Research Article

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Isolation, screening and optimization of alkaliphilic cellulolytic fungi for production of cellulase

https://doi.org/10.1515/gps-2023-0153
received August 11, 2023; accepted December 27, 2023

Abstract: This study concerns with the production and partial characterization of alkaline cellulase from alkaliphilic cellulolytic (AC) fungi isolated from soil in Perlis, Malaysia. The best fungi strain was selected on the basis of producing the highest cellulase at high pH conditions. Cellulase from the selected fungi strain was further characterized under saccharification but varies in operating parameters. Finally, the kinetic model describing the growth of the AC fungi strain was studied by employing the logistic model. Among the tested fungi strains, Basidiomycetes strain (BK1) showed high potentiality for the production of maximum alkaline cellulase production at pH 9 after 72 h of incubation at 30°C containing 6 g·L\(^{-1}\) carboxyl methyl cellulose. The saccharification process showed that the enzyme favour high alkaline condition and proves thermotolerant properties, while 15% (v/v) enzyme loading and 1% substrate concentration recorded the highest glucose production at about 1.2–1.3 mg·mL\(^{-1}\). The novelty of the study is to identify and optimize a unique indigenous fungi that emit alkaliphilic cellulase as alternative usage in biotechnology industries due to its capacity to adapt to the extreme conditions of specific industrial processes. There are revolutionary options for use in biotechnological businesses that involve high pH and therefore have substantial biotechnological promise.

Keywords: alkaliphilic cellulolytic fungi, alkaline cellulase, enzyme, saccharification, logistic growth model

1 Introduction

Cellulose has been identified as a rich and limitless supply of raw material for a range of bioproducts. It is the most common organic polymer, making up roughly \(1.5 \times 10^{12}\) tonnes of the entire annual biomass production \([1,2]\). It is a main component of raw materials such as cotton and wood, as well as the plant cell wall’s principal structural component. Agricultural and forestry wastes, municipal solid waste, herbaceous and woody plants, and unused standing forest are all examples of cellulosic biomass that contain cellulose. Agricultural waste primarily consists of cellulotic matter, which is easily dispatched by a range of
physical, chemical, and biological processes [3]. Cellulase is a family of enzymes that can hydrolyze cellulose on 1,4-D-glucan linkages and create glucose, cellobiose, and cello-oligosaccharides as major products. Exoglucanase (EC 3.2.1.91), endoglucanase (EC 3.2.1.4), and β-glucosidase (EC 3.2.1.21) are the three key components of this multicomponent enzyme system [4]. The majority of the cellulases studied and applied in industries are mainly from nonextremophiles, while only a few reports [5,6] deal with the production of the cellulases from extremophiles. Alkaliphiles research has led to the development of a variety of enzymes with unique characteristic. Over the last two decades, the usage of alkaline cellulase has grown significantly in industries such as agriculture, kraft pulping, textile, food and beverages, animal feed, detergent industry, and biorefinery since the enzymes can increase the products quality [8,9]. Indeed, the demand for these enzymes is increasing at a faster rate than that of yeast and bacteria. Cellulases from fungal sources are preferred for commercial enzyme production because the enzymes produced are of high quality [10]. Most of cellulases produced by fungi and bacteria are well known to perform optimally under alkaline and neutral conditions [11]. The study of extremophiles and extremozymes began with alkaliphiles, and Horikoshi [10] is well known to perform optimally under alkaline and neutral conditions. Alkaliphiles research has led to the discovery of AC fungi strains that can emit cellulase enzymes and its isolation and identification, as well as the enhancement of cellulase enzyme activity under optimal operating conditions, constitute this research’s uniqueness.

2 Materials and methods

2.1 Sample collection

Sampling points were chosen from three different alkaline areas at UniMAP Agrotech (6.6522°N, 100.2609°E), Institute at Sungai Chuchuh, (6.6493°N, 100.2574°E), Bukit Keteri (6.5174°N, 100.2450°E), and Sungai Jernih (6.5439°N, 100.2684°E). Samples were taken from 10 to 15 cm depth and were sieved through a 2 mm sieve. The samples were maintained at 4°C in sterile sampling bags for future use.

