CHARACTERIZATION of THE α-AMYLASE GENE FROM *Bacillus* sp. BBM1

CARACTERIZACIÓN DEL GEN DE α-AMILASADE LA CEPA NATIVA Bacillussp. BBM1

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ABSTRACT

Starch degrading enzymes represent about 30% of the enzyme world and they are used in the production of glucose, maltose and oligosaccharides, which can be further processed to produce fructose and dextrose syrups. The resulting glucose can also be fermented for the production of ethanol, amino acids and organic acids. α -amylases are also used as an alternative to the addition of malt in the beer industry, the improvement of flour in the baking industry, the removal of starch in the paper and textile industry, and as a detergent additive. In this paper, the complete nucleotide sequence of the α -amylases BBM1 produced by the native strain *Bacillus* sp. BBM1 is reported. The deduced aminoacid sequence shows that this enzyme is translated as a 659 a.a. protein, which after the secretion cleaves to generate a 618 mature enzyme of 68 kDa. The BBM1 α -amylases is transcribed as a monocistronic mRNA, as it is suggested by the presence of inverted repeat elements upstream and downstream of the protein coding region. The expression of the BBM1 α -amylase is under the control of the *AmyR1* allele from the *AmyO* operator sequence, which is recognized by the regulatory protein CcpA. In contrast to most α -amylases, BBM1 seems to possess two additional carbohydrate-binding domains, which probably increase its efficiency in the degradation of granular starch. A homology model of the enzyme is presented and its interaction with calcium ions and substrate is analyzed.

Keywords: Starch, polysaccharides, glucan 1,4-alpha-Glucosidases, enzymology.

RESUMEN

Las enzimas degradadoras del almidón representan cerca del 30% del mercado mundial de enzimas y son utilizadas en la producción de glucosa, maltosa y oligosacáridos; los cuales pueden ser transformados posteriormente en jarabes de fructosa y dextrosa. La glucosa también puede ser utilizada en la producción de etanol, aminoácidos y ácidos orgánicos. La α -amilasa también puede ser utilizada como una alternativa a la adición de malta en la industria de la cerveza, el mejoramiento de harinas y la remoción de almidón en la industria papelera y textil y como aditivo de detergentes. En este trabajo reportamos la secuenciación completa del gen codificante para la α -amilasa BBM1 producida por la cepa nativa *Bacillus* sp. BBM1, incluyendo sus regiones reguladoras 3' y 5'. La secuencia de aminoácidos corresponde a una proteína de 659 residuos que, luego de ser secretada y procesada post-traduccionalmente, da origen a una enzima madura

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de 618 a.a con un peso de 68 kDa. La amilasa BBM1 es transcrita como un mRNA monocistrónico, tal como lo sugiere la presencia de estructuras terminadoras de la transcripción. Su expresión está regulada por el factor CcpA cuya secuencia operadora corresponde al alelo *AmyR1*. A diferencia de la mayoría de las amilasas estudiadas, BBM1 parece poseer dos dominios adicionales de unión a carbohidratos, lo cual indica que esta enzima puede ser más eficiente en la degradación de almidón granular. Finalmente, se presenta un modelo por homología para esta enzima que indica las posibles interacciones con iones de calcio y el sustrato.

Palabras clave: almidón, polisacáridos, glucano 1,4-alfa-Glucosidasa, enzimología

INTRODUCTION

 α -amylases (EC 3.2.1.1) are enzymes that catalyze the hydrolysis of the a-1,4 glycosidic bonds that are present in starch, glycogen and other polysaccharides. Starch is composed by two different glucose polymers: amylose, linked by a-1,4 bonds; and amylopectin that, in addition to a-1,4 bonds, has a-1,6 branch sites (1). Starch degrading enzymes represent about 30% of the enzyme world in commercial products such as Ultra-thin® (Valley Research/Diversa), Multifect AA 21L[®] (Genencor), Termamyl[®] and Liquozyme[®] (Novozymes) (2). The starch industry is the main user of α -amylases for the production of glucose, maltose and oligosaccharides, which can be further processed to produce fructose and dextrose syrups. The resulting glucose can also be fermented for the production of ethanol, amino acids and organic acids (3). α -amylases are also used as an alternative to the addition of malt in the beer industry, the improvement of flour in the baking industry, the removal of starch in the paper and textile industry, and as a detergent additive (4). Previously, we reported the purification and biochemical characterization of an extracellularaamylase produced by Bacillus sp. BBM1, a native strain that has been isolated by our research group (5). This enzyme has an optimal temperature of 60°C, but it can be used with 80% of efficiency at temperatures up to 72°C. In this paper, the analysis of the complete nucleotide sequence of α -amylase BBM1, including the 5' and 3' intergenic regions, is reported. A homology model was constructed and used to understand the interactions of this enzyme with the ions and the substrate.

