	0	3	0	5	4	0	8	2
SACTE_1137		ACTE_Peptidase S45, penicillin amidase	0	0	0	4	9	4	0	2
SACTE_5455		ACTE_Amidohydrolase 1	0	10	0	0	0	3	0	2
SACTE_2645		ACTE_Penicillin/cephalosporin acylase	0	13	0	2	0	0	0	2
SACTE_5859		ACTE_hypothetical protein	0	13	0	6	66	27	51	0
SACTE_5231	GH31	ACTE_Glycoside hydrolase, family 31	0	2	0	41	2	4	21	0
SACTE_1073		ACTE_Proteasome, alpha subunit	0	0	0	0	2	2	11	0
SACTE_1901	GH51	ACTE_Alpha-L-arabinofuranosidase, C-terminal	0	0	0	0	8	9	7	0
SACTE_1074		ACTE_Proteasome, subunit alpha/beta	0	0	0	0	7	0	7	0
SACTE_6558		ACTE_Quinoprotein amine dehydrogenase, beta chain-like	6	16	0	11	17	24	5	0
SACTE_4718		ACTE_Delta-1-pyrroline-5-carboxylate dehydrogenase 1	3	0	0	3	14	4	5	0
SACTE_0841		ACTE_Winged helix-turn-helix transcription repressor DNA-binding	0	3	0	0	8	0	5	0
SACTE_1897		ACTE_hypothetical protein	0	0	0	0	4	0	5	0
SACTE_4728		ACTE_Aminotransferase, class IV	0	0	0	0	0	0	5	0
SACTE_1619		ACTE_Glutamine synthetase, beta-Grasp	0	0	0	0	0	0	5	0
SACTE_6303		ACTE_Serine/cysteine peptidase, trypsin-like	0	0	0	0	4	0	4	0
SACTE_5482		ACTE_Aldo/keto reductase, conserved site	5	4	0	3	0	0	4	0
SACTE_1650		ACTE_Bacterial extracellular solute-binding, family 1	6	2	0	0	0	0	4	0
SACTE_0782		ACTE_Transcription regulator PadR N-terminal-like	0	0	0	0	0	0	4	0
SACTE_1895		ACTE_Mandelate racemase/muconate lactonizing enzyme, N-terminal	0	0	0	2	9	6	3	0

Figure 18 (Continued)

SACTE_3219		ACTE_Methionyl-tRNA synthetase, class Ia, N-terminal	14	7	0	0	2	2	3	0
SACTE_6051		ACTE_Catalase	0	8	0	0	0	0	3	0
SACTE_1473		ACTE_Dienelactone hydrolase	0	0	0	0	0	0	3	0
SACTE_0534		ACTE_Bacterial extracellular solute-binding, family 1	0	0	0	0	0	0	3	0
SACTE_1136		ACTE_Peptidoglycan recognition protein	3	1	0	11	5	0	2	0
SACTE_1344		ACTE_Pyridoxal phosphate-dependent transferase, major domain	13	0	0	0	2	0	2	0
SACTE_1859		ACTE_Bacterial stress protein	9	395	0	298	77	38	0	0
SACTE_2065		ACTE_Peptidase S8/S53, subtilisin/kexin/sedolisin	31	1219	0	121	26	37	0	0
SACTE_3700		ACTE_Bacterial stress protein	162	207	0	198	68	28	0	0
SACTE_4624	GH16	ACTE_Concanavalin A-like lectin/glucanase	0	26	0	0	17	11	0	0
SACTE_1250		ACTE_Peptidase S8/S53, subtilisin, active site	7	8	0	50	13	8	0	0
SACTE_5685	GH13	ACTE_Glycoside hydrolase, subgroup, catalytic core	0	10	0	5	14	5	0	0
SACTE_0364	GH87	ACTE_Galactose-binding domain-like	0	0	0	5	9	4	0	0
SACTE_1312	PL1	ACTE_Pectate lyase/Amb allergen	0	0	0	5	4	3	0	0
SACTE_5682		ACTE_Galactose-binding domain-like	0	44	0	14	4	2	0	0
SACTE_5606		ACTE_D-hydantoinase	0	1	0	3	3	2	0	0
SACTE_3777		ACTE_Aldo/keto reductase	0	4	0	0	15	0	0	0
SACTE_5880		ACTE_Twin-arginine translocation pathway, signal sequence	0	19	0	7	12	0	0	0
SACTE_5741		ACTE_NAD(P)-binding domain	0	0	0	0	8	0	0	0
SACTE_2762		ACTE_hypothetical protein	0	0	0	0	8	0	0	0
SACTE_1162		ACTE_Lipase, class 2	0	0	0	0	7	0	0	0
SACTE_1640	GH13	ACTE_Glycoside hydrolase, carbohydrate-binding	0	0	0	0	7	0	0	0
SACTE_2544		ACTE_Thiolase, C-terminal	0	5	0	0	6	0	0	0
SACTE_2213		ACTE_NAD(P)-binding domain	0	0	0	0	5	0	0	0
SACTE_4459		ACTE_Peptidoglycan binding-like	0	0	0	15	4	0	0	0
SACTE_4102		ACTE_Fumarate reductase/succinate dehydrogenase, FAD-binding site	20	12	0	10	4	0	0	0
SACTE_5740		ACTE_Twin-arginine translocation pathway, signal sequence	0	0	0	2	3	0	0	0
SACTE_0133	CBM35,6	ACTE_Galactose-binding domain-like	0	0	0	2	3	0	0	0
SACTE_4566		ACTE_ATPase, F1 complex, alpha subunit, C-terminal	72	0	0	0	3	0	0	0
SACTE_4568		ACTE_ATPase, F1 complex, beta subunit	32	0	0	0	3	0	0	0
SACTE_6063		ACTE_Protein of unknown function DUF336	0	0	0	0	3	0	0	0
SACTE_3962		ACTE_Peptidase S1A, chymotrypsin	0	0	0	0	3	0	0	0
SACTE_2518		ACTE_Acyl-CoA dehydrogenase/oxidase, N-terminal	0	0	0	0	3	0	0	0
SACTE_1738		ACTE_N-acetyl-gamma-glutamyl-phosphate reductase	0	0	0	0	3	0	0	0
SACTE_3227		ACTE_Peptidase M18, aminopeptidase I	0	24	0	5	2	0	0	0
SACTE_1995		ACTE_Gamma-glutamyl phosphate reductase GPR	0	0	0	3	2	0	0	0
SACTE_5109		ACTE_Luciferase-like	0	29	0	0	2	0	0	0
SACTE_5881		ACTE_Multicopper oxidase, type 2	0	0	0	0	2	0	0	0
SACTE_3741		ACTE_Twin-arginine translocation pathway, signal sequence	0	0	0	0	2	0	0	0
SACTE_1638		ACTE_Tautomerase	0	0	0	0	2	0	0	0
SACTE_0528	GH115	ACTE_Twin-arginine translocation pathway, signal sequence	0	0	0	50	0	0	0	0
SACTE_3197		ACTE_Heat shock protein DnaI-like protein dJIA	241	0	0	44	0	0	0	0
SACTE_4483		ACTE_Cyclic nucleotide-binding-like	0	0	0	24	0	0	0	0
SACTE_5630		ACTE_Aldo/keto reductase	0	0	0	17	0	0	0	0
SACTE_1603		ACTE_Peptidase M17, leucyl aminopeptidase, C-terminal	0	20	0	16	0	0	0	0
SACTE_1239		ACTE_Enoyl-[acyl-carrier-protein] reductase (NADH)	0	0	0	14	0	0	0	0
SACTE_0169		ACTE_Glyceraldehyde 3-phosphate dehydrogenase, active site	0	41	0	12	0	0	0	0
SACTE_3335		ACTE_Single-strand DNA-binding	7	35	0	12	0	0	0	0
SACTE_0365		ACTE_Peptidase S8/S53, subtilisin/kexin/sedolisin	0	15	0	11	0	0	0	0
SACTE_1434		ACTE_5'-Nucleotidase/apyrase	12	14	0	11	0	0	0	0
SACTE_1680		ACTE_Cupredoxin	0	3	0	10	0	0	0	0
SACTE_1003		ACTE_NAD(P)-binding domain	0	0	0	10	0	0	0	0
SACTE_0549	GH16	ACTE_Concanavalin A-like lectin/glucanase	0	14	0	9	0	0	0	0
SACTE_1325		ACTE_Dihydrodipicolinate synthetase	0	0	0	9	0	0	0	0

Figure 18 (Continued)

SACTE_1324		ACTE_hypothetical protein	0	0	0	9	0	0	0	0
SACTE_0222		ACTE_Peptidase M4, propeptide, PepSY	0	21	0	8	0	0	0	0
SACTE_2068		ACTE_Peptidase M1, aminopeptidase N actinomycete-type	6	15	0	8	0	0	0	0
SACTE_1281		ACTE_Surface protein from Gram-positive cocci	0	3	0	8	0	0	0	0
SACTE_4038		ACTE_Cystathionine beta-synthase, core	5	0	0	8	0	0	0	0
SACTE_4081		ACTE_NAD(P)-binding domain	13	7	0	7	0	0	0	0
SACTE_2819		ACTE_Chaperonin C1pA/B, conserved site	5	0	0	7	0	0	0	0
SACTE_3589		ACTE_S'-Nucleotidase/apyrase	0	0	0	7	0	0	0	0
SACTE_6144		ACTE_Twin-arginine translocation pathway, signal sequence	0	2	0	6	0	0	0	0
SACTE_4739	GH3	ACTE_Na-Ca exchanger/integrin-beta4	0	1	0	6	0	0	0	0
SACTE_4768		ACTE_Glutamyl/glutaminyl-tRNA synthetase, class Ic, catalytic domain	0	0	0	6	0	0	0	0
SACTE_4860		ACTE_Aminotransferase class-III	17	5	0	5	0	0	0	0
SACTE_3765		ACTE_hypothetical protein	0	4	0	5	0	0	0	0
SACTE_0687		ACTE_Protein of unknown function DUF885, bacterial	7	0	0	5	0	0	0	0
SACTE_1715		ACTE_Intermediate filament, C-terminal	0	45	0	4	0	0	0	0
SACTE_0325		ACTE_Bacterial stress protein	0	7	0	4	0	0	0	0
SACTE_5146		ACTE_FMN-binding split barrel, related	0	0	0	4	0	0	0	0
SACTE_4910		ACTE_Dihydrodipicolinate reductase, N-terminal	0	0	0	4	0	0	0	0
SACTE_0213		ACTE_Luciferase-like	0	0	0	4	0	0	0	0
SACTE_1311	CE8	ACTE_Pectinesterase, catalytic	0	0	0	4	0	0	0	0
SACTE_1313	CE12	ACTE_Lipase, GDSL	0	0	0	4	0	0	0	0
SACTE_6512	CBM13	ACTE_Cadherin-like	0	15	0	3	0	0	0	0
SACTE_5114		ACTE_Ornithine carbamoyltransferase	0	1	0	3	0	0	0	0
SACTE_2698		ACTE_hypothetical protein	0	1	0	3	0	0	0	0
SACTE_4618		ACTE_Phosphofructokinase, pyrophosphate dependent	5	0	0	3	0	0	0	0
SACTE_1240		ACTE_NAD(P)-binding domain	3	0	0	3	0	0	0	0
SACTE_6206		ACTE_hypothetical protein	0	0	0	3	0	0	0	0
SACTE_4814		ACTE_Ribosomal protein L19	0	0	0	3	0	0	0	0
SACTE_2318		ACTE_HAD-superfamily hydrolase, subfamily IIA	0	0	0	3	0	0	0	0
SACTE_1862		ACTE_Transketolase, C-terminal/Pyruvate-ferredoxin oxidoreductase, dom	0	0	0	3	0	0	0	0
SACTE_5493		ACTE_Periplasmic binding protein/LacI transcriptional regulator	0	12	0	2	0	0	0	0
SACTE_1002		ACTE_Twin-arginine translocation pathway, signal sequence	0	4	0	2	0	0	0	0
SACTE_2381		ACTE_Exoribonuclease, phosphorolytic domain 2	0	1	0	2	0	0	0	0
SACTE_1703		ACTE_Periplasmic binding protein	2	81	0	0	0	0	0	0
SACTE_5235	GH92	ACTE_Galactose-binding domain-like	0	25	0	0	0	0	0	0
SACTE_2966		ACTE_Pyridoxal phosphate-dependent transferase, major domain	0	24	0	0	0	0	0	0
SACTE_2238		ACTE_Putative agmatinase	5	21	0	0	0	0	0	0
SACTE_1694		ACTE_NAD(P)-binding domain	0	21	0	0	0	0	0	0
SACTE_0323	CBM32	ACTE_Galactose-binding domain-like	0	16	0	0	0	0	0	0
SACTE_4616		ACTE_Acetate/butyrate kinase	5	15	0	0	0	0	0	0
SACTE_5312		ACTE_Peptidase M24, structural domain	0	15	0	0	0	0	0	0
SACTE_4503		ACTE_NAD(P)-binding domain	0	13	0	0	0	0	0	0
SACTE_4366		ACTE_4Fe-4S ferredoxin, iron-sulphur binding domain	0	13	0	0	0	0	0	0
SACTE_0275		ACTE_Aldehyde/histidinol dehydrogenase	0	13	0	0	0	0	0	0
SACTE_0800		ACTE_Glycine cleavage system P-protein, N-terminal	7	10	0	0	0	0	0	0
SACTE_5452		ACTE_Transthyretin/hydroxysourate hydrolase	0	10	0	0	0	0	0	0
SACTE_0378		ACTE_Twin-arginine translocation pathway, signal sequence	0	10	0	0	0	0	0	0
SACTE_1426		ACTE_Extracellular ligand-binding receptor	6	9	0	0	0	0	0	0
SACTE_2063		ACTE_Peptidase M1, aminopeptidase N actinomycete-type	0	9	0	0	0	0	0	0
SACTE_1081		ACTE_tRNA methyltransferase complex GCD14 subunit	0	9	0	0	0	0	0	0
SACTE_6392		ACTE_Peptidase S33, prolyl aminopeptidase	0	8	0	0	0	0	0	0
SACTE_5353		ACTE_NAD(P)-binding domain	0	7	0	0	0	0	0	0
SACTE_3611		ACTE_L-asparaginase II	0	7	0	0	0	0	0	0
SACTE_3235		ACTE_Peptidase S11, D-alanyl-D-alanine carboxypeptidase A	0	7	0	0	0	0	0	0

Figure 18 (Continued)

SACTE_1582		ACTE_Cytochrome c oxidase subunit II C-terminal	0	7	0	0	0	0	0	0
SACTE_5319	GH13	ACTE_Malto-oligosyltrehalose synthase	0	7	0	0	0	0	0	0
SACTE_6361		ACTE_Molybdenum-pterin binding	0	6	0	0	0	0	0	0
SACTE_5555		ACTE_Beta-lactamase-like	0	6	0	0	0	0	0	0
SACTE_5395		ACTE_Barstar (barnase inhibitor)	0	6	0	0	0	0	0	0
SACTE_4606		ACTE_Cobalamin (vitamin B12)-dependent enzyme, catalytic subdomain	0	6	0	0	0	0	0	0
SACTE_4371		ACTE_Conserved hypothetical protein CHP00730	0	6	0	0	0	0	0	0
SACTE_1827		ACTE_Protease inhibitor I4, serpin	0	6	0	0	0	0	0	0
SACTE_5743	CBM6	ACTE_PKD/Chitinase domain	0	6	0	0	0	0	0	0
SACTE_4399		ACTE_Protein of unknown function DUF3107	14	5	0	0	0	0	0	0
SACTE_3023		ACTE_hypothetical protein	0	5	0	0	0	0	0	0
SACTE_1131		ACTE_N-acetylmuramoyl-L-alanine amidase, family 2	0	5	0	0	0	0	0	0
SACTE_0847		ACTE_Twin-arginine translocation pathway, signal sequence	0	5	0	0	0	0	0	0
SACTE_4591		ACTE_Thiolase-like	20	4	0	0	0	0	0	0
SACTE_1151		ACTE_Cupin, RmIC-type	2	4	0	0	0	0	0	0
SACTE_6170		ACTE_Domain of unknown function DUF1996	0	4	0	0	0	0	0	0
SACTE_5742		ACTE_Xylose isomerase-like, TIM barrel domain	0	4	0	0	0	0	0	0
SACTE_5113		ACTE_Arginine deiminase	0	4	0	0	0	0	0	0
SACTE_4590		ACTE_Glyoxalase/bleomycin resistance protein/dioxygenase	0	4	0	0	0	0	0	0
SACTE_4101		ACTE_Ferredoxin	0	4	0	0	0	0	0	0
SACTE_2614		ACTE_Deoxyribonuclease, TatD Mg-dependent, prokaryote	0	4	0	0	0	0	0	0
SACTE_2342		ACTE_hypothetical protein	0	4	0	0	0	0	0	0
SACTE_1575		ACTE_Rieske [2Fe-2S] iron-sulphur domain	0	4	0	0	0	0	0	0
SACTE_1224		ACTE_Band 7 protein	0	4	0	0	0	0	0	0
SACTE_0983		ACTE_Arginine biosynthesis protein ArgJ	0	4	0	0	0	0	0	0
SACTE_0872		ACTE_Cystathionine beta-synthase, core	0	4	0	0	0	0	0	0
SACTE_6011	GH27	ACTE_Glycoside hydrolase, family 27	0	4	0	0	0	0	0	0
SACTE_6458	GH25	ACTE_Glycoside hydrolase, family 25 subgroup	0	4	0	0	0	0	0	0
SACTE_6165		ACTE_NAD(P)-binding domain	0	3	0	0	0	0	0	0
SACTE_5944		ACTE_FMN-binding split barrel, related	0	3	0	0	0	0	0	0
SACTE_5619		ACTE_Isocitrate dehydrogenase NADP-dependent, monomeric type	0	3	0	0	0	0	0	0
SACTE_5406		ACTE_Aldehyde oxidase/xanthine dehydrogenase, molybdopterin binding	0	3	0	0	0	0	0	0
SACTE_4968		ACTE_Bacterial stress protein	0	3	0	0	0	0	0	0
SACTE_3045		ACTE_Lipase, GDSL	0	3	0	0	0	0	0	0
SACTE_2702		ACTE_Luciferase-like	0	3	0	0	0	0	0	0
SACTE_2272		ACTE_Acyl-CoA dehydrogenase/oxidase, N-terminal	0	3	0	0	0	0	0	0
SACTE_1479		ACTE_Imidazoleglycerol-phosphate dehydratase, conserved site	0	3	0	0	0	0	0	0
SACTE_5320	GH13	ACTE_Immunoglobulin-like fold	0	3	0	0	0	0	0	0
SACTE_5226	GH1	ACTE_Glycoside hydrolase, family 1	0	3	0	0	0	0	0	0
SACTE_1368		ACTE_Phosphoglycerate kinase	13	2	0	0	0	0	0	0
SACTE_4205		ACTE_Cys/Met metabolism, pyridoxal phosphate-dependent enzyme	7	2	0	0	0	0	0	0
SACTE_5135		ACTE_Concanavalin A-like lectin/glucanase	0	2	0	0	0	0	0	0
SACTE_4745		ACTE_Periplasmic binding protein	0	2	0	0	0	0	0	0
SACTE_4639		ACTE_Galactose-binding domain-like	0	2	0	0	0	0	0	0
SACTE_3883		ACTE_Ferredoxin	0	2	0	0	0	0	0	0
SACTE_2202		ACTE_Metallophosphoesterase	0	2	0	0	0	0	0	0
SACTE_0583		ACTE_Serine/threonine-protein kinase, active site	0	2	0	0	0	0	0	0
SACTE_0561	GH43	ACTE_Glycoside hydrolase, family 43	0	2	0	0	0	0	0	0
SACTE_4106	GH20	ACTE_Glycoside hydrolase, family 20	0	2	0	0	0	0	0	0
SACTE_5065	CE4	ACTE_Glycoside hydrolase/deacetylase, beta/alpha-barrel	0	2	0	0	0	0	0	0
SACTE_2290	CBM2	ACTE_Cellulose-binding domain, family II, bacterial type	0	2	0	0	0	0	0	0
SACTE_4042		ACTE_FAD-dependent glycerol-3-phosphate dehydrogenase	0	1	0	0	0	0	0	0
SACTE_3354		ACTE_Cys/Met metabolism, pyridoxal phosphate-dependent enzyme	0	1	0	0	0	0	0	0
SACTE_3102		ACTE_DeoxyUTP pyrophosphatase domain	0	1	0	0	0	0	0	0

Figure 18 (Continued)

SACTE_2853	ACTE_Protein of unknown function DUF2342	0	1	0	0	0	0	0	0
SACTE_2831	ACTE_Pyridoxal phosphate-dependent transferase, major domain	0	1	0	0	0	0	0	0
SACTE_2532	ACTE_Peptidase M20, dimerisation	0	1	0	0	0	0	0	0
SACTE_2215	ACTE_Peptidase M7, snapalysin	0	1	0	0	0	0	0	0
SACTE_1681	ACTE_Nitrilase/cyanide hydratase, conserved site	0	1	0	0	0	0	0	0
SACTE_1509	ACTE_Predicted pyridoxal phosphate-dependent enzyme, YBL036C type	0	1	0	0	0	0	0	0
SACTE_1424	ACTE_Aspartate decarboxylase-like fold	0	1	0	0	0	0	0	0
SACTE_1355	ACTE_Transketolase, C-terminal/Pyruvate-ferrredoxin oxidoreductase, dom	0	1	0	0	0	0	0	0
SACTE_2877	GH64 ACTE_Twin-arginine translocation pathway, signal sequence	0	1	0	0	0	0	0	0
SACTE_4708	CBM35 ACTE_Soluble quinoprotein glucose/sorbose dehydrogenase	0	1	0	0	0	0	0	0
SACTE_1419	ACTE_Ribosomal protein S1, RNA-binding domain	126	0	0	0	0	0	0	0
SACTE_1888	ACTE_Acyl carrier protein-like	60	0	0	0	0	0	0	0
SACTE_2468	ACTE_beta-acterial NAD-glutamate dehydrogenase	36	0	0	0	0	0	0	0
SACTE_3896	STRACTE_03857	35	0	0	0	0	0	0	0
SACTE_3716	STRACTE_03676	28	0	0	0	0	0	0	0
SACTE_4281	STRACTE_04237	27	0	0	0	0	0	0	0
SACTE_3955	STRACTE_03916	25	0	0	0	0	0	0	0
SACTE_4565	STRACTE_04519	23	0	0	0	0	0	0	0
SACTE_4031	ACTE_Chaperonin Cpn60	22	0	0	0	0	0	0	0
SACTE_3086	ACTE_hypothetical protein	22	0	0	0	0	0	0	0
SACTE_4376	STRACTE_04332	19	0	0	0	0	0	0	0
SACTE_4194	ACTE_Twin-arginine translocation pathway, signal sequence	18	0	0	0	0	0	0	0
SACTE_1934	ACTE_ATP-grasp fold, subdomain 2	18	0	0	0	0	0	0	0
SACTE_4501	STRACTE_04456	17	0	0	0	0	0	0	0
SACTE_4959	STRACTE_04913	16	0	0	0	0	0	0	0
SACTE_2103	ACTE_Protein of unknown function DUF756	16	0	0	0	0	0	0	0
SACTE_4795	ACTE_SCP-like extracellular	15	0	0	0	0	0	0	0
SACTE_4073	ACTE_NAD(P)-binding domain	15	0	0	0	0	0	0	0
SACTE_2558	STRACTE_02525	15	0	0	0	0	0	0	0
SACTE_4224	STRACTE_04182	13	0	0	0	0	0	0	0
SACTE_3067	STRACTE_03033	13	0	0	0	0	0	0	0
SACTE_4462	STRACTE_04417	12	0	0	0	0	0	0	0
SACTE_3361	STRACTE_03325	12	0	0	0	0	0	0	0
SACTE_3995	ACTE_RNA polymerase, alpha subunit, C-terminal	11	0	0	0	0	0	0	0
SACTE_0514	STRACTE_00503	11	0	0	0	0	0	0	0
SACTE_6342	ACTE_Biotin/lipoyl attachment	10	0	0	0	0	0	0	0
SACTE_4610	STRACTE_04564	10	0	0	0	0	0	0	0
SACTE_1068	STRACTE_01051	10	0	0	0	0	0	0	0
SACTE_4926	ACTE_Ferritin-related	9	0	0	0	0	0	0	0
SACTE_4830	STRACTE_04784	9	0	0	0	0	0	0	0
SACTE_3392	STRACTE_03356	9	0	0	0	0	0	0	0
SACTE_3037	ACTE_Tellurium resistance	9	0	0	0	0	0	0	0
SACTE_2403	STRACTE_02370	9	0	0	0	0	0	0	0
SACTE_5983	STRACTE_05928	8	0	0	0	0	0	0	0
SACTE_4283	STRACTE_04239	8	0	0	0	0	0	0	0
SACTE_3385	STRACTE_03349	8	0	0	0	0	0	0	0
SACTE_1328	STRACTE_01307	8	0	0	0	0	0	0	0
SACTE_4415	STRACTE_04372	7	0	0	0	0	0	0	0
SACTE_2431	STRACTE_02398	7	0	0	0	0	0	0	0
SACTE_1586	STRACTE_01564	7	0	0	0	0	0	0	0
SACTE_1201	STRACTE_01182	7	0	0	0	0	0	0	0
SACTE_5081	ACTE_5-methyltetrahydropteroyltrimethylglutamate--homocysteine S-methyltran	6	0	0	0	0	0	0	0
SACTE_4873	STRACTE_04827	6	0	0	0	0	0	0	0
SACTE_3960	ACTE_Translation elongation factor EFG/EF2	6	0	0	0	0	0	0	0

Figure 18 (Continued)

SACTE_2756	STRACTE_02721	6	0	0	0	0	0	0	0	0
SACTE_3956	ACTE_DNA-directed RNA polymerase, beta subunit, bacterial-type	5	0	0	0	0	0	0	0	0
SACTE_3948	STRACTE_03909	5	0	0	0	0	0	0	0	0
SACTE_2801	STRACTE_02766	5	0	0	0	0	0	0	0	0
SACTE_0810	ACTE_Alpha-D-phosphohexomutase, alpha/beta/alpha domain II	5	0	0	0	0	0	0	0	0
SACTE_0669	STRACTE_00659	5	0	0	0	0	0	0	0	0
SACTE_5028	STRACTE_04982	4	0	0	0	0	0	0	0	0
SACTE_4567	ACTE_ATPase, F1 complex, gamma subunit	4	0	0	0	0	0	0	0	0
SACTE_4397	ACTE_Ferritin/ribonucleotide reductase-like	4	0	0	0	0	0	0	0	0
SACTE_4191	ACTE_L-Aspartase-like	4	0	0	0	0	0	0	0	0
SACTE_4030	ACTE_Chaperonin Cpn10, subgroup	4	0	0	0	0	0	0	0	0
SACTE_3961	STRACTE_03922	4	0	0	0	0	0	0	0	0
SACTE_3438	ACTE_Uracil phosphoribosyl transferase	4	0	0	0	0	0	0	0	0
SACTE_3371	STRACTE_03335	4	0	0	0	0	0	0	0	0
SACTE_3088	ACTE_Chaperonin ClpB	4	0	0	0	0	0	0	0	0
SACTE_2755	STRACTE_02720	4	0	0	0	0	0	0	0	0
SACTE_2729	STRACTE_02694	4	0	0	0	0	0	0	0	0
SACTE_2036	ACTE_Aminoacyl-tRNA synthetase, class 1a, anticodon-binding	4	0	0	0	0	0	0	0	0
SACTE_1285	STRACTE_01266	4	0	0	0	0	0	0	0	0
SACTE_1006	STRACTE_00988	4	0	0	0	0	0	0	0	0
SACTE_0548	ACTE_Aldo/keto reductase	4	0	0	0	0	0	0	0	0
SACTE_6004	STRACTE_05949	3	0	0	0	0	0	0	0	0
SACTE_5818	STRACTE_05765	3	0	0	0	0	0	0	0	0
SACTE_5021	STRACTE_04975	3	0	0	0	0	0	0	0	0
SACTE_5002	ACTE_DNA topoisomerase, type IIA, subunit A/C-terminal	3	0	0	0	0	0	0	0	0
SACTE_4828	ACTE_Translation elongation factor EFTs/EF1B, dimerisation	3	0	0	0	0	0	0	0	0
SACTE_4295	STRACTE_04251	3	0	0	0	0	0	0	0	0
SACTE_4186	ACTE_Acetyl-coenzyme A carboxyltransferase, C-terminal	3	0	0	0	0	0	0	0	0
SACTE_3957	ACTE_DNA-directed RNA polymerase, subunit beta-prime	3	0	0	0	0	0	0	0	0
SACTE_3895	ACTE_Polyprenyl synthetase	3	0	0	0	0	0	0	0	0
SACTE_3892	ACTE_Fumarylacetoacetase, N-terminal	3	0	0	0	0	0	0	0	0
SACTE_3284	ACTE_Serine/threonine-protein kinase, active site	3	0	0	0	0	0	0	0	0
SACTE_2533	STRACTE_02500	3	0	0	0	0	0	0	0	0
SACTE_2035	STRACTE_02006	3	0	0	0	0	0	0	0	0
SACTE_1872	ACTE_Beta-lactamase-related	3	0	0	0	0	0	0	0	0
SACTE_1045	ACTE_Periplasmic binding protein	3	0	0	0	0	0	0	0	0
SACTE_5618	STRACTE_05563	2	0	0	0	0	0	0	0	0
SACTE_3950	ACTE_Transcription antitermination protein, NusG	2	0	0	0	0	0	0	0	0
SACTE_3301	STRACTE_03267	2	0	0	0	0	0	0	0	0
SACTE_0900	ACTE_Chorismate synthase	2	0	0	0	0	0	0	0	0

Figure 19

>SACTE_0237|I, 4-beta cellobiohydrolase|GH6 (SEQ ID NO:17)

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>SACTE_0236|glycoside hydrolase family 48|GH48 (SEQ ID NO:18)

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Figure 19 (continued)

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Figure 19 (continued)

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Figure 19 (continued)

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Figure 19 (continued)

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Figure 19 (continued)

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Figure 19 (continued)

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Figure 19 (continued)

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Figure 19 (continued)

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Figure 20

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NPTNPACRDALSQSGANALYNWF AVLDSNAGGRGAGYVPDGLCSAGDRSPYDFSAY
NAARADWPRTHLTSGATLKVQYSNWA AHPGDFRVYLTKPGWAPTSELAWDDLQLVQ
TVSNPPQQGGAGTNGGHYYWDLALPSGRSGDALMFIQWVRSDSQENFFSCSDIVFDGG
NGEVTGIGGTGTPPTPTPTPTPTPTDPEHSGSCMAVYNNVSSWAGGFQASVEVMNHGT
EPRNGWAVQWKP GSGTQINSVWNGSLSTGSDGTVTVRVDVHN RV IAPDGSVTFGFTAT
STGNDYPAGTIGCVTS*

Figure 20 (continued)

VKRFLALLATCATVLGLTALTGPQAVAAAGCTADYTITSQWQGGFQAAVKVTNLGTPV
TGWKLFTFLPDAGQKVQGWNAAWSQSGSAVTAAGADWNGTLATGASAEAGFVGSF
TGANPPPATAFALNGVACTGSTGEPAGSDGGTPVDVNGQLHVCVNLNQNQYDRPVQLR
GMSTHGIQWFDACYDAASLDALANDWKSDDLRIAMYVQEDGYETDPAGFTRRVNDLV
DMAEARGMYALIDFHLLTPGDPNVNLDRAKTFASVAARNAGKKNVIYEIANEPNGVT
WTAVKSYAEQVIPVIRAADPDVAVVIVGTRGWSSLGVSDESEVVNSPVNATNIMYAF
HFYAASHKDAYRSTLSRAAARLPLFVTEFGTVSATGGGAMDRASTTAWLDDLQDKIS
YANWTVSDAPESSAARPGTCGGGDYSGSGVLTESGALLKNRISTPDSFPTG*

>SACTE_0265|glycoside hydrolase family 10|GH10|GI:344313522 (SEQ ID NO:5)
MAKKIPARARRALSVLTAGVLAAGVVSAGTAEAAAGTLGDAAAAAGRYFGTAVAAAN
HLGEAPYASTLDAQFDSVTPENEMKWDAVEGSRNSFTFTAADQIVSHAQSKGMKVRG
HTLVWHSQLPGWVGGGLGATDLRAAMNNHITQVMTHYKGGKIHSWDVNEAFQDGNNG
ARRSSPFQDKLGDGFIEEAFRTARTVDPTAKLCYNDYNTDGRNAKSDAVYAMAKDFKQ
RGVPIDCVGFQSHFNSNSPVPSDYRANLQRFADLGLDVQITELDIEGSGSAQAANYTSVV
NACLAVTRCTGLTVWGVTDKYSWRSSGTPLLFDGDYNNKKPAYDAVLAALGGTPDGGG
DDGGGDNNGGNTGSCATYTTQTATWNGGYNGEVTVKAGSSGITTWSVPVTVPSQQV
SALWNGAPTWNAGNTVMTVKPTYNGTLAAGASTSFGFTVMTNGNTSAPAVGACTAS*

>SACTE_2347|cellulose-binding family II|GH5,CE3|GI:344315549 (SEQ ID NO:6)
VRTAIRTARRPQPLALLLRGLAAFLGLALAGALGPATARAADLPQRAEARAAGLHISDG
RLVEGNGNDFVMRGINHAHTWYPGETQSLADIKATGANTVRVVLSDGYRWSNSPED
VASHARCKAERLICVLEVHDTTGYGEDAAAGTLDHAADYWIGLKDVLGDGEEDYVVINI
GNEPWGNADPAGWTAPTTAAIQKLRAAGFAHTIMVDAPNWGQDWEGVMRADARSVY
DADPTGNLIFSIIHMYSVYDTAAKVTDYLNFAFVDAGLPLLIGEFGGPADQYGDPEDTM
MATAEELGLGLYLAWSWSGNTDPVLDLVLDFDPTRLSSWGERVLHGPDGITETSREATV
FGGGQGGGDEAPTAPGTPTASGVTATSVTLGWSAATDDVGVTA YDVVRVTVGGSETK
VASSAATSVTVTGLSAGTAYSFAVYARDAAGNRSARSQTVSVTTDEGGSVPGGACSVG
YRVIGEWPGGFQGEITLRNTGAAAVDGTWTLGFAFADGQTVTNMWGGTATQSGGAVSV
TPASYTSTIAAGGSVTVGFTGTTLTGANAAPAAFTLNGATCTAA*

>SACTE_0357|polysaccharide deacetylase|CE4|GI:344313612 (SEQ ID NO:7)
MSITPRPSLRAMVTGLAVAASALAGGAVTAAPARAAACNGYVGLTFDDGPSAAQTPAL
LSALKQNGLRATMFNQGNYAASNPAQVKAQVDAGMWVGNHSYSHPHLTQQSQAQM
DSEISRTQQAIAAGGGGTPKLFPPYGETNATLRSVEAKYGLTEVIWDVDSQDWNGAST
DAIVQAVSRLTAGQVILMHEWPANTLAAIPRIAQTL SAKGLCSGMISPQTGRAVAPDGG
GNGGGGGGGGCTATLSAGEKWGDYRNLNVAVSGSSNWTVTMNVPSGERVMTTWN
VSASYPSAQVLVAKPNGSGNNWGATI QANGNWTWPTVSCCTS*

>SACTE_0358|Endo-1,4-beta-xylanase|GH11|GI:344313613 (SEQ ID NO:8)
MNPLVYTERRRRGRSLTSLAGSVCALVLA AAAAMLLPGTASADTVVTTNQTGNNGGY
YSFWTDGGGQVSMNLA SGGSYSTSWTNTGNFVAGKGWSTGGRKSVTYSGTFNPSGNA
YLTLYGWSTNPLVEYYIVDNWGTYRPTGTGFKGTVSSDGGTYDIYETTRTNAPSIEGKTF

Figure 20 (continued)

KQFWSVRQSKRTGGTTTTGNHFDAWARNGMNLGTMNYMILATEGYQSSGSSNITVSEG
GSGGGGDNGGGGGGGGGCTATLSAGEKWGDYRNLNVAVSGSSNWTVTMNVPSAEKV
LSTWNISASYPSSQVLVAKPNGSGNNWGATIQUANGNWTWPTVSCCTS*

>SACTE_1310|Pectate lyase|PL3|GI:344314542 (SEQ ID NO:9)
MSERAASPRTHRRRPGRRRIATALTAALGLTGAALATGVMLQPAGAATTAIPAWPSAT
GSQSVSKTIEVSGTYDGGKLRFTGSGDLGDDGGQDEGQDPIFKLKDGIATIKNVILGTPAAD
GIHCSGSCITQNVWVEDVGEDAASFSGTSTSSVYTVYGGGAKKASDKVVFQNGAGKL
VVTKFQVADFGKLVRSNGCNSKQYKREIIVNDVDVTAPGKSLVGINTNYGDTAALRSV
RVHGDSSKKIKPCVRYTGNSTGAEPKETGSGPDGTYCKYTASDLSYD*

>SACTE_3717|glycoside hydrolase family 9|GH9|GI:344316877 (SEQ ID NO:10)
MWCHPYLRLRTSGRKVSSVNALPPPARPAPVRPRSRYGRRVLGMSAAALLCAGALAVP
GTAMADDAEPGPGPEQITNGDFATGTSAPWWWTPNASAAVSEGRLCVEVPAGTANAW
DVIVGQNDVPIVAGESYELSYTARSTVPLTVQTRVQEAVEPYTTVLATADPVGAEDTRV
ARTFTASVDQPAASVQLQIGGGERATTFCLDDVSLRGGAEPPVYVPDGTSPVRVNQVG
YLPRGPKSGTVVTDAAEAPLTWTVKAEDGSTAATGTTVPRGEDPSSRRRVHTFDFGDLTT
AGDGYTVEVDGEVSEPFISIRGDLYDSLRS DALAYFYHNRSGIEIDADLVGEQYARPAGHI
GVAPNKGDTDVPCRPGVCDYRLDVSGGWYDAGDHGKYV VNGGISVAQLMATYERTL
TAPDAESAELGDGALRVPERDNGVPDILDEARWEMDFLIKMQVPAGEQLAGMVVHHKM
HDAEWTGLPMKPHLDPPQRELHPPSTAATLNLAATAAQCARLYAPFDADFADRCLRA
AETAWDAAKRHPDVLADPNDGIGGGAYNDDDDVSDEFYWAAAEELFTTTGKDIYRQAVL
SSAWHGDAGAVFPAGGGISWGSTAGLGVLT LATVPNALTS DQLAQVRTV VTEGADRY
AAQSREQAYGLPYAPRGEDYVWGSNSQVLNMMVVLATAHDLTGDAAYQDAVLRGAD
YLLGRNPLNQSYVTGYGERDSHNQHHRFWAHQNDP SLPNPAPGSIAGGPNLTAIASGDP
VAAEKLSGCAPAMCYVDDIGSWATNEITINWNAFLAFIASYLDDAGEGGQTAARTCQ
VTYSSHPWNSGSTVTVRVENTGSDPVSPWALTWLLPGEQRLSHTWSAEFDQHGRTVSA
RPLSWNRTLAPGAAVDFGFNTSAAGSSPEPGAFLKLN GRACCSAG*

>SACTE_4638|conserved hypothetical protein||GI:344317777 (SEQ ID NO:11)
MRTGSIARVLGLAAALAALLTTAFMAPAMAGKH DATDSPSAAAAPASFTHPGVLVSRP
QLDFVRGKVQAGAQPWK GAYDQMLASPYASLSRTAKPRAVVECGSYSNPNGCTDER
EDALAAAYTSLAWYISQDGRYAQKAIQIMDAWSGVIKDHTNSNAPLQTGWAGSSWPR
AAEIIKYTYGNWPASGRFGTMLRDVYLPKVANGSNSNGNWELSMTEAAIGIAVFLEDR
GAYDRAVAKFRGRVPAIYVTADGSLPKAAPGSLDTREKIINYWQQQSTFVDGLSQET
CRDLTHTGYGLSAISHIAETSRIQQDLYPEVADRRLHALGLHAKYQLGEKVPSSLCGGS
LKDSLGPVTEVGFNALHNRMGYAMTNTQTLTERQRPAASNNL FVAWETLTHADNPN*

>SACTE_4738|glycoside hydrolase family 16|GH16|GI:344317876 (SEQ ID NO:12)
MPSRTTLIATTAALVALAAPMAFAAPAPDPAVEAAAAAWDTDRAASAYAANPAAV
TASGSENPASGPGAATDGDATTRWSSDFADNAWIRVDL GSTIRINQVKLEWEAAYGKK
YVLEVSKDGTNWTPFYTEDAGTGGTVTAHTYPQEV TGRYVVRMRGVERATAWGYSLFS

Figure 20 (continued)

FQVYGGEPAPASTTRS NLALNHPAYGDLYQHAGNSPAFVTDGGWPADLKADRSRWSS
DWNADRWVGVDLGATSTINSVDLYWEAAYAVDYEIQVSDDNRTWRTVHRPSAAEVA
ARRADVKAPEAEVGRHDTINLPTPATGRYVRMLGKERRSFYNPAPSTAQFGYSLYEFQ
VWGTGGSADAAYPALPKNPGGAYRTTFFDDFTGSGGLDRSKWRVVRTGTEMGPVNGES
QAYVDSPDNIRTENGALVLESKYCKGCTPTPNGTFFDFTSGRVDTNTKFDFTYGKVSARM
KLPVGDGFWPAFWLLGSDVDDPAVSWPGSGETDIMENIGYGDWTSSGLHGGPGYSADG
NIGASQTYPNGGRADEWHTYGVWEWTPPEGMFTTVDDR VVQQTSRQKLESTRGKWVFDH
NQYVILNLALGGAYPGGYNQVTQPYWGLPQSSVDRIAQGGIKA EIDWVRVEQK*

>SACTE_4755|conserved hypothetical protein|GH64|GI:344317893 (SEQ ID NO:13)
VISRRMFLTGAAASATALTYPLWGTALSPRTSAAAATCELALENRSLPGTVHAYVTGHE
QGTDSWVLLRADGSVYRPESPGAPQTPLPVDCAIPLNGAGAGPVVLTLPQMYGARVYF
VRDDKLDLYLNPGPSLVEPAFATPTDPNYGRTWSFCEFTFNQQLYANISYVDLVTALPI
GLTLEGDSHTVAPLPDGAVQRIADDLTAQAAADGQPWDKLVTRGSDGQVLRVVSPQ
NLMAPYFDRPDEMPFRDLFAAQIDEVWEKYRSTDLRIDLQGGRTLAGRVSGDTLTFE
GGHTFSKPTSKDIFTCNHGPFTNPNPSDSDDKKALLARIAAGFNRSIMLSHPSQPNGTVA
DYYQDAVTNHWSRVVHANSPIGYAFPYDDVRPDGEPDVSGAANDGNPRRFTVSVGS*

>SACTE_5457|Chitosanase|GH46|GI:344318578 (SEQ ID NO:14)
VLPHNRRTARRTTRLTRTGGLAAAALGLALMALPVTAHAGAPTQPAAHHLEAAATGL
DDPAKKDIAMQLVSSAENSTLDWKAQYGYIEDIGDGRGYTAGIIGFCSGTGDMLALVER
YDTRSPGNVLA SYL PALREVDGTDSDHGLDPGFPRDWAEAAKDPVFQQAQNDERDRV
YFDPAVRQAKDDGLGTLGQFAYYDAIVMHGGGGDSTSFGSIRQRALAEAEPPSRGGDE
VAYLDAFLDARVWAMRQEEAHS DTSRVDTAQRVFLRDGNLNLDPPLDWQVYGD SFHI
G*

>SACTE_5647|coagulation factor 5/8 type domain protein|GH87|GI:344318749 (SEQ ID
NO:15)
MTPPHRHRLFRRSVSASLSLALTA VGTAAAVVLAGAPAAQAAAVPAPSPVGISGRGAA
VPFTEQEA EYAATNGTLIGPDRRYGSLPSEASGRQAVTLDAAGEYVEFTLTAPANAMTF
RYSLPD NAAAGTRDASLDLRVNGSVLKSVPVTSKYGWYGGYPFN NNPGDTPHFFY
DETRTMFGSTLPAGTKVRLQVASTAGSPSFTVDLADFEQVAAPV GKPSGALDVVSDFG
ADPTGAADSTAKIQAAVDAGRTQ GKVVYIPQGT FQVRDHIVVDQVTLRGAGPWYSVLT
GRHPTDRSKAVGVY GK YSAQGGSRNVTLKDFAIIGDIQERVDNDQVNAIGGAMSDSVV
DNVWMQHTKCGAWMDGPM DNFTIKNSRILDQTADGVNFHYGVTNSTVTNTFVRNTG
DDGLAMWAENVPNVKNKFTFNTVILPILANNIVTYGGKDITISDNVMADTITNGGGLHI
ANRYPGVNSGQGTAVAGHTHTAARNTLIRTGNSDFNWNFGV GAIWFSGLNEPISNATINI
TDSEVLDSSYAAIHLIEGASNGLHFKNVKIDGAGTYALQIQAPGTATFENVVATHIAQSN
PIHNCVSGSFQITRSGNSGWYADPPACTGVWPDVWTN GGVPGGGGPTNPTDPTDPT
DPTDPTDPEETGNLARGRTVTETSHTDVYGAANTVDGNADTYWESRNNAFPQSVTV D
LGA AKAVKR VV LKLPAAAWATRTQ T LSVSGSTDNGTYNSLKASAGYTFNPSSGNTAT
VSLPGTPVRYLRLTFTQNTGWPA AQLSELEAYTS*

>SACTE_5978|Pectate lyase/Amb allergen|PL1|GI:344319072 (SEQ ID NO:16)

Figure 20 (continued)

PANYDIIAVSFADAAGTPGAVTFNLDSAGLNGYTVAQFKADIKAKQAAGKNVIISVGGGE
KGTVSVNSDASANAFADSLYTLIQEYGFNGVDIDLLENGLNSTYMTKALRSLSSKVGSGL
VITMAPQTIDMQSTSGEYFKTALNIKDILTVVNMQYYNSGSM LGCDGKVYSQGSVDFLT
ALACIQLEGLAPSQVGLGVPASTRGAGSGYVAPSVVNAALDCLAKGTGCGSFKPSRT
YPDIRGAMTWSNWDATAGNAWSNAVGVPHVHGLP*

>SACTE_3064|Chitinase|GH19|GI:344316244 (SEQ ID NO:37)
VIRVMGLLTALAAVVATLVFLPAATASAAATCAPAWNASSVYTGGGSASYNGHNWSA
KWWTQNERPGTSDVWADQGACGSGGGTDPNPSGFVVSEAQFNQMFPSRNSFYTYSG
LTAALSAYPAFANTGSDTVKKQEAFLANVSHETGGLVHIVEQNTANYPHYCDTSQS
YGCAPAGQAAYYGRGPIQLSWNFNYKAAGDALGIDLLGNPWQVEQNASVAWKTGLWY
WNTQSGPGTMTPHNAIVNGSGFGETIRSINGSIECNGGNPGQVQSRVNTYQSFVQILGTT
PGSNLSC*

>SACTE_5764|Chitinase|GH18|GI:344318865 (SEQ ID NO:38)
MRRRSVRALVTAAVTTVAAAGMAVLGSGTAQAATPLPDHVFAPYFESWTGESPAAM
AAESGAKHLTMAFLQTTAKGSCPTYWNGDTGLPIAQASFGADIDTIQAGGGDVIPSGGG
YTADTTGTEIADSDTDVQIAAAQKVVTTYDVSRDMDIEVDSLDDTAGIDRRNKAIK
KLQDWADANGRDLEISYTLPTTTRGLASSGLAVLRNAVTVNGARVDV VNLMTFDYYDN
ASHDMAADTETAAQGLYDQLAKLYPGRATQLWSMVGVTMPGVDDFGPAETFTLAN
AARVYDWAVAKGINTLSFWALQRDNGGCPGPAADDCSGIQQNTWDFTRVFAPFTSG
TTAPDDDFSVTATPASGTVTAGGSATTTVKTAVTKGAAQQVGLTVSGVPAGVTASLSPS
SVTAGGRSTLTLATTQAAVSGTYRISVTGTSPSGSHATA YTLTVTGGTGSQCTAGPWAG
GTVYTGGQQVSYKGH TWKAKWWTGEEP GTTGEWGVWQDLGAC*

>SACTE_4439|Catalase||GI:344317584 (SEQ ID NO:39)
VTQGPLTTEAGAPVADNQNSETAGPGGPVLVQDQALLEKLAHFNRRERIPERVVHARGA
GAYGTFTLTRDVSQWTRAKFLSEVGKETETFLRFSTVAGNLGSADAARDPRGWALKFY
TEEGNYDLVGNNTPVFFIKDAIKFPDFIHTQKRDPYTGSEADNVWDFWGLSPESTHQV
TWLFGDRGIPASFRHMNGYGSHTFQWNNEAGEVFWKYHFKTDQGIKNTTTEEA VRLS
GVDPDSHQRDRESIERGDFPTWTVQVQIMPAAEAA TYRFNPFDLTKVWPHEDYPPIEIG
KLELNRNPNENIFAEVEQSIFSPAHFVPGIGSPDKMLQGR LFAYGDAHRYRVGINADHLP
VNRPHATEARTNSRDGYLYDGRHKGTKNYEPNSFGGPVQTD RPLWQPVSVTGGTGNH
EAAVHAEDNDFVQAGNLYRLMSEDEKGR LIDNLAGFIAKVS RDDIADRINFRQADA
DFGKRLEVAVQALRG*

>SACTE_0562|cellulose-binding family II|GH74|GI:344313814 (SEQ ID NO:40)
VYAMPSTAPAAVQSGEDAPVRSSPRPFAALLAALALTAGLSLIGTPAVARSDEAPAATE
ASDVSAADTYTWKNARIDGGGFVPGIVFNRSKLN LAYARTDIGGAYRWDQSGKQWKP
LLDWVDWDRWGWTVGVVSLASDTVDPDNVYAAVGT YTNSWDPTDGAVLRSSDRGAS
WKAATLPFKLGGNMPGRGMGERLAVDPNKNSVLYL GAPSGNGLWRSTDAGVSWSEV
TAFPNPNGNYAQDPSDTSYGNNDNQIVWVTFDE RSGSAGSATQDIYVGVADKENTVYR
STDGGATWSRIPGQPTGYLAHKGVLDSATGHLYL TSLSDTGGPYDGGKGRIWRYDTASG
AWQDVSPVAEADAYYGFSGLSVDRQKPGTLMATA YSSWWPDTQIFRSTDSGATWTQA
WDYTYGPNRSNRYTLDVSSVPWLSWGASPAPPETAPKLGWMTEALEIDPFSDRMMY

Figure 20 (continued)

GTGATVYGTEDLTSWDSGGTFRITPMVKGIEETAVNDLASPPSGAPLLSALGDIGGFRHT
DLDAVPDLMYTSPNLDSTTSLDFAESSPGTVVRVGNDAAPHIGFSTDNGANWFQSE
SGVTGGGTVAAAADGSGFVWSPEGAGVHHTTGFGTSWTASTGIPAGATVESDRKNPEK
FYGFAGTFYVSTDDGATFTAETGLPAEGNVRFQALPGTEGDIWLAGGSDTGAYGLW
RSTDSGATFTKSAGVEQADSVGFGKAAPGASYRTVFVSAKIGGVRGIFRSTDAGASWTR
INDDAHQWGWGTGAAITGDPRVYGRVYVSTNGRGIQVGETSDSGGGGTDPGTDPGTDPG
TDPGPEQPADAACAVTYAVTNQWPGGFQADVTVTNTGDAAYNWGWKLGSFPGGQQIS
QIWNASHRQDGVKVTVTDAGWNGTVAPGSSAGFGFTGSWAGSNAEPAFTLDGQACT
VG*

>SACTE_4343|extracellular solute-binding protein family 5||GI:344317489 (SEQ ID NO:41)
MRGAKSAKWVAGAAIIALAATACGGGSDSDNGAKGAVDADGIFSVEVGEQNPQLQP
ANTMESNGSIVTDAIFSQLVDYDPDGKLEMINAESVETDTSKLWTVKLKDKWKFHDGT
PVTADSYVKAWNWAANIENAQTNASWFADIKGYADVHPDGEGAKPKSDAMSGLKKV
DDYFTTIELNSAVPYFSYKLGTVFSPLPESFYADPKAAGEKPVGNNGAYKFFVSWDHKKQ
IKVVRNDDYKGPDKAKNGGVIFKNYTTLETAYEDLKSGNVVDVLRQIGPKDLPVYRADL
EDRAVDKAYSQVTLGVAMYTDQWKNTDPKVLQGLSMAIDRDTITKTVLQGTREPAT
GWVAKGVVLYQENVAGDVTKYDPAKAKALIKEGGGVPNGEIQFNADGGHKEWIEA
VCNSITQATGVKCTGDSKADFQADLNARDAKQVKSFYRSGWVLDYPVNFANFISDLFRT
GAAGNNGFFSNKDLDAKIKAAADSAASLDDSVKAYQEIEKELVNYMPSIPLWYYKVNAG
YSENVKNVDYAQDGDPILETVQVIK*

>SACTE_1546|bacterioferritin||GI:344314774 (SEQ ID NO:42)
MQGDPEVLEFLNEQLTAELTAINQYFLHAKMQDHRGWTKLAKHTRAESFDEMKAHEIL
TDRILLLDGLPNYQRLFHVRVGQTVTEMFQADRQVEVEAIDRLRRGVLDLMRAKSDITSA
NIFERILEDEEHIDYLDTQLELIEKLGEPYLAQVIEQVEL*

>SACTE_3590|phosphatidylinositol-specific phospholipase C X region||GI:344316754 (SEQ ID
NO:43)
MSPYTATRRFTLTGALAAATGVVVGTPALAAPARVLGTQDWMGALADSTPLRRLTIP
GTHNAGARYGGPWTECQNTTVAEQLGSGIRFLDVRCRITGDAFAIHHGASYQNLMFQD
VLIACRDFLAHPSETVLMRVKQEYSESDAAFRQIFDLYLDGKGWRPLFRLDPTLPDL
GGARGKVLLADNGLPGVRYADPAVFDIQDDYMAEPFGKYPKIEAQFRKAAQQPGK
LFMNYVSTAALLPPRSNADRLNPQVHTFLDGSEAAGWTGLGIVPLDYPATRPGLVESLI
RHNPVA*

>SACTE_2172|citrate synthase I||GI:344315379 (SEQ ID NO:44)
VSEHTNNAVVLRYGDDEYTPVIDSTVGDKGFDIGKLRANTGLVTLDSGYGNTAAYKS
AITYLDGEQGILRYRGYPIEQLAESSTFLEVAYTLINGDLPKVDELSAFKNEITQHTLLHE
DVKRFFDGFPRDAHPMAMLLSSVVSALSTFYQDSHNPFDDEQRHLSTIRLLAKLPTIAAYA
YKKSIGHPFVYPRNDLGYVENFLRMTFSVPAQYVVPDPIVVSALAKLLILHADHEQNCST
STVRLVGSSQANMFASISAGISALWGPLHGGANQSVLEMLEGIQANGGDVDSFIQKVK
KEDGVRLMGFGHRVYKSFDPRAKIIKAAAHDVLSLGSDELLELDIALKLEEHALSDDYF

Figure 20 (continued)

VSRNLYPNVDFYTG LIYRAMGFPTMFTVLFALGRLPGWIAQWHEMIKEPGSRIGRPRQI
YTGEVLRDFVPVESR*

>SACTE_5668[TAP domain protein][GI:344318769 (SEQ ID NO:45)

MTKRAGILVAVGATVAGLVTAVPSAASTAPGAPGAAAPLKWTACGTKAYPTQQCATV
RAPLDHDRPSGRQVTLALARIPHTAKTSQGPLLVPNGGPGGSGLSMAGFVASSLPAKLA
AQYDVIGFDPRGVGRSSPALDCVPHFDPVRPDTVPGSPRDERTNRERAASFADACGEK
HGDLLPFMDTVSTAKDLDVIRRALGARQINYFGYSYGTYLGA VYAKLFFPERVRRLLVD
SIVDPDG VVYEDNLGQDYAFDARHKAFAAWVAKNDATYRLGTDPAKVEAAWYRMR
AAVKKHFAAGKVG PSELEDTF LGGYNGYWPQLAEFAAYVNDKDEDALATAYDDF
AAVDASGDNGYSVYTA VQCRDTGWPKSWTTWRNDTWQ AHRKAPFMSWNNTWYNAP
CATWPVAPLRPVRVTNREIPPALLFQATDDAATPYEGGLSMHRKCLKGSRLVVEEGGN
HGISLSGNDCLDAHLIAYLTDGTLPRSGGSGADAVCDALPEPEAAATAKAKAATGQKGS
TLHSLLGFRG*

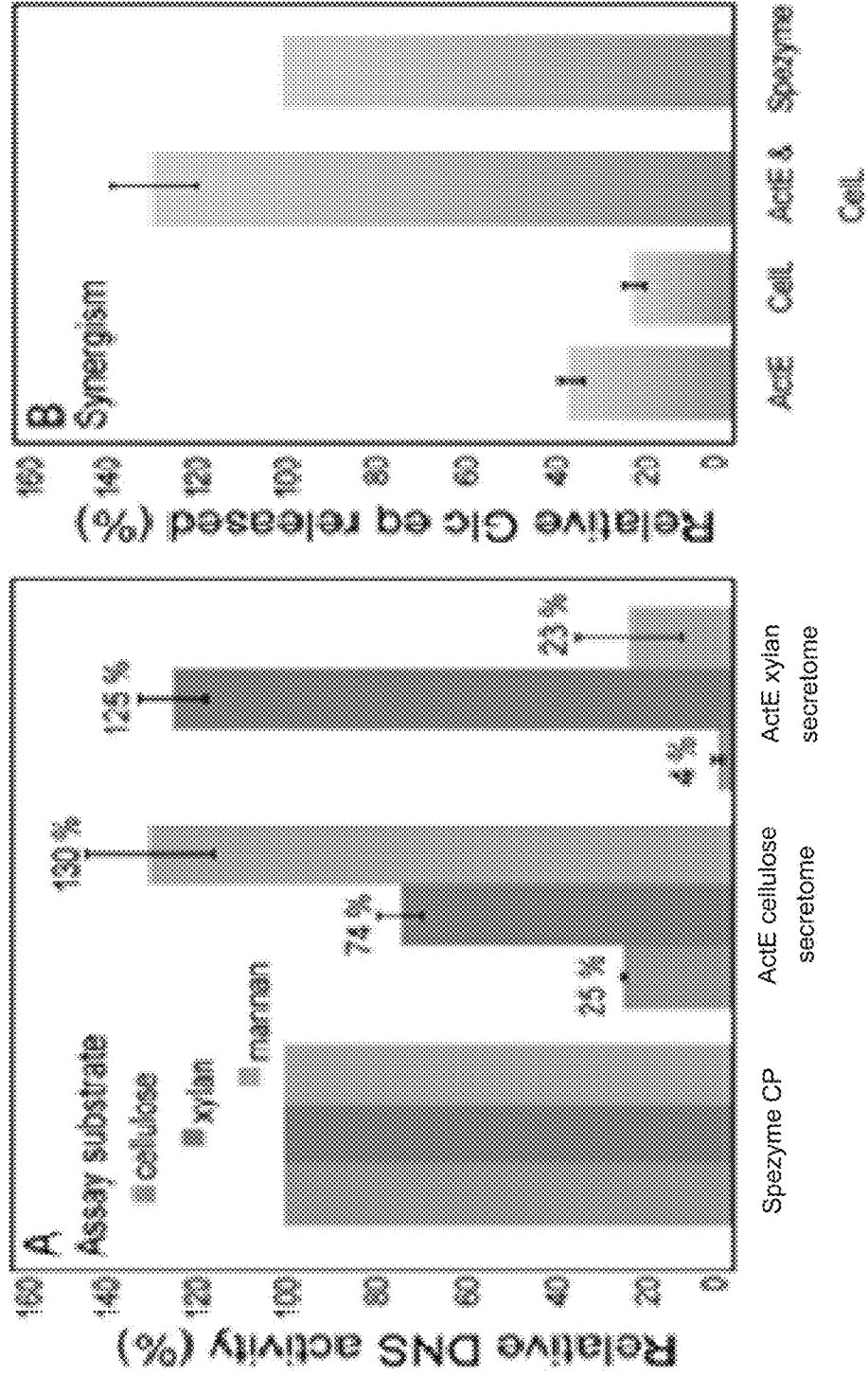
>SACTE_6428[chitin-binding domain 3 protein][CBM33][GI:344319509 (SEQ ID NO:46)

MNCHDRINLRGWTTRLSGLFVA AVLCLLPWTGTAEAHGSVVDPASRNYGCWLRWGS
D FQNPAMAQEDPMCWQAWQADPNAMWNWNGLYRNE SAGNFP AVIPDGQLCSGGRTE
GGRYNALDTVGAWQATDITDDFTVRLEDQASHGADYFRVYVTEQGFDP TAQPLTWGA
LDLVAETGRYGPSTSYEIPVSTSGYTGRHVYTIWQASHMDQTYFLCSDVNFG*

>SACTE_0366[alpha-L-rhamnosidase][GH78][GI:344313621 (SEQ ID NO:47)

VISRRRLSTTAATAALAAVSSPAARAAAPADTAAGR LRVGTGPTVEYVRRPLGLDVS
RPLSWPLASDHPDHGQSA YQVRVATSPDRLARPDVWDSGKVVSP TSVLVPYAGPALVSR
TRYHWSVRVWDQGRVSAWSEPSWWTGLLDEADWSAGWIGAPAALTSSPSLEAAS
WIWFPEDPAV GAPAATR WFRGRVEIPEGVTRARLVMTADDGFTALVDG VQVARTEP
DGPAENWRRPVVDVTAHLSPGSRVVAVTATNAV DGPAGLLGALELTTADGAVTLAT
GTGWRA TDREPDGDWASGGYDDTGWPAAAVLAPWGS GPWGEVRAALSPATQLRTEF
RLGRKRVARARLYSTALGLYEVFLNGARVGEDRLAPGWTDYRKR VQYQTYDVTALLR
SGGNALGVT LAPGWYAGNIAWFGPHQYGERPAVLAQLEVTFTDGSIERVLSGTGWAAA
TGPVTATDLMAGEEYDARLETDGWSRAGFDASGWLAAEA VEGVTA VPVAAVDGACR
VERELTAREVTEPEPGVYVFDLGQNMVGTVRLLVSGPAGTTVRLRHAEVLNPDGTYT
ANLRTARATDTYTLRGGGPETYEP RFTFHGFRYVEVTGFPGRPGPDAVVGRVIHTSAPF
TMAFSTDVPM LDRLHSNITWGQRGNFLSPTDTPARDERL GWTDINVFAPTAAYTME
SARFLGKWLQDLRDDQLADGAPNVAPDLPGVGS GAAGWGDAGVTPWALYQAYGD
VRVLEQSWSSMVAWLEYLQAHS DGLLRPADGYGDWLNIEDETPKDVIGTAYFAHSAD
LTARTAEVLGKDPGPYRTL SGRVRDAFRAAYVGDGGRVKGDTQTAYVLALSMDLLEP
GDRAPAADRLVALIEAKDWHLSTGFLGTPRLLPVLTD TGHTDVAYRLLTRRTFPSWGY
QIDRGATTMWERWDSVRPDGGFQDAGMNSFNHYAYG SVGEW MYANIAGIAPAAPGF
REIRVRPRPGGVHRAEARFDSL YGPVTTTRWTS DGGGFALRVVLPANTTAEVWVPGD
GRSSVRGTAVFLRREDGCAVFAAGSGIHRFTAPA*