2.2 Isolation of AC fungi

Serial dilutions were used to isolate cellulolytic fungi from soil samples on Czapek-Dox agar medium. The medium consisted of (g·L\(^{-1}\)) 30 g sucrose, 2 g NaNO\(_3\), 1 g K\(_2\)HPO\(_4\), 0.05 g MgSO\(_4\), 0.5 g KCl, 0.01 g FeSO\(_4\), 15 g agar, and 1% of carboxymethyl cellulose (CMC). In a sterile centrifuge tube, 1 g of soil sample was added to 10 mL of sterile distilled water and vortex for 2 min on high speed. Dilutions up to 10\(^{-6}\) were prepared, and 100 µL of suspension spreaded on Czapek-Dox agar plates with varying pH values (pH 8, 9, 10, and 11) for 7 days at 30°C. The fungi produced on plates were labelled as BK1 (Bukit Keteri 1), BK2 (Bukit Keteri 2), SC1 (Sg. Chuchuh 1), SC2 (Sg. Chuchuh 2), and BJ1 (Bukit Jernih 1) and inoculated on Potato Dextrose Agar (PDA) plates at 30°C for 7 days to obtain a single colony and stored under 4°C for future use.
2.3 Screening of AC fungi

The hydrolysis capacity (HC) test with Gram’s iodine was used to screen AC fungi [18]. In this approach, a week-old culture fungi was immersed in 15 mL of sterile distilled water. In the middle of Czapek-Dox agar plates (pH 8, 9, 10, and 11), a tiny well was formed, dilutions were made up to 10^{-6}, and 100 µL of spore suspension was injected. The plates were then incubated at 30°C for 48 h. After that, agar plates were flooded for 3–5 min with Gram’s iodine mix, which contained 2.0 g KI and 1.0 g iodine in 300 mL of distilled water. The diameter of the cleared zone was measured and used as an indicator of cellulase activity.

\[
HC = \frac{\text{Diameter of clearing zone}}{\text{Diameter of colony}}
\]  

(1)

2.4 Optimization of cellulose hydrolysis using one-factor-at-a-time (OFAT) method

Typically, the saccharification process was carried out in 50 mL Erlenmeyer flasks that included microcrystalline cellulose (MCC), enzyme extracts, and certain buffers based on the pH. After that, flasks were incubated with shaking at 180 rpm, and 200 L hydrolysis samples was taken once every 8 h from each flask. Precisely, the OFAT method was used to determine the optimal level of parameters for cellulose hydrolysis. Total enzyme loading, microcrystalline cellulose (MCC) concentration, pH, and temperature were the four key characteristics that could affect cellulase hydrolysis in this study. The optimum total enzyme loading of crude cellulase for the hydrolysis of MCC in 0.05 M glycine–NaOH buffer (pH 9.0) was determined by incubating the mixture of 1% (w/v) MCC with different concentrations of crude enzyme ranging from 0.3 to 1.2 U at 50°C and 180 rpm. To optimize MCC concentration, the study varied from 1% to 4% (w/v). 0.05 M phosphate buffer (pH 7), 0.05 M Tris-HCl buffer (pH 8), and 0.05 M glycine–NaOH buffer (pH 9 and pH 10) were utilised to examine the effects of pH (7–10). The temperature effect was evaluated by varying the saccharification temperature from 30°C to 90°C. All the experimental runs were monitored their sugar production for 48 h (1, 3, 6, 12, 24, 36, and 48 h). The glucose standard curve was constructed according to the study by Miller et al. [19]. Then, cellulase activity was assayed by the DNS method.

2.5 Analytical methods

2.5.1 Determination of fungi biomass

The best four strains resulted from the HC test were grown at pH 9.0, temperature 30°C, and agitated at 150 rpm in the potato dextrose broth (PDB) medium. Each strain was added to broth medium with 10% (v/v) of inoculum concentration with 10^7 of total spore. The saccharification media was recovered every 24 h and underwent 10 min centrifugal process at 10,000 rpm under 4°C environments. The residual cell (residue) was dried at 70°C for 24 h, and cell’s dry weight was measured to determine the growth activity.