MATERIALS AND METHODS

DNA extraction and PCR

Bacillus sp. BBM1 was isolated from a clay soil sample at Universidad Nacional de Colombia

sede Medellín with the following characteristics: pH 5.4, 36 g/kg organic matter, 7 mg/kg NO₃, 16 mg/kg NH4⁺, and 20 mg/kg phosphate. Bacillus sp. BBM1 grows optimally at 30°C and pH 7.0 in M9 minimal media and LB. A neighbor-joining analysis revealed a phylogenetic affinity of Bacillus sp. BBM1 with B. methylotrophicus, a recently characterized species isolated from the rhizosphere (5). Bacillus sp. BBM1 was grown in LB medium at 30°C and DNA was purified with the DNeasy Blood & Tissue kit (QIAGEN), following the manufacturer's protocol. A total of six primers were used for the amplification of the AmyE gene, as it is shown in figure 1 and table 1. Primers ycgb-F, ldh-R and ldh2-R annealed to flanking genes ycgB and ldh. Primers AmyE1I-F, AmyE2I-R and amy3-F annealed to the internal regions of AmyE. The 5' and 3' regions of AmyE were amplified with primers ycgB-F/AmyE2I-R and AmyE1I-F/ ldh-R, respectively. Sequence gaps were completed with the primer combination amy3-F/ldh2-R. Each PCR reaction consisted of a total volume of 50 mL containing 0.05 u/ml of Taq DNA polymerase, 1X PCR buffer, 2.5 mM MgCl₂, 0.2 mM of each deoxynucleotidetriphosphate, 0.5 mM for each primer, and 50 ng/ml of template DNA. The PCR conditions were the following: initial denaturation at 94°C for 1 min, 30 cycles of amplification consisting of a 30 second denaturation step at 94°C, a 30 s annealing step, and a 3 min extension step at 72°C. The annealing temperature was set according to the primer melting temperature reported in table 1. DNA amplicons were visualized in 1% agarose gel electrophoresis in TBE buffer. DNA was sequenced by Macrogen Inc. (Seoul, Korea) and assembled with the CAP Contig assembly module from BIOEDIT (6). The complete sequence of the AmyE gene was deposited in GenBank with the accession code GU947099.

Primer	Sequence (5'-3')	Tm (°C)	Amplicon size (pb)
amyE1I-F	TGATGCAGGATATACAGCCATTCAGAC	50.9	2822
ldh-R	CACCGTATTGCCCGTCC	54	
ycgB-F	TGCTTACGATGTACGACAGG	61.2	2231
amyE2I-R	GGCAGCGTCAGCGTGTAAATTCC	63.5	
amy3-F	ACGAACGGCGAGGGTGCAGC	66.1	1072
ldh2-R	GCCGCTGCCGATGACGCG	67.6	1075

Table 1. Primers used for sequencing the α-amylase gene of *Bacillus* sp. BBM1.

Sequence annotation

The coding region for the *AmyE* gene from the *Bacillus* sp. BBM1 was identified by means of the BLASTX software (7). Transcription terminators were predicted with RNAfold (8). The Shine-Dalgarnosequence (SD) was identified through a sequence complementarity to the 3' region of the 16S RNA of *B. amyloliquefaciens* FZB42 (NC_009725.1) and *B. subtillis* subsp.subtilis str. 168 (NC_000964.3), using a Perl routine written by the authors. The promotor region was identified with a custom designed program that used a Position Specific Scoring Matrix (PSSM) derived from the promotor regions of the α -amylase gene from *B.subtilis, B. stearothermophilus, B. licheniformis and B. amyloliquefaciens* (9).

Phylogenetic Analysis

A set of 136 non-redundant α -amylases sequences from fungi and bacteria were downloaded from the SwissProt/TrEMBL database (10). A multiple alignment was performed with CLUSTALX using the Gonnet substitution matrix and penalties of 10 and 0.2 for gap insertion and extension, respectively (11). Phylogenetic trees were built using MEGA4 (12). Sequences in the trees were labeled using their corresponding entry name at SwissProt/TrEMBL.

Homology modeling

The N-terminal domains (A, B and C) were modeled using the crystallographic structure of *Bacillus subtilis* α -amylase as a template (PDBid: 1BAG) (13). The template for the C-terminus was chosen with the iterative BLAST protocol PSI-BLAST (14) that identified the cyclodextrin glucanotransferases from *Bacillus circulans* and *Geobacillus stearothermophilus* as distant homologues of α -amylase BBM1 (15). The homology model was generated with MODELLER (16).