Figure 21 A-B



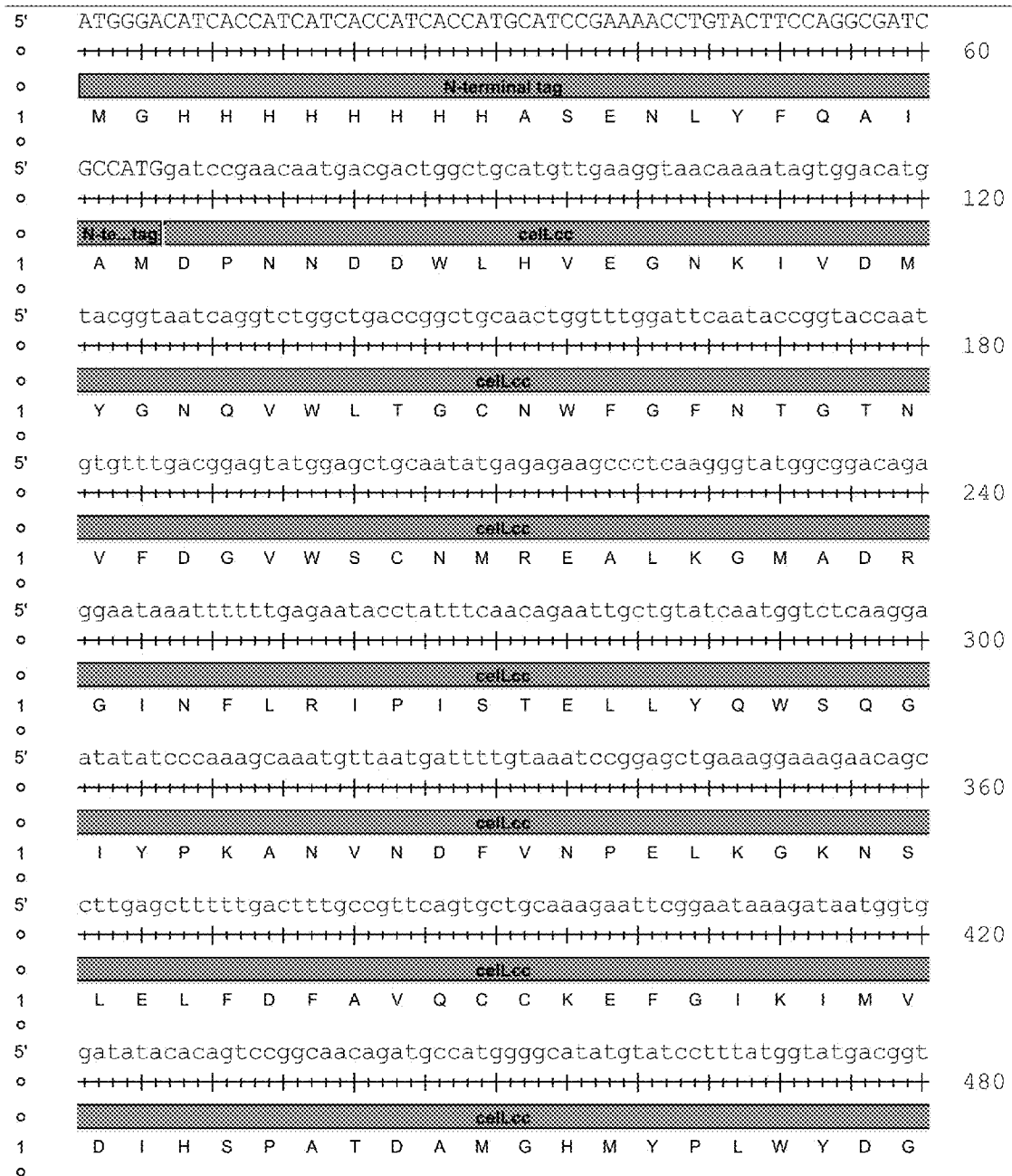


Figure 22 (SEQ ID NOs: 63 & 64)

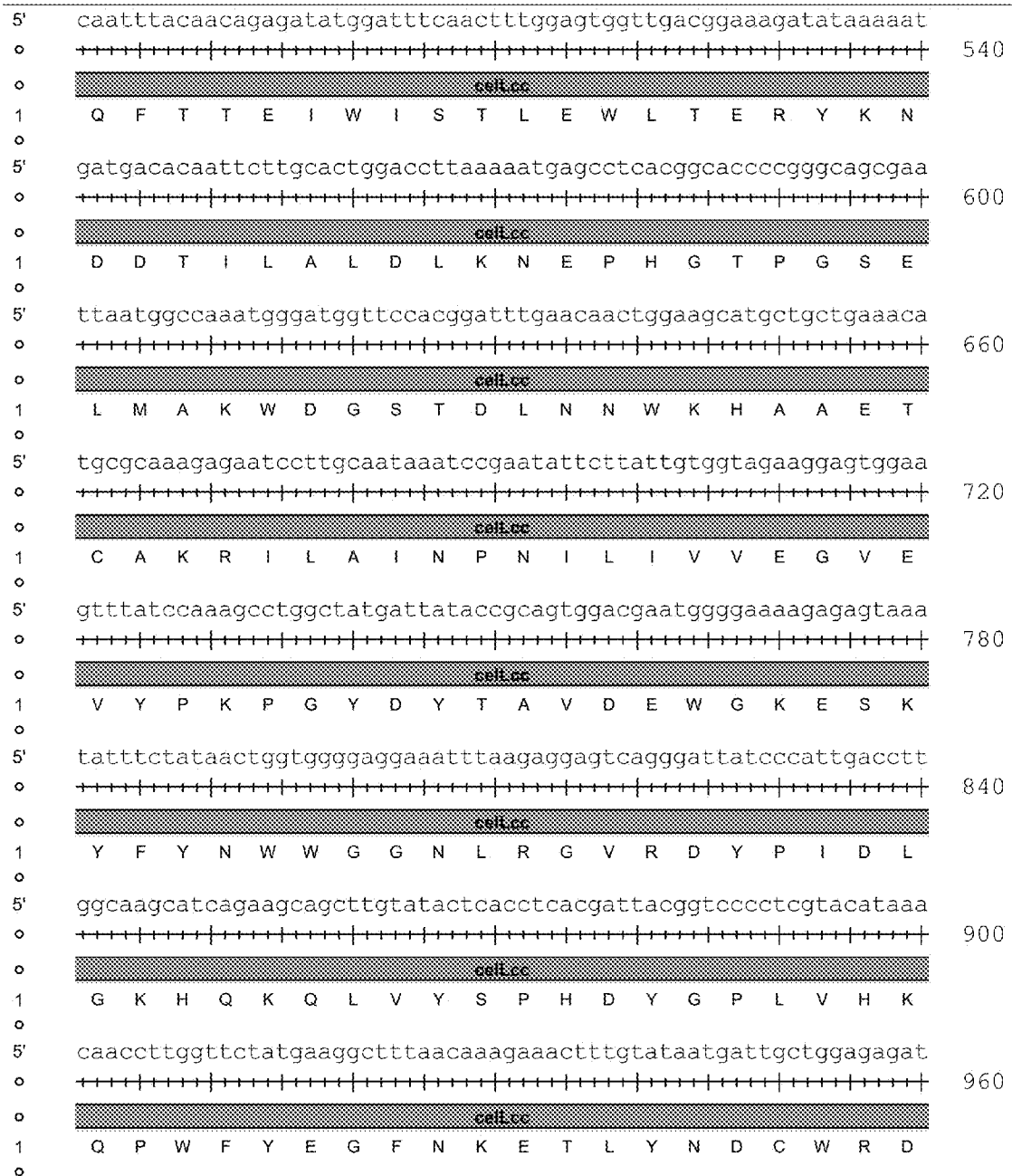


Figure 22 cont'd (SEQ ID NOs: 63 & 64)

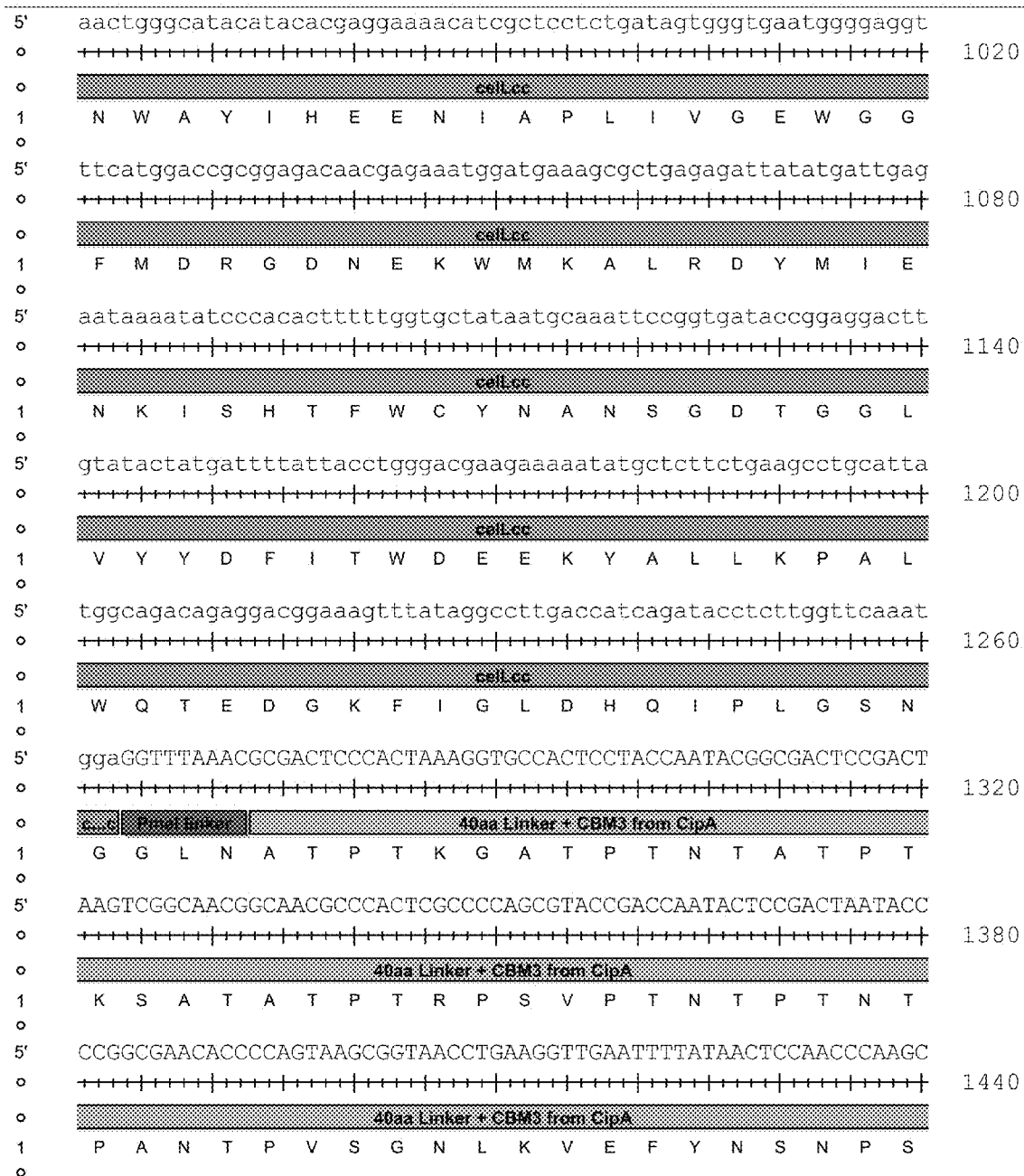


Figure 22 cont'd (SEQ ID NOs: 63 & 64)

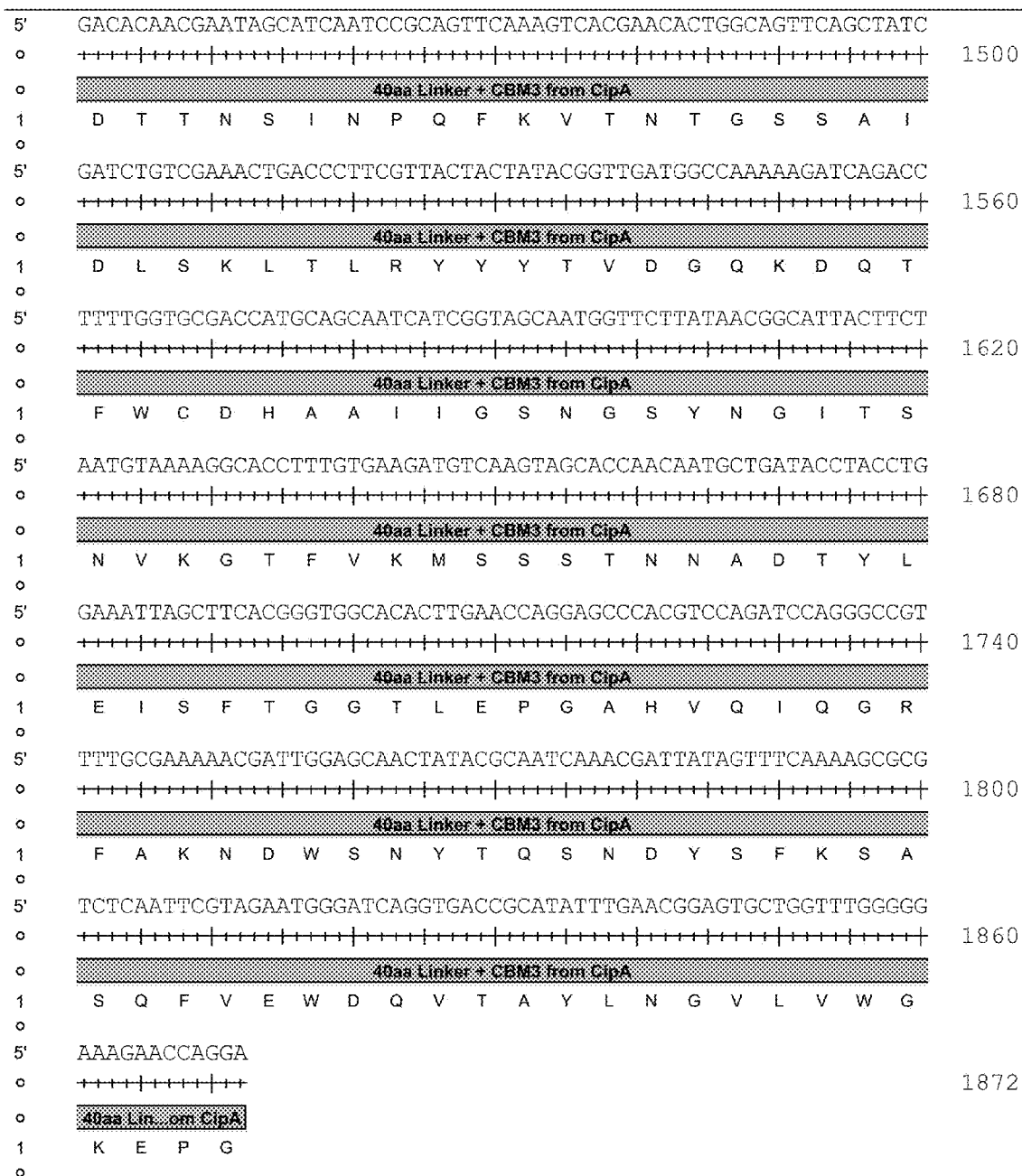


Figure 22 cont'd (SEQ ID NOs: 63 & 64)

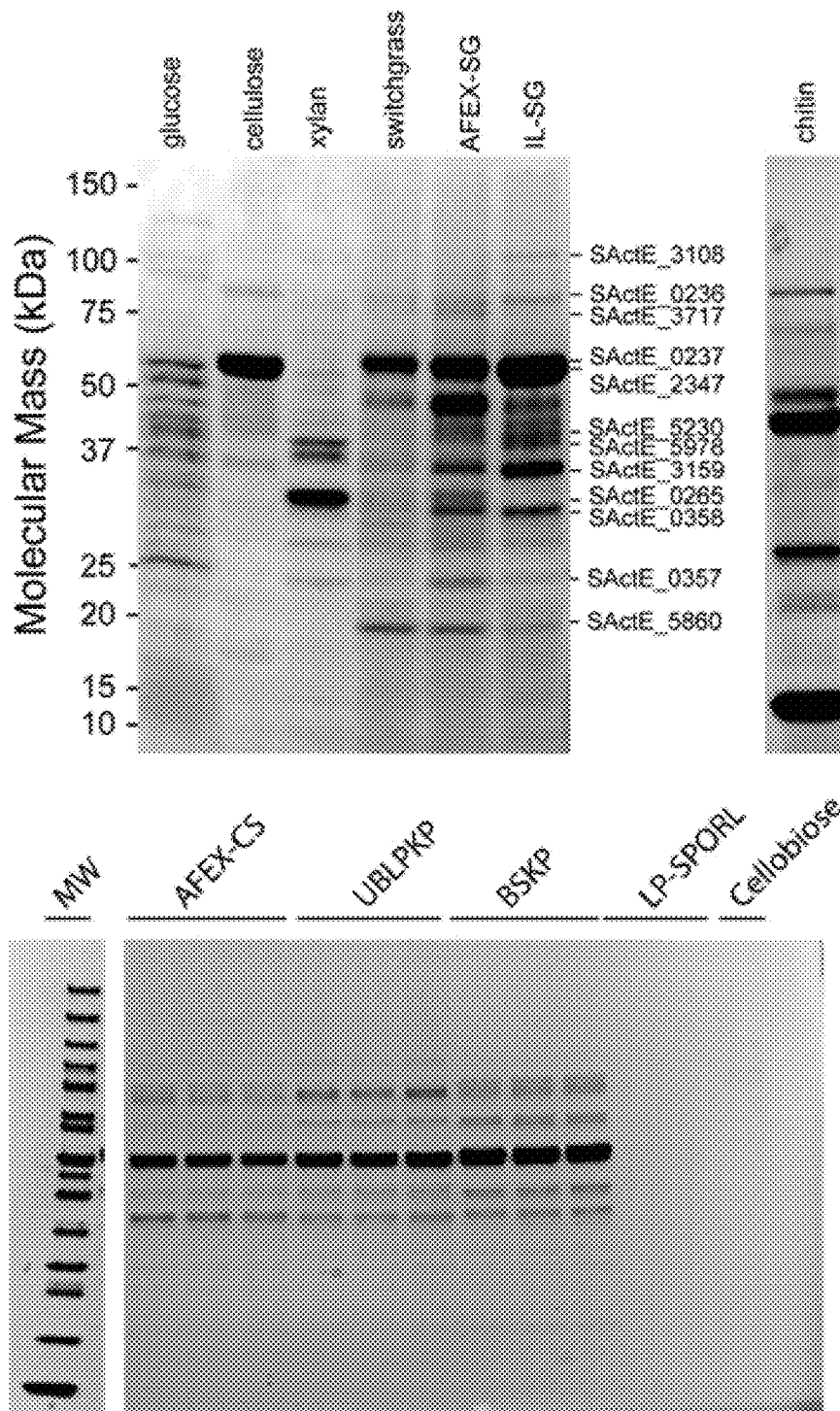


Figure 23

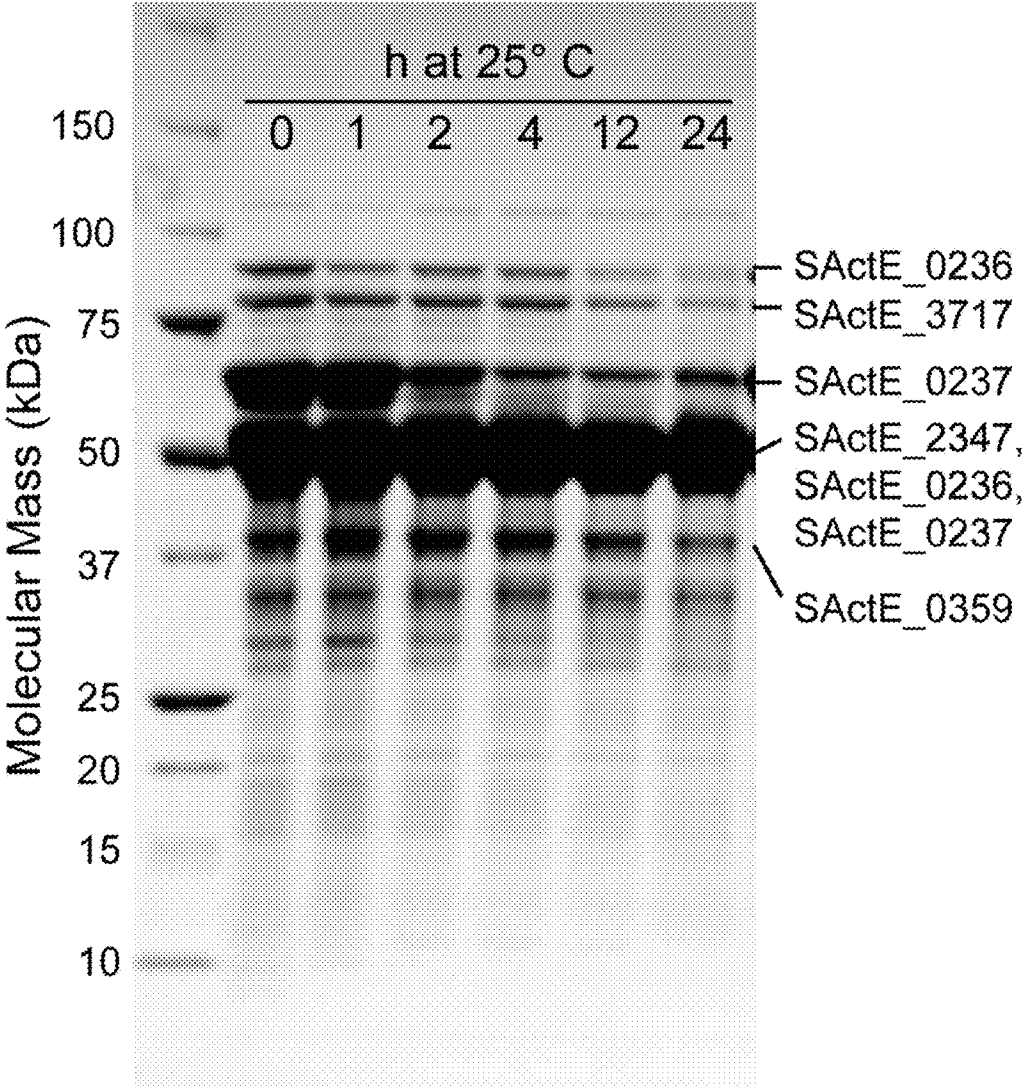
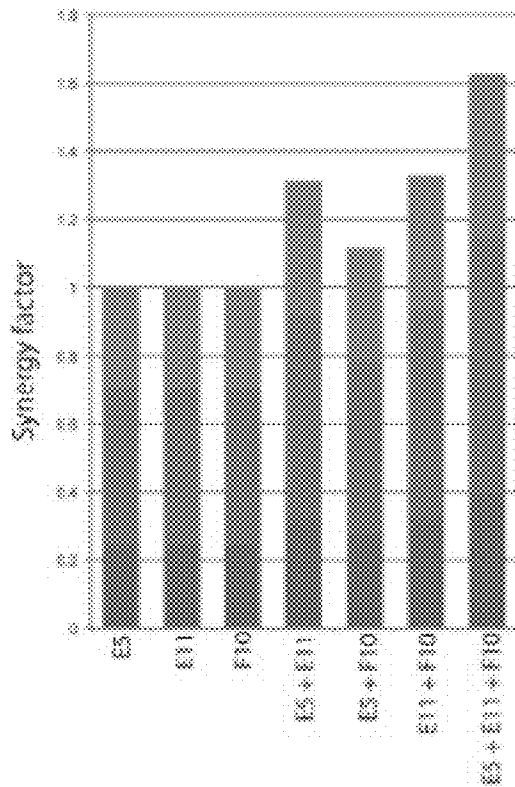


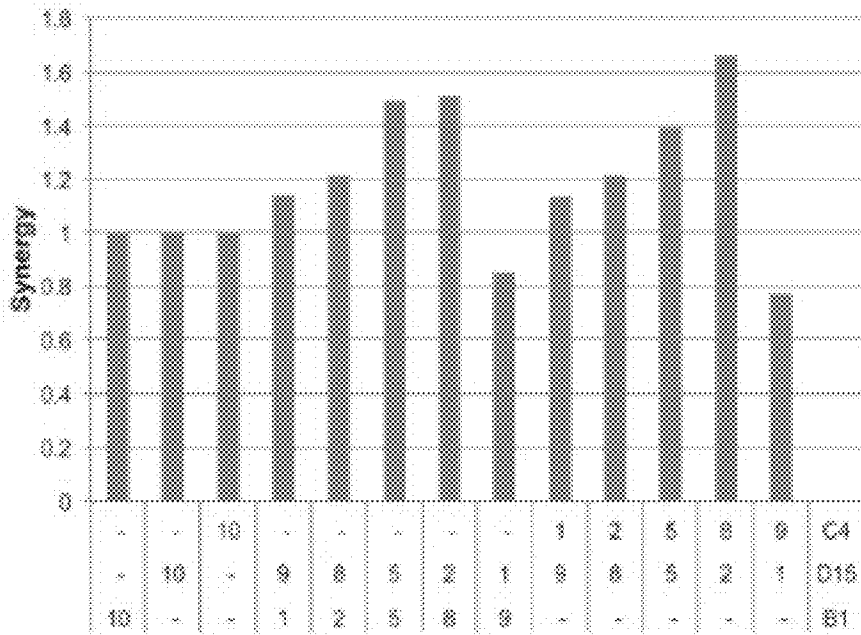
Figure 24

A



Fraction Number from Figure 28

B



Fraction Number from Figure 7

Figure 25 A-B

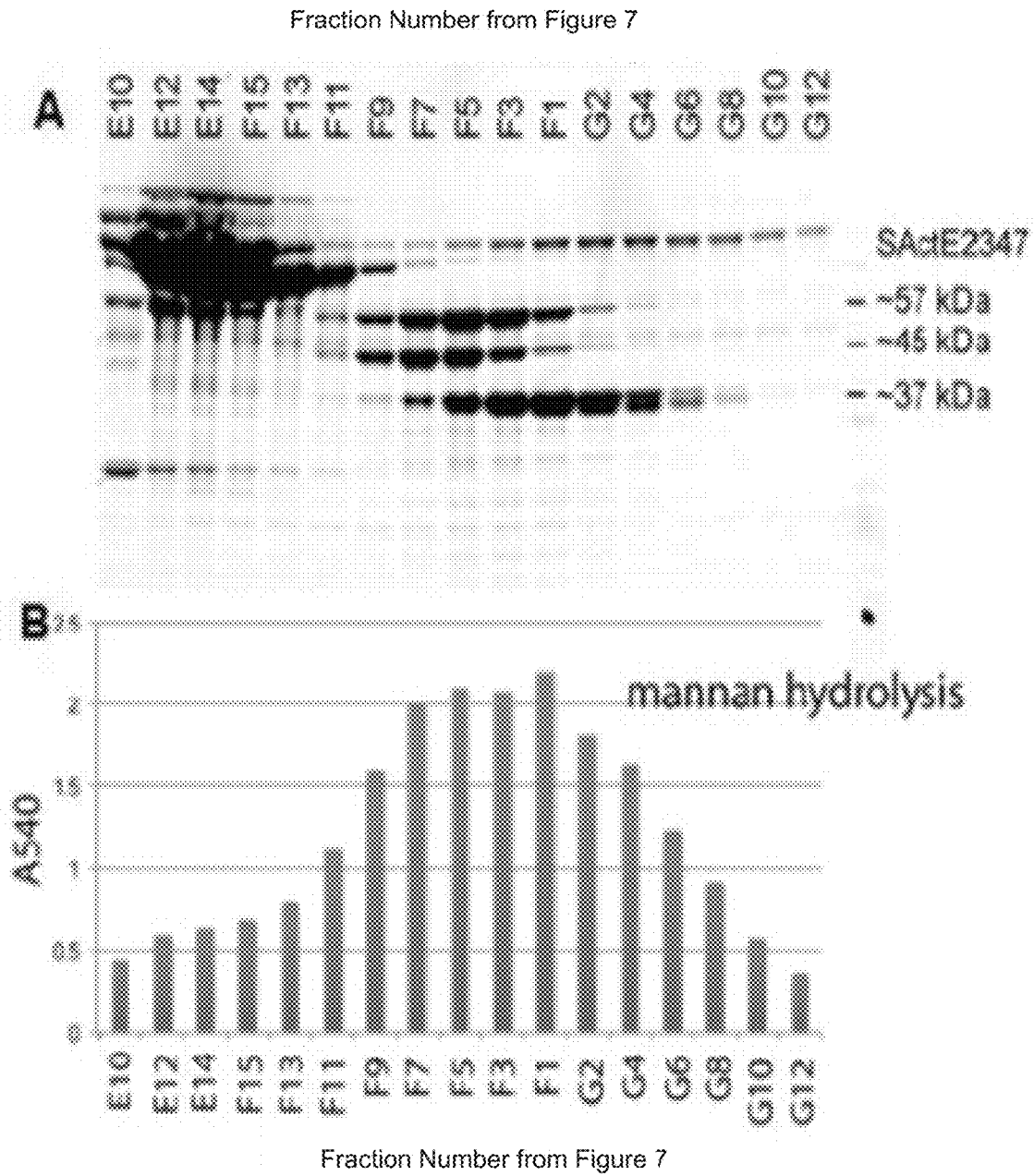


Figure 26 A-B

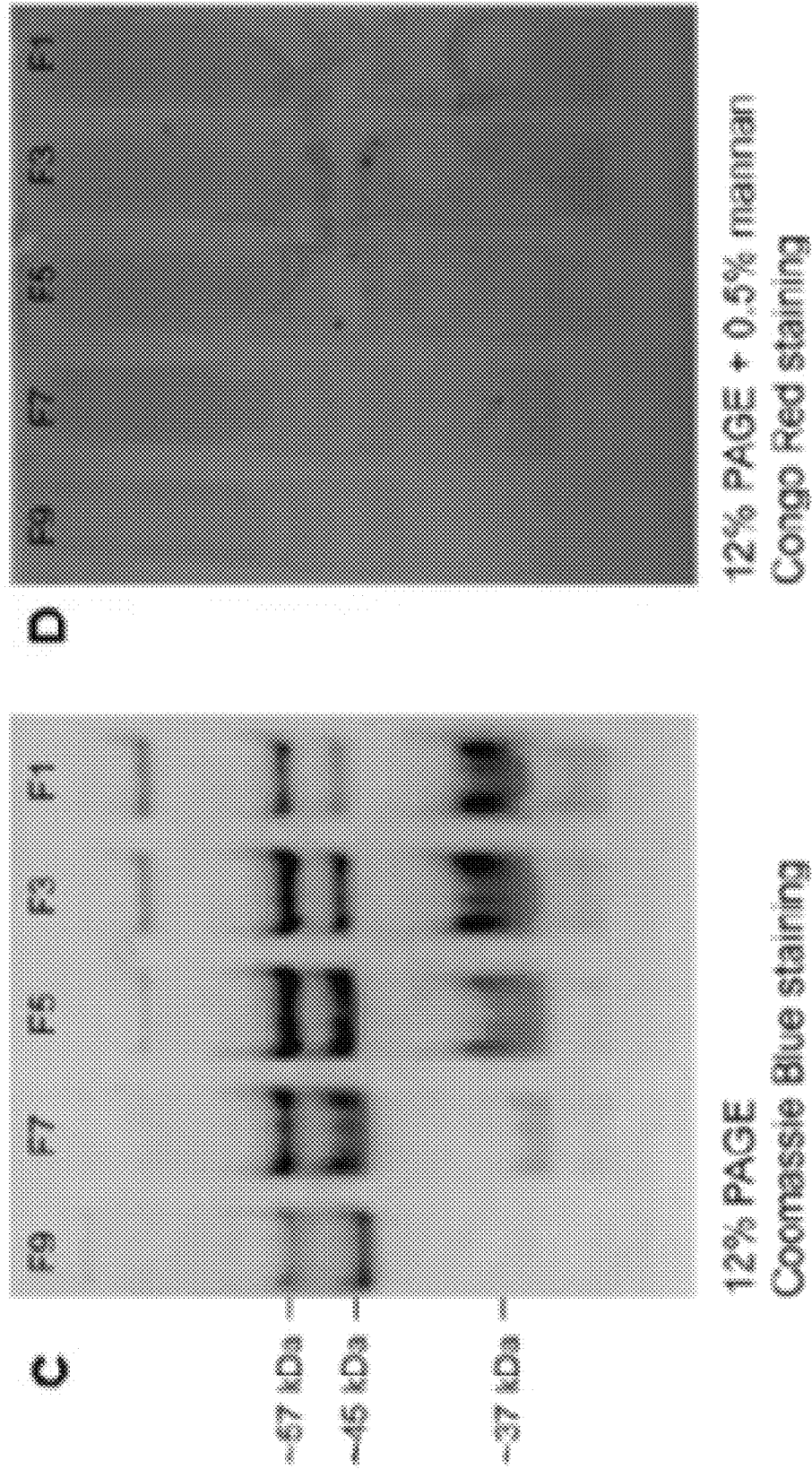


Figure 26 C-D

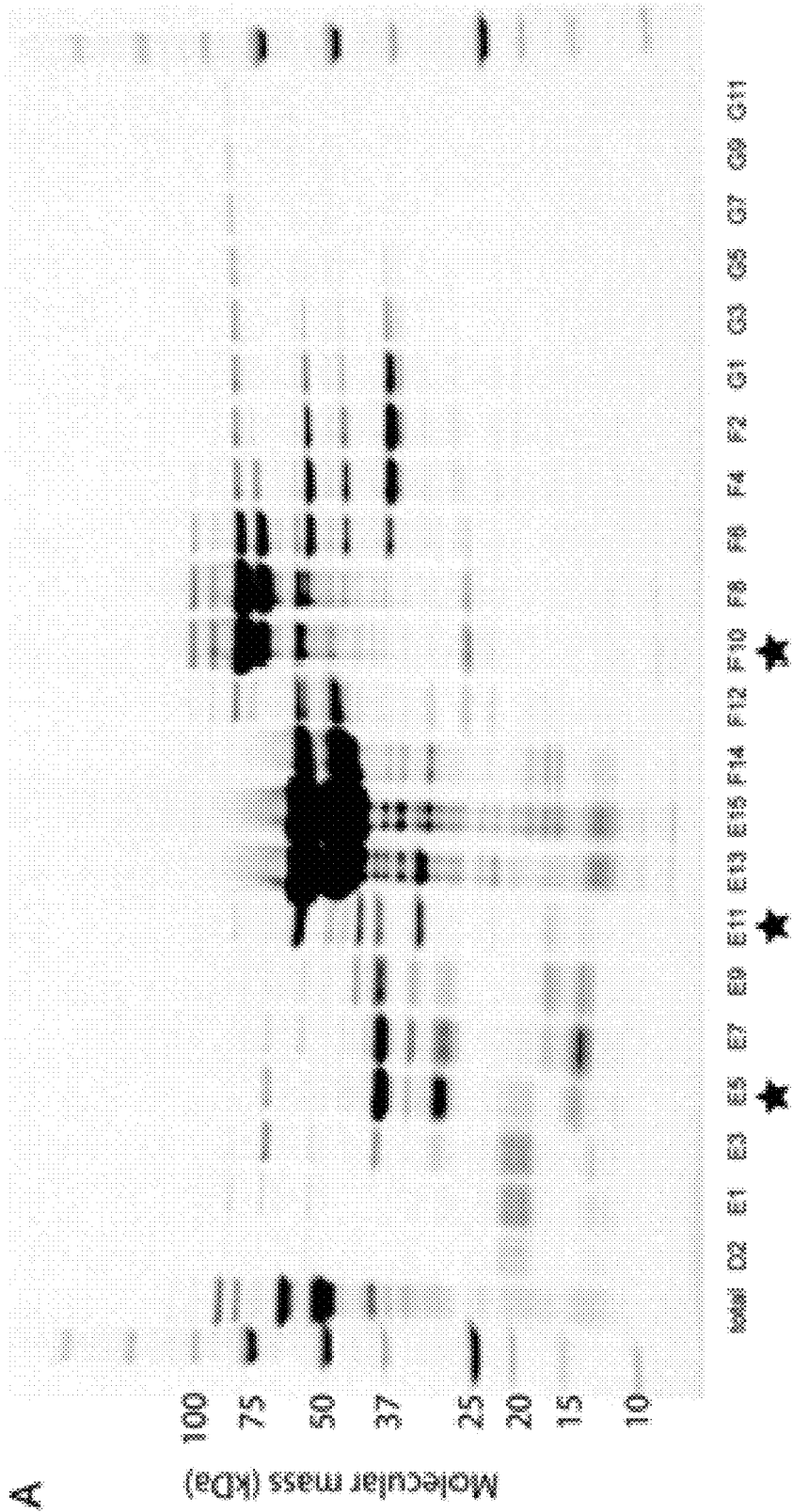


Figure 27A

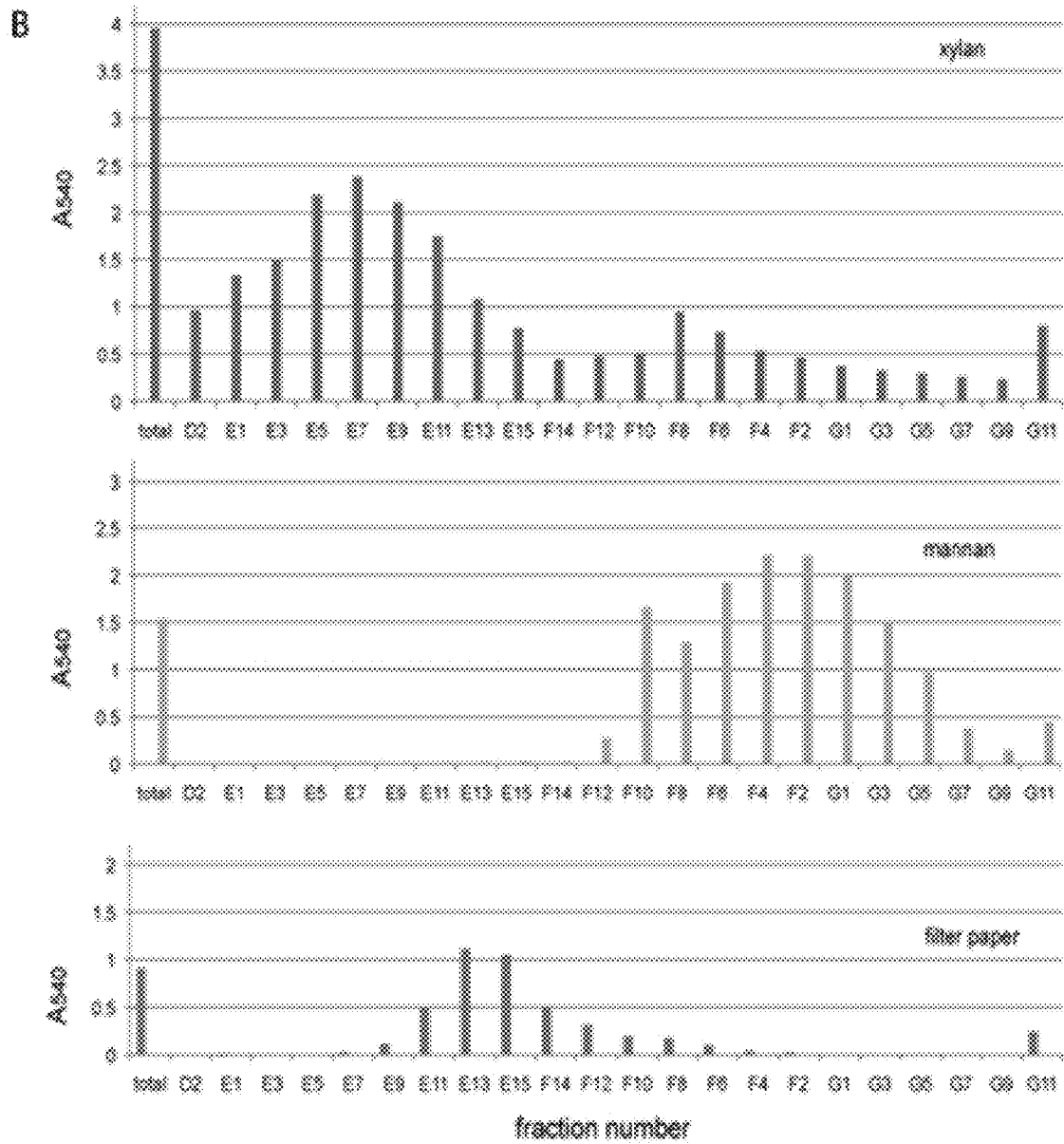


Figure 27B

Figure 28A

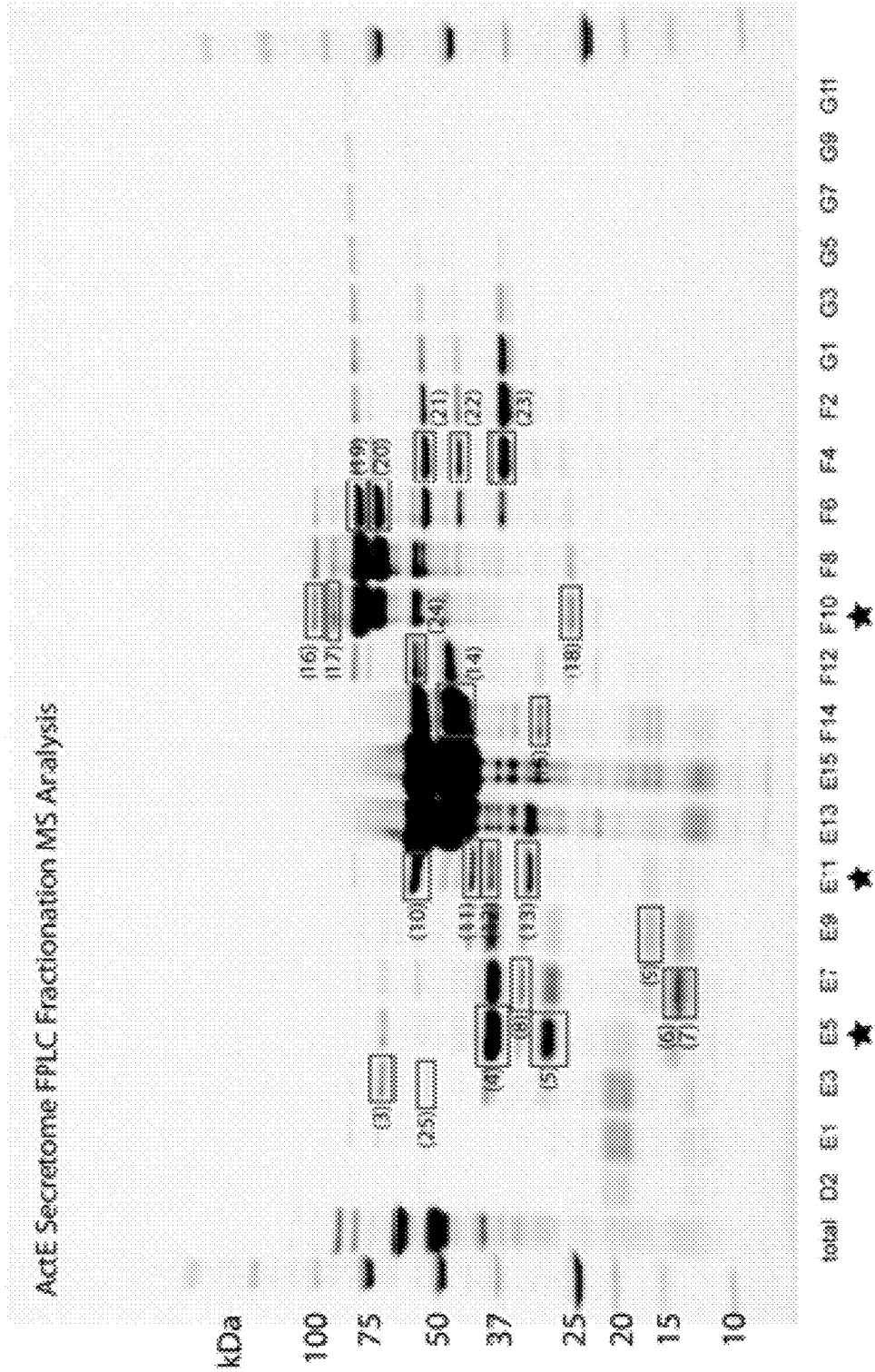
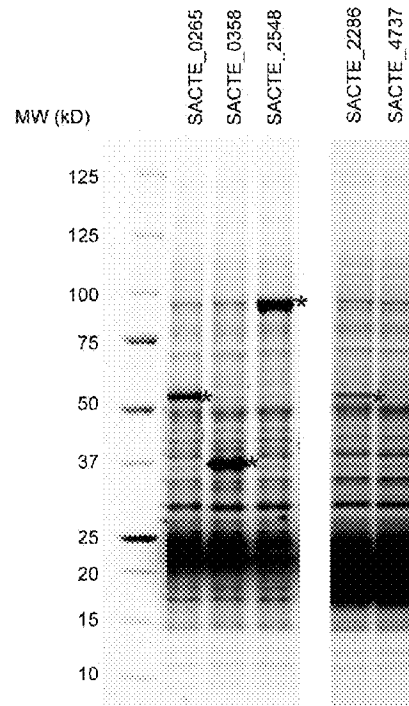


Figure 28B

MS #	ID	Function	CAZY	CBM	AA	Best BLAST
3	SACTE_4738	glycoside hydrolase family 16	GH16	CBM32	627	beta-1,3-glucanase
4	SACTE_3159	chitin-binding domain 3 protein	CBM33.2	CBM33.2	363	cellulose oxygenase
5	SACTE_3159	chitin-binding domain 3 protein	CBM33.2	CBM33.2	363	
6	SACTE_3159	chitin-binding domain 3 protein	CBM33.2	CBM33.2	363	
7	SACTE_3159	chitin-binding domain 3 protein	CBM33.2	CBM33.2	363	
8	SACTE_0265	glycoside hydrolase family 10	GH10	CBM2	459	xyfanase
9	SACTE_5083	putative RNA polymerase	#N/A	#N/A	418	
10	SACTE_0237	1, 4-beta cellobiohydrolase	GH6	CBM2	587	1,4-beta cellobiohydrolase
11	SACTE_0482	glycoside hydrolase family 5	GH5	CBM2	457	endo-1,4-beta-glucanase
12	SACTE_4755	beta-1,3-glucanase	GH64		409	beta-1,3-glucanase
13	SACTE_0482	glycoside hydrolase family 5	GH5	CBM2	457	
14	SACTE_0237	1, 4-beta cellobiohydrolase	GH6	CBM2	587	1,4-beta cellobiohydrolase
15	SACTE_0549	glucan endo-1,3-beta-D-glucosidase	GH16	CBM54	307	beta-1,3-glucanase
16	SACTE_0236	glycoside hydrolase family 48	GH48	CBM2,37	955	cellulose 1,4-beta-cellobiosidase
17	SACTE_0236	glycoside hydrolase family 48	GH48	CBM2,37	955	
18	SACTE_5457	chitosanase	GH46		290	chitosanase
19	SACTE_0236	glycoside hydrolase family 48	GH48	CBM2,37	955	cellulose 1,4-beta-cellobiosidase
20	SACTE_3717	carbohydrate-binding, Ceric-like	GH9	CBM2,4	909	endo-1,4-beta-glucanase
21	SACTE_2347	cellulose-binding family II	GH5,CE3	CBM2,37	563	secreted beta-mannosidase
22	SACTE_2347	cellulose-binding family II	GH5,CE3	CBM2,37	563	secreted beta-mannosidase
23	SACTE_2347	cellulose-binding family II	GH5,CE3	CBM2,37	563	secreted beta-mannosidase
24	SACTE_5629	Ricin B lectin	GH93	CBM42,13	593	exo-alpha-L-1,5-arabinanase
25	SACTE_4363	putative secreted protein	GH55	CBM56,54,57	606	endo-beta-1,3-glucanase

A



B

Gene Locus	CAZy	MW (kDa)	Microarray rank ^a			Present in secretomes	Diagnostic substrate			
			cellulose	xylan	chitin		MUG	MUC	MUM	MUX2
SACTE_0265	GH10	49.8	20	519	3530	yes	-	-	-	+
SACTE_0358	GH11	37.2	13	160	593	yes	-	-	-	+
SACTE_2548	GH1	90.8	4197	4135	5330	no	-	-	-	-
SACTE_2286	GH2	55.3	28	2533	3012	no	+	-	-	-
SACTE_4737	GH1	52	702	791	1718	no	+	-	-	-

^a Out of 6152 genes total, ranking by transcript intensity, with highest rank equal 1.

Figure 29 A-B

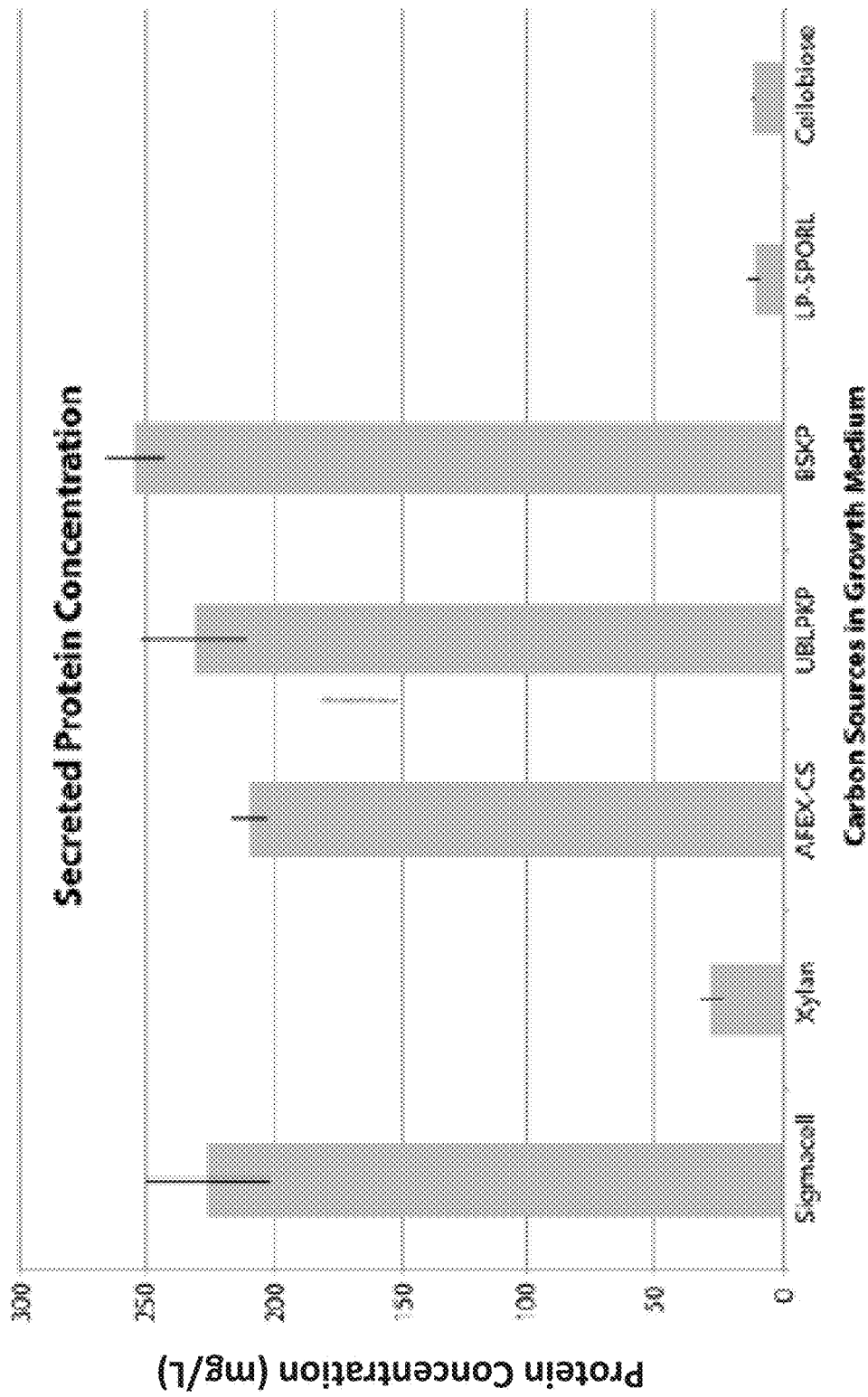


Figure 30

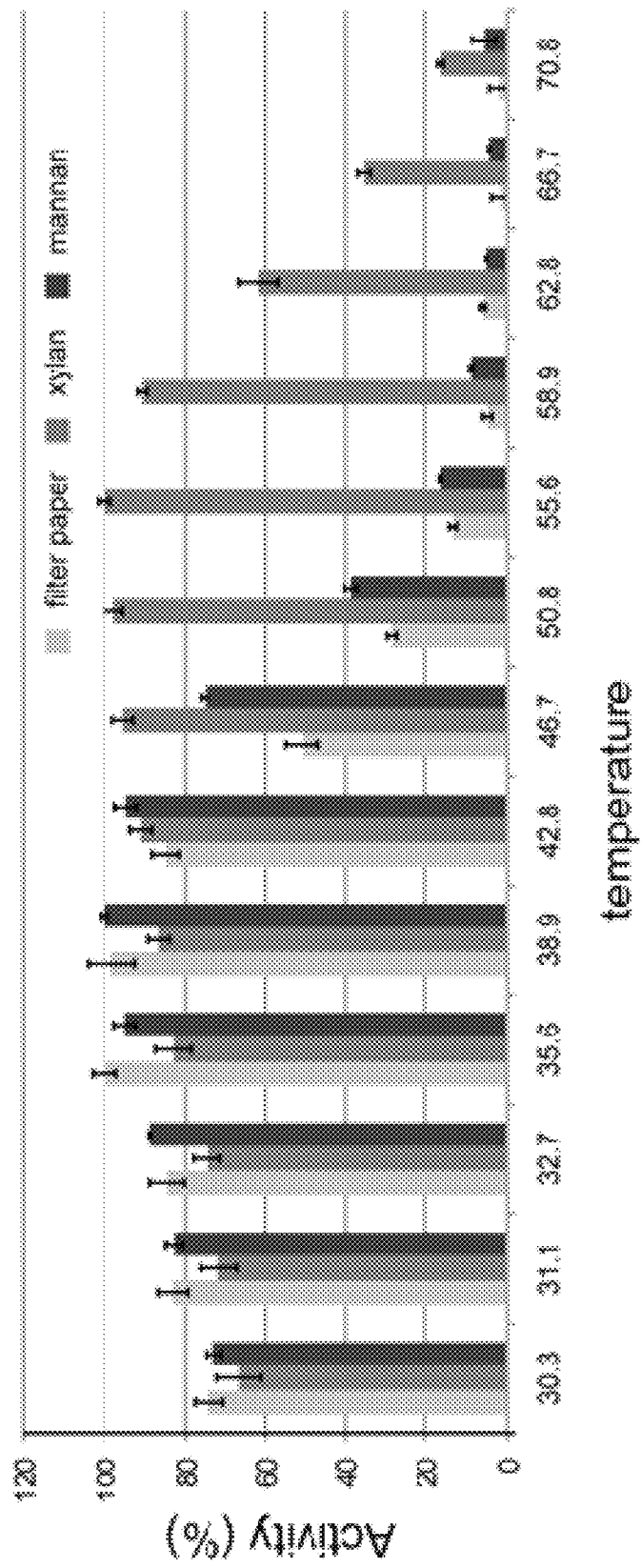


Figure 31

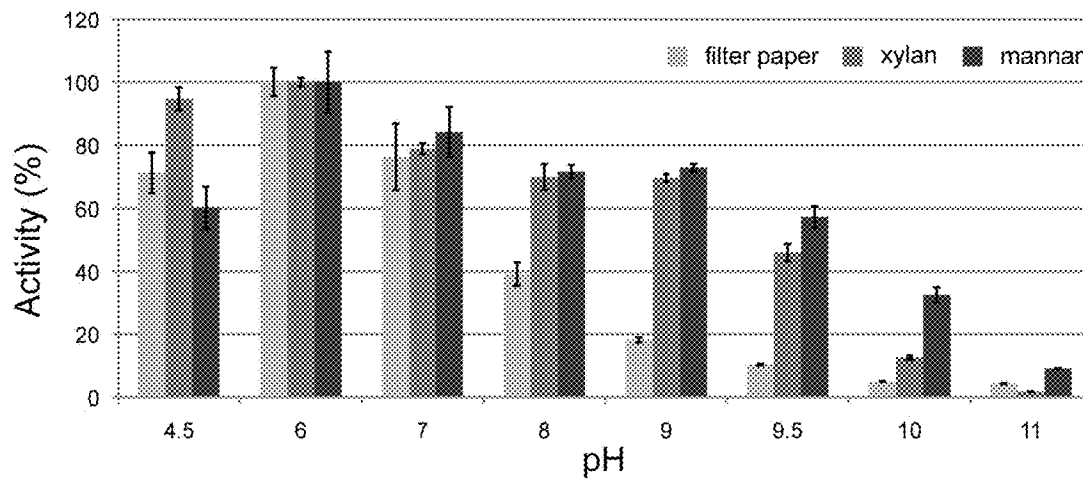


Figure 32

Cellulose/Glucose
(percent of total carbon)

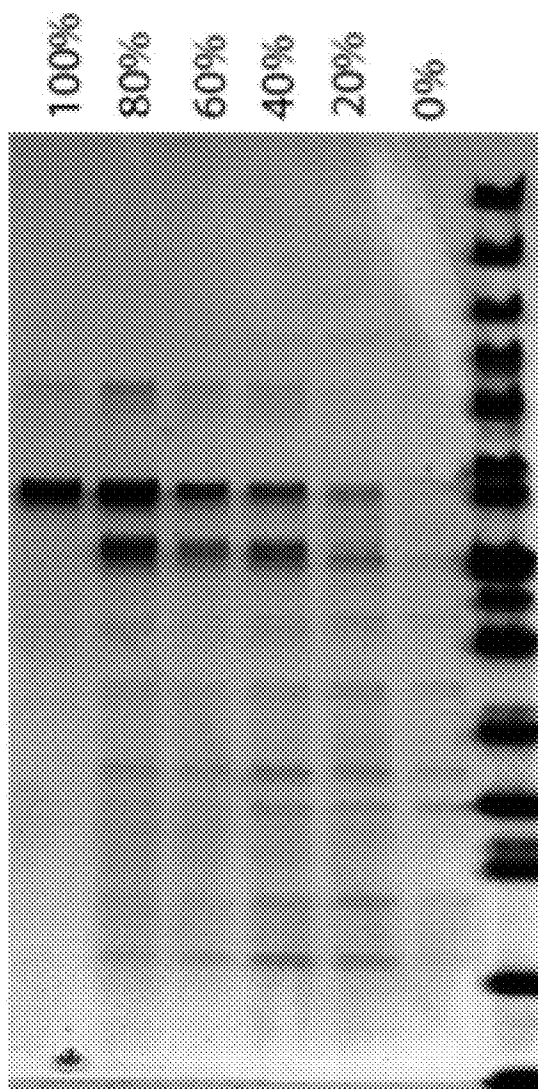
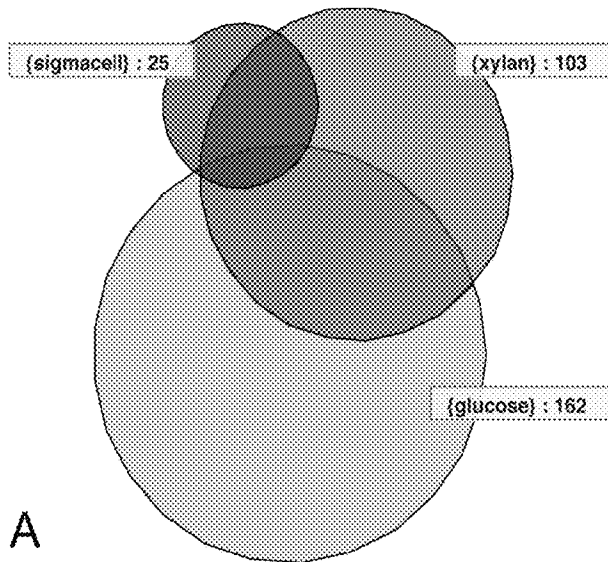


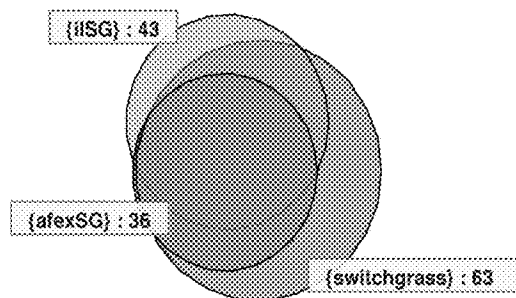
Figure 33



$$\begin{aligned} \text{glucose} \cap \text{sigma-cell} &= 4 \\ \text{glucose} \cap \text{xylan} &= 45 \\ \text{sigma-cell} \cap \text{xylan} &= 16 \\ \text{glucose} \cap \text{sigma-cell} \cap \text{xylan} &= 4 \end{aligned}$$

$$\begin{aligned} \text{glucose} / (\text{sigma-cell} \cup \text{xylan}) &= 117 \\ \text{sigma-cell} / (\text{xylan} \cup \text{glucose}) &= 9 \\ \text{xylan} / (\text{glucose} \cup \text{sigma-cell}) &= 46 \end{aligned}$$

A



$$\begin{aligned} \text{switchgrass} \cap \text{afexSG} &= 36 \\ \text{switchgrass} \cap \text{iISG} &= 35 \\ \text{afexSG} \cap \text{iISG} &= 27 \\ \text{switchgrass} \cap \text{afexSG} \cap \text{iISG} &= 27 \end{aligned}$$

$$\begin{aligned} \text{switchgrass} / (\text{afexSG} \cup \text{iISG}) &= 19 \\ \text{afexSG} / (\text{iISG} \cup \text{switchgrass}) &= 0 \\ \text{iISG} / (\text{switchgrass} \cup \text{afexSG}) &= 8 \end{aligned}$$

B

Figure 34A-B

METHOD AND COMPOSITIONS FOR IMPROVED LIGNOCELLULOSIC MATERIAL HYDROLYSIS

CROSS-REFERENCE TO RELATED APPLICATIONS

This application claims benefit from U.S. Provisional Application 61/579,301 filed Dec. 22, 2011 and U.S. Provisional Application 61/579,897 filed Dec. 23, 2011, both of which are incorporated herein by reference for all purposes.

STATEMENT REGARDING FEDERALLY SPONSORED RESEARCH OR DEVELOPMENT

This invention was made with government support under DE-FC02-07ER64494 awarded by the US Department of Energy and GM094584 awarded by the National Institutes of Health. The government has certain rights in the invention.

BACKGROUND

Cellulose is the most abundant organic polymer on Earth and represents a vast source of renewable energy. Most of this energy is stored in the recalcitrant polysaccharide cellulose, which is difficult to hydrolyze because of the highly crystalline structure, and in hemicellulose, which presents challenges because of its structural diversity and complexity. Plant cell walls are approximately composed in pinewood of lignin (30% by weight), hemicellulose (glucomanan, 20%, arabinoxylan, 10%), and crystalline cellulose (40%), which presents a major barrier to efficient use. In terrestrial ecosystems, cellulolytic microbes help drive carbon cycling through the deconstruction of biomass into simple sugars. The deconstruction is largely accomplished through the action of combinations of secreted glycoside hydrolases (GHs), carbohydrate esterases (CEs), polysaccharide lyases (PLs), and carbohydrate binding modules (CBMs) (Baldrian and Valaskova, 2008; Cantarel, et al., 2009; Lynd, Weimer, et al., 2002; Schuster and Schmol, 2010). Consequently, organisms from many lignocellulose-rich environments and their enzymes are being studied for new insights into overcoming this barrier.

In order to obtain the hydrolysis of crystalline cellulose, enzymes must cleave three types of glycosidic bonds. These enzymes are endocellulases, which cleave beta-1,4 glycosidic bonds that reside within intact cellulose strands in the crystalline face, non-reducing-end exocellulases, which remove cellobiose units from the non-reducing end of cellulose strands, and reducing-end exocellulases, which remove glycosyl units from the reducing-end of a cellulose strand. The endocellulolytic reaction is essential because it creates the non-reducing and reducing ends that serve as the starting point for exocellulolytic reactions. The exocellulolytic reactions are essential because they remove glycosyl groups in a processive manner from the breakages in the cellulose strand introduced by the endocellulases, thus amplifying the single initiating reaction of the endocellulases.

Trichoderma reesei and *Clostridium thermocellum* are well-characterized cellulose-utilizing organisms (Merino and Cherry, 2007; Bayer et al., 2008; Wilson, 2011). *T. reesei* is a slow-growing eukaryote fungus that secretes enzymes containing glycoside hydrolase (GH) domains fused to carbohydrate binding domains, while *C. thermocellum* is a strictly anaerobic prokaryote that predominantly assembles

GHs and carbohydrate-binding molecules (CBMs) into a large complex called the cellulosome. Enzymes from these free-living organisms cleave polysaccharides using general acid-base catalyzed hydrolytic reactions (Vuong and Wilson, 2010). Moreover, fungal and microbial communities associated with termites (Scharf et al., 2011) shipworms (Luyten et al., 2006), and rumen (Hess et al., 2011) contribute these types of hydrolytic enzymes to their respective anaerobic niches.

Some free-living aerobes such as *Cellvibrio japonicus* (Ueda 107) (DeBoy et al., 2008), *Streptomyces* (Schloch-termeier et al., 1992; Wilson, 1992; Forsberg et al., 2011), *Thermoascus aurantiacus* (Langston et al., 2011; Quinlan et al., 2011) and *Serratia marcescens* (Vaaje-Kolstad et al., 2010) also grow on biomass polysaccharides. Recent work with some of these organisms has identified that the structurally related fungal GH61 (Langston et al., 2011; Quinlan et al., 2011) and bacterial CBM33 (Forsberg et al., 2011) families of proteins catalyze a previously unrecognized oxidative breakage of glycosidic bonds. This reaction is thought to be an endo-cleavage, with the oxidation reaction yielding gluconate and keto-sugars instead of the typically observed reducing and non-reducing sugars obtained from hydrolytic cellulases.