2.5.2 Cellulase assay

The cellulase activity was measured using crude cellulase extracted from clear saccharification supernatant. The cellulase activity was measured using Ghose’s technique [20]. In a test tube, a rolled strip of Whatmen No.1 filter paper measuring 1.0 × 6.0 cm (50 mg) was dipped in 1 mL of glycine–NaOH buffer (0.05 M, pH 9.0). The mixture was then incubated at 50°C for 60 min with 0.5 mL of crude cellulase. After removing the tubes from the water bath, the reaction was halted by adding 3 mL of 3,5-dinitrosalicylic acid reagent (DNS) to each test tube. The tubes were then boiled for 5 min at 95°C for the colour development. After boiling, all the tubes were transferred into a cold-water bath. All reaction mixtures were diluted with distilled water appropriately (0.2 mL of reaction mixture added with 2.5 mL distilled water in a spectrophotometer cuvette). The absorbance of triplicates colour-formed reaction mixtures was measured against spectrometer zero at 540 nm wavelength, and the average value was taken.

2.5.3 Cell phenotypic identification

After 2 days of incubation, the colony was observed under macroscopic characterizations which are the structural of hyphae and basidiospore. One drop of sterile distilled water and a loop of colony were transferred onto a microscope glass slide. The glass slide was heat fixed on Bunsen burner flames several times. Methylene blue was added onto the smear for staining and washed under slow running tap water. The slide was left to air dry; the colony was observed under high-performance microscope for mor-
phological identification. Later, portions of fungi mycelia were taken from the agar dish to a 1.5 mL microcentrifuge tube containing 50 L autoclaved-ultrapure water using a sterile toothpick. The cell suspension was mixed by the vortex method vigorously for 5 min and then incubated at 95°C for 10 min. After that, the microcentrifuge tube was quickly frozen at −20°C for 10 min. For PCR amplification, the suspension was centrifuged at 11,000 rpm for 5 min, and the 5–10 µL of supernatant was used as the DNA template.

2.5.4 Cell molecular and phylogenetic analysis

For PCR amplification in cell molecular and phylogenetic analysis, the test began by designing specific primers for the target gene or region. In a 25 µL reaction, 1 µL of DNA template was combined with forward and reverse primers (0.5 µL each), Taq DNA polymerase (0.25 µL), dNTP mix (1 µL), and 10× PCR buffer (2.5 µL), with MgCl₂ adjusted accordingly. Thirty-five cycles of denaturation at 95°C for 30 s, primer-specific annealing for 30 s, and extension at 72°C were carried out for a time based on the expected amplicon size. The process concluded with a final extension at 72°C for 10 min. The PCR products were then verified on an agarose gel, purified using a DNA kit, and sequenced using the ABI 3,730/3,730 DNA analyzer. Taxonomic identities were confirmed with Basic Local Alignment Search Tool for Nucleotide (BLASTN) against the National Center for Biotechnology Information (NCBI) nonredundant (nr) database, and phylogenetic analysis was conducted using NCBI BLAST Tree View with the Fast Minimum Evolution tree method. TIANquick Mini Purification Kit (TIANGEN, China) was used to purify PCR products as directed by the manufacturer, while ABI 3,730/3,730 × 1 DNA Analyzer was used for DNA sequencing (Life Technologies, USA). The taxonomic identities of the acquired sequences were confirmed against the nr nucleotide database of the NCBI using the BLASTN databases based on nucleotide query 2.23 1+. Phylogenetic analysis was conducted using NCBI BLAST Tree View with the fast minimum evolution tree method.

2.6 Kinetic modelling of alkaliphilic cellulolytic fungi

2.6.1 Logistic kinetic models

The logistic equation has been used to describe the cell development in several saccharification processes, including the breakdown of cellulose by the AC fungi. Some derivations have been developed to employ the logistic model (Table 1) in the Polymath software, and the model encompasses both the exponential and stationary stages, as shown in Eq. 2 [12]:

$$X(t) = \frac{X_m e^{\mu m t}}{1 - \frac{X_m}{X_m} (1 - e^{\mu m t})}$$

(2)

where $X$ represents the biomass concentration (mg g⁻¹ substrate) at time $t$ (h), $m$ indicates the maximum specific growth rate (1/h), $X_m$ denotes the beginning biomass concentration, and $X_m$ marks the maximum biomass concentration. Non-linear regression was used to fit the model to the experimental data. Samsudin and Mat Don et al. [21], Don and Shoparwe [22], Pazouki et al. [23], Elibol and Mavituna [24] also employed these models. The objective is to use mathematical modelling to convert a complex (biological) entity to a simplified (mathematical) version that can be studied more thoroughly with significant properties established. To estimate the initial condition value, linearization was done for the logistic model (Table 1). Experimental result was analyzed using the linearized equations of the model.

