A Novel Thermostable Keratinase from *Deinococcus geothermalis* with Potential Application in Feather Degradation

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Abstract: Keratinase can specifically attack disulfide bridges in keratin to convert them from complex to simplified forms. Keratinase thermal stability has drawn attention to various biotechnological industries. In this study, a keratinase DgeKer was identified from a slightly thermophilic species, *D. geothermalis*. The in silico analysis showed that DgeKer is composed of signal peptide, N-terminal propeptide, mature domain, and C-terminal extension. DgeKer and its C-terminal extension-truncated enzyme (DgeKer-C) were cloned and expressed in *E. coli*. The purified DgeKer and DgeKer-C showed maximum activity at 70 °C and pH 9. The thermal stability assay (60 °C) showed that the half-life value of DgeKer and DgeKer-C were 103.45 min and 169.10 min, respectively. DgeKer and DgeKer-C were stable at the range of pH from 9 to 11 and showed good tolerance to some metal ions, surfactants and organic solvent. Furthermore, DgeKer could degrade feathers at 70 °C for 60 min. However, the medium became turbid with obvious softening of barbules after being treated with DgeKer-C, which might be due to C-terminal extension. In summary, a thermostable keratinase DgeKer with high efficiency degradation of feathers may have great potential in industry.

Keywords: *Deinococcus geothermalis*; keratinase; thermostability; feather degradation; C-terminal extension

1. Introduction

Keratinase (EC 3.4.21/24/99.11) is a class of enzymes that can specifically degrade keratin, which is a structural protein used by animals for mechanical protection with sulfur-containing compounds and disulfide bonds. It can degrade feathers and other difficult-to-degrade keratinous substances with low energy consumption and environmental friendliness, which has application potential in husbandry [1, 2], medical [3– 5], cosmetics, biological materials [6] and other industries. Keratin is widely found in agricultural waste, especially bird feathers from poultry processing plants [7], and urgently needs to be processed and recycled. In the process of feather degradation, higher temperatures are favorable for the destruction of disulfide bonds in keratin, which improves hydrolysis efficiency. However, high temperature will destroy the non-covalent interaction of proteins and eventually lead to enzyme inactivation. Keratinolytic thermophilic or hyperthermophilic bacteria and archaea have been known to degrade keratin at ≥70 °C [8]. At present, many keratinases have been identified from microorganisms, but the number of keratinases used in the market is very small, which might be related to the thermal stability of keratinases [9]. Therefore, it is very important to find and identify high temperature-adaptive keratinases.

In order to find thermally stable keratinases, researchers tend to isolate keratinase bacteria from a high-temperature environment, such as hyperthermophilic archaeon *Thermococcus kodakaraensis* [10], the thermophile strain *Meiothermus taiwanensis* WR-220 (isolated...
from hot springs) [11], etc. The optimal temperature and pH of keratinase MtaKer from *M. taiwanensis* WR-220 are 65 °C and pH 10, respectively. MtaKer has keratinase activity in a larger temperature and pH range. In extreme environments, the protein encoded by extremophiles should evolve into functional proteins to adapt to the environment. The strong tolerance of MtaKer may be related to the extreme living environment of *M. taiwanensis* WR-Many recent studies have focused on extremozymes, especially the high temperature-resistant enzymes, because adjustment to industrial harsh conditions needs the activity and stability of the applied enzymes in such conditions [12,13]. Since evolution and natural selection have occurred in the environment for billions of years, extreme bacteria may produce customized enzymes with some unique properties required for industrial applications. *Deinococcus* bacteria are famous for their radiation tolerance, however, the peptidase genes present in their genome are ignored. *D. geothermalis* was discovered in hot springs, and the optimal growth temperature of *D. geothermalis* is 50 °C, higher than the growth temperature of medium-temperature bacteria. The enzymes of *D. geothermalis* might have good tolerance to extreme industrial conditions, especially thermostability.

In this study, the keratinase gene *Dgeo_0978*, named dgeKer, was identified from the *D. geothermalis* genome. The keratinase DgeKer (GenBank No. WP_011530115.1) from *D. geothermalis* was successfully expressed and purification in *E. coli* BL21-codonplus (DE3) strains. The biochemical properties of the recombinant enzyme and the ability in degrading feathers were examined. The results showed that DgeKer had high activity under high temperature and pH conditions, which making it a promising enzyme for industry.

2. Materials and Methods

2.1. Strains, Plasmids and Materials for Gene Cloning and Expression

*Deinococcus geothermalis* DSM 11300 was cultured at 50 °C for 24 h in TGY media (10 g/L typtone, 1 g/L glucose, 5 g/L yeast extract) [14]. *E. coli* BL21-codonplus (DE3) was cultured in Luria-Bertani (LB) at 37°C. The vector pET22b was used for expression of keratinase genes in *E. coli* BL21-codonplus. All the enzymes for DNA manipulations were purchased from vazyme (Nanjing, China). All chemicals used in this study were of reagent grade.

2.2. Sequence Analysis

The gene sequence of DgeKer was analyzed by the NCBI (National Center for Biotechnology Information) and InterPro. The signal peptide prediction was performed by the SignalP 5.0 server. The alignment analysis of amino acid sequences was performed by ESPript3. The phylogenetic tree of protein sequences was constructed by MEGA 6.0 with the neighbor-joining (NJ) method.

2.3. Plasmids Construction

The genome of *D. geothermalis* was used as a DNA template for the DgeKer gene cloning. The keratinase genes without the signal sequence were PCR amplified using 2 × phanta max master mix with the instructions and the primers in Table 1. The amplified fragments were cloned into a pET22b expression vector (Novagen) containing the C-terminal His-tag using vazyme ClonExpress Ultra One Step Cloning Kit. The correctly recombinant plasmids were transformed into *E. coli* BL21-codonplus (DE3) competent cells for gene expression. The construction of the keratinase without C-terminal extension (DgeKer-C) was the same as DgeKer.
Table 1. Primers used in this study.

| Genes     | Primer Sequences (5′ to 3′)                                                                 |
|-----------|---------------------------------------------------------------------------------------------|
| dgeKer    | F1: 5′tgcagctcagcagcaagtATTGTCCGGACACCGGGAATGCGGACAGCAGGCGAATTTGTA                      |
|           | R1: 5′gtgcagctcagcagcaagtATTGTCCGGACACCGGGAATGCGGACAGCAGGCGAATTTGTA                    |
| dgeKer-C  | F1: 5′tgcagctcagcagcaagtATTGTCCGGACACCGGGAATGCGGACAGCAGGCGAATTTGTA                    |
|           | R2: 5′gtgcagctcagcagcaagtATTGTCCGGACACCGGGAATGCGGACAGCAGGCGAATTTGTA                    |

The lowercase letters indicate pET22b vector homology arms.

2.4. Expression and Purification of Keratinases

The induction and purification of keratinases was carried out according to Cheng Zhou et al. [15] with some modification. *E. coli* BL21-codonplus harboring pET22b-DgeKer was used for recombinant keratinase production. The recombinant *E. coli* was cultured overnight at 37 °C in 20 mL of LB medium containing 50 μg/mL of ampicillin (Amp) and 34 μg/mL of chloroamphenicol (Cm). The cultures were transferred into 150 mL of fresh medium and induced with 0.1 mM isopropyl-β-D-thiogalactopyranoside (IPTG) at OD600 = 0.6–0.8, followed by overnight expression at 16 °C. The cells were harvested by centrifugation at 4000 × g for 15 min and resuspended with Tris-HCl, and disrupted on ice by an ultrasonicator. The debris was removed by centrifugation at 12,000 × g for 30 min. The recombinant keratinases were purified by Ni-affinity chromatography (Qiagen) and eluted with an increasing imidazole gradient. The molecular weight of purified DgeKer was determined by sodium dodecyl sulfate polyacrylamide gel electrophoresis (SDS-PAGE).

2.5. Keratinolytic Activity Assay

Keratinolytic activity was measured with soluble keratin through the modified method by Jin-Song Gong [16]. Soluble keratin was purchased from Tokyo Chemical Industry (Tokyo, Japan). The reaction mixture, which contained 20 μL enzyme, 80 μL buffer and 100 μL soluble keratin (1%, w/v), was incubated at 50 °C for 20 min. A blank control was conducted by adding 200 μL of 20 % (w/v) trichloroacetic acid (TCA) instead of soluble keratin. The reaction was terminated by the addition of 200 μL TCA, and centrifuged at 12,000 rpm for 10 min. We added 100 μL soluble keratin to the control mixture. Then, 200 μL supernatant was pipetted into another tube with 1 mL 0.4 mol/L Na2CO3 and 200 μL Folin-phenol regent. Finally, the mixtures were incubated at 40 °C for 20 min to develop the color. Proteolytic products in the mixture were measured spectrophotometrically at 680 nm. One unit (U) of keratinase activity was defined as an increase of 0.01 absorbance unit at 680 nm under the aforementioned conditions. All assays were performed in triplicate.

The determination of feather powder activity was carried out according to Lian Yang et al. [17] with some modification. The reaction system including 4 mL buffer, 0.01 g feather powder and 200 μL enzyme with 5 mmol/L dithiothreitol (DTT) was incubated at 70 °C for 60 min and subsequently terminated by adding 2.0 mL of 20% TCA. After centrifugation at 12,000 × g for 5 min, the supernatant was gathered for measuring the absorbance at 280 nm. TCA (20%) was added before the enzymatic reaction as the control. One unit (U) of feather powder activity was defined as an increase of 0.01 absorbance unit at 280 nm. All assays were performed in triplicate. Protein concentration was determined using the Bradford method with bovine serum albumin as a standard.

2.6. Effects of pH, Temperature, and Reagents on Enzyme Activity and Stability

The optimal reaction temperature of DgeKer and DgeKer-C was investigated in the temperature range 30–90 °C, and the optimal pH experiment was carried out over a pH range of 3–11 (pH 3.0–6.0, 50 mmol/L citrate buffer, pH 7–9, 50 mmol/L Tris-HCl, pH 10–11, 50 mmol/L Gly-NaOH), using 1% soluble keratin. In thermostability assays, enzymes were pre-incubated at temperatures of 30 °C to 80 °C for 1 h, and residual enzyme activities was determined. The pH stability was measured by pre-incubating DgeKer and DgeKer-C in different pH solutions at 4 °C for 1 h, and then the residual activities were detected.
The effect of metal ions on enzyme activity was analyzed by assaying the relative activity with addition of 1.0 mmol/L and 5.0 mmol/L of metal ions (K⁺, Li⁺, Mn²⁺, Cu²⁺, Fe²⁺, Ca²⁺, Co²⁺, Mg²⁺, Ni²⁺, Cd²⁺, Cr³⁺). Similarly, the effect of some surfactants and organic solvent on enzyme activity were also analyzed. We added 1% dimethyl sulfoxide (DMSO), 1% β-mercaptoethanol, 1% tween-20, 1% tween-80 and 1% sodium dodecyl sulfate (SDS) (v/v) for the relative activity evaluation.

2.7. Chicken Feather Degradation In Vitro

Chicken feathers degradation was carried out according to Yaxin Ren [18] with some modifications. Chicken feather was previously washed and cleaned with detergent and tap water, followed by soaking in 70% ethanol for 1 h and drying. After that, it was autoclaved at 121 °C for 30 min to remove microorganisms for following use. The intact chicken feathers were incubated with 0.06 mg enzyme at 70 °C with 5 mmol/L DTT. The degradation process of feathers was recorded regularly.

2.8. Statistical Analysis

The results were analyzed with GraphPad Prism 8.0.2 and SPSS Statistics 25 software. Differences in means between groups were compared for statistical significance at

3. Results

3.1. Sequence Analysis of DgeKer

The gene dgeKer (1545 bp) encodes a protein comprising 514 amino acids with a calculated mass of 50.0 kDa. The amino acid sequence of DgeKer was compared with those of keratinases, which were KerA from Bacillus licheniformis PWD-1, MtaKer from M. taiwanensis WR-220 and aqualysin-I from Thermus aquaticus YT-1 in Figure 1. It was proposed that DgeKer consists of four parts, including signal peptide (Pre), N-terminal pro-peptide (N-pro), mature domain with a three highly conserved catalytic triad residues which is formed by Asp171, His203 and Ser354 (black frame). Additionally, DgeKer has a C-terminal extension like the thermophilic alkaline serine protease aqualysin-I, while the classic keratinase KerA does not contain a C-terminal extension.

Phylogenetic analysis of DgeKer and other homologous proteins was constructed based on DgeKer protein sequence. As shown in Figure 2, the DgeKer showed significant relatedness to peptidase from Deinococcus bacteria: they were localized at the same branch in the phylogenetic tree, indicating a common evolutionary origin. In addition, DgeKer had a certain homology with the keratinase from the Thermus, such as aqualysin I (WP_053768483.1) derived from Thermus aquaticus, with 53.49% identity. The protein sequence of DgeKer displayed similarity to MtaKer from M. taiwanensis WR-220 (GenBank No. AWR86689.1) and KerA (GenBank No. AAB34259.1) from Bacillus licheniformis with 58.96% and 40.20% identity, respectively. Based on the phylogenetic tree and sequence identity analysis, we concluded that DgeKer is a keratinase.

3.2. Expression and Purification of Recombinant DgeKer and DgeKer-C

E. coli is an ideal strain for protein heterologous expression. In order to obtain purified DgeKer and DgeKer-C (truncated form of the DgeKer without the C-terminal extension), recombinant plasmids carrying keratinase genes without signal peptide were constructed for E. coli expression system. As shown in Figure 3, DgeKer and DgeKer-C were expressed in E. coli BL21-codonplus (DE3) strains and purified successfully. The cloned DgeKer and DgeKer-C have expected molecular weight of 50.0 kDa and 38.4 kDa, respectively. However, the protein bands were detected in the SDS-PAGE after purification by Ni-affinity chromatography and migrated at approximately 38 kDa and 27 kDa, which correspond to the molecular mass of the mature keratinases without N-propeptide. Beside the major bands, protein bands at lower molecular weights (approximately 10–15 kDa) were detected.
Figure 1. Alignment of the amino acid sequences of DgeKer. The keratinases used were KerA from *B. licheniformis* PWD-1, MtaKer from *M. taiwanensis* WR-220, etc. Sequence alignment showing structural elements of DgeKer were generated with ESPript3. The identical residues of all aligned proteins are shaded red, conserved residues are presented in red, and conserved regions are presented in blue boxes. α, β, η, TT above the sequences represent α helix, β sheet, 310-helices and β-turns, respectively. Positions of the starting residues of the signal peptide (pre), N-terminal pro-peptide (N-pro), mature protease (mature), and C-terminal pro-peptide (C-pro) are marked by black arrows at the bottom of the sequences.
Figure 2. Phylogenetic analysis of amino acid sequence of keratinases. The keratinases used were KerA from *B. licheniformis* PWD-1, MtaKer from *M. taiwanensis* WR-220 and other homologous protein based on amino acid sequences. The name and GenBank accession No. for each protease are in front of the Latin name of each strain. The phylogenetic tree derived by the neighbor-joining method. The percentages of replicate trees in which the associated taxa clustered together in the bootstrap test (1000 replicates) are shown next to the branches.

![Phylogenetic tree](image)

**Figure 3.** Sodium dodecyl sulfate polyacrylamide gel electrophoresis (SDS-PAGE) analysis of DgeKer (a) and DgeKer-C (b). (a) M, protein MW maker; lane 1, *E. coli* BL21-codonplus (DE3) harboring pET22b for control; lane 2, pET22b-DgeKer for control; lane 3, DgeKer crude extract; lane 4, purified keratinase DgeKer (arrow). (b) M, protein MW maker; lane 1, *E. coli* BL21-codonplus (DE3) harboring pET22b for control; lane 2, pET22b-DgeKer-C for control; lane 3, DgeKer-C crude extract; lane 4, purified keratinase DgeKer-C (arrow).
3.3. Effects of Temperature and pH on Activity and Stability

In order to characterize the recombinant keratinases, the effects of temperature and pH on the recombinant keratinases was determined in this study. We used 1% soluble keratin to measure the enzyme activity. As shown in Figure 4, the optimal temperature of DgeKer and DgeKer-C were 70 °C. Moreover, the thermostability of DgeKer and DgeKer-C was also characterized. The residual activity of DgeKer after being incubated at 60 °C for 90 min was approximately 54%, and that of DgeKer-C was 69%, indicating that both of them have thermal stability. As shown in Table 2, the half-life ($t_{1/2}$) of DgeKer at 60 °C was 103.45 min, which was longer than that of KerA (26.46 min) but shorter than that of DgeKer-C (169.10 min). At 70 °C, DgeKer-C also exhibited better temperature stability than DgeKer. Unfortunately, KerA completely lost its enzymatic activity after being incubated at 70 °C for 20 min (Figure 4c). In terms of pH, the effects of pH to DgeKer and DgeKer-C was about the same (Figure 4b). DgeKer and DgeKer-C showed maximum activity at around pH 9–DgeKer and DgeKer-C were sensitive in the range of pH 3–7, while they were stable in range of 8–DgeKer and DgeKer-C had approximately the whole activity after being treated at pH 11 for 60 min. The data appears to suggest that DgeKer and DgeKer-C have potential industry application, due to their outstanding thermostability and pH stability.

| Keratinases  | $t_{1/2}$ (min) |
|--------------|-----------------|
| DgeKer       | 103.45 ± 1.26 b |
| DgeKer-C     | 169.10 ± 8.20 c |
| KerA         | 26.46 ± 0.22 a  |

Superscript letter $^{a,b,c}$ down the column were used to indicate significant difference at $p < 0.05$.  

![Figure 4](image-url)  
Figure 4. Effects of temperature and pH on keratinase activity of DgeKer, DgeKer-C and KerA. (a) The effects of temperature for DgeKer, DgeKer-C and KerA. (b) The effects of pH for DgeKer, DgeKer-C and KerA. (c) The thermostability for DgeKer, DgeKer-C and KerA. (d) The pH stability for DgeKer, DgeKer-C and KerA. The filled triangle represents DgeKer, and the filled square represents DgeKer-C. Error bars represent standard deviations.
As shown in Table 3, recombinant keratinases DgeKer and DgeKer-C showed the same capabilities to hydrolyze soluble keratin and feather powder. With soluble keratin as substrate, DgeKer and DgeKer-C showed highest specific activity of 40,192.30 ± 230.48 U/mg and 40,408.89 ± 342.73 U/mg, respectively. DgeKer and DgeKer-C had almost the same enzyme activity on feather powder, approximately 1450 U/mg.

Table 3. Activities of different keratinases on several proteinaceous substrates.

| Substrate      | Activity of Different Keratinases (U/mg) |
|----------------|-----------------------------------------|
|                | DgeKer                  | DgeKer-C                 |
| Soluble keratin| 40,192.30 ± 230.48        | 40,408.89 ± 342.73       |
| Feather powder | 1494.78 ± 25.78           | 1462.54 ± 71.19          |

3.4. Effects of Metal Ions and Chemical Agents on Keratinases Stability

The effects of various metal ions with different concentrations (1 mmol/L and 5 mmol/L) on keratinolytic activity were studied and the results were shown in Table 4. Mg^{2+} and Li^{+} had stimulatory effect on residual activity of DgeKer; K^{+} caused increased activity of DgeKer (104.11 ± 0.14%). By contrast, other tested metal ions prompted lower stability of DgeKer. 1 mmol/L and 5 mmol/L Ca^{2+} affected the DgeKer stability with residual activity of 94.07 ± 0.72% and 83.06 ± 0.36%, respectively. DgeKer-C was stable (101.07 ± 1.37%) at 5 mmol/L Co^{2+}. The other tested metal ions had a different decrease on the stability of DgeKer-C with residual activity of 88.25 ± 0.58% after pretreatment with 5 mmol/L K^{+}. Unfortunately, DgeKer and DgeKer-C showed complete inhibition with 5 mmol/L Cr^{3+}. There was no significant difference (p > 0.05) between DgeKer and DgeKer-C after pretreatment with 1 mmol/L K^{+}, Ca^{2+}, Mn^{2+}, Ni^{2+} and 5 mmol/L Cd^{2+}. DgeKer showed better tolerance than DgeKer-C to Li^{+} and Mg^{2+}. However, DgeKer-C was more stable than DgeKer with the pretreatment Co^{2+} and Cu^{2+}. In general, DgeKer and DgeKer-C had good tolerance to some metal ions.

Table 4. Effects of various metal ions on keratinolytic activity of DgeKer and DgeKer-C.

| Metal Ions | Concentration (mmol/L) | Residual Activity (%) |
|------------|------------------------|-----------------------|
| Control    |                        | DgeKer                | DgeKer-C               |
| K^{+}      | 1                      | 100.00 ± 0.27          | 100.00 ± 3.11          |
|            | 91.77 ± 0.49           | 87.45 ± 1.41           |
| Li^{+}     | 1                      | 104.11 ± 0.14          | 88.60 ± 0.56           |
|            | 102.11 ± 0.59          | 86.60 ± 0.43           |
| Mg^{2+}    | 1                      | 113.97 ± 0.62          | 88.81 ± 1.72           |
|            | 103.35 ± 1.07          | 93.80 ± 1.12           |
| Ca^{2+}    | 1                      | 121.53 ± 0.75          | 87.09 ± 0.71           |
|            | 94.07 ± 0.72           | 92.13 ± 1.18           |
| Co^{2+}    | 1                      | 83.06 ± 0.36           | 90.65 ± 0.88           |
|            | 91.20 ± 1.66           | 99.20 ± 3.37           |
| Mn^{2+}    | 1                      | 92.82 ± 0.49           | 101.07 ± 1.37          |
|            | 85.07 ± 0.89           | 80.94 ± 0.45           |
| Ni^{2+}    | 1                      | 64.69 ± 0.27           | 60.39 ± 0.96           |
|            | 88.33 ± 0.82           | 86.67 ± 0.87           |
| Cd^{2+}    | 1                      | 74.93 ± 0.62           | 86.34 ± 0.83           |
|            | 63.16 ± 1.02           | 65.32 ± 1.23           |
Table 4. Cont.

| Metal Irons | Concentration (mmol/L) | Residual Activity (%) |
|-------------|------------------------|-----------------------|
|             |                        | DgeKer | DgeKer-C |
| Cu²⁺        | 1                      | 79.33 ± 0.14 f         | 86.36 ± 0.09 f* |
|             | 5                      | 57.42 ± 0.41 b         | 67.21 ± 0.71 c* |
| Cr³⁺        | 1                      | 86.41 ± 0.85 h         | 73.24 ± 0.31 d* |
|             | 5                      | 0 ± 0 a                | 0 ± 0 a          |

Each value was repeated in triple independent measurements. Superscript letter(s) a–q down the column were used to indicate significant difference at p < 0. The * represents a significant difference (p < 0.05) between DgeKer and DgeKer-C with the same pretreatment.

As shown in Table 5, all the surfactants and organic solvent employed negatively impacted on the stability of DgeKer and DgeKer-C, with more drastic effect obtained in the presence of β-mercaptoethanol. The surfactant (tween-80 and tween-20) only slightly inhibited DgeKer and DgeKer-C activity. The good tolerance of DgeKer and DgeKer-C to organic solvent DMSO was discovered. The residual activity of DgeKer and DgeKer-C was approximately retained 90%. Furthermore, DgeKer and DgeKer-C showed good stability in 1% SDS with residual activity of 83.16 ± 0.54% and 93.85 ± 0.56%, respectively. The good tolerance to surfactants as well as organic solvent make DgeKer and DgeKer-C potential enzymes for industrial application. DgeKer was more stable to tween-80 and tween-20 than DgeKer-C, but it was opposite to 1% SDS and DMSO.

Table 5. Effects of surfactants and organic solvent on keratinolytic activity.

| Chemical Reagents          | Concentration (v/v) | Residual Activity (%) |
|----------------------------|---------------------|-----------------------|
|                            |                     | DgeKer | DgeKer-C |
| Control                    |                     | 100.00 ± 0.27 d       | 100.00 ± 3.11 d |
| SDS                        | 1%                  | 83.16 ± 0.54 b        | 93.85 ± 0.56 c,d,* |
| Tween-80                   | 1%                  | 86.70 ± 0.62 c        | 64.59 ± 4.98 b,c,* |
| Tween-20                   | 1%                  | 82.68 ± 0.81 b        | 71.54 ± 1.00 b,c,* |
| DMSO                       | 1%                  | 87.75 ± 0.95 c        | 92.63 ± 0.56 c,d,* |
| β-mercaptoethanol          | 1%                  | 0 ± 0 a               | 15.36 ± 5.14 a,c,* |

Each value represents the mean of triple independent measurements. Superscript letter(s) a–d down the column were used to indicate significant difference at p < 0. The * represents a significant difference (p < 0.05) between DgeKer and DgeKer-C with the same pretreatment.

3.5. Chicken Feather Degradation of DgeKer and DgeKer-C In Vitro

The necessary steps for keratinase to degrade feathers are the breaking of disulfide bonds and subsequent enzymatic hydrolysis. In this study, a reducing agent of DTT (5 mmol/L) was added to the degradation system to reduce disulfide bonds of feathers. In addition, we detected the degradation of feathers by keratinases at a high temperature of 70 °C. Tris-HCl buffer without keratinase was used as a control. Figure 5 shows that DTT (Tris-HCl with DTT) did not have the ability to degrade intact feathers. Interestingly, the barbules on the feather were degraded by DgeKer after being incubated for 60 min in the presence of 5 mmol/L DTT, and significantly degraded after being incubated for 120 min. In contrast, the medium became turbid with obvious softening of barbules after being treated at the same time with DgeKer-C, indicating that DgeKer had stronger feather hydrolysis ability at 70 °C.
Each value represents the mean of triple independent measurements. Superscript letter(s) a–d down the column were used to indicate significant difference at $p < 0.05$. The * represents a significant difference ($p < 0.05$) between DgeKer and DgeKer-C with the same pretreatment.

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Figure 5. Visual observation of disintegration of intact feathers by the DgeKer and DgeKer-C. Tris-HCl is a control without keratinase. DTT is a reducing chemical reagent dithiothreitol, which was added to feather degradation system to reduce feather disulfide bonds. (a), The feather degradation in the absence of DTT. “−DTT” represents the system without DTT. (b), The feather degradation in the presence of DTT. “+DTT” represents the system with 5 mmol/L DTT. The reaction systems were incubated at 70 °C with shaking.

4. Discussion

Extremophile microorganisms are a source of enzymes with good tolerances to extreme conditions [19]. Many keratinases from extremophiles have attractive properties, such as Nocardiopsis [20] and Salicola Marasensis [21], etc. M. twanensis WR-220 is a thermophilic microorganism with a growth temperature of 55–65 °C, and the keratinase in this bacterium also has thermal stability [11]. Many studies have demonstrated the presence of peptidase in the extreme Deinococcus microorganisms [22], such as serine peptidase from D. geothermalis [23] and Deinococcus sp. D7000 [24]. The degrading feather temperature of Deinococcus fucus [25] (feather-degrading bacterium) and D. radiodurans R1 [22] was 30 °C, which is limited for industrial applications. In this study, DgeKer gene was found
in *D. geothermalis*, which grow at 50 °C [14]. According to Figure 2, DgeKer has high homology with peptidases from *Calidithermus* and *Thermus*. These bacteria were isolated from extreme environments such as hot springs and are known for their thermal stability. *Thermus* and *Calidithermus* bacteria generally grow at medium and high temperature, and their keratinases might have high temperature range and temperature stability, such as MtaKer [11] and aqualysin I [26]. The amylolucrase from *Calidithermus timidus* has high thermostability with a half-life of 1.09 h at 70 °C [27]. As mentioned above, DgeKer has a long C-terminal extension, as with aqualysin-I [26] and VPR from *Vibrio* sp. PA44 [28]. As Wan-Ling Wu, etc., reported, C-terminal extension only exists in some keratinases from *Thermus*, *Thermococcus*, *Acinetobacter*, *Vibrio* and *Deinococcus*, which are generally mesophilic and thermophilic bacteria or extreme [11] bacteria. Therefore, it is valuable to find high temperature-resistant keratinase from *D. geothermalis*.

For industrial needs, it is necessary to overexpress keratinase in *E. coli* and other cells [7,29]. In general, keratinase consists of signal peptide, N-terminal pro-peptide and catalytic domains, and some also carry an extra C-terminal extension [11,30]. The folding of keratinase requires the N-pro-peptide (intramolecular chaperones) to facilitate correct folding and regulate the activation of catalytic domains [31]. Many studies indicate that the N-pro-peptide function as an inhibitor of subtilisin to prevent premature protease activity. The peptide bonds between the intramolecular chaperones and the protease domain are automatically hydrolyzed after the protein folding is complete. However, intramolecular chaperones still bind to proteases in the form of inhibitory complexes [31]. As shown in Figure 3, DgeKer and DgeKer-C could autonomously cleave the N-terminal propeptide to become mature keratinase in *E. coli* which was consistent with previous [11,31] reports, and the minor bands below the main band might result from processing or autohydrolysis of the protease. However, the N-terminal cleavage process is currently unclear. Studies have shown that the cleavage process is related to the amino acid preference of the cleavage site [32]. The C-terminal extension is usually located at the C-terminal of some secreted bacterial peptidase, some proteins from metalloproteinase families M4, M9 and M28 and serine protease family S8 were found to contain the C-terminal extension. A phylogenetic tree was constructed based on the C-terminal extended sequence of DgeKer and those of other keratinases. As shown in Figure S1, the C-terminal extension of DgeKer was homologous with those of keratinases from *Deinococcus*. Obviously, the C-terminal extension was different from PPC (bacterial pre-peptidase C-terminal domain) and PKD (polycystic kidney disease). The modeled structure of C-terminal extension of DgeKer was also built (Figure S2) based on the structure of a keratinase (ProN-TK-SP) from *Thermococcus kodakaraensis* (3D structure data in Protein Data Bank, PDB ID: 3AFG) [33] which exhibited about 33% sequence identity with C-terminal extension of DgeKer in the PDB database. The special structure of C-terminal domain of DgeKer was characterized as β-jelly roll with six β-strands and loops (Figure S2). According to previous studies, PPC and PKD domains may inhibit keratinase activity [34], enhance heat resistance [10,33], and bind insoluble substrates [35]. However, there are also studies that have the opposite outcome [15]. In general, at present, the mechanism of C-terminal extension is still unclear [36]. Therefore, we constructed the C-terminal extension deletion mutant protein DgeKer-C to explore the effect of C-terminal extension on DgeKer thermal stability and pH stability.

As shown in Figure 4, DgeKer and DgeKer-C exhibited maximal activity at 70 °C, which was higher than most of the other keratinases [18,32], such as KerA-derived (50 °C) [30] and MtaKer (65 °C) [11]. In terms of thermostability, DgeKer-C performed better than DgeKer (Figure 4). As incubation time extended, the residual enzyme activity of DgeKer-C was higher than DgeKer. DgeKer-C was stable with half-life (*t*1/2) of 169.10 min, which was 6.4-fold more than KerA. In general, DgeKer and DgeKer-C have high optimal temperature and thermal stability. Feather degradation fermentation is a continuous process and the temperature stability of keratinase is very important for feather degradation. These properties give DgeKer and DgeKer-C greater industrial application potential.
The optimal pH of DgeKer and DgeKer-C was 9–10, which was higher than that of most keratinases, such as KerA (pH 8). In the pH range of 3–7, DgeKer and DgeKer-C were greatly affected by pH, and in the pH range of 7–11, neither of them was significantly affected by pH, indicating that DgeKer and DgeKer-C were stable in the pH range of 7–11. Studies have shown that the feather degradation process involves sulfitolysis and deamination [1,37], which leads to an elevated pH in the fermentation environment [38–40]. The pH of the feather degradation system increased to 8.8 [41]. Therefore, reaction pH and pH stability are important factors in industrial applications. The activity and stability of DgeKer under alkaline conditions give this keratinase a certain potential for industrial applications.

DgeKer and DgeKer-C retained high activity over several metal ions, surfactants, reducing agents and organic solvents. In generally, keratinase is stimulated by some metal ions (Ca\(^{2+}\) and Mg\(^{2+}\)) [3,42,43] which might maintain the active enzyme conformation and stabilize the enzyme–substrate complex [38]. Mg\(^{2+}\) increased the enzymatic activity of DgeKer, which is consistent with previous studies. Ca\(^{2+}\) had no positive effect on the enzymatic activity of DgeKer and DgeKer-C, although two hypothetical Ca\(^{2+}\) ion binding sites were predicted in their sequence by NCBI. In general, both DgeKer and DgeKer-C had good metal ion tolerance.

At present, there are several degradation hypotheses about the mechanism of microbial degradation of keratin, but the key step is the degradation of disulfide bonds, so that keratinase can better hydrolyze keratin. Reducing agent DTT can improve the activity of some keratinases [30]. It might be the reducing agent reacting with the substrate rather than keratinases [38]. DTT [44], Na\(_2\)SO\(_3\) [45], L-cysteine [16] can accelerate the degradation of feathers. In this experiment, 5 mmol/L DTT was always added to the feather degradation system to promote feather degradation. As shown in Figure 5, it could degrade feathers only when keratinase and DTT exit at the same time, which was consistent with the published feather degradation mechanism. Firstly, the disulfide bond of the feather is opened by the reducing agent, and then the keratinase will degrade [46]. Research has shown that C-terminal extension can facilitate enzymes attach to insoluble substrates to promote degradation [47]. DgeKer-C had the same ability to hydrolyze feather powder as DgeKer (Table 3). However, DgeKer degraded intact feathers faster and better than DgeKer-C, which may be attributed to C-terminal extension. Accordingly, the C-terminal residues of MatKer was buried in the active-site cavity and formed a hydrophobic interaction with S1 binding pocket [11], indicating that the C-terminal residues are related to the conformation of the S1 binding pocket. DgeKer and DgeKer-C resulted in different feather degradation. It is possible that the truncated C-terminal extension was too long in this study, which affected the confirmation of DgeKer-C S1 binding pocket. In short, more experiments are needed to explore the function of C-terminal extension.

5. Conclusions

In conclusion, the DgeKer and its C-terminal extension truncated enzyme DgeKer-C from slightly thermophilic species D. geothermophilis showed excellent stability over the alkaline range of pH and at a temperature of 60 °C, and also displayed excellent tolerance to some metal ions, surfactants and organic solvent, holding promise for industrial applications, such as feather composting. In addition, The C-terminal extension of DgeKer could promote enzyme degradation of insoluble substrates, but has no effect on thermal stability. Indeed, further research is needed to ferment DgeKer in B. subtilis or yeast.

Supplementary Materials: The following are available online at https://www.mdpi.com/article/10.3390/app11073136/s1, Figure S1: Phylogenetic analysis of C-terminal extension of DgeKer and other keratinases, Figure S2: Modeled structure of C-terminal extension of DgeKer using Pymol software.
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