Actinobacteria in the genus *Streptomyces* are an ecologically important group, especially in soil environments, where they are considered to be vital players in the decomposition of cellulose and other biomass polymers (Cantarel et al., 2009; Crawford et al., 1978; Goodfellow and Williams, 1983; McCarthy and Williams, 1992). *Streptomyces* are able to utilize a wide range of carbon sources, form spores when resources are depleted, and produce antimicrobial secondary metabolites to reduce competition (Goodfellow and Williams, 1983; Schlatter et al., 2009).

Although a large number of *Streptomyces* species can grow on biomass, only a small percentage (14%) have been shown to efficiently degrade crystalline cellulose (Wachinger, Bronnenmeier, et al., 1989). Furthermore, the secreted cellulolytic activities of only a few species have been biochemically characterized, and still fewer species have been examined to identify key biomass degrading enzymes (Ishaque and Kluepfel, 1980; Semedo et al., 2004). *Streptomyces reticuli* is one of the best-studied cellulose- and chitin-degrading soil-dwelling *Streptomyces*; functional analyses of several important cellulases and other hydrolytic enzymes have been reported (Wachinger, Bronnenmeier, et al., 1989; Schlochtermeier, Walter, et al., 1992; Walter and Schrepf, 1996).

Furthermore, polysaccharide monooxygenase (PMO) activity with cellulose was identified using the CBM33 protein from *Streptomyces coelicolor* (Forsberg, et al., 2011), which suggests *Streptomyces* may use both hydrolytic and oxidative enzymes to deconstruct biomass. With the tremendous amount of sequence data collected in the past few years, and despite the view that *Streptomyces* make important contributions to cellulose degradation in the soil, genome-wide analyses of cellulolytic *Streptomyces* have not been reported.

In addition to their putative roles in carbon cycling in the soil, *Streptomyces* may also potentiate biomass deconstruction in insects through symbiotic associations (Bignell, Anderson, et al., 1991; Pasti and Belli, 1985; Pasti, Pometto, et al., 1990; Schafer, et al., 1996). Recent work has identified cellulose degrading *Streptomyces* associated with the pine-boring woodwasp *Sirex noctilio*, including *Streptomyces* sp. SirexAA-E (ActE) (Adams, et al., 2011). *S. noctilio* is a highly destructive wood-feeding insect that is found

throughout forests in Eurasia and North Africa and is spreading invasively in North America and elsewhere (Bergeron, et al., 2011). While the wasp itself does not produce cellulolytic enzymes, evidence supports the role of a symbiotic microbial community that secretes biomass-degrading enzymes to facilitate nutrient acquisition for developing larvae in the pine tree (Kukor and Martin, 1983).

The white rot fungus, *Amylostereum areolatum*, is the best-described member of this community, and the success of *Sirex* infestations is thought to arise from the insect's association with this cellulolytic fungal mutualist. However, work with pure cultures has suggested that ActE and other *Sirex*-associated *Streptomyces* are more cellulolytic than *A. areolatum* (Adams, et al., 2011).

Optimal activity in the CBM33 enzymes apparently requires the addition of a transition metal ion such as Cu(II), Fe(III), or Mn(II) and an external reducing agent. In the laboratory, the reducing agent can be provided by ascorbate. In natural systems, the reducing function is most likely provided by another redox active protein such as cellobiose dehydrogenase (Langston et al., 2011; Quinlan et al., 2011) or some other presently unknown protein.

Needed in the art are improved compositions and organisms for digestion of lignocellulosic materials.

BRIEF SUMMARY

The invention relates generally to methods and compositions for digesting lignocellulosic material and more particularly to methods that involve exposing the material to secretome derived from *Streptomyces* sp. ActE.

In a first aspect, the present invention is summarized as a method of digesting a lignocellulosic material comprising the step of exposing the material to an effective amount of *Streptomyces* sp. ActE secretome preparation such that at least partial lignocellulosic digestion occurs.

In some embodiments of the first aspect, the preparation is a supernatant preparation obtained from a *Streptomyces* sp. ActE culture. In some embodiments of the first aspect, the preparation is obtained from *Streptomyces* sp. ActE grown on a substrate wherein at least 40%, preferably 85%, of *Streptomyces* sp. ActE's carbon source in the substrate is derived from a material selected from the group consisting of cellulose, cellulose/hemicelluloses mixture, hemicelluloses, xylan, non-wood biomass, wood biomass and chitin. In some embodiments of the first aspect, the lignocellulosic material is selected from the group consisting of materials that comprise at least 75% cellulose, cellulose/hemicelluloses, xylose, biomass and chitin.

In a second aspect, the present invention is summarized as a purified preparation comprising the *Streptomyces* sp. ActE secretome.

In some embodiments of the second aspect, the preparation is a supernatant preparation obtained from a *Streptomyces* sp. ActE culture. In some embodiments of the second aspect, *Streptomyces* sp. ActE is grown on a substrate wherein at least 40%, preferably 85%, of *Streptomyces* sp. ActE's carbon source in the substrate is derived from a material selected from the group consisting of cellulose, cellulose/hemicelluloses mixture, hemicelluloses, xylan, non-wood biomass, wood biomass and chitin.

In a third aspect, the present invention is summarized as a composition useful for digesting lignocellulosic material comprising SActE_0237 (GH6) (SEQ ID NO:1) gene or expression product thereof.

In a fourth aspect, the present invention is summarized as a composition useful for digesting lignocellulosic material comprising SActE_0236 (GH48) (SEQ ID NO:2) gene or expression product thereof.

In a fifth aspect, the present invention is summarized as a composition useful for digesting lignocellulosic material comprising SActE_3159 (CBM33) (SEQ ID NO:3) gene or expression product thereof.

In a sixth aspect, the present invention is summarized as a composition useful for digesting lignocellulosic material comprising SActE_0482 (GH5) (SEQ ID NO:4) gene or expression product thereof.

In a seventh aspect, the present invention is summarized as a composition useful for digesting lignocellulosic material comprising SActE_0265 (GH10) (SEQ ID NO:5) gene or expression product thereof.

In an eighth aspect, the present invention is summarized as a composition useful for digesting lignocellulosic material comprising SActE_2347 (GH5) (SEQ ID NO:6) gene or expression product thereof.

In a ninth aspect, the present invention is summarized as a composition useful for digesting lignocellulosic material comprising SActE_0237 (GH6) (SEQ ID NO:1), SActE_0236 (GH48) (SEQ ID NO:2), SActE_3159 (CBM33)(SEQ ID NO:3), SActE_0482 (GH5) (SEQ ID NO:4) and gene or expression product thereof.

In some embodiments of the third, fourth, fifth, sixth, seventh, eighth, and ninth aspects, the composition is optimized for cellulose utilization. In these embodiments the composition can additionally comprise at least one member selected from SActE_0265 (GH10) (SEQ ID NO:5) and SActE_2347 (GH5) (SEQ ID NO:6) genes or expression products thereof. In a preferred embodiment, the composition comprises at least three or four of the genes or expression products.

In some embodiments of the third, fourth, fifth, sixth, seventh, eighth, and ninth aspects, the composition is optimized for xylan release. By "release," we mean degradation, such as hydrolysis, and release of an important or desired product. In these embodiments the composition can additionally comprise at least one member selected from SActE_0265 (GH10) (SEQ ID NO:5), SActE_0358 (GH11) (SEQ ID NO:8), SActE_0357 (CE4) (SEQ ID NO:7), SActE_5978 (PL1)(SEQ ID NO:16) and SActE_5230 (xylose isomerase) genes or expression products thereof. In a preferred embodiment, the composition comprises at least three or four of the genes or expression products.

In some embodiments of the third, fourth, fifth, sixth, seventh, eighth, and ninth aspects, the composition is optimized for chitin release. In these embodiments the composition can additionally comprise at least one member selected from SActE_4571 (GH18), SActE_2313 (CBM33), SActE_4246 (GH18), SActE_3064 (GH19) and SActE_5764 (GH18) genes or expression products thereof. In a preferred embodiment, the composition comprises at least three or four of the genes or expression products.

In some embodiments of the third, fourth, fifth, sixth, seventh, eighth, and ninth aspects, the composition is optimized for biomass degradation. In these embodiments the composition can additionally comprise SActE_5457 (GH46) (SEQ ID NO:14) gene or expression products thereof.

In some embodiments of the third, fourth, fifth, sixth, seventh, eighth, and ninth aspects, the composition is optimized for mannan release. In these embodiments the composition can additionally comprise SActE_2347 (GH5) (SEQ ID NO:6) gene or expression products thereof.

In some embodiments of the third, fourth, fifth, sixth, seventh, eighth, and ninth aspects, the composition is optimized for beta-1,3-glucan release. In these embodiments the composition can additionally comprise at least one member selected from SActE_4755 (GH64) (SEQ ID NO:13) and SActE_4738 (GH16) (SEQ ID NO:12) genes or expression products thereof. In a preferred embodiment, the composition comprises both of the genes or expression products.

In some embodiments of the third, fourth, fifth, sixth, seventh, eighth, and ninth aspects, the composition is optimized for pectin cleavage. In these embodiments the composition can additionally comprise SActE_1310 (PL3) (SEQ ID NO:9) gene or expression products derived thereof.

In some embodiments of the third, fourth, fifth, sixth, seventh, eighth, and ninth aspects, the composition is optimized for alginate release. In these embodiments the composition can additionally comprise SActE_4638 (SEQ ID NO:11) gene or expression products derived thereof.

In some embodiments of the third, fourth, fifth, sixth, seventh, eighth, and ninth aspects, the composition is optimized for galactose release. In these embodiments the composition can additionally comprise SActE_5647 (GH87) (SEQ ID NO:15) gene or expression products derived thereof.

In a tenth aspect, the present invention is summarized as a composition useful for xylan degradation comprising SActE_0265 (GH10) (SEQ ID NO:5) and SActE_0358 (GH11) (SEQ ID NO:8) gene or expression products thereof.

In some embodiments of the tenth aspect, the composition additionally comprises SActE_0265 (GH10) (SEQ ID NO:5), SActE_0358 (GH11) (SEQ ID NO:8), SActE_0357 (CE4) (SEQ ID NO:7), SActE_5978 (PL1) (SEQ ID NO:16), and SActE_5230 (xylose isomerase) genes or expression products thereof. In a preferred embodiment, the composition comprises at least three or four of the genes or expression products.

In an eleventh aspect, the present invention is summarized as a composition useful for biomass degradation comprising SActE_0237 (GH6) (SEQ ID NO:1), SActE_0482 (GH5) (SEQ ID NO:4), SActE_3159 (CBM33)(SEQ ID NO:3), SActE_0236 (GH48) (SEQ ID NO:2), SActE_3717 (GH9) (SEQ ID NO:10), SActE_0265 (GH10) (SEQ ID NO:5), SActE_0358 (GH11) (SEQ ID NO:8), SActE_2347 (GH5) (SEQ ID NOs:6) and SActE_1310 (PL1) genes or expression products thereof. In a preferred embodiment, the composition comprises at least three or four of the genes or expression products.

In a twelfth aspect, the present invention is summarized as a composition useful for cellulose degradation comprising SActE_0237 (GH6) (SEQ ID NO:1), SActE_0482 (GH5) (SEQ ID NO:4), SActE_3159 (CBM33) (SEQ ID NO:3) SActE_0236 (GH48) (SEQ ID NO:2), SActE_2347 (GH5) (SEQ ID NO:6), and SActE_0265 (GH10) (SEQ ID NO:5) genes or expression products thereof. In a preferred embodiment, the composition comprises at least three or four of the genes or expression products.

In a thirteenth aspect, the present invention is summarized as a method for digesting a lignocellulosic material, comprising exposing the material to a sufficient amount of a composition of any one of the third to eighth aspects of the invention, wherein the exposed material is at least partially digested.

In a fourteenth aspect, the present invention is summarized as a purified preparation of *Streptomyces* sp. ActE, wherein the *Streptomyces* sp. ActE has been grown on a substrate wherein at least 40%, preferably 85%, of *Strepto-*

myces sp. ActE's carbon source in the substrate is derived from a material selected from the group consisting of cellulose, cellulose/hemicelluloses mixture, hemicelluloses, xylan, non-wood biomass, wood biomass, and chitin.

In a fifteenth aspect, the present invention is summarized as a purified preparation of *Streptomyces* sp. ActE, wherein the *Streptomyces* sp. ActE has been grown on a substrate wherein at least 40%, preferably 85%, of *Streptomyces* sp. ActE's carbon in the substrate is derived from pretreated lignocellulosic material.

In some embodiments of the fifteenth aspect, the pretreated material has been exposed to pretreatment selected from the group consisting of acid hydrolysis, steam explosion, ammonia fiber expansion (AFEX), organosolve, sulfite pretreatment to overcome recalcitrance of lignocellulose (SPORL), ionic liquids, metal-catalyzed hydrogen peroxide, alkaline wet oxidation and ozone pretreatment. In some embodiments of the fifteenth aspect, the pretreated material is wood.

These and other features, objects, and advantages of the present invention will become better understood from the description that follows. In the description, reference is made to the accompanying drawings, which form a part hereof and in which there is shown by way of illustration, not limitation, embodiments of the invention. The description of preferred embodiments is not intended to limit the invention to cover all modifications, equivalents and alternatives. Reference should therefore be made to the claims recited herein for interpreting the scope of the invention.

BRIEF DESCRIPTION OF THE DRAWINGS

The patent or application file contains at least one drawing executed in color. Copies of this patent or patent application publication with color drawing(s) will be provided by the Office upon request and payment of the necessary fee.

The present invention will be better understood and features, aspects and advantages other than those set forth above will become apparent when consideration is given to the following detailed description thereof. Such detailed description makes reference to the following drawings, wherein:

FIG. 1 is a set of pictures showing growth of ActE in minimal medium containing filter paper as the sole carbon source. (A) Growth of ActE, *Streptomyces coelicolor*, and *Streptomyces griseus* in minimal medium for 7 days at 30° C. and pH 6.9. The expanded image shows small colonies of *S. coelicolor* and *S. griseus* forming on the surface of the paper. (B) Growth of ActE and *Trichoderma reesei* Rut-C30 for 7 days at 30° C. and pH 6.0.

FIG. 2 is a set of graphs demonstrating reactions of ActE secretomes and Spezyme CP. (A) HPLC of sugars released from cellulose (1, cellobiose; 2, cellobiose; 3, glucose) and quantification of glucose equivalent (insert). (B) Reducing sugars released from xylan and mannan by the secretomes of ActE grown on cellulose and xylan. (C) Total reducing sugar released from ionic liquid-switchgrass (IL-SG) or AFEX-switchgrass (AFEX-SG) in reactions of the ActE cellulose, AFEX-SG, and IL-SG secretomes and Spezyme CP. Data represent the mean±s.d. from three experiments; * indicates P<0.01 compared with Spezyme CP.

FIG. 3 is a table illustrating composition of ActE secretomes identified by LC-MS/MS. (A) CAZy genes account for 2.6% of the 6357 predicted protein-coding sequences in the ActE genome. (B) Identity of most abundant proteins in the cellulose secretome proteins is sorted according to

decreasing spectral counts (accounting for 95% of total spectral counts); corresponding spectral counts from other secretomes are also shown.

FIG. 4 is a systematic diagram showing genome-wide changes in expression during growth of ActE on AFEX-treated switchgrass (AFEX-SG) versus glucose. Nodes are genes (circles) or KEGG/CAZy functional categories (yellow triangles); edges indicate that the gene belongs to the indicated functional group as defined by either KEGG or CAZy analysis. Gene node sizes reflect expression intensity determined by microarray from growth on AFEX-SG as a \log_2 ratio, where the genome-wide average transcriptional intensity was ~ 10.5 for both substrates. Node colors represent expression changes as the \log_2 ratio of AFEX-SG/glucose transcript intensities.

FIG. 5 is a diagram with a table showing expression of ActE CAZy genes on various carbon sources. (A) Hierarchical clustering of expression for 167 CAZy genes from the ActE genome during growth on the indicated substrates. (B) Identity of CAZy genes with distinct changes in expression observed in group 1 CAZy genes during growth in different carbon sources.

FIG. 6 is a set of scanning electron microscopy (SEM) images showing ActE grown on different carbon sources including glucose, cellulose, xylan, switchgrass, ammonia fiber expansion-treated switchgrass (AFEX-SG) and ionic liquid-treated switchgrass (IL-SG). ActE cells were grown in minimum medium with the indicated substrate as a sole carbon source for 7 days at 30° C. The scale bar indicates 5 μm .

FIG. 7 is a set of graphs demonstrating fractionation of the ActE cellulose secretome and assays of reactions with different polysaccharides. (A) Anion exchange chromatography was performed using the ActE cellulose secretome, and fractions were collected and analyzed by SDS-PAGE. Lowercase letters indicate protein identified by MALDI-TOF MS shown in FIG. 17. (B) Results from hydrolysis assays for reaction with filter paper (FP), xylan, mannan and beta-1,3 glucan as detected by DNS assay of each fraction. The percentage reactivity relative to the maximum activity observed for each substrate is shown. Error bars indicate the standard deviation, with $n=3$ for technical replicates.

FIG. 8 is a set of diagrams showing temperature and pH profiles of the ActE secretome obtained from growth on AFEX-treated corn stover. (A) The effect of temperature on the deconstruction of AFEX-treated switchgrass (AFEX-SG) and ionic liquid-treated switchgrass (IL-SG). The relative activity of the ActE secretome was compared to the maximal rates determined for reaction with AFEX-SG (blue star), and IL-SG (red star) at pH 6.0. (B) The effect of pH on the AFEX-SG and IL-SG deconstruction activities in the indicated ActE secretomes. The maximal rates observed for AFEX-SG and IL-SG were at pH 7.0 (blue star) and pH 8 (red star), respectively. Reactions were carried out at 40° C. and the 0.1 M buffers used were citrate (pH 4.5), phosphate (pH 6-8), CHES (pH 9-10), and CAPS (pH 11). The reaction was performed for 20 h and the reducing sugar content was measured by DNS assay.

FIG. 9 is a systematic diagram showing genome-wide changes in expression during growth of ActE on substrate cellobiose versus glucose visualized as a Cytoscape interaction network. Nodes are genes (circles) or KEGG/CAZy functional categories (yellow triangles); edges indicate that the gene belongs to the indicated functional group as defined by either KEGG or CAZy analysis. Gene node sizes reflect expression intensity determined by microarray from growth on substrate as a \log_2 ratio. Node colors represent expres-

sion changes as the \log_2 ratio of substrate/glucose transcript intensities, where the genome-wide average transcriptional intensity was ~ 10.5 for both substrate and glucose. Transcripts with less than two-fold changes in expression intensity are colored white; transcripts with greater than two-fold increase in expression intensity during growth on substrate are shown as a red gradient; transcripts with greater than two-fold increase in expression intensity during growth on glucose are shown as a blue gradient.

FIG. 10 is a systematic diagram showing genome-wide expression changes for growth on the substrate cellulose versus glucose visualized as a Cytoscape interaction network. Other information is the same as that described in FIG. 9.

FIG. 11 is a systematic diagram showing genome-wide expression changes for growth on the substrate xylan versus glucose visualized as a Cytoscape interaction network. Other information is the same as that described in FIG. 9.

FIG. 12 is a systematic diagram showing genome-wide expression changes for growth on the substrate switchgrass versus glucose visualized as a Cytoscape interaction network. Other information is the same as that described in FIG. 9.

FIG. 13 is a systematic diagram showing genome-wide expression changes for growth on the substrate IL-treated switchgrass versus glucose visualized as a Cytoscape interaction network. Other information is the same as that described in FIG. 9.

FIG. 14 is a systematic diagram showing genome-wide expression changes for growth on the substrate chitin versus glucose visualized as a Cytoscape interaction network. Other information is the same as that described in FIG. 9.

FIG. 15 is a diagram with a table showing expression of 167 predicted CAZy genes in ActE, highlighting group 2 genes. These genes showed no signal above the average genomic expression intensity ($\log_2=10.5$). (A) Clustering of genes with similar expression profiles. (B) Additional information on group 2 genes including expression profile, SACTE_locus ID, CAZy family, and annotated function.

FIG. 16 is a diagram with a table showing expression of 167 predicted CAZy genes in ActE, highlighting group 3 genes. (A) Clustering of genes with similar expression profiles. (B) Additional information on group 3 genes including expression profile, SACTE_locus ID, CAZy family, and annotated function.

FIG. 17 is a table illustrating proteins separated by ion exchange chromatography and identified by mass spectrometry.

FIG. 18 is a table showing spectra count of proteins identified on each substrate, where top 95% spectra covered were highlighted green, light purple, purple, blue, orange, pink, light blue and yellow on glucose, cellobiose, cellulose, xylan, switchgrass, AFEX-SG, IL-SG and chitin, respectively.

FIG. 19 shows the nucleic acid sequences of the ActE genes.

FIG. 20 shows the amino acid sequences of the ActE genes.

FIG. 21 is a graph illustrating a comparison of specific activities of *Streptomyces* sp. ActE secretomes with Spezyme CP. FIG. 21A depicts relative specific activity of ActE secretomes prepared from growth on cellulose or xylan and Spezyme CP (100%) for reducing sugar release from xylan or mannan. FIG. 21B depicts relative activity (pH 6.0, 40° C.) of ActE cellulose secretome and Cellcc_CBM3a, an

engineered *C. thermocellum* endo/exoglucanase, compared to Spezyme CP. Total amounts of protein included in all reactions were equivalent.

FIG. 22 illustrates nucleotide (SEQ ID NO:63) and amino acid (SEQ ID NO:64) sequence of CeLcc_CBM3a. Construct described in US Patent Application Publication No.: US2010/037094 (Fox and Elsen).

FIG. 23 is a graph illustrating SDS-PAGE of *Streptomyces* sp. ActE secretomes obtained from growth on minimal medium containing different substrates (SG, switchgrass; CS, corn stover; UBLPKP, unbleached lodgepole pine kraft pulp; BSKP, bleached spruce kraft pulp; LP-SPORL, lodgepole pine pretreated by sulfite pretreatment to overcome recalcitrance of lignocellulose (SPORL)). Culture secretomes were separated after 7 days of growth at 30° C. by centrifugation and concentrated by ultrafiltration. Sample loading was normalized to total protein. The identities of proteins were determined from samples extracted from the SDS-PAGE gel. Among the 162 proteins accounting for 95% of spectral counts from the glucose secretome, most were intracellular proteins originating from cell lysis during growth, and were not detected in the polysaccharide secretomes.

FIG. 24 is a graph illustrating SDS-PAGE of time-dependent changes in the *Streptomyces* sp. ActE secretome obtained from growth on minimal medium containing cellulose. Culture secretomes were collected after 7 days by centrifugation and concentrated by ultrafiltration. The concentrated secretomes were incubated at 25° C. for the indicated times and analyzed. Protein bands with time-dependent decrease in intensity were excised from the gel and identified by LC-MS/MS.

FIG. 25 illustrates synergy of recombinant fractions from ion exchange chromatography. All reactions were prepared to contain the same total amount of protein.

FIG. 26 is a set of graphs illustrating mannanase activity demonstrated in fractions containing various naturally truncated versions of SACTE_2347 (GH5). FIGS. 26A-B depict proteins found in previous assayed fractions. FIG. 26C depicts Coomassie Blue staining of 12% polyacrylamide gel (PAGE) separation of different mannanase isoforms. Three polypeptide bands corresponding to SACTE_2347 (GH5) with molecular masses of ~57, ~45, and ~37 kDa. FIG. 26D depicts a zymogram performed in the presence of 0.5% mannan. The strong clearing zone in fraction F1 associated with the ~37 kDa isoform demonstrates how size reduction can increase the specific activity of a protein.

FIG. 27 is a set of graphs illustrating ion exchange fractionation of *Streptomyces* sp. ActE secretome. FIG. 27A depicts an SDS-PAGE analysis of the fractionation of an ActE secretome by ion exchange chromatography. FIG. 27B depicts catalytic assays of the separate fractions at 40° C. for 20 h in 0.1 M phosphate buffer, pH 6.0, showing different enzymes are capable of reacting with xylan, mannan, and cellulose. The reactivity of fractions marked with stars is also described in FIG. 25A.

FIG. 28 is a SDS-PAGE graph and a list illustrating mass spectral assignment of polypeptides from the *Streptomyces* sp. ActE secretome separated by ion exchange chromatography. FIG. 28A depicts an SDS PAGE of separated fractions annotated with identities of polypeptides determined by LC-MS analysis. FIG. 28B depicts information on the identified proteins including gene locus, function, CAZY GH and CBM assignments, number of amino acid (AA) residues, and best BLAST result for relationship to another known enzyme. The reactivity of fractions marked with stars is also described in FIG. 25A.

FIG. 29 is a SDS-PAGE graph and a table that demonstrates the existence of xylanases from *Streptomyces* sp. ActE. Five ActE proteins were produced using cell-free translation as described in US Patent Application Publication No.: US2010/037094 (Fox and Elsen). FIG. 29A depicts a stain-free gel image of proteins produced by wheat germ cell-free translation (indicated by asterisks). FIG. 29B depicts a summary of protein information, expression and secretion data, and diagnostic assay results. Small molecule assays (MUG, methylumbelliferyl glucoside; MUC, methylumbelliferyl cellobioside; MUM, methylumbelliferyl mannoside and MUX2, methylumbelliferyl xylobioside) were performed in 0.1 M phosphate buffer, pH 6.0, at 30° C. SACTE_0265 and SACTE_0358, highly expressed and secreted proteins during growth on xylan, are confirmed by these assays to be xylanases. Results from three other non-secreted ActE enzymes are provided as controls.

FIG. 30 is a graph illustrating quantification of total secreted protein obtained from *Streptomyces* sp. ActE grown on different substrates (AFEX-CS, AFEX corn stover; UBLPKP, unbleached lodgepole pine kraft pulp; BSKP, bleached spruce kraft pulp; LP-SPORL, lodgepole pine pretreated by SPORL).

FIG. 31 is a graph illustrating the temperature versus activity profile of the *Streptomyces* sp. ActE secretome obtained from growth on cellulose. Hydrolysis activities were measured by DNS assay. Greater than 80% of maximal rates for cellulase and mannanase activity were observed at the range of 31-43° C., while greater than 80% of maximal rate for xylanase activity was observed in the range of 35-59° C.

FIG. 32 is a graph illustrating the pH versus activity profile of the *Streptomyces* sp. ActE secretome obtained from growth on cellulose. The maximal rate was observed at approximately pH 6. Buffers used in this study were 0.1 M citrate (pH 4.5), phosphate (pH 6-8), CHES (pH 9-10) and CAPS (pH 11).

FIG. 33 is a SDS-PAGE graph illustrating ActE induction in medium containing as little as 20% cellulose.

FIG. 34 is a set of Venn diagrams representing 95% of total proteins identified in LC-MS/MS analyses generated using VennMaster-0.37.5 (Kestler et al., 2008). FIG. 8A depicts secretomes obtained from growth on glucose, Sigmacell™, and xylan. FIG. 8B depicts secretomes obtained from growth on switchgrass, ammonia fiber expansion (AFEX)-SG, and IL-SG. For clarification, glucose ∩ Sigmacell=4 represents the intersection of the two sets, while glucose/(Sigmacell ∪ xylan)=117 represents the proteins uniquely associated with growth on glucose as compared to Sigmacell. Other results are interpreted in a similar manner.

While the present invention is susceptible to various modifications and alternative forms, exemplary embodiments thereof are shown by way of example in the drawings and are herein described in detail. It should be understood, however, that the description of exemplary embodiments is not intended to limit the invention to the particular forms disclosed, but on the contrary, the intention is to cover all modifications, equivalents and alternatives falling within the spirit and scope of the invention as defined by the appended claims.

DESCRIPTION OF EXEMPLARY EMBODIMENTS

In General

The present invention comprises many embodiments. In one embodiment, the invention is a method of digesting a

lignocellulosic material, comprising the step of exposing the material to an effective amount of *Streptomyces* sp. ActE secretome preparation such that at least partial lignocellulosic digestion occurs. In one embodiment of that method, the preparation is a supernatant preparation obtained from a *Streptomyces* sp. ActE culture. In another embodiment of that method, the preparation is obtained from *Streptomyces* sp. ActE grown on a substrate wherein at least 40%, preferably 85%, of *Streptomyces* sp. ActE's carbon source in the substrate is derived from a material selected from the group consisting of cellulose, cellulose/hemicelluloses mixture, hemicelluloses, xylan, non-wood biomass, wood biomass, and chitin. In another embodiment of that method, the lignocellulosic material is selected from the group consisting of materials that comprise at least 75% cellulose, cellulose/hemicelluloses, xylose, biomass and chitin.

In one embodiment, the invention is a purified preparation comprising the *Streptomyces* sp. ActE secretome. In one embodiment, the preparation is a supernatant preparation obtained from a *Streptomyces* sp. ActE culture. In another embodiment of the preparation, *Streptomyces* sp. ActE is grown on a substrate wherein at least 40%, preferably 85%, of *Streptomyces* sp. ActE's carbon source in the substrate is derived from a material selected from the group consisting of cellulose, cellulose/hemicelluloses mixture, hemicelluloses, xylan, non-wood biomass, wood biomass, and chitin.

In one embodiment, the invention is a composition useful for digesting lignocellulosic material comprising one gene or expression product thereof selected from the group consisting of SActE_0237 (GH6) (SEQ ID NO:1), SActE_0236 (GH48) (SEQ ID NO:2), SActE_3159 (CBM33) (SEQ ID NO:3), SActE_0482 (GH5) (SEQ ID NO:4), SActE_0265 (GH10) (SEQ ID NO:5), and SActE_2347 (GH5) (SEQ ID NO:6) genes or expression products thereof. In one embodiment, the composition additionally comprises at least one member selected from the group consisting of SActE_0357 (CE4) (SEQ ID NO:7), SActE_0358 (GH11) (SEQ ID NO:8), SActE_1310 (PL3) (SEQ ID NO:9), SActE_3717 (GH9) (SEQ ID NO:10), SActE_4638 (SEQ ID NO:11), SActE_4738 (GH16) (SEQ ID NO:12), SActE_4755 (GH64) (SEQ ID NO:13), SActE_5457 (GH46) (SEQ ID NO:14), SActE_5647 (GH87) (SEQ ID NO:15), and SActE_5978 (PL1) (SEQ ID NO:16) genes or expression products derived thereof.

In one embodiment, the invention is a composition useful for cellulose degradation comprising SActE_0236 (GH48) (SEQ ID NO:2), SActE_3159 (CBM33) (SEQ ID NO:3), SActE_0482 (GH5) (SEQ ID NO:4) and SActE_0237 (GH6) (SEQ ID NO:1) genes or expression product thereof. In one embodiment, the composition additionally comprises at least one member selected from the group consisting of SActE_0357 (CE4) (SEQ ID NO:7), SActE_0358 (GH11) (SEQ ID NO:8), SActE_1310 (PL3) (SEQ ID NO:9), SActE_3717 (GH9) (SEQ ID NO:10), SActE_4638 (SEQ ID NO:11), SActE_4738 (GH16) (SEQ ID NO:12), SActE_4755 (GH64) (SEQ ID NO:13), SActE_5457 (GH46) (SEQ ID NO:14), SActE_5647 (GH87) (SEQ ID NO:15), and SActE_5978 (PL1) (SEQ ID NO:16) genes or expression products derived thereof.

In one embodiment, the invention is a method for digesting a lignocellulosic material, comprising exposing the material to a sufficient amount of a composition of any combinations of genes or expression products derived thereof as disclosed above, wherein the exposed material is at least partially digested.

In one embodiment, the invention is a purified preparation of *Streptomyces* sp. ActE, wherein the *Streptomyces* sp. ActE

has been grown on a substrate wherein at least 40%, preferably 85%, of *Streptomyces* sp. ActE's carbon source in the substrate is derived from a material selected from the group consisting of cellulose, cellulose/hemicelluloses mixture, hemicelluloses, xylan, non-wood biomass, wood biomass and chitin.

In one embodiment, the invention is a purified preparation of *Streptomyces* sp. ActE, wherein the *Streptomyces* sp. ActE has been grown on a substrate wherein at least 40%, preferably 85%, of *Streptomyces* sp. ActE's carbon in the substrate is derived from pretreated lignocellulosic material. In one embodiment of the preparation, the pretreated material has been exposed to pretreatment selected from the group consisting of acid hydrolysis, steam explosion, ammonia fiber expansion (AFEX), organosolve, sulfite pretreatment to overcome recalcitrance of lignocellulose (SPORL), ionic liquids (IL), metal-catalyzed hydrogen peroxide treatment, alkaline wet oxidation and ozone pretreatment. In another embodiment of the preparation, the pretreated material is wood.

Specific Embodiments

Applicants have been interested in insects that utilize plant biomass and their associated microbial and fungal communities. *Sirex noctilio*, a wood boring wasp, is found in pine forests throughout Eurasia and North Africa and is spreading throughout North America and elsewhere (Bergeron et al., 2011). Although the destructive nature of the *Sirex* infestation is generally considered to arise from a symbiotic relationship between *S. noctilio* and *Amylostereum areolatum*, a white rot basidiomycete (Kukor and Martin, 1983; Klepzig et al., 2009; Bergeron et al., 2011), the role of cellulolytic microbes has not been previously considered in the context of the infestation or symbiosis. *Streptomyces* sp. SirexAA-E [*Streptomyces* sp. ActE, also referred to herein as "ActE" (Adams et al., ISME J. 5:1321-1231, 2011)], was isolated from the ovipositor mycangium of *S. noctilio* (Adams et al., 2011). Applicants hypothesized that ActE is inoculated into insect feeding tunnels upon infestation along with the symbiotic fungus. Thus, Applicants were interested to learn how ActE might contribute to the *Sirex* community.

The present invention will be more fully understood upon consideration of the following non-limiting Examples. All papers and patents disclosed herein are hereby incorporated by reference as if set forth in their entirety.

As used herein, the term "ActE" refers to *Streptomyces* sp. SirexAA-E, as described in Adams et al., ISME J. 5:1321-1231, 2011. A representative sample of *Streptomyces* sp. ActE has been deposited according to the Budapest Treaty for the purpose of enabling the present invention. The repository selected for receiving the deposit is the American Type Culture Collection (ATCC) having an address at 10801 University Boulevard, Manassas, Va. USA, Zip Code 20110. The ATCC repository has assigned the patent deposit designation PTA-12245 to the *Streptomyces* sp. ActE strain.

As used herein, the term "secretome" refers to the plurality of secreted enzymes. For example, ActE secretome refers to the secreted enzymes from *Streptomyces* sp. SirexAA-E.

As used herein, the term "lignocellulosic material" refers to any material that is composed of cellulose, hemicellulose, and lignin, wherein the carbohydrate polymers (cellulose and hemicelluloses) are tightly bound to the lignin.

As used herein, the term "biomass" refers to a renewable energy source, is biological material from living or recently

living organisms. As an energy source, biomass can either be used directly, or converted into other energy products such as biofuel. Biomass includes plant or animal matter that can be converted into fibers or other industrial chemicals, including biofuels. Industrial biomass can be grown from numerous types of plants, including miscanthus, switchgrass, hemp, corn, poplar, willow, sorghum, sugarcane, bamboo, and a variety of tree species, ranging from eucalyptus to oil palm (palm oil). Thus, biomass can include wood biomass and non-wood biomass.

The present invention has multiple embodiments. All embodiments are related to Applicants' discovery of improved lignocellulosic digestion and utilization using proteins and genes obtained from the *Streptomyces* sp. ActE secretome.

ActE Isolates and Secretomes

Streptomyces sp. SirexAA-E may be isolated from ovipositor mycangia of *S. noctilio*. In Adams, et al, *S. noctilio* were collected from a population in Pennsylvania, USA. Infested trees were cut and transported to USDA Pest Survey, Detection, and Exclusion Lab in Syracuse, N.Y., USA (Zylstra et al. (2010) Agric. Forest. Entomol. in press). Four adult females and six larvae from the Pennsylvania population were sampled, and cultures of bacteria derived from these insect samples were screened for cellulose degradation.

Prior to sampling for bacteria, all insects were typically surface sterilized in 95% ethanol for 1 minute and then rinsed twice in sterile phosphate-buffered solution (1×PBS). Larval guts and adult ovipositors and mycangia were removed surgically. These segments and the body were ground separately in 1 ml 1×PBS using a sterilized mortar and pestle. 50 µl of three 100-fold dilutions of each insect part were plated onto yeast and malt extract agar (Becton, Dickinson and Company, Sparks, Md., USA), acidified yeast malt extract agar (for gut dissections only), 10% tryptic soy agar (Becton, Dickinson and Company, Sparks, Md., USA), and agar supplemented with chitin (MP Biomedicals, Solon, Ohio). Petri dishes were stored at room temperature in darkness for at least three days until visible colonies formed, except for Petri dishes with chitin agar that were stored for at least one month.

All isolates were typically screened for production of cellulolytic enzymes on carboxymethyl cellulose (CMC) (Teather R M, Wood P J (1982); incorporated herein by reference as if set forth in its entirety). Isolates that tested positive on CMC were then studied further. Assays on CMC, AFEX-treated corn stover at three pH levels, and microcrystalline cellulose were typically performed to assess growth and degradation ability of each insect-derived bacterial isolate. Isolates capable of degrading CMC were further analyzed genomically to identify isolates with high CAZy content relative to one another and relative to known organisms. *Streptomyces* sp. ActE was selected based on its CMC degradation and CAZy gene profile.

In one embodiment, secretomes from ActE would be used alone in a first reaction to convert biomass into a hydrolyzed solution of sugars that would be used in a second reaction with a fermentation organism to convert the sugars into usable biofuels. The first and second reaction could occur simultaneously.

In a second embodiment, secretomes from ActE would be combined with secretomes from other organisms, or with enzymes or enzyme compositions, such as Spezyme CP, to increase the activity of both preparations by synergy of the enzymes contained in each preparation.

Preferably, the ActE secretomes would be prepared as supernatants from ActE preparations.

In one embodiment, the supernatant is prepared by centrifugation of the ActE culture for 10 min at 3,000×g, which will pellet the remaining insoluble polysaccharide and adhered ActE cells. The supernatant fraction is filter-sterilized, preferably using a 0.22 µm filter, in order to remove any remaining cells. The supernatant is concentrated, preferably using a 3 kDa cut-off ultrafiltration membrane. The concentration of total protein is determined by Bradford assay (Bradford, 1976). In one preferred embodiment, the proteomic composition of the ActE secretome is that described in FIG. 3 or FIG. 18.

The secretomes obtained from growth on specific lignocellulosic materials, such as cellulose, xylan, cellulosic hemi-cellulosic biomass, and chitin, will have distinct compositions of individual enzymes and also distinct reactivity with different polysaccharides. The cellulosic hemi-cellulosic biomass may be non-wood biomass or wood biomass. For example, the secretome prepared from ActE grown on cellulose has unique enzymes and enhanced reactivity with cellulose and mannan. Also, the secretome prepared from ActE grown on xylan possesses high xylan degradation activity, whereas the secretome from ActE grown on chitin possesses uniquely high chitin degradation activity. Example A discloses the specific secretomes.

When ActE is grown on switchgrass, AFEX-pretreated switchgrass or ionic liquid pretreated switchgrass, the secretome has a protein composition that partially matches that obtained from growth on either cellulose or xylan. However, switchgrass, AFEX-pretreated switchgrass or ionic liquid pretreated switchgrass elicit the appearance of new proteins in the secretome that enhance the degradative ability of the secretome for the plant biomass materials. Applicants envision that the present invention would also apply to other pretreatment methods comprising acid hydrolysis, steam explosion, organosolve, sulfite pretreatment to overcome recalcitrance of lignocellulose (SPORL), metal-catalyzed hydrogen peroxide treatment, alkaline wet oxidation and ozone pretreatment.

The inventors' preliminary data shows synergistic filter paper degrading activity between the ActE secretome and other cellulases from a different organism. Also, addition of a beta-glucosidase to the secretome helps to break down the oligosaccharides (e.g., cellotetraose, cellobiose and cellobiose) released from filter paper into simpler sugars.

Preferably, the secretome would be prepared as a concentrated solution by ultrafiltration. The concentrated material would be mixed with the substrate at weight percentages varying from 0.1% to 20% w/w, with the remainder of the solution containing a buffer substance that controls pH. Trace metals would be added to the reaction. The material would be incubated at the appropriate temperature to allow the reaction to occur, with mixing of the reaction materials. The sample might be equilibrated with air or O₂ gas throughout the reaction time period.

The secretome obtained from growth of ActE on cellulose provides all necessary enzymes for most efficient breakdown of cellulose to cellobiose and mannan to mannose. Weak reaction is observed for breakdown of xylan to xylose and a mixture of mannobiose and mannose.

The secretome obtained from growth of ActE on xylan provides all necessary enzymes for most efficient breakdown of xylan to xylobiose and xylose. Weak reaction is observed for breakdown of cellulose to cellobiose and for breakdown of mannan to mannose.

The secretome obtained from growth of ActE on chitin provides all necessary enzymes for most efficient breakdown of chitin to N-acetylglucosamine. Weak reaction is observed for breakdown of xylan to xylose. Weak reaction is observed for breakdown of cellulose to cellobiose and for breakdown of mannan to mannose.

The secretome obtained from growth of ActE on switchgrass biomass provides all of the necessary enzymes for breakdown of cellulose, xylan, and mannan contained in switchgrass to the constituent monosaccharides and disaccharides. Growth of ActE on switchgrass exposed to different chemical pretreatments changes the composition of enzymes present, which alters the rate of production and yield of the constituent monosaccharides and disaccharides.

The secretome obtained from growth of ActE on cellulose provides the necessary enzymes for breakdown of cellulose to cellobiose. ActE uses cellobiose as the growth substrate, so no enzymes are present to convert cellobiose to glucose.

In order to obtain glucose, a cellobiase or beta-glucosidase would be added. This is a standard practice in biofuels enzymology.

In order to convert cellobiose to glucose, a cellobiase or beta-glucosidase would be added. Addition of cellulases from other organisms can improve the rate of hydrolysis of cellulose, e.g., addition of Cellcc_CBM3a, an engineered enzyme from *C. thermocellum* covered in Fox and Elsen Patent Application No.: PCT/US2010/037094.

The secretome obtained from growth of ActE on cellulose provides all of the necessary enzymes for breakdown of cellulose to cellobiose in a soluble form. One skilled in the art might purify these proteins directly from the secretome without use of tags or recombinant approaches.

As previously noted, the dominance of cellobiose as a product of cellulose deconstruction by ActE might help to channel cellulolytic activity to only a subset of the diverse microbes found in the *Sirex* community. Exploiting this community interaction, along with establishing control of the highly regulated patterns of gene expression observed in ActE provides the basis for a new biotechnological route for lignocellulosic digestion. For example, use of ActE secretomes to produce cellobiose will restrict the use of cellulose as a fermentation substrate to only those organisms capable of cellobiose uptake followed by intracellular conversion to glucose and subsequent glycolytic pathway intermediates. This might be achieved by coupling ActE enzymes with a yeast fermentation strain engineered to contain a specific cellobiose transporter and an intracellular cellobiose phosphorylase, leading to the intracellular production of glucose and glucose-1-phosphate.

ActE secretomes can be mixed with cellulosic biomass to convert it to cellobiose and xylose, as in the biofuels industry. For example, one might (1) mix the secretome with paper waste to convert it to a mixture of readily fermentable oligo-, di-, and monosaccharides; (2) mix with animal feeds to increase the digestibility of the biomass to promote animal growth; (3) mix with cotton-based textiles for smoothing or other refinements; (4) mix with waste from the shrimp industry to process solid chitin to soluble constituents; (5) mix with mannan-enriched materials to convert them to mannose and mannanose. One would also find the secretome useful for commercial food processing or treatment of cellulosic bezoar found in the human stomach.

One embodiment of the present invention is an isolation or purified preparation of *Streptomyces* sp. ActE.

An isolation of ActE was originally reported by Adams et al., (2011) ISME j doi:10.1038/ismej.2011.14, where it was stated that "*Sirex noctilio* were collected from infested scots

pine, *Pinus sylvestris* L., in Onondaga County, N.Y., USA in 2008", and "Microbial isolates were obtained from four adult females and six larvae collected in 2008, and were screened for cellulase activity." These isolates were screened for cellulolytic ability by growing them on CMC, AFEX-treated corn stover, and microcrystalline cellulose.

Applicants envision that one would wish to prepare ActE isolates on specific nutrient sources for optimization for particular digestion profiles. Therefore, one may wish to prepare ActE on substrates wherein at least 40%, preferably 85% of *Streptomyces* sp. ActE's carbon source in the substrate is derived from a material selected from the group consisting of cellulose, cellulose/hemicelluloses mixture, hemicelluloses, xylan, non-wood biomass, wood biomass, and chitin.

In a preferred embodiment, ActE would be grown aerobically to maximize the secretion of enzymes that include both oxidative and hydrolytic enzymes capable of the rapid deconstruction of biomass. Since ActE cannot utilize mannose for growth, but efficiently liberates mannose from biomass, mannose would become available for growth of the inoculum of a fermentation organism in co-culture. The likely fact that ActE produces at least one antibiotic that would help maintain culture sterility is another possible advantage to establishment of an effective co-culture.

The high capacity for mannan hydrolysis coupled with the inability of ActE to use mannose as a growth substrate offers unique potential opportunity for expansion of deconstruction enzymology to the use of woody substrates. The deconstruction of woody substrates is considered to be more challenging for biofuels production despite the fact that woody substrates are also considerably more highly enriched in mannan than grass substrates. This unique potential opportunity will be enhanced by ongoing plant engineering research efforts to redefine the proportion of xylan and mannan in plant hemicellulose. The availability of plant material enriched in mannan will be coupled to vigorous conversion to mannose by ActE secretomes, providing a targeted, simply fermented C6 sugar for exclusive use by the fermentation organism.

When sufficient titer of enzymes and fermentation organism have been achieved, facilitated by the vigorous, obligate aerobic growth of ActE and corresponding deconstruction of biomass, the fermentation could be initiated by removal of the air source from the culture vessel. In the anoxic conditions, ActE would cease to grow, and perhaps even lyse to become a protein source for the fermentation organism, which will continue to grow on biomass that is simultaneously being deconstructed by the loading of highly active hydrolytic enzymes originally produced by ActE during the aerobic growth phase.

Applicants envision adding an ActE isolate directly to biomass slurry. More preferably an ActE isolate would be added to the pretreated biomasses in the enzyme hydrolysis step, because ActE is able to grow at wide range of pH. ActE can be genetically modified so that the proteolysis proof secretome will be achieved. Growth on switchgrass elicits the appearance of new proteins in the secretome that enhance the degradative ability of the secretome for the plant biomass materials. Applicants envision that the present invention would apply to the biomasses pretreated by many pretreatment methods comprising AFEX, ionic liquid pretreated, acid hydrolysis, steam explosion, organosolve, sulfite pretreatment to overcome recalcitrance of lignocellulose (SPORL), metal-catalyzed hydrogen peroxide, alkaline wet oxidation and ozone pretreatment.

In one preferred embodiment of the present invention, at least one key enzyme in the secretome can be overexpressed by genetic modification of the ActE strain. Table 1 provides various combinations of genes that can be overexpressed. For example, one may wish to overexpress core cellulose deconstructing enzymes, SACTE_0237, SACTE_0482, SACTE_0236, or SACTE_3159 together with one or more of SACTE_2347, and SACTE_0265. One may wish to overexpress core xylan deconstructing enzymes, SACTE_0265, SACTE_0358, SACTE_0357, SACTE_5978, and SACTE_5230. One may wish to overexpress core mannan deconstruction enzymes, such as SACTE_2347. Additionally, SACTE_4755 and SACTE_4738 may be overexpressed for beta-1,3-glucan deconstruction. One may also overexpress all or some of the aforementioned genes for efficient biomass deconstruction.

In another embodiment of the present invention, at least one key enzyme in the secretome can be overexpressed and secreted by genetic modification of a different microbial host such as *Streptomyces lividans*, which is used for industrial secretion of proteins (Anne and Van Mellaert. (1993)), or *T. reesei*, which is used for secretion of enzymes in the biofuels industry (Saloheimo and Pakula, Microbiology, Epub date 2011 Nov. 5).

In another embodiment of the present invention, at least one key enzyme in the secretome can be overexpressed by genetic modification of a different microbial host such as *S. cerevisiae* or *E. coli* such that the expressed protein will be retained inside of the host cell. The host cells would then be harvested and used as a delivery agent without need for purification of the entrained enzyme, as described in (Wood et al., 1997). This version of the invention may be useful in the enzymatic pretreatment of agricultural crop materials for consumption by ruminant animals.

Combinations of ActE Genes and Expression Products

Selected minimal genes in each subset were chosen based on the combination of genomic, transcriptomic and secretomic results (See Examples and Table 1). For example, in the cellulose minimal gene set, expression of these genes was relatively enriched in cellulose grown cells, compared to glucose grown cells, also corresponding proteins were highly secreted in response to the cellulose in culture medium. Elected minimal genes were annotated to have cellulose utilization function. A larger set of genes for cellulose utilization were selected based on the enrichment of gene expression in cellulose-grown cells relative to glucose-grown cells, and a functional annotation supports cellulose utilization of these genes. Additionally, neighborhood genes to these selected genes on genome were included as genes regulated under same promoter. Similarly, both minimal and a large set of genes for xylan, chitin, and biomasses were elected.

In one embodiment, the present invention is a composition useful for digesting lignocellulosic material comprising genes or expression products thereof selected from the group consisting of: (a) SACTE_0237 (SEQ ID NO:1), SACTE_0236 (SEQ ID NO:2), SACTE_3159 (SEQ ID NO:3), SACTE_0482 (SEQ ID NO:4), SACTE_0265 (SEQ ID NO:5), and SACTE_2347 (SEQ ID NO:6), and (b) SACTE_0357 (CE4) (SEQ ID NO:7), SACTE_0358 (GH11) (SEQ ID NO:8), SACTE_1310 (PL3) (SEQ ID NO:9), SACTE_3717 (GH9) (SEQ ID NO:10), SACTE_4638 (SEQ ID NO:11), SACTE_4738 (GH16) (SEQ ID NO:12), SACTE_4755 (GH64) (SEQ ID NO:13), SACTE_5457 (GH46) (SEQ ID NO:14), SACTE_5647 (GH87) (SEQ ID NO:15), and SACTE_5978 (PL1) (SEQ ID NO:16). In a preferred embodi-

ment, the composition comprises at least three or four of the genes or expression products.

In one embodiment, one would use at least one member of (a) to digest a preferred lignocellulosic material.

In another embodiment, one would use at least the first four members [SACTE_0237 (SEQ ID NO:1), SACTE_0236 (SEQ ID NO:2), SACTE_3159 (SEQ ID NO:3), and SACTE_0482 (SEQ ID NO:4)] of (a) to digest a preferred lignocellulosic material.

In another embodiment, one would use at least one member of (a) and at least one member from (b), to digest a preferred lignocellulosic material.

In a preferred embodiment, one would use all the members of (a) and (b), to digest a preferred lignocellulosic material.

In other embodiments, the combination of genes or expression products thereof in the present invention is dependent on the specific lignocellulosic material to be digested. In one embodiment, a composition optimized for cellulose utilization may include any combinations of ActE genes and expression products disclosed above with at least one member selected from SACTE_0265 (GH10) (SEQ ID NO:5) and SACTE_2347 (GH5) (SEQ ID NO:6) genes or expression products thereof.

In another embodiment, a composition optimized for xylan utilization may include any combinations of ActE genes and expression products disclosed above with at least one member selected from SACTE_0265 (GH10) (SEQ ID NO:5), SACTE_0358 (GH11) (SEQ ID NO:8), SACTE_0357 (CE4) (SEQ ID NO:7), SACTE_5978 (PL1) (SEQ ID NO:16) and SACTE_5230 (xylose isomerase) genes or expression products thereof. In a preferred embodiment, the composition comprises at least three or four of the genes or expression products.

In another embodiment, a composition optimized for chitin utilization may include any combinations of ActE genes and expression products disclosed above with at least one member selected from SACTE_4571 (GH18), SACTE_2313 (CBM33), SACTE_4246 (GH18), SACTE_3064 (GH19), and SACTE_5764 (GH18) genes or expression products thereof. In a preferred embodiment, the composition comprises at least three or four of the genes or expression products.

In another embodiment, a composition optimized for biomass utilization may include any combinations of ActE genes and expression products disclosed above with SACTE_5457 (GH46) (SEQ ID NO:14) genes or expression products thereof.

In another embodiment, a composition optimized for mannan utilization may include any combinations of ActE genes and expression products disclosed above with SACTE_2347 (GH5) (SEQ ID NO:6) genes or expression products thereof.

In another embodiment, a composition optimized for beta-1,3-glucan utilization may include any combinations of ActE genes and expression products disclosed above with at least one member selected from SACTE_4755 (GH64) (SEQ ID NO:13) and SACTE_4738 (GH16) (SEQ ID NO:12) genes or expression products thereof.

In another embodiment, a composition optimized for pectin release utilization may include any combinations of ActE genes and expression products disclosed above with SACTE_1310 (PL3) (SEQ ID NO:9) gene or expression products derived thereof.

In another embodiment, a composition optimized for alginate release utilization may include any combinations of

ActE genes and expression products disclosed above with SActE_4638 (SEQ ID NO:11) gene or expression products derived thereof.

In another embodiment, a composition optimized for galactose release utilization may include any combinations of ActE genes and expression products disclosed above with SActE_5647 (GH87) (SEQ ID NO:15) gene or expression products derived thereof.

In another embodiment, the present invention is summarized as a composition useful for xylan degradation comprising SActE_0265 (GH10) (SEQ ID NO:5) and SActE_0358 (GH11) (SEQ ID NO:8) genes or expression products thereof.

In another embodiment, the present invention is summarized as a composition useful for xylan degradation comprising SActE_0265 (GH10) (SEQ ID NO:5), SActE_0358 (GH11) (SEQ ID NO:8), SActE_0265 (GH10) (SEQ ID NO:5), SActE_0358 (GH11) (SEQ ID NO:8), SActE_0357 (CE4) (SEQ ID NO:7), SActE_5978 (PL1) (SEQ ID NO:16), and SActE_5230 (xylose isomerase) genes or expression products thereof. In a preferred embodiment, the composition comprises at least three or four of the genes or expression products.

In another embodiment, the present invention is summarized as a composition useful for biomass degradation comprising SActE_0237 (GH6) (SEQ ID NO:1), SActE_0482 (GH5) (SEQ ID NO:4), SActE_3159 (CBM33) (SEQ ID NO:3), SActE_0236 (GH48) (SEQ ID NO:2), SActE_3717 (GH9) (SEQ ID NO:10), SActE_0265 (GH10) (SEQ ID NO:5), SActE_0358 (GH11) (SEQ ID NO:8), SActE_2347 (GH5) (SEQ ID NO:6) and SActE_1310 (PL1) genes or expression products thereof. In a preferred embodiment, the composition comprises at least three or four of the genes or expression products.

In one embodiment, the present invention is a composition useful for digesting lignocellulosic material comprising genes or expression products thereof selected from the group consisting of: (a) SActE_0237 (SEQ ID NO:1), SActE_0236 (SEQ ID NO:2), SActE_3159 (SEQ ID NO:3), SActE_0482 (SEQ ID NO:4), SActE_0265 (SEQ ID NO:5), and SActE_2347 (SEQ ID NO:6) (for cellulose); (b) SActE_0265 (SEQ ID NO:5), SActE_0357 (SEQ ID NO:7), SActE_0358 (SEQ ID NO:8), SActE_5230 and SActE_5978 (for xylan); (c) SActE_2313, SActE_3064, SActE_4246, SActE_4571 and SActE_5764 (for chitin); (d) SActE_2347 (for mannan); and (e) SActE_0236 (SEQ ID NO:2), SActE_0237 (SEQ ID NO:1), SActE_0265 (SEQ ID NO:5), SActE_0358, SActE_1310, SActE_2347 and SActE_3159 (for biomass). In a preferred embodiment, the composition comprises at least three or four of the genes or expression products.

In one embodiment, one would use at least two members of (a), (b), (c), (d) or (e) to digest a preferred lignocellulosic material.

In another embodiment, one would use at least three members.

In a preferred embodiment, one would use all members of (a), (b), (c), (d) or (e).

In another embodiment, one would add gene expression products from the list in Table 1 to a substrate to be digested. For example, for preferred cellulose digestion, one would select at least two members of (a), as described above, and at least one member of the "additional useful genes" in Table 1.

In the case of cellulose degradation, the inventors believe SACTE_3159, SACTE_0237, SACTE_0482, and SACTE_0236 act cooperatively to create nicks and hydrolyze cellobiose units from crystalline cellulose.

ActE key genes can be transferred into known cellulolytic organisms in order to enhance the cellulolytic ability of these organisms. A cellulolytic fungus, *T. reesei*, has been studied for industrial applications, and can be genetically modified. Applicants' data support synergism of cellulolytic ability of enzymes from different species. A chromosomal gene transfer can be performed into *T. reesei* by protoplast transformation with a high copy plasmid carrying one or more of the ActE cellulolytic key genes.

A chromosomal or a non-chromosomal gene transfer can be made into a yeast species such as *Saccharomyces cerevisiae*. For non-chromosomal gene transfer, a high copy plasmid carrying a cassette of five minimal genes (SACTE_0236, SACTE_0237, SACTE_0482, SACTE_3717 and SACTE_3159) would be used to confer cellulolytic and mannanolytic capability to the yeast strain. Similar approaches could be used to confer xylanolytic and chitinolytic capability using combinations of the genes described herein.

One might wish to recombinantly express the disclosed enzymes in *E. coli* in order to achieve high yield of each enzyme. As is shown in the synergistic result in Example 18, cellulose degradation can be improved by combination of ActE enzymes to enzymes from other organisms.

FIG. 18 shows Spectra count of proteins identified on each substrate, where top 95% most abundant proteins were highlighted green, light purple, purple, blue, orange, pink, light blue and yellow on glucose, cellobiose, cellulose, xylan, switchgrass, AFEX-SG, IL-SG and chitin, respectively.

Applicants envision that one would use a composition comprising at least one member of the abundant proteins, e.g., those highlighted proteins in FIG. 18, for digesting the corresponding lignocellulosic materials. For example, to digest a cellulose material, one would choose at least one gene or expression products thereof selected from the group consisting of SACTE_0237, SACTE_0236, SACTE_2347, SACTE_3159, SACTE_0482, SACTE_0265, SACTE_0357, SACTE_4439, SACTE_0562, SACTE_0358, SACTE_4343, SACTE_1546, SACTE_1310, SACTE_4638, SACTE_5668, SACTE_3717, SACTE_3590, SACTE_2172, SACTE_4571, SACTE_5978, SACTE_6428, SACTE_2313, and SACTE_0366. In a preferred embodiment, the composition comprises at least three or four of the genes or expression products.

In one preferred embodiment, one would use all the highlighted proteins for digesting the corresponding lignocellulosic materials.

In another embodiment, one would add gene expression products from the list in Table 1 to a substrate to be digested. For example, for preferred cellulose digestion, one would select at least one member of the abundant proteins, as described above, and at least one member of the "additional useful genes" in Table 1.

TABLE 1

ActE genes or expression products useful for lignocellulosic degradation.		
Gene or Expression Product Combinations	Preferred subsets	Additional Useful Genes
SACTE_0236, SACTE_0237, SACTE_3159, SACTE_0482, SACTE_2347, and SACTE_0265.	Cellulose degradation	SACTE_0229, SACTE_0230, SACTE_0231, SACTE_0232, SACTE_0233, SACTE_0234, SACTE_0235, SACTE_0480, SACTE_0481, SACTE_0483, SACTE_0562, SACTE_0563, SACTE_0733, SACTE_0734, SACTE_2286, SACTE_2287, SACTE_2288, SACTE_2289, SACTE_3158, SACTE_4737, and SACTE_6428
SACTE_0265, SACTE_0357, SACTE_0358, SACTE_5230 and SACTE_5978	Xylan degradation	SACTE_0364, SACTE_0365, SACTE_0366, SACTE_0368, SACTE_0369, SACTE_0370, SACTE_0527, SACTE_0528, SACTE_5227, SACTE_5228, SACTE_5229, SACTE_5858, and SACTE_5859
SACTE_2313, SACTE_3064, SACTE_4246, SACTE_4571 and SACTE_5764	Chitin degradation	SACTE_0080, SACTE_0081, SACTE_0844, SACTE_0846, SACTE_0860, SACTE_3063, SACTE_4858, SACTE_6493 and SACTE_6494
SACTE_2347	Mannan degradation	
SACTE_1310	Pectin degradation	
SACTE_4638	Alginate release	
SACTE_5647	Galactose release	SACTE_5648
SACTE_4738 and SACTE_4755	Beta-1,3-glucan degradation	SACTE_4737, SACTE_4739 and SACTE_4756
SACTE_0236, SACTE_0237, SACTE_0265, SACTE_0358, SACTE_0482, SACTE_1310, SACTE_2347, SACTE_3159 and SACTE_3717	Cellulose and hemicelluloses degradation	SACTE_3065, SACTE_4730, SACTE_4755, and SACTE_5166

In one embodiment, the present invention is a method for digesting a lignocellulosic material, comprising exposing the material to a sufficient amount of a composition of enzymes, wherein the exposed material is at least partially digested. The enzymes may be ActE secretomes, and ActE secretomes may be prepared and isolated using the methods described above.

In another embodiment, the composition of enzymes for a method for digesting a lignocellulosic material may include ActE secretomes in a combination with secretomes from other organisms, or with enzymes or enzyme compositions, such as Spezyme CP, to increase the activity of both preparations by synergy of the enzymes contained in each preparation.

In another embodiment, the composition of enzymes for a method for digesting a lignocellulosic material may be any combinations of ActE genes and expression products as described above.

EXAMPLES

Materials and Methods

Genome Analysis. The complete genome sequence of *Streptomyces* sp. SirexAA-E (ActE, taxonomy ID 862751) was determined by the Joint Genome Institute, project ID 4086644. Gene annotation models were predicted using Prodigal (Hyatt, et al., 2010), examined using Artemis (Rutherford, et al., 2000), and are available at NCBI with the following accession numbers, GenBank: CP002993.1; RefSeq: NC_015953.1. Carbohydrate-active enzymes were

annotated by comparison of all translated open-reading frames to the CAZy database (Cantarel, et al., 2009). We collected CAZy annotated genes from the CAZy database (www.cazy.org). We then used BLASTP to compare all ActE protein-coding sequences to the CAZy database and to the pfam database (ftp://ftp.ncbi.nih.gov/pub/mmdb/cdd/little_endian/Pfam_LE.tar.gz). These two annotations were then crosschecked, and proteins annotated by both databases were identified as our final CAZy annotation. Secreted proteins were identified by SignalP, TatP, and SecretomeP analyses. BLAST was used to identify sequence orthologs in other organisms. Secondary metabolite gene clusters were identified by AntiSmash analysis (Medema, et al., 2011). CebR boxes were identified by using BLAST comparison of the *S. griseus* CebR box sequence to the ActE genome (Marushima, Ohnishi, et al., 2009). Networks of expression and functional categories were visualized using Cytoscape (Shannon, et al., 2003)

Biomass Substrates. Switchgrass and AFEX-treated switchgrass were obtained from Great Lakes Bioenergy Research Center. Extensively washed ionic liquid-treated switchgrass was the generous gift of Dr. Masood Hadi (Joint BioEnergy Institute). Wood kraft pulp preparations were the generous gift of Dr. Xuejun Pan (University of Wisconsin Department of Biosystems Engineering).

The complete genome sequence of *Streptomyces* sp. SirexAA-E (ActE, taxonomy ID 862751) was determined by the Joint Genome Institute, project ID 4086644. Gene annotation models were predicted using Prodigal (Hyatt, et al., 2010), examined using Artemis (Rutherford, et al., 2000),

and are available at NCBI with the following accession numbers, GenBank: CP002993.1; Ref Seq: NC_015953.1. Carbohydrate-active enzymes were annotated by comparison of all translated open-reading frames to the CAZY database (Cantarel, et al., 2009). We collected CAZY annotated genes from the CAZY database (See CAZY's website for detail information). We then used BLASTP to compare all ActE protein-coding sequences to the CAZY database and to the pfam database (the pfam database can be found in the website of the National Institute of Health (NIH)). These two annotations were then crosschecked, and proteins annotated by both databases were identified as our final CAZY annotation. Secreted proteins were identified by SignalP, TatP, and SecretomeP analyses. BLAST was used to identify sequence orthologs in other organisms. Secondary metabolite gene clusters were identified by AntiSmash analysis (Medema, et al., 2011). CebR boxes were identified by using BLAST comparison of the *S. griseus* CebR box sequence to the ActE genome (Marushima, Ohnishi, et al., 2009). Networks of expression and functional categories were visualized using Cytoscape (Shannon, et al., 2003)

RNA microarray. ActE was grown in minimal medium plus the indicated substrate for 7 days. The cell pellet was separated from the culture medium by centrifugation for 10 min at 3000×g. Microarray experiments were carried out as reported previously (Riederer, et al., 2011). The total RNA was extracted from the cell pellet and purified. The University of Wisconsin Gene Expression Center carried out the syntheses of cDNA and array hybridizations. Four-plex arrays were constructed by Nimblegen and hybridized with 10 µg of labeled cDNA. ArrayStar (v4.02, DNASTAR, Madison, Wis.) was used to quantify and visualize data. All analyses were based on three or more biological replicates per carbon source. Quantile normalization and robust multi-array averaging (RMA) were applied to the entire data set. Unless otherwise specified, expression levels are based on log₂ values and statistical analysis of the datasets were performed using the moderated t-test.