3 Results and discussion

3.1 Isolation and screening of AC fungi

The soil sample was taken from three different identified limestone areas which were Sungai Chucuh, Bukit Keteri, and Bukit Jernih in Perlis. The soil pH of these alkaline areas was tested by the soil pH meter. The pH obtained was above 9.0. The fungi from the soils were grown on the petri dish containing CMC agar. After 7 days of incubation, there were five different species observed. To have a pure strain of the species, the isolation method for the samples was done and cultured on the PDA. The collected samples

<table>
<thead>
<tr>
<th>Table 1: Kinetic growth models selected and their linear form</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Logistic model</strong></td>
</tr>
<tr>
<td>$\frac{dX}{dt} = \mu_m \left(1 - \frac{X}{X_m}\right)X$</td>
</tr>
<tr>
<td>Integration from $t = 0, X = X_0$</td>
</tr>
<tr>
<td>$X = \frac{X_m e^{\mu m t}}{1 - \frac{X_m}{X_m} e^{\mu m t}}$</td>
</tr>
<tr>
<td>Linear form</td>
</tr>
<tr>
<td>$\ln \left(\frac{X}{X_m - X}\right) = \mu m t - \ln \frac{X_m}{X_m - 1}$</td>
</tr>
</tbody>
</table>
were named based on areas which were BK – Bukit Keteri, BJ – Bukit Jernih, and SC – Sg. Chucuh. These five sample strains were used for further study. Figure 1 shows the observation of AC cultivation on the petri dish.

### 3.2 Evaluation of AC fungi in producing alkaline cellulases

Determination of AC strain was done by calculating the HC on different pH of CMC agar plates. The higher value of HC shows the ability of the strain tolerant with a high pH medium. In this study, the collected sample strain was tested, and *Aspergillus niger* (strain from Bioprocess Laboratory, UniMAP) was used as the comparison factor. These five sample strains were then exposed to plate screening on CMC agar media at pH 8, 9, 10, and 11 using Gram’s iodine as a colouring agent. When Gram’s iodine was added to colonies, they became discoloured, as shown in Figure 2. The ability of the strain to release cellulase enzyme at high pH was demonstrated by the hydrolyzed zone around the colonies. The clearing zone around the colonies, which corresponds to the zone of CMC degradation, can also be utilized.

![Figure 1: Isolated fungi from CMC agar plates.](image-url)
to estimate HC as shown in Eq. 1 for further comparative testing between the strains. The HC of isolated fungi is depicted in Table 2.

Based on the results obtained, there are only five sample strains that give positive results. These five strains (BK1, BK2, SC1, SC2, and BJ1) exhibit higher HC values compared to A. niger (commercial strain) within the studied range of pH levels. This exemplifies the alkalophilic traits of the fungi, which favours high pH conditions for both growth and activity [25]. Furthermore, the breakdown of cellulose at high pH conditions suggests that these fungi have the capacity to release extracellular alkaline-tolerant cellulase. Nevertheless, Ahamed and Vemette claim that this qualitative screening was insufficient to draw a conclusion regarding the success of alkaline cellulase production because only a small number of cellulolytic microorganisms were capable of depolymerizing crystalline cellulose effectively [26]. Furthermore, Liang et al. found no real relationship between cellulolytic indicator and cellulase activity in a variety of liquid media containing CMC and other cellulosic components. Liang and his colleagues also stated that the enzyme generated by these strains was insufficient to detect when cultured in the liquid medium or that the fungi’s ability to produce CMC was ineffective when cultivated on agar plates [27]. Agrawal, on the other hand, discovered that the length of the hydrolyzing domain may not correctly reflect the actual cellulase activity [28]. A much more precise quantitative cellulase screening is required to determine the enzymatic degradation of cellulose into fermentable sugar.

### 3.3 Production of alkaline cellulases by saccharification

The best AC fungi strains then were further screened by analyzing alkaline cellulases produced under the saccharification process. The sample AC fungi strains were

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**Table 2: HC of isolated fungi in different pH**

<table>
<thead>
<tr>
<th>Strains</th>
<th>CMC media pH 8</th>
<th>CMC media pH 9</th>
<th>CMC media pH 10</th>
<th>CMC media pH 11</th>
</tr>
</thead>
<tbody>
<tr>
<td>BK 1</td>
<td>1.6778 ± 0.32</td>
<td>1.8074 ± 0.19</td>
<td>1.5297 ± 0.25</td>
<td>1.3148 ± 0.23</td>
</tr>
<tr>
<td>BK 2</td>
<td>1.0815 ± 0.21</td>
<td>0.6963 ± 0.09</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>SC 1</td>
<td>1.2735 ± 0.18</td>
<td>1.2272 ± 0.30</td>
<td>1.2276 ± 0.16</td>
<td>1.1218 ± 0.11</td>
</tr>
<tr>
<td>SC 2</td>
<td>1.2124 ± 0.21</td>
<td>1.2284 ± 0.16</td>
<td>0.8335 ± 0.27</td>
<td>1.2367 ± 0.19</td>
</tr>
<tr>
<td>BJ 1</td>
<td>1.1484 ± 0.17</td>
<td>1.0906 ± 0.22</td>
<td>1.1415 ± 0.16</td>
<td>1.0682 ± 0.31</td>
</tr>
</tbody>
</table>

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**Figure 2:** Clearing zone on CMC agar plate.

**Figure 3:** Average value of cellulase activity (U·mL\(^{-1}\)) of strains under 2 g·L\(^{-1}\) of CMC concentration after triplicate.
cultivated at 30°C for 7 days in PDB media with a pH of 9 and a mixture of CMC as a carbon source, with varying concentrations of CMC ranging from 1% to 4% employed in the saccharification medium. Figures 3–5 prove that BK1 strain was the best cellulose-forming fungi where with all the tested conditions, BK1 recorded the highest cellulase activity with 0.03 U·mL$^{-1}$ (second day on 2 g·L$^{-1}$ CMC concentration), 0.0463 U·mL$^{-1}$ (fourth day on 4 g·L$^{-1}$ CMC concentration), and 0.0532 U·mL$^{-1}$ (third day on 6 g·L$^{-1}$ CMC concentration). Furthermore, Zahra et al. investigated the optimal CMCase activity and filter paper cellulase activity after 4 days of incubation at 30°C in media containing 5% substrate concentration, 0.05% peptone, and 0.5% KH$_2$PO$_4$ concentration [29]. This also explains the behaviour and the role of CMC as a carbon source in improving the enzyme activity. It was reported that CMC as a substrate induced the production of cellulases by activating cellulase regulatory protein called the cellulase activator molecule [30]. While BJ1 recorded the lowest performance among all the strains with the highest for BJ1 were 0.029 U·mL$^{-1}$ (2nd day on 2 g·L$^{-1}$ CMC concentration), 0.0250 U·mL$^{-1}$ (2nd day on 4 g·L$^{-1}$ CMC concentration), and 0.02 U·mL$^{-1}$ (2nd day on 6 g·L$^{-1}$ CMC concentration). The observed low cellulase production in the study may be attributed to a combination of factors, including inherent variability in cellulase production capabilities among different fungal strains (BK1 and BJ1), suboptimal concentrations of carboxymethyl cellulose (CMC) as a substrate, and potential limitations in the experimental conditions. The strain-specific genetic backgrounds of BK1 and BJ1 could influence cellulase gene

![Figure 4: Average value cellulase activity (U·mL$^{-1}$) of strains under 4 g·L$^{-1}$ of CMC concentration after triplicate.](https://example.com/f4.png)

![Figure 5: Average value cellulase activity (U·mL$^{-1}$) of strains under 6 g·L$^{-1}$ of CMC concentration after triplicate.](https://example.com/f5.png)
expression, impacting enzymatic productivity [31]. In addition, the chosen CMC concentrations may not have been optimal for cellulase induction, and further substrate optimization might be necessary. The relatively short cultivation period of 7 days might not provide sufficient time for some strains to reach peak cellulase production. Nutrient availability in the PDB medium and the absence of essential micronutrients could also play a role.