RESULTS AND DISCUSSION

Sequence analysis

Bacillus sp. BBM1 has a phylogenetic affinity with *B. methylotrophicus*, a species closely related to *B. amyloliquefaciens* and *B. subtilis*. In these species, the α -amylase gene is flanked by *ycgB*, a gene of unknown function, and *ldh*, coding for lactate dehydrogenase (17, 18). Based on the complete genome sequence Bacillus amyloliquefaciens FZB42 (NC_009725.1) and Bacillus subtilis subsp.subtilis str. 168 (NC_000964.3), a set of primers for the amplification and sequencing of 2971nt (comprising the *AmyE* gene and the 3' and 5' intergenic regions) were designed (see table 1 and figure 1).



Figure 1. Genetic context of the *AmyE* gene and location of the primers used in this work.

The BLASTX analysis showed that α-amylase BBM1 has 99 and 98% of similarity with its homologues in *Bacillus* spB-5 (gb:ADB81848.1) and *B. amyloliquefaciens* FZB42 (gb:ABS72721.1), respectively. The expressed protein has a secretion signal that spans the first 27 residues (MFEKRFKT-SLLPLFAGFLLLFHLVLSG), and a cleavage site at position 41 (NK/VT) that gives rise to a mature protein of 618 amino acids with a predicted molecular weight of 68051.7 Da, as it is shown in figure 2. According to our predictions, the ribosomal binding site (RBS) corresponds to the AATAAGGAGT sequence located eight nucleotides upstream of the start codon.