Preparation of Secretomes. Supernatants obtained from different culture media were prepared by centrifugation of the culture medium for 10 min at 3000×g, which removed the remaining insoluble polysaccharide and adhered cells. The supernatant fraction was then passed through a 0.22-µm filter in order to remove any remaining cells. For enzymatic assays, the secretomes were concentrated using a 3-kDa cut off ultrafiltration membrane. The concentration of secretome protein was determined by Bradford assay, and the typical yield was ~150-300 mg of total secreted protein per liter of culture medium.

Extracellular Protein Profiles. Extracellular proteins from culture secretomes were precipitated with trichloroacetic acid (TCA), resuspended in denaturing sample buffer (SDS and 2-mercaptoethanol), and separated by SDS-PAGE in 4-20% gels. Protein bands of interest were excised from the gel, digested with trypsin, desalted with C18 pipette tips (Millipore, Billerica, Mass.) and identified by MALDI-TOF (MDS SCIEX 4800 MALDI TOF/TOF, Applied Biosystems, Foster City, Calif.). Additional samples from the same culture secretomes were analyzed by LC-MS/MS to identify highly abundant proteins in the sample.

Ion exchange separation of the ActE secretome. The ActE cellulose secretome was diluted with cold deionized water until the ionic strength was less than 50 mM. The diluted sample was loaded onto an AKTApurifier™ chromatography station equipped with a 16/10 MonoQ FF ion exchange

proteins were eluted in a linear, 200 mL gradient of NaCl from 0 to 0.8 M in the same buffer. Fractions from the gradient elution were collected and separated by SDS PAGE. The proportional contribution of individual proteins in each fraction was estimated from SDS PAGE. Individual protein bands from each fraction were cut from the gel and submitted for LC-MS/MS analysis to confirm their identities.

LC-MS/MS Analyses. These experiments were performed at the University of Wisconsin Biotechnology Center. Samples were prepared by TCA precipitation of 100 ng of total secreted protein from 7-day old culture supernatants. Protein samples were digested with trypsin (sequencing grade trypsin, Promega, Madison, Wis.) and were desalted using C18 pipette tips (Millipore, Billerica, Mass.). High-energy collision dissociation (HCD) MS analyses employing a capillary LC-MS/MS were performed on an electrospray ionization FT/ion-trap mass spectrometer (LTQ Orbitrap XL, Thermo Fisher Scientific, San Jose, Calif.). The MS and MS/MS spectra were searched against the spectra obtained from the ActE proteome by using Scaffold (Scaffold_3_00_06, Proteome Software, Portland, Oreg.).

Enzyme Activity Measurements. Reducing sugar assays were carried out by mixing secretome preparations with polysaccharide-containing substrates including cellulose (either Whatman #1 filter paper or Sigmacell-20 as indicated), xylan, chitin, mannan, switchgrass, AFEX pretreated switchgrass, or ionic-liquid pretreated switchgrass²⁴. After incubation in 0.1 M sodium phosphate, pH 6 at 40° C. for 20 h, the reducing sugar content was detected by dinitrosalicylic acid assay (Miller, 1959) and calibrated by using glucose, xylose, or mannose as standards. Purified polysaccharide preparations had negligible background response in the absence of added enzymes. Cellobionic and gluconic acids were assayed by a coupled enzyme assay (K-GATE system, Megazyme, Bray Ireland). Spezyme CP was obtained from Genencor with batch number #4901522860. The distributions of soluble sugar oligomers obtained from secretome reactions were determined using a Shimadzu Liquid Chromatograph HPLC system (Shimadzu Scientific Instruments, Columbia, Md.) equipped with a refractive index detector (RID-10A) and a Phenomenex Rezex RPM-monosaccharide column. The temperature was maintained at 85° C. and Milli-Q water was used as the mobile phase at 0.6 mL min⁻¹ flow rate. Glucose, cellobiose, cellobiose, cellobiose, cellopentaose, and cellohexaose (Sigma) were used as standards. The integrated areas of peaks were analyzed by EZ start 7.2 SP1 software (Shimadzu).

Fractions obtained from the ion exchange separation of the ActE cellulose secretome were combined as unary, binary, ternary, and quaternary assemblies where the total protein concentration was fixed and the individual fractions contributed all, halves, thirds, or quarters of the total protein. The most active fraction was assembled from a ternary combination of fractions containing the following enzymes: fraction 1, SACTE_3159 (CBM33/CBM2 oxidative endocellulase, 95%) and SACTE_4738 (GH16 β-1,3 endoglucanase, 5%); fraction 2, SACTE_0237 (GH6 exocellulase, 60%), SACTE_0482 (GH5 endocellulase, 25%), SACTE_0237 (β-1,3 glucanase, 10%) and SACTE_3159 (oxidative endocellulase, <5%); and fraction 3, SACTE_0236 (GH48 exocellulase, 75%), SACTE_3717 (GH9 endocellulase, 20%) and SACTE_5457 (GH46 chitinase, 5%).

Cellobionic and gluconic acids were assayed by a coupled enzyme assay (K-GATE system, Megazyme, Bray Ireland),

either with or without the addition of a large excess of β -glucosidase (Cat. No. 31571, Lucigen, Middleton, Wis.).

Two lots of Spezyme CP were obtained from Genencor (#4900901244, Jan. 27, 2010 and #4901522860, Sep. 2, 2011). The specific activity of these two preparations was indistinguishable.

HPLC Analysis. The distributions of soluble sugar oligomers obtained from secretome reactions without and with the addition of excess β -glucosidase (Lucigen) were determined using a Shimadzu Liquid Chromatograph HPLC system (Shimadzu Scientific Instruments, Columbia, Md.) equipped with a refractive index detector (RID-10A) and a Phenomenex Rezex RPM-monosaccharide column. The temperature was maintained at 85° C. and milli-Q water was used as the mobile phase at 0.6 mL min⁻¹ flow rate. Glucose, cellobiose, cellotriose, cellotetraose, and cellopentaose (Sigma) were used as standards. The integrated areas of peaks were analyzed by EZ start 7.2 SP1 software (Shimadzu).

For the experiments shown in FIG. 21, the ActE secretome (1 μ g total protein); CelLcc_CBM3a (1 μ g); ActE secretome (0.5 μ g) and CelLcc_CBM3a (0.5 μ g); or Spezyme CP (1 μ g total protein) were used. The products of the enzyme reactions detected by HPLC were: ActE secretome, 95% cellobiose, 5% glucose; CelLcc_CBM3a reaction, 90% cellobiose, 10% glucose; ActE & CelLcc_CBM3a, 5% cellotetraose, 80% cellotriose, 15% cellobiose; Spezyme CP, 33% cellobiose, 67% glucose. All products could be converted to glucose in the presence of excess β -glucosidase.

CelLcc_CBM3a. The nucleotide and amino acid sequence of CelLcc_CBM3a is shown in FIG. 22. CelRcc_CBM3a is an engineered exoglucanase composed of the catalytic core of *C. thermocellum* CelL (Cthe_0405, residues 32 to 429) fused to a *C. thermocellum*-derived linker sequence and the CBM3a domain from Cthe_3077, the CipA scaffoldin. This construct was created to better understand the performance of enzymes that are normally targeted to the clostridial cellulosome. The replacement of the dockerin domain in Cthe_0405 with the CBM3a domain abrogates the need for a cellulosomal attachment to obtain maximal catalytic activity from CelLcc_CBM3a on solid substrates. The indicated nucleotide sequence was sub-cloned into wheat germ cell-free translation (Makino et al., 2010) and *E. coli* expression vectors (Blommel et al., 2009) for protein production. CelLcc_CBM3a was purified by standard immobilized metal (Ni²⁺) chromatography. There was no difference in the specific activity of the protein prepared by these two methods.

Example 1

ActE Exhibits High Cellulolytic Activity Relative to other Cellulolytic Organisms

Prokaryotes such as *Streptomyces* are often easier to grow than eukaryotes (i.e., fungi such as *T. reesei*), and aerobes are often easier and more energetically efficient to grow than anaerobes. *Streptomyces* may also have an advantage of producing antibiotics that limit the ability of other organisms to contaminate the culture medium during growth (Galm et al., 2011; Susi et al., 2011). This may be of advantage during large-scale culture with non-sterile biomass materials such as will be encountered in the biofuels industry.

When compared to other cellulolytic organisms (FIG. 1 and FIG. 6), ActE grows well on pure cellulose substances including amorphous cellulose (cellulose treated with phos-

phoric acid so as to remove all crystalline structure), filter paper (containing a mixture of amorphous and crystalline cellulose) and Sigmacell (primarily in the crystalline state as determined by X-ray powder diffraction), as well as other polysaccharides such as beta-1,3-glucan (callose), xylan, and chitin. ActE also grows well on biomass samples such as corn stover, ammonia-fiber expansion pretreated corn stover, switchgrass, ammonia-fiber expansion pretreated switchgrass, ionic liquids pretreated switchgrass, bleached spruce wood kraft pulp, and unbleached lodgepole pine kraft pulp.

FIG. 1 compares the ability of ActE, *S. coelicolor* A3(2) (NCBI taxonomy ID 100226) and *S. griseus* (NCBI CP002993.1; RefSeq: NC_015953.1) to grow in minimal medium containing filter paper as the only carbon and energy source. These images demonstrate the considerably different capabilities of the three ostensibly cellulolytic organisms. Thus ActE completely destroys the filter paper and achieves high cell density, while the two other, reputedly highly cellulolytic strains are only capable of weak colony formation attached to the filter paper. This result establishes that ActE has uniquely high cellulolytic capacity relative to other *Streptomyces* strains reported to also have this capability (Forsberg et al., 2011). In fact, the images of FIG. 1 and FIG. 6 demonstrate ActE has cellulolytic capacity rivaling that of *T. reesei* strain Rut-C30, which is widely acknowledged to be the industrial benchmark for cellulolytic capacity (Merino and Cherry, Adv. Biochem. Eng. Biotechnol. 108:95-120, 2007).

Example 2

Pretreatments Useful for Generating Fermentable Sugars

In the biofuels arena, the desired cellulose fractions of plant biomass are protected by the crystalline packing of the individual cellulose strands, and by the surrounding coating of hemicellulose and lignin. In order to most efficiently access the cellulose, chemical pretreatments are required to “loosen up” the plant cell wall structure. In this context, “loosen up” may mean removal of the lignin fraction, partial hydrolysis of feruloyl and acetyl esters present in hemicellulose, and changes in the crystallinity of the cellulose. An optimal pretreatment retains all fractions of biomass (i.e., lignin, hemicellulose and cellulose) in physical states that can be subsequently used by microbes and enzymes as substrates.

Ammonia-fiber expansion is a pretreatment that uses a combination of ammonia gas, low pressure, and low temperature to effect the loosening process (Balan et al., 2009; Chundawat et al., 2011; International Patent Publication No.: WO 2010/125679). It is particularly effective with grasses, and retains all fractions of the biomass for subsequent valorization without introducing water or salts into the biomass. Ionic liquids pretreatment comprises mixing a charged chemical substance (i.e., the ionic liquid) in equal mass proportions with the biomass material. Interactions between the ionic liquid substance and the biomass cause the crystalline structure of cellulose to convert to an amorphous state (Cheng et al., 2011; Li et al., 2011) but the biomass also becomes heavily contaminated with the ionic liquid during this pretreatment, requiring extensive washing with water, a valuable resource in many localities. Kraft pulping is a method for production of paper from wood that involves treatment of the biomass material with strong alkali, sodium sulfite and moderate temperature, resulting in destruction of

the lignin and hemicellulose from the desired cellulose fraction; the final biomass material is also heavily contaminated with salts that also requires extensive washing with water to remove. Acid pretreatments retain the lignin and cellulose but destroy the hemicellulose fraction, and in doing so create toxic substances derived from the decomposition of hemicellulose. Because of the need to neutralize the acid, this pretreatment generates a large contamination of salt that also requires extensive washing with water. SPORL is an acidic pretreatment that uses sulfuric acid, elevated

(CAZy) database. Among the 6357 predicted protein-coding genes, 167 have one or more domains assigned to CAZy families, including 119 glycoside hydrolases (GHs), 29 carbohydrate esterases (CEs), 6 polysaccharide lyases (PLs) and 85 carbohydrate binding modules (CBMs). ActE contains 45 different types of GH families, 4 PL families, 7 CE families, and 21 CBM families. The number of total CAZy domains and diversity of CAZy families is comparable to other highly cellulolytic organisms.

TABLE 2

	ActE	<i>S. coelicolor</i>	<i>S. griseus</i>	<i>C. thermocellum</i>	<i>C. japonicus</i>
Genome size (nt)	7414440	8667507	8545929	3843301	4576573
Proteome size	6357	8153	7136	3173	3750
Total CAZy	167	221	132	103	183
Proteins					
% CAZy Proteins ^a	2.6%	2.7%	1.8%	3.2%	4.9%
Total GH ^b	119	154	80	70	124
Total PL ^c	6	11	4	6	14
Total CE ^d	29	36	23	20	28
Total CBM ^e	85	98	68	121	134
antiSMASH clusters ^f	22	24	37	3	4
Genes in clusters	620	718	1139	89	111
% antiSMASH	9.8%	8.8%	16.0%	2.8%	3.0%

^aProteins classified as Carbohydrate Active Enzymes (CAZy).

^bGH, glycoside hydrolase.

^cPL, pectate lyase.

^dCE, carbohydrate esterase.

^eCBM, carbohydrate binding module.

^fPutative antibiotic producing gene cluster.

temperature, and sodium bisulfite to effect the pretreatment (Wang et al., 2009; Tian et al., 2011). In SPORL, the lignin and hemicellulose are destroyed and cellulose is recovered, but the cellulose is again heavily contaminated with salts and toxic substances derived from chemical decomposition of hemicellulose.

ActE secretomes are highly effective for degradation of lignocellulosic material pre-treated with AFEX. ActE secretomes are also effective for degrading lignocellulosic material pretreated with ionic liquids, Kraft pulping, acid or SPORL and for degrading untreated lignocellulosic material.

Example 3

ActE Genome has High Content of Genes Encoding Carbohydrate Active enZymes (CAZy) Relative to other Cellulolytic Organisms

Protein-coding sequences of the ActE genome (Hyatt et al., 2010) were analyzed by BLAST comparison (Altschul et al., 1990) to the Carbohydrate Active enZyme (CAZy) database (Cantarel et al., 2009).

Table 2 compares the genomic characteristics of ActE with well-known soil-isolated *Streptomyces* that produce antibiotics and with two model cellulolytic bacteria, *Clostridium thermocellum* and *Cellvibrio japonicus* (Lynd, Weimer, et al., 2002; Deboy, et al., 2008; Riederer, et al., 2011). Putative biomass-degrading protein-coding sequences from ActE were identified by BLAST analysis of the finished genome to the Carbohydrate Active enZyme

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Nearly all publically available *Streptomyces* genomes encode a relatively high percentage of genes for putative cellulolytic enzymes. Interestingly, ActE and the antibiotic producing *Streptomyces*, *S. griseus* and *S. coelicolor*, shown in Table 2 have similar numbers and compositions of CAZy families, but substantially different genome sizes. However, these antibiotic-producing *Streptomyces* are not highly cellulolytic (FIG. 1). Relative to *S. griseus* and *S. coelicolor*, the ActE genome contains two unique CAZy families but does not possess 16 CAZy families present in these species. However, ActE contains more representatives in 13 CAZy families. Enrichment of certain CAZy families was observed in other highly cellulolytic organisms. For example, *C. thermocellum* contains 16 genes in the GH9 family alone. It is interesting to consider whether the reduction in total genome size and differences in CAZy composition between ActE and other closely related soil-dwelling *Streptomyces* might have arisen from evolutionary specialization of ActE, perhaps driven by association with the *Sirex*-fungal symbiosis.

ActE contained 12 CAZy families not found in the other model cellulolytic organisms shown in FIG. 3, including GHs, CBMs, and PLs. Seven other CAZy categories, primarily hemicellulases, were shared only with *T. reesei*. ActE had 23 GH, 10 CBM and 2 PL not found in *Thermobifida fusca*, another cellulolytic *Actinomycetales*, which had only 1 GH and 1 CBM not found in ActE. The genome sequence revealed *C. japonicus* (strain Ueda 107) is highly enriched in GH43 enzymes required for hemicellulose utilization, but is missing a key reducing end exocellulase (bacterial GH48) required for robust growth on cellulose [e.g., see page 5459

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of (DeBoy et al., 2008)]; both of these enzyme families are present in highly cellulolytic ActE. Furthermore, ActE also contained 6 genes from the CBM33 family, recently shown to catalyze oxidative cleavage of chitin (Vaaje-Kolstad et al., 2010) and cellulose (Forsberg et al. 2011). Thus, ActE has genomic composition overlapping other cellulolytic organisms, but with notable expansion in the CAZy composition for both hydrolytic and oxidative enzymes and the presence of the complete set of enzymes required for efficient cellulose deconstruction.

Example 4

Genome-wide Gene Expression Analysis of ActE CAZy Gene

Gene expression profiles were determined for ActE grown on purified polysaccharides and plant biomass by whole genome microarrays (FIGS. 4 and 5, FIGS. 9 to 14). Genome-wide gene expression was analyzed as a functional annotation network composed of ActE genes (circles) connected to predicted functional groups (triangles; KEGG or CAZy). In FIG. 4, the network was annotated with genome-wide microarray expression data to indicate genes that were differentially expressed when ActE was grown on either AFEX-SG or glucose, and further annotated to indicate normalized expression levels observed during growth on AFEX-SG. While many aspects of metabolism are modestly changed in response to these different carbon sources, the CAZy and ABC transporter categories were substantially enriched in differentially expressed genes (FIG. 4, green circles). Furthermore, pentose sugar metabolism, sulfur metabolism, and some amino acid biosynthesis pathways (e.g., aromatic amino acids) were also highly induced during growth on AFEX-SG relative to other carbon sources (FIGS. 9-14). In contrast, ribosomal, secondary metabolite, and DNA repair genes showed little change in expression across the conditions examined. Within the CAZy functional group, there was a large induction of genes that contained both a GH domain and a CBM2 domain. Among the 11 genes in the ActE genome that contain a CBM2 domain, 6 were induced greater than 4-fold during growth on AFEX-SG. Furthermore, 9 of the 11 CBM2 containing proteins were identified in the secreted proteome (FIG. 3).

Example 5

ActE CAZy Gene Expression is Dependent on ActE Growth Substrate

Given the large number of differentially expressed CAZy genes identified in the network analysis, Applicants analyzed the expression of this group of genes in cultures grown

on different carbon sources (FIG. 5, FIG. 15 and FIG. 16). As with other cellulolytic organisms, there was strong correlation between the content of the secreted proteomes and the most highly expressed genes. Of the 167 ActE genes containing CAZy domains, 68 genes (FIG. 5, group 1) showed distinct increases in expression when grown on different polymeric substrates, 14 genes (FIG. 15, group 2) did not show any appreciable level of expression, and 85 genes (FIG. 16, group 3) showed moderate changes in expression with the different substrates. A significant fraction of these genes contained translocation signals for either the Sec or twin-arginine translocation pathways, and genes encoding structural polypeptides for these translocation pathways were also highly expressed. Besides correlation with secreted proteins, the transcriptomic studies also gave insight into co-regulated gene clusters that potentially encode functional units for utilization of different polysaccharides by ActE. In the following, the 130 genes with normalized expression intensities in the top 2% of all genes are described.

During growth on cellulose, four CAZy genes (SACTE_0236, SACTE_0237, SACTE_3159, and SACTE_0482) showed >15-fold increase in transcript abundance (FIG. 5), and the corresponding proteins were highly enriched in the secreted proteome. None of these four were obviously placed in a gene cluster, and the two most highly expressed genes, SACTE_0236 and SACTE_0237, while adjacent on the chromosome, were transcribed in opposite directions. Nevertheless, these four most highly expressed genes and three others that showed >5-fold increase in transcript abundance (SACTE_3717, SACTE_6428, SACTE_2347, Table 3) were associated with a conserved 14 bp palindromic promoter sequence, TGGGAGCGCTCCCA (the CebR binding element). CebR proteins are LacI/GalR-like transcriptional regulators shown to provide transcriptional control of gene expression in response to the presence of cellobiose or other small oligosaccharides in *S. griseus*, *S. reticuli*, and *Thermobifida fusca* (Marushima, Ohnishi, et al., 2009; Water and Schrempf, 1996; Deng and Fong, 2010). Likewise, the genes (SACTE_2285 to SACTE_2289) encoding a CebR regulator (SACTE_2285), a GH1 protein (β -glucosidase), a two-protein cellobiose transporter system, and an extracellular solute binding protein were associated with a CebR binding element and were also among the most highly expressed genes during growth on cellulose. These latter five genes have 75% or greater sequence identity with the cellobiose utilization operon identified in *S. griseus* and *S. reticuli* (Marushima, Ohnishi, et al., 2009; Schlosser and Schrempf, 1996). There were only 15 genes annotated as hypothetical or domain of unknown function (12% up-regulated during growth on cellulose, a considerably smaller percentage of these than in the entire genome (27%).

TABLE 3

Analysis of upstream DNA sequence elements in ActE genes upregulated during growth on cellulose.

Locus	Catalytic domain	CBM Annotated function	Sequence ^a	Rank ^b	Fold change ^b
SACTE_0236	GH48	CBM2 1,4-beta cellobiohydrolase	TGGGAGCGCTC CCA	1	21.7
SACTE_0237	GH6	CBM2 1,4-beta cellobiohydrolase	TGGGAGCGCTC CCA	2	17.3

TABLE 3-continued

Analysis of upstream DNA sequence elements in ActE genes upregulated during growth on cellulose.						
Locus	Catalytic domain	CBM	Annotated function	Sequence ^a	Rank ^b	Fold change ^b
SACTE_3159	CBM33	CBM2	Cellulose-binding domain	TGGGAGCGCTC CCA	3	16.2
SACTE_0482	GH5	CBM2	Endo-1,4-beta-glucosidase	TGGGAGCGCTC CCA	4	15.4
SACTE_2288			Transport systems inner membrane component	TGGGAGCGCTC CCA	5	11.2
SACTE_3717	GH9	CBM2	1,4-beta cellobiohydrolase	TGGGAGCGCTC CCA	6	9.7
SACTE_6428	CBM33		Chitin-binding, domain 3	GGGAGCGCTCC CA	9	7.9
SACTE_2347	GH5	CBM2	Beta-mannosidase	TGGGAGCGCTC CCA	11	5.0
SACTE_2287			Transport systems inner membrane component	TGGGAGCGCTC CCA	15	4.3
SACTE_2289			Family 1 extracellular solute-binding protein	TGGGAGCGCTC CCA	19	3.9
SACTE_0352			GCN5-related N-acetyltransferase	TGGGAGCGCTC CCA	22	3.6
SACTE_2286	GH1		Glycoside hydrolase 1	GGGAGCGCTCC CA	27	3.4
SACTE_0483		CBM2	Cellulose-binding family protein	GGGAGCGCTCC CA	503	1.6
SACTE_0562	GH74	CBM2	Secreted cellulase (endo)	TGGGAGCGCTC CCA	5759	0.7
SACTE_2285			LacI family transcriptional regulator (CebR)	TGGGAGCGCTC CCA	6229	0.6

^aPredicted binding sequence element found upstream from gene locus.

^bRanking and fold change in expression intensity detected by microarray for ActE genes when grown on cellulose relative to glucose.

Several characteristics distinguished expression during growth on either xylan or chitin. First, unique sets of genes were induced, as there was only 14% and 10% overlap, respectively, when compared to cellulose. Second, ~33% of the top 2% of genes expressed during growth on either xylan or chitin were annotated as hypothetical or domain of unknown function, which greatly exceeds the unknown fraction in the cellulose secretome. During growth on xylan, two clusters of genes were up-regulated. One extended from SACTE_0357 to SACTE_0370, encoding proteins from the GH11, GH13, GH42, GH43, GH78, GH87, and CE4 families, a LacI-like transcriptional regulator, a secreted peptidase, and two sets of inner membrane transporters and associated solute binding proteins. Alternatively, during growth on chitin, three CBM33 proteins were up-regulated (SACTE_0080, SACTE_2313, SACTE_6493), and two of these had an immediately adjacent gene encoding a GH18 (SACTE_6494) or GH19 (SACTE_0081) that was up-regulated.

When ActE was grown on biomass samples, 14 additional CAZy genes were uniquely up regulated, and the corre-

sponding proteins were identified in the proteomic analysis of biomass secretomes (FIGS. 3 and 4). A gene cluster extending from SACTE_5858 to SACTE_5864 was uniquely up regulated during growth on biomass. Among these genes, SACTE_5860 and SACTE_5862 are annotated as a twin-arginine translocation pathway protein and an ABC transporter, respectively, while the rest are annotated either as hypothetical protein or as domain of unknown function.

Eight CAZy genes were >4-fold up-regulated during growth on cellulose, including endoglucanases, reducing and non-reducing end exoglucanases, xylanase and CBM33 proteins (FIG. 5, Table 4). During growth on xylan, eight CAZy genes were elevated >4-fold relative to glucose, including exoglucanase, xylanase, pectate lyase and other hemicellulases (Table 4). Furthermore, chitin-grown cells contained 2 up-regulated genes from CAZy families including chitinase (SACTE_4571) and a CBM33 protein [SACTE_2313, an ortholog of oxidative chitin oxidase from *S. marcescens* (Vaaje-Kolstad et al., 2010)]. Thus on a

genome-wide basis ActE selectively expresses small, distinct sets of CAZy genes during growth on pure polysaccharides, which is distinct from the larger numbers of CAZy

genes expressed by *T. reesei* (Herpoel-Gimbert et al., 2008), *C. thermocellum* (Raman et al., 2009; Riederer et al., 2011), and *T. fusca* (Chen and Wilson, 2007).

TABLE 4

<i>Streptomyces</i> sp. ActE genes with >4-fold expression increase during growth on pure polysaccharides.					
CAZy	Annotation	Fold increase			
		Sigmacell: glc	xylan: glc	chitin: glc	
<u>Sigmacell</u>					
SACTE_6428	CBM33	Chitin-binding, domain 3	7.06	1.64	1.81
SACTE_3159	CBM33, 2	Cellulose-binding domain, family II, bacterial type	13.03	1.90	1.29
SACTE_0358	GH11, CBM60, 36	Glycoside hydrolase, family 11, active site	6.28	4.01	2.12
SACTE_0236	GH48, CBM2, 37	Glycoside hydrolase, 48F	19.00	4.93	3.91
SACTE_0482	GH5, CBM2	Cellulose-binding family II/chitinase, carbohydrate-binding domain	11.84	3.01	2.00
SACTE_2347	GH5, CE3, CBM2, 37	Cellulose-binding family II/chitinase, carbohydrate-binding domain	4.46	1.17	0.99
SACTE_0237	GH6, CBM2	1,4-beta cellobiohydrolase	15.33	1.12	0.77
SACTE_3717	GH9, CBM4, 2	Carbohydrate-binding, CenC-like	8.03	2.61	1.55
SACTE_2288		Binding-protein-dependent transport systems inner membrane component	11.05	4.76	3.26
SACTE_0168		Transcription regulator LuxR, C-terminal	7.55	1.53	1.37
SACTE_0169		Glyceraldehyde 3-phosphate dehydrogenase, active site	5.01	0.75	1.08
SACTE_3594		Peptidase S1C, HrtA/DegP2/Q/S	4.52	3.36	2.70
SACTE_5228		Binding-protein-dependent transport systems inner membrane component	4.20	4.35	3.24
<u>Xylan</u>					
SACTE_4029	CE4	Glycoside hydrolase/deacetylase, beta/alpha-barrel	1.07	4.35	2.22
SACTE_0358	GH11, CBM60, 36	Glycoside hydrolase, family 11, active site	6.28	4.01	2.12
SACTE_0382	GH2, CBM42	Galactose-binding domain-like	1.79	4.18	2.46
SACTE_1230	GH23	Lytic transglycosylase-like, catalytic	1.29	5.64	3.70
SACTE_0816	GH31	Glycoside hydrolase, family 31	1.53	4.51	3.27
SACTE_0236	GH48, CBM2, 37	Glycoside hydrolase, 48F	19.00	4.93	3.91
SACTE_1290	GH53, CBM61	Galactose-binding domain-like	1.43	4.73	2.40
SACTE_5978	PL1, CBM35	Galactose-binding domain-like	2.00	6.86	2.12
SACTE_5325		Binding-protein-dependent transport systems inner membrane component	1.78	8.26	3.76
SACTE_6023		Galactose-binding domain-like	1.92	7.84	3.34
SACTE_1834		Alkaline phosphatase D-related	1.78	7.73	3.98
SACTE_6100		Sulfate transporter	2.07	7.45	4.75
SACTE_5361		hypothetical protein	1.77	7.20	3.94
SACTE_5163		Lambda repressor-like, DNA-binding	1.47	6.89	3.29
SACTE_6365		Isocitrate lyase/phosphorylmutase	1.88	6.82	4.01
SACTE_0254		Thiolase-like	2.13	6.76	5.02
SACTE_6478		FAD-dependent pyridine nucleotide-disulfide oxidoreductase	2.00	6.72	4.46
SACTE_3570		hypothetical protein	1.61	6.71	3.72
SACTE_0590		Polyketide cyclase/dehydrase	1.55	6.67	4.42
SACTE_3152		Twin-arginine translocation pathway, signal sequence	1.41	6.60	2.98

TABLE 4-continued

<i>Streptomyces</i> sp. ActE genes with >4-fold expression increase during growth on pure polysaccharides.					
CAZy	Annotation	Fold increase			
		Sigmacell: glc	xylan: glc	chitin: glc	
SACTE_5285	Bacterial bifunctional deaminase-reductase, C-terminal	1.71	6.54	3.33	
SACTE_1383	Glycerophosphoryl diester phosphodiesterase	1.08	6.50	3.51	
SACTE_4333	Binding-protein-dependent transport systems inner membrane component	1.37	6.46	3.58	
SACTE_3876	hypothetical protein	1.21	6.42	2.73	
SACTE_6340	Monoxygenase, FAD-binding	2.82	6.27	3.69	
SACTE_4237	hypothetical protein	1.82	6.27	2.91	
SACTE_5136	NAD(P)-binding domain	2.20	6.27	2.87	
SACTE_6561	hypothetical protein	2.92	6.06	5.65	
SACTE_0686	Transcription regulator AsnC-type	0.88	6.04	2.72	
SACTE_0817	NUDIX hydrolase, conserved site	1.96	6.03	3.19	
SACTE_3004	Type II secretion system F domain	1.67	6.01	4.18	
SACTE_1835	DoxX	1.66	5.97	3.30	
SACTE_1933	hypothetical protein	0.93	5.96	2.77	
SACTE_6290	Glyoxalase/bleomycin resistance protein/dioxygenase	1.86	5.95	4.10	
SACTE_5583	hypothetical protein	1.33	5.87	4.56	
SACTE_0586	hypothetical protein	1.40	5.81	2.90	
SACTE_0046	NADH: flavin oxidoreductase/NADH oxidase, N-terminal	2.48	5.75	4.56	
SACTE_1096	Mandelate racemase/muconate lactonizing enzyme, N-terminal	1.19	5.73	3.32	
SACTE_2897	hypothetical protein	1.18	5.73	3.81	
SACTE_5359	Rhs repeat-associated core	1.30	5.70	2.41	
SACTE_0200	hypothetical protein	1.34	5.67	3.64	
SACTE_0018	hypothetical protein	1.67	5.63	3.58	
SACTE_5542	hypothetical protein	2.03	5.61	3.52	
SACTE_3137	hypothetical protein	1.46	5.61	3.91	
SACTE_0017	DNA helicase, UvrD/REP type	2.32	5.58	4.26	
SACTE_0672	hypothetical protein	1.53	5.54	3.20	
SACTE_1393	Urease, beta subunit	2.08	5.53	3.67	
SACTE_0064	Transcription regulator PadR N-terminal-like	2.17	5.52	3.07	
SACTE_1168	Peptidase S1/S6, chymotrypsin/Hap	0.98	5.51	3.36	
SACTE_6371	hypothetical protein	1.37	5.51	3.44	
SACTE_4334	Binding-protein-dependent transport systems inner membrane component	1.46	5.50	3.35	
SACTE_2457	CDP-glycerol glycerophosphotransferase	1.07	5.48	3.79	
SACTE_4734	Binding-protein-dependent transport systems inner membrane component	1.21	5.44	3.31	
SACTE_3661	hypothetical protein	1.76	5.44	3.25	
SACTE_0036	hypothetical protein	1.75	5.43	2.99	
SACTE_6005	Citrate synthase-like, core	1.01	5.38	2.90	
SACTE_6562	hypothetical protein	2.34	5.36	3.37	
SACTE_1937	Major facilitator superfamily MFS-1	0.88	5.34	3.02	
SACTE_6220	Dodecin flavoprotein	2.13	5.32	5.08	
SACTE_0778	FMN-binding split barrel	1.13	5.28	2.72	
SACTE_5672	Acytransferase 3	1.33	5.28	3.09	
SACTE_5989	Cysteine-rich domain	1.40	5.24	3.11	
SACTE_5296	HTH transcriptional regulator, MarR	1.42	5.22	2.96	
SACTE_2021	hypothetical protein	1.44	5.17	2.54	
SACTE_1845	Transposase, IS4-like	1.69	5.16	3.30	
SACTE_1771	Phage T4-like virus tail tube gp19	1.55	5.10	1.71	
SACTE_2583	hypothetical protein	1.38	5.10	3.11	

TABLE 4-continued

CAZy	Annotation	Fold increase		
		Sigmacell: glc	xylan: glc	chitin: glc
SACTE_5957	Helix-turn-helix, HxIR type	2.38	5.09	3.95
SACTE_4642	hypothetical protein	1.31	5.08	3.05
SACTE_3695	Aminoglycoside/hydroxyurea antibiotic resistance kinase	1.41	5.03	3.76
SACTE_0079	ATPase-like, ATP-binding domain	2.21	5.01	2.98
SACTE_0727	hypothetical protein	2.54	5.00	3.88
SACTE_0019	hypothetical protein	1.37	5.00	2.40
SACTE_6422	<i>Streptomyces</i> cyclase/dehydrase	2.40	4.99	3.57
SACTE_4348	Bacterial extracellular solute-binding protein, family 5	1.60	4.97	3.06
SACTE_5318	Forkhead-associated (FHA) domain	1.50	4.93	2.84
SACTE_5413	Urease accessory protein UreF	1.94	4.93	2.52
SACTE_5434	Glutathione S-transferase, C-terminal-like	2.41	4.93	2.96
SACTE_6061	Glyoxalase/bleomycin resistance protein/dioxygenase	1.61	4.92	2.18
SACTE_0025	hypothetical protein	1.58	4.92	4.22
SACTE_5552	Transposase, IS4-like	1.94	4.92	3.26
SACTE_4156	HTH transcriptional regulator, LysR	1.57	4.86	2.81
SACTE_5600	hypothetical protein	1.78	4.83	2.01
SACTE_5331	Conserved hypothetical protein CHP03086	1.56	4.82	2.96
SACTE_0784	hypothetical protein	1.43	4.80	2.65
SACTE_0045	NAD(P)-binding domain	1.74	4.78	3.35
SACTE_5426	Twin-arginine translocation pathway, signal sequence	0.80	4.77	2.68
SACTE_2654	4Fe—4S ferredoxin, iron-sulfur binding domain	1.30	4.77	2.68
SACTE_2288	Binding-protein-dependent transport systems inner membrane component	11.05	4.76	3.26
SACTE_2324	Membrane insertion protein, OxaA/YidC, core	0.91	4.75	2.58
SACTE_0142	Amidohydrolase 2	1.28	4.71	2.65
SACTE_0787	hypothetical protein	1.66	4.70	2.93
SACTE_5790	hypothetical protein	1.28	4.69	2.83
SACTE_6291	hypothetical protein	1.25	4.68	3.13
SACTE_6499	hypothetical protein	1.66	4.67	3.29
SACTE_6548	Lytic transglycosylase-like, catalytic	1.97	4.66	3.20
SACTE_3087	Major facilitator superfamily MFS-1	1.30	4.66	3.26
SACTE_5512	hypothetical protein	1.79	4.64	3.48
SACTE_0491	hypothetical protein	2.44	4.63	2.71
SACTE_0312	Thiamine pyrophosphate enzyme, C-terminal TPP-binding	2.32	4.60	3.49
SACTE_6130	hypothetical protein	1.47	4.55	2.64
SACTE_3787	Helix-turn-helix type 3	1.38	4.53	2.73
SACTE_0040	hypothetical protein	1.64	4.52	4.80
SACTE_2461	Macrocin-O-methyltransferase	1.07	4.51	3.00
SACTE_5041	hypothetical protein	1.50	4.49	3.25
SACTE_5540	Transposase, IS204/IS1001/IS1096/IS1165	1.79	4.49	2.99
SACTE_0776	Protein of unknown function DUF6, transmembrane	1.34	4.48	2.52
SACTE_0785	Bacterial ThiB	1.67	4.43	2.93
SACTE_0360	Binding-protein-dependent transport systems inner membrane component	1.70	4.43	2.39
SACTE_3569	Protein of unknown function DUF1023	1.00	4.42	2.78
SACTE_2986	hypothetical protein	1.62	4.42	2.96
SACTE_4732	Twin-arginine translocation pathway, signal sequence	2.08	4.41	2.72

TABLE 4-continued

<i>Streptomyces</i> sp. ActE genes with >4-fold expression increase during growth on pure polysaccharides.					
CAZy	Annotation	Fold increase			
		Sigmacell: glc	xylan: glc	chitin: glc	
SACTE_5228	Binding-protein-dependent transport systems inner membrane component	4.20	4.35	3.24	
SACTE_0406	Binding-protein-dependent transport systems inner membrane component	1.34	4.35	2.52	
SACTE_6516	Binding-protein-dependent transport systems inner membrane component	2.24	4.34	3.41	
SACTE_1781	hypothetical protein	1.16	4.34	2.56	
SACTE_5936	Radical SAM	1.43	4.33	2.23	
SACTE_0819	Protein of unknown function DUF962	1.50	4.33	2.83	
SACTE_4539	NERD	1.42	4.32	3.98	
SACTE_0532	Binding-protein-dependent transport systems inner membrane component	3.47	4.31	2.42	
SACTE_3300	hypothetical protein	1.68	4.31	2.59	
SACTE_6277	hypothetical protein	2.24	4.31	3.11	
SACTE_0941	Twin-arginine translocation pathway, signal sequence	1.32	4.30	2.63	
SACTE_1115	GntR, C-terminal	1.57	4.29	2.63	
SACTE_6105	Fatty acid hydroxylase	1.63	4.29	2.78	
SACTE_4407	Spherulation-specific family 4	1.19	4.29	4.15	
SACTE_5387	hypothetical protein	1.24	4.27	3.08	
SACTE_5053	NmrA-like	1.23	4.27	3.05	
SACTE_5562	Amino acid ABC transporter, permease protein, 3-TM domain, His/Glu/Gln/Arg/opine family	1.37	4.26	3.75	
SACTE_5522	Galactose-binding domain-like	1.82	4.26	2.62	
SACTE_5484	Transcription regulator, TetR-like, DNA-binding, bacterial/archaeal	1.45	4.21	3.24	
SACTE_6526	Restriction endonuclease, type IV-like, Mrr	2.31	4.20	2.40	
SACTE_4164	hypothetical protein	1.06	4.19	2.48	
SACTE_4979	Transcription regulator, TetR-like, DNA-binding, bacterial/archaeal	1.20	4.19	2.34	
SACTE_0952	hypothetical protein	1.33	4.18	2.02	
SACTE_1785	hypothetical protein	1.25	4.17	1.94	
SACTE_3454	hypothetical protein	1.46	4.16	2.32	
SACTE_1271	Class II aldolase/adducin, N-terminal	1.77	4.16	2.65	
SACTE_1760	hypothetical protein	1.38	4.13	2.07	
SACTE_0035	hypothetical protein	1.93	4.13	3.13	
SACTE_0247	Protein of unknown function DUF2241	1.30	4.10	2.77	
SACTE_3796	F420-dependent enzyme, PPOX class, family Rv2061, putative	1.43	4.10	3.33	
SACTE_4641	hypothetical protein	1.43	4.09	2.60	
SACTE_4816	Peptidase S26, conserved region	1.17	4.09	2.77	
SACTE_2331	Major facilitator superfamily MFS-1	1.15	4.08	2.20	
SACTE_1666	hypothetical protein	1.44	4.07	2.46	
SACTE_5867	Mammalian cell entry, mce1C	1.79	4.07	2.92	
SACTE_2705	AMP-binding, conserved site	1.38	4.07	2.75	
SACTE_6014	Binding-protein-dependent transport systems inner membrane component	0.89	4.07	2.51	
SACTE_2018	Putative DNA binding domain	1.05	4.06	2.63	
SACTE_5690	Gluconate transporter	1.00	4.05	2.29	
SACTE_3243	hypothetical protein	0.91	4.05	2.23	
SACTE_0786	Polynucleotidyl transferase, ribonuclease H fold	1.81	4.03	2.98	
SACTE_6450	Rhamnose isomerase related	2.72	4.02	2.90	
SACTE_0097	Beta-lactamase-related	1.70	4.02	2.52	

TABLE 4-continued

<i>Streptomyces</i> sp. ActE genes with >4-fold expression increase during growth on pure polysaccharides.					
CAZy	Annotation	Fold increase			
		Sigmacell: glc	xylan: glc	chitin: glc	
SACTE_6341	FMN-binding split barrel, related	1.82	4.01	2.45	
SACTE_1483	hypothetical protein	0.82	4.01	2.75	
SACTE_0754	Uncharacterised protein family UPF0060	1.21	4.00	2.51	
SACTE_5308	Winged helix-turn-helix transcription repressor DNA-binding	1.33	4.00	1.56	
SACTE_5862	ABC transporter, conserved site	1.87	4.00	3.05	
Chitin					
SACTE_2313	CBM33	1.08	1.24	4.77	
SACTE_4571	GH18, CBM57, 2	0.88	1.37	4.08	
SACTE_5381	hypothetical protein	1.31	3.09	10.06	
SACTE_5386	hypothetical protein	0.96	1.59	8.49	
SACTE_1949	Peptidase M4, thermolysin	1.30	2.16	7.57	
SACTE_6519	Binding-protein-dependent transport systems inner membrane component	2.00	3.04	7.36	
SACTE_0243	Protein kinase-like domain	1.68	2.55	6.89	
SACTE_6520	ABC transporter, conserved site	1.03	1.18	6.25	
SACTE_5384	hypothetical protein	1.16	2.39	5.99	
SACTE_6463	hypothetical protein	1.28	2.52	5.85	
SACTE_6561	hypothetical protein	2.92	6.06	5.65	
SACTE_5383	hypothetical protein	1.06	1.69	5.28	
SACTE_6518	hypothetical protein	1.66	1.91	5.21	
SACTE_4797	hypothetical protein	2.22	0.34	5.19	
SACTE_6170	Domain of unknown function DUF1996	1.47	3.49	5.12	
SACTE_6220	Dodecin flavoprotein	2.13	5.32	5.08	
SACTE_0254	Thiolase-like	2.13	6.76	5.02	
SACTE_2678	Protein of unknown function DUF397	1.40	1.13	5.02	
SACTE_5968	hypothetical protein	1.58	1.31	4.90	
SACTE_4757	Acetyl-coenzyme A carboxyltransferase, C-terminal	1.64	0.59	4.86	
SACTE_0040	hypothetical protein	1.64	4.52	4.80	
SACTE_6100	Sulfate transporter	2.07	7.45	4.75	
SACTE_1833	Twin-arginine translocation pathway, signal sequence	1.64	1.56	4.64	
SACTE_5583	hypothetical protein	1.33	5.87	4.56	
SACTE_0046	NADH: flavin oxidoreductase/NADH oxidase, N-terminal	2.48	5.75	4.56	
SACTE_5398	hypothetical protein	1.45	1.73	4.55	
SACTE_6144	Twin-arginine translocation pathway, signal sequence	1.21	1.13	4.52	
SACTE_6478	FAD-dependent pyridine nucleotide-disulfide oxidoreductase	2.00	6.72	4.46	
SACTE_0590	Polyketide cyclase/dehydrase	1.55	6.67	4.42	
SACTE_2112	Homeodomain-like	1.44	1.33	4.40	
SACTE_0017	DNA helicase, UvrD/REP type	2.32	5.58	4.26	
SACTE_5841	Protein of unknown function, ATP binding	1.90	3.09	4.24	
SACTE_0025	hypothetical protein	1.58	4.92	4.22	
SACTE_3004	Type II secretion system F domain	1.67	6.01	4.18	
SACTE_4407	Spherulation-specific family 4	1.19	4.29	4.15	
SACTE_0307	Protein of unknown function DUF320, <i>Streptomyces</i> species	1.13	1.79	4.15	
SACTE_6290	Glyoxalase/bleomycin resistance protein/dioxygenase	1.86	5.95	4.10	
SACTE_5286	hypothetical protein	1.33	3.34	4.07	
SACTE_5953	Protein of unknown function, ATP binding	1.35	2.11	4.05	

TABLE 4-continued

<i>Streptomyces</i> sp. ActE genes with >4-fold expression increase during growth on pure polysaccharides.					
CAZy	Annotation	Fold increase			
		Sigmacell: glc	xylan: glc	chitin: glc	
SACTE_6365	Isocitrate lyase/phosphorylmutase	1.88	6.82	4.01	

Example 6

Composition of ActE Secretome is Dependent on ActE Growth Substrate

To identify secreted proteins, supernatants from ActE cultures grown on glucose, cellobiose, cellulose, xylan, chitin, switchgrass, AFEX-SG, and IL-SG were analyzed by LC-MS/MS (FIG. 3 and FIG. 18). The proteins were sorted into a descending rank according to spectral counts, and sets whose spectral counts summed to 95% of the total protein in each secretome are shown. FIG. 3A summarizes the percentages of CAZy families in the detected proteins. The glucose secretome had a protein concentration of ~0.03 g/L of culture medium, and among the 136 proteins identified only 3% had a CAZy annotation. Indeed, the majority (>90%) likely originated from cell lysis. In contrast, the polysaccharide secretomes had a protein concentration of ~0.3 g/L of culture medium, a ~10-fold increase from the glucose secretome. Pectate lyase (SACTE_1310), chondroitin/alginate lyase (SACTE_4638), an extracellular solute binding protein (SACTE_4343), bacterioferritin (SACTE_1546), and catalase (SACTE_4439) were observed in all polysaccharide secretomes. The first two proteins, SACTE_1310 and SACTE_4638, have signal peptides and are thus secreted as part of the response needed for growth on polysaccharides.

FIG. 3 and FIG. 18 further demonstrate that 22 proteins accounted for 95% of the total spectral counts during growth on cellulose; two-thirds were from CAZy families. The five most abundant proteins, in order and representing ~85% of the total spectral counts, were reducing and non-reducing exoglucanases (SACTE_0236 and SACTE_0237), a CBM33 polysaccharide monooxygenase (SACTE_3159), an endoglucanase (SACTE_0482), and a β -mannosidase (SACTE_2347). The first four proteins encode a non-redundant set of enzymes that likely provide the essential activities required for utilization of crystalline cellulose (Deboy, et al., 2008). Among the 22 most abundant proteins, there were representatives from 9 different GH families, two CE families, two PL families, and two additional CMB33 proteins. Collectively, these secreted proteins represent ~20% of the CAZy composition in the ActE genome.

There were substantial differences in the composition of the xylan and chitin secretomes as compared to the cellulose secretome (FIG. 3 and FIG. 18). In the xylan secretome, 92 proteins comprise 95% of the detected spectral counts. Twenty GHs from 18 different CAZy families were included, along with 1 CE4 and 2 PL family proteins. Thus, growth on xylan elicits secretion of representatives from half of the total CAZy families found in the ActE genome. The broad distribution of hemicellulolytic enzymes in the xylan secretome contrasts with the considerably less diverse composition of the chitin secretome, which consists of 7 representatives from GH18 (e.g., chitinase, endo beta-N-acetylglucosaminidase), 2 from GH19 (e.g., chitinase, lysozyme),

and 1 chitinolytic CBM33 (FIG. 18). While chitinolytic CAZy families account for two-thirds of the proteins secreted during growth on chitin, they represent only ~6% of the diversity of CAZy families found in the genome. These results document the substantially different substrate-specific responses of ActE during growth on different polysaccharides.

The secretomes isolated from cells grown on switchgrass, AFEX-SG, and IL-SG contained the highly abundant secreted proteins identified in the purified cellulose and xylan experiments and some additional proteins. These additional proteins likely reflect cellular response to the more complex composition of polysaccharides present in the biomass samples. The increased diversity of proteins present in the biomass secretome also increased the efficiency of reaction with plant biomass (FIG. 2C). In total, the biomass secretomes contained 31 different CAZy families that contributed to the total spectral counts (~70% of the CAZy families present in the ActE genome), thus representing coordinated and extensive use of CAZyme families present in the ActE genome for biomass utilization.

The gene loci of the 117 proteins observed only in the glucose secretome are: SACTE_0494; SACTE_0514; SACTE_0541; SACTE_0548; SACTE_0604; SACTE_0669; SACTE_0687; SACTE_0800; SACTE_0810; SACTE_0899; SACTE_1006; SACTE_1045; SACTE_1068; SACTE_1069; SACTE_1111; SACTE_1201; SACTE_1240; SACTE_1285; SACTE_1328; SACTE_1344; SACTE_1368; SACTE_1419; SACTE_1426; SACTE_1506; SACTE_1522; SACTE_1586; SACTE_1650; SACTE_1861; SACTE_1888; SACTE_1934; SACTE_2036; SACTE_2049; SACTE_2068; SACTE_2238; SACTE_2403; SACTE_2468; SACTE_2431; SACTE_2558; SACTE_2645; SACTE_2729; SACTE_2755; SACTE_2756; SACTE_2801; SACTE_2819; SACTE_3012; SACTE_3037; SACTE_3067; SACTE_3086; SACTE_3088; SACTE_3097; SACTE_3219; SACTE_3327; SACTE_3361; SACTE_3371; SACTE_3385; SACTE_3389; SACTE_3392; SACTE_3414; SACTE_3438; SACTE_3511; SACTE_3604; SACTE_3716; SACTE_3896; SACTE_3948; SACTE_3955; SACTE_3956; SACTE_3960; SACTE_3961; SACTE_3989; SACTE_3995; SACTE_4030; SACTE_4031; SACTE_4038; SACTE_4039; SACTE_4073; SACTE_4081; SACTE_4083; SACTE_4145; SACTE_4191; SACTE_4194; SACTE_4205; SACTE_4224; SACTE_4281; SACTE_4283; SACTE_4376; SACTE_4397; SACTE_4399; SACTE_4415; SACTE_4462; SACTE_4501; SACTE_4550; SACTE_4565; SACTE_4566; SACTE_4567; SACTE_4568; SACTE_4591; SACTE_4610; SACTE_4616; SACTE_4618; SACTE_4652; SACTE_4718;

SACTE_4768; SACTE_4791; SACTE_4795;
 SACTE_4830; SACTE_4860; SACTE_4873;
 SACTE_4926; SACTE_4959; SACTE_5028;
 SACTE_5081; SACTE_5192; SACTE_5267;
 SACTE_5482; SACTE_5519; SACTE_5983; and
 SACTE_6342.

The gene loci of the 9 proteins observed only in the Sigmacell secretome are: SACTE_0236; SACTE_0482; SACTE_0562; SACTE_2313; SACTE_2347; SACTE_3590; SACTE_3717; SACTE_4571; and SACTE_6428.

The gene loci of the 46 proteins observed only in the xylan secretome are: SACTE_0081; SACTE_0169; SACTE_0365; SACTE_0379; SACTE_0383; SACTE_0464; SACTE_0528; SACTE_0549; SACTE_0634; SACTE_0880; SACTE_1003; SACTE_1130; SACTE_1239; SACTE_1324; SACTE_1325; SACTE_1356; SACTE_1364; SACTE_1367; SACTE_1603; SACTE_1680; SACTE_1858; SACTE_1949; SACTE_2768; SACTE_3064; SACTE_4231; SACTE_4246; SACTE_4363; SACTE_4459; SACTE_4483; SACTE_4515; SACTE_4607; SACTE_4612; SACTE_4624; SACTE_4730; SACTE_4755; SACTE_4858; SACTE_5166; SACTE_5230; SACTE_5231; SACTE_5418; SACTE_5457; SACTE_5630; SACTE_5647; SACTE_5682; SACTE_5751; and SACTE_6439.

In the xylan secretome, five proteins accounted for half of the total secreted protein. These were xylanases (GH10 and GH11, respectively; SACTE_0265, 9.7% and SACTE_0358, 8.1%), extracellular xylose isomerase (SACTE_5230, 12.7%), acetyl xylan esterase (CE4; SACTE_0357, 11.7%), and pectate lyase (PL1, SACTE_5978, 6.6%). Among the remaining 98 proteins, there were numerous GH families. Given the complexity of hemicellulose, which is enriched in xylan but also contains many other sugars and many different bonding linkages between these sugars, it is noted that these additional proteins represent many GH families associated with unique hemicellulolytic activities.

Although not analyzed in FIG. 34, the chitin secretome contained ten proteins from the chitinase GH18 (49% of total protein) and GH19 (21%) families. In addition, the CBM33 protein SACTE_2313, having 50% primary sequence identity with the CBP21 chitin oxygenase from *S. marcescens*, was also detected (3.9%). Insect molt and fungal hyphae provide abundant chitin, likely accounting for the utility of these enzymes in the natural environment. There were 50 other proteins (63 total) that comprised 95% of the chitin secretome. Relative to the glucose, Sigmacell, and xylan secretomes, the following 15 proteins were observed only in the chitin secretome: SACTE_0746, SACTE_0844, SACTE_0860, SACTE_1702, SACTE_2033, SACTE_2059, SACTE_2062, SACTE_2384, SACTE_3685, SACTE_4468, SACTE_4472, SACTE_4727, SACTE_5330, SACTE_5764, and SACTE_6494.

The gene loci of the 19 proteins observed only in the switchgrass secretome are: SACTE_0642; SACTE_1130; SACTE_1250; SACTE_1858; SACTE_2033; SACTE_3012; SACTE_3777; SACTE_4198; SACTE_4571; SACTE_4624; SACTE_4669; SACTE_4676; SACTE_4718; SACTE_4738; SACTE_5220; SACTE_5418; SACTE_5685; SACTE_5751; and SACTE_5880.

The gene loci of the 8 proteins observed only in the IL-SG secretome are: SACTE_0132; SACTE_0880; SACTE_2556; SACTE_4246; SACTE_4515; SACTE_4702; SACTE_5231; and SACTE_5330.

There were no proteins observed only in the AFEX-SG secretome when compared to either the switchgrass or IL-SG secretomes.

Example 7

Minimized Size of ActE Enzymes Increases Specific Activity

When ActE is grown on Sigmacell, AFEX-SG, IL-SG, AFEX-CS, unbleached lodgepole pine kraft pulp (UBLPKP) or bleached spruce wood kraft pulp (BSKP), the characteristic secretome consists of the proteins that permit deconstruction of these substrates into sugars that can be used for growth (FIG. 23). Interestingly, ActE is not capable of growing on lodgepole pine pretreated by SPORL, indicating this pretreatment produces toxins that inhibit the growth of highly cellulolytic microbes. When ActE is grown on cellobiose, which it does readily and rapidly, it produces a secretome that is distinct from those obtained from ActE grown on cellulose, xylan or biomass substrates, demonstrating that ActE has highly specific responses to different polymeric substances that are present in biomass. This behavior is distinct from that observed for *T. fusca*, another cellulolytic *Actinomyces*, and from *C. thermocellum*, where each organism produced similar sets of secreted proteins during growth on either cellulose or cellobiose (Chen and Wilson, 2007; Riederer et al., 2011). This result indicates ActE contains a unique regulatory mechanism for controlling cellulose deconstruction genes that can provide exquisite control of their production under desired circumstances.

For a single enzyme from a secretome, (Segel, Enzyme kinetics: behavior and analysis of rapid equilibrium and steady state enzyme systems. Wiley, New York, 1993) the specific activity ($\mu\text{mol}/\text{min}/\text{mg}$) is defined as mol of product formed per unit time (i.e., $\mu\text{mol}/\text{min}$) per unit mass of enzyme (i.e., mg). Specific activity is the parameter that must be used in making comparisons of catalytic properties between enzymes with different molecular masses. If two enzyme isoforms yield the same $\mu\text{mol}/\text{min}$, the isoform with the smaller molecular weight will, by definition, have the higher specific activity. In this application, it is relevant to consider the implications of a 10% or more reduction in the mass of an enzyme required to treat gigatonnes of biomass.

In the cellulose secretome, five proteins contributed ~85% of the total spectral counts. These were reducing and non-reducing end exoglucanases, endoglucanases, and CBM33 (SACTE_0237, SACTE_0236, SACTE_2347, SACTE_0482 and SACTE_3159); xylanase, another endoglucanase, and another CBM33 were also abundant (SACTE_0265, SACTE_3717 and SACTE_6428). According to the definition provided above, size minimization is a way to achieve the desired increases in specific activity. Interestingly, the set of ActE enzymes described above are on average 10% smaller in mass than their closest orthologs from *T. fusca* (Chen and Wilson, 2007), suggesting size minimization may have occurred in ActE (Table 5). These enzymes also provide all of the requisite catalytic reactions needed for the deconstruction of crystalline cellulose.

TABLE 5

ActE cellulose secretome proteins and corresponding best match
in *T. fusca*. The single protein SACTE_0237 is the best match to both Cel6A and Cel6B
suggesting one protein from ActE might replace two proteins from another organism.

ActE Gene locus	CAZy	residues	MW	identity	coverage	<i>T. fusca</i> Gene locus	Protein name	residues	MW
SACTE_0237	GH6	586		49	80	Tfu_1074	Cel6A	441	45844
SACTE_0237	GH6	586	61062	62	93	Tfu_0620	Cel6B	596	63548
SACTE_0236	GH48	954	100726	57	95	Tfu_1959	Cel48A	984	107127
SACTE_2347	GH8	562	57753	45	23	Tfu_2176	Cel9A	880	95203
SACTE_3159	CBM33	362	37787	42	71	Tfu_1665	E8	438	46808
SACTE_0482	GH5	456	47654	51	97	Tfu_0901	Cel5A	466	49807
SACTE_0265	GH10	458	47683	44	95	Tfu_2923	Xyl10A	491	53185
SACTE_3717	GH9	909	96338	61	82	Tfu_1627	Cel9B	998	107045
SACTE_6428	CBM33	222	24668	62	99	Tfu_1268	E7	222	25372
Average identity, coverage				53	82				
Sum ActE		4509	473671			with Cel6A		4920	530391
		4509	473671			with Cel6B		5075	548095
Percentage with Cel6A		92%	89%			with Cel6A, B		5516	593939
Percentage with Cel6B		89%	86%						
Percentage with Cel6A, B		82%	80%						

Example 8

ActE Secretome Specific Activity is Comparable to that of Spezyme CP™

The enzymatic activities of ActE secretomes were compared with a commercial secretome, Spezyme CP. The enzyme cocktail of Spezyme CP was prepared from *T. reesei* Rut-C30, thus providing a useful, routinely available reference point for the capabilities of other cellulolytic organisms. HPLC analysis showed that the ActE cellulose secretome released cellobiose as the primary product during reaction with cellulose (FIG. 2A, 95% of products), which is distinct from the higher proportion of glucose produced by the *T. reesei* secretome. Similarly, the primary products from xylan and mannan were xylobiose and mannobiose, respectively. Upon accounting for total glucose equivalents released, the ActE secretome obtained from growth on pure cellulose had specific activity that was about half of that provided by Spezyme CP (FIG. 2A, inset). Interestingly, the ActE secretome obtained from growth on pure cellulose had higher specific activity for deconstruction of pure mannan than Spezyme CP (FIG. 2B). Additionally, the ActE secretome obtained from growth on pure xylan had higher specific activity for reaction with pure xylan than Spezyme CP. Cellulose, xylan, and mannan are all abundant in pinewood, thus accounting for the necessity of each of the major catalytic activities detected.

Anion exchange chromatography was performed to fractionate the ActE secretome obtained from cells grown on cellulose as the sole carbon source. We identified fractions that hydrolyzed pure polysaccharides by biochemical assays (FIG. 7), and confirmed the identity of the protein or proteins contained in these fractions by mass spectrometry (FIG. 17). Where multiple polypeptides were present, the identity of each was confirmed by mass spectrometry to correspond to the indicated gene locus. In several cases, these most likely arise from proteolysis of a single protein found in the secretome. Fractions containing the maximum cellulase activity were highly enriched in SACTE_0236 and SACTE_0237, reducing and non-reducing end cellobiohydrolases from the GH6 and GH48 families, respectively. SACTE_0265 and SACTE_2347 were identified as the major proteins present in fractions associated with xylan and mannan hydrolysis, respectively. A CBM33 polysaccharide

monoxygenase (SACTE_3159) was also identified in the ion exchange profile. Moreover, beta-1,3 glucanase activity was identified in fractions that were enriched in SACTE_4755.

When ActE was grown on either ammonia fiber expansion-treated switchgrass (AFEX-SG) (Li, C. et al., 2011) or ionic liquid-treated switchgrass (IL-SG), the secretomes had ~2-fold increase in specific activity relative to the cellulose secretome and were equivalent to Spezyme CP for reaction with both the AFEX- and IL-treated biomass (FIG. 2C) (Li, C. et al., 2011). The ActE secretomes retained greater than 60% of maximal activity for the hydrolysis of AFEX- and IL-SG from 30 to 55° C. and 35 to 47° C., respectively, which is comparable to recent reports on the temperature profile of secretomes from thermophilic biomass-degrading fungi (Tolonen et al., 2011) (FIG. 8A). The secretomes showed a pH optimum of ~7 for reaction with AFEX-SG and a pH optimum of ~8 for reaction with IL-SG. Moreover, these secretomes retained greater than 60% of maximal activity over the ranges of pH 4.5 to 8.0, and pH 7.0 to 8.0, respectively (FIG. 8B). These optimal pH values are considerably higher than observed for Spezyme CP.

Example 9

ActE Produces Cellobiose as the Primary Extracellular Product of Cellulose Utilization

The isolated ActE secretomes contained substantial ability to release reducing sugars from pure polysaccharides. Cellobiose accounted for ~95% of soluble sugar released from pure cellulose and glucose represented the remainder; celotriose and cellobionic acid were not detected. Neither cellobiosidase nor β -glucosidase was detected in the ActE secretome. Thus ActE produces cellobiose as the primary extracellular product of cellulose utilization and also grows vigorously on this. Dominance of cellobiose may help to channel cellulolytic activity to only a subset of the *Sirex* community. Since genes encoding cellobiose oxidase and cellobiose dehydrogenase (Eastwood et al., 2011; Langston et al., 2011) were not present in ActE, biological reduction systems for the CBM33 proteins may be provided by other members of the *Sirex* community, in analogy to that

described for the heterologous complex of *T. aurantiacus* GH61 and *Humicola insolens* cellobiose dehydrogenase (Langston et al., 2011).

Example 10

Enzymatic Activity of the ActE Secretome can be Improved by Adding One or More Enzymes from other Organisms or Sources

In the ActE secretome, enzymes SACTE_0236, SACTE_0237, and SACTE_3717 (GH48, GH6, and GH9, respectively) showed decreases in content of the native forms over a 24 h period, and SACTE_0236 and SACTE_0237 were converted into ~50 kDa fragments (FIG. 24). SACTE_0359 (CBM33) also showed a time-dependent decrease. The reactions could be slowed but not eliminated by addition of phenylmethylsulfonyl fluoride (1 mM), a possible inhibitor of serine proteases (Turini et al., 1969), but not by EDTA (10 mM), a possible inhibitor of metalloproteases (Trop and Birk, 1970).

SACTE_5668, a serine protease, was detected in all pure polysaccharide secretomes (FIG. 3), while another metallopeptidase, SACTE_3389 annotated as peptidase M24B, X-Pro dipeptidase/aminopeptidase P, was detected in all secretomes at low level (0.026%). The protease SACTE_5530 (peptidase S1/S6, chymotrypsin/Hap, 0.1%) was also present in all polysaccharide and biomass samples. The proteases SACTE_5668 (annotated secreted peptidase, 0.3%) and SACTE_4231 (serine/cysteine peptidase, trypsin-like, 0.039%) were also detected in all pure polysaccharide secretomes, and the protease SACTE_6303 (serine/cysteine peptidase, trypsin-like, 0.039%) was also present in all biomass samples. Elimination of one or more of these proteases may impart stabilization of the enzymatic activity in the secreted proteome.

Addition of Cellcc_CBM3a (SEQ ID Nos:63 and 64), an engineered exoglucanase (FIG. 22) that produces cellobiose with low specific activity alone (FIG. 21), gave a synergistic increase in the activity of the ActE cellulose secretome. This result demonstrates the potential for heterologous supplementation of the ActE secretome to improve its performance by replacing an enzyme activity that is lost to proteolysis.

Example 11

ActE Cellulolytic Activity Requires a Minimal Set of Enzymes

When the ActE secretome obtained from growth on cellulose was fractionated by ion exchange chromatography (FIG. 7), several fractions were obtained that could be tested in unary, binary, ternary and quaternary combinations for reconstitution of cellulose hydrolysis and other enzymatic activities (FIG. 25). SDS PAGE and LC-MS/MS analysis showed that these fractions contained the following polypeptides in the approximate weight percentages: fraction 1, SACTE_3159 (CBM33/CBM2 oxidative endocellulase, 95%) and SACTE_4738 (GH16 β -1,3 endoglucanase, 5%); fraction 2, SACTE_0237 (GH6 non-reducing end exocellulase, 60%), SACTE_0482 (GH5 endocellulase, 25%), SACTE_4755 (GH64 β -1,3 glucanase, 10%) and SACTE_3159 (oxidative endocellulase, <5%); and fraction 3, SACTE_0236 (GH48 reducing end exocellulase, 75%), SACTE_3717 (GH9 endocellulase, 20%) and SACTE_5457 (GH46 chitinase, 5%). These results demonstrate that SACTE_3159 (oxidative endocellulase) provides a comple-

mentary activity to SACTE_0482 and SACTE_3717 (hydrolytic endocellulolytic activity). Evidently, the oxidative reaction provides breaks in the cellulose strands that can be readily used by non-reducing and reducing end exocellulases also present in the secretome to processively deconstruct the polymeric material.

According to the current understanding of reactions required for hydrolysis of crystalline cellulose, SACTE_3159 (CBM33/CBM2 oxidative endocellulase), SACT_0482 (GH5), and SACTE_3717 provide endocellulolytic activities, while SACTE_0237 (GH6) provides non-reducing end exocellulase reaction and SACTE_0236 (GH48) provides reducing end exocellulase activity.

FIG. 16 shows that the secretome contains beta-1,3 endoglucanase activity. The majority of this activity corresponds to the fractions containing SACTE_4738 and SACTE_4755. These enzymes hydrolyze callose, a cellulose-like material that is typically produced by plants in response to wounding by invasive insects and other trauma.

The proteins described here constitute a naturally evolved and matched set specialized for the hydrolysis of cellulosic substrates.

Example 12

ActE Mannanase Specific Activity Increases as Mannanase Molecular Weight Decreases

FIG. 26 shows that the mannanase activity present in the ActE secretome is associated with fractions containing various naturally truncated variants of SACTE_2347 (GH5) with molecular weights of ~57, ~45, and ~37 kDa. Fractions F9 through F1 from ion exchange chromatographic separation of the ActE secretome were examined for mannan-deconstruction activity by Zymogram assay. The basis of the Zymogram assay is as follows: Congo Red stain interacts with the polysaccharide fraction (mannan) incorporated into the gel and imparts a red color. When an enzyme's activity hydrolyzes the mannan, the interaction of Congo Red with the polysaccharide is broken and the gel takes on a dark grey appearance. Of note, the strongest mannanase activity was observed in fraction F1, which primarily contains the 37 kDa truncated variant. Corresponding to the definition of specific activity given above, the 37 kDa variant has an ~35% increase in specific activity relative to the 57 kDa variant. This provides a naturally produced example of how size reduction may contribute to increased specific activity of enzymes.

Example 13

Recombination of ActE Secretome Fractions Provides Synergistic Cellulolytic Activity

FIG. 25 shows synergy of reaction obtained by recombining fractions obtained from ion exchange fractionation. In FIG. 25A, reactions were obtained from combinations of the fractions indicated by stars in FIG. 27 and FIG. 28. Combination of fractions E5 (oxidative endocellulase) and E11 (hydrolytic endo- and exocellulases) gave a ~30% increase in product yield over that expected from the arithmetic sum of reactions of E5 and E11 alone, i.e., synergy in reaction. Combination of fractions E5, E11 and F10 (hydrolytic endo- and exocellulases) gave ~60% increase in reactivity. In FIG. 25B, reactions were obtained from recombining fractions shown in FIG. 16. Titration of fraction B1 (full-length oxidative endocellulase) into D15 (hydrolytic

endo- and exocellulases) shows an optimal reactivity at ~1:1 ratio of proteins from the two fractions, while an excess of B1 relative to D15 causes decrease in reaction because of depletion of required exocellulase activities. Titration of fraction C4 (truncated oxidative endocellulase and beta-1,3-endocellulase) with D15 gave maximal stimulation (62% increase) at an 80:20 proportion. These results indicate both forms of oxidative endocellulase SACTE_3159 are catalytically active, with the smaller form providing a higher synergistic response, again corresponding to a specific activity increase associated with size minimization.

Example 14

The Function of ActE Xylanases can be Assigned by Functional Assay of Proteins Produced by Using Cell-free Translation

FIG. 29 shows that both of the xylanases identified in the fractions of ActE secretomes obtained from ion exchange chromatography can also be expressed using cell-free translation and demonstrated to be xylanases by catalytic activity assays. These proteins are SACTE_0265 and SACTE_0358. Other proteins that are not secreted were successfully expressed (SACTE_2548, SACTE_2286, SACTE_437) as control proteins, and as expected from their predicted intracellular localization, none of these controls exhibited xylanase activity. The negative result with the control proteins also demonstrates that the wheat germ extract used for cell-free translation of novel cellulolytic enzymes does not have an endogenous xylanase activity, as established in US Patent Application Publication No.: US2010/037094 (Fox and Elsen).

Example 15

Total Protein Secreted by ActE can be Increased

A minimal set of enzymes for biomass deconstruction can be defined by combining the additional enzymes whose expression is elicited during growth on biomass (Table 1) with enzymes uniquely expressed during growth on cellulose and xylan.

Besides assembling the proper enzymatic constituents, the level of total protein secreted is an important biotechnological constraint for industrial enzyme production. FIG. 30 shows the non-optimized level of secreted protein obtained from growth of ActE on different biomass substrates. By use of lignocellulosic substrates for growth, secreted protein levels up to 0.25 g per liter of culture medium can be readily obtained. Growth on non-polymeric substrates such as cellobiose does not elicit a secreted protein response. FIG. 15, FIG. 16 and Table 1 indicate that the twin-arginine pathway (Tat) is used during growth, thus identifying this pathway as playing a key role in the secretion of enzymes required for extracellular deconstruction of biomass polysaccharides (Natale et al., 2008; Chater et al., 2010). Methods to increase the titer of secreted proteins are known, and have been highly effectively when applied to *Streptomyces* and other organisms (Cereghino et al., 2002; Zhang et al., 2006; Nijland and Kuipers, 2008; Chater et al., 2010; Schuster and

Schmoll, 2010). These established methods can be applied to ActE to obtain more concentrated secretome preparations.

Example 16

ActE Enzymatic Activity Corresponds with Optimal Growth Conditions of Fermentation Organisms

FIG. 31 shows the temperature versus activity profile for ActE secretomes for reaction with cellulose, xylan and mannan. These profiles are well matched to the growth optima range for mesophilic fermentation organisms such as *Saccharomyces cerevisiae*, *Zymomonas mobilis*, *Escherichia coli* or others (Jarboe et al., 2010; Peralta-Yahya and Keasling, 2010), which are widely used for ethanol production from sugar hydrolysates. These hydrolysates are produced from biomass by the enzymatic action of highly cellulolytic secretomes, such as those described here from ActE. These optima are also well matched with the conditions found in the rumen, where the efficiency of conversion of animal feed, which is a biomass material, can be improved by addition of enzymes.

FIG. 32 shows the pH versus activity profiles for ActE secretomes for reaction with cellulose, xylan and mannan. These profiles are well matched to the growth optima range for fermentation organisms such as *S. cerevisiae*, *Z. mobilis*, *E. coli* or other organisms (Jarboe et al., 2010; Peralta-Yahya and Keasling, 2010) which are widely used for ethanol production from sugar hydrolysates such as might be produced from biomass by a highly cellulolytic secretome, such as those described here from ActE. These optima are also well matched with the conditions found in the rumen, where the efficiency of conversion of animal feed, which is a biomass material, can be improved by addition of enzymes. The ActE secretome retains high specific activity (>80% of maximal) at pH 7, which closely approximates that of the rumen. Secretomes from fungi such as *T. reesei* are considerably less active at neutral pH, rendering them less effective at neutral pH.

The high cellulolytic capacity of ActE, and its corresponding secretomes, coupled with the temperature and pH optima described above permit assembly of two-part systems to effect the simultaneous deconstruction of biomass and fermentation to fuels.

Example 17

ActE Induction in Medium Containing Various Percentages of Cellulose

To determine ActE's growth profile on cellulose as a carbon source ActE was grown in M63 media plus 5 g/L carbon. The carbon source ratio was adjusted from 100% cellulose to 100% glucose, total carbon in each culture was equal. Cells were grown for 6 days at 30 degrees. Supernatant was harvested, filtered, and separated by 4-20% SDS-PAGE. Results suggest that ActE is induced in media containing as little as 20% cellulose, with optimal induction in medium containing between 80%-100% cellulose (FIG. 33).

Example 18

Discussion

The work presented here provides the first genome-wide insight into how an aerobic microbe deconstructs polysac-

charides. ActE achieves efficient utilization of cellulose by a simple combination of well-understood hydrolytic reactions with newly identified oxidative reactions. The two required exoglucanases are each encoded by a single gene, which also represents the only example of their respective GH families in the genome. The proteins encoded by these genes provide reactions that are complementary to the reactions of other enzymes in the secretome, and provide cellobiose as the major product of reaction. We have discovered that many of the highly abundant enzymes secreted by ActE during growth on cellulose have reduced size relative to their orthologs from closely related organisms. This novel finding suggests natural evolution to improve specific activity has already occurred in ActE in response to growth in the highly specialized insect association. Additional specializations of ActE were identified by demonstrating the secretion of a unique set of proteins in response to biomass. In addition, this work defines how simple new combinations of improved biomass deconstruction enzymes can be assembled according to the propensities of the naturally evolved system.

The present work also indicates that insect-associated microbes such as ActE are important contributors to the vigorous attack on biomass by insects. The 'highly invasive' designation given to *Sirex* has been generally attributed to the combined action of wasp and fungus (Tabata and Abe, 2000; Bergeron et al., 2011). Species convergence is now recognized in the microbial communities associated with insects (Suen et al., 2010; Hulcr et al., 2011). Given the ubiquitous presence of *Streptomyces* in these communities, the enzymatic properties described here also contribute a potential risk to pine forests, including those used for industrial purposes.

The invention has been described in connection with what are presently considered to be the most practical and preferred embodiments. However, the present invention has been presented by way of illustration and is not intended to be limited to the disclosed embodiments. Accordingly, those skilled in the art will realize that the invention is intended to encompass all modifications and alternative arrangements within the spirit and scope of the invention as set forth in the appended claims.

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				565					570						575
Glu	Leu	Met	Lys	Asn	Ala	Tyr	Pro	Ala	Leu						
			580						585						

<210> SEQ ID NO 2

<211> LENGTH: 954

<212> TYPE: PRT

<213> ORGANISM: Streptomyces sp. ACTE

<400> SEQUENCE: 2

Val	Ala	Ala	Leu	Ala	Leu	Pro	Leu	Gly	Met	Thr	Ala	Ala	Ala	Gly	Thr
1				5					10					15	
Glu	Ala	Gln	Ala	Ala	Ala	Val	Ala	Cys	Ser	Val	Asp	Tyr	Thr	Thr	Ser
			20						25					30	

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Asp Trp Gly Ser Gly Phe Thr Thr Glu Leu Thr Leu Thr Asn Arg Gly
 35 40 45
 Ser Ala Ala Ile Asp Gly Trp Thr Leu Thr Tyr Asp Tyr Ala Gly Asn
 50 55 60
 Gln Gln Leu Thr Ser Gly Trp Ser Gly Thr Trp Ser Gln Ser Gly Lys
 65 70 75 80
 Thr Val Ser Val Lys Asn Ala Ala Trp Asn Gly Ala Ile Ala Ala Gly
 85 90 95
 Ala Ala Val Thr Thr Gly Ala Gln Phe Thr Tyr Ser Gly Ala Asn Thr
 100 105 110
 Ala Pro Thr Thr Phe Ala Val Asn Gly Thr Val Cys Ala Gly Ala His
 115 120 125
 Gln Pro Pro Ile Ala Val Leu Thr Ser Pro Ala Ala Gly Ala Val Phe
 130 135 140
 Ser Ala Gly Asp Pro Val Pro Leu Ala Ala Thr Ala Ala Ala Ala Asp
 145 150 155 160
 Gly Ala Thr Ile Ser Lys Val Glu Phe Tyr Asp Asp Thr Thr Leu Leu
 165 170 175
 Gly Thr Asp Thr Thr Ser Pro Tyr Ser Tyr Glu Ala Gly Gln Leu Ala
 180 185 190
 Ala Gly Ser His Ser Val Tyr Ala Arg Ala Tyr Asp Ser Leu Gly Ala
 195 200 205
 Ser Ala Asp Ser Pro Pro Ala Gly Ile Thr Val Val Thr Gly Pro Ala
 210 215 220
 Val Val Val Ser Pro Ala Gln Leu Gly Val Gln Gln Gly Arg Ser Gly
 225 230 235 240
 Thr Phe Asp Val Ser Leu Ser Thr Ala Pro Ala Ala Asp Val Thr Val
 245 250 255
 Thr Ala Ala Arg Ser Ala Gly Asn Thr Gly Leu Ser Val Thr Gly Gly
 260 265 270
 Ser Thr Leu Thr Phe Thr Pro Ala Asn Trp Ser Thr Pro Gln Lys Val
 275 280 285
 Thr Val Thr Ala Asp Gly Ser Gly Thr Gly Ala Ala Thr Phe Thr Val
 290 295 300
 Thr Ala Pro Gly His Gly Lys Ala Glu Val Thr Val Thr Gln Leu Ala
 305 310 315 320
 Ala Ala Lys Glu Tyr Asp Ala Arg Phe Leu Asp Leu Tyr Gly Lys Ile
 325 330 335
 Thr Asp Pro Ala Asn Gly Tyr Phe Ser Pro Glu Gly Ile Pro Tyr His
 340 345 350
 Ser Val Glu Thr Leu Ile Val Glu Ala Pro Asp His Gly His Glu Thr
 355 360 365
 Thr Ser Glu Ala Tyr Ser Tyr Leu Ile Trp Leu Gln Ala Met Tyr Gly
 370 375 380
 Lys Ile Thr Gly Asp Trp Thr Lys Phe Asn Gly Ala Trp Asp Thr Met
 385 390 395 400
 Glu Thr Tyr Met Ile Pro Thr His Ala Asp Gln Pro Thr Asn Ser Phe
 405 410 415
 Tyr Asp Ala Ser Lys Pro Ala Thr Tyr Ala Pro Glu His Asp Thr Pro
 420 425 430
 Asn Glu Tyr Pro Ala Val Leu Asp Gly Ser Ala Ser Ser Gly Ser Asp
 435 440 445
 Pro Ile Ala Ala Glu Leu Lys Ser Ala Tyr Gly Thr Asp Asp Ile Tyr

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450			455			460									
Gly	Met	His	Trp	Ile	Gln	Asp	Val	Asp	Asn	Val	Tyr	Gly	Tyr	Gly	Asn
465				470					475						480
Ala	Pro	Gly	Thr	Cys	Ala	Ala	Gly	Pro	Thr	Gln	Ala	Gly	Pro	Ser	Tyr
				485					490						495
Ile	Asn	Thr	Phe	Gln	Arg	Gly	Ser	Gln	Glu	Ser	Val	Trp	Glu	Thr	Val
			500					505						510	
Thr	His	Pro	Thr	Cys	Asp	Asn	Phe	Thr	Tyr	Gly	Gly	Ala	Asn	Gly	Tyr
		515					520					525			
Leu	Asp	Leu	Phe	Thr	Gly	Asp	Ser	Ser	Tyr	Ala	Lys	Gln	Trp	Lys	Phe
530						535					540				
Thr	Asn	Ala	Pro	Asp	Ala	Asp	Ala	Arg	Ala	Val	Gln	Ala	Ala	Tyr	Trp
545					550					555					560
Ala	Asp	Val	Trp	Ala	Lys	Glu	Gln	Gly	Lys	Ala	Gly	Glu	Val	Ala	Asp
				565					570						575
Thr	Val	Gly	Lys	Ala	Ala	Lys	Met	Gly	Asp	Tyr	Leu	Arg	Tyr	Ser	Met
			580					585						590	
Phe	Asp	Lys	Tyr	Phe	Lys	Lys	Ile	Gly	Asp	Cys	Val	Gly	Pro	Thr	Thr
		595					600					605			
Cys	Pro	Ala	Gly	Ser	Gly	Lys	Asp	Ser	Ala	His	Tyr	Leu	Met	Ser	Trp
610						615					620				
Tyr	Tyr	Ala	Trp	Gly	Gly	Ala	Thr	Asp	Thr	Ser	Ala	Gly	Trp	Ser	Trp
625					630					635					640
Arg	Ile	Gly	Ser	Ser	His	Ala	His	Gly	Gly	Tyr	Gln	Asn	Pro	Met	Ala
				645					650						655
Ala	Tyr	Ala	Leu	Ser	Ser	Val	Ala	Asp	Leu	Lys	Pro	Lys	Ser	Ala	Thr
			660					665						670	
Gly	Ala	Gln	Asp	Trp	Ala	Lys	Ser	Leu	Asp	Arg	Gln	Leu	Asp	Phe	Tyr
		675					680					685			
Gln	Trp	Leu	Gln	Ser	Asp	Glu	Gly	Ala	Ile	Ala	Gly	Gly	Ala	Thr	Asn
690						695					700				
Ser	Trp	Lys	Gly	Ser	Tyr	Ala	Gln	Pro	Pro	Ala	Gly	Thr	Pro	Thr	Phe
705					710					715					720
Tyr	Gly	Met	Tyr	Tyr	Asp	Glu	Lys	Pro	Val	Tyr	His	Asp	Pro	Pro	Ser
				725					730						735
Asn	Gln	Trp	Phe	Gly	Phe	Gln	Ala	Trp	Ser	Met	Glu	Arg	Val	Ala	Glu
			740					745						750	
Tyr	Tyr	His	Glu	Ser	Gly	Asp	Ala	Gln	Ala	Lys	Ala	Val	Leu	Asp	Lys
		755					760						765		
Trp	Val	Asp	Trp	Ala	Leu	Ser	Glu	Thr	Thr	Val	Asn	Pro	Asp	Gly	Thr
	770					775					780				
Tyr	Leu	Met	Pro	Ser	Thr	Leu	Gln	Trp	Ser	Gly	Ala	Pro	Asp	Thr	Trp
785					790					795					800
Asn	Ala	Ser	Asn	Pro	Gly	Ala	Asn	Ala	Gln	Leu	His	Val	Thr	Val	Ala
				805					810						815
Asp	Tyr	Thr	Asp	Asp	Val	Gly	Val	Ala	Gly	Ala	Tyr	Ala	Arg	Thr	Leu
			820					825						830	
Thr	Tyr	Tyr	Ala	Ala	Lys	Ser	Gly	Asp	Thr	Glu	Ala	Glu	Ala	Thr	Ala
		835					840							845	
Glu	Ala	Leu	Leu	Asp	Gly	Met	Trp	Gln	His	His	Gln	Asp	Asp	Ala	Gly
	850					855						860			
Val	Ala	Val	Pro	Glu	Thr	Arg	Ala	Asp	Tyr	Asn	Arg	Phe	Asp	Asp	Pro
865					870					875					880

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Val Tyr Val Pro Gly Gly Trp Thr Gly Ala Met Pro Asn Gly Asp Thr
 885 890 895

Val Asp Glu Asp Ser Thr Phe Leu Ser Ile Arg Ser Phe Tyr Lys Asp
 900 905 910

Asp Pro Asn Trp Pro Gln Val Gln Ala Tyr Leu Asp Gly Gly Ala Ala
 915 920 925

Pro Val Phe Thr Tyr His Arg Phe Trp Ala Gln Ala Asp Ile Ala Leu
 930 935 940

Ala Leu Gly Ala Tyr Ala Asp Leu Leu Glu
 945 950

<210> SEQ ID NO 3
 <211> LENGTH: 362
 <212> TYPE: PRT
 <213> ORGANISM: Streptomyces sp. ACTE

<400> SEQUENCE: 3

Met Ala Arg Arg Ser Arg Leu Ile Ser Leu Ala Ala Val Leu Ala Thr
 1 5 10 15

Leu Leu Gly Ala Leu Gly Leu Thr Ala Leu Trp Pro Gly Lys Ala Glu
 20 25 30

Ala His Gly Val Ala Met Thr Pro Gly Ser Arg Thr Tyr Leu Cys Gln
 35 40 45

Leu Asp Ala Leu Ser Gly Thr Gly Ala Leu Asn Pro Thr Asn Pro Ala
 50 55 60

Cys Arg Asp Ala Leu Ser Gln Ser Gly Ala Asn Ala Leu Tyr Asn Trp
 65 70 75 80

Phe Ala Val Leu Asp Ser Asn Ala Gly Gly Arg Gly Ala Gly Tyr Val
 85 90 95

Pro Asp Gly Ser Leu Cys Ser Ala Gly Asp Arg Ser Pro Tyr Asp Phe
 100 105 110

Ser Ala Tyr Asn Ala Ala Arg Ala Asp Trp Pro Arg Thr His Leu Thr
 115 120 125

Ser Gly Ala Thr Leu Lys Val Gln Tyr Ser Asn Trp Ala Ala His Pro
 130 135 140

Gly Asp Phe Arg Val Tyr Leu Thr Lys Pro Gly Trp Ala Pro Thr Ser
 145 150 155 160

Glu Leu Ala Trp Asp Asp Leu Gln Leu Val Gln Thr Val Ser Asn Pro
 165 170 175

Pro Gln Gln Gly Gly Ala Gly Thr Asn Gly Gly His Tyr Tyr Trp Asp
 180 185 190

Leu Ala Leu Pro Ser Gly Arg Ser Gly Asp Ala Leu Met Phe Ile Gln
 195 200 205

Trp Val Arg Ser Asp Ser Gln Glu Asn Phe Phe Ser Cys Ser Asp Ile
 210 215 220

Val Phe Asp Gly Gly Asn Gly Glu Val Thr Gly Ile Gly Gly Thr Gly
 225 230 235 240

Thr Pro Thr Pro Thr Pro Thr Pro Thr Pro Thr Pro Thr Asp
 245 250 255

Pro Glu His Ser Gly Ser Cys Met Ala Val Tyr Asn Val Val Ser Ser
 260 265 270

Trp Ala Gly Gly Phe Gln Ala Ser Val Glu Val Met Asn His Gly Thr
 275 280 285

Glu Pro Arg Asn Gly Trp Ala Val Gln Trp Lys Pro Gly Ser Gly Thr

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290					295					300					
Gln	Ile	Asn	Ser	Val	Trp	Asn	Gly	Ser	Leu	Ser	Thr	Gly	Ser	Asp	Gly
305					310					315					320
Thr	Val	Thr	Val	Arg	Asp	Val	Asp	His	Asn	Arg	Val	Ile	Ala	Pro	Asp
				325					330					335	
Gly	Ser	Val	Thr	Phe	Gly	Phe	Thr	Ala	Thr	Ser	Thr	Gly	Asn	Asp	Tyr
				340				345						350	
Pro	Ala	Gly	Thr	Ile	Gly	Cys	Val	Thr	Ser						
		355					360								
<210> SEQ ID NO 4															
<211> LENGTH: 456															
<212> TYPE: PRT															
<213> ORGANISM: Streptomyces sp. ACTE															
<400> SEQUENCE: 4															
Val	Lys	Arg	Phe	Leu	Ala	Leu	Leu	Ala	Thr	Cys	Ala	Thr	Val	Leu	Gly
1				5					10					15	
Leu	Thr	Ala	Leu	Thr	Gly	Pro	Gln	Ala	Val	Ala	Ala	Ala	Gly	Cys	Thr
			20					25					30		
Ala	Asp	Tyr	Thr	Ile	Thr	Ser	Gln	Trp	Gln	Gly	Gly	Phe	Gln	Ala	Ala
		35					40					45			
Val	Lys	Val	Thr	Asn	Leu	Gly	Thr	Pro	Val	Thr	Gly	Trp	Lys	Leu	Thr
		50				55					60				
Phe	Thr	Leu	Pro	Asp	Ala	Gly	Gln	Lys	Val	Val	Gln	Gly	Trp	Asn	Ala
65				70					75					80	
Ala	Trp	Ser	Gln	Ser	Gly	Ser	Ala	Val	Thr	Ala	Ala	Gly	Ala	Asp	Trp
			85						90					95	
Asn	Gly	Thr	Leu	Ala	Thr	Gly	Ala	Ser	Ala	Glu	Ala	Gly	Phe	Val	Gly
			100					105					110		
Ser	Phe	Thr	Gly	Ala	Asn	Pro	Pro	Pro	Thr	Ala	Phe	Ala	Leu	Asn	Gly
		115				120					125				
Val	Ala	Cys	Thr	Gly	Ser	Thr	Gly	Glu	Pro	Pro	Ala	Gly	Ser	Asp	Gly
	130					135					140				
Gly	Thr	Pro	Val	Asp	Val	Asn	Gly	Gln	Leu	His	Val	Cys	Gly	Val	Asn
145				150					155					160	
Leu	Cys	Asn	Gln	Tyr	Asp	Arg	Pro	Val	Gln	Leu	Arg	Gly	Met	Ser	Thr
			165					170						175	
His	Gly	Ile	Gln	Trp	Phe	Asp	Ala	Cys	Tyr	Asp	Ala	Ala	Ser	Leu	Asp
		180						185					190		
Ala	Leu	Ala	Asn	Asp	Trp	Lys	Ser	Asp	Leu	Leu	Arg	Ile	Ala	Met	Tyr
		195				200						205			
Val	Gln	Glu	Asp	Gly	Tyr	Glu	Thr	Asp	Pro	Ala	Gly	Phe	Thr	Arg	Arg
	210					215					220				
Val	Asn	Asp	Leu	Val	Asp	Met	Ala	Glu	Ala	Arg	Gly	Met	Tyr	Ala	Leu
225				230					235					240	
Ile	Asp	Phe	His	Thr	Leu	Thr	Pro	Gly	Asp	Pro	Asn	Val	Asn	Leu	Asp
			245					250						255	
Arg	Ala	Lys	Thr	Phe	Phe	Ala	Ser	Val	Ala	Ala	Arg	Asn	Ala	Gly	Lys
			260					265						270	
Lys	Asn	Val	Ile	Tyr	Glu	Ile	Ala	Asn	Glu	Pro	Asn	Gly	Val	Thr	Trp
		275					280					285			
Thr	Ala	Val	Lys	Ser	Tyr	Ala	Glu	Gln	Val	Ile	Pro	Val	Ile	Arg	Ala
	290					295					300				

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Ala Asp Pro Asp Ala Val Val Ile Val Gly Thr Arg Gly Trp Ser Ser
305 310 315 320

Leu Gly Val Ser Asp Gly Ser Asp Glu Ser Glu Val Val Asn Ser Pro
325 330 335

Val Asn Ala Thr Asn Ile Met Tyr Ala Phe His Phe Tyr Ala Ala Ser
340 345 350

His Lys Asp Ala Tyr Arg Ser Thr Leu Ser Arg Ala Ala Ala Arg Leu
355 360 365

Pro Leu Phe Val Thr Glu Phe Gly Thr Val Ser Ala Thr Gly Gly Gly
370 375 380

Ala Met Asp Arg Ala Ser Thr Thr Ala Trp Leu Asp Leu Leu Asp Gln
385 390 395 400

Leu Lys Ile Ser Tyr Ala Asn Trp Thr Tyr Ser Asp Ala Pro Glu Ser
405 410 415

Ser Ala Ala Phe Arg Pro Gly Thr Cys Gly Gly Gly Asp Tyr Ser Gly
420 425 430

Ser Gly Val Leu Thr Glu Ser Gly Ala Leu Leu Lys Asn Arg Ile Ser
435 440 445

Thr Pro Asp Ser Phe Pro Thr Gly
450 455

<210> SEQ ID NO 5
 <211> LENGTH: 458
 <212> TYPE: PRT
 <213> ORGANISM: Streptomyces sp. ACTE

<400> SEQUENCE: 5

Met Ala Lys Lys Ile Pro Ala Arg Ala Arg Ala Leu Ser Val Leu
1 5 10 15

Thr Ala Gly Val Leu Ala Ala Ala Gly Val Val Ser Leu Ala Gly Thr
20 25 30

Ala Glu Ala Ala Gly Thr Leu Gly Asp Ala Ala Ala Ala Lys Gly Arg
35 40 45

Tyr Phe Gly Thr Ala Val Ala Ala Asn His Leu Gly Glu Ala Pro Tyr
50 55 60

Ala Ser Thr Leu Asp Ala Gln Phe Asp Ser Val Thr Pro Glu Asn Glu
65 70 75 80

Met Lys Trp Asp Ala Val Glu Gly Ser Arg Asn Ser Phe Thr Phe Thr
85 90 95

Ala Ala Asp Gln Ile Val Ser His Ala Gln Ser Lys Gly Met Lys Val
100 105 110

Arg Gly His Thr Leu Val Trp His Ser Gln Leu Pro Gly Trp Val Gly
115 120 125

Gly Leu Gly Ala Thr Asp Leu Arg Ala Ala Met Asn Asn His Ile Thr
130 135 140

Gln Val Met Thr His Tyr Lys Gly Lys Ile His Ser Trp Asp Val Val
145 150 155 160

Asn Glu Ala Phe Gln Asp Gly Asn Ser Gly Ala Arg Arg Ser Ser Pro
165 170 175

Phe Gln Asp Lys Leu Gly Asp Gly Phe Ile Glu Glu Ala Phe Arg Thr
180 185 190

Ala Arg Thr Val Asp Pro Thr Ala Lys Leu Cys Tyr Asn Asp Tyr Asn
195 200 205

Thr Asp Gly Arg Asn Ala Lys Ser Asp Ala Val Tyr Ala Met Ala Lys
210 215 220

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Asp Phe Lys Gln Arg Gly Val Pro Ile Asp Cys Val Gly Phe Gln Ser
 225 230 235 240
 His Phe Asn Ser Asn Ser Pro Val Pro Ser Asp Tyr Arg Ala Asn Leu
 245 250 255
 Gln Arg Phe Ala Asp Leu Gly Leu Asp Val Gln Ile Thr Glu Leu Asp
 260 265 270
 Ile Glu Gly Ser Gly Ser Ala Gln Ala Ala Asn Tyr Thr Ser Val Val
 275 280 285
 Asn Ala Cys Leu Ala Val Thr Arg Cys Thr Gly Leu Thr Val Trp Gly
 290 295 300
 Val Thr Asp Lys Tyr Ser Trp Arg Ser Ser Gly Thr Pro Leu Leu Phe
 305 310 315 320
 Asp Gly Asp Tyr Asn Lys Lys Pro Ala Tyr Asp Ala Val Leu Ala Ala
 325 330 335
 Leu Gly Gly Thr Pro Asp Gly Gly Gly Asp Asp Gly Gly Gly Asp Asn
 340 345 350
 Gly Gly Gly Asn Thr Gly Ser Cys Thr Ala Thr Tyr Thr Gln Thr Ala
 355 360 365
 Thr Trp Asn Gly Gly Tyr Asn Gly Glu Val Thr Val Lys Ala Gly Ser
 370 375 380
 Ser Gly Ile Thr Thr Trp Ser Val Pro Val Thr Val Pro Ser Ser Gln
 385 390 395 400
 Gln Val Ser Ala Leu Trp Asn Gly Ala Pro Thr Trp Asn Ala Gly Asn
 405 410 415
 Thr Val Met Thr Val Lys Pro Thr Tyr Asn Gly Thr Leu Ala Ala Gly
 420 425 430
 Ala Ser Thr Ser Phe Gly Phe Thr Val Met Thr Asn Gly Asn Thr Ser
 435 440 445
 Ala Pro Ala Val Gly Ala Cys Thr Ala Ser
 450 455

<210> SEQ ID NO 6

<211> LENGTH: 562

<212> TYPE: PRT

<213> ORGANISM: Streptomyces sp. ACTE

<400> SEQUENCE: 6

Val Arg Thr Ala Ile Arg Thr Ala Arg Arg Pro Gln Pro Leu Ala Leu
 1 5 10 15
 Leu Leu Arg Gly Leu Ala Ala Phe Leu Gly Leu Ala Leu Ala Gly Ala
 20 25 30
 Leu Gly Pro Ala Thr Ala Arg Ala Ala Asp Leu Pro Gln Arg Ala Glu
 35 40 45
 Ala Arg Ala Ala Gly Leu His Ile Ser Asp Gly Arg Leu Val Glu Gly
 50 55 60
 Asn Gly Asn Asp Phe Val Met Arg Gly Ile Asn His Ala His Thr Trp
 65 70 75 80
 Tyr Pro Gly Glu Thr Gln Ser Leu Ala Asp Ile Lys Ala Thr Gly Ala
 85 90 95
 Asn Thr Val Arg Val Val Leu Ser Asp Gly Tyr Arg Trp Ser Glu Asn
 100 105 110
 Ser Pro Glu Asp Val Ala Ser Ile Ile Ala Arg Cys Lys Ala Glu Arg
 115 120 125
 Leu Ile Cys Val Leu Glu Val His Asp Thr Thr Gly Tyr Gly Glu Asp

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130			135			140									
Ala	Ala	Ala	Gly	Thr	Leu	Asp	His	Ala	Ala	Asp	Tyr	Trp	Ile	Gly	Leu
145					150					155					160
Lys	Asp	Val	Leu	Asp	Gly	Glu	Glu	Asp	Tyr	Val	Val	Ile	Asn	Ile	Gly
			165						170						175
Asn	Glu	Pro	Trp	Gly	Asn	Ala	Asp	Pro	Ala	Gly	Trp	Thr	Ala	Pro	Thr
			180					185						190	
Thr	Ala	Ala	Ile	Gln	Lys	Leu	Arg	Ala	Ala	Gly	Phe	Ala	His	Thr	Ile
		195					200						205		
Met	Val	Asp	Ala	Pro	Asn	Trp	Gly	Gln	Asp	Trp	Glu	Gly	Val	Met	Arg
210						215					220				
Ala	Asp	Ala	Arg	Ser	Val	Tyr	Asp	Ala	Asp	Pro	Thr	Gly	Asn	Leu	Ile
225					230					235					240
Phe	Ser	Ile	His	Met	Tyr	Ser	Val	Tyr	Asp	Thr	Ala	Ala	Lys	Val	Thr
				245					250						255
Asp	Tyr	Leu	Asn	Ala	Phe	Val	Asp	Ala	Gly	Leu	Pro	Leu	Leu	Ile	Gly
			260					265						270	
Glu	Phe	Gly	Gly	Pro	Ala	Asp	Gln	Tyr	Gly	Asp	Pro	Asp	Glu	Asp	Thr
		275					280						285		
Met	Met	Ala	Thr	Ala	Glu	Glu	Leu	Gly	Leu	Gly	Tyr	Leu	Ala	Trp	Ser
290						295					300				
Trp	Ser	Gly	Asn	Thr	Asp	Pro	Val	Leu	Asp	Leu	Val	Leu	Asp	Phe	Asp
305					310					315					320
Pro	Thr	Arg	Leu	Ser	Ser	Trp	Gly	Glu	Arg	Val	Leu	His	Gly	Pro	Asp
				325					330						335
Gly	Ile	Thr	Glu	Thr	Ser	Arg	Glu	Ala	Thr	Val	Phe	Gly	Gly	Gly	Gln
			340					345							350
Gly	Gly	Gly	Asp	Thr	Glu	Ala	Pro	Thr	Ala	Pro	Gly	Thr	Pro	Thr	Ala
		355					360						365		
Ser	Gly	Val	Thr	Ala	Thr	Ser	Val	Thr	Leu	Gly	Trp	Ser	Ala	Ala	Thr
370						375					380				
Asp	Asp	Val	Gly	Val	Thr	Ala	Tyr	Asp	Val	Val	Arg	Val	Thr	Gly	Gly
385					390					395					400
Ser	Glu	Thr	Lys	Val	Ala	Ser	Ser	Ala	Ala	Thr	Ser	Val	Thr	Val	Thr
				405					410						415
Gly	Leu	Ser	Ala	Gly	Thr	Ala	Tyr	Ser	Phe	Ala	Val	Tyr	Ala	Arg	Asp
			420					425						430	
Ala	Ala	Gly	Asn	Arg	Ser	Ala	Arg	Ser	Gly	Thr	Val	Ser	Val	Thr	Thr
		435					440							445	
Asp	Glu	Gly	Gly	Ser	Val	Pro	Gly	Gly	Ala	Cys	Ser	Val	Gly	Tyr	Arg
450						455					460				
Val	Ile	Gly	Glu	Trp	Pro	Gly	Gly	Phe	Gln	Gly	Glu	Ile	Thr	Leu	Arg
465					470					475					480
Asn	Thr	Gly	Ala	Ala	Ala	Val	Asp	Gly	Trp	Thr	Leu	Gly	Phe	Ala	Phe
				485					490						495
Ala	Asp	Gly	Gln	Thr	Val	Thr	Asn	Met	Trp	Gly	Gly	Thr	Ala	Thr	Gln
			500					505						510	
Ser	Gly	Gly	Ala	Val	Ser	Val	Thr	Pro	Ala	Ser	Tyr	Thr	Ser	Thr	Ile
		515						520						525	
Ala	Ala	Gly	Gly	Ser	Val	Thr	Val	Gly	Phe	Thr	Gly	Thr	Leu	Thr	Gly
		530						535					540		
Ala	Asn	Ala	Ala	Pro	Ala	Ala	Phe	Thr	Leu	Asn	Gly	Ala	Thr	Cys	Thr
545					550					555					560

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Ala Ala

<210> SEQ ID NO 7
 <211> LENGTH: 328
 <212> TYPE: PRT
 <213> ORGANISM: Streptomyces sp. ACTE

<400> SEQUENCE: 7

Met Ser Ile Thr Pro Arg Pro Ser Leu Arg Ala Met Val Thr Gly Leu
 1 5 10 15
 Ala Val Ala Ala Ser Ala Leu Ala Gly Gly Ala Val Thr Ala Ala Pro
 20 25 30
 Ala Arg Ala Ala Ala Cys Asn Gly Tyr Val Gly Leu Thr Phe Asp Asp
 35 40 45
 Gly Pro Ser Ala Ala Gln Thr Pro Ala Leu Leu Ser Ala Leu Lys Gln
 50 55 60
 Asn Gly Leu Arg Ala Thr Met Phe Asn Gln Gly Asn Tyr Ala Ala Ser
 65 70 75 80
 Asn Pro Ala Gln Val Lys Ala Gln Val Asp Ala Gly Met Trp Val Gly
 85 90 95
 Asn His Ser Tyr Ser His Pro His Leu Thr Gln Gln Ser Gln Ala Gln
 100 105 110
 Met Asp Ser Glu Ile Ser Arg Thr Gln Gln Ala Ile Ala Ala Gly Gly
 115 120 125
 Gly Gly Thr Pro Lys Leu Phe Arg Pro Pro Tyr Gly Glu Thr Asn Ala
 130 135 140
 Thr Leu Arg Ser Val Glu Ala Lys Tyr Gly Leu Thr Glu Val Ile Trp
 145 150 155 160
 Asp Val Asp Ser Gln Asp Trp Asn Gly Ala Ser Thr Asp Ala Ile Val
 165 170 175
 Gln Ala Val Ser Arg Leu Thr Ala Gly Gln Val Ile Leu Met His Glu
 180 185 190
 Trp Pro Ala Asn Thr Leu Ala Ala Ile Pro Arg Ile Ala Gln Thr Leu
 195 200 205
 Ser Ala Lys Gly Leu Cys Ser Gly Met Ile Ser Pro Gln Thr Gly Arg
 210 215 220
 Ala Val Ala Pro Asp Gly Gly Gly Asn Gly Gly Gly Gly Gly Gly
 225 230 235 240
 Gly Gly Cys Thr Ala Thr Leu Ser Ala Gly Glu Lys Trp Gly Asp Arg
 245 250 255
 Tyr Asn Leu Asn Val Ala Val Ser Gly Ser Ser Asn Trp Thr Val Thr
 260 265 270
 Met Asn Val Pro Ser Gly Glu Arg Val Met Thr Thr Trp Asn Val Ser
 275 280 285
 Ala Ser Tyr Pro Ser Ala Gln Val Leu Val Ala Lys Pro Asn Gly Ser
 290 295 300
 Gly Asn Asn Trp Gly Ala Thr Ile Gln Ala Asn Gly Asn Trp Thr Trp
 305 310 315 320
 Pro Thr Val Ser Cys Thr Thr Ser
 325

<210> SEQ ID NO 8
 <211> LENGTH: 335
 <212> TYPE: PRT
 <213> ORGANISM: Streptomyces sp. ACTE

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<400> SEQUENCE: 8

Met Asn Pro Leu Val Tyr Thr Glu Arg Arg Arg Arg Gly Arg Leu Thr
 1 5 10 15
 Ser Leu Ala Gly Ser Val Cys Ala Leu Val Leu Ala Ala Ala Ala
 20 25 30
 Met Leu Leu Pro Gly Thr Ala Ser Ala Asp Thr Val Val Thr Thr Asn
 35 40 45
 Gln Thr Gly Asn Asn Asn Gly Tyr Tyr Tyr Ser Phe Trp Thr Asp Gly
 50 55 60
 Gly Gly Gln Val Ser Met Asn Leu Ala Ser Gly Gly Ser Tyr Ser Thr
 65 70 75 80
 Ser Trp Thr Asn Thr Gly Asn Phe Val Ala Gly Lys Gly Trp Ser Thr
 85 90 95
 Gly Gly Arg Lys Ser Val Thr Tyr Ser Gly Thr Phe Asn Pro Ser Gly
 100 105 110
 Asn Ala Tyr Leu Thr Leu Tyr Gly Trp Ser Thr Asn Pro Leu Val Glu
 115 120 125
 Tyr Tyr Ile Val Asp Asn Trp Gly Thr Tyr Arg Pro Thr Gly Thr Phe
 130 135 140
 Lys Gly Thr Val Ser Ser Asp Gly Gly Thr Tyr Asp Ile Tyr Glu Thr
 145 150 155 160
 Thr Arg Thr Asn Ala Pro Ser Ile Glu Gly Thr Lys Thr Phe Lys Gln
 165 170 175
 Phe Trp Ser Val Arg Gln Ser Lys Arg Thr Gly Gly Thr Ile Thr Thr
 180 185 190
 Gly Asn His Phe Asp Ala Trp Ala Arg Asn Gly Met Asn Leu Gly Thr
 195 200 205
 Met Asn Tyr Met Ile Leu Ala Thr Glu Gly Tyr Gln Ser Ser Gly Ser
 210 215 220
 Ser Asn Ile Thr Val Ser Glu Gly Gly Ser Gly Gly Gly Asp Asn
 225 230 235 240
 Gly Gly Gly Gly Gly Gly Gly Gly Gly Cys Thr Ala Thr Leu Ser Ala
 245 250 255
 Gly Glu Lys Trp Gly Asp Arg Tyr Asn Leu Asn Val Ala Val Ser Gly
 260 265 270
 Ser Ser Asn Trp Thr Val Thr Met Asn Val Pro Ser Ala Glu Lys Val
 275 280 285
 Leu Ser Thr Trp Asn Ile Ser Ala Ser Tyr Pro Ser Ser Gln Val Leu
 290 295 300
 Val Ala Lys Pro Asn Gly Ser Gly Asn Asn Trp Gly Ala Thr Ile Gln
 305 310 315 320
 Ala Asn Gly Asn Trp Thr Trp Pro Thr Val Ser Cys Thr Thr Ser
 325 330 335

<210> SEQ ID NO 9

<211> LENGTH: 280

<212> TYPE: PRT

<213> ORGANISM: Streptomyces sp. ACTE

<400> SEQUENCE: 9

Met Ser Glu Arg Ala Ala Ser Pro Arg Thr His Arg Arg Arg Pro Gly
 1 5 10 15
 Arg Arg Arg Ile Ala Thr Ala Leu Thr Ala Ala Leu Gly Leu Thr Gly
 20 25 30

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Ala Ala Leu Ala Thr Gly Val Met Leu Gln Pro Ala Gly Ala Ala Thr
 35 40 45
 Thr Ala Ile Pro Ala Trp Pro Ser Ala Thr Gly Ser Gln Ser Val Ser
 50 55 60
 Lys Thr Ile Glu Val Ser Gly Thr Tyr Asp Gly Gly Leu Lys Arg Phe
 65 70 75 80
 Thr Gly Ser Gly Asp Leu Gly Asp Gly Gly Gln Asp Glu Gly Gln Asp
 85 90 95
 Pro Ile Phe Lys Leu Lys Asp Gly Ala Thr Ile Lys Asn Val Ile Leu
 100 105 110
 Gly Thr Pro Ala Ala Asp Gly Ile His Cys Ser Gly Ser Cys Thr Ile
 115 120 125
 Gln Asn Val Trp Trp Glu Asp Val Gly Glu Asp Ala Ala Ser Phe Lys
 130 135 140
 Gly Thr Ser Thr Ser Ser Val Tyr Thr Val Tyr Gly Gly Gly Ala Lys
 145 150 155 160
 Lys Ala Ser Asp Lys Val Phe Gln Phe Asn Gly Ala Gly Lys Leu Val
 165 170 175
 Val Thr Lys Phe Gln Val Ala Asp Phe Gly Lys Leu Val Arg Ser Cys
 180 185 190
 Gly Asn Cys Ser Lys Gln Tyr Lys Arg Glu Ile Ile Val Asn Asp Val
 195 200 205
 Asp Val Thr Ala Pro Gly Lys Ser Leu Val Gly Ile Asn Thr Asn Tyr
 210 215 220
 Gly Asp Thr Ala Ala Leu Arg Ser Val Arg Val His Gly Asp Ser Ser
 225 230 235 240
 Lys Lys Ile Lys Pro Cys Val Arg Tyr Thr Gly Asn Ser Thr Gly Ala
 245 250 255
 Glu Pro Lys Glu Thr Gly Ser Gly Pro Asp Gly Thr Tyr Cys Lys Tyr
 260 265 270
 Thr Ala Ser Asp Leu Ser Tyr Asp
 275 280

<210> SEQ ID NO 10
 <211> LENGTH: 909
 <212> TYPE: PRT
 <213> ORGANISM: Streptomyces sp. ACTE

<400> SEQUENCE: 10

Met Trp Cys His Pro Tyr Leu Arg Leu Arg Thr Ser Gly Arg Lys Val
 1 5 10 15
 Ser Ser Val Asn Ala Leu Pro Pro Pro Ala Arg Pro Ala Pro Val Arg
 20 25 30
 Pro Arg Ser Arg Tyr Gly Arg Arg Val Leu Gly Met Ser Ala Ala Ala
 35 40 45
 Leu Leu Cys Ala Gly Ala Leu Ala Val Pro Gly Thr Ala Met Ala Asp
 50 55 60
 Asp Ala Glu Pro Gly Pro Gly Pro Glu Gln Ile Thr Asn Gly Asp Phe
 65 70 75 80
 Ala Thr Gly Thr Ser Ala Pro Trp Trp Trp Thr Pro Asn Ala Ser Ala
 85 90 95
 Ala Val Ser Glu Gly Arg Leu Cys Val Glu Val Pro Ala Gly Thr Ala
 100 105 110
 Asn Ala Trp Asp Val Ile Val Gly Gln Asn Asp Val Pro Ile Val Ala

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115					120					125					
Gly	Glu	Ser	Tyr	Glu	Leu	Ser	Tyr	Thr	Ala	Arg	Ser	Thr	Val	Pro	Leu
130					135					140					
Thr	Val	Gln	Thr	Arg	Val	Gln	Glu	Ala	Val	Glu	Pro	Tyr	Thr	Thr	Val
145					150					155					160
Leu	Ala	Thr	Ala	Asp	Pro	Val	Gly	Ala	Glu	Asp	Thr	Arg	Val	Ala	Arg
				165					170					175	
Thr	Phe	Thr	Ala	Ser	Val	Asp	Gln	Pro	Ala	Ala	Ser	Val	Gln	Leu	Gln
				180					185					190	
Ile	Gly	Gly	Gly	Glu	Arg	Ala	Thr	Thr	Phe	Cys	Leu	Asp	Asp	Val	Ser
				195					200					205	
Leu	Arg	Gly	Gly	Ala	Glu	Pro	Pro	Val	Tyr	Val	Pro	Asp	Thr	Gly	Ser
				210					215					220	
Pro	Val	Arg	Val	Asn	Gln	Val	Gly	Tyr	Leu	Pro	Arg	Gly	Pro	Lys	Ser
				225					230					235	
Gly	Thr	Val	Val	Thr	Asp	Ala	Glu	Ala	Pro	Leu	Thr	Trp	Thr	Val	Lys
				245					250					255	
Ala	Glu	Asp	Gly	Ser	Thr	Ala	Ala	Thr	Gly	Thr	Thr	Val	Pro	Arg	Gly
				260					265					270	
Glu	Asp	Pro	Ser	Ser	Arg	Arg	Arg	Val	His	Thr	Phe	Asp	Phe	Gly	Asp
				275					280					285	
Leu	Thr	Thr	Ala	Gly	Asp	Gly	Tyr	Thr	Val	Glu	Val	Asp	Gly	Glu	Val
				290					295					300	
Ser	Glu	Pro	Phe	Ser	Ile	Arg	Gly	Asp	Leu	Tyr	Asp	Ser	Leu	Arg	Ser
				305					310					315	
Asp	Ala	Leu	Ala	Tyr	Phe	Tyr	His	Asn	Arg	Ser	Gly	Ile	Glu	Ile	Asp
				325					330					335	
Ala	Asp	Leu	Val	Gly	Glu	Gln	Tyr	Ala	Arg	Pro	Ala	Gly	His	Ile	Gly
				340					345					350	
Val	Ala	Pro	Asn	Lys	Gly	Asp	Thr	Asp	Val	Pro	Cys	Arg	Pro	Gly	Val
				355					360					365	
Cys	Asp	Tyr	Arg	Leu	Asp	Val	Ser	Gly	Gly	Trp	Tyr	Asp	Ala	Gly	Asp
				370					375					380	
His	Gly	Lys	Tyr	Val	Val	Asn	Gly	Gly	Ile	Ser	Val	Ala	Gln	Leu	Met
				385					390					395	
Ala	Thr	Tyr	Glu	Arg	Thr	Leu	Thr	Ala	Pro	Asp	Ala	Glu	Ser	Ala	Glu
				405					410					415	
Leu	Gly	Asp	Gly	Ala	Leu	Arg	Val	Pro	Glu	Arg	Asp	Asn	Gly	Val	Pro
				420					425					430	
Asp	Ile	Leu	Asp	Glu	Ala	Arg	Trp	Glu	Met	Asp	Phe	Leu	Ile	Lys	Met
				435					440					445	
Gln	Val	Pro	Ala	Gly	Glu	Gln	Leu	Ala	Gly	Met	Val	His	His	Lys	Met
				450					455					460	
His	Asp	Ala	Glu	Trp	Thr	Gly	Leu	Pro	Met	Lys	Pro	His	Leu	Asp	Pro
				465					470					475	
Gln	Gln	Arg	Glu	Leu	His	Pro	Pro	Ser	Thr	Ala	Ala	Thr	Leu	Asn	Leu
				485					490					495	
Ala	Ala	Thr	Ala	Ala	Gln	Cys	Ala	Arg	Leu	Tyr	Ala	Pro	Phe	Asp	Ala
				500					505					510	
Asp	Phe	Ala	Asp	Arg	Cys	Leu	Arg	Ala	Ala	Glu	Thr	Ala	Trp	Asp	Ala
				515					520					525	
Ala	Lys	Arg	His	Pro	Asp	Val	Leu	Ala	Asp	Pro	Asn	Asp	Gly	Ile	Gly
				530					535					540	

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Gly Gly Ala Tyr Asn Asp Asp Asp Val Ser Asp Glu Phe Tyr Trp Ala
 545 550 555 560
 Ala Ala Glu Leu Phe Thr Thr Thr Gly Lys Asp Ile Tyr Arg Gln Ala
 565 570 575
 Val Leu Ser Ser Ala Trp His Gly Asp Ala Gly Ala Val Phe Pro Ala
 580 585 590
 Gly Gly Gly Ile Ser Trp Gly Ser Thr Ala Gly Leu Gly Val Leu Thr
 595 600 605
 Leu Ala Thr Val Pro Asn Ala Leu Thr Ser Asp Gln Leu Ala Gln Val
 610 615 620
 Arg Thr Val Val Thr Glu Gly Ala Asp Arg Tyr Ala Ala Gln Ser Arg
 625 630 635 640
 Glu Gln Ala Tyr Gly Leu Pro Tyr Ala Pro Arg Gly Glu Asp Tyr Val
 645 650 655
 Trp Gly Ser Asn Ser Gln Val Leu Asn Asn Met Val Val Leu Ala Thr
 660 665 670
 Ala His Asp Leu Thr Gly Asp Ala Ala Tyr Gln Asp Ala Val Leu Arg
 675 680 685
 Gly Ala Asp Tyr Leu Leu Gly Arg Asn Pro Leu Asn Gln Ser Tyr Val
 690 695 700
 Thr Gly Tyr Gly Glu Arg Asp Ser His Asn Gln His His Arg Phe Trp
 705 710 715 720
 Ala His Gln Asn Asp Pro Ser Leu Pro Asn Pro Ala Pro Gly Ser Ile
 725 730 735
 Ala Gly Gly Pro Asn Leu Thr Ala Ile Ala Ser Gly Asp Pro Val Ala
 740 745 750
 Ala Glu Lys Leu Ser Gly Cys Ala Pro Ala Met Cys Tyr Val Asp Asp
 755 760 765
 Ile Gly Ser Trp Ala Thr Asn Glu Ile Thr Ile Asn Trp Asn Ala Pro
 770 775 780
 Leu Ala Phe Ile Ala Ser Tyr Leu Asp Asp Ala Gly Glu Gly Gly Gln
 785 790 795 800
 Thr Ala Ala Ala Arg Thr Cys Gln Val Thr Tyr Ser Ser His Pro Trp
 805 810 815
 Asn Ser Gly Ser Thr Val Thr Val Arg Val Glu Asn Thr Gly Ser Asp
 820 825 830
 Pro Val Ser Pro Trp Ala Leu Thr Trp Leu Leu Pro Gly Glu Gln Arg
 835 840 845
 Leu Ser His Thr Trp Ser Ala Glu Phe Asp Gln His Gly Arg Thr Val
 850 855 860
 Ser Ala Arg Pro Leu Ser Trp Asn Arg Thr Leu Ala Pro Gly Ala Ala
 865 870 875 880
 Val Asp Phe Gly Phe Asn Thr Ser Ala Ala Gly Ser Ser Pro Glu Pro
 885 890 895
 Gly Ala Phe Lys Leu Asn Gly Arg Ala Cys Ser Ala Gly
 900 905

<210> SEQ ID NO 11

<211> LENGTH: 405

<212> TYPE: PRT

<213> ORGANISM: Streptomyces sp. ACTE

<400> SEQUENCE: 11

Met Arg Thr Gly Ser Ile Ala Arg Val Leu Gly Leu Ala Ala Ala Leu

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1	5	10	15
Ala Ala Leu	Leu Thr Thr	Ala Phe Met	Ala Pro Ala Met Ala Gly Lys
	20	25	30
His Asp Ala	Thr Asp Ser Pro	Ser Ala Ala Ala	Ala Pro Ala Ser Phe
	35	40	45
Thr His Pro	Gly Val Leu	Val Ser Arg Pro	Gln Leu Asp Phe Val Arg
	50	55	60
Gly Lys Val	Gln Ala Gly	Ala Gln Pro Trp	Lys Gly Ala Tyr Asp Gln
	65	70	75
Met Leu Ala	Ser Pro Tyr Ala	Ser Leu Ser Arg	Thr Ala Lys Pro Arg
	85	90	95
Ala Val Val	Glu Cys Gly	Ser Tyr Ser Asn	Pro Asn Asn Gly Cys Thr
	100	105	110
Asp Glu Arg	Glu Asp Ala	Leu Ala Ala Tyr	Thr Leu Ser Leu Ala Trp
	115	120	125
Tyr Ile Ser	Gln Asp Gly	Arg Tyr Ala Gln	Lys Ala Ile Gln Ile Met
	130	135	140
Asp Ala Trp	Ser Gly Val	Ile Lys Asp His	Thr Asn Ser Asn Ala Pro
	145	150	155
Leu Gln Thr	Gly Trp Ala	Gly Ser Ser Trp	Pro Arg Ala Ala Glu Ile
	165	170	175
Ile Lys Tyr	Thr Tyr Gly	Asn Trp Pro Ala	Ser Gly Arg Phe Gly Thr
	180	185	190
Met Leu Arg	Asp Val Tyr	Leu Pro Lys Val	Ala Asn Gly Ser Asn Ser
	195	200	205
Asn Gly Asn	Trp Glu Leu	Ser Met Thr Glu	Ala Ala Ile Gly Ile Ala
	210	215	220
Val Phe Leu	Glu Asp Arg	Gly Ala Tyr Asp	Arg Ala Val Ala Lys Phe
	225	230	235
Arg Gly Arg	Val Pro Ala	Tyr Ile Tyr Val	Thr Ala Asp Gly Ser Leu
	245	250	255
Pro Lys Ala	Ala Pro Gly	Ser Gly Leu Asp	Thr Arg Glu Lys Ile Ile
	260	265	270
Asn Tyr Trp	Gln Gly Gln	Ser Thr Phe Val	Asp Gly Leu Ser Gln Glu
	275	280	285
Thr Cys Arg	Asp Leu Thr	His Thr Gly Tyr	Gly Leu Ser Ala Ile Ser
	290	295	300
His Ile Ala	Glu Thr Ser	Arg Ile Gln Gly	Gln Asp Leu Tyr Pro Glu
	305	310	315
Val Ala Asp	Arg Leu Arg	His Ala Leu Gly	Leu His Ala Lys Tyr Gln
	325	330	335
Leu Gly Glu	Lys Val Pro	Ser Ser Leu Cys	Gly Gly Ser Leu Lys Asp
	340	345	350
Ser Leu Gly	Pro Val Thr	Glu Val Gly Phe	Asn Ala Leu His Asn Arg
	355	360	365
Met Gly Tyr	Ala Met Thr	Asn Thr Gln Thr	Leu Thr Glu Arg Gln Arg
	370	375	380
Pro Ala Ala	Ser Asn Asn	Leu Phe Val Ala	Trp Glu Thr Leu Thr His
	385	390	395
Ala Asp Asn	Pro Asn		
	405		

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<211> LENGTH: 626

<212> TYPE: PRT

<213> ORGANISM: Streptomyces sp. ACTE

<400> SEQUENCE: 12

Met Pro Ser Arg Thr Thr Leu Ile Ala Thr Thr Ala Ala Leu Val Ala
 1 5 10 15

Leu Ala Ala Pro Met Ala Phe Ala Ala Pro Ala Pro Ala Pro Asp Pro
 20 25 30

Ala Val Glu Ala Ala Ala Ala Ala Trp Asp Thr Asp Arg Ala Ala Ser
 35 40 45

Ala Tyr Ala Ala Asn Pro Ala Ala Val Thr Ala Ser Gly Ser Glu Asn
 50 55 60

Pro Ala Ser Gly Pro Gly Ala Ala Thr Asp Gly Asp Ala Thr Thr Arg
 65 70 75 80

Trp Ser Ser Asp Phe Ala Asp Asn Ala Trp Ile Arg Val Asp Leu Gly
 85 90 95

Ser Thr Ile Arg Ile Asn Gln Val Lys Leu Glu Trp Glu Ala Ala Tyr
 100 105 110

Gly Lys Lys Tyr Val Leu Glu Val Ser Lys Asp Gly Thr Asn Trp Thr
 115 120 125

Pro Phe Tyr Thr Glu Asp Ala Gly Thr Gly Gly Thr Val Thr Ala His
 130 135 140

Thr Tyr Pro Gln Glu Val Thr Gly Arg Tyr Val Arg Met Arg Gly Val
 145 150 155 160

Glu Arg Ala Thr Ala Trp Gly Tyr Ser Leu Phe Ser Phe Gln Val Tyr
 165 170 175

Gly Gly Glu Pro Ala Pro Ala Ser Thr Thr Arg Ser Asn Leu Ala Leu
 180 185 190

Asn His Pro Ala Tyr Gly Asp Leu Tyr Gln His Ala Gly Asn Ser Pro
 195 200 205

Ala Phe Val Thr Asp Gly Gly Trp Pro Ala Asp Leu Lys Ala Asp Arg
 210 215 220

Ser Arg Trp Ser Ser Asp Trp Asn Ala Asp Arg Trp Val Gly Val Asp
 225 230 235 240

Leu Gly Ala Thr Ser Thr Ile Asn Ser Val Asp Leu Tyr Trp Glu Ala
 245 250 255

Ala Tyr Ala Val Asp Tyr Glu Ile Gln Val Ser Asp Asp Asn Arg Thr
 260 265 270

Trp Arg Thr Val His Arg Pro Ser Ala Ala Glu Val Ala Ala Arg Arg
 275 280 285

Ala Asp Val Lys Ala Pro Ala Glu Ala Val Gly Arg His Asp Thr Ile
 290 295 300

Asn Leu Pro Thr Pro Ala Thr Gly Arg Tyr Val Arg Met Leu Gly Lys
 305 310 315 320

Glu Arg Arg Ser Phe Tyr Asn Pro Ala Pro Ser Thr Ala Gln Phe Gly
 325 330 335

Tyr Ser Leu Tyr Glu Phe Gln Val Trp Gly Thr Gly Gly Ser Ala Asp
 340 345 350

Ala Ala Tyr Pro Ala Leu Pro Lys Asn Pro Gly Gly Ala Tyr Arg Thr
 355 360 365

Thr Phe Phe Asp Asp Phe Thr Gly Ser Gly Leu Asp Arg Ser Lys Trp
 370 375 380

Arg Val Val Arg Thr Gly Thr Glu Met Gly Pro Val Asn Gly Glu Ser

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385	390	395	400
Gln Ala Tyr Val Asp Ser Pro Asp Asn Ile Arg Thr Glu Asn Gly Ala			
	405	410	415
Leu Val Leu Glu Ser Lys Tyr Cys Lys Gly Cys Thr Pro Thr Pro Asn			
	420	425	430
Gly Thr Phe Asp Phe Thr Ser Gly Arg Val Asp Thr Asn Thr Lys Phe			
	435	440	445
Asp Phe Thr Tyr Gly Lys Val Ser Ala Arg Met Lys Leu Pro Val Gly			
	450	455	460
Asp Gly Phe Trp Pro Ala Phe Trp Leu Leu Gly Ser Asp Val Asp Asp			
	465	470	475
Pro Ala Val Ser Trp Pro Gly Ser Gly Glu Thr Asp Ile Met Glu Asn			
	485	490	495
Ile Gly Tyr Gly Asp Trp Thr Ser Ser Gly Leu His Gly Pro Gly Tyr			
	500	505	510
Ser Ala Asp Gly Asn Ile Gly Ala Ser Gln Thr Tyr Pro Asn Gly Gly			
	515	520	525
Arg Ala Asp Glu Trp His Thr Tyr Gly Val Glu Trp Thr Pro Glu Gly			
	530	535	540
Met Thr Phe Thr Val Asp Asp Arg Val Val Gln Gln Thr Ser Arg Gln			
	545	550	555
Lys Leu Glu Ser Thr Arg Gly Lys Trp Val Phe Asp His Asn Gln Tyr			
	565	570	575
Val Ile Leu Asn Leu Ala Leu Gly Gly Ala Tyr Pro Gly Gly Tyr Asn			
	580	585	590
Gln Val Thr Gln Pro Tyr Trp Gly Leu Pro Gln Ser Ser Val Asp Arg			
	595	600	605
Ile Ala Gln Gly Gly Ile Lys Ala Glu Ile Asp Trp Val Arg Val Glu			
	610	615	620
Gln Lys			
625			
<210> SEQ ID NO 13			
<211> LENGTH: 408			
<212> TYPE: PRT			
<213> ORGANISM: Streptomyces sp. ACTE			
<400> SEQUENCE: 13			
Val Ile Ser Arg Arg Met Phe Leu Thr Gly Ala Ala Ala Ser Ala Thr			
1	5	10	15
Ala Leu Thr Tyr Pro Leu Trp Gly Thr Ala Leu Ser Pro Arg Thr Ser			
	20	25	30
Ala Ala Ala Ala Thr Cys Glu Leu Ala Leu Glu Asn Arg Ser Leu Pro			
	35	40	45
Gly Thr Val His Ala Tyr Val Thr Gly His Glu Gln Gly Thr Asp Ser			
	50	55	60
Trp Val Leu Leu Arg Ala Asp Gly Ser Val Tyr Arg Pro Glu Ser Pro			
	65	70	75
Gly Ala Pro Gln Thr Pro Leu Pro Val Asp Cys Ala Ile Pro Leu Asn			
	85	90	95
Gly Ala Gly Ala Gly Pro Val Val Leu Thr Leu Pro Gln Met Tyr Gly			
	100	105	110
Ala Arg Val Tyr Phe Val Arg Asp Asp Lys Leu Asp Phe Tyr Leu Asn			
	115	120	125

