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1 **Supplementary Information**

2 **Figure legends**

3 **Fig. S1**

4 Rarefaction analysis of metatranscriptome sequencing depth from sugarcane bagasse composting community by two methods. The first is based
5 on the assumption that the sequencing depth affects the statistics. Therefore, when sequencing becomes redundant, the statistics will be stable
6 [94]. The second method extracts k-mers from each read and checks if it has been seen before. For each 25,000 reads, a point is plotted with the
7 percentage of new reads versus the number of reads processed. The sequencing is saturated after zero is reached. **a** The predicted expression
8 level using the entire and rarefacted libraries were compared at different sequencing depths. At 90% rarefaction, most of the genes have less
9 than 10% fragments per kilobase of transcript per million (FPKM) relative error, but there are still genes with more than 90% relative error. **b**
10 Percentage of unique k-mers as more reads are sequenced. Based on both methods, the sequencing saturation was not reached.

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12 **Fig. S2**

13 Phylogenetic assignment of the expressed CAZymes in sugarcane bagasse composting community through time using the Lowest Common
14 Ancestor algorithm. **a** Relative expression of bacterial phyla. The abundance of genes assigned to Bacteroidetes showed an increase from 29% to
15 44% during 5-week trail, in contrast to genes originating from Proteobacteria that showed opposite trend by decreasing from 42% to 24%. The

16 phylum Firmicutes showed a gradual increase from 1% to 5%. **b** Eukaryotic kingdoms. The expression of CAZymes from non-fungal kingdoms
17 highly grew over time. The total expression of each domain is represented by the gray line.

18

19 **Fig. S3**

20 Biochemical characterization of the compost7_GH6, compost13_GH10 and compost21_GH11 proteins derived from sugarcane bagasse
21 composting community. Effect of **a** pH and **b** temperature on enzyme activity. **c** Substrate specificity examined towards an array of
22 polysaccharides. **d** Residual activity after incubation in the studied temperature.

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24 **Fig. S4**

25 Thermal stability of compost7_GH6 protein examined at different pH values as assessed by ThermoFluor.

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31 **Tables**

32 **Table S1**

Growth weeks	Relative percentage of fungi to bacteria rDNA
0	11 ± 2
1	4.8 ± 0.4
2	9 ± 1
3	21 ± 3
4	20 ± 3
5	22 ± 1

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34 Relative abundance of rDNA amplified from fungal and bacterial specific regions.

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36 **Table S2**

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ID	Length (AA)	Completeness	e-value	Alignment parameters with best hit				Expression	Characterized
				Length	Identity/Gap (%)	Accession number	Taxonomy		
Compost 1_GH5	326	3' partial	0.0	326	52 / 1	gi 919149142	<i>Teredinibacter sp.</i>	N	-
Compost 2_GH5	239	internal	3.2E-08	233	37 / 1	gi 775268352	<i>Acidisphaera rubrifaciens</i>	N	-

Compost 3_GH5	325	5' partial	6.4E-24	341	47 / 7	gi 737251030	<i>Acidobacteriaceae bacterium</i>	N	-
Compost 4_GH5	366	5' partial	4.7E-22	330	48 / 1	gi 931376366	<i>Coxiella sp.</i>	N	-
Compost 5_GH5_5	355	5' partial	5.1E-24	335	43 / 2	gi 931376366	<i>Coxiella sp.</i>	N	-
Compost 6_GH6	284	3' partial	7.7E-30	264	77 / 1	gi 653077963	<i>Marinimicrobium agarilyticum</i>	Y	N
Compost 7_GH6	390	5' partial	0.0	373	49 / 3	gi 1005329896	<i>Sorangium cellulosum</i>	Y	Y
Compost 8_GH6	273	3' partial	2.2E-20	246	74 / 1	gi 653077963	<i>Marinimicrobium agarilyticum</i>	Y	N
Compost 9_GH6	324	internal	0.0	326	48 / 3	gi 546309190	<i>Chondrus crispus</i>	Y	N
Compost 10_GH6_5	377	internal	0.0	380	48 / 1	gi 546309190	<i>Chondrus crispus</i>	N	-
Compost 11_GH7	445	5' partial	0.0	438	68 / 0	gi 761948412	<i>Cylindrobasidium torrendii</i>	N	-
Compost 12_GH9	514	5' partial	7.3E-28	456	49 / 2	gi 797005938	<i>Teredinibacter sp.</i>	N	-
Compost 13_GH10	287	5' partial	0.0	285	91 / 0	gi 769243366	<i>Sorangium cellulosum</i>	Y	Y
Compost 14_GH10	334	5' partial	0.0	327	95 / 0	gi 1005175543	<i>Sorangium cellulosum</i>	N	-
Compost 15_GH10	274	complete	2.0E-44	269	50 / 5	gi 797008181	<i>Teredinibacter sp.</i>	Y	N

Compost 16_GH10	306	internal	0.0	295	38 / 8	gi 1310760	<i>Clostridium thermocellum</i>	N	-
Compost 17_GH10_5	258	internal	0.0	264	52 / 4	gi 161162172	<i>Sorangium cellulosum</i>	N	-
Compost 18_GH11	253	complete	0.0	256	78 / 2	gi 902716143	<i>Cellvibrio sp.</i>	N	-
Compost 19_GH11	244	complete	0.0	239	85 / 0	gi 902716143	<i>Cellvibrio sp.</i>	N	-
Compost 20_GH11	183	5' partial	1.4E-31	184	38 / 5	gi 595588127	<i>Neocallimastix patriciarum</i>	N	-
Compost 21_GH11	227	internal	9.8E-45	229	77 / 0	gi 653077723	<i>Marinimicrobium agarilyticum</i>	Y	Y
Compost 22_GH12	263	5' partial	6.3E-18	269	25 / 21	gi 496168814	<i>Haloterrigena salina</i>	N	-
Compost 23_GH12	250	complete	1.1E-11	364	29 / 43	gi 797011013	<i>Teredinibacter sp.</i>	N	-
Compost 24_GH12	203	internal	1.1E-18	162	27 / 30	gi 493937532	<i>Halosimplex carlsbadense</i>	N	-
Compost 25_GH45	310	5' partial	0.0	241	46 / 6	gi 121816	<i>Cellvibrio japonicus</i>	N	-
Compost 26_GH45	200	5' partial	0.0	222	49 / 10	gi 665990613	<i>Alteromonadaceae bacterium</i>	N	-
Compost 27_GH48	449	internal	0.0	452	96 / 0	gi 502883342	<i>Cellulomonas flavigena</i>	N	-

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40 Parameters of the 27 targets selected for cloning. Some targets had one or both ends missing during sequencing/assembly. However, the
41 predicted domain was fully present. The genes expressed in *E. coli* soluble fraction that were successfully characterized are highlighted.

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