VITAE

92	TAAAAAAGAAATCATCTTGACAGGTGGTTTCTTTTTTTGTTTG
182	TCTGTCACGTTGCTTGATAGAGAGAGTGTGTGATAAGTTGAAAGTTGAAAGCGTTAACAGAATTCTCCCAGTCTTCACAACAGTTGAAAGGAGC
272 1	TGAAGAAAGAGTAAGAGGAATTTTTGACTCCGCAGTCAGT
362	ACCTCTTTACTGCCGTTATTCGCCGGATTTTTATTGCTGTTTCATTGGTTTTGTCAGGCCCGGCGGCTGCAAACGCTGAAACTGCAAAC
8	<u>T S L L^VP L P A G F L L L F H L V L S G</u> P A A A N A E T A N
452	AAATCGAATAAGGTGACCGCGTCATCGGTCAAAAACGGGACCATCCTTCATGCATG
38	K S N K V T A S S V K N G T I L H A W N W S F N T L T Q N M
542	AAAGATATTCGTGATGCGGGCTATGCAGCCATTCAGACGTCTCCGATTAACCAAGTAAAGGAAGG
68	K D I R D A G Y A A I Q T S P I N Q V K E G N Q G D K S M R
632	AACTGGTACTGGCTGTATCAGCCGACATCGTACCAAATCGGCAACCGTTACTTAGGCACTGAACAAGAATTTAAGGACATGTGTGCAGCC
98	N W Y W L Y Q P T S Y Q I G N R Y L G T E Q E F K D M C A A
122	GCGGGAAAAATATGGCGTAAAAGTCATTGTCGATGGGGTTATCATCATCATCATCATCATCATGGCGGGATTATGGCGCGGATTATGGCGGGATTATGGCGGGATTATGGCGGATTATGGCGGATTATGGCGGGATTATGGCGGGATTATGGCGGATTGGGGGATTATGGCGGGATTATGGCGGGATTATGGCGGGATTATGGCGGGATTATGGCGGGATTATGGCGGGATTATGGCGGGATTATGGCGGGATTATGGCGGGATTATGGCGGGATTATGGCGGGATTATGGCGGGATTATGGCGGGATGGGGATTATGGCGGGATTATGGCGGGATTATGGCGGGATTATGGCGGGATTATGGCGGGATTATGGCGGGATTATGGCGGGATGGGGATTATGGCGGGATTATGGCGGGATTATGGCGGGATTATGGCGGGGATTATGGCGGGATTATGGCGGGATTATGGCGGGGATTATGGCGGGATTATGGCGGGATGGGAGGAGGAGGAGGAGGAGGAGGAGGAGGAGGAG
912	
158	
902	TGGAATACACAGAATACTGAGGTGCAGGTCTATCTTGAAAGAGGAATGAAGAGCAGAGGAGCGGAGCGGATCCGCCTATGAT
188	W N T O N T E V O V Y L K R F L E R A L N D G A D G F R Y D
992	GCCGCCAAGCATATAGAGCTTCCGGATGATGGGAATTACGGCAGCCAATTTTGGCCGAATATCACAAAATATATCGGCAGAGTTCCAATAC
218	A A K H I E L P D D G N Y G S Q F W P N I T N I S A E F Q Y
1082	GGAGAAATCCTGCAAGACAGCGCGTCCAGAGATACTGCTTATGCGAATTATATGAATGTGACGGCTTCTAACTACGGGCATTCCATCAGA
248	G E I L Q D S A S R D T A Y A N Y M N V T A S N Y G H S I R
1172	TCCGCTTTAAAGAACCGTAATCTGAGTGTGTCGAATATTTCCCATTATGCATCTGATGTGTCTGCGGGACAAGTTGGTCACATGGGTGGAA
278	S A L K N R N L S V S N I S H Y A S D V S A D K L V T W V E
1262	TCACATGATACGTATGCCAATGATGACGAAGAGTCCACATGGATGAGCGATGACGATATTCGTTTAGGATGGGCAGTGATTGGTTCCCGT
308	S H D T Y A N D D E E S T W M S D D D I R L G W A V I G S R
1352	TCAGGAAGTACGCCTCTTTTCCTTTTCCAGACCTGAGGGCGGAAGGAA
338	
7442	
1532	
398	G N N O I F M N O R G S K G V V L A N A G S S S V T I N T S
1622	ACGAAATTACCTGACGGCAGGTATGATAATAGGGCCGGCGCGCGC
428	T K L P D G R Y D N R A G A G S F O V T N G K L T G T I N A
1712	AGATCCGCGGCTGTTCTTTATCCTGACAACATTGGAAATGCGCCTCATGTCTTTCTT
458	R S A A V L Y P D N I G N A P H V F L E N Y Q T G A V H S F
1802	AATGATCAGCTGACGGTCACCCTGCGTGCAAATGCGAAAACAACAACAAAGCCGTTTACCAAATCAATAATGGGCAGCAGCAGCAGCATTTAAG
488	N D Q L T V T L R A N A K T T K A V Y Q I N N G Q Q T A F K
1892	GATGGAGACCGACTAACGATCGGGAAAGGAGATCCAATCGGCACGACATACAACATCAAATTAACCGGAACGAAC
518	D G D R L T I G K G D P I G T T Y N I K L T G T N G E G A A
1982	AGAACCCAAGAATACACGTTTGTCAAAAAAGACCCGTCCCAAACCAACATCATTGGCTATCAAAATCCGGACCAGTGGGGCCCAGGTAAAT
548	
2072	COTTATATCATGATGAGGGGGGGGGGGGGGGGGGGGGGGG
2162	ALL A R D O O R A L D D I O D R P O R A R T R N A D O M
608	Y T L T L P E N A D T A N A K V I F N N G S A O V P G O N O
2252	CCCGGCTTTGATTATGTGCAGAATGGTTTGTATAACAACTCCGGTTTAAACGGTTATCTTCCGCATTAATGGATAGGGCCATCATCACAT
638	PGPDYVQNGLYNNSGLNGYLPH*
2342	CAGCGTGCCGGCAAACCCCCGCTTCTGGGAAAGGGACAGCGCGTCTTCAATCACGCTGAAAAGATGACAACATCCTAAAACGGTTCCGTT
2432	TTAGGATGTTGTCAACAATCCGGAAACCATCACCGATATTTTTT

Figure 2. Complete nucleotide and amino acid sequence of α-amylase BBM1.

In order to prove that the expression of the *AmyE* gene is coupled to the transcription of the YcgA-YcgB operon, an analysis of attenuators in the 5' intergenic region was performed. Secondary structure predictions suggest the presence of a 93 nt sequence downstream of the *YcgB* termination codon, which can form a hairpin structure with a free energy of -14.80 kcal/mol. This fact implies an independent transcription from the YcgA-YcgB operon (shown

in figures 2 and 3). Another termination signal with a free energy of -28.40 kcal/mol was detected 85 nt downstream the termination codon, suggesting that the transcription of *AmyE* is not coupled to *ldh* either. Therefore, the translation of *AmyE* BBM1 is probably caused by a monocistronic mRNA, a fact that can be observed in other α -amylases from the genus *Bacillus* (19-22).



Figure 3. Predicted transcription termination signals flanking the *AmyE* BBM1 gene.