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Pro Gly Pro Ser Leu Val Glu Pro Ala Phe Ala Thr Pro Thr Asp Pro
 130          135          140

Asn Tyr Gly Arg Thr Trp Ser Phe Cys Glu Phe Thr Phe Asn Pro Gln
 145          150          155          160

Gln Leu Tyr Ala Asn Ile Ser Tyr Val Asp Leu Val Thr Ala Leu Pro
          165          170          175

Ile Gly Leu Thr Leu Glu Gly Asp Ser Thr His Thr Val Ala Pro Leu
          180          185          190

Pro Asp Gly Ala Val Gln Arg Ile Ala Asp Asp Leu Thr Ala Gln Ala
 195          200          205

Ala Ala Asp Gly Gln Pro Trp Asp Lys Leu Val Thr Arg Gly Ser Asp
 210          215          220

Gly Gln Val Leu Arg Val Val Ser Pro Gln Asn Leu Met Ala Pro Tyr
 225          230          235          240

Phe Asp Arg Pro Asp Glu Met Pro Phe Arg Asp Leu Phe Ala Ala Gln
          245          250          255

Ile Asp Glu Val Trp Glu Lys Tyr Arg Ser Thr Asp Leu Arg Ile Asp
          260          265          270

Leu Gln Gly Gly Arg Gly Thr Leu Ala Gly Arg Val Ser Gly Asp Thr
          275          280          285

Leu Thr Phe Glu Gly Gly His Thr Phe Ser Lys Pro Thr Ser Lys Asp
 290          295          300

Ile Phe Thr Cys Asn His Gly Pro Phe Thr Asn Asn Pro Ser Asp Ser
 305          310          315

Asp Asp Lys Lys Ala Leu Leu Ala Arg Ile Ala Ala Gly Phe Asn Arg
          325          330          335

Ser Ile Met Leu Ser His Pro Ser Gln Pro Asn Gly Thr Ser Val Ala
          340          345          350

Asp Tyr Tyr Gln Asp Ala Val Thr Asn His Trp Ser Arg Val Val His
          355          360          365

Ala Asn Ser Pro Ile Gly Tyr Ala Phe Pro Tyr Asp Asp Val Arg Pro
 370          375          380

Asp Gly Glu Pro Asp Val Ser Gly Ala Ala Asn Asp Gly Asn Pro Arg
 385          390          395          400

Arg Phe Thr Val Ser Val Gly Ser
          405

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<210> SEQ ID NO 14
<211> LENGTH: 289
<212> TYPE: PRT
<213> ORGANISM: Streptomyces sp. ACTE

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<400> SEQUENCE: 14

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Val Leu His Pro His Asn Arg Thr Ala Arg Arg Thr Thr Arg Leu Thr
 1          5          10          15

Arg Thr Gly Gly Leu Ala Ala Ala Ala Leu Gly Leu Ala Leu Met Ala
          20          25          30

Leu Pro Val Thr Ala His Ala Gly Ala Pro Thr Gln Pro Ala Ala His
          35          40          45

His Leu Glu Ala Ala Ala Thr Gly Leu Asp Asp Pro Ala Lys Lys Asp
          50          55          60

Ile Ala Met Gln Leu Val Ser Ser Ala Glu Asn Ser Thr Leu Asp Trp
 65          70          75          80

Lys Ala Gln Tyr Gly Tyr Ile Glu Asp Ile Gly Asp Gly Arg Gly Tyr
          85          90          95

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Thr Ala Gly Ile Ile Gly Phe Cys Ser Gly Thr Gly Asp Met Leu Ala
 100 105 110
 Leu Val Glu Arg Tyr Thr Asp Arg Ser Pro Gly Asn Val Leu Ala Ser
 115 120 125
 Tyr Leu Pro Ala Leu Arg Glu Val Asp Gly Thr Asp Ser His Asp Gly
 130 135 140
 Leu Asp Pro Gly Phe Pro Arg Asp Trp Ala Glu Ala Ala Lys Asp Pro
 145 150 155 160
 Val Phe Gln Gln Ala Gln Asn Asp Glu Arg Asp Arg Val Tyr Phe Asp
 165 170 175
 Pro Ala Val Arg Gln Ala Lys Asp Asp Gly Leu Gly Thr Leu Gly Gln
 180 185 190
 Phe Ala Tyr Tyr Asp Ala Ile Val Met His Gly Gly Gly Gly Asp Ser
 195 200 205
 Thr Ser Phe Gly Ser Ile Arg Gln Arg Ala Leu Ala Glu Ala Glu Pro
 210 215 220
 Pro Ser Arg Gly Gly Asp Glu Val Ala Tyr Leu Asp Ala Phe Leu Asp
 225 230 235 240
 Ala Arg Val Trp Ala Met Arg Gln Glu Glu Ala His Ser Asp Thr Ser
 245 250 255
 Arg Val Asp Thr Ala Gln Arg Val Phe Leu Arg Asp Gly Asn Leu Asn
 260 265 270
 Leu Asp Pro Pro Leu Asp Trp Gln Val Tyr Gly Asp Ser Phe His Ile
 275 280 285

Gly

<210> SEQ ID NO 15
 <211> LENGTH: 790
 <212> TYPE: PRT
 <213> ORGANISM: Streptomyces sp. ACTE

<400> SEQUENCE: 15

Met Thr Pro Pro His Arg His Arg Leu Phe Arg Arg Ser Val Ser Ala
 1 5 10 15
 Ser Leu Ser Leu Ala Leu Thr Ala Val Gly Thr Ala Ala Ala Val Val
 20 25 30
 Leu Ala Gly Ala Pro Ala Ala Gln Ala Ala Ala Val Pro Ala Pro Ser
 35 40 45
 Pro Val Gly Ile Ser Gly Arg Gly Ala Ala Val Pro Phe Thr Glu Gln
 50 55 60
 Glu Ala Glu Tyr Ala Ala Thr Asn Gly Thr Leu Ile Gly Pro Asp Arg
 65 70 75 80
 Arg Tyr Gly Ser Leu Pro Ser Glu Ala Ser Gly Arg Gln Ala Val Thr
 85 90 95
 Leu Asp Ala Ala Gly Glu Tyr Val Glu Phe Thr Leu Thr Ala Pro Ala
 100 105 110
 Asn Ala Met Thr Phe Arg Tyr Ser Leu Pro Asp Asn Ala Ala Gly Thr
 115 120 125
 Gly Arg Asp Ala Ser Leu Asp Leu Arg Val Asn Gly Ser Val Leu Lys
 130 135 140
 Ser Val Pro Val Thr Ser Lys Tyr Gly Trp Tyr Tyr Gly Gly Tyr Pro
 145 150 155 160
 Phe Asn Asn Asn Pro Gly Asp Thr Asn Pro His His Phe Tyr Asp Glu
 165 170 175

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Thr Arg Thr Met Phe Gly Ser Thr Leu Pro Ala Gly Thr Lys Val Arg
 180 185 190
 Leu Gln Val Ala Ser Thr Ala Gly Ser Pro Ser Phe Thr Val Asp Leu
 195 200 205
 Ala Asp Phe Glu Gln Val Ala Ala Pro Val Gly Lys Pro Ser Gly Ala
 210 215 220
 Leu Asp Val Val Ser Asp Phe Gly Ala Asp Pro Thr Gly Ala Ala Asp
 225 230 235 240
 Ser Thr Ala Lys Ile Gln Ala Ala Val Asp Ala Gly Arg Thr Gln Gly
 245 250 255
 Lys Val Val Tyr Ile Pro Gln Gly Thr Phe Gln Val Arg Asp His Ile
 260 265 270
 Val Val Asp Gln Val Thr Leu Arg Gly Ala Gly Pro Trp Tyr Ser Val
 275 280 285
 Leu Thr Gly Arg His Pro Thr Asp Arg Ser Lys Ala Val Gly Val Tyr
 290 295 300
 Gly Lys Tyr Ser Ala Gln Gly Gly Ser Arg Asn Val Thr Leu Lys Asp
 305 310 315 320
 Phe Ala Ile Ile Gly Asp Ile Gln Glu Arg Val Asp Asn Asp Gln Val
 325 330 335
 Asn Ala Ile Gly Gly Ala Met Ser Asp Ser Val Val Asp Asn Val Trp
 340 345 350
 Met Gln His Thr Lys Cys Gly Ala Trp Met Asp Gly Pro Met Asp Asn
 355 360 365
 Phe Thr Ile Lys Asn Ser Arg Ile Leu Asp Gln Thr Ala Asp Gly Val
 370 375 380
 Asn Phe His Tyr Gly Val Thr Asn Ser Thr Val Thr Asn Thr Phe Val
 385 390 395 400
 Arg Asn Thr Gly Asp Asp Gly Leu Ala Met Trp Ala Glu Asn Val Pro
 405 410 415
 Asn Val Lys Asn Lys Phe Thr Phe Asn Thr Val Ile Leu Pro Ile Leu
 420 425 430
 Ala Asn Asn Ile Val Thr Tyr Gly Gly Lys Asp Ile Thr Ile Ser Asp
 435 440 445
 Asn Val Met Ala Asp Thr Ile Thr Asn Gly Gly Gly Leu His Ile Ala
 450 455 460
 Asn Arg Tyr Pro Gly Val Asn Ser Gly Gln Gly Thr Ala Val Ala Gly
 465 470 475 480
 Thr His Thr Ala Ala Arg Asn Thr Leu Ile Arg Thr Gly Asn Ser Asp
 485 490 495
 Phe Asn Trp Asn Phe Gly Val Gly Ala Ile Trp Phe Ser Gly Leu Asn
 500 505 510
 Glu Pro Ile Ser Asn Ala Thr Ile Asn Ile Thr Asp Ser Glu Val Leu
 515 520 525
 Asp Ser Ser Tyr Ala Ala Ile His Leu Ile Glu Gly Ala Ser Asn Gly
 530 535 540
 Leu His Phe Lys Asn Val Lys Ile Asp Gly Ala Gly Thr Tyr Ala Leu
 545 550 555 560
 Gln Ile Gln Ala Pro Gly Thr Ala Thr Phe Glu Asn Val Val Ala Thr
 565 570 575
 His Ile Ala Gln Ser Asn Pro Ile His Asn Cys Val Gly Ser Gly Phe
 580 585 590

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Gln Ile Thr Arg Gly Ser Gly Asn Ser Gly Trp Tyr Ala Asp Pro Pro
 595 600 605
 Ala Cys Thr Gly Val Trp Pro Asp Pro Val Trp Thr Asn Gly Gly Val
 610 615 620
 Pro Gly Gly Gly Gly Pro Thr Asn Pro Thr Asp Pro Thr Asp Pro Thr
 625 630 635 640
 Asp Pro Thr Asp Pro Thr Asp Pro Pro Glu Glu Thr Gly Asn Leu Ala
 645 650 655
 Arg Gly Arg Thr Val Thr Glu Thr Ser His Thr Asp Val Tyr Gly Ala
 660 665 670
 Ala Asn Thr Val Asp Gly Asn Ala Asp Thr Tyr Trp Glu Ser Arg Asn
 675 680 685
 Asn Ala Phe Pro Gln Ser Val Thr Val Asp Leu Gly Ala Ala Lys Ala
 690 695 700
 Val Lys Arg Val Val Leu Lys Leu Pro Pro Ala Ala Ala Trp Ala Thr
 705 710 715 720
 Arg Thr Gln Thr Leu Ser Val Ser Gly Ser Thr Asp Asn Gly Thr Tyr
 725 730 735
 Asn Ser Leu Lys Ala Ser Ala Gly Tyr Thr Phe Asn Pro Ser Ser Gly
 740 745 750
 Asn Thr Ala Thr Val Ser Leu Pro Gly Thr Pro Val Arg Tyr Leu Arg
 755 760 765
 Leu Thr Phe Thr Gln Asn Thr Gly Trp Pro Ala Ala Gln Leu Ser Glu
 770 775 780
 Leu Glu Ala Tyr Thr Ser
 785 790

<210> SEQ ID NO 16

<211> LENGTH: 514

<212> TYPE: PRT

<213> ORGANISM: Streptomyces sp. ACTE

<400> SEQUENCE: 16

Met Arg Arg Pro Val Ala Leu Arg Leu Ser Ala Ala Gly Ala Thr Leu
 1 5 10 15
 Ala Leu Ala Ala Ala Thr Gly Ala Leu Met Ala Met Pro Glu Ala Ala
 20 25 30
 Ser Ala Ala Thr Gly Gly Val Thr Gly Tyr Ala Thr Gln Asn Gly Gly
 35 40 45
 Thr Thr Gly Gly Ala Gly Gly Gln Thr Val Arg Ala Thr Thr Gly Thr
 50 55 60
 Ala Ile His Ala Ala Leu Cys Gly Arg Ala Ser Ser Ser Thr Pro Leu
 65 70 75 80
 Thr Ile Gln Val Glu Gly Thr Ile Asn His Gly Asn Thr Asp Lys Val
 85 90 95
 Ser Gly Ser Ser Cys Asn Thr Ala Ala Gly Val Ile Glu Leu Lys Gln
 100 105 110
 Ile Ser Asn Val Thr Ile Val Gly Val Gly Gly Gly Ala Val Phe Asp
 115 120 125
 Gln Val Gly Ile His Val Arg Glu Ser Ser Asn Ile Ile Ile Gln Asn
 130 135 140
 Val Thr Val Lys Asn Val Lys Lys Ser Gly Ser Pro Thr Ser Asn Gly
 145 150 155 160
 Gly Asp Ala Ile Gly Met Glu Lys Asp Val Arg Asn Val Trp Val Asp
 165 170 175

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His Thr Thr Leu Glu Ala Ser Gly Gly Glu Ser Glu Gly Phe Asp Gly
 180 185 190
 Leu Phe Asp Met Lys Ala Gly Thr Gln Tyr Val Thr Leu Ser Tyr Ser
 195 200 205
 Ile Leu Arg Asn Ser Gly Arg Gly Gly Leu Val Gly Ser Ser Glu Ser
 210 215 220
 Asp Leu Ser Asn Gly Phe Ile Thr Tyr His His Asn Leu Tyr Glu Asn
 225 230 235 240
 Ile Asp Ser Arg Ala Pro Leu Leu Arg Gly Gly Val Ala His Ile Tyr
 245 250 255
 Asn Asn His Tyr Val Gly Leu Ser Lys Ser Gly Ile Asn Ser Arg Ala
 260 265 270
 Gly Ala Arg Ala Lys Val Asp Asn Asn Tyr Phe Glu Asp Ser Lys Asp
 275 280 285
 Val Leu Gly Thr Phe Tyr Thr Asp Ala Ala Gly Tyr Trp Gln Val Ser
 290 295 300
 Gly Asn Val Phe Asp Asn Val Thr Trp Ser Gly Arg Ser Ser Asp Asn
 305 310 315 320
 Asn Pro Ala Gly Pro Asp Pro Gln Ser Asn Thr Ser Val Ser Ile Pro
 325 330 335
 Tyr Ala Tyr Thr Leu Asp Gly Ala Asn Cys Val Pro Ser Val Val Ser
 340 345 350
 Arg Thr Ala Gly Ala Asn Thr Gly Leu Lys Val Ser Asp Gly Ser Cys
 355 360 365
 Ser Pro Gln Thr Pro Asp Pro Thr Asp Pro Thr Pro Asp Pro Thr Pro
 370 375 380
 Asp Pro Thr Asp Pro Thr Pro Pro Thr Gly Thr Asn Leu Ser Leu Gly
 385 390 395 400
 Ala Gly Ser Asp Gly Ser Ser Lys Ala Ser Gly Thr Ser Tyr Gly Asp
 405 410 415
 Val Arg Asp Gly Asp Met Ser Thr Tyr Trp Ser Pro Ser Gly Ser Thr
 420 425 430
 Gly Ser Val Ser Ile Lys Trp Ser Ser Ala Thr Thr Val Ser Lys Ile
 435 440 445
 Asn Val Arg Glu Ala Ala Gly Ser Thr Gly Ser Ile Thr Ser Trp Lys
 450 455 460
 Val Gly Asn Ala Asp Thr Gly Ala Val Leu Ala Ser Gly Ser Gly Ala
 465 470 475 480
 Gly Val Ile Thr Phe Pro Gln Thr Ser Leu Arg Lys Ile Thr Phe Glu
 485 490 495
 Ile Thr Gly Ser Thr Gly Thr Pro Lys Val Ala Glu Phe Glu Thr Tyr
 500 505 510
 Ala Gly

<210> SEQ ID NO 17

<211> LENGTH: 1761

<212> TYPE: DNA

<213> ORGANISM: Streptomyces sp. ACTE

<400> SEQUENCE: 17

atgagccgca cgagccgcac caccctgcgc cgatcccgaa cagcaactcat ggcggcgggc 60

gccctcgtcg ccgcagccgc gggctccgcc gcagccgcgg cacccttcgg tgccaccgcc 120

gccgcggcgg ccggctgcac cgtcgactac aagatccaga accagtggaa cggcgggctc 180

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accgcctcgg tgagcgtcac caacaacggg gacgccatct ccggctggca gctccagtgg	240
agcttcgccc gcgccgagca ggtcagccag gggaggaaacg ccaccgtctc tcagagcggc	300
tccgcctca cgcacaagga cgcggctac aacgcgccc tggccaccgg ggcatcggcc	360
tccttcggtt tcaacgcgac gggcaacggc aacagcgtcg tccccgcgac gttcaagctg	420
aacggcgtca cctgcaacgg cggcaccacg ggcccgaccg atccccagga ccccacggac	480
ccgacggacc cgaccgaccc gcccgcgggc aaccgtgtgg acaaccctca ccagggagcc	540
aaggtctatg tgaaccggga gtggtcggcg aacgcgccc cagagccggg cggcgacaga	600
atcgccgacc agcccacggc cgtctggctg gaccgcatcg ccgcatcga gggcgcaac	660
ggttcgatgg gtctgcgca ccatctcgac gaggccctga cgcagaaggg ctccgcgaa	720
ctcgtctcc aggtcgtcat ctacaacctg cccggcgag actgcccggc gctggcctcc	780
aacggtgagc tcggaccgac cgagatcggc cgctacaaga ccgagtacat cgaccgatc	840
gcggagatcc tcggcagccc gaagtacggc ggctcgcga tcgtcaccac ggtcgagatc	900
gactcgtgc cgaacctcgt caccacgccc ggcggcccgc ccaaggccac tccggcctgt	960
gacgtcatga aggccaacgg caactacgtc aaggcgtcg gctacgcgt caacaagctc	1020
ggcgacgcgc ccaacgtcta caactacatc gacgcggccc accacggctg gatcggtgg	1080
gacgacaact tcggcgctc cgcggagatc ttccacgagg ccgacgcgc cgaggcgcg	1140
accgtcaacg acgtgacgg cttcatcacc aacaccgcca actacagcgc gctgaaggag	1200
gagaacttct ccacgcagca cgcctgaac ggcacgtcgg tccggcagtc gaagtgggtc	1260
gactggaacc gctacagga cgagctgtcc ttcgcgagg ccttcgcaa cgagctggtc	1320
tccgtcggct tcaactcggc catcggcatg ctcatcgaca cctcccgcaa cggctggggc	1380
ggcgcaacc gcccgagcgg accggcgcg aacaccagcg tcgacaccta tgtggacggc	1440
gggcgctaag accccgcat ccacctgggc aactggtgca accaggcagg agcgggtctc	1500
ggcgaacggc cgcagggcgc ccccgagcgg gggatcgacg cgtacgtctg gatgaagccc	1560
ccgggggagt ccgacggtc cagctcggag atcccgaacg acgagggcaa gggattcgac	1620
cggatgtcgc acccgaccta caggggtaac gcccgtaaca acaacaacat gtcggggcg	1680
ctgggtggcg ccccgctctc cgggaagtgg ttctcggccc agttccagga gctcatgaag	1740
aacgcctacc cggcgtctca g	1761

<210> SEQ ID NO 18

<211> LENGTH: 2865

<212> TYPE: DNA

<213> ORGANISM: Streptomyces sp. ACTE

<400> SEQUENCE: 18

gtggccgccc tcgccctccc cttgggaatg accgcagcgg ccggcaccga ggcccaggcc	60
gccgcctcgg cgtgcagcgt cgactacacg accagtgact ggggatcggg gttcaccacc	120
gaactcacc tgaccaaccg gggctcggc gcgatcgacg gctggaccct gacgtacgac	180
tacgcccggg accagcagct cagcagcggc tggagcggca cctggctcca gtcaggcaag	240
accgtcagcg tgaagaacgc agcctggaac ggtgcgatcg ccgcccgtgc cgcgctcacg	300
accggcgcgc agttcaccta cagcggcgcc aacaccgac cgaccacctt cgcgctcaac	360
ggcagcgtct cgcggggggc ccaccagcgg ccgatcggc tcctcacctc cccggcgcg	420
ggcgccgtct tctccgccc ggaccgggt ccgctggcgg cgaaccgcgc ggcccgggac	480

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ggggcgacga tcagcaaggt cgagttctac gacgacacga ccctcctcgg caccgacacc	540
acctccccgt acagctacga ggccgggcaa ctggcgggccg gcagccactc cgtgtacgcc	600
agggcctacg acagcctcgg cgcctccgcg gattccccgc ccgcccgcac caccgtcgtc	660
accggccccg cggtcgtcgt cccccccgt caactcggcg tccagcaggg caggtcggga	720
accttcgacg tctcgtcgtc caccgcgccc gcggcgagcgc tcaccgtcac ggccgccccg	780
tccgcggtga acaccgggct gagcgtcacc gcggggtcga ccctcacctt cccccccg	840
aactgggtcca caccacagaa ggtgacgcgc acggccgacg gctccggcac cggggccg	900
accttcaccg tcaccggcccc cggccacggc aaggccgagc tcaccgtcac ccagctggcg	960
gcggcggaag agtacgacgc ccgtttctc gacctctacg ggaagatcac cgatccccg	1020
aacggctact tctcgcggga ggaatcccc taccactccg tcgagacgct gatcgtcgag	1080
gcgccccgacc acgggacgca gaccacctcg gaggcctaca gctacctgat ctggctgcag	1140
gcgatgtacg gcaagatcac cggcgactgg accaagttca acggtgcgtg ggacaccatg	1200
gagacgtaca tgatccccac ccacgcccac cagcccacga actccttcta cgacgcgtcc	1260
aagcccccca cctacgcgcc cgagcacgac accccgaacg agtaccgccg ggtgctcgac	1320
ggctccgcct cctccggctc cgacccgatc gcggcagagc tgaagagcgc gtacggcacc	1380
gacgacatct acggcatgca ctggatccag gacgtcgaca acgtctacgg atacggcaac	1440
gcgccccgca cgtcgcgggc cggccccacc caggccggtc cgtcctacat caacaccttc	1500
cagcgcggct cgcaggagtc ggtctgggag accgtcaccc acccgacctg cgacaacttc	1560
acgtacggcg gcgccaacgg ctacctcgac ctgttcaccg gggactcctc gtacggcaag	1620
cagtggaaag tcaccaacgc ccccgacgcc gacgcccgcg ccgtgcaggc cgctactgg	1680
gcccgcgtct gggcgaaagga gcagggggaag gcgggogaag tcgcccacac cgtcggcaag	1740
gcggcgaaga tgggtgacta cctgcgctac tccatgttcg acaagtactt caagaagatc	1800
ggcgactcgc tcggccccgac cacctgcgcc gccggctccg gcaaggacag cgcgcactac	1860
ctgatgtcct ggtactacgc ctggggcggc gccaccgaca cctcggccgg ctggctcctgg	1920
cggatcggct ccagccacgc ccacggggga taccagaacc cgatggcggc ctacgcgctg	1980
agctccgtgg ccgacctcaa gcccaagtgc gccaccggag cgcaggactg ggccaagagc	2040
ctggaccgcc aactggactt ctaccagtgg ctccagtcgg acgaggggtc catcgcgggc	2100
ggtgcgacca acagctggaa gggcagctac gccagcccc cggccggcac gccgaccttc	2160
tacggcatgt actacgacga gaagccccgtg taccacgacc cgccgtccaa ccagtggttc	2220
ggcttccagg cgtgggtccat ggagcgcgtc gccgagtact accacgagtc ggggtgacgcc	2280
caggcgaagg ccgtgctcga caagtgggtc gactgggccc tgtccgagac gaccgtcaac	2340
ccggacggca cctatctgat gccctccacc ctccagtggc cgggcgcgcc ggacacctgg	2400
aacgcctcga acccccgtgc caacgcccag ctccaagtca cggctgccga ctacaccgac	2460
gacgtcggcg tggccggcgc gtacgcccgg acaactgacct actacgccgc caagtccggt	2520
gacacggagg ccgaggccac cgcgaggcgc ctgctcgacg gcatgtggca gcaccaccag	2580
gacgacgcgc gcgtggcggc gcccgagacc cgcgcccact acaaccggtt cgacgacccg	2640
gtctacgtcc ccggtggctg gacgggccc atgcccacg gtgacaccgt cgacgaggac	2700
tcgacgttcc tctccatccg ctctctctac aaggacgacc cgaactggcc ccaggtgcag	2760
gcgtacctgg acggcgggtc cccccggtc ttcacctacc accggttctg ggcgcaggcc	2820
gacatcgcac tggccctggg ggcgtagccc gacctcctgg agtga	2865

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<210> SEQ ID NO 19
 <211> LENGTH: 1089
 <212> TYPE: DNA
 <213> ORGANISM: Streptomyces sp. ACTE

<400> SEQUENCE: 19

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atggctagac gcagcagact catctcctg gcagcgggtg tggccaccct gctcggggcg      60
ctcggcctca ccgactctg gccgggcaag gcgaggcg acggtgtcgc gatgaccccc      120
ggatcgcgta cctatctgtg ccagctcgac gccctgtccg gcaccggcgc gctgaacccc      180
acgaacccgg cctgccggga cgcgctgagc cagagcggcg cgaacgcgct gtacaactgg      240
ttcggcctgc tcgactccaa cgcgggcggc cgcggcggcg gatatgtgcc ggacggcagc      300
ctgtgcagtg ccggtgaccg ctecccgtag gacttctccg cctacaacgc cgcgcgcgcc      360
gactggcccc ggacacatct gacctccggt gcgacgctca aggtgcagta cagcaactgg      420
gccgcccacc ccggtgactt ccgggtctac ctgaccaagc cgggctgggc acccacgtcc      480
gaactcgctt gggacgacct tcagttggta cagaccgtaa gcaacccgcc gcagcagggc      540
ggggcgggca ccaacggcgg gcaactactac tgggaacctg cgctgccgtc gggccgttcc      600
ggtgacgcgc tgatgttcat ccagtggtg cgttcggaca gtcaggagaa cttcttctcc      660
tgctcggaca tcgtcttoga cggcggcaac ggcgaggtga cgggaatcgg cggcacgggc      720
acccccacc cactccgac cccgactccg accccgacc cgcacggacc ggagcactcc      780
ggttctctga tggccgtcta caacgtcgtc agctcctggg ccggtggctt ccaggcctcc      840
gtcagagtgta tgaaccacgg tacggaaccg cgcacggctt gggccgtgca gtggaagccc      900
ggttccggga cgcagatcaa cagcgtgtgg aacggctccc tctccaccgg gtccgacggc      960
accgtgacgg tgcgcgacgt ggaccacaac cgtgtcatcg ccccgacgg cagtgtgacc     1020
ttcgggttca ccgccacctc cacgggcaac gactaccccg ccgggacgat cgggtgtgtg     1080
acgtcctag                                     1089
  
```

<210> SEQ ID NO 20
 <211> LENGTH: 1371
 <212> TYPE: DNA
 <213> ORGANISM: Streptomyces sp. ACTE

<400> SEQUENCE: 20

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gtgaaacgct ttctggcctt actggccacc tgcgcgacgg tcctgggctt cacggcactg      60
accggcccc aggcgggtgg cgcgcggggc tgcacggccg actacacgat caccagccag     120
tggcagggcg gcttccaggc cgcggtgaag gtcaccaacc tgggaacccc cgtgaccggg     180
tggaaagctca cgttcacctt gccggacgcg ggacagaagg tcgtccaggg ctggaacgcc     240
gcctgggtcgc agtcgggttc cgcggtcacc gccgcggcgc ccgactggaa cggcacactg     300
gccaccggcg cgtcggccga ggcgggcttc gtgggtcctt tcacgggcgc caaccgcctt     360
cccacggcgt tcgcgctcaa cgtgttcgcc tgtacgggct ccacgggaga acccccggcc     420
ggctccgacg gggcaccccc cgtggaacgt aacgggcagc tccacgtctg cggggtgaa      480
ctctgcaacc agtacgacc gcccggtcag ctgcggggta tgagcacgca cggcatccag     540
tggttcgacg cctgctacga cgcgcctcc ctggaacgcgc tggcgaacga ctggaagtcg     600
gacctgctgc gcatcgccat gtacgtgcag gaggacggtt acgagaccga cccggcgggc     660
ttcaccggcg gcgtgaacga cctcgtcgac atggccgagg cccgcggcat gtacgcgttg     720
  
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atcgacttcc acaccctgac cccgggcgac ccgaacgtca acctcgaccg cgccaagacg	780
ttcttcgcgt ccgtcgccgc gcgcaacgcc ggcaagaaga acgtgatcta cgagatcgcc	840
aacgagccca acggcgtgac ctggacggcc gtcaagagct acgccgagca ggtcatcccg	900
gtgatccggg ccgccgaacc ggacgccgtc gtcacgtctg gcaccccgcg ctggtcctcg	960
ctggcgctct cggacggctc cgacgagagc gaggtcgtca acagccccgt caatgccacc	1020
aacatcatgt acgcgttoca cttctacgca gcgagccaca aggaagccta ccgctccacg	1080
ctgagccggg cggcggcgcg gcttccgctc ttcgtcaccg agttcggcac ggtgagcgcc	1140
accggcggcg gggcgatgga cggggcgagc accacggcct ggctggacct gctcgaccag	1200
ctgaagatca gctatgcaaa ctggacctat tccgacgcgc ccgagagcag cgcggcgctc	1260
cggccgggca cctgcccggg cggcgactac agcggcagcg gcgtgctgac cgagtccggg	1320
gcgctgctca agaaccggat cagcaccccc gattccttcc ccacggctg a	1371

<210> SEQ ID NO 21

<211> LENGTH: 1377

<212> TYPE: DNA

<213> ORGANISM: Streptomyces sp. ACTE

<400> SEQUENCE: 21

atggccaaga aaatccccgc ccgtgccaga cgggcactct ccgtcctgac ggcgggctg	60
ctcgcgccgc cggcgtcgt ctcgctcgcc ggcacggccg aggcagcagg caccctgggt	120
gacgcggcgg cggcgaaggg ccggtaactc ggcaccgcgg tcgcggcgaa ccacctggc	180
gaggcaccgt acgcgtccac gctggacgcc cagttcgact cggtcacccc ggagaacgag	240
atgaagtggg acgcggtcga gggcagccgc aactccttca ccttcacggc cgcgaccag	300
atcgtcagtc acgcccagag caagggaaatg aaggtgcccg ggcacacct ggtgtggcac	360
tcgcagctgc cgggctgggt cggcggcctg ggcgccaccg acctccgcgc ggcgatgaa	420
aaccacatca cccaggatgat gacgcactac aagggcaaga tccattcctg ggacgtgggt	480
aacgaggcct tccaggacgg caacagcggg gcccgggcga gctctccctt ccaggacaag	540
ctgggtgacg gcttcatcga ggaggcgttc cgcaccgccc gtacggtcga tccgaccgcg	600
aagctctggt acaacgacta caacaccgac ggcgggaacg cgaagagcga cgcggtctac	660
gccatggcga aggacttcaa gcagcgcggg gtgccgatcg actgctggg cttccagtcc	720
cacttcaaca gcaactcccc cgtgccctcc gactaccggg ccaatctcca gcgcttcgcc	780
gacctcggtc tcgacgtcca gatcaccgaa ctggacatcg agggttccgg ctcgcccag	840
gccgcgaact acacgagcgt cgtgaaacgc tgcctggccc tgaccgcctg caccggcctc	900
accgtctggg gtgtaccga caagtaactc tggcgcagca gcggcaccgc gctgctcttc	960
gacggcgact acaacaagaa gccggcgtac gacgcgggtc tcgccgcgct cggcggcacc	1020
cccgcgggtg gcgggtgacga cggcggcggc gacaaacggc gcgggaaacac cggcagctgc	1080
acggcgcagt acacgcagac cggccagtgg aacggcgggt acaacgggtg ggtgacggtc	1140
aaggcaggct cctccgcat caccacctgg tcggtgcccg tgaccgtgcc ctctccag	1200
caggtctccg ccctctggaa cggcgcctcc acgtggaacg ccggcaaacac cgtgatgacg	1260
gtgaagccca cctacaacgg gaccctggcg gccgggtgctt cgacgagctt cgggttcacc	1320
gtcatgacga acggcaaacac ctggcgccc gccgtcggcg cctgcaccgc ctctgta	1377

<210> SEQ ID NO 22

<211> LENGTH: 1689

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<212> TYPE: DNA

<213> ORGANISM: Streptomyces sp. ACTE

<400> SEQUENCE: 22

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gtgagaacag cgatacgcac agcacgacga ccacagcccc tggcccttct gctgagaggt    60
ctggccgcct tcctggggct cgcctcgcc ggagccctcg gcccggccac cgcgcgggcc    120
gcggaacctgc cccagcgggc ggaggcgcgg gccgcgggcc tccacatcag cgacgggggc    180
ctggtcgaag gcaacggcaa cgacttcgtc atgcgcggca tcaaccacgc ccacacctgg    240
tatccgggag agaccagtc cctcgccgac atcaaggcga ccggcgcgaa cacggtccgc    300
gtggtgctgt ccgacggcta ccgctggagc gagaacagcc ccgaggacgt cgcctcgatc    360
atcggccggt gcaaggccga gcggctcatc tgcgtcctgg aggtccacga caccaccggg    420
tacggggagg acgcccgcgc cggaaacctc gaccacgcgg ccgaactactg gatcggcctg    480
aaggacgtac tcgacggcga ggaggactac gtcgtcatca acatcggcaa cgagccctgg    540
ggcaacgcgc atccgggggg ctggaccgcc cccacgacgg ccgcgatcca gaagctgcgc    600
gccgcgggtt tcgccacac gatcatggtg gacgcgccca actggggcca ggactgggag    660
ggcgtcatgc gggccgacgc ccggagcgtg tacgacgccg acccgaccgg caatctgatc    720
ttctcgatcc acatgtacag cgtctacgac accgcccgca aggtccacga ctacctcaac    780
gccttcgtcg acgcccgaact tcccctgctc atcgccgagt tcggcggccc cgcggaccag    840
tacggcgacc cggacgagga cacgatgatg gccaccgccg aggagttggg gctcggttac    900
ctggcctggt cctggagcgg caacacggat ccggctcctcg acctggtcct cgacttcgac    960
cccccccggc tcagctcgtg gggcgcgcgc gtcctccacg gcccgcacgg catcacccgag    1020
acgtcccgtg aggccaccgt cttcggcggc gggcagggcg ggggcgacac cgaggccccg    1080
accgcaccgg gcaccccgc ggcctccggg gtgacggcga cctccgtcac cctcggctgg    1140
agtgcggcca ccgacgacgt cggcgtcacc gcgtacgacg tggtcgcggt gaccggcggc    1200
tccgagacga aggtcgctc ctccgcggcc acctcggtea ccgtgaccgg tctgagcggc    1260
ggcaccgcgt acagcttcgc cgtctacgcc cgggacgcgg ccggcaaccg ttcggcgcgc    1320
tccggcacgg tgtcggtaac caccgacgag ggcggcagcg tgcccggggg cgcctgctcc    1380
gtgggctacc gggatgatcg cgagtggcgg gccggttcc agggggagat cacctccgg    1440
aacaccggcg ccgcccgcgt cgacggctgg acgctgggct tcgccttcgc cgacgggacg    1500
accgtcacga acatgtgggg cggcacccgc acgcagagcg ggggcgcggg gagcgtcacc    1560
ccggcctcgt acacctccac gatcgccgcc ggcggtcgg tcaccgtcgg ctccaccggc    1620
accctgactg gcgcgaacgc cgcgccggcg gccttcacgc tcaacggcgc cacctgcacc    1680
gcggcctga                                     1689

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<210> SEQ ID NO 23

<211> LENGTH: 987

<212> TYPE: DNA

<213> ORGANISM: Streptomyces sp. ACTE

<400> SEQUENCE: 23

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tccgccctgg cgggcggcgc cgtcacccgc gcaccggccc gggccgcgcg ttgcaacggc    120
tacgtcgggc tcaccttcga cgacggaccg tcggcggccc agaccccggc cctgctgtcc    180
gcgctcaagc agaacggcct gcggggccacc atgttcaacc agggcaacta cgcgcctccc    240

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aaccceggccc aggtcaaggc ccaggtcgac gccggcatgt gggtcggcaa ccacagctac 300
agccaccgcg acctgacceca gcagagccag gcgcagatgg actccgagat ctcccggacc 360
cagcaggcca tcgccgccgg aggcggcggc acaccgaaac tgttcgccc gccgtacggc 420
gagaccaacg ccacgctgcg gtcggtcgag gcgaagtag gtctcaccga ggcatctgg 480
gacgtcgact cgcaggactg gaacggcgcg agcaccgacg cgatcgtgca ggcggctccc 540
cggtccaccg ccggtcaggt catcctgatg cagcagtgcc ccgccaacac cctcgccgcg 600
atcccgcgca tcgcccagac cctgtccgcc aaggggttgt gttccggcat gatctccccg 660
cagaccggcc gcgccgctgc tcccagcggc ggcggcaacg gtggaggggg cggtgccggt 720
ggcgggtgca ccgcgacggt gtcggcgggt gagaagtggg gtgaccggta caacctgaac 780
gtggcggtag gcggtccag caactggacg gtgacgatga acgtgccgtc gggcgagagg 840
gtcatgacga cctggaacgt cagcgcgagt tatccgagcg cgcaggtcct ggtcgccaag 900
ccgaacggga gcgggaacaa ctgggggtcg acgatccagg ccaacggcaa ctggacctgg 960
ccgaccgtct cctgcaccac gagctga 987

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<210> SEQ ID NO 24

<211> LENGTH: 1008

<212> TYPE: DNA

<213> ORGANISM: Streptomyces sp. ACTE

<400> SEQUENCE: 24

```

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agcgtctgcg ccctggtaact ggccgcccgc gccgcgatgc tgctgcccgg caccggccagt 120
gccgacacgg tcgtcacgac gaaccagacc ggcaacaaca acggctacta ctactcgttc 180
tggaccgacg gcggcggcca ggtctccatg aacctggcct ccggcggcag ctacagcacc 240
tcgtggacga acaccggcaa cttcgtcgcc ggcaagggtc ggagcacggg cggccgtaag 300
agcgtcaact actcgggcac cttcaaccgg tccggcaacg cctacctgac gctgtacgga 360
tggtcgacga acccgtctgt cgagtactac atcgtggaca actggggcac ctaccggccc 420
accggtacgt tcaagggcac ggtctccagc gacggcggca cgtacgacat ctacgagacc 480
accgcacca acgccccctc catcgagggt acgaagacct tcaagcagtt ctggagcgtc 540
cggcagtcga agcgggaccg cggcaccatc accaccggca accacttcga cgctcggccc 600
cgcaacggca tgaacctcgg caccatgaac tacatgatcc tcgcccacga gggctaccag 660
agcagcggca gctccaacat caccgtgacg gagggcggat ccggtggtgg cggcgacaac 720
ggtagggggg gcggtggcgg tggcgggtgc accgccacgt tgtcggcggg tgagaagtgg 780
ggtgaccggt acaacctgaa cgtggcggtg agcggctcca gcaactggac ggtgacgatg 840
aacgtgccgt cggcgggagaa ggtgctgtcg acctggaaca tcagcgcgag ttatccgagc 900
tcccaggtcc tggtgcgcaa gccgaacggg agcgggaaca actgggggtgc gacgatccag 960
gccaacggca actggacgtg gccgaccgtc tctgcacca cgagctga 1008

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<210> SEQ ID NO 25

<211> LENGTH: 843

<212> TYPE: DNA

<213> ORGANISM: Streptomyces sp. ACTE

<400> SEQUENCE: 25

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atgagtgaaa gagccgcac cccacgtacc caccggcgcc gcccccggcg ccggcgcac 60
gccaccgccc tgaccggcgc actgggctc accggcggcg cactggccac cggcgtgatg 120

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ctccagccgg cggcgcgggc caccaccgcg atccccgct ggcctccgc cacgggcagc	180
cagtccgtct cgaagaccat cgaggctcc gggacgtacg acggcggctt gaagcgcttc	240
accggcagcg gtgacctggg cgacgggtgc caggacgagg gccaggaccc gatcttcaag	300
ctgaaggacg gggcgacgat caagaacgtc atcctgggca ctccggcgc cgacggcatc	360
cactgctccg gcagctgcac gatccagaac gtctggtggg aggacgtcgg cgaggacgcc	420
gcgtccttca agggcacctc cacgtcgtcc gtgtacacgg tgtacggcgg cggcgcgaa	480
aaggcctccg acaaggtott ccagttcaac ggcgcgggca agctggtcgt gacgaagttc	540
caggtcgcgg acttcggcaa gctggtccgc tcgtgcggca actgctcca gacgtacaag	600
cgcgagatca tcgtcaacga cgtcgcagtc acggcgccgg gcaagtcctt ggtcggcatc	660
aacaccaact acggggacac cgcgcgctg cgctcggtc gcgtccacgg cgacagcagc	720
aagaagatca agccctcgtt ccgctacacc ggcaacagca cggcgcgga accgaaggag	780
acgggcagcg gtccggacgg cacgtactgc aagtacacc cctcggacct gagctacgac	840
tag	843

<210> SEQ ID NO 26

<211> LENGTH: 2730

<212> TYPE: DNA

<213> ORGANISM: Streptomyces sp. ACTE

<400> SEQUENCE: 26

atgtggtgtc acccgtacct ccgcctccgc acgtccggac gaaaggtttc ctcggtgaa	60
gcccttccac cccccgccc gcccgcaccc gtccgaccac ggtcccggta cggcgggcgc	120
gtgctcggga tgcggccgc cgccctgctg tgcgcagggg ccctggccgt gcccggtacg	180
gccatggccg acgacgccga acccggaccc ggccccgagc agatcaccaa cggcgacttc	240
gccaccggta cctcagcccc gtggtggtgg acgcccgaac cctcggcgc cgtgtccgag	300
ggccggctct gcgtggaggt gcccgccggc acggccaacg cctgggacgt catcgtcggc	360
cagaacgacg taccgatcgt cgcgggcgag agctacgagc tgcctacac ggcgcgttcg	420
accgtgcccc tgaccgttca gaccgggtc caggaggcgg tggagcccta cacgacggtg	480
ctggcgacgg cggatccggt gggcgcgag gacacgcggg tcgcccgcac gttcacggcc	540
tcggtggacc agccccgcgc gtcggtgacg ttgcagatcg gtggcgggga gcgggcgacg	600
acgttctgcc tggacgacgt gtcgctcggg ggcggggccg agccgccctt gtacgtaccg	660
gacaccgget cgcgggtccg cgtcaaccag gtcgggtatc tgcccccgcg tcccagagc	720
ggcaccgtgg tcaccgacgc cgaggcgcgg ctgacctgga cggtaaaagc cgaggacggt	780
tcgacggccg ccaccggtac gaccgttccg cgaggtgagg accccagctc gcgccgacgg	840
gtccacacct tcgacttcgg cgacctacc acggcggggg acggctacac cgtggaggtc	900
gacggtgagg tgagcgagcc gttctcgatc cgcggggacc tgtacgactc cctgcgctcg	960
gacgcgctgg cgtacttcta ccacaaccgc agcggcatcg agatcgacgc ggacctcgtc	1020
ggtgagcagt acgcgcgcc gcccggtcac atcggcgtcg cgcccacaa gggcgacacg	1080
gacgtgccgt gccgacctgg ggtctgcgac taccggtgg acgtgtcggg cggtcgttac	1140
gacgcggggc accacggcaa gtacgtggtc aacggcggga tctcggggc ccagctgatg	1200
gccacgtacg agcggacct caccgccccg gacgcggagt cggccgagct cggcgacggc	1260
gcgctcgggg tgccccgagc cgacaacggg gtgcgggaca tcctggacga ggcgcgctgg	1320

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gagatggact tcctcatcaa gatgcaggtc ccggcgggcg agcagctggc ggggatggtc	1380
caccacaaga tgcacgacgc cgagtggacc gggctgccga tgaagccgca cctggacccg	1440
cagcagcgcg agctgcaccc gccgtcgacg gccgccacac tcaacctcgc cgccacggcc	1500
gcccagtgcg cccggctcta cgcgcccttc gacgcggact tcgctggacc ctgectgctg	1560
gccgcccaga ccgctgggga ccgcccgaag ccgaccccgg acgtgctcgc cgaccccgaac	1620
gacggcatcg gccggctgtc gtacaacgac gacgacgtct cggacgagtt ctactgggctg	1680
gccgcccagc tcttcaccac gacgggcaag gacatctacc ggcagggcgt gctctcctcc	1740
gcatggcacg gtgacgcggg ccgctctctc ccggcgggctg gcggaatctc ctggggctcc	1800
acggcccggc tcggcgtgct caccctggcc accgtgccca acgcccctgac gtcgatcag	1860
ctcggcccagg tgcgcacggg ggtgaccgag gccgcccacc gctacgcccgc gcagtcccgt	1920
gagcagggct acgggctgcc gtacgcgcc ccggggggagg actacgtctg ggggtccaac	1980
agtcaggtgc tcaacaacat ggtcgtctct gccaccgccc acgacctgac cggtgacgcc	2040
gcctaccagg acgcccgtgt ccggggcgcc gactatctgc tgggcccga cccgctgaac	2100
cagtcgtacg tcaccggcta ccgagcggc gactcgcaca accagcaca ccgcttctgg	2160
gcgaccaga acgaccccag cctgccgaac ccggcggccc gttcgatcgc gggcgcccc	2220
aacctcacg cgatcgcctc ccgtgacccg gtggcggcgg agaagctcag ccgctgcgcg	2280
cccgccatgt gctacgtcga cgacatcgcc tcttggcgca ccaacgagat caccatcaac	2340
tggaacgcac cgctgcctt catcgcctcc tacctggacg acgcccggca gggcgggcag	2400
accgcccggg cccgcacctg ccaggtcacg tactcctcgc acccgtggaa cagcgggtcg	2460
acggtgacgg tacgggtcga gaacacccgc tcggatcccg tctcgcctg gccgctgacc	2520
tggctgctcc ccggcgagca gccgctgagc cacacgtgga gcgcccaggt cgaccagcac	2580
ggccgtacgg tcagcggccc gccgctgctg tggaaaccga ccctggcacc ccgcccggcg	2640
gtcgacttcg gcttcaacac ctcggcggcg ggtcctcgc ccgagcccgg ccggttcaag	2700
ctgaacggcc gggcctgctc agcgggctga	2730

<210> SEQ ID NO 27

<211> LENGTH: 1218

<212> TYPE: DNA

<213> ORGANISM: Streptomyces sp. ACTE

<400> SEQUENCE: 27

atgctgaccg gatccatcgc gccgctcctg ggcctcgcgc ccgcccctggc cgcactgctc	60
accacggcct tcattggccc ggccatggcc ggcaaacacg acgcccaccga ctcccctcc	120
gccgcccggc ccccgccgct cttcaccac cccgcccctc tggtcagccc gccgcagctc	180
gacttcgtac gccgcaaggc ccagggcggg gccacgccc ggaagggggc gtacgaccag	240
atgctggcca gtcctcagc ctcgctctcg cggaccccga agccccgcgc cgtcgtggag	300
tgcggctcgt actccaacc caacaacggc tgcaccgacg agcgcgagga cgcgctggcc	360
gcgtacaccc tctcgtggc ctggatcacc agccaggacg gccgctacgc ccagaaggcg	420
atccagatca tggacgcctg gtcgggctg atcaaggacc acaccaacag caacgcccc	480
ctgcagacgg gctgggcccg ctctcctcgg ccgcccggcg ccgagatcat caagtacacg	540
tacggcaact ggcggcgctc ccgccccttc gccaccatgc tgcgtgacgt ctacctgcc	600
aaggtcgcga accgctcga cagcaacggc aactgggaac tctccatgac cgaggcccgcg	660
atcggcatcg ccgctctcct ggaggaccg gccgcccacg acagggcccgt cgccaagttc	720

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cgcggccgcg tccccgcgta catatacgtg accgcgcgacg gatcgctgcc gaaggccgcg	780
cccggcagcg gtctcgacac gcgggaaaag atcatcaact actggcaggg ccagtcgacc	840
ttcgtggacg ggctctcgca ggagacctgc cgcgacctca cccacaccgg ctacgggctc	900
tccgcgatct cccacatcgc cgagaccagc cggatccagg gccaggacct ctaccggag	960
gtcggccgacc ggctccgtca cgcgctgggg ctgcacgcc agtaccagct gggggagaag	1020
gtcccgtcct ccctgtgcgg cggctcgtc aaggacagcc tcggcccgg caccgaggtc	1080
ggcttcaacg ccctgcacaa ccgcatgggt tacgccatga cgaacaccca gaccctcacc	1140
gagcggcagc ggccccgcgc ctcgaacaac ctgttcgtgg cctgggagac cctgacgcac	1200
gccgacaacc cgaactga	1218

<210> SEQ ID NO 28

<211> LENGTH: 1881

<212> TYPE: DNA

<213> ORGANISM: Streptomyces sp. ACTE

<400> SEQUENCE: 28

atgccctccc gtacgacgtt gatcgccacc accgcggccc tggtcgcct cgccgcccc	60
atggccttcg cggctcccgc ccccgcccc gaccccgccg tcgaggccgc cgccgaggcc	120
tgggacaccg accgcgcggc gtcgcctac gggcggaacc ccgcgcctg caccgcgtcc	180
ggcagcgaga accccgcctc cggaccgggc gccgcccaccg acggcgacgc caccaccgc	240
tggtcacgcg acttcgcgca caacgcctgg atacgcgtcg acctcgctc caccatccgg	300
atcaaccagg tgaagctgga gtgggaggcc gcctacggca agaagtacgt cctggaagtc	360
tccaaggacg gcaccaactg gaccccctc tacacggagg acgcgggac cgcgggcacc	420
gtcaccgccc acacctacc gcaggaggtc accggccgct acgtgaggat gcgcggcgtc	480
gaacgcgcca cggcctgggg ctactccctc ttctccttc aggtctacgg gggcgagccg	540
gccccgcct cgaccaccg cagcaacctc gccctcaacc acccgccta cggcgacctc	600
taccgacagc ccggcaactc gcccgcatc gtcaccgacg gcggctggcc cgccgacctg	660
aaggcggacc gctcccgtg gtctctcgac tggaaacgcg accgctgggt cggcgtcgac	720
ctcggcgcga cctccaccat caacagcgtc gacctctact gggaggcggc ctacgcctc	780
gactacgaga tccaggtgtc cgacgacaac cggacctggc ggacctcca ccgcccctcc	840
gccgcccagg tcgcccag acgcgcgac gtcaaggccc cggccgaggc cgtcggacgc	900
cacgacacca tcaacctgcc caccggccc accggccgct acgtccggat gctgggcaag	960
gagcgcggtt cctctacaa cccggcacc tccaccgccc agttcggcta ctgctctac	1020
gagttccagg tgtggggcgc cggcggcagc gcggacgccc cctaccccgc cctgcccagg	1080
aaccggcg gcgcctaccg caccacctc ttcgacgact tcaccggctc cggcctggac	1140
cgctccaagt ggcgctgggt gcgcaccggt acggagatgg gcccggtcaa cggggagtcc	1200
caggcctacg tcgactgcc ggacaacatc cgtaccgaga acggcgcctt ggtcctggag	1260
tccaagtact gcaagggtg cccccccag cccaacggca ccttcgactt cacctgggc	1320
cgcgtcgaca ccaacaccaa gttcgacttc acctacggca aggtgagcgc ccgtatgaag	1380
ctcccggtcg gcgacggtt ctggccggcg ttctggctgc tgggcagcga cgtcgcgac	1440
ccggcgtct cctggcccgc ctcggcgag acggacatca tggagaacat cggctacggc	1500
gactggacca gctccggcct gcacggacc ggctactccg cagacggcaa catcggcgcc	1560

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tcccagacct acccgaacgg cggccgggccc gacgagtgge acacctacgg cgtcgaatgg 1620
acccccgaag gcatgacctt caccgtogac gaccgogtgg tgcagcagac ctcccgccag 1680
aagctggagt ccaccgcggg caagtgggtc ttcgaccaca accagtaagt gatcctcaac 1740
ctggccctcg gcgggcgcta cccggggcga tacaaccagg tcaccagcc ctactggggc 1800
cttccgcagt ccagcgtoga ccgcatogca cagggcgcca tcaaggcgga gatcgactgg 1860
gtacgggtcg agcagaagta a 1881

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<210> SEQ ID NO 29
<211> LENGTH: 1227
<212> TYPE: DNA
<213> ORGANISM: Streptomyces sp. ACTE

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<400> SEQUENCE: 29
gtgatttcgc gcagaatggt cctgaccggc gccgcgcct ccgacaccgc gctcaacctat 60
cgctctggg gcaccgcct gagccgcgc acgtggggcg cggccgccac gtggaactg 120
gccctcgaga accgttcggt gcccggtacg gtgacgcct acgtaccgg tcacgagcag 180
ggcaccgaca gctgggtgct gctgcccggc gacggcagcg tgtaccgcc cgagtcgcc 240
ggcgtccgc agaccctct gccggtggac tgcgccatcc cgctgaacgg cgcggcgcc 300
ggcccggtcg tcctgacgct gcccagatg tacggcgcg ggtctactt cgtccgtgac 360
gacaagctgg acttctacct gaaccgggc cctcgctgg tcgagccggc ctccgcgac 420
cccaccgacc cgaactacgg gcgcacctgg tegtctgg agttacact caaccgcag 480
cagctgtagc cgaacatcag ctacgtogac ctggtaccg ccctgcccag cggcctgacc 540
ctggagggcg actccacca caccgtogcc ccgctcccgg acggcgccgt gcagcgcac 600
gccgacgacc tgacggccca ggcggcgccc gacgggcagc cgtgggacaa gctggtcacc 660
cgtggctcgg acggccaggt gctgcccgtc gtctcgcgc agaacctgat ggcgcccgtac 720
ttcgaccggc ccgacgagat gccgttcgg gacctgttcg cggcccagat cgacgaggtc 780
tgggagaagt accgctccac cgacctggg atcgacctc agggcgggcg gggcacctg 840
gcccggccgg tcagcgggga cacgctgacc ttcgagggcg gacacacct ctccaagccc 900
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gacgacaaga aggcgctgct ggcacggatc gggcggggt tcaaccggtc gatcatgctg 1020
agccacccca gccagccgaa cggcacctcg gtggcggact actaccagga cgcggtgacc 1080
aaccactggt cgcgggtcgt ccacgcgaac tccccatcg ggtacgcgtt cccgtacgac 1140
gacgtacgcc ccgacggtga gccggacgtc tggggcgcg cgaacgacgg caacccccgg 1200
cgcttcacgg tgagcgtggg ttctga 1227

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<210> SEQ ID NO 30
<211> LENGTH: 870
<212> TYPE: DNA
<213> ORGANISM: Streptomyces sp. ACTE

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<400> SEQUENCE: 30
gtgcttcacc cccacaaccg caccgcacgt cgcaccactc ggctcaccgc cacggcggt 60
ctcgccgcgc cggccctcgg gctcgcgctc atggcgctcc ccgtcaccgc tcaccggcg 120
gccccacgc agccggcgcg tcatcatctg gaggcgcgc cgaccggact ggaagatccc 180
gcgaagaagg acatcgccat gcagttggtc tccagcgcgg agaactccac gctggactgg 240
aaggcgcagt acggctacat cgaggacatc ggcgacggac gcggctacac cgcggcacc 300

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atcggcttct gctcggggac cggagacatg ctgcacctgg tgcagcgcta cacggaccgc	360
tcaccgggca acgtactggc gtcgtacctg cccgccctgc gcgaggtcga cgggaccgac	420
tcgcacgacg ggctcgaccc cggcttcccc cgggactggg ccgagggcgc gaaggaccgc	480
gtgttcacgc aggcgcagaa cgacgagcgg gaccgggtgt acttcgaccc ggcgggtgcg	540
caggccaagg acgacgggct ggggacgctc ggccagtctc cgtactacga cgccatcgtc	600
atgcacggag gcggcgggga cagcacgagc ttcgggtcca tccggcagcg cgcgctcgcg	660
gaggcggaac cgccctcgcg gggcgggtgac gaggtcgctt acctcgacgc gttcctggac	720
gcgcgggtct gggcgatgcg gcaggaggag gcccaactcg acaccagccg ggtcgacacc	780
gcgcagcgcg tcttctcgcg cgacgggaat ctgaacctgg atccgcccgt ggactggcag	840
gtgtacggcg acagcttcca catcggctga	870

<210> SEQ ID NO 31

<211> LENGTH: 2373

<212> TYPE: DNA

<213> ORGANISM: Streptomyces sp. ACTE

<400> SEQUENCE: 31

atgacccac cgcacagaca ccgcctgttc aggcgctcgg tgtccgcttc cctctcgctg	60
gccttcaccg ccgtcggcac cgcgcgcgcg gtcgtcctgg ccggtgcccc ggcggcccag	120
gccgcccggg tccccgcacc ctccccggtc ggcatatccg gccggggcgc cgcctgcccg	180
ttcacggagc agggaggcca gtacgcgcgc accaacggca cgctcatcgg cccggaccgg	240
cgctacggct cactgccttc ggaggcgtcc ggcggcagc ccgtcacgct cgacgcggcc	300
ggtgagtagc tggagttcac cctcacgcc cccgccaacg cgatgacctt ccgtattcgc	360
ctgccggaca acgcccgcgg gacgggcccg gacgcctctc tcgacctgcg ggtgaacggc	420
tcggctctca agagcgtgcc ggtgacctcg aagtaaggct ggtactacgg gggttacccc	480
ttcaacaaca accccgggga caccaaccg caccatttct acgacgagac ccggaccatg	540
ttcggctcga ccctgcccgc cggtaacgaag gtcgggtgct aggtggcgtc caccgcccgc	600
tcgccctcgt tcaccgtcga cctggccgac ttcgagcagg tggccgcgcc cgtcggcaag	660
ccgtccggcg cactggaagt ggtgagcgac ttcggggccg acccgaccgg ggcggccgac	720
tccaccgcca agatccaggc ggcggctgac gcggggcgc cccagggcaa ggtcgtctac	780
atcccgcagg ggaacctcca ggtgctgac cacatcgtcg tggaccaggc gacgctgcg	840
ggcgcggccc cctggtacag cgtgctgac gggcgtcacc ccacggaccg gagcaaggcg	900
gtcgggtgct acgggaagta ctcggcgcag ggcggcagca ggaacgtcac cctcaaggac	960
ttcggcatca tcggcgacat ccaggagcgt gtcggacaac accaggtcaa cgccatcggc	1020
ggggccatgt ccgactcggc cgtcgacaac gtctggatgc agcacaccaa gtgoggcgcc	1080
tggatggacg gcccgatgga caatttcacc atcaagaaca gtcgcatcct ggaccagacc	1140
gcggacggcg tgaacttcca ctacggggtc acgaactcga ccgtcacgaa caccttcgtc	1200
cgcaacaccg gtgacgacgg cctggccatg tgggcggaga acgtcccga cgtgaagaac	1260
aagttcacgt tcaacacggt gatcctgccg atcctggcca acaacatcgt gacgtacggc	1320
ggcaaggaca tcacgatctc cgacaacgct atggcggaca ccatcaccaa cggcggcggg	1380
ctgcacatcg ccaaccgcta cccgggcgtc aactcggggc aggggacggc cgtcgcgggg	1440
acgcacacgg ccgcgcgcaa caccctgatc cgtaccgca acagcgactt caactggaac	1500

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ttcggcgctcg gggcgatctg gttcagcggg ctcaacgaac cgatcagcaa cgccaccatc	1560
aacatcaccg acagcggagt cctggacagc tctacgccc cgatccacct gatcgagggt	1620
gcgagcaacg ggctgcactt caagaacgtc aagatcgacg gggcgggtac ctacgcctg	1680
cagatccagg caccgggac ggccacctc gagaacgctg tggccacca catcgcccag	1740
tccaaccga tccacaactg tgtcggcagc ggcttcaga tccccggg cagcggcaac	1800
tccggctggt acgcccacc gcccgctgc accggggtct ggcccgacc ggtgtggacc	1860
aacggcggcg tccccggagg cggcgggtcc accaaccga ccgacccac cgaccccacc	1920
gacccgacgg accccaccga cccgcctgag gagacgggca acctcgccc gggacgcacc	1980
gtcaccgaga ccagccacac ggacgtgtac ggcggggcca acacgtoga cggcaacgcg	2040
gacacgtact gggagagccg caacaacgcc tccccgagt ccgtcacctg cgacctggc	2100
gtgccaagg cggtgaagcg ggtggtgctg aagctcccgc cggccgcgcg gtggcgacc	2160
cgcaecgaga cgctctcctg gtcggcagc accgacaacg ggacgtaaa ctcgctgaag	2220
gcgtcggcgg gttacacctt caaccgctc agcggcaaca ccgcgacggt ctccctccc	2280
gggacgccgg tccggtacct gcggctgacc ttcaccaga acaccgggtg gcccgccgc	2340
cagctgtccg aactggaggc ctacaccagc tga	2373

<210> SEQ ID NO 32

<211> LENGTH: 1545

<212> TYPE: DNA

<213> ORGANISM: Streptomyces sp. ACTE

<400> SEQUENCE: 32

atgaggagac cagtcgcct gcgactcagc gcggcggggg ccaccctggc cctggtgcc	60
gcgaccggcg cactgatggc gatgccgag gcggcgtcgg cagcgcacgg cggcgtcacc	120
ggatacgcga ccagaaacgg cggcaccacc ggcggcggcg gcgggcagac ggtgcggggc	180
accaccggga ccgcgatcca gcggccctg tgcggggcgg ccagcagctc cacccectc	240
accatccagg tcgaggggac catcaaccac ggcaaacccg acaaggtctc gggcagcagc	300
tgcaaacccg ccgcccggagt catcgagctg aagcagatca gcaacgtcac gatcgtcggc	360
gtggcggcgg gcgcccctt cgaccaagta ggcattccag tccgcgagtc cagcaacatc	420
atcatccaga acgtcaccgt caagaacgtc aagaagtccg gctcggcccac gtccaacggc	480
ggtgacgcca tcggcatgga gaaggacgtc cgcaacgtct ggggtgacca caccacctg	540
gaggcctcgg gcggcgagtc ggagggcttc gacggcctct tcgacatgaa ggccggcacc	600
cagtacgtga cgctgtccta cagcatctc gcgaaactcc gccggggagg cctcgtcggc	660
tccagcgaga gcgacctctc gaacggcttc atcacctacc accacaacct gtacgagaac	720
atcgactccc gcgcccctct gctgcggggc ggcgtcggcc acatctacaa caaccctac	780
gtgggactca gcaagtggg catcaactcc cgggcccggc cccgcgcca ggtggacaac	840
aactacttcg aggactccaa ggacgtctc ggcacctct acaccgacgc ggccggctac	900
tggcaggtca gcgcaacgt cttcgacaac gtgacgtggt ccggcccag cagcgacaac	960
aaccccggcg gcccggacc gcagtcaca acctcggta gcatccccta cgctacacc	1020
ctcgacgggg cgaactcgt accgtccgtc gtgagccgga cggcggggc gaacacgggg	1080
ctgaagggtg cggacggcag ctgctcggcg cagacggcgg acccgacga ccccacccc	1140
gacccgacgc cggaccggac cgaccccact cggcccacc ggaccaacct cagcctcggg	1200
gccggctcgg acggtccag caaggcagc gggaccagct acggcgacgt gcgggacggt	1260

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gacatgagca cctactggtc accgtccggc tegaccgggtt ccgtctcgat caagtgggagc 1320
tccgccacca ccgtctccaa gatcaacgtg cgcgaggcgg cgggctccac gggtccatc 1380
acctctgga aggtcggcaa cgcgcacacc ggcgcgctcc tggcctccgg cagcggggcg 1440
ggcgtcatca cgttcccgca gacctcgctg cgcaagatca cgttcgagat caccgggctcg 1500
acgggcacgc cgaaggtcgc cgagttcgag acgtacgccc gctga 1545

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<210> SEQ ID NO 33

<211> LENGTH: 389

<212> TYPE: PRT

<213> ORGANISM: Streptomyces sp. ACTE

<400> SEQUENCE: 33

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Met Pro Glu Arg Phe Thr Pro Thr Pro Glu Asp Lys Phe Thr Phe Gly
 1          5          10          15
Leu Trp Thr Val Gly Trp Arg Gly Asn Asp Pro Phe Gly Glu Pro Thr
          20          25          30
Arg Pro Val Leu Asp Pro Val Glu Ser Val Glu Arg Leu Ala Glu Leu
          35          40          45
Gly Ala His Gly Val Thr Phe His Asp Asp Asp Leu Ile Pro Phe Gly
          50          55          60
Ser Asp Asp Arg Glu Arg Ala Arg Leu Val Gly Arg Phe Arg Glu Ala
 65          70          75          80
Leu Glu Arg Thr Gly Leu Lys Val Pro Met Ala Thr Thr Asn Leu Phe
          85          90          95
Thr His Pro Val Phe Lys Asp Gly Gly Phe Thr Ser Asn Asp Arg Asp
          100          105          110
Val Arg Arg Phe Ala Leu Arg Lys Val Ile Arg Asn Ile Asp Leu Ala
          115          120          125
Val Glu Leu Gly Ala Gln Thr Tyr Val Ala Trp Gly Gly Arg Glu Gly
          130          135          140
Ala Glu Ser Gly Ala Ala Lys Asp Val Arg Ser Ala Leu Asp Arg Met
 145          150          155          160
Lys Glu Ala Phe Asp Leu Leu Gly Asp Tyr Val Thr Glu Gln Gly Tyr
          165          170          175
Asp Leu Arg Phe Ala Ile Glu Pro Lys Pro Asn Glu Pro Arg Gly Asp
          180          185          190
Ile Leu Leu Pro Thr Ile Gly His Ala Leu Ala Phe Ile Glu Arg Leu
          195          200          205
Glu Arg Pro Glu Leu Val Gly Val Asn Pro Glu Thr Gly His Glu Gln
          210          215          220
Met Ala Gly Leu Asn Phe Pro His Gly Ile Ala Gln Ala Leu Trp Ala
 225          230          235          240
Gly Lys Leu Phe His Ile Asp Leu Asn Gly Gln Ser Gly Ile Lys Tyr
          245          250          255
Asp Gln Asp Phe Arg Phe Gly Ala Gly Asp Leu Arg Gln Ala Phe Trp
          260          265          270
Leu Val Asp Leu Leu Glu Thr Ala Gly Trp Asp Gly Ser Arg His Phe
          275          280          285
Asp Phe Lys Pro Val Arg Thr Asp Gly Ile Asp Gly Val Trp Glu Ser
          290          295          300
Ala Lys Asn Cys Met Arg Asn Tyr Leu Ile Leu Lys Glu Arg Ala Ala
 305          310          315          320

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Val Lys Asn Leu Val Thr Ser Gly Ser Ala Glu Lys Ile Thr His Ile
 305 310 315 320
 Asn Tyr Ala Phe Gly Asn Val Gln Gly Gly Lys Cys Thr Ile Gly Asp
 325 330 335
 Ser Tyr Ala Asp Tyr Asp Lys Ala Tyr Thr Ala Asp Gln Ser Val Asp
 340 345 350
 Gly Val Ala Asp Thr Trp Asp Gln Pro Leu Arg Gly Asn Phe Asn Gln
 355 360 365
 Leu Arg Lys Leu Lys Ala Lys Tyr Pro His Ile Lys Val Ile Trp Ser
 370 375 380
 Phe Gly Gly Trp Thr Trp Ser Gly Gly Phe Gly Ala Ala Ala Gln Asn
 385 390 395 400
 Pro Ala Ala Phe Ala Gln Ser Cys Tyr Asp Leu Val Glu Asp Pro Arg
 405 410 415
 Trp Ala Asp Val Phe Asp Gly Ile Asp Ile Asp Trp Glu Tyr Pro Asn
 420 425 430
 Ala Cys Gly Leu Thr Cys Asp Thr Ser Gly Pro Ala Ala Leu Lys Asn
 435 440 445
 Leu Ser Ser Ala Leu Arg Ala Lys Phe Gly Ala Lys Asn Leu Val Thr
 450 455 460
 Ala Ala Ile Thr Ala Asp Gly Ser Asp Gly Gly Lys Ile Asp Ala Ala
 465 470 475 480
 Asp Tyr Ala Gly Ala Ala Gln Ser Phe Asp Trp Tyr Asn Val Met Thr
 485 490 495
 Tyr Asp Phe Phe Gly Ala Trp Glu Ala Lys Gly Pro Thr Ala Pro His
 500 505 510
 Ser Pro Leu Asn Ala Tyr Ala Gly Ile Pro Gln Asp Gly Phe Asn Ser
 515 520 525
 Ala Ala Ala Ile Ala Lys Leu Lys Ala Lys Gly Val Pro Ala Ser Lys
 530 535 540
 Leu Leu Leu Gly Ile Gly Phe Tyr Gly Arg Gly Trp Thr Gly Val Thr
 545 550 555 560
 Gln Ala Ala Pro Gly Gly Thr Ala Thr Gly Ala Ala Pro Gly Thr Tyr
 565 570 575
 Glu Ala Gly Ile Glu Asp Tyr Lys Val Leu Lys Thr Ser Cys Pro Ala
 580 585 590
 Thr Gly Thr Ile Ala Gly Thr Ala Tyr Ala His Cys Gly Thr Asn Trp
 595 600 605
 Trp Ser Tyr Asp Thr Pro Ala Thr Ile Thr Ser Lys Met Ala Trp Ala
 610 615 620
 Asn Ser Gln Gly Leu Gly Gly Ala Phe Phe Trp Glu Phe Ser Gly Asp
 625 630 635 640
 Thr Ala Asn Gly Glu Leu Val Ser Ala Met Asp Ser Gly Leu Asn
 645 650 655

<210> SEQ ID NO 35

<211> LENGTH: 196

<212> TYPE: PRT

<213> ORGANISM: Streptomyces sp. ACTE

<400> SEQUENCE: 35

Met Arg Lys Arg Ala Ser Ala Ala Val Ile Gly Leu Ala Ile Ala Gly
 1 5 10 15

Val Ser Met Phe Ala Thr Ser Ser Ala Ser Ser His Gly Tyr Thr Asp

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20				25				30							
Ser	Pro	Ile	Ser	Arg	Gln	Lys	Leu	Cys	Ala	Asn	Gly	Thr	Val	Thr	Gly
	35					40					45				
Cys	Gly	Asn	Ile	Gln	Trp	Glu	Pro	Gln	Ser	Val	Glu	Gly	Pro	Lys	Gly
	50				55						60				
Phe	Pro	Ala	Ala	Gly	Pro	Ala	Asp	Gly	Lys	Ile	Cys	Ala	Gly	Gly	Asn
65					70					75					80
Ser	Ser	Phe	Ala	Ala	Leu	Asp	Asp	Pro	Arg	Gly	Gly	Asn	Trp	Pro	Ala
			85						90					95	
Thr	Gln	Val	Thr	Gly	Gly	Gln	Gly	Tyr	Asn	Phe	Arg	Trp	Gln	Phe	Thr
			100						105					110	
Ala	Arg	His	Ala	Thr	Thr	Asp	Phe	Arg	Tyr	Tyr	Ile	Thr	Lys	Asp	Gly
			115				120							125	
Trp	Asp	Ser	Thr	Lys	Pro	Leu	Thr	Arg	Ala	Ala	Leu	Glu	Ser	Gln	Pro
	130					135					140				
Phe	Met	Thr	Val	Pro	Tyr	Gly	Asn	Gln	Gln	Pro	Pro	Ala	Thr	Leu	Thr
145					150					155					160
His	Gln	Gly	Thr	Ile	Pro	Thr	Gln	Lys	Ser	Gly	Lys	His	Ile	Ile	Leu
			165						170					175	
Ala	Val	Trp	Asn	Val	Ala	Asp	Thr	Ala	Asn	Ala	Phe	Tyr	Ala	Cys	Ser
			180						185					190	
Asp	Val	Lys	Phe												
		195													

<210> SEQ ID NO 36

<211> LENGTH: 556

<212> TYPE: PRT

<213> ORGANISM: Streptomyces sp. ACTE

<400> SEQUENCE: 36

Val	Ala	Ala	Leu	Ala	Ala	Gly	Ala	Leu	Thr	Val	Thr	Gly	Leu	Val	Gly
1					5					10				15	
Thr	Ala	Gln	Ala	Ala	Asp	Ile	Asn	Val	Ala	Lys	Asn	Ala	Gly	Phe	Glu
			20				25						30		
Ser	Gly	Leu	Ser	Gly	Trp	Thr	Cys	Thr	Gly	Gly	Ser	Gly	Ala	Thr	Val
		35					40						45		
Ser	Ser	Pro	Val	His	Gly	Gly	Ser	Ala	Ala	Leu	Lys	Ala	Thr	Pro	Ser
		50				55					60				
Gly	Gln	Asp	Asn	Ala	Lys	Cys	Thr	Gln	Thr	Val	Ala	Val	Lys	Pro	Asn
65					70					75					80
Ser	Thr	Tyr	Ala	Leu	Ser	Ser	Trp	Val	Gln	Gly	Gly	Tyr	Ala	Tyr	Leu
			85						90					95	
Gly	Ala	Ser	Gly	Thr	Gly	Thr	Thr	Asp	Val	Ser	Thr	Trp	Thr	Pro	Gly
			100						105					110	
Ser	Thr	Gly	Trp	Thr	Gln	Leu	Arg	Thr	Ser	Phe	Thr	Thr	Gly	Pro	Ser
			115				120							125	
Thr	Thr	Ser	Val	Gln	Val	Tyr	Thr	His	Gly	Trp	Tyr	Gly	Gln	Ala	Ala
			130			135								140	
Tyr	Tyr	Ala	Asp	Asp	Val	Ala	Val	Thr	Gly	Pro	Asp	Gly	Gly	Gly	Gly
145					150					155					160
Thr	Glu	Glu	Pro	Gly	Pro	Ala	Ile	Pro	Gly	Ala	Pro	Ala	Gly	Leu	Ala
					165					170				175	
Val	Gly	Thr	Thr	Thr	Ser	Ser	Ser	Val	Ala	Leu	Ser	Trp	Asn	Ala	Val
			180							185				190	

-continued

Ser Gly Ala Thr Gly Tyr Thr Val Tyr Arg Asp Gly Thr Lys Ala Thr
 195 200 205

Thr Thr Thr Gly Thr Ser Ala Thr Val Ser Gly Leu Ala Ala Asp Thr
 210 215 220

Ala Tyr Gln Phe Ser Val Ser Ala Thr Asn Ala Ala Gly Glu Ser Val
 225 230 235 240

Arg Ser Ala Thr Val Ser Gly Arg Thr Ala Lys Lys Asp Glu Thr Gly
 245 250 255

Pro Gly Pro Ser Thr Ser Val Pro Lys His Ala Val Thr Gly Tyr Trp
 260 265 270

Gln Asn Phe Asn Asn Gly Ala Ala Val Gln Lys Leu Ser Asp Val Pro
 275 280 285

Ala Asn Tyr Asp Ile Ile Ala Val Ser Phe Ala Asp Ala Ala Gly Thr
 290 295 300

Pro Gly Ala Val Thr Phe Asn Leu Asp Ser Ala Gly Leu Asn Gly Tyr
 305 310 315 320

Thr Val Ala Gln Phe Lys Ala Asp Ile Lys Ala Lys Gln Ala Ala Gly
 325 330 335

Lys Asn Val Ile Ile Ser Val Gly Gly Glu Lys Gly Thr Val Ser Val
 340 345 350

Asn Ser Asp Ala Ser Ala Asn Ala Phe Ala Asp Ser Leu Tyr Thr Leu
 355 360 365

Ile Gln Glu Tyr Gly Phe Asn Gly Val Asp Ile Asp Leu Glu Asn Gly
 370 375 380

Leu Asn Ser Thr Tyr Met Thr Lys Ala Leu Arg Ser Leu Ser Ser Lys
 385 390 395 400

Val Gly Ser Gly Leu Val Ile Thr Met Ala Pro Gln Thr Ile Asp Met
 405 410 415

Gln Ser Thr Ser Gly Glu Tyr Phe Lys Thr Ala Leu Asn Ile Lys Asp
 420 425 430

Ile Leu Thr Val Val Asn Met Gln Tyr Tyr Asn Ser Gly Ser Met Leu
 435 440 445

Gly Cys Asp Gly Lys Val Tyr Ser Gln Gly Ser Val Asp Phe Leu Thr
 450 455 460

Ala Leu Ala Cys Ile Gln Leu Glu Gly Gly Leu Ala Pro Ser Gln Val
 465 470 475 480

Gly Leu Gly Val Pro Ala Ser Thr Arg Gly Ala Gly Ser Gly Tyr Val
 485 490 495

Ala Pro Ser Val Val Asn Ala Ala Leu Asp Cys Leu Ala Lys Gly Thr
 500 505 510

Gly Cys Gly Ser Phe Lys Pro Ser Arg Thr Tyr Pro Asp Ile Arg Gly
 515 520 525

Ala Met Thr Trp Ser Thr Asn Trp Asp Ala Thr Ala Gly Asn Ala Trp
 530 535 540

Ser Asn Ala Val Gly Pro His Val His Gly Leu Pro
 545 550 555

<210> SEQ ID NO 37
 <211> LENGTH: 295
 <212> TYPE: PRT
 <213> ORGANISM: Streptomyces sp. ACTE

<400> SEQUENCE: 37

Val Ile Arg Arg Val Met Gly Leu Leu Thr Ala Leu Ala Ala Val Val
 1 5 10 15

-continued

Ala Thr Leu Val Phe Leu Pro Ala Ala Thr Ala Ser Ala Ala Thr Cys
 20 25 30
 Ala Pro Ala Trp Asn Ala Ser Ser Val Tyr Thr Gly Gly Gly Ser Ala
 35 40 45
 Ser Tyr Asn Gly His Asn Trp Ser Ala Lys Trp Trp Thr Gln Asn Glu
 50 55 60
 Arg Pro Gly Thr Ser Asp Val Trp Ala Asp Gln Gly Ala Cys Gly Ser
 65 70 75 80
 Gly Gly Gly Gly Thr Asp Pro Asn Pro Ser Gly Phe Val Val Ser Glu
 85 90 95
 Ala Gln Phe Asn Gln Met Phe Pro Ser Arg Asn Ser Phe Tyr Thr Tyr
 100 105 110
 Ser Gly Leu Thr Ala Ala Leu Ser Ala Tyr Pro Ala Phe Ala Asn Thr
 115 120 125
 Gly Ser Asp Thr Val Lys Lys Gln Glu Ala Ala Ala Phe Leu Ala Asn
 130 135 140
 Val Ser His Glu Thr Gly Gly Leu Val His Ile Val Glu Gln Asn Thr
 145 150 155 160
 Ala Asn Tyr Pro His Tyr Cys Asp Thr Ser Gln Ser Tyr Gly Cys Pro
 165 170 175
 Ala Gly Gln Ala Ala Tyr Tyr Gly Arg Gly Pro Ile Gln Leu Ser Trp
 180 185 190
 Asn Phe Asn Tyr Lys Ala Ala Gly Asp Ala Leu Gly Ile Asp Leu Leu
 195 200 205
 Gly Asn Pro Trp Gln Val Glu Gln Asn Ala Ser Val Ala Trp Lys Thr
 210 215 220
 Gly Leu Trp Tyr Trp Asn Thr Gln Ser Gly Pro Gly Thr Met Thr Pro
 225 230 235 240
 His Asn Ala Ile Val Asn Gly Ser Gly Phe Gly Glu Thr Ile Arg Ser
 245 250 255
 Ile Asn Gly Ser Ile Glu Cys Asn Gly Gly Asn Pro Gly Gln Val Gln
 260 265 270
 Ser Arg Val Asn Thr Tyr Gln Ser Phe Val Gln Ile Leu Gly Thr Thr
 275 280 285
 Pro Gly Ser Asn Leu Ser Cys
 290 295

<210> SEQ ID NO 38

<211> LENGTH: 507

<212> TYPE: PRT

<213> ORGANISM: Streptomyces sp. ACTE

<400> SEQUENCE: 38

Met Arg Arg Ser Arg Ser Val Arg Ala Leu Val Thr Ala Ala Val Thr
 1 5 10 15
 Thr Val Ala Ala Ala Gly Met Ala Val Leu Gly Ser Gly Thr Ala Gln
 20 25 30
 Ala Ala Thr Pro Leu Pro Asp His Val Phe Ala Pro Tyr Phe Glu Ser
 35 40 45
 Trp Thr Gly Glu Ser Pro Ala Ala Met Ala Ala Glu Ser Gly Ala Lys
 50 55 60
 His Leu Thr Met Ala Phe Leu Gln Thr Thr Ala Lys Gly Ser Cys Thr
 65 70 75 80
 Pro Tyr Trp Asn Gly Asp Thr Gly Leu Pro Ile Ala Gln Ala Ser Phe

-continued

85				90				95							
Gly	Ala	Asp	Ile	Asp	Thr	Ile	Gln	Ala	Gly	Gly	Gly	Asp	Val	Ile	Pro
			100								105				110
Ser	Phe	Gly	Gly	Tyr	Thr	Ala	Asp	Thr	Thr	Gly	Thr	Glu	Ile	Ala	Asp
			115				120								125
Ser	Cys	Thr	Asp	Val	Asp	Gln	Ile	Ala	Ala	Ala	Tyr	Gln	Lys	Val	Val
			130				135				140				
Thr	Thr	Tyr	Asp	Val	Ser	Arg	Leu	Asp	Met	Asp	Ile	Glu	Val	Asp	Ser
			145				150				155				160
Leu	Asp	Asp	Thr	Ala	Gly	Ile	Asp	Arg	Arg	Asn	Lys	Ala	Ile	Lys	Lys
			165								170				175
Leu	Gln	Asp	Trp	Ala	Asp	Ala	Asn	Gly	Arg	Asp	Leu	Glu	Ile	Ser	Tyr
			180								185				190
Thr	Leu	Pro	Thr	Thr	Thr	Arg	Gly	Leu	Ala	Ser	Ser	Gly	Leu	Ala	Val
			195				200								205
Leu	Arg	Asn	Ala	Val	Thr	Asn	Gly	Ala	Arg	Val	Asp	Val	Val	Asn	Leu
			210				215				220				
Met	Thr	Phe	Asp	Tyr	Tyr	Asp	Asn	Ala	Ser	His	Asp	Met	Ala	Ala	Asp
			225				230				235				240
Thr	Glu	Thr	Ala	Ala	Gln	Gly	Leu	Tyr	Asp	Gln	Leu	Ala	Lys	Leu	Tyr
			245								250				255
Pro	Gly	Arg	Thr	Ala	Thr	Gln	Leu	Trp	Ser	Met	Val	Gly	Val	Thr	Glu
			260								265				270
Met	Pro	Gly	Val	Asp	Asp	Phe	Gly	Pro	Ala	Glu	Thr	Phe	Thr	Leu	Ala
			275				280								285
Asn	Ala	Ala	Arg	Val	Tyr	Asp	Trp	Ala	Val	Ala	Lys	Gly	Ile	Asn	Thr
			290				295				300				
Leu	Ser	Phe	Trp	Ala	Leu	Gln	Arg	Asp	Asn	Gly	Gly	Cys	Pro	Gly	Gly
			305				310				315				320
Pro	Ala	Ala	Asp	Asp	Cys	Ser	Gly	Ile	Gln	Gln	Asn	Thr	Trp	Asp	Phe
			325								330				335
Thr	Arg	Val	Phe	Ala	Pro	Phe	Thr	Ser	Gly	Thr	Thr	Ala	Pro	Asp	Asp
			340								345				350
Asp	Phe	Ser	Val	Thr	Ala	Thr	Pro	Ala	Ser	Gly	Thr	Val	Thr	Ala	Gly
			355				360								365
Gly	Ser	Ala	Thr	Thr	Thr	Val	Lys	Thr	Ala	Val	Thr	Lys	Gly	Ala	Ala
			370				375				380				
Gln	Gln	Val	Gly	Leu	Thr	Val	Ser	Gly	Val	Pro	Ala	Gly	Val	Thr	Ala
			385				390				395				400
Ser	Leu	Ser	Pro	Ser	Ser	Val	Thr	Ala	Gly	Gly	Arg	Ser	Thr	Leu	Thr
			405								410				415
Leu	Ala	Thr	Thr	Gln	Ala	Ala	Val	Ser	Gly	Thr	Tyr	Arg	Ile	Ser	Val
			420								425				430
Thr	Gly	Thr	Ser	Pro	Ser	Gly	Ser	His	Ala	Thr	Ala	Tyr	Thr	Leu	Thr
			435				440								445
Val	Thr	Gly	Gly	Thr	Gly	Ser	Gln	Cys	Thr	Ala	Gly	Pro	Trp	Ala	Gly
			450				455				460				
Gly	Thr	Val	Tyr	Thr	Gly	Gly	Gln	Gln	Val	Ser	Tyr	Lys	Gly	His	Thr
			465				470				475				480
Trp	Lys	Ala	Lys	Trp	Trp	Thr	Thr	Gly	Glu	Glu	Pro	Gly	Thr	Thr	Gly
			485								490				495
Glu	Trp	Gly	Val	Trp	Gln	Asp	Leu	Gly	Ala	Cys					
			500								505				

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<210> SEQ ID NO 39
 <211> LENGTH: 483
 <212> TYPE: PRT
 <213> ORGANISM: Streptomyces sp. ACTE
 <400> SEQUENCE: 39

Val Thr Gln Gly Pro Leu Thr Thr Glu Ala Gly Ala Pro Val Ala Asp
 1 5 10 15
 Asn Gln Asn Ser Glu Thr Ala Gly Pro Gly Gly Pro Val Leu Val Gln
 20 25 30
 Asp Gln Ala Leu Leu Glu Lys Leu Ala His Phe Asn Arg Glu Arg Ile
 35 40 45
 Pro Glu Arg Val Val His Ala Arg Gly Ala Gly Ala Tyr Gly Thr Phe
 50 55 60
 Thr Leu Thr Arg Asp Val Ser Gln Trp Thr Arg Ala Lys Phe Leu Ser
 65 70 75 80
 Glu Val Gly Lys Glu Thr Glu Thr Phe Leu Arg Phe Ser Thr Val Ala
 85 90 95
 Gly Asn Leu Gly Ser Ala Asp Ala Ala Arg Asp Pro Arg Gly Trp Ala
 100 105 110
 Leu Lys Phe Tyr Thr Glu Glu Gly Asn Tyr Asp Leu Val Gly Asn Asn
 115 120 125
 Thr Pro Val Phe Phe Ile Lys Asp Ala Ile Lys Phe Pro Asp Phe Ile
 130 135 140
 His Thr Gln Lys Arg Asp Pro Tyr Thr Gly Ser Gln Glu Ala Asp Asn
 145 150 155 160
 Val Trp Asp Phe Trp Gly Leu Ser Pro Glu Ser Thr His Gln Val Thr
 165 170 175
 Trp Leu Phe Gly Asp Arg Gly Ile Pro Ala Ser Phe Arg His Met Asn
 180 185 190
 Gly Tyr Gly Ser His Thr Phe Gln Trp Asn Asn Glu Ala Gly Glu Val
 195 200 205
 Phe Trp Val Lys Tyr His Phe Lys Thr Asp Gln Gly Ile Lys Asn Leu
 210 215 220
 Thr Thr Glu Glu Ala Val Arg Leu Ser Gly Val Asp Pro Asp Ser His
 225 230 235 240
 Gln Arg Asp Leu Arg Glu Ser Ile Glu Arg Gly Asp Phe Pro Thr Trp
 245 250 255
 Thr Val Gln Val Gln Ile Met Pro Ala Ala Glu Ala Ala Thr Tyr Arg
 260 265 270
 Phe Asn Pro Phe Asp Leu Thr Lys Val Trp Pro His Glu Asp Tyr Pro
 275 280 285
 Pro Ile Glu Ile Gly Lys Leu Glu Leu Asn Arg Asn Pro Glu Asn Ile
 290 295 300
 Phe Ala Glu Val Glu Gln Ser Ile Phe Ser Pro Ala His Phe Val Pro
 305 310 315 320
 Gly Ile Gly Pro Ser Pro Asp Lys Met Leu Gln Gly Arg Leu Phe Ala
 325 330 335
 Tyr Gly Asp Ala His Arg Tyr Arg Val Gly Ile Asn Ala Asp His Leu
 340 345 350
 Pro Val Asn Arg Pro His Ala Thr Glu Ala Arg Thr Asn Ser Arg Asp
 355 360 365
 Gly Tyr Leu Tyr Asp Gly Arg His Lys Gly Thr Lys Asn Tyr Glu Pro

-continued

370					375					380					
Asn	Ser	Phe	Gly	Gly	Pro	Val	Gln	Thr	Asp	Arg	Pro	Leu	Trp	Gln	Pro
385					390					395					400
Val	Ser	Val	Thr	Gly	Gly	Thr	Gly	Asn	His	Glu	Ala	Ala	Val	His	Ala
				405					410					415	
Glu	Asp	Asn	Asp	Phe	Val	Gln	Ala	Gly	Asn	Leu	Tyr	Arg	Leu	Met	Ser
			420						425					430	
Glu	Asp	Glu	Lys	Gly	Arg	Leu	Ile	Asp	Asn	Leu	Ala	Gly	Phe	Ile	Ala
		435					440						445		
Lys	Val	Ser	Arg	Asp	Asp	Ile	Ala	Asp	Arg	Ala	Ile	Asn	Asn	Phe	Arg
	450					455							460		
Gln	Ala	Asp	Ala	Asp	Phe	Gly	Lys	Arg	Leu	Glu	Val	Ala	Val	Gln	Ala
465					470					475					480
Leu	Arg	Gly													
<210> SEQ ID NO 40															
<211> LENGTH: 926															
<212> TYPE: PRT															
<213> ORGANISM: Streptomyces sp. ACTE															
<400> SEQUENCE: 40															
Val	Tyr	Ala	Met	Pro	Ser	Thr	Ala	Pro	Ala	Ala	Val	Gln	Ser	Gly	Glu
1				5					10					15	
Asp	Ala	Pro	Val	Arg	Ser	Ser	Pro	Arg	Pro	Phe	Ala	Ala	Leu	Leu	Ala
			20					25					30		
Ala	Leu	Ala	Leu	Thr	Ala	Gly	Leu	Ser	Leu	Ile	Gly	Thr	Pro	Ala	Val
		35					40					45			
Ala	Arg	Ser	Asp	Glu	Ala	Pro	Ala	Ala	Thr	Glu	Ala	Ser	Asp	Val	Ser
	50					55					60				
Ile	Ala	Ala	Asp	Thr	Tyr	Thr	Trp	Lys	Asn	Ala	Arg	Ile	Asp	Gly	Gly
65					70					75				80	
Gly	Phe	Val	Pro	Gly	Ile	Val	Phe	Asn	Arg	Ser	Glu	Lys	Asn	Leu	Ala
				85					90					95	
Tyr	Ala	Arg	Thr	Asp	Ile	Gly	Gly	Ala	Tyr	Arg	Trp	Asp	Gln	Ser	Gly
			100					105					110		
Lys	Gln	Trp	Lys	Pro	Leu	Leu	Asp	Trp	Val	Asp	Trp	Asp	Arg	Trp	Gly
		115					120					125			
Trp	Thr	Gly	Val	Val	Ser	Leu	Ala	Ser	Asp	Thr	Val	Asp	Pro	Asp	Asn
	130					135					140				
Val	Tyr	Ala	Ala	Val	Gly	Thr	Tyr	Thr	Asn	Ser	Trp	Asp	Pro	Thr	Asp
145					150					155					160
Gly	Ala	Val	Leu	Arg	Ser	Ser	Asp	Arg	Gly	Ala	Ser	Trp	Lys	Ala	Ala
				165					170					175	
Thr	Leu	Pro	Phe	Lys	Leu	Gly	Gly	Asn	Met	Pro	Gly	Arg	Gly	Met	Gly
			180					185					190		
Glu	Arg	Leu	Ala	Val	Asp	Pro	Asn	Lys	Asn	Ser	Val	Leu	Tyr	Leu	Gly
		195					200					205			
Ala	Pro	Ser	Gly	Asn	Gly	Leu	Trp	Arg	Ser	Thr	Asp	Ala	Gly	Val	Ser
	210					215					220				
Trp	Ser	Glu	Val	Thr	Ala	Phe	Pro	Asn	Pro	Gly	Asn	Tyr	Ala	Gln	Asp
225					230					235					240
Pro	Ser	Asp	Thr	Ser	Gly	Tyr	Gly	Asn	Asp	Asn	Gln	Gly	Ile	Val	Trp
				245					250					255	
Val	Thr	Phe	Asp	Glu	Arg	Ser	Gly	Ser	Ala	Gly	Ser	Ala	Thr	Gln	Asp

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260					265					270					
Ile	Tyr	Val	Gly	Val	Ala	Asp	Lys	Glu	Asn	Thr	Val	Tyr	Arg	Ser	Thr
	275						280					285			
Asp	Gly	Gly	Ala	Thr	Trp	Ser	Arg	Ile	Pro	Gly	Gln	Pro	Thr	Gly	Tyr
	290					295					300				
Leu	Ala	His	Lys	Gly	Val	Leu	Asp	Ser	Ala	Thr	Gly	His	Leu	Tyr	Leu
	305					310					315				320
Thr	Leu	Ser	Asp	Thr	Gly	Gly	Pro	Tyr	Asp	Gly	Gly	Lys	Gly	Arg	Ile
				325					330					335	
Trp	Arg	Tyr	Asp	Thr	Ala	Ser	Gly	Ala	Trp	Gln	Asp	Val	Ser	Pro	Val
				340					345					350	
Ala	Glu	Ala	Asp	Ala	Tyr	Tyr	Gly	Phe	Ser	Gly	Leu	Ser	Val	Asp	Arg
				355					360					365	
Gln	Lys	Pro	Gly	Thr	Leu	Met	Ala	Thr	Ala	Tyr	Ser	Ser	Trp	Trp	Pro
	370					375					380				
Asp	Thr	Gln	Ile	Phe	Arg	Ser	Thr	Asp	Ser	Gly	Ala	Thr	Trp	Thr	Gln
	385					390					395				400
Ala	Trp	Asp	Tyr	Thr	Gly	Tyr	Pro	Asn	Arg	Ser	Asn	Arg	Tyr	Thr	Leu
				405					410					415	
Asp	Val	Ser	Ser	Val	Pro	Trp	Leu	Ser	Trp	Gly	Ala	Ser	Pro	Ala	Pro
				420					425					430	
Pro	Glu	Thr	Ala	Pro	Lys	Leu	Gly	Trp	Met	Thr	Glu	Ala	Leu	Glu	Ile
				435					440					445	
Asp	Pro	Phe	Asp	Ser	Asp	Arg	Met	Met	Tyr	Gly	Thr	Gly	Ala	Thr	Val
	450					455					460				
Tyr	Gly	Thr	Glu	Asp	Leu	Thr	Ser	Trp	Asp	Ser	Gly	Gly	Thr	Phe	Arg
	465					470					475				480
Ile	Thr	Pro	Met	Val	Lys	Gly	Ile	Glu	Glu	Thr	Ala	Val	Asn	Asp	Leu
				485					490					495	
Ala	Ser	Pro	Pro	Ser	Gly	Ala	Pro	Leu	Leu	Ser	Ala	Leu	Gly	Asp	Ile
				500					505					510	
Gly	Gly	Phe	Arg	His	Thr	Asp	Leu	Asp	Ala	Val	Pro	Asp	Leu	Met	Tyr
				515					520					525	
Thr	Ser	Pro	Asn	Leu	Asp	Ser	Thr	Thr	Ser	Leu	Asp	Phe	Ala	Glu	Ser
	530					535					540				
Ser	Pro	Gly	Thr	Val	Val	Arg	Val	Gly	Asn	Ser	Asp	Ala	Ala	Pro	His
	545					550					555				560
Ile	Gly	Phe	Ser	Thr	Asp	Asn	Gly	Ala	Asn	Trp	Phe	Gln	Gly	Ser	Glu
				565					570					575	
Pro	Ser	Gly	Val	Thr	Gly	Gly	Gly	Thr	Val	Ala	Ala	Ala	Ala	Asp	Gly
				580					585					590	
Ser	Gly	Phe	Val	Trp	Ser	Pro	Glu	Gly	Ala	Gly	Val	His	His	Thr	Thr
				595					600					605	
Gly	Phe	Gly	Thr	Ser	Trp	Thr	Ala	Ser	Thr	Gly	Ile	Pro	Ala	Gly	Ala
	610								615					620	
Thr	Val	Glu	Ser	Asp	Arg	Lys	Asn	Pro	Glu	Lys	Phe	Tyr	Gly	Phe	Glu
	625					630					635				640
Ala	Gly	Thr	Phe	Tyr	Val	Ser	Thr	Asp	Gly	Gly	Ala	Thr	Phe	Thr	Ala
				645					650					655	
Glu	Ala	Thr	Gly	Leu	Pro	Ala	Glu	Gly	Asn	Val	Arg	Phe	Gln	Ala	Leu
				660					665					670	
Pro	Gly	Thr	Glu	Gly	Asp	Ile	Trp	Leu	Ala	Gly	Gly	Ser	Asp	Thr	Gly
				675					680					685	

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Ala Tyr Gly Leu Trp Arg Ser Thr Asp Ser Gly Ala Thr Phe Thr Lys
690 695 700

Ser Ala Gly Val Glu Gln Ala Asp Ser Val Gly Phe Gly Lys Ala Ala
705 710 715 720

Pro Gly Ala Ser Tyr Arg Thr Val Phe Val Ser Ala Lys Ile Gly Gly
725 730 735

Val Arg Gly Ile Phe Arg Ser Thr Asp Ala Gly Ala Ser Trp Thr Arg
740 745 750

Ile Asn Asp Asp Ala His Gln Trp Gly Trp Thr Gly Ala Ala Ile Thr
755 760 765

Gly Asp Pro Arg Val Tyr Gly Arg Val Tyr Val Ser Thr Asn Gly Arg
770 775 780

Gly Ile Gln Val Gly Glu Thr Ser Asp Ser Gly Gly Gly Gly Thr Asp
785 790 795 800

Pro Gly Thr Asp Pro Gly Thr Asp Pro Gly Thr Asp Pro Gly Pro Glu
805 810 815

Gln Pro Ala Asp Ala Ala Cys Ala Val Thr Tyr Ala Val Thr Asn Gln
820 825 830

Trp Pro Gly Gly Phe Gln Ala Asp Val Thr Val Thr Asn Thr Gly Asp
835 840 845

Ala Ala Tyr Asn Gly Trp Lys Leu Gly Trp Ser Phe Pro Gly Gly Gln
850 855 860

Gln Ile Ser Gln Ile Trp Asn Ala Ser His Arg Gln Asp Gly Val Lys
865 870 875 880

Val Thr Val Thr Asp Ala Gly Trp Asn Gly Thr Val Ala Pro Gly Ser
885 890 895

Ser Ala Gly Phe Gly Phe Thr Gly Ser Trp Ala Gly Ser Asn Ala Glu
900 905 910

Pro Ala Ala Phe Thr Leu Asp Gly Gln Ala Cys Thr Val Gly
915 920 925

<210> SEQ ID NO 41

<211> LENGTH: 543

<212> TYPE: PRT

<213> ORGANISM: Streptomyces sp. ACTE

<400> SEQUENCE: 41

Met Arg Gly Ala Lys Ser Ala Lys Trp Val Ala Gly Ala Ala Ile Ile
1 5 10 15

Ala Leu Ala Ala Thr Ala Cys Gly Gly Gly Asp Ser Asp Ser Asp Asn
20 25 30

Gly Ala Lys Gly Ala Val Asp Ala Asp Gly Ile Phe Ser Val Glu Val
35 40 45

Gly Glu Pro Gln Asn Pro Leu Gln Pro Ala Asn Thr Met Glu Ser Asn
50 55 60

Gly Ser Ile Val Thr Asp Ala Ile Phe Ser Gln Leu Val Asp Tyr Asp
65 70 75 80

Pro Asp Gly Lys Leu Glu Met Ile Asn Ala Glu Ser Val Glu Thr Thr
85 90 95

Asp Ser Lys Leu Trp Thr Val Lys Leu Lys Lys Asp Trp Lys Phe His
100 105 110

Asp Gly Thr Pro Val Thr Ala Asp Ser Tyr Val Lys Ala Trp Asn Trp
115 120 125

Ala Ala Asn Ile Glu Asn Ala Gln Thr Asn Ala Ser Trp Phe Ala Asp

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130			135			140									
Ile	Lys	Gly	Tyr	Ala	Asp	Val	His	Pro	Asp	Gly	Glu	Gly	Ala	Lys	Pro
145					150					155					160
Lys	Ser	Asp	Ala	Met	Ser	Gly	Leu	Lys	Lys	Val	Asp	Asp	Tyr	Thr	Phe
				165						170					175
Thr	Ile	Glu	Leu	Asn	Ser	Ala	Val	Pro	Tyr	Phe	Ser	Tyr	Lys	Leu	Gly
			180					185						190	
Tyr	Thr	Val	Phe	Ser	Pro	Leu	Pro	Glu	Ser	Phe	Tyr	Ala	Asp	Pro	Lys
		195						200					205		
Ala	Ala	Gly	Glu	Lys	Pro	Val	Gly	Asn	Gly	Ala	Tyr	Lys	Phe	Val	Ser
		210					215				220				
Trp	Asp	His	Lys	Lys	Gln	Ile	Lys	Val	Val	Arg	Asn	Asp	Asp	Tyr	Lys
225					230						235				240
Gly	Pro	Asp	Lys	Ala	Lys	Asn	Gly	Gly	Val	Ile	Phe	Lys	Asn	Tyr	Thr
				245						250					255
Thr	Leu	Glu	Thr	Ala	Tyr	Glu	Asp	Leu	Lys	Ser	Gly	Asn	Val	Asp	Val
			260					265						270	
Leu	Arg	Gln	Ile	Gly	Pro	Lys	Asp	Leu	Pro	Val	Tyr	Arg	Ala	Asp	Leu
		275						280						285	
Glu	Asp	Arg	Ala	Val	Asp	Lys	Ala	Tyr	Ser	Ala	Val	Gln	Thr	Leu	Gly
		290					295				300				
Val	Ala	Met	Tyr	Thr	Asp	Gln	Trp	Lys	Asn	Thr	Asp	Pro	Lys	Val	Leu
305					310						315				320
Gln	Gly	Leu	Ser	Met	Ala	Ile	Asp	Arg	Asp	Thr	Ile	Thr	Lys	Thr	Val
				325						330					335
Leu	Gln	Gly	Thr	Arg	Glu	Pro	Ala	Thr	Gly	Trp	Val	Ala	Lys	Gly	Val
			340					345						350	
Leu	Gly	Tyr	Gln	Glu	Asn	Val	Ala	Gly	Asp	Val	Thr	Lys	Tyr	Asp	Pro
		355						360						365	
Ala	Lys	Ala	Lys	Ala	Leu	Ile	Lys	Glu	Gly	Gly	Gly	Val	Pro	Gly	Asn
		370						375			380				
Glu	Ile	Phe	Ile	Gln	Phe	Asn	Ala	Asp	Gly	Gly	His	Lys	Glu	Trp	Ile
385					390						395				400
Glu	Ala	Val	Cys	Asn	Ser	Ile	Thr	Gln	Ala	Thr	Gly	Val	Lys	Cys	Thr
				405						410					415
Gly	Asp	Ser	Lys	Ala	Asp	Phe	Gln	Ala	Asp	Leu	Asn	Ala	Arg	Asp	Ala
				420				425						430	
Lys	Gln	Val	Lys	Ser	Phe	Tyr	Arg	Ser	Gly	Trp	Val	Leu	Asp	Tyr	Pro
		435						440						445	
Val	Asn	Ala	Asn	Phe	Ile	Ser	Asp	Leu	Phe	Arg	Thr	Gly	Ala	Ala	Gly
		450						455			460				
Asn	Asn	Gly	Phe	Phe	Ser	Asn	Lys	Asp	Leu	Asp	Ala	Lys	Ile	Lys	Ala
465					470						475				480
Ala	Asp	Ser	Ala	Ala	Ser	Leu	Asp	Asp	Ser	Val	Lys	Ala	Tyr	Gln	Glu
				485						490					495
Ile	Glu	Lys	Glu	Leu	Val	Asn	Tyr	Met	Pro	Ser	Ile	Pro	Leu	Trp	Tyr
				500				505						510	
Tyr	Lys	Val	Asn	Ala	Gly	Tyr	Ser	Glu	Asn	Val	Lys	Asn	Val	Asp	Tyr
		515						520						525	
Ala	Gln	Asp	Gly	Asp	Pro	Ile	Leu	Thr	Glu	Val	Gln	Val	Ile	Lys	
		530						535						540	

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<211> LENGTH: 159

<212> TYPE: PRT

<213> ORGANISM: Streptomyces sp. ACTE

<400> SEQUENCE: 42

Met Gln Gly Asp Pro Glu Val Leu Glu Phe Leu Asn Glu Gln Leu Thr
 1 5 10 15
 Ala Glu Leu Thr Ala Ile Asn Gln Tyr Phe Leu His Ala Lys Met Gln
 20 25 30
 Asp His Arg Gly Trp Thr Lys Leu Ala Lys His Thr Arg Ala Glu Ser
 35 40 45
 Phe Asp Glu Met Lys His Ala Glu Ile Leu Thr Asp Arg Ile Leu Leu
 50 55 60
 Leu Asp Gly Leu Pro Asn Tyr Gln Arg Leu Phe His Val Arg Val Gly
 65 70 75 80
 Gln Thr Val Thr Glu Met Phe Gln Ala Asp Arg Gln Val Glu Val Glu
 85 90 95
 Ala Ile Asp Arg Leu Arg Arg Gly Val Asp Leu Met Arg Ala Lys Ser
 100 105 110
 Asp Ile Thr Ser Ala Asn Ile Phe Glu Arg Ile Leu Glu Asp Glu Glu
 115 120 125
 His His Ile Asp Tyr Leu Asp Thr Gln Leu Glu Leu Ile Glu Lys Leu
 130 135 140
 Gly Glu Pro Leu Tyr Leu Ala Gln Val Ile Glu Gln Val Glu Leu
 145 150 155

<210> SEQ ID NO 43

<211> LENGTH: 297

<212> TYPE: PRT

<213> ORGANISM: Streptomyces sp. ACTE

<400> SEQUENCE: 43

Met Ser Pro Tyr Thr Ala Thr Arg Arg Thr Phe Leu Thr Gly Ala Leu
 1 5 10 15
 Ala Ala Ala Thr Gly Val Val Leu Gly Gly Thr Pro Ala Leu Ala Ala
 20 25 30
 Pro Ala Arg Val Leu Gly Thr Gln Asp Trp Met Gly Ala Leu Ala Asp
 35 40 45
 Ser Thr Pro Leu Arg Arg Leu Thr Ile Pro Gly Thr His Asn Ala Gly
 50 55 60
 Ala Arg Tyr Gly Gly Pro Trp Thr Glu Cys Gln Asn Thr Thr Val Ala
 65 70 75 80
 Glu Gln Leu Gly Ser Gly Ile Arg Phe Leu Asp Val Arg Cys Arg Ile
 85 90 95
 Thr Gly Asp Ala Phe Ala Ile His His Gly Ala Ser Tyr Gln Asn Leu
 100 105 110
 Met Phe Gly Asp Val Leu Ile Ala Cys Arg Asp Phe Leu Ala Ala His
 115 120 125
 Pro Ser Glu Thr Val Leu Met Arg Val Lys Gln Glu Tyr Ser Glu Glu
 130 135 140
 Ser Asp Ala Ala Phe Arg Gln Ile Phe Asp Leu Tyr Leu Asp Gly Lys
 145 150 155 160
 Gly Trp Arg Pro Leu Phe Arg Leu Asp Pro Thr Leu Pro Asp Leu Gly
 165 170 175
 Gly Ala Arg Gly Lys Val Val Leu Leu Ala Asp Asn Gly Gly Leu Pro
 180 185 190

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Gly Val Arg Tyr Ala Asp Pro Ala Val Phe Asp Ile Gln Asp Asp Tyr
 195 200 205

Met Ala Glu Pro Phe Gly Lys Tyr Pro Lys Ile Glu Ala Gln Phe Arg
 210 215 220

Lys Ala Ala Gln Gln Pro Gly Lys Leu Phe Met Asn Tyr Val Ser Thr
 225 230 235 240

Ala Ala Leu Leu Pro Pro Arg Ser Asn Ala Asp Arg Leu Asn Pro Gln
 245 250 255

Val His Thr Phe Leu Asp Gly Ser Glu Ala Ala Gly Trp Thr Gly Leu
 260 265 270

Gly Ile Val Pro Leu Asp Tyr Pro Ala Thr Arg Pro Gly Leu Val Glu
 275 280 285

Ser Leu Ile Arg His Asn Pro Val Ala
 290 295

<210> SEQ ID NO 44

<211> LENGTH: 432

<212> TYPE: PRT

<213> ORGANISM: Streptomyces sp. ACTE

<400> SEQUENCE: 44

Val Ser Glu His Thr Asn Asn Ala Val Val Leu Arg Tyr Gly Asp Asp
 1 5 10 15

Glu Tyr Thr Tyr Pro Val Ile Asp Ser Thr Val Gly Asp Lys Gly Phe
 20 25 30

Asp Ile Gly Lys Leu Arg Ala Asn Thr Gly Leu Val Thr Leu Asp Ser
 35 40 45

Gly Tyr Gly Asn Thr Ala Ala Tyr Lys Ser Ala Ile Thr Tyr Leu Asp
 50 55 60

Gly Glu Gln Gly Ile Leu Arg Tyr Arg Gly Tyr Pro Ile Glu Gln Leu
 65 70 75 80

Ala Glu Ser Ser Thr Phe Leu Glu Val Ala Tyr Thr Leu Ile Asn Gly
 85 90 95

Asp Leu Pro Lys Val Asp Glu Leu Ser Ala Phe Lys Asn Glu Ile Thr
 100 105 110

Gln His Thr Leu Leu His Glu Asp Val Lys Arg Phe Phe Asp Gly Phe
 115 120 125

Pro Arg Asp Ala His Pro Met Ala Met Leu Ser Ser Val Val Ser Ala
 130 135 140

Leu Ser Thr Phe Tyr Gln Asp Ser His Asn Pro Phe Asp Glu Glu Gln
 145 150 155 160

Arg His Leu Ser Thr Ile Arg Leu Leu Ala Lys Leu Pro Thr Ile Ala
 165 170 175

Ala Tyr Ala Tyr Lys Lys Ser Ile Gly His Pro Phe Val Tyr Pro Arg
 180 185 190

Asn Asp Leu Gly Tyr Val Glu Asn Phe Leu Arg Met Thr Phe Ser Val
 195 200 205

Pro Ala Gln Glu Tyr Val Pro Asp Pro Ile Val Val Ser Ala Leu Glu
 210 215 220

Lys Leu Leu Ile Leu His Ala Asp His Glu Gln Asn Cys Ser Thr Ser
 225 230 235 240

Thr Val Arg Leu Val Gly Ser Ser Gln Ala Asn Met Phe Ala Ser Ile
 245 250 255

Ser Ala Gly Ile Ser Ala Leu Trp Gly Pro Leu His Gly Gly Ala Asn

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260					265					270					
Gln	Ser	Val	Leu	Glu	Met	Leu	Glu	Gly	Ile	Gln	Ala	Asn	Gly	Gly	Asp
		275					280					285			
Val	Asp	Ser	Phe	Ile	Gln	Lys	Val	Lys	Asn	Lys	Glu	Asp	Gly	Val	Arg
	290					295					300				
Leu	Met	Gly	Phe	Gly	His	Arg	Val	Tyr	Lys	Ser	Phe	Asp	Pro	Arg	Ala
305					310					315					320
Lys	Ile	Ile	Lys	Ala	Ala	Ala	His	Asp	Val	Leu	Ser	Ser	Leu	Gly	Lys
			325						330					335	
Ser	Asp	Glu	Leu	Leu	Asp	Ile	Ala	Leu	Lys	Leu	Glu	Glu	His	Ala	Leu
		340						345					350		
Ser	Asp	Asp	Tyr	Phe	Val	Ser	Arg	Asn	Leu	Tyr	Pro	Asn	Val	Asp	Phe
		355					360					365			
Tyr	Thr	Gly	Leu	Ile	Tyr	Arg	Ala	Met	Gly	Phe	Pro	Thr	Glu	Met	Phe
370						375					380				
Thr	Val	Leu	Phe	Ala	Leu	Gly	Arg	Leu	Pro	Gly	Trp	Ile	Ala	Gln	Trp
385					390					395					400
His	Glu	Met	Ile	Lys	Glu	Pro	Gly	Ser	Arg	Ile	Gly	Arg	Pro	Arg	Gln
				405					410					415	
Ile	Tyr	Thr	Gly	Glu	Val	Leu	Arg	Asp	Phe	Val	Pro	Val	Glu	Ser	Arg
			420					425					430		

<210> SEQ ID NO 45

<211> LENGTH: 527

<212> TYPE: PRT

<213> ORGANISM: Streptomyces sp. ACTE

<400> SEQUENCE: 45

Met	Thr	Lys	Arg	Ala	Gly	Ile	Leu	Val	Ala	Val	Gly	Ala	Thr	Val	Ala
1				5					10					15	
Gly	Leu	Val	Thr	Ala	Val	Pro	Ser	Ala	Ala	Ser	Thr	Ala	Pro	Gly	Ala
			20					25					30		
Pro	Gly	Ala	Ala	Ala	Pro	Leu	Lys	Trp	Thr	Ala	Cys	Gly	Thr	Lys	Ala
		35					40					45			
Tyr	Pro	Thr	Gln	Gln	Cys	Ala	Thr	Val	Arg	Ala	Pro	Leu	Asp	His	Asp
	50					55					60				
Arg	Pro	Ser	Gly	Arg	Gln	Val	Thr	Leu	Ala	Leu	Ala	Arg	Ile	Pro	His
65					70					75				80	
Thr	Ala	Lys	Thr	Ser	Gln	Gly	Pro	Leu	Leu	Val	Asn	Pro	Gly	Gly	Pro
				85					90					95	
Gly	Gly	Ser	Gly	Leu	Ser	Met	Ala	Gly	Phe	Val	Ala	Ser	Ser	Leu	Pro
			100					105						110	
Ala	Lys	Leu	Ala	Ala	Gln	Tyr	Asp	Val	Ile	Gly	Phe	Asp	Pro	Arg	Gly
		115					120					125			
Val	Gly	Arg	Ser	Ser	Pro	Ala	Leu	Asp	Cys	Val	Pro	Lys	His	Phe	Asp
	130					135					140				
Pro	Val	Arg	Pro	Asp	Thr	Val	Pro	Gly	Ser	Pro	Arg	Asp	Glu	Arg	Thr
145					150					155					160
Asn	Arg	Glu	Arg	Ala	Ala	Ser	Phe	Ala	Asp	Ala	Cys	Gly	Glu	Lys	His
				165					170					175	
Gly	Asp	Leu	Leu	Pro	Phe	Met	Asp	Thr	Val	Ser	Thr	Ala	Lys	Asp	Leu
			180					185						190	
Asp	Val	Ile	Arg	Arg	Ala	Leu	Gly	Ala	Arg	Gln	Ile	Asn	Tyr	Phe	Gly
		195					200						205		

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Tyr Ser Tyr Gly Thr Tyr Leu Gly Ala Val Tyr Ala Lys Leu Phe Pro
 210 215 220
 Glu Arg Val Arg Arg Leu Val Leu Asp Ser Ile Val Asp Pro Asp Gly
 225 230 235 240
 Val Trp Tyr Glu Asp Asn Leu Gly Gln Asp Tyr Ala Phe Asp Ala Arg
 245 250 255
 His Lys Ala Phe Ala Ala Trp Val Ala Lys Asn Asp Ala Thr Tyr Arg
 260 265 270
 Leu Gly Thr Asp Pro Ala Lys Val Glu Ala Ala Trp Tyr Arg Met Arg
 275 280 285
 Ala Ala Val Lys Lys His Pro Ala Ala Gly Lys Val Gly Pro Ser Glu
 290 295 300
 Leu Glu Asp Thr Phe Leu Pro Gly Gly Tyr Tyr Asn Gly Tyr Trp Pro
 305 310 315 320
 Gln Leu Ala Glu Ala Phe Ala Ala Tyr Val Asn Asp Lys Asp Glu Asp
 325 330 335
 Ala Leu Ala Thr Ala Tyr Asp Asp Phe Ala Ala Val Asp Ala Ser Gly
 340 345 350
 Asp Asn Gly Tyr Ser Val Tyr Thr Ala Val Gln Cys Arg Asp Thr Gly
 355 360 365
 Trp Pro Lys Ser Trp Thr Thr Trp Arg Asn Asp Thr Trp Gln Ala His
 370 375 380
 Arg Lys Ala Pro Phe Met Ser Trp Asn Asn Thr Trp Tyr Asn Ala Pro
 385 390 395 400
 Cys Ala Thr Trp Pro Val Ala Pro Leu Arg Pro Val Arg Val Thr Asn
 405 410 415
 Arg Glu Ile Pro Pro Ala Leu Leu Phe Gln Ala Thr Asp Asp Ala Ala
 420 425 430
 Thr Pro Tyr Glu Gly Gly Leu Ser Met His Arg Lys Leu Lys Gly Ser
 435 440 445
 Arg Leu Val Val Glu Glu Gly Gly Gly Asn His Gly Ile Ser Leu Ser
 450 455 460
 Gly Asn Asp Cys Leu Asp Ala His Leu Ile Ala Tyr Leu Thr Asp Gly
 465 470 475 480
 Thr Leu Pro Arg Ser Gly Gly Ser Gly Ala Asp Ala Val Cys Asp Ala
 485 490 495
 Leu Pro Glu Pro Glu Ala Ala Ala Thr Ala Lys Ala Lys Ala Ala Thr
 500 505 510
 Gly Gln Lys Gly Ser Thr Leu His Ser Leu Leu Gly Phe Arg Gly
 515 520 525

<210> SEQ ID NO 46

<211> LENGTH: 222

<212> TYPE: PRT

<213> ORGANISM: Streptomyces sp. ACTE

<400> SEQUENCE: 46

Met Asn Cys His Asp Arg Ile Asn Leu Arg Gly Trp Thr Thr Arg Leu
 1 5 10 15
 Ser Gly Leu Phe Val Ala Ala Val Leu Cys Leu Leu Pro Trp Thr Gly
 20 25 30
 Thr Ala Glu Ala His Gly Ser Val Val Asp Pro Ala Ser Arg Asn Tyr
 35 40 45
 Gly Cys Trp Leu Arg Trp Gly Ser Asp Phe Gln Asn Pro Ala Met Ala
 50 55 60

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Gln Glu Asp Pro Met Cys Trp Gln Ala Trp Gln Ala Asp Pro Asn Ala
 65 70 75 80
 Met Trp Asn Trp Asn Gly Leu Tyr Arg Asn Glu Ser Ala Gly Asn Phe
 85 90 95
 Pro Ala Val Ile Pro Asp Gly Gln Leu Cys Ser Gly Gly Arg Thr Glu
 100 105 110
 Gly Gly Arg Tyr Asn Ala Leu Asp Thr Val Gly Ala Trp Gln Ala Thr
 115 120 125
 Asp Ile Thr Asp Asp Phe Thr Val Arg Leu Glu Asp Gln Ala Ser His
 130 135 140
 Gly Ala Asp Tyr Phe Arg Val Tyr Val Thr Glu Gln Gly Phe Asp Pro
 145 150 155 160
 Thr Ala Gln Pro Leu Thr Trp Gly Ala Leu Asp Leu Val Ala Glu Thr
 165 170 175
 Gly Arg Tyr Gly Pro Ser Thr Ser Tyr Glu Ile Pro Val Ser Thr Ser
 180 185 190
 Gly Tyr Thr Gly Arg His Val Val Tyr Thr Ile Trp Gln Ala Ser His
 195 200 205
 Met Asp Gln Thr Tyr Phe Leu Cys Ser Asp Val Asn Phe Gly
 210 215 220

<210> SEQ ID NO 47

<211> LENGTH: 1065

<212> TYPE: PRT

<213> ORGANISM: Streptomyces sp. ACTE

<400> SEQUENCE: 47

Val Ile Ser Arg Arg Arg Leu Leu Ser Thr Thr Ala Ala Thr Ala Ala
 1 5 10 15
 Leu Ala Ala Val Ser Ser Pro Ala Ala Arg Ala Ala Ala Pro Ala Asp
 20 25 30
 Thr Ala Ala Gly Arg Leu Arg Val Thr Gly Pro Thr Val Glu Tyr Val
 35 40 45
 Arg Arg Pro Leu Gly Leu Asp Val Ser Arg Pro Arg Leu Ser Trp Pro
 50 55 60
 Leu Ala Ser Asp His Pro Asp His Gly Gln Ser Ala Tyr Gln Val Arg
 65 70 75 80
 Val Ala Thr Ser Pro Asp Arg Leu Ala Arg Pro Asp Val Trp Asp Ser
 85 90 95
 Gly Lys Val Val Ser Pro Thr Ser Val Leu Val Pro Tyr Ala Gly Pro
 100 105 110
 Ala Leu Val Ser Arg Thr Arg Tyr His Trp Ser Val Arg Val Trp Asp
 115 120 125
 Gln Asp Gly Arg Val Ser Ala Trp Ser Glu Pro Ser Trp Trp Glu Thr
 130 135 140
 Gly Leu Leu Asp Glu Ala Asp Trp Ser Ala Gly Trp Ile Gly Ala Pro
 145 150 155 160
 Ala Ala Leu Thr Ser Ser Pro Ser Leu Glu Ala Ala Ser Trp Ile Trp
 165 170 175
 Phe Pro Glu Gly Asp Pro Ala Val Gly Ala Pro Ala Ala Thr Arg Trp
 180 185 190
 Phe Arg Gly Arg Val Glu Ile Pro Glu Gly Val Thr Arg Ala Arg Leu
 195 200 205
 Val Met Thr Ala Asp Asp Gly Phe Thr Ala Leu Val Asp Gly Val Gln

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210			215			220									
Val	Ala	Arg	Thr	Glu	Pro	Asp	Gly	Pro	Ala	Glu	Asn	Trp	Arg	Arg	Pro
225					230					235					240
Val	Val	Val	Asp	Val	Thr	Ala	His	Leu	Ser	Pro	Gly	Ser	Arg	Val	Val
			245						250					255	
Ala	Val	Thr	Ala	Thr	Asn	Ala	Val	Asp	Gly	Pro	Ala	Gly	Leu	Leu	Gly
			260					265					270		
Ala	Leu	Glu	Leu	Thr	Thr	Ala	Asp	Gly	Ala	Val	Thr	Leu	Ala	Thr	Gly
		275					280					285			
Thr	Gly	Trp	Arg	Ala	Thr	Asp	Arg	Glu	Pro	Asp	Gly	Asp	Trp	Ala	Ser
290						295					300				
Gly	Gly	Tyr	Asp	Asp	Thr	Gly	Trp	Pro	Ala	Ala	Ala	Val	Leu	Ala	Pro
305					310					315					320
Trp	Gly	Ser	Gly	Pro	Trp	Gly	Glu	Val	Arg	Ala	Ala	Leu	Ser	Pro	Ala
				325					330					335	
Thr	Gln	Leu	Arg	Thr	Glu	Phe	Arg	Leu	Gly	Arg	Lys	Arg	Val	Ala	Arg
			340					345					350		
Ala	Arg	Leu	Tyr	Ser	Thr	Ala	Leu	Gly	Leu	Tyr	Glu	Val	Phe	Leu	Asn
		355					360					365			
Gly	Ala	Arg	Val	Gly	Glu	Asp	Arg	Leu	Ala	Pro	Gly	Trp	Thr	Asp	Tyr
370					375						380				
Arg	Lys	Arg	Val	Gln	Tyr	Gln	Thr	Tyr	Asp	Val	Thr	Ala	Leu	Leu	Arg
385				390						395					400
Ser	Gly	Gly	Asn	Ala	Leu	Gly	Val	Thr	Leu	Ala	Pro	Gly	Trp	Tyr	Ala
			405						410					415	
Gly	Asn	Ile	Ala	Trp	Phe	Gly	Pro	His	Gln	Tyr	Gly	Glu	Arg	Pro	Ala
			420					425					430		
Val	Leu	Ala	Gln	Leu	Glu	Val	Thr	Phe	Thr	Asp	Gly	Ser	Ile	Glu	Arg
		435					440					445			
Val	Leu	Ser	Gly	Thr	Gly	Trp	Ala	Ala	Ala	Thr	Gly	Pro	Val	Thr	Ala
450					455						460				
Thr	Asp	Leu	Met	Ala	Gly	Glu	Glu	Tyr	Asp	Ala	Arg	Leu	Glu	Thr	Asp
465				470						475					480
Gly	Trp	Ser	Arg	Ala	Gly	Phe	Asp	Ala	Ser	Gly	Trp	Leu	Ala	Ala	Glu
			485					490						495	
Ala	Val	Glu	Gly	Val	Thr	Ala	Val	Pro	Val	Ala	Ala	Val	Asp	Gly	Ala
			500					505					510		
Cys	Arg	Val	Glu	Arg	Glu	Leu	Thr	Ala	Arg	Glu	Val	Thr	Glu	Pro	Glu
		515					520						525		
Pro	Gly	Val	Tyr	Val	Phe	Asp	Leu	Gly	Gln	Asn	Met	Val	Gly	Thr	Val
530					535						540				
Arg	Leu	Leu	Val	Ser	Gly	Pro	Ala	Gly	Thr	Thr	Val	Arg	Leu	Arg	His
545				550						555					560
Ala	Glu	Val	Leu	Asn	Pro	Asp	Gly	Thr	Leu	Tyr	Thr	Ala	Asn	Leu	Arg
			565					570						575	
Thr	Ala	Arg	Ala	Thr	Asp	Thr	Tyr	Thr	Leu	Arg	Gly	Gly	Gly	Pro	Glu
			580					585						590	
Thr	Tyr	Glu	Pro	Arg	Phe	Thr	Phe	His	Gly	Phe	Arg	Tyr	Val	Glu	Val
		595					600					605			
Thr	Gly	Phe	Pro	Gly	Arg	Pro	Gly	Pro	Asp	Ala	Val	Val	Gly	Arg	Val
610					615						620				
Ile	His	Thr	Ser	Ala	Pro	Phe	Thr	Met	Ala	Phe	Ser	Thr	Asp	Val	Pro
625					630					635					640

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Ala Gly Ser Gly Ile His Arg Phe Thr Ala Pro Ala
 1055 1060 1065

<210> SEQ ID NO 48
 <211> LENGTH: 1170
 <212> TYPE: DNA
 <213> ORGANISM: Streptomyces sp. ACTE

<400> SEQUENCE: 48

atgccggagc gtttcaactcc cactcctgag gacaagtcca cgttcgggtct gtggaccgtg 60
 ggctggcggg gcaacgaccc gttcgggtgag cgcagcgcgc cgggtgctgga cccgggtggag 120
 tcggtcgagc ggctggcgga gctcgggtgcg cacgggggtga cgttccatga cgacgacctg 180
 attccgttcg ggtcgggacga ccgtgagcgg gcgcggtgg tcgggcggtt caggggaggcg 240
 ctggagcgta ccgggtctaa ggtgccgatg gcgacgacga acctgttcac gcacccgggtg 300
 ttcaaggacg gcgggttcac ctccaacgac cgtgacgtgc ggcgggttcgc gctgcgcaag 360
 gtgatccgca acatcgatct cgcgggtggag ctccggcgcgc agacgtatgt ggctgggggc 420
 gggcgtgagg gcgcccagtc cgtgcccggc aaggacgtgc ggtcggccct ggaccggatg 480
 aaggaggcct tcgacctgct gggcgactac gtcaccgagc agggctacga cctgcggttc 540
 gcgatcgagc ccaagcccaa cgagccccgc ggtgacatcc tgctgcccac gatcggggcac 600
 gcgctggcct tcacgagcg cctggagcgc cccgagctgg tcgggggtgaa cccggagacc 660
 gggcacgagc agatggccgg gctgaacttc ccccacggca tcgcccaggc cctgtggggcg 720
 ggcaagctct tccacatcga cctcaacggc cagtcgggga tcaagtacga ccaggacttc 780
 cgcttcggcg ccgggtgaact gcgccaggcg ttctggctcg tggacctcct ggagacggcc 840
 ggctgggacg gctcacgcca cttcgacttc aagccggtag gcaccgacgg catcgacggg 900
 gtgtgggagt ccgcaagaa ctgcatgcgc aactacctca tcctcaagga gcgcccggcc 960
 gccttcgcg ccgacccgc cgtccaggag gccctcacgc cctcccgcct cgacgaactc 1020
 gcccgcccc ccccgacga cggcctcaag gcaactcctc cgcaccgcac cgctacgag 1080
 gacttcgacg ccaccgcgc gcgccaacgc tccatggcct tcgaagccct cgaccagctc 1140
 gccatggacc acctcctcaa cgtccgctga 1170

<210> SEQ ID NO 49
 <211> LENGTH: 1968
 <212> TYPE: DNA
 <213> ORGANISM: Streptomyces sp. ACTE

<400> SEQUENCE: 49

atgacaagcg cgctcagggc gacgcagggt ttgcagtcca cgaaccaccc ccgtttgctg 60
 gacctcacc caggagcacc gttgagcact gaatcccccc gaagaagtcc ccgtctcaga 120
 tggagactcg gcccggggcg ggcaccccg gccaaaggcg tcgcccggctt caccgcactg 180
 ctgctgccgc tcgcccgat ggtcggcctg gcgtcccccg cccaggccgc gacctcggcg 240
 accgccacct acctcaagaa gtcggactgg ggcagcggct tcgagggcca gtggacgggtg 300
 aagaacaccg gcaccaccgc cctgtcctcc tggacgatcg agtgggactt cccctccggc 360
 accgcggctg gctccgcctg ggacgcctcc gtgaccagct ccggcaccca ctggaccgcc 420
 aagaacctcg gctggaacgg tacggtcgcc ccgggtgcca gcatcagctt cggttcaac 480
 ggcaccggat ccggctcccc caccggctgc aagctgaacg gtgcctcctg tgacggcggc 540
 ggcacggctc ccggcgacag cgcctcctcc aagcccggca cccccaccgc gagcggcatc 600

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accgacacct cgggtgaagct ctcctggagc gcagccaccg acgacaaggg catcaagaac 660
tacgacgtcc tgcgcgacgg cgccaaggtc gcgacggta ccacgacgac gtacaccgac 720
accggcctca ccaagggcac ggactactcc tactcgtgc aggcccgga cacgccgac 780
cagaccggac cggtcagcgg cgcggtggcc gtgcccacca cgggcccggaa cgacaaccg 840
ggccccgca cggcagcaa ggtcaacctc ggctacttca ccaactgggg cgtctacggg 900
cgcaactacc acgtcaagaa cctggtgacc tggggtcgg ccgagaagat cacgcacatc 960
aactacgcct tcggcaacgt ccagggcggc aagtgcacca tcggcgactc ctacgccgac 1020
tacgacaagg cctacaccgc cgaccagtgc gtcgacggcg tcgcccacac gtgggaccag 1080
ccgctgcgcg gcaacttcaa ccagctgcgc aagctcaagg cgaagtacc gcacatcaag 1140
gtgatctggt cgttcggcgg ctggacctgg tccggcggct tcggtgccgc ggcgcagaac 1200
ccggcccgct tcgcccagtc ctgctacgac ctggtggagg acccccctg ggccgatgtc 1260
ttcgacggca tcgacatcga ctgggagtag cccaacgcct gcggcctgac ctgtgacacc 1320
agcggccccg ccgcgctgaa gaacctgtcc tccgcgctcc gcgccaagtt cggcgcgaag 1380
aacctggtca ccgcccgcat caccgcggac ggctcggacg gcggcaagat cgacgccgcc 1440
gactacgcgg gcgccgcga gtccttcgac tggtaacaag tgatgacgta cgactttctc 1500
ggcgcctggg aggcgaaggg tccgacggcc ccgcactccc cgctgaacgc gtacgcccgc 1560
atccccgagg acgcttcaa ctcgccgcc gccatcgcca agctgaaggc caagggcgtc 1620
ccggcctcga agctgctgct cggcatcggc ttctacggcc gcggctggac gggcgtgacc 1680
caggcggcac cgggcggcac cgccaccggc gcggccccgg gcacgtacga ggcgggcatc 1740
gaggactaca aggtcctcaa gaccagctgc ccggccaccg gcacgatcgc cggcacccgc 1800
tacgcgcact gcggcaccaa ctggtggagc tacgacacc cggcgaccat cacctccaag 1860
atggcctggg cgaacagcca gggcctcggc ggtgcgttct tctgggagtt cagcggcgac 1920
accgccaacg gcgagctcgt gagcgccatg gacagcggcc tcaactag 1968

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<210> SEQ ID NO 50
<211> LENGTH: 591
<212> TYPE: DNA
<213> ORGANISM: Streptomyces sp. ACTE

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<400> SEQUENCE: 50

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atgcgaaaaa gggcaagcgc ggccgtcata ggctggcga tcgcccggct ctgatgttc 60
gccaccagca gtgccagcag ccacggctac accgattccc ccatcagcag acagaagctg 120
tgtgccaacg gcaccgtcac cggctcgggc aacatccagt gggagccgca gagcgtcgag 180
ggcccgaagg gcttcccggc ggcaggtccg gcggacggca agatctgcgc cggcggaaac 240
agctccttcg ccgcgctcga cgaccgcgc gggggcaact ggcccgccac ccaggtcacc 300
ggcggccagg gctacaactt ccgctggcag ttcaccgccc gccacgccac gaccgacttc 360
cggtaactaca tcaccaagga cggctgggac tccaccaagc cgctcaccag ggccgcctg 420
gagtcgcagc ccttcatgac ggtgccgtac ggaaccagc agccccggc gaccctgacc 480
caccagggca ccatccccac ccagaagtcc ggcaagcaca tcatcctggc cgtctggaac 540
gtggctgaca ccgccaacgc gttctacgcy tgctcggacg tgaagttctg a 591

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<210> SEQ ID NO 51
<211> LENGTH: 1671
<212> TYPE: DNA
<213> ORGANISM: Streptomyces sp. ACTE

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<400> SEQUENCE: 51

gtggcgcgcc	tcgcggccgg	cgccctgacc	gtgaccggtc	tggtcggcac	cgcacaggcg	60
gccgacatca	acgtcgccaa	gaacgcggg	ttcgagagcg	gcctcagcgg	ctggacctgt	120
accggcggca	gcgggccac	cgtctctcc	cccgtagcag	gcggtccgc	cgccctcaag	180
gccaccccga	gcgccagga	caacgcgaag	tgaccccaga	ccgtggccgt	gaagcccaac	240
tccacctatg	cgctcagttc	ctgggtgacg	ggcgggtacg	cctacctcgg	ggcgagcggc	300
accggcacca	ccgacgtctc	cacctggacc	cccggcagca	ccggctggac	ccagctgcgc	360
acgagcttca	ccaccggccc	gtccaccacc	tcggtgacgg	tctacacca	cggtcggtac	420
ggccaggcgg	cctactacgc	ggacgacgtc	gcggtcaccg	gaccgcagcg	cgcgcgcggt	480
acggaggagc	ccggcccggc	gatccccggc	gcccccgccg	gtctggccgt	cggcaccacc	540
acgtcctcct	cggtggccct	gtcgtggaac	gcggtctccg	gcgccaccgg	ctacaccgtc	600
taccgggacg	gcaccaaggc	gaccaccacc	accggcacct	ccgcgacggg	gagcggcctg	660
gccgcccaca	ccgcgtacca	gttctcggtg	agcccaccaca	acgcgcggcg	tgagtccgtc	720
aggtcggcga	ccgtgagcgg	acgtacggcc	aagaaggacg	agaccggccc	gggccctcctg	780
acctccgtgc	ccaagcacgc	cgtgaccggc	tactggcaga	acttcaacaa	cgcgcgggcc	840
gtccagaagc	tcagcgacgt	gcccgcgaac	tacgacatca	tcgccgtctc	cttcgcggac	900
gccgcccgta	ccccgggtgc	cgtcaccttc	aacctcgact	cgccgggccc	gaacggctac	960
accgtcgcgc	agttcaaggc	cgacatcaag	gcccaagcagg	ccgcgggcaa	gaacgtcatc	1020
atctccgtcg	gcgcgagaaa	gggcaccgtc	tcggtcaaca	gcgacgcctc	ggcgaaacgcg	1080
ttcgcggact	cgctgtacac	gctgatccag	gagtacggct	tcaacggcgt	cgacatcgac	1140
ctggagaacg	gcctcaactc	cacctacatg	acgaaggccc	tgcggtcgct	gtcctcgaag	1200
gtgggctccg	gtctcgtcat	cacgatggcg	ccgcagacga	tcgacatgca	gtcgacgtcg	1260
ggtgagtact	tcaagacggc	gctcaacatc	aaggacatcc	tgaccgtcgt	caacatgcag	1320
tactacaaca	gcggttcgat	gctgggctgc	gacggcaagg	tctactcgca	gggctcggtg	1380
gacttctctc	ccgcgctcgc	ctgcatccag	ttggagggcg	gcctcgcgcc	gtcccaggtc	1440
ggcctcggtg	tgcccgcctc	caccgcgggc	gcgggcagcg	gctacgtcgc	cccgtcggtc	1500
gtgaacgcgg	ccctggactg	cctggccaag	ggcaccggct	gcggttcctt	caagccgtcc	1560
aggacgtacc	cggacatccg	tggtgcatg	acctggctga	cgaactggga	cgccacggcg	1620
ggcaacgcct	ggtccaacgc	ggtcggcccg	cacgtccacg	gccttccgta	a	1671

<210> SEQ ID NO 52

<211> LENGTH: 888

<212> TYPE: DNA

<213> ORGANISM: Streptomyces sp. ACTE

<400> SEQUENCE: 52

gtgatcagac	gcgtcatggg	cctgctcacc	gcgctggccg	cggtcgtcgc	gacgctcgtc	60
ttctccccg	ccgccacggc	ctcggcggcc	acctgcgccc	cggcctggaa	cgctcgtcc	120
gtgtacacgg	gcgcggtctc	cgctcgtac	aacgggcaca	actggtcggc	gaagtgggtg	180
acgcagaaag	agcgtccggg	cacctcggac	gtctggggccg	accagggcgc	ctgcggttcg	240
ggcgcgggcg	gcaccgaccc	gaaccctcgc	ggcttcgtcg	tcagcgaggc	gcagtccaac	300
cagatgttcc	cgagccggaa	ctcctctac	acctacagcg	ggctcaccgc	cgcgctgagc	360

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gcctaccocg ccttcgcca caccggcagc gacaccgtga agaagcagga ggcggcgccg	420
ttcctcgcca acgtcagcca tgagaccggc ggcctggctc acatcgtgga gcagaacacc	480
gccaaactacc cgcactactg cgacaccagc cagtcctacg gctgcccggc cggccaggcc	540
gcctactacg gccggggccc catccagctc agctggaact tcaactacaa ggcggccggc	600
gacgcctcgc gcatcgacct gctgggcaac cctcggcagg tggagcagaa cgctccgtg	660
gcttgggaaga cggcctctg gtactggaac acccagtcgg gcccggcac catgacgccc	720
cacaacgcca tcgtcaacgg ctccggattc ggtgagacca tccggctcat caacggcagc	780
atcgagtcca acggcgccaa ccccggccag gtccagagcc gcgtcaaacac ctaccagtcg	840
ttcgtccaga tcctcggtac caccggcgcc tcgaaacctga gctgctga	888

<210> SEQ ID NO 53

<211> LENGTH: 1524

<212> TYPE: DNA

<213> ORGANISM: Streptomyces sp. ACTE

<400> SEQUENCE: 53

atgagacgct cacgatccgt ccgcgcgctg gtgacggcgg ccgtcaccac ggtggcccg	60
gcaggcatgg ccgtgctggg ctccggcacc gccaggcggc cgaccccgtg gcccgaccac	120
gtcttcgccc cctacttoga gtcgtggacc ggagagagcc cggcggccat ggcggccgag	180
tccggggcga aacacctgac catggcgctc ctccagacga cggccaaggc ctctgcacg	240
ccgtactgga acggcgacac cggcctgccc atcggcccagg cgtccttcgg cgcgcacatc	300
gacacgatcc aggccggagg cggcgacgtc atcccgtcgt tcggcggcta caccgaggac	360
accaccggca cggagatcgc cgacagctgc accgacgtcg accagatcgc cgcggcctac	420
cagaaggctg tcacgacgta cgacgtctcg cggctcgaca tggacatcga ggtcgactcc	480
ctcgacgaca ccgcccggat cgaccggcgg aacaaggcca tcaagaagct ccaggactgg	540
gcggacgcga acggccgtga cctggagatc tcctacaagc ttccgacgac caccgcggga	600
ctggcctcca gggcctcgc cgtgctgcgc aacgcctgta ccaacggggc acgggtcgac	660
gtcgtgaacc tgatgacggt cgactactac gacaacgcgt cccacgacat ggcgcgcgac	720
accgagaccg ccgcccaggc cctgtacgac cagctcgcga agctgtaccc gggcaggacc	780
gccacccagc tgtggctccat ggtcggcgtc accgagatgc ccggcgtcga cgacttcggc	840
ccggccgaga ccttcacgct cgccaacgcc gcccggtgt acgactgggc ggtggccaag	900
ggcatcaaca ccctgtcctt ctggggcgtc cagcgcgaca acggcggctg ccccggcgcc	960
ccggccgccc acgactgctc cggcatccag cagaacacct gggacttcac ccgctcttc	1020
gcgcccttca ccagcggcac cacggcgccg gacgacgact tctcggtgac ggccacgccc	1080
gcctccggga cggtgaccgc gggcggttcg gccaccacca cgggtgaagac cgcctgacc	1140
aagggcgccg cacagcaggt cggcctcagc gtcagcgggg tcccggccgg tgtcaccgcc	1200
tcccctagcc cctcctcggc gaccggggc ggccggctca cgtcaccct cgccacgacc	1260
caggccgccc tctcgggcac gtaccggatc agcgtcaccg gtacgagccc gtcgggcagc	1320
cacgcgacgg cctacacgct gaccgtcacc ggcggcaccg gcagccagtg caccggggg	1380
ccgtggggcg gcgggagcgt ctacaccggc ggcagcagg tctcgtacaa gggccacacc	1440
tggaaaggcca agtgggtggc gacggcgag gagcccggca ccaccgggta gtggggcgtc	1500
tggcaggacc tgggcgcctg ctga	1524

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<210> SEQ ID NO 54

<211> LENGTH: 1452

<212> TYPE: DNA

<213> ORGANISM: *Streptomyces* sp. ACTE

<400> SEQUENCE: 54

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gtgacgcagg gaccgctcac cacggaggcc ggcgcgccgg tagccgacaa ccagaacagt    60
gagaccgcag gccccgggtg accggttctc gttcaggacc aggcgcttct ggagaagctg    120
gcccacttca accgggagcg catcccggag cgcgtcgtgc atgccccggg agccggcgcg    180
tacggcacgt tcacgtgac cctgacgtc tcgcagtggc cgcgtgcgaa gttcctctcg    240
gaggtcggca aggagaccga gaccttctg cgtttctcca ccgtcgcggg caacctcggc    300
tcggccgacg cggcgcgtga cccgcgcggc tgggcgctga agttctacac cgaagagggc    360
aactacgacc tcgtcggcaa caacaccccg gtgttcttca tcaaggacgc catcaagttc    420
cccgaactta tccacacca gaagcgcgac ccgtacacgg gctcccagga ggcggacaaac    480
gtctgggact tctggggcct gtcgcgggaa tccaccacc aggtgacctg gctcttcggt    540
gaccgcggca tcccggctc gttccgtcac atgaacggct acggctcgca cacgttccag    600
tggaacaaag aggccggcga ggtcttctgg gtcaagtacc acttcaagac cgaccagggc    660
atcaagaacc tcaccacca ggaggcgcgc cgcctctccg gcgtcgaccc ggacagccac    720
cagcgcgata tgcgtgagtc catcgcgcgc ggtgacttcc cgacctggac ggtgcaggtc    780
cagatcatgc cggcggccga ggcggccacg taccgcttca acccgttcga cctgaccaag    840
gtgtggccgc acgaggacta cccgcgcgac gagatcgca agctggagct caaccgcaac    900
ccggagaaca tcttcgcoga ggtcgcgagc tcgatcttca gcccggcgca cttcgtaccc    960
ggcatcggcc cgtccccgga caagatgctc cagggccgcc tggtcgcgta cggcgaagcc    1020
caccgctacc gcgtcggcat caacgcgcgc cacctgcggg tgaaccgtcc gcacgccacc    1080
gaggcgcgta ccaacagccg tgacggctac ctgtaagcgc gccggcacia gggcacgaaag    1140
aactacgagc cgaacagctt cggcggcccg gtccagaccg acaggccgct ctggcagccc    1200
gtctccgtca cggcgggtac gggcaaccac gaggccgcgc tccacgcgga ggacaacgac    1260
ttcgtgcagg ccggcaatct ctaccggctg atgtcggagg acgagaaggg ccggctgatc    1320
gacaacctgg ccgggttcat cgcgaaggtg tcgcgcgacg acatgcgca tcgocgcatc    1380
aacaacttcc gtcaggccga cgcggacttc ggcaagcggc tggaggtcgc ggtccaggcc    1440
ctgcgcggct ga                                     1452

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<210> SEQ ID NO 55

<211> LENGTH: 2781

<212> TYPE: DNA

<213> ORGANISM: *Streptomyces* sp. ACTE

<400> SEQUENCE: 55

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gtgtatgcca tgccctccac cgccctcgc gcggtccagt ccggagagga cgctcccgtg    60
cgttcaagcc ccagaccctt cgcgcgccctg ctggcggcgc tcgccctgac cgcagggttg    120
tcactcatcg gaaccctcgc cgtggcgcgc tccgacgagg cacctgctgc gacagaagca    180
tcggatgtgt ccatagccgc ggacacctac acctggaaga acgccccgat cgacggcggc    240
ggcttcgtcc ccgggatcgt cttcaaccgg tccgagaaga acctgccta cgcccgacc    300
gacatcggcg gcgcctaccg ctgggaccag tccggcaagc agtggaaagcc cctgctggac    360
tgggtggact gggaccgctg gggctggaag ggcgtggtga gcctcgcctc cgacaaggctc    420

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gacccccgaca acgtgtaacgc cgcctgtgggg acgtacacca acagctggga cccgaccgac	480
ggcgcgggtcc tgcgctctctc ggaccggggc gcctcctgga aggcggccac cctcccgttc	540
aagctcggcg gcaacatgcc cggacgcggc atgggggagc ggctcgcggt cgaccgaa	600
aagaactccg tgctctaact gggcgcgcc agcggcaacg gcctctggcg gtccaccgac	660
gcgggagtca gctggtcoga ggtgacggcc tcccccaacc ccgggaacta cgcgcaggac	720
ccgtcggaca ccagcggtc cggcaacgac aaccaggga tcgtctgggt gacctcgac	780
gagcgttccg gcagcgggg cagcgcacc caggacatct acgtcggggt cgccgacaag	840
gagaacaccg tctaccgtc cacggacggc ggccaccct ggctcgcggt cccgggcca	900
ccccccgct acctcgcga caagggcgta ctcgactccg cgaccggcca cctctatctg	960
acgctgagcg acacggggcg cccctacgac ggccgcaagg gccgatctg gcggtacgac	1020
acggcgtccg gcgctggca ggacgtcagc ccggtggcg aggcgcagc ctactacggc	1080
ttcagcgggc tctccgtgga ccggcagaag cccggcacc tgatggccac cgcctacagc	1140
tcttgggtgc ccgacacca gatcttccgc tccacggaca gcggtgccac ctggaccag	1200
gcctgggact acaccggcta cccgaaccgc tccaaccgt acacgctgga cgtctcctcc	1260
gtgcctggc tctcctggg cgttcccc gcaccggcg agaccgccc gaagctgggc	1320
tggatgacgg aggcgctgga gatcgaccg ttcgactcgg accggatgat gtacggcacc	1380
ggagcgcagg tctacggcac cgaggacct acgtcctggg actccggcg cacgttcagg	1440
atcccccca tggtaaggg gatcgaggag acggcgcgta acgacctggc cagcccgcc	1500
tccggggcac cgttgctgag cgcactcggg gacatcgggg gcttccggca caccgacctc	1560
gacgccgtgc cggacctgat gtacacctc ccgaacctc actcgaccac cagcctggac	1620
ttcgcggaga gctcgcggc caccggtcgc cgggtcggca actccgagc cgcgccccac	1680
atcggcttct ccaccgaaa cggggccaac tggttccagg gctcggagcc ttcgggctc	1740
accggcggcg gcacggctgc ggcggcgcg gacggcagc gcttcgtgtg gagcccggag	1800
ggcgcggggc tccaccacac caccggcttc ggcacctct ggaccgcctc caccggcatc	1860
ccggccggtg ccacggctga gtccgaccg aagaacccc agaagtcta cggattcgag	1920
gcgggcaact tctacgtctc gaccgacggc ggggcgacct tcaccgcca ggccaccggg	1980
ctgcccggcg agggcaacgt ccgcttccag gcaactgccg ggacggaggg cgacatctgg	2040
ctcgcggcg gctccgacac cggggcgta ggtctgtggc gctccaccga ctccggggcg	2100
acgttcacga agtccgccc cgtcagcag gcggacagc tgggcttcgg caaggccgcc	2160
ccgggcgcct cgtaccggac ggtgttcgta agcgcgaaga tcggcggggg gcgcgccatc	2220
ttccgggtcca ccgacggcg ggcgagctgg accaggatca acgacgagc ccaccagtgg	2280
ggctggaccg gcgcccgat cacgggcgac cccagggtct acgggcgct ctacgtctcc	2340
accaacgggc gcgggatcca ggtgggagag acctccgaca gcggcggcg aggcacggac	2400
cccggcaccg atcccggcac cgatcccgc accgatccc gtcggagca gcccgcgac	2460
gccgcctgtg cggtagcga cgcggtcacc aaccagtggc cggcggctt ccaggccgat	2520
gtgacggta ccaaacggg tgacgcggc tacaacggct ggaagctcgg ctggtcgttc	2580
cccggcgggc agcagatcag ccagatctgg aacgcctcgc accggcagga cggggtgaag	2640
gtcacctgca cggacggcg ctggaacggc acggtggcgc ccggtctgctc ggccggcttc	2700
ggcttaccg gcagttggc ggggagcaac gccgaaccgg ccgccttcac cctggacggc	2760
caggcctgca ccgtgggtg a	2781

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<210> SEQ ID NO 56
<211> LENGTH: 1632
<212> TYPE: DNA
<213> ORGANISM: Streptomyces sp. ACTE

<400> SEQUENCE: 56
atgCGcgggtg ccaagagcgc caagtgggtc gCGggagcgg caatcatcgc cctggccgcg      60
accgcctgtg gtggcggcga cagcgcacgc gacaacgggtg ccaagggcgc cgtcgcgcgcg      120
gacggcatat tctcgcgcga ggtcgggtgag cgcgagaacc cgtcgcgcgc ggccaacacg      180
atggagtcga acggcagcat cgtcaccgac gccatcttct cgcagctcgt cgactacgac      240
cccgcggca agctcgcgat gatcaacgcc gagtcgcgtc agacgcgcga cagcaagctg      300
tggacggtea agctcaagaa ggactggaag ttccacgcgc gcacccccgt caccgcgcgc      360
tcctacgtca aggcctggaa ctgggcgcgc aacatcgaga acgcgcgcgc gaacgcctcc      420
tggttcgcgc acatcaaggg ctacgcgcgc gtccaccccc acggcgcggg cgccaagccg      480
aagtcgcgcg ccatgtcgcg cctgaagaag gtggacgcgc acaccttcac catcgcgcgc      540
aactcgcgcg tcccgtactt ctcgtaaac ctcggtcaca cggctctctc gccgctgccc      600
gagtccttct acgcggcccc gaaggcgcgc ggtgagaagc cggtcgcgca cggcgcgcgc      660
aagttcgcga gctgggacca caagaagcgc atcaaggtcg tccgcaacga cgactacaag      720
ggccccgaca aggcgaagaa cggtggtgtg atcttcaaga actacaccac cctcgcgcgc      780
gcctacgcgc acctcaagtc cggcaacgcgc gacgtgctcc gccagatcgc cccgaaggac      840
ctccccggtc accgtgcgca cctcgcgcgc cgcgcgcgtg acaaggccta ctccgcgcgc      900
cagacgcgcg gtgtcgcgat gtacaccgac cagtggaaga acacggcccc gaaggctctc      960
cagggcctgt cgatggccat cgaccgggac acgatcacca agacgggtgt ccagggcacc      1020
cgcgcgcgcg ccacgggctg ggtcgcgcaag ggcgtcctcg gttaccagga gaacgcgcgc      1080
ggtgacgcga ccaagtaaga cccggcgaag gccaaagccc tcatcaagga gggtggcggt      1140
gttccgggca acgagatctt catccagttc aacgcgcgcg gcggccacaa ggagtggtat      1200
gaggcgcgct gcaacagcat cagcgcgcgc accggcgcgc agtgacccgg cgactcgaag      1260
gccgacttcc aggcgcacct gaacgcgcgc gacgccaagc aggtgaagtc gttctaccgc      1320
agtggctggg tcctcgcacta cccggtcaac gccaaactca tcagcgcacct gttcgcgcgc      1380
ggtgcgcgcg gcaacaacgc cttcttctcc aacaaggacc tcgacgcgga gatcaaggcc      1440
gcgcgcgcgc cgcgcgcgcct cgcgcgcgcgt gtaaggcct accaggagat cgagaaggag      1500
ctggtcaact acatgcgccg catccccgctc tggctactaca aggtcaacgc cggctactcg      1560
gagaacgcga agaacgtgga ctacgcgcgc gacggcgcgc cgcgcgcgcgc cgaagtcgcg      1620
gtcatcaagt aa                                                    1632

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<210> SEQ ID NO 57
<211> LENGTH: 480
<212> TYPE: DNA
<213> ORGANISM: Streptomyces sp. ACTE

<400> SEQUENCE: 57
atgcagggcg accccgaggt cctcgcgcgc ctgaacgaac agctgaccgc cgaattgact      60
gccatcaatc agtacttctc gcacgcgaag atgcaggatc accgcgcgctg gaccaagctc      120
gccaaacaca cccgggcccga gtcggtcgcgc gagatgaagc acgcgcgcgc cctgaccgcgc      180

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cggatcctgc tgctggacgg cctgcccac taticagcggc tgttccacgt gcgggtgggc	240
cagaccgtca cggagatggt ccaggccgac cggcaggctc aggtcagggc gatcgaccga	300
ctgcggcgcg gtgtcgatct gatgcgcgcc aagagcgaca tcacgtccgc caacatcttc	360
gaacggatcc tggaggacga ggagcaccac atcgactatc tcgacacca gctggagctg	420
atcgagaagc tcggggagcc gctctacctc gcccaggtea tcgagcaggt cgagctctga	480

<210> SEQ ID NO 58

<211> LENGTH: 894

<212> TYPE: DNA

<213> ORGANISM: Streptomyces sp. ACTE

<400> SEQUENCE: 58

atgagcccgt acaccgccac gcgccggacc ttcctcaccg gcgccctggc gcgccccacc	60
ggagtcgtcc tcggtggtac gcccgccctc gccgcccccg cgagagtccct ggggaccag	120
gactggatgg gggccctcgc cgactccacc ccgtgctgac gcctcaccat ccccggcacc	180
cacaacgcgg gggcccgcga cggcggacc tggaccgagt gccagaacac cacggtggcc	240
gagcagctcg gcagcggcat ccgcttccctg gacgtgcgct gccggatcac cggcgacgcg	300
ttcgcgatcc accacggcgc ctctgaccag aacctgatgt tcggggacgt cctcatcgcc	360
tgccgggact tcctggccgc gcaccctcc gagacgggtc tgatgcgggt caagcaggag	420
tactcggagg agagcgacgc cgcgttcocg cagatcttcg acctgtacct cgacggcaag	480
ggctggcgcc cgtctctccg cctcgacccc accctgccgg acctcggcgg cggccggggc	540
aaggtcgtgc tcctcgcgga caacggcggc ctgcccgggg tccggtaacg cgaaccggcg	600
gtcttcgaca tccaggacga ctacatggcc gagcccttcg gcaagtacct caagatcgag	660
gcgaggttcc gcaaggccgc ccagcagccc ggcaagctct tcatgaacta cgtgtccacc	720
gctgccctgc tgccgcgcg ctcgaaagcc gaccggctca acccgagggt ccacacgttc	780
ctcgacggct ccgaggcggc gggctggacc ggcctcggaa tcgtcccgtt ggactatccg	840
gcgacccgcc ccggcctggt cgagtcgctg atcaggcaca acccggtggc ctga	894

<210> SEQ ID NO 59

<211> LENGTH: 1299

<212> TYPE: DNA

<213> ORGANISM: Streptomyces sp. ACTE

<400> SEQUENCE: 59

gtgagcgagc acaccaacaa cgctgtagta ctgcggtagc gcgatgacga gtacacctac	60
ccggtgatcg acagcaccgt cggcgcacaag ggcttcgaca tcgggaagct cggggccaat	120
acgggcctgg tcacgctgga cagcggatac ggcaacaccg ccgcctataa atccgccatc	180
acctatctcg acggcgaaca gggcatcctg cgctaccgcg gctaccgat cgagcagctc	240
gcgagagact cgacgttccg cgaggtcgcc tacacgctga tcaacggcga ccttcccga	300
gtcgacgagc tgtcggcctt caagaacgag atcaccagc acacgctgct gcacgaggac	360
gtcaagcgtc tcttcgacgg cttcccgcgc gacgcccacc cgatggccat gctgtcctcg	420
gtcgtcagcg cgctgtccac gttctaccag gacagccaca acccgttcga cgaggagcag	480
cgtcacctct cgaagatccg gctgctggcc aagctcccga cgatcgcgcg gtacgcgtac	540
aagaagtcca tcggtaaccc gttcgtctac ccgcgcaacg acctcgggta cgtcgagaac	600
ttcctgcgca tgacctctc ggtcccggcc caggagtacg tgccggaccc gatcgtcgtc	660
tcggcgctcg agaagctgct catcctgcac cgggaccacg agcagaactg ttcgacctcc	720

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accgtgctc tggtcggctc ctgcaggcc aacatgttcg cctccatctc cgcggcacc 780
tcggcgtgt ggggcccgct gcacgggtgc gccaacaccg cgggtgctgga gatgctgaa 840
ggcatccagg ccaacggcgg cgacgtcgac tcttccatcc agaaggtcaa gaacaaggag 900
gacggcgtcc gcctgatggg cttcggccac cgggtgtaca agtccttcga cccgcggccc 960
aagatcatca aggccggcgg ccacgaagtc ctctcctcgc tcggcaagtc cgacgagctg 1020
ctggacatcg cgctcaagct ggaggagcac gcgctctccg acgactactt cgtctcggcc 1080
aacctctacc ccaacgtgga cttctacacg gccctgatct accgggccat gggcttcccg 1140
accgagatgt tcaccgtgct cttcgcgctc gcccgctccc cgggtggat cgctcagtg 1200
cacgagatga tcaaggagcc gggttcccgc atcggcccgc cgcgccagat ctacaccggc 1260
gaggtcctgc gcgacttcgt ccccgtcgag agccgctga 1299

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<210> SEQ ID NO 60

<211> LENGTH: 1584

<212> TYPE: DNA

<213> ORGANISM: Streptomyces sp. ACTE

<400> SEQUENCE: 60

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atgacgaaac gtgcaggcat tctggtcgca gtcggcgcca cggtcgcccg gctggtcacc 60
gcggttccgt ccgcggctc caccgcgcc ggggccctg gggccggcgc gccgctgaag 120
tggaccgctt gcgggacgaa ggcgatccg acccagcagt gcgcaaccgt tcgcccacca 180
ctggaccatg acaggccgctc aggacggcag gtcacgctcg ccctgcgccg gatcccgcac 240
acggcgaaga cctgcgaggg tccgctgctg gtcaaccccg gcggcccggg cggcagcggg 300
ctctcgatgg ccggtctcgt ggcgtcctcg ctgccggcga agctcggcgc ccagtaagac 360
gtgatcggct tcgaccgctc cggggtcggc aggagcagcc cggcgtgga ctgcgtaccg 420
aagcacttcg acccggtaag ccccagacc gtgccggct ccccgcgga cgagcggacc 480
aaccgggaac gcgcccgtc cttcggcag gcgtgcccgc agaagcagc ggaacctgctg 540
ccgttcatgg acacggctcag caccgcaag gacctggacg tgatccgccc ggcctcggc 600
gcacggcaga tcaactactt cggctactcc tacggcacct acctgggccc cgtctacgcc 660
aagctgttcc cggagcggct gcggcgcctg gtgctcgact cgatcgtcga cccggacggc 720
gtctggtaag aggacaacct cggccaggac tacgccttcg acgcccgtca caaggcgttc 780
gccgcctggg tggcgaagaa cgacgcccacc taccggctcg gcaccgacc ggcgaaggtc 840
gaagccgcct ggtaccggat gcgggcccgc gtgaagaagc acccggcggc gggcaaggtc 900
ggcccagcgc agctggagga cacctcctc cccggcggct actacaacgg ctactggccc 960
caactggccc aggcgttcgc cgcgtacgtg aacgacaagg acgaggacgc gctggccacc 1020
gcgtacgacg acttcggccc ggtcgaagcg agcggggaca acgctactc cgtctacacg 1080
gccgtccagt gcccgacac gggctggccc aagtcctgga ccacctggcg caacgacacc 1140
tggcaggcgc accgcaaggc gccgttcctg tcttggaaac acacctggta caacgcgccc 1200
tgcgccacct ggcggctcgc accgctgccc cgggtgcccg tcaccaaccg cgagatcccg 1260
ccggcgtccc tcttccaggc caccgacgac gcggcgacc cgtacgaggg cggcctgagc 1320
atgacccgca agctcaaggc ctcgcgctg gtcgtcgagg agggcggcgg caaccacggc 1380
atcagcctga gcggcaacga ctgcctcgac gcgcacctga tcgcctacct caccgacggc 1440
accctgcccc gctccggcgg cagcggcggc gacgcggctc gcgacgctc ccccgagccc 1500

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gaggcggcgg cgaccgcgaa ggcgaaggcc gctacgggcc agaagggcag caccctgcac 1560
 agcctgctcg gcttccgggg ctga 1584

<210> SEQ ID NO 61
 <211> LENGTH: 669
 <212> TYPE: DNA
 <213> ORGANISM: Streptomyces sp. ACTE

<400> SEQUENCE: 61

atgaattgtc atgatcgcat caacttacgc ggctggacga cacggctgag cggctctgttc 60
 gtcgcccggc tgctctgtct gctcccgtgg acgggcacgg ccgaggccca cggctcggtc 120
 gtcgaccggc cgteccgcaa ctacggctgc tggctccgct ggggcagcga cttccagaac 180
 cccgccatgg cgcaggaaga ccccatgtgc tggcaggcat ggcaggccga cccgaacgcc 240
 atgtggaact ggaacggcct gtaccgcaac gagtcggcgg gcaacttccc ggcagtgatc 300
 cccgacgggc agctgtgcag cggcggccgg accgagggcg gccggtacaa cgcgctggac 360
 accgtggggc cctggcaggc cacggacatc acggacgact tcaccgtgag gctggaggac 420
 caggccagcc acggcgcgca ctacttccgg gtgtacgtca ccgagcaggg cttcgacccc 480
 actgtcagc ccctgacctg gggcgcactc gacctggtgg cggagaccgg acgttacggt 540
 cccagtacga gctacgagat ccccgtagt acgtcggggg acaccggccg ccatgtcgtc 600
 tacacgatct ggcaggcctc gccatggac cagacgtact tcctgtgcag tgacgtgaac 660
 ttcggctga 669

<210> SEQ ID NO 62
 <211> LENGTH: 3198
 <212> TYPE: DNA
 <213> ORGANISM: Streptomyces sp. ACTE

<400> SEQUENCE: 62

gtgatcagca gaagacgact gctcagcacc accgcccga ccgcccctt cgcgcggtc 60
 tcctcgcccg ccgcccggc cgcgcccgg gccgacacc cggccggctc gctccgctc 120
 accggcccga ccgtggagta cgtacgccgc ccgctcggcc tcgacgtctc ccgcccggc 180
 ctgagctggc ccctcgctc ggaccaccgc gaccacggcc agtcgccta ccaggctcgg 240
 gtcgcccact cgcgggacc cctggcccgc cccgacgtct gggacagcgg caaggctcgt 300
 tcccgcagc cgggtcgtgt cccgtaacgc gcccggcgc tggctcctcc taagcgtac 360
 cactggctcg tgccgctgtg ggaccaggac ggacgcgtct cggcctggag ctagcctcc 420
 tgggtgggaga cggggtcctt ggacgaggcc gactggctcg cggggtggat cggcgcgcc 480
 gccgcgctga cctcctcacc ctccctggag gcccgcctct ggatctggtt cccggagggc 540
 gatccggcgg tggcgctcc ggcggccacc cgggtggtcc gcggcgggtt ggagatccc 600
 gaggcgctca ccccgcccgg cctggctcat accgcccagc acggcttcac cgcctcgtc 660
 gacggtgtcc aggtggcccg taccgagccc gacggcccgg cggagaactg gcgtcgtccc 720
 gtggtggtgg acgtgacggc gacacctctc cccgctccc gggctcgtcg cgtgacggcc 780
 accaacgcgg tggacggccc ggcgggtctg ctccggggc tggagctgac caccgcccga 840
 ggtgcggtca cactcgccac gggaaaccgga tggcgggcca ccgaccggga gccggacggg 900
 gactgggctg ccggcgcta cgcgacacc ggctggccc cgcagcgggt cctcgcccgg 960
 tggggttccg gccctcgggg ctaggtacgg gcccctctt ccccggccac ccagctcgcg 1020
 accgaattcc ggctggggcg caagcgcgtc gcgcccggcc ggctgtactc gaccgctc 1080

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ggcctgtaag aggtgttctt gaacggcgca cgtgtcggcg aggaccggct cgcgcccgcc 1140
tggaccgact accgcaagcg cgtccagtag cagacgtacg acgtgacggc actgcttcgg 1200
tccggcggca acgctctcgg ggtcaccctc gcgcgggggt ggtacgcggg gaacatcgcc 1260
tggttcggac cgcaccagta cggcgaacgt ccggcgtac tggcccagtt ggaggtcacc 1320
ttcaccgaag ggtcgatcga gcgggtgctg tcgggcaccg gctgggcccgc cgcgaccggg 1380
cccgtcaccg ccaccgacct catggcaggc gaggagtacg acgcccggct ggagaccgac 1440
ggctggagcc gcgcgggatt cgacgcgtcg gggggctcg cggcagaagc ggtggaaggg 1500
gtcacggccg tgccggctgc cgcgggtggc ggggcctgcc gtgtcgagcg cgagctgacg 1560
gccccgaggg tgaccgaacc cgaccggggg gtctacgtgt tcgacctcgg acagaacatg 1620
gtgggcaccg tacggctcct tgtctcgggg ccggcgggca cgacgggtcg gctgcgccac 1680
gccgaggtgc tgaaccggga cggcaccctc tacacggcca acctgcgcac cgcaccggcc 1740
accgacacct acacgctcag gggcggcgga ccggagacgt acgagccccg cttcaccttc 1800
cacggtttcc gctacgtcga ggtgacgggc ttccggggcc gcccccggcc ggacgcggtg 1860
gtgggcccggg tcatccacac ctccggcgcg ttcaccatgg cctctctgac cgacgtcccc 1920
atgctcgacc ggctccacag caacatcacc tgggggcagc gcggcaactt cctctccgtc 1980
cggaccgaca cgcgccgcgc cgacgaacgc ctccggctga ccggcgacat caacgtcttc 2040
gcgcccaccg ccgcgtacac gatggagtcg gcccgcttcc tcggcaagtg gctccaggac 2100
ctgcgcgacg accagctggc cgaccggccc ttcccgaacg tcgccccgga cctcccgggc 2160
gtcggcagcg gggcggccgg ctggggcgac gccgggggta cggctccgtg ggccctgtac 2220
caggcgtacg gggacgtgcg ggtgctggag cagtcctggt cgtcgatggt ggcctggctg 2280
gagtacctcc aggcgcacag cgaccgtctc ctgcggccgg ccgatgggta cggggactgg 2340
ctcaacatcg aggacgagac acccaaggac gtcacgggca cggcgtaact cgcaccacag 2400
gccgacctca cggcccggac cgcgaggtg ctgggcaagg accccgggccc ctaccgcacg 2460
ctgtccggcc ggggtgcgca cgcgttcocg gcgcggtacg tgggcgacgg cgggcgggtg 2520
aaggggcaca cgcagaccgc gtagctcctg gccctgtcga tggacctgct ggagccgggc 2580
gaccgcgcac cggctgcgga caggctggtc gcgctgatcg aggcgaagga ctggcaactg 2640
tcgacggggg tcctcggcac accgcgcctg ctgcgggtgc tgaccgacac cgggcacacg 2700
gacgtgcctt accggctgct gacgcggcgg acgttcccga gctgggggta ccagatcgac 2760
cggggtgcca ccacgatgtg ggagcgtgg gactccgtgc ggcgggacgg cggtttccag 2820
gacgccggga tgaactcctt caaccactac gcctacgggt cgggtgggca gtggatgtac 2880
gcgaacatcg cgggcatcgc cccggcggcg cccggcttcc gcgagatccg ggtgcttccg 2940
cgtccggggg gcgggggtgca ccgggcccag gcccggttcg actccctgta cgggcccggc 3000
accacccgct ggacctcgga cgggggcggc ttcgcgcttc ggggtgctct gccgcccaac 3060
acgacggccg aggtgtgggt gccgggcggt gacgggagga gctccgtccg gggcacccgc 3120
gtgttctcgc ggcgggagga cgggtgcgcg gtcttcgccc cgggctcggg catccaccgc 3180
ttcaccgcgc cggcctga 3198

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<210> SEQ ID NO 63

<211> LENGTH: 1872

<212> TYPE: DNA

<213> ORGANISM: Artificial Sequence

<220> FEATURE:

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<223> OTHER INFORMATION: CelLcc_CBM3a DNA

<400> SEQUENCE: 63

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atgggacatc accatcatca ccatcaccat gcatccgaaa acctgtactt ccaggcgatc    60
gccatggatc cgaacaatga cgactggctg catgttgaag gtaacaaaat agtggacatg    120
tacggtaatc aggtctggct gaccggctgc aactggtttg gattcaatac cggtagcaat    180
gtgtttgacg gagtatggag ctgcaaatag agagaagccc tcaagggtat ggcggacaga    240
ggaataaatt ttttgagaat acctatttca acagaattgc tgtatcaatg gtctcaagga    300
atatatccca aagcaaatgt taatgatttt gtaaatccgg agctgaaagg aaagaacagc    360
cttgagcttt ttgactttgc cgttcagtgc tgcaaagaat tcggaataaa gataatgggtg    420
gatatacaca gtccggcaac agatgccatg gggcatatgt atcctttatg gtatgacggt    480
caatttaca cagagatgat gatttcaact ttggagtggt tgacggaaag atataaaaat    540
gatgacacaa ttcttgcaact ggaccttaaa aatgagcctc acggcacccc gggcagcgaa    600
ttaatggcca aatgggatgg ttccacggat ttgaacaact ggaagcatgc tgctgaaaca    660
tgcgcaaaga gaatccttgc aataaatccg aatattctta ttgtgtgtaga aggagtggaa    720
gtttatccaa agcctggcta tgattatacc gcagtggacg aatggggaaa agagagttaa    780
tatttctata actggtgggg aggaaattta agaggagtca gggattatcc cattgacctt    840
ggcaagcatc agaagcagct tgtatactca cctcacgatt acgggtcccct cgtacataaa    900
caaccttggt tctatgaagg cttaacaaa gaaactttgt ataatgattg ctggagagat    960
aactgggcat acatacacga ggaaaacatc gctcctctga tagtgggtga atggggaggt   1020
ttcatggacc gcgagacaa cgagaaatgg atgaaagcgc tgagagatta tatgattgag   1080
aataaaatat cccacacttt ttggtgctat aatgcaaatt ccggtgatac cggaggactt   1140
gtatactatg attttattac ctgggacgaa gaaaaaatg ctcttctgaa gcctgcatta   1200
tggcagacag aggacggaaa gtttataggc cttgaccatc agatacctct tggttcaaat   1260
ggaggtttaa acgcgactcc cactaaaggt gccactccta ccaatacggc gactccgact   1320
aagtcggcaa cggcaacgcc cactcgcgcc agcgtaccga ccaatactcc gactaatacc   1380
ccggcgaaca cccagtaag cggtaacctg aaggttgaat tttataactc caaccaagc    1440
gacacaacga atagcatcaa tccgcagttc aaagtcacga aactggcag ttcagctatc   1500
gatctgtcga aactgacctc tcgttactac tatacggttg atggcaaaa agatcagacc   1560
ttttggtgcg accatgcagc aatcatcggt agcaatggtt cttataacgg cattacttct   1620
aatgtaaaag gcacctttgt gaagatgtca agtagcacca acaatgctga tacctacctg   1680
gaaattagct tcacgggtgg cacacttgaa ccaggagccc acgtccagat ccagggccgt   1740
tttgcgaaaa acgattggag caactatacg caatcaaacg attatagttt caaaagcgcg   1800
tctcaattcg tagaatggga tcaggtgacc gcataattga acggagtgtc ggtttggggg   1860
aaagaaccag ga                                                    1872
    
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<210> SEQ ID NO 64

<211> LENGTH: 1872

<212> TYPE: PRT

<213> ORGANISM: Artificial Sequence

<220> FEATURE:

<223> OTHER INFORMATION: CelLcc_CBM3a Amino acids

<400> SEQUENCE: 64

Ala Thr Gly Gly Gly Ala Cys Ala Thr Cys Ala Cys Cys Ala Thr Cys

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1	5	10	15
Ala Thr Cys	Ala Cys Cys	Ala Thr Cys	Ala Thr Gly Cys
	20	25	30
Ala Thr Cys	Cys Gly Ala	Ala Ala Ala	Ala Cys Cys
	35	40	45
Thr Thr Cys	Cys Ala Gly	Gly Cys Gly	Ala Thr Cys
	50	55	60
Thr Gly Gly	Ala Thr Cys	Cys Gly Ala	Ala Ala Thr
	65	70	80
Cys Gly Ala	Cys Thr Gly	Gly Cys Thr	Gly Cys Cys
	85	90	95
Gly Ala Ala	Gly Gly Thr	Ala Ala Cys	Ala Ala Thr
	100	105	110
Thr Gly Gly	Ala Cys Ala	Thr Gly Thr	Ala Cys Gly
	115	120	125
Thr Cys Ala	Gly Gly Thr	Cys Thr Gly	Gly Ala Cys
	130	135	140
Gly Gly Cys	Thr Gly Cys	Ala Ala Cys	Thr Thr Thr
	145	150	160
Gly Ala Thr	Thr Cys Ala	Ala Thr Ala	Cys Cys Gly
	165	170	175
Cys Ala Ala	Thr Gly Thr	Gly Thr Thr	Thr Gly Ala
	180	185	190
Gly Thr Ala	Thr Gly Gly	Ala Gly Cys	Thr Gly Cys
	195	200	205
Thr Gly Ala	Gly Ala Gly	Ala Ala Gly	Cys Cys Cys
	210	215	220
Gly Gly Gly	Thr Ala Thr	Gly Gly Cys	Gly Gly Ala
	225	230	235
Gly Gly Ala	Ala Thr Ala	Ala Ala Thr	Thr Thr Thr
	245	250	255
Gly Ala Ala	Thr Ala Cys	Cys Thr Ala	Thr Thr Cys
	260	265	270
Ala Gly Ala	Ala Thr Thr	Gly Cys Thr	Gly Thr Ala
	275	280	285
Thr Gly Gly	Thr Cys Thr	Cys Ala Ala	Gly Gly Ala
	290	295	300
Ala Thr Cys	Cys Cys Ala	Ala Ala Gly	Cys Ala Ala
	305	310	315
Thr Ala Ala	Thr Gly Ala	Thr Thr Thr	Thr Thr Thr
	325	330	335
Cys Cys Gly	Gly Ala Gly	Cys Thr Gly	Ala Ala Gly
	340	345	350
Ala Gly Ala	Ala Cys Ala	Gly Cys Cys	Thr Thr Gly
	355	360	365
Thr Thr Thr	Thr Gly Ala	Cys Thr Thr	Thr Gly Cys
	370	375	380
Cys Ala Gly	Thr Gly Cys	Thr Gly Cys	Ala Ala Gly
	385	390	395
Thr Cys Gly	Gly Ala Ala	Thr Ala Ala	Gly Ala Ala
	405	410	415
Gly Gly Thr	Gly Gly Ala	Thr Ala Thr	Ala Cys Ala
	420	425	430

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Cys Cys Gly Gly Cys Ala Ala Cys Ala Gly Ala Thr Gly Cys Cys Ala
 435 440 445
 Thr Gly Gly Gly Gly Cys Ala Thr Ala Thr Gly Thr Ala Thr Cys Cys
 450 455 460
 Thr Thr Thr Ala Thr Gly Gly Thr Ala Thr Gly Ala Cys Gly Gly Thr
 465 470 475 480
 Cys Ala Ala Thr Thr Thr Ala Cys Ala Ala Cys Ala Gly Ala Gly Ala
 485 490 495
 Thr Ala Thr Gly Gly Ala Thr Thr Thr Cys Ala Ala Cys Thr Thr Thr
 500 505 510
 Gly Gly Ala Gly Thr Gly Gly Thr Thr Gly Ala Cys Gly Gly Ala Ala
 515 520 525
 Ala Gly Ala Thr Ala Thr Ala Ala Ala Ala Ala Thr Gly Ala Thr Gly
 530 535 540
 Ala Cys Ala Cys Ala Ala Thr Thr Cys Thr Thr Gly Cys Ala Cys Thr
 545 550 555 560
 Gly Gly Ala Cys Cys Thr Thr Ala Ala Ala Ala Ala Thr Gly Ala Gly
 565 570 575
 Cys Cys Thr Cys Ala Cys Gly Gly Cys Ala Cys Cys Cys Cys Gly Gly
 580 585 590
 Gly Cys Ala Gly Cys Gly Ala Ala Thr Thr Ala Ala Thr Gly Gly Cys
 595 600 605
 Cys Ala Ala Ala Thr Gly Gly Gly Ala Thr Gly Gly Thr Thr Cys Cys
 610 615 620
 Ala Cys Gly Gly Ala Thr Thr Thr Gly Ala Ala Cys Ala Ala Cys Thr
 625 630 635 640
 Gly Gly Ala Ala Gly Cys Ala Thr Gly Cys Thr Gly Cys Thr Gly Ala
 645 650 655
 Ala Ala Cys Ala Thr Gly Cys Gly Cys Ala Ala Ala Gly Ala Gly Ala
 660 665 670
 Ala Thr Cys Cys Thr Thr Gly Cys Ala Ala Thr Ala Ala Ala Thr Cys
 675 680 685
 Cys Gly Ala Ala Thr Ala Thr Thr Cys Thr Thr Ala Thr Thr Gly Thr
 690 695 700
 Gly Gly Thr Ala Gly Ala Ala Gly Gly Ala Gly Thr Gly Gly Ala Ala
 705 710 715 720
 Gly Thr Thr Thr Ala Thr Cys Cys Ala Ala Ala Gly Cys Cys Thr Gly
 725 730 735
 Gly Cys Thr Ala Thr Gly Ala Thr Thr Ala Thr Ala Cys Cys Gly Cys
 740 745 750
 Ala Gly Thr Gly Gly Ala Cys Gly Ala Ala Thr Gly Gly Gly Gly Ala
 755 760 765
 Ala Ala Ala Gly Ala Gly Ala Gly Thr Ala Ala Ala Thr Ala Thr Thr
 770 775 780
 Thr Cys Thr Ala Thr Ala Ala Cys Thr Gly Gly Thr Gly Gly Gly Gly
 785 790 795 800
 Ala Gly Gly Ala Ala Ala Thr Thr Thr Ala Ala Gly Ala Gly Gly Ala
 805 810 815
 Gly Thr Cys Ala Gly Gly Gly Ala Thr Thr Ala Thr Cys Cys Cys Ala
 820 825 830
 Thr Thr Gly Ala Cys Cys Thr Thr Gly Gly Cys Ala Ala Gly Cys Ala
 835 840 845

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Thr Cys Ala Gly Ala Ala Gly Cys Ala Gly Cys Thr Thr Gly Thr Ala
 850 855 860

Thr Ala Cys Thr Cys Ala Cys Cys Thr Cys Ala Cys Gly Ala Thr Thr
 865 870 875 880

Ala Cys Gly Gly Thr Cys Cys Cys Cys Thr Cys Gly Thr Ala Cys Ala
 885 890 895

Thr Ala Ala Ala Cys Ala Ala Cys Cys Thr Thr Gly Gly Thr Thr Cys
 900 905 910

Thr Ala Thr Gly Ala Ala Gly Gly Cys Thr Thr Thr Ala Ala Cys Ala
 915 920 925

Ala Ala Gly Ala Ala Ala Cys Thr Thr Thr Gly Thr Ala Thr Ala Ala
 930 935 940

Thr Gly Ala Thr Thr Gly Cys Thr Gly Gly Ala Gly Ala Gly Ala Thr
 945 950 955 960

Ala Ala Cys Thr Gly Gly Gly Cys Ala Thr Ala Cys Ala Thr Ala Cys
 965 970 975

Ala Cys Gly Ala Gly Gly Ala Ala Ala Ala Cys Ala Thr Cys Gly Cys
 980 985 990

Thr Cys Cys Thr Cys Thr Gly Ala Thr Ala Gly Thr Gly Gly Gly Thr
 995 1000 1005

Gly Ala Ala Thr Gly Gly Gly Gly Ala Gly Gly Thr Thr Thr Cys
 1010 1015 1020

Ala Thr Gly Gly Ala Cys Cys Gly Cys Gly Gly Ala Gly Ala Cys
 1025 1030 1035

Ala Ala Cys Gly Ala Gly Ala Ala Ala Thr Gly Gly Ala Thr Gly
 1040 1045 1050

Ala Ala Ala Gly Cys Gly Cys Thr Gly Ala Gly Ala Gly Ala Thr
 1055 1060 1065

Thr Ala Thr Ala Thr Gly Ala Thr Thr Gly Ala Gly Ala Ala Thr
 1070 1075 1080

Ala Ala Ala Ala Thr Ala Thr Cys Cys Cys Ala Cys Ala Cys Thr
 1085 1090 1095

Thr Thr Thr Thr Gly Gly Thr Gly Cys Thr Ala Thr Ala Ala Thr
 1100 1105 1110

Gly Cys Ala Ala Ala Thr Thr Cys Cys Gly Gly Thr Gly Ala Thr
 1115 1120 1125

Ala Cys Cys Gly Gly Ala Gly Gly Ala Cys Thr Thr Gly Thr Ala
 1130 1135 1140

Thr Ala Cys Thr Ala Thr Gly Ala Thr Thr Thr Thr Ala Thr Thr
 1145 1150 1155

Ala Cys Cys Thr Gly Gly Gly Ala Cys Gly Ala Ala Gly Ala Ala
 1160 1165 1170

Ala Ala Ala Thr Ala Thr Gly Cys Thr Cys Thr Thr Cys Thr Gly
 1175 1180 1185

Ala Ala Gly Cys Cys Thr Gly Cys Ala Thr Thr Ala Thr Gly Gly
 1190 1195 1200

Cys Ala Gly Ala Cys Ala Gly Ala Gly Gly Ala Cys Gly Gly Ala
 1205 1210 1215

Ala Ala Gly Thr Thr Thr Ala Thr Ala Gly Gly Cys Cys Thr Thr
 1220 1225 1230

Gly Ala Cys Cys Ala Thr Cys Ala Gly Ala Thr Ala Cys Cys Thr
 1235 1240 1245

Cys Thr Thr Gly Gly Thr Thr Cys Ala Ala Ala Thr Gly Gly Ala

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1250	1255	1260
Gly Gly Thr Thr Thr Ala Ala Ala Cys Gly Cys Gly Ala Cys Thr	1265 1270	1275
Cys Cys Cys Ala Cys Thr Ala Ala Ala Gly Gly Thr Gly Cys Cys	1280 1285	1290
Ala Cys Thr Cys Cys Thr Ala Cys Cys Ala Ala Thr Ala Cys Gly	1295 1300	1305
Gly Cys Gly Ala Cys Thr Cys Cys Gly Ala Cys Thr Ala Ala Gly	1310 1315	1320
Thr Cys Gly Gly Cys Ala Ala Cys Gly Gly Cys Ala Ala Cys Gly	1325 1330	1335
Cys Cys Cys Ala Cys Thr Cys Gly Cys Cys Cys Ala Gly Cys	1340 1345	1350
Gly Thr Ala Cys Cys Gly Ala Cys Cys Ala Ala Thr Ala Cys Thr	1355 1360	1365
Cys Cys Gly Ala Cys Thr Ala Ala Thr Ala Cys Cys Cys Cys Gly	1370 1375	1380
Gly Cys Gly Ala Ala Cys Ala Cys Cys Cys Cys Ala Gly Thr Ala	1385 1390	1395
Ala Gly Cys Gly Gly Thr Ala Ala Cys Cys Thr Gly Ala Ala Gly	1400 1405	1410
Gly Thr Thr Gly Ala Ala Thr Thr Thr Thr Ala Thr Ala Ala Cys	1415 1420	1425
Thr Cys Cys Ala Ala Cys Cys Cys Ala Ala Gly Cys Gly Ala Cys	1430 1435	1440
Ala Cys Ala Ala Cys Gly Ala Ala Thr Ala Gly Cys Ala Thr Cys	1445 1450	1455
Ala Ala Thr Cys Cys Gly Cys Ala Gly Thr Thr Cys Ala Ala Ala	1460 1465	1470
Gly Thr Cys Ala Cys Gly Ala Ala Cys Ala Cys Thr Gly Gly Cys	1475 1480	1485
Ala Gly Thr Thr Cys Ala Gly Cys Thr Ala Thr Cys Gly Ala Thr	1490 1495	1500
Cys Thr Gly Thr Cys Gly Ala Ala Ala Cys Thr Gly Ala Cys Cys	1505 1510	1515
Cys Thr Thr Cys Gly Thr Thr Ala Cys Thr Ala Cys Thr Ala Thr	1520 1525	1530
Ala Cys Gly Gly Thr Thr Gly Ala Thr Gly Gly Cys Cys Ala Ala	1535 1540	1545
Ala Ala Ala Gly Ala Thr Cys Ala Gly Ala Cys Cys Thr Thr Thr	1550 1555	1560
Thr Gly Gly Thr Gly Cys Gly Ala Cys Cys Ala Thr Gly Cys Ala	1565 1570	1575
Gly Cys Ala Ala Thr Cys Ala Thr Cys Gly Gly Thr Ala Gly Cys	1580 1585	1590
Ala Ala Thr Gly Gly Thr Thr Cys Thr Thr Ala Thr Ala Ala Cys	1595 1600	1605
Gly Gly Cys Ala Thr Thr Ala Cys Thr Thr Cys Thr Ala Ala Thr	1610 1615	1620
Gly Thr Ala Ala Ala Ala Gly Gly Cys Ala Cys Cys Thr Thr Thr	1625 1630	1635
Gly Thr Gly Ala Ala Gly Ala Thr Gly Thr Cys Ala Ala Gly Thr	1640 1645	1650

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Ala	Gly	Cys	Ala	Cys	Cys	Ala	Ala	Cys	Ala	Ala	Thr	Gly	Cys	Thr
1655						1660						1665		
Gly	Ala	Thr	Ala	Cys	Cys	Thr	Ala	Cys	Cys	Thr	Gly	Gly	Ala	Ala
1670						1675					1680			
Ala	Thr	Thr	Ala	Gly	Cys	Thr	Thr	Cys	Ala	Cys	Gly	Gly	Gly	Thr
1685						1690					1695			
Gly	Gly	Cys	Ala	Cys	Ala	Cys	Thr	Thr	Gly	Ala	Ala	Cys	Cys	Ala
1700						1705					1710			
Gly	Gly	Ala	Gly	Cys	Cys	Cys	Ala	Cys	Gly	Thr	Cys	Cys	Ala	Gly
1715						1720					1725			
Ala	Thr	Cys	Cys	Ala	Gly	Gly	Gly	Cys	Cys	Gly	Thr	Thr	Thr	Thr
1730						1735					1740			
Gly	Cys	Gly	Ala	Ala	Ala	Ala	Ala	Cys	Gly	Ala	Thr	Thr	Gly	Gly
1745						1750					1755			
Ala	Gly	Cys	Ala	Ala	Cys	Thr	Ala	Thr	Ala	Cys	Gly	Cys	Ala	Ala
1760						1765					1770			
Thr	Cys	Ala	Ala	Ala	Cys	Gly	Ala	Thr	Thr	Ala	Thr	Ala	Gly	Thr
1775						1780					1785			
Thr	Thr	Cys	Ala	Ala	Ala	Ala	Gly	Cys	Gly	Cys	Gly	Thr	Cys	Thr
1790						1795					1800			
Cys	Ala	Ala	Thr	Thr	Cys	Gly	Thr	Ala	Gly	Ala	Ala	Thr	Gly	Gly
1805						1810					1815			
Gly	Ala	Thr	Cys	Ala	Gly	Gly	Thr	Gly	Ala	Cys	Cys	Gly	Cys	Ala
1820						1825					1830			
Thr	Ala	Thr	Thr	Thr	Gly	Ala	Ala	Cys	Gly	Gly	Ala	Gly	Thr	Gly
1835						1840					1845			
Cys	Thr	Gly	Gly	Thr	Thr	Thr	Gly	Gly	Gly	Gly	Gly	Ala	Ala	Ala
1850						1855					1860			
Gly	Ala	Ala	Cys	Cys	Ala	Gly	Gly	Ala						
1865						1870								

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We claim:

1. A method for digesting a non-wood biomass lignocel-
lulosic material, wherein the method comprises:

recombinantly expressing in a non-native microbial host
cell SActE_0237 (GH6) (SEQ ID NO: 1), SActE_0265
(GH10) (SEQ ID NO: 5) and SActE_0236 (GH48)
(SEQ ID NO: 2),

isolating SActE_0237 (GH6) (SEQ ID NO: 1),
SActE_0265 (GH10) (SEQ ID NO: 5) and SActE_0236
(GH48) (SEQ ID NO: 2) from the host cell, and

exposing the non-wood biomass lignocellulosic material
to a sufficient amount of a composition comprising the
isolated SActE_0237, SActE_0265, and SActE_0236,
wherein the exposed lignocellulosic material is at least
partially digested.

2. The method of claim 1, wherein the microbial host is
selected from the group consisting of *Streptomyces lividans*,
Trichoderma reesei, and *Escherichia coli*.

3. The method of claim 1, wherein the method addition-
ally comprises recombinantly expressing in a microbial host
at least one member selected from the group consisting of
SActE_0357 (CE4) (SEQ ID NO:7), SActE_0358 (GH11)
(SEQ ID NO:8), SActE_1310 (PL3) (SEQ ID NO:9),
SActE_3717 (GH9) (SEQ ID NO:10), SActE_4638 (SEQ
ID NO:11), SActE_4738 (GH16) (SEQ ID NO:12),
SActE_4755 (GH64) (SEQ ID NO:13), SActE_5457

(GH46) (SEQ ID NO:14), SActE_5647 (GH87) (SEQ ID
NO:15), and SActE_5978 (PL1) (SEQ ID NO:16) to pro-
duce a composition comprising isolated SActE 0237,
SActE_0265, SActE_0236, and the at least one member, and
exposing the at least 85% non-wood biomass lignocellulosic
material to a sufficient amount of the composition compris-
ing isolated SActE_0237, SActE_2065, SActE_0236, and
the at least one member.

4. A method for digesting a non-wood biomass lignocel-
lulosic material, wherein the method comprises:

genetically modifying an ActE strain to overexpress
SActE_0237 (GH6) (SEQ ID NO:1). SActE_0265
(GH10) (SEQ ID NO:5), and SActE_0236 (GH48)
(SEQ ID NO:2), and

exposing the non-wood biomass lignocellulosic material
to the genetically modified ActE strain,
wherein the exposed lignocellulosic material is at least
partially digested.

5. The method of claim 4, wherein the ActE strain is
further genetically modified to overexpress least one mem-
ber selected from the group consisting of SActE_0357
(CE4) (SEQ ID NO:7), SActE_0358 (GH11) (SEQ ID
NO:8), SActE_1310 (PL3) (SEQ ID NO:9), SActE_3717
(GH9) (SEQ ID NO:10), SActE_4638 (SEQ ID NO:11),
SActE_4738 (GH16) (SEQ ID NO:12), SActE_4755
(GH64) (SEQ ID NO:13), SActE_5457 (GH46) (SEQ ID

NO:14), SActE_5647 (GH87) (SEQ ID NO:15), and SActE_5978 (PL1) (SEQ ID NO:16).

6. A method for enzymatically pretreating agricultural crop materials for consumption by ruminant animals, wherein the method comprises:

recombinantly expressing in a non-native microbial host cell SActE_0237 (GH6) (SEQ ID NO:1), SActE_0265 (GH10) (SEQ ID NO:5) and SActE_0236 (GH48) (SEQ ID NO:2),

harvesting said host cells, and

exposing agricultural crop materials for consumption by ruminant animals to the host cells,

wherein the agricultural crop materials are at least partially digested.

7. The method of claim 6, wherein the host cell is *Saccharomyces cerevisiae* or *Escherichia coli*.

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