3.4 Optimization of cellulase activity (HC) using OFAT method

3.4.1 Effect of different total enzyme loading

The effect of total enzyme loading (5–20%) on the hydrolyzing cellulose by AC BK1 fungi was studied (Figure 6). The lowest glucose concentration was observed (0.2–0.6 mg·mL⁻¹) after 60 h saccharification using 5% and 20% enzyme loading. This might be because the enzyme's ability was insufficient to fully saccharify the cellulose, resulting in a diluted concentration of the enzyme available for saccharification. While saccharification with total enzyme loading at 15% illustrated a high rate of cellulose conversion to glucose around 1.2–1.3 mg·mL⁻¹. A similar trend was observed for parameter 10% enzyme loading, and this was due to the enzyme having the sufficient amount of active site to interact with the substrate. However, the conversion of cellulose using enzyme loading at 10% recorded around 0.8–0.9 mg·mL⁻¹ of glucose. The results showed that alkaline cellulase degraded cellulose, producing oligosaccharides that may enter the cell and induce cellulase expression and glucose production [31]. It was proven that high total enzyme loading affected the glucose production as the more enzymes used was not efficient due to high competition between the enzyme and thus leading to ineffective substrate–enzyme complex.

3.4.2 Effect of different microcrystalline cellulose concentration

The effect of microcrystalline cellulose concentration (1–4%) of the characterization on the hydrolyzing cellulase by AC BK1 fungi was studied (Figure 7). Using 4% concentration of microcrystalline cellulose rapidly boosted glucose concentration at 0.9 mg·mL⁻¹ within 4 h of the saccharification process. This may be due to the huge availability of cellulosic material in media, and thus, hydrolyzing the rate becomes effective. However, after 8 h of saccharification, using 4% concentration microcrystalline had decreased to 0.8 mg·mL⁻¹ glucose concentration conversion. This may be due to the saturated active site of the enzyme, and thus, low performance of the conversion occurred [32]. While using 3% MCC concentration produced the least glucose concentration (around 0.6 mg·mL⁻¹) within 4 h of the saccharification process. In addition, the saccharification using 1% microcrystalline cellulose concentration led to the production of glucose once beyond the 8 h characterization. This was most probably because the enzyme still had an empty active site for the substrate to attach and simplify it to become glucose [33].

3.4.3 Effect of different pH

The effect of pH (7–10) of the enzymatic saccharification medium on the hydrolyzing cellulose by AC BK1 fungi was studied (Figure 8). At the pH values of 7.0 and 8.0,
they recorded the lowest glucose concentration around 1.15–1.2 mg·mL⁻¹, which was observed after 24 h of saccharification at pH of the medium between 7 and 8. As for the saccharification medium at pH 9, it produced a high rate of cellulose conversion to glucose (1.2–1.25 mg·mL⁻¹; 24% increment), after 4–8 h of the saccharification process. But it slightly dropped at 12 h of saccharification. It may be due to the limited active site of enzyme alkaline cellulase. Saccharification with pH 10 obtained highest glucose concentration with 1.35 mg·mL⁻¹ after 24 h. This proves that this AC fungi favours and is active under alkaline condition [29]. Therefore, it is important that the saccharification is operated at the optimum pH condition to obtain the highest production (glucose). A similar observation was reported by other researchers who used alkaline cellulose from fungi strains.

According to Vega et al. [25], the cellulolytic fungi extracted from the Peru tropical rain forest shows the activity performing well at pH 9.4 \((\text{Penicillium sp. LM-HP06} (12.9 \text{ U·g}^{-1}\text{·h}^{-1}), \text{Penicillium sp. LM-HP34} (13.8 \text{ U·g}^{-1}\text{·h}^{-1})).\) It demonstrates how these species have a strong aptitude for cellulose conversion under alkaline conditions. When compared to other
alkaline cellulases, the endo-ß-glucanase from the yeast *Rhodotorula glutinis* was generally stable at a pH range of 2–9, whereas alkaliphilic *Bacillus* sp No 1139 was stable up to pH 11 [34].

### 3.4.4 Effect of different temperature

The effect of saccharification temperature (30–60°C) on the hydrolyzing cellulose by AC BK1 fungi was studied (Figure 9). There were gradual increases in the production of glucose concentration for all the runs. The highest generation of glucose concentration was obtained at temperature of 50°C with 1.28 mg·mL⁻¹ after 24 h characterization. This proves the ability of the cellulase to tolerate high temperature [35]. Additionally, when applied to difficult industrial environments, the extremophile’s ability to stabilise under extreme conditions adds value. Other than the above, saccharification of alkaline cellulase BK1 at 30°C illustrates low production of glucose concentration (0.73 mg·mL⁻¹). This could be due to the need for temperature to boost the reaction rate and hydrolyzing rate of alkaline cellulase. Too high temperatures do not ensure the increased cellulase conversion activity. Temperature-sensitive enzyme components, such as protein, may suffer irreparable damage. This is described in the saccharification graph at 60°C. According to Kasana et al. [36], *R. glutinis* may grow at temperatures ranging from 4°C until slightly below 30°C. The best temperature for growth was 20°C, but its cellulase was active at temperatures ranging from 4°C to 70°C, with a peak at 50°C which was proven in Bai et al.’s research, thermophilic cellulase from Penicillium sp. CR316 has an optimal temperature of 65°C, and the activity is steady at that temperature for 3 h [37]. In comparison to other alkaline cellulases, the optimum temperature of cellulase activity was discovered to vary depending on the organism. While alkaline cellulases from thermophilic species functioned better at higher temperatures, cellulases from *Bacillus* sp. KSM-635 displayed highest the activity at 40°C, identical to the cellulase in this experiment [38]. Aside from that, alkaline cellulases from thermophilic organisms performed best at high temperatures, such as 60°C for both *Bacillus stearothermophilus* and *Clostridium josui* alkaline cellulases [39].

### 3.5 Kinetic modelling of alkaliphilic cellulolytic fungi

#### 3.5.1 Logistic kinetic model

For the kinetic expression, the logistic mathematical model was applied as shown in Figure 10. It was adjusted to the AC fungi biomass experimental results to investigate the BK1 growth. The model accurately illustrated the lag, exponential, and stationary phases. To calculate the EB’s specific growth rate (µ), the highest biomass, *Xₘ* = 0.4261 mg biomass·g⁻¹ sample from the experimental observations was used, and a plot of linear (Eq. 5) provided *µ* = 1.053 (day⁻¹). Then, the initial lag period of the current

![Figure 9: Glucose production at different temperatures.](image-url)
investigation was extended to 6–12 h. The extent of the lag phase can be influenced by a variety of factors, including medium composition (broth), the types and maturities of the AC fungi strain, the number of cells, and physical variables such as temperature and pH [27–29]. The AC fungi initiate an exponential phase after a lag phase, which begins between 1 and 5 days. The maximal EB biomass was estimated to be 29.65 mg·g\(^{-1}\) by the logistic model. The AC fungi biomass ceased expanding exponentially once the limiting cellulose ingredient in the broth began to diminish, followed by the stationary phase. The \(X_m - X_0\) of BK1 was 0.399 after 7 days saccharification, and HC proved that it had the highest ability for degradation of cellulose. It could be concluded that the production of enzyme secreted by AC BK1 fungi was very conducive.

Table 3: Kinetic parameter obtained from the Logistic models for AC fungi growth

<table>
<thead>
<tr>
<th>Kinetic parameters</th>
<th>AC, BK1</th>
</tr>
</thead>
<tbody>
<tr>
<td>(X_0)</td>
<td>0.027</td>
</tr>
<tr>
<td>(u_m)</td>
<td>1.053</td>
</tr>
<tr>
<td>(X_m)</td>
<td>0.4261</td>
</tr>
<tr>
<td>(X_m - X_0)</td>
<td>0.399</td>
</tr>
<tr>
<td>(R^2)</td>
<td>0.9963</td>
</tr>
<tr>
<td>RMSE</td>
<td>0.0032</td>
</tr>
</tbody>
</table>

Figure 10: Simulated and experimental data for alkaline cellulase-producing (AC) BK1 fungi growth.

Figure 11: Phylogenetic trees for BK species (Query_10143).
and effective. The POLYMATH software’s data prediction likewise coincided with the experimental output, with $R^2$ and RMSE values of 0.9963 and 0.32, respectively, as shown in Table 3.

### 3.5.2 Phylogenetic analysis

Four main samples of isolated AC fungi were sent to Microbiology Company, Genomics BioSci and Tech, Ltd., Taiwan, and the determined species were to be from the genus *Basidiomycetes* (BK1), *Trametes* (SC1), *Trametes* (SC2), and *Penicillium* (BJ1). Researchers believe that the presence of *Basidiomycetes* (BK1) species in the saccharification of cellulose can help the hydrolysis mechanism, as the secretion of enzyme cellulolytic enhanced the conversion cellulose to glucose, while for *Trametes* (SCI) and (SC2), it had proven by the researcher that *Trametes* secretes laccase which helps in hydrolyzing the cellulose and hemicellulose. Meanwhile, for *Penicillium* (BJ1) species, research managed to discover their potential for the cellulosic activity by comparing *Penicillium* with other fungi such as *Trichoderma reesei* species [40]. The four species’ phylogenetic trees are presented in Figures 11–14. The presence of these four species in this study proved that they had promising abilities that could be explored deeper in the future study.

Figure 12: Phylogenetic trees for BJ species (Query_58523).
4 Conclusions

The present work attempted to obtain optimum alkaline cellulase conditions using batch saccharification. The objectives of these studies were achieved by combining the experimental work, kinetic studies, and the modelling approach. A pure AC fungus strain with capability to produce the highest alkaline cellulase among the four positive samples was successfully isolated from Bukit Keteri (BK1) area. It consisted of highest amount of HC (pH 8: 1.6778, pH 9: 1.8889, pH 10: 1.5297, pH 11: 1.3148) result compared to Sg Chuchuh SC1 (pH 8: 1.2735, pH 9: 1.2273, pH 10: 1.2276, pH 11: 1.1218), SC2 (pH 8: 1.2124, pH 9: 1.2284, pH 10: 1.2305, pH 11: 1.2367), and Bukit Jernih (pH 8: 1.1484, pH 9: 1.0906, pH 10: 1.1415, pH 11: 1.0682), respectively. Diverse microbial communities of fungi are capable of secreting alkaline cellulase (ability to degrade the cellulose), and phylogenetic analysis proved the presence of genus *Basidiomycetes* (BK1), *Trametes* (SC1), *Trametes* (SC2), and *Penicillium* (BJ1). They actively secreted cellulase which plays an important role in hydroxylation of cellulose in the saccharification. The one-factor-at-one-time (OFAT) method was used to identify the ideal conditions for cellulase activity in the saccharification flask utilising AC fungi. The cellulase enzyme isolated from the BK1 AC fungus strain favoured and was active at high alkaline conditions (pH 9 and pH 10) according to the research results. In addition, the enzyme was still active at high temperatures, supporting its ability to withstand extreme heat. The greatest glucose generation from the saccharification process was also seen at 15% (v/v) enzyme loading and 1% microcrystalline cellulose concentration. The logistic model accurately represented the growth behaviour of AC fungi in saccharification, with $R^2$ varying from 0.9923 to 0.9987 and RMSE ranging from 0.0018 to 0.0066. From the findings, alkaline cellulase produced from the newly isolated alkaphilic cellulolytic fungi strain may have considerable potential for industrial application especially when operating in extreme conditions owing to its useful properties.

Figure 13: Phylogenetic trees for SC1 species (Query_55781).
Acknowledgement: The authors would like to thank the Ministry of Education for financing the research under the Fundamental Research Grant Scheme (FRGS/1/2018/STG01/UNIMAP/03/3), Universiti Malaysia Perlis for the financial support of this study via the Research University Grant, a scholarship (MyMasters) for the first author from the Ministry of Higher Education Malaysia. The authors are grateful to the Researchers Supporting Project Number (RSP2023R326), King Saud University, Riyadh, Saudi Arabia.

Funding information: The research was supported by the Fundamental Research Grant Scheme (FRGS/1/2018/STG01/UNIMAP/03/3). The authors are grateful to the Researchers Supporting Project Number (RSP2023R326), King Saud University, Riyadh, Saudi Arabia.

Author contributions: Conceptualization, NIZ, MMZM, AANG, SCBG; writing – original draft preparation, NIZ, MMZM; writing – review and editing, AAA, KP, MRS, MA, MR; supervision, MMZM, AANG, MR; funding acquisition, MMZM and MR. All authors have read and agreed to the submitted version of the manuscript.

Conflict of interest: The authors state no conflict of interest.

Data availability statement: The datasets generated during and/or analysed during the current study are available from the corresponding author on reasonable request.

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