The analysis of the promotor region is important for understanding the regulatory mechanisms involved in the regulation of the protein expression (23-25). In this case, the -35 promotor region has the sequence TTGATA, and it is identical to the one present in the α -amylase hyperproducing strain of B. subtilis (22). The -10 sequence corresponds to TTGAAA, which is slightly different from the consensus TANAAT observed in other α-amylase genes. In *B. subtilis*, the expression of α -amylase is regulated in *cis* by the operator sequence *amyO* (TGT/AAANCGNTNA/TCA), which is recognized by protein factor CcpA that is a repressor of the LacI-GaR family (26). In our case, we found the sequence TGTAAGCGTTAACA, which is identical to the amyR1 allele present in B. subtillis

168 (27). In amylase overproducing strains such as *B. amyloliquefaciens* and *Bacillus natto* IAM 121, the *amyO* sequence has a CG substitution at position 7, which presents a weaker affinity with the repressor (28). This condition suggests that the production of α -amylase by *Bacillus* sp. BBM1 is under a tighter control, and that it is produced at lower levels than the corresponding hyperproducing strains.

Comparison with other microbial α -amylases

Using the UPGMA clustering method, we found that α -amylases can be divided into five groups as it is shown in figure 4:

- Group 1: Mostly fungal α-amylases.
- Group 2: Bacterial α-amylases from the genera *Aliivibrio*, *Streptococcus*, *Bacillus* and the *Enterobacteriacea* family.
- Group 3: Proteobacterial α-amylases from the genera *Vibrio*, *Aeromonas* and *Xanthomonas*.
- Group 4: Bacterial α-amylases from the phyla *Proteobacteria* and *Actinobacteria*.
- Group 5: Bacterial α-amylases from the genera *Clostridium, Streptococcus* and *Bacillus.*

With the exception of *B. megaterium* (AMY_BACME) and *B. circulans* (AMY_BACCI), all α -amylases from the *Bacillus* genus are members of the groups 2 or 5. α -amylase BBM1 belongs to *B.amyloliquefaciens* FZB42, *B. amyloliquefaciens* TB2, and *B. subtilis* X-23. Group 5 α -amylases have an approximate size of 660 residues, in contrast to enzymes from group 2, which have an average size of 520 residues.



Figure 4. UPGMA clustering of microbial α-amylases.



Figure 5. Sequence and structure comparison between *B. subtilis* (BSU), *B. licheniformis* (BLA), and *Bacillus* sp. BBM1 α -amylases.

Most α -amylases comprise three domains: Domain A has a TIM barrel structure and corresponds to the catalytic center. Domain B originates from a protuberance in the region that connects the third helix and the strand from domain A. The structure of domain B is very variable and its function seems to be associated with substrate binding and thermostability. Domain C has a b-sandwich structure, but the information regarding its function is scarce (29). To understand

VITAE

the interaction of the α -amylase BBM1 with the ions and the substrate, a homology model was built. Domains A, B and C were modeled using the structure *B. subtilis* α -amylase as template (PDBid: 1BAG). The selection of templates for the C-terminal extension was more complex because no structures with a significant similarity to BBM1 were found using a regular BLAST search. However, an iterative search using the PSI-BLAST software found a homology between α-amylase BBM1 and the cyclodextrin glycosiltransferases (CGTases) from Bacillus circulans and Geobacillus stearothermophilus (15); this alignment included the C-terminal extension (data not shown). This fact is not surprising as both CGTases and α -amylases belong to the family 13 of glycosyl hydrolases.

A comparison between the BBM1 model and the structure of B. licheniformis amylase (group 2) is shown in figure 5. According to our model, the substrate binding site of α-amylase BBM1 can accommodate at least five glucose units between domains A (Blue) and B (green), as it is shown in figure 5B. α -amylases interact with the substrate through multiple binding sites in the enzyme surface, as it is shown in figure 5A. Black and white circles correspond to polar and hydrophobic interactions respectively. The only difference between the binding sites of α-amylase BBM1 and its homolog in *B. subtilis* is the substitution F103S, which interacts with the first glucose unit (30, 31). Catalytic residues Asp174, Glu206 y Asp267 are completely conserved (shown in figure 5A, with asterisks). Glu206 is responsible for the protonation of the glycosidic oxygen. Asp 174 is involved in the following step and its function consists in performing a nucleophilic attack on C1 on the sugar position at subsite -1 (29). The restitution of the active site is achieved by water activated by deprotonated Glu206. Asp267 is an essential residue involved in raising the pKa of Glu206 (29).

Most α -amylases have calcium-binding sites that are important for the structural stability of the enzyme. *B. subtilis* α -amylase has three calcium ions associated to its structure, while *B. licheniformis* has an extra sodium ion that forms an anionic triad together with two calcium ions (shown in figure 5B in green) (32). The binding of the first calcium ion is due to interactions with Asