Microbiome and Biocatalytic Bacteria in Monkey Cup (Nepenthes Pitcher) Digestive Fluid

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Tropical carnivorous plant, Nepenthes, locally known as “monkey cup”, utilises its pitcher as a passive trap to capture insects. It then secretes enzymes into the pitcher fluid to digest the insects for nutrients acquisition. However, little is known about the microbiota and their activity in its pitcher fluid. Eighteen bacteria phyla were detected from the metagenome study in the Nepenthes pitcher fluid. Proteobacteria, Bacteroidetes and Actinobacteria are the dominant phyla in the Nepenthes pitcher fluid. We also performed culturomics approach by isolating 18 bacteria from the Nepenthes pitcher fluid. Most of the bacterial isolates possess chitinolytic, proteolytic, amyloytic, and cellulolytic and xylanolytic activities. Fifteen putative chitinase genes were identified from the whole genome analysis on the genomes of the 18 bacteria isolated from Nepenthes pitcher fluid and expressed for chitinase assay. Of these, six clones possessed chitinase activity. In conclusion, our metagenome result shows that the Nepenthes pitcher fluid contains vast bacterial diversity and the culturomic studies confirmed the presence of biocatalytic bacteria within the Nepenthes pitcher juice which may act in symbiosis for the turn over of insects trapped in the Nepenthes pitcher fluid.

Results

Metagenomic Studies. A total of 0.5 million reads (585,046) was generated from metagenomic sequencing. The average read length after quality trimming is 351 bp. Based on the rarefaction curve (Supplementary Figure S1), the alpha diversity of Nepenthes pitcher fluid sample H1 is 31.44. These metagenomic dataset was deposited in sequence read archive (SRA) with the accession number SRR916131.

Microbiome in Nepenthes Digestive Fluid. Metagenome result shows that Nepenthes pitcher fluid contained vast bacterial diversity. Eighteen phyla (Fig. 1) including 29 classes, 53 orders, 112 families and 238 genera
and 616 species were identified by metagenome analysis. The dominant bacteria phyla in *Nepenthes* pitcher fluid sample H1 were *Proteobacteria* followed by *Bacteroidetes*, *Actinobacteria*, *Verrucomicrobia*, and *Planctomycetes* (Fig. 1). A total of 27% of the sequencing read does not match to any DNA sequence in the RDP database. Thus, these reads were listed as “unclassified” (Fig. 1).

*Acidobacteria* which is commonly found in environment with lower pH, was not the dominant bacterial group in this slightly acidic *Nepenthes* pitcher fluid (pH6). It contributed only 0.3% of the total bacteria in *Nepenthes* fluid. A total of three members from the phylum *Acidobacteria* were identified within the *Nepenthes* fluid, namely *Terriglobis saanensis* (98%), *Acidobacterium capsulatum* (0.9%) and *Candidatus Koribacter versatilis* (0.8%).

**Bacteria Identity.** Eighteen bacteria were isolated from *Nepenthes* pitcher fluid sample H1. These bacteria were identified by MALDI-TOF MS Biotyper analysis on the bacterial protein profile and the phylogenetic analysis of the bacteria 16S rDNA gene. The bacteria identity obtained from both methods is listed in Table 1. The DNA sequences of 16S rDNA genes were deposited in GenBank with the accession numbers listed in Table 1.

**Biocatalytic Activity of the Bacterial Isolates.** We conducted the amylolytic, proteolytic, cellulolytic, xylanolytic and chitinolytic screenings and the results are shown in Table 2. *Bacillus* sp. strains H1a and H1m, *P. aeruginosa* strain H1l, *Sphingobacterium* sp. strain H1ai, *S. marcescens* strain PH1a and H1q, *Pseudomonas* strain...
PH1b, *Myroides odoratimimus* strain H1bi and *Microbacterium paraoxydans* strain DH1b possessed proteolytic activity, specifically caseinase activity, as indicated by the formation of cleared zones on the skim milk agars. The ability of the 18 isolates in the degradation of polysaccharide of different degree of complexity was tested using starch, cellulose and xylan as substrates. Among the tested isolates, *Bacillus* strains H1a, H1m and *Sphingobacterium* sp. strain H1ai shows amylolytic activity. The degradation of the iodine/starch complex by the bacteria released iodine into the medium forming yellow zone around the colony on the dark blue background. On the other hand, the cellulose degradation by cellulolytic bacteria was identified by the Congo red discolouration which resulted in the formation of light yellow zone or clear zone around the colony on cellulose agar. Yellow zone surrounding the bacterial colony on cellulose agars was observed upon inoculation of *Bacillus* sp. strains H1a and H1m, *Sphingobacterium* sp. strain H1ai, *Pseudomonas* sp. strains H1h and PH1b, *M. odoratimimus* strain H1bi, *S. marcescens* strain PH1a and *M. paraoxydans* strain DH1b, indicating cellulolytic activity.

The screening for xylanase activity was conducted by fluorescence-based protocol with EnzChek Ultra Xylanase Assay. The xylanase substrate was labeled with fluorescence dye as described by the manufacturer.

| Strain | Identity | Proteolytic | Amylolytic | Cellulolytic | Xylanolytic | Chitinolytic |
|--------|----------|-------------|------------|-------------|-------------|-------------|
| H1a    | *Bacillus* sp. | + | + | + | − | − |
| H1g    | *Klebsiella oxytoca* | − | − | − | − | + |
| H1h    | *Pseudomonas* sp. | − | − | + | − | − |
| H1k    | *Lysinibacillus fusiformis* | − | − | − | − | − |
| H1l    | *Pseudomonas aeruginosa* | + | − | − | − | − |
| H1m    | *Bacillus* sp. | + | + | + | − | − |
| H1n    | *Serratia fonticola* | − | − | − | − | − |
| H1q    | *Serratia marcescens* | + | − | − | − | + |
| H1r    | *Morganella morganii* | − | − | − | − | − |
| H1w    | *Serratia fonticola* | − | − | − | − | − |
| H1ai   | *Sphingobacterium* sp. | + | + | + | − | − |
| H1aii  | *Leifsonia aquatica* | − | − | − | − | − |
| H1bi   | *Myroides odoratimimus* | + | − | − | − | − |
| DH1b   | *Microbacterium paraoxydans* | − | − | − | − | − |
| DH1f   | *Achromobacter* sp. | − | − | − | − | − |
| PH1a   | *Serratia marcescens* | − | − | − | − | − |
| PH1b   | *Pseudomonas* sp. | − | − | − | − | − |
| PH1c   | *Leucobacter* sp. | − | − | − | − | − |

Table 1. The identity of bacteria isolated from *Nepenthes* pitcher fluid H1. Bacteria were identified by 16S rDNA gene sequences phylogenetic analysis and MALDI-TOF MS analysis.

| Strain | Identity | Proteolytic | Amylolytic | Cellulolytic | Xylanolytic | Chitinolytic |
|--------|----------|-------------|------------|-------------|-------------|-------------|
| H1a    | *Bacillus* sp. | + | + | + | − | − |
| H1g    | *Klebsiella oxytoca* | − | − | − | − | + |
| H1h    | *Pseudomonas* sp. | − | − | + | − | − |
| H1k    | *Lysinibacillus fusiformis* | − | − | − | − | − |
| H1l    | *Pseudomonas aeruginosa* | + | − | − | − | − |
| H1m    | *Bacillus* sp. | + | + | + | − | − |
| H1n    | *Serratia fonticola* | − | − | − | − | − |
| H1q    | *Serratia marcescens* | + | − | − | − | + |
| H1r    | *Morganella morganii* | − | − | − | − | − |
| H1w    | *Serratia fonticola* | − | − | − | − | − |
| H1ai   | *Sphingobacterium* sp. | + | + | + | − | − |
| H1aii  | *Leifsonia aquatica* | − | − | − | − | − |
| H1bi   | *Myroides odoratimimus* | + | − | − | − | − |
| DH1b   | *Microbacterium paraoxydans* | − | − | − | − | − |
| DH1f   | *Achromobacter* sp. | − | − | − | − | − |
| PH1a   | *Serratia marcescens* | − | − | − | − | − |
| PH1b   | *Pseudomonas* sp. | − | − | − | − | − |
| PH1c   | *Leucobacter* sp. | − | − | − | − | − |

Table 2. Screening of amylolytic, proteolytic, cellulolytic, xylanolytic and chitinolytic activities from bacteria isolated from *Nepenthes* pitcher fluid H1. +: positive result, −: negative result.
Xylanase-producing bacteria cleaved the xylosidic linkages of the xylanase substrate and released the xylose(s) as well as the fluorophore from the mentioned xylanase substrate. The fluorescence level was compared with the positive control and the fluorescence reference control. Results show that *Pseudomonas* sp. strains *H1i*, *P. aeruginosa* strain *H1l*, *P. fluorescens* strain *H1g*, and *Pseudomonas* sp. strains *PH1b* exhibited xylanolytic activity.

In this study, β-N-acetylglucosaminidase, chitobiosidase and endochitinase activities were tested by supplying bacterial isolates with chito oligosaccharides (4-nitrophenyl N-acetyl-β-D-glucosaminide, 4-nitrophenyl N,N,N′,N′′-tetraacytylchitotriose) with nitrophenol added to the substrate's terminal non-reducing end. The chitinase produced by the bacterial isolates hydrolysed the β-(1→4) glycosidic bond of the chitinase substrate supplied released p-nitrophenol into the solution. The p-nitrophenol was ionized by the stop solution (sodium carbonate) into p-nitrophenylate ion which appeared as yellow color to human eye and absorbed at the absorption 405 nm.

Analysis shows that *S. marcescens* strain *H1g* and *PH1a* exhibited the highest β-N-acetylglucosaminidase, chitobiosidase and endochitinase activities (Supplementary Table S1). These bacteria digested the substrate 4-nitrophenyl N-acetyl-β-D-glucosaminide at the kinetic of releasing 2.15 nmole min⁻¹ p-nitrophenol per min in 1 ml of solution, 4-nitrophenyl N,N,N′,N′′-tetraacytylchitotriose at the releasing kinetic of 1.52 and 4.11 nmole min⁻¹ respectively (Supplementary Table S1). Additionally, other *Serratia* spp. isolates possessed insignificant β-N-acetylglucosaminidase, chitobiosidase and endochitinase activities.

On the other hand, *Klebsiella oxytoca* strain *H1g* and *Pseudomonas* sp. strain *PH1b* exhibited relatively high chitobiosidase activity by releasing p-nitrophenol from N,N,N′,N′′-tetraacytylchitotriose at the 2.38 and 3.79 nmole min⁻¹ ml⁻¹, respectively (Supplementary Table S1). In addition, *Pseudomonas* sp. *PH1b* also displayed a relatively significant endochitinase activity in which it releases p-nitrophenol from 4-nitrophenyl β-(1→3)-N,N′,N′′-tetraacytylchitotriose at the rate of 3.79 and 1.41 nmole min⁻¹ ml⁻¹, respectively (Supplementary Table S1).

**Bacterial Genome.** Genomic study was carried out on each of the bacterial isolated from the *Nepenthes* fluid. The estimated genome size, average content, G + C content, number of contigs and GenBank accession number of each bacteria are listed in Table 3. The size of bacteria genome in this study range from 3.1 to 7.4 Mbps and the genome G + C content ranges from 34% to 71.3% (Table 3). This Whole Genome Shotgun project has been deposited at DDBJ/EMBL/GenBank under the accession number of AYME0000000-AYMV000000. The version described in this paper is version AYME01000000-AYMV01000000.

**Chitinase Gene Determination.** A total of 15 bioinformatically predicted chitinase-encoding genes were identified from the mentioned bacterial genomes and the sequences were deposited at DDBJ/EMBL/GenBank with the accession numbers KT921876-KT921890 (Supplementary Table S2). The amino acid sequences of the predicted genes were compared against NCBI-nr database followed by categorization of sequences according to members of glycoside hydrolases (GH). Thirteen of these predicted chitinase genes were grouped under the GH 18 family chitinase (Fig. 2A) and 2 of the genes were grouped into GH 19.

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**Table 3. Bacteria Genome Sequence Analysis.** Bacteria were sequenced to the average coverage of at least 30-fold and assembled less than 250 contigs. The draft genomes sequences were deposited at DDBJ/EMBL/GenBank.
family chitinase (Fig. 2B). The predicted chitinase gene sequences from S. marcescens H1q and PH1a were identical. Therefore, only chitinase genes from S. marcescens H1q were selected for subsequence studies. Similarly, chitinase gene sequences of S. fonticola strain H1n and strain H1w were identical. Hence only chitinase gene from S. fonticola H1n was selected for further studies. The selections were based on alphabetical order of the strain name.

These predicted chitinase genes activity were validated empirically where the predicted chitinase genes were heterologously cloned and expressed in Escherichia coli BL21 followed by chitinase assay on these transformants. A total of 6 out of 11 transformants exhibited chitinolytic activity.

β-N-acetylglucosaminidase cloned from contig 15 of K. oxytoca H1g and contig 36 of S. fonticola H1n showed hydrolysis of 4-nitrophenyl N-acetyl-β-D-glucosaminide and thus releasing p-nitrophenol at 1.58 and 1.45 nmole min⁻¹ ml⁻¹, respectively (Supplementary Table S1). The transformants which carried the chitobiosidase-encoding genes of Pseudomonas H1h, P. aeruginosa H1l, Bacillus sp. H1m, and S. marcescens H1q were capable to hydrolyse the β-1,4-glycosidic bond of N,N′,N″-triacetyl-β-D-chitotriose and thus releasing the p-nitrophenol at the rate of 0.99, 1.81, 0.83 and 0.93 nmole min⁻¹ ml⁻¹, respectively (Supplementary Table S1).

Discussion

Several literatures have reported the low bacterial diversity within the Nepenthes pitcher fluid. The low bacterial diversity is often associated with the low pH of the Nepenthes juice and the presence of antibacterial and antifungal compounds in the Nepenthes digestive fluid which created an unfavourable environment for bacteria’s growth. It is noteworthy that most studies conducted on Nepenthes were based on the cultured bacteria, therefore the actual bacterial diversity is yet to be discovered especially the uncultured bacteria. Contrary to this, the diverse microbial distribution observed in this study suggested that conventional culture-dependent methods have indeed undermined the microbial diversity in the pitcher fluids. In addition to this, 27% of...
the bacteria from *Nepenthes* pitcher fluid sample H1 were unclassified, as these sequences do not match to any of the sequences in the database. This suggests that the microbial diversity of *Nepenthes* pitcher fluid has a relatively higher complexity. However, the microbiota community within this quadrant remains unknown at the moment due to limitation of resources. The absence of these sequences from the database is because these bacteria were either yet to be discovered or still in the progress of characterization. Therefore, their 16S rDNA sequences are not available in the database.

Our data show that the dominant bacterial phyla in this *Nepenthes* pitcher fluid are *Proteobacteria*, *Bacteroidetes*, *Actinobacteria*, *Verrucomicrobia* and *Planctomycetes*. Former studies have reported their role in the decomposition of organic material in various environments including the gut of eukaryotes. For example, the phylum *Proteobacteria* (*Enterobactericeae*) decomposes monosaccharides and recycle nutrient of forest soil35. On the other hand, the phylum *Bacteroidetes* which is the major composition of human guts microbiota plays an important role in breaking down complex glycan, while the phylum *Actinobacteria* degrades starch and amylopectin in human gastrointestinal16,17. *Verrucomicrobia*, which was found in the guts of sea cucumber and termites, has been reported to be the key player in degrading complex polysaccharide18. The presence of these polysaccharides-degrading bacteria suggest there is a high probability that these microbes play a crucial role in the carbon recycling system within the *Nepenthes* pitcher. These microbes convert complex carbohydrates into simple sugar for the absorption of the plant via the pitcher organ. However, further study is needed in order to determine their contribution within *Nepenthes* digestive fluid.

The culture-dependent approach was conducted to confirm the presence of bacteria in the *Nepenthes* pitcher fluid. MALDI-TOF MS identification was adopted for rapid bacterial identification as it has short turn around time. Results of both identification methods, 16S rDNA and MALDI-TOF biotyping, agreed with each other (Table 1), suggests that the environmental bacterial identification by MALDI-TOF MS Biotyper is reliable. Hence, we would like to recommend the usage of MALDI-TOF MS biotyping for environmental bacterial identification owing to its fast turn over, low operating cost and simple workflow19,20.

In general, most of the *Nepenthes* live on barren soil that lacks of nitrogen source, which is an essential macro-nutrient for plant growth21. The absorption of nutrients from *Nepenthes* captive by the pitcher stimulates the nutrient uptake activity via root, and thus the later activity promotes the growth of the plant22. This interconnect mechanism display the importance of the pitcher for the vital plant growth23.

In this work, screening of the bacterial biocatalytic activities shows that a total of 12 bacterial isolates were able to degrade complex carbohydrate such as starch, xylan and cellulose. The degradation of complex polysaccharides such as tree bark, plant debris, and leaves that fallen into the *Nepenthes* pitcher fluid releases carbon as well as nitrogen into the *Nepenthes* pitcher fluid, thus nourishing the plant as well as the microbes inside the fluid24.

The *Nepenthes* pitcher serves as a passive trap that lures insects, which are organisms armoured by fibrous chitin and filled with protein. Therefore, the degradation of the insect cadavers involves complex proteolytic and chitinolytic activities25. The nitrogenous protein stored within the insect fibrous chitin and proteins embedded within the insect’s exoskeleton made up of 10% of insect’s biomass44–46. Among the 18 bacterial isolates, 9 proteases-producing and 4 chitinases-producing bacteria strains were identified as the potential contributors in insect decomposition processes. Previous study has indicated 61.5% ± 7.6% of the nitrogen within *Nepenthes* plant contributed by the decomposition of insects27.

Previous studies have confirmed that *Nepenthes* secretes class I, III, and IV chitinases into its digestive pitcher fluid48,28,29. The upregulation of the *Nepenthes* chitinase gene expression in the presence of insect or chitin and the ability to hydrolyse β-1–4 glycosidic bond of chitin has led to the suggestion that the plant chitinases plays an important role in insect degradation in *Nepenthes* pitcher6,7,29. The exoskeleton of insect is the main captive in *Nepenthes* pitcher which is made up of primarily chitin, a polymer of N-acetyl-β-D-glucosamine30,31. The chitin degradation involves the cleavage of its β-1–4 linkage from the non-reducing end by exochitinase and the random cleavage of chitin β-1–4 linkage within the chitin chain by the endochitinase30,32–35. The degradation of chitin involved 2 major steps, which is the cleavage of chitin into chitin oligosaccharides followed by the further cleavage to N-acetylglicosamine by chitobiases31,36.

In this study, 4 chitin oligosaccharides degradation bacteria were isolated from the *Nepenthes* digestive fluid. The whole genome analysis on the bacterial genome has determined 15 endo- and exochitinases. These chitinase genes were further classified into GH18 and GH19 families based on their amino acid sequences37,38. Before the whole genome analysis on the bacterial genome has determined 15 endo- and exochitinases. These chitinase genes were further classified into GH18 and GH19 families based on their amino acid sequences37,38. Before the documentation of GH19 family chitinase gene in *Streptomyces griseus* HUT6037, GH19 family was only found on higher plants29. However, at the point when this report was written, 1799 bacteria GH19 were documented in carbohydrate active enzyme (CAZY) database. Further screening has confirmed the β-N-acetylglicosaminidase, chitobiosidase and endochitinase activities of the chitinase genes. These suggest that the bacteria inhabit in *Nepenthes* digestive fluid cooperate with the *Nepenthes* plant for the complete digestion of the insect captive. The *Nepenthes* chitinase breakdown the chitin in the insect exoskeleton converting the chitin into chitin oligosaccharides, while the bacteria aid in the chitin oligosaccharides degradation, accelerating the nutrient recycling process.

The production of a wide range of biocatalytic enzymes by both the *Nepenthes* and bacteria inhabit in the *Nepenthes* pitcher digestive fluid contrives a dynamic environment in which both work in synchrony for the decomposition of insects, benefiting both the plant and its microbiota. This symbiotic system ensures constant essential nutrient supply from *Nepenthes* pitcher to the entire plant, thus enables the *Nepenthes* plant to survive and thrive in a nutrient-deplete environment. However, further studies need to be conducted in order to prove this hypothesis.
Methods

Sample Source. *Nepentes* pitcher fluid (Sample H1) was collected from wild *Nepentes* in Mossy Forest, Pahang, Malaysia (N 04°31′, E 101°22′), at the altitude of 1970 m above sea level. The *Nepentes* digestive fluid was transported to laboratory and processed immediately.

Metagenome DNA Extraction. Total genomic DNA was extracted from 35 ml of *Nepentes* pitcher fluid by modified cetyltrimethylammonium bromide (CTAB) DNA extraction protocol. Briefly, cells were mechanically lysed with glass bead followed by chemical lysis with CTAB lysis buffer (100 mM Tris–HCl, 100 mM EDTA, 100 mM K2HPO4, 1.5 M NaCl, 1.0% (w/v) SDS and 1.0% (w/v) CTAB)40,41. Enzymatically lysis was performed by addition of lysozyme (100 μg/ml) and incubated at 37°C for 30 min. After Proteinase K (40μg/ml) and RNase A treatment, the DNA was purified with phenol/chloroform/isoamyl alcohol mixtures (25:24:1) and precipitated according to the protocol provided by manufacture at 25°C. The presence of xylanase was detected by measuring the fluorescence when xylanase breaks the xylosidic linkages of the xylanase. The fluorescence was measured at excitation/emission 360/465 using Infinite® F200 PRO (Tecan, Switzerland).

Targeted Metagenomic Sequencing. Total DNA extracted from the *Nepentes* fluid was subjected to 16S rDNA genes amplification with forward primer (MID1_530F, 5′-ACG AGT GCG TGT GCC AGC AGC MGC NGC G-G-3′) and reverse primer (MID1_1100modR, 5′-ACG AGT GCG TGG GTT NCG NTC GTT RC -3′)42. Gene amplification was performed with gradient annealing temperature from 55°C to 65°C. The amplicon sequencing was performed on GS-FLX Titanium platform (Roche, USA).

Taxonomic Assignment of Metagenomic Sequences. Sequences were trimmed with CLC genomics workbench and annotated with MG-RAST (v3.3) against RDP database21,43,44. Rarefaction curve and alpha diversity were constructed and calculated by comparing the data to RDP database using default setting (maximum e-value cutoff: 1e−5, minimum identity cutoff: 60.0% and minimum alignment cutoff: 15) in MG-RAST user interface. The bacteria abundance was calculated based on the data generated from MG-RAST.

Culturomics Bacterial Identification. Bacteria were isolated from highland *Nepentes* pitcher fluid (sample H1) and maintained using Luria-Bertani (LB) medium at 28°C. The identity of the culturable bacteria was identified using MALDI-TOF MS (Bruker, Germany) equipped with Bruker FlexControl software version 3.3 and Bruker MALDI Biotype Real Time Classification (RTC) version 3.1. This bacterial identification was performed according to the direct transfer procedure from Bruker45. Protein from the bacterial cell was measured by MALDI-TOF MS and the spectra generated were compared to the reference database for bacterial identification. Results with the log (score) value equal or higher than 2 indicate high confidence identification and is tabulated in Table 1.

The bacterial identities were confirmed molecularly by phylogenetic analysis on their 16S rDNA genes sequences. Bacterial genomic DNA were extracted using QIAamp DNA mini kit according to manufacturer instruction. 16S rDNA genes were amplified with 27F and 1525R primer pair46. Molecular identities of the isolates were identified based on the phylogenetic analysis of the 16S rDNA gene sequences using MEGA (v6.06)47.

Biocatalytic Assays. Amylolytic activity of the 18 bacterial isolates was screened using diluted LB agar supplied with 0.5% (w/v) soluble starch. The utilization of starch by the bacteria was observed by flooding the starch agar with iodine. Iodine reacts with starch giving a dark blue substrate. Amylolytic bacteria degrade starch and release iodine from iodine/starch complex into the medium which results yellow halo around bacteria colonies on dark blue background.

The bacterial proteolytic activity was screened using diluted LB agar supplied with 5% (v/v) skim milk. The bacterial proteolytic activity was screened using diluted LB agar supplied with 5% (v/v) skim milk. Skim milk contributed to the opaque properties on the agar. Clear-zone around the bacterial colony will be observed on dark blue background. The bacterial proteolytic activity was screened using diluted LB agar supplied with 5% (v/v) skim milk.

Cellulolytic activity was tested with basal salt medium supplied with 0.15% (w/v) microcrystalline cellulose, adjusted to pH 7.0. This agar was stained red with 0.02% (w/v) Congo red. The cellulose degradation by cellulolytic bacteria was identified by the Congo red discolourization which results in the formation of light yellow zone or clear zone around the colony on medium9.

Xylanolytic activity was screened using EnzChek® ultra xylanase assay kit (Invitrogen, USA). The xylanase substrate (synthetic hemicellulose polysaccharides) was tagged with fluorescence dye and supplied by the kit. The structure of the xylanase is proprietary product to the manufacture. Xylanase by *Trichoderma viride* (Sigma, US) was used as the positive control for the xylanolytic assay. Bacterial samples were prepared by resuspending the cell pellet from 1 ml of bacteria culture into 100μl of sterile culture medium. Working buffer (100μl) was added to the resuspended cell before loading the bacteria cells (50μl) into the 96 wells black plate. The assay was performed according to the protocol provided by manufacture at 25°C. The presence of xylanase was detected by measuring the fluorescence level and compared to the standard curve. Fluorophore was released from the xylanase substrate when xylanase breaks the xylosidic linkage of the xylanase. The fluorescence was measured at excitation/emission 360/465 using Infinite® F200 PRO (Tecan, Switzerland).

Chitinase, in particular, β-N-acetylglucosaminidase, chitobiosidase and endochitinase activities of the bacterial isolates were tested using chitinase assay kit (Sigma, US) as described by manufacture48. Substrate solution was prepared by dissolving 1 mg of appropriate substrate (4-nitrophenyl N-acetyl-β-D-glucosaminide, 4-nitrophenyl N,Ndiacetyl-β-D-chitobiose or 4-nitrophenylβ-D-N,N′,N″-triacetyltchitotirome) in 1 ml of assay buffer. Chitinase control enzyme was diluted to 10μg/ml with phosphate buffered saline before used. Diluted chitinase control enzyme (10μl) was added to 90μl of substrate solution and this reaction served as positive control in the assay. Blank reaction was prepared by loading 100μl of substrate solution to a well in microtitre plate. Each reaction tested by adding 10μl of overnight bacteria culture to 90μl of substrate solution. The reaction was incubated for 24 hours followed by the addition of 200μl of stop solution. The assay was carried out at 28°C for 30min.
bacterial culture and 37 °C for *E. coli* transformant carrying chitinase gene. β-N-acetylglucosaminidase, chitobiosidase and endochitinase producing bacteria hydrolyzed the chitinase substrate and released p-nitrophenol. The p-nitrophenol released was ionized by the stop solution (sodium carbonate) into p-nitrophenylate ion which was yellow color. The yellow tint measurement was taken at the absorption 405 nm using Infinite® M200 (Tecan, Switzerland) and compared with the reading from chitinase control enzyme.

**Bacteria Whole Genome Sequencing.** Genomic DNA of the bacterial isolates was extracted using QIAamp DNA mini kit (Qiagen, Germany) as described by the manufacturer. The quality and quantity of the DNA was checked with Nanodrop 2000c (Thermo Scientific, USA) and Qubit 2.0 fluorometer (Invitrogen, USA), respectively. Sequencing libraries were prepared with Nextera sample prep kit (Illumina, USA) and the indexing QIAamp DNA mini kit (Qiagen, Germany) as described by the manufacturer. The quality and quantity of the genomes was deposited on DDBJ/EMBL/GenBank.

**Bacteria Draft Genome Analysis and Gene Prediction.** The quality of the sequencing raw data was checked with FastQC[29]. Subsequently, the paired-end reads were trimmed and de novo assembled with CLC genomic workbench version 6 (CLC, Denmark)[31]. The coding DNA sequence (CDS) was predicted using prodigal (v2.6.0)[39]. Draft genomes were annotated by BLASTX against NCBI non-redundant (nr) database[51]. The draft ing library was loaded into MiSeq sequencing cartridge (2 × 250 bp, 500 cycles) (Illumina, USA). The sequencing was performed on MiSeq (Illumina, USA) sequencing platform.

**Chitinase Gene Analysis and Cloning.** The putative chitinase genes sequences with e-value of 0.00, indicating that the sequence alignment was not occurred by chance, were preceded for further analysis[52]. The low e-value indicates that the match of the query gene against the database was significant. The chitinase genes were grouped by phylogenetic analysis using MEGA (v6.06) by its amino acid sequences[47]. These genes were synthesized and inserted into pET−23(+) expression vector by service provider, Genscript. Subsequently, the vector was transformed into *Escherichia coli* BL21 and preceded to chitinolytic assays.

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Author Contributions
X.Y.C., K.W.H. and W.E.Y. performed the experiments and analysed the data. X.Y.C. and K.W.H. prepared the figures. X.Y.C., K.W.H., W.E.Y. and K.G.C. collected the samples and prepared the manuscripts. W.E.Y. procured the chemicals, instruments and reagents for this work. K.G.C. conceived the idea, obtained the funding and proofread the draft.

Additional Information
Accession Codes: Bacteria 16S rDNA sequences are available at DDBJ/EMBL/GenBank with the accession number KF557585–KF557587, KF557591–KF557597, KF557599–KF557602 and KF742682–K742685. The bacteria whole genome sequences are available at DDBJ/EMBL/GenBank with the accession number AYME00000000 – AYM00000000. The putative chitinase genes sequences were deposited in DDBJ/EMBL/GenBank and accessible with the accession numbers KT921876 – KT921890 KT921890 KT921890. KT921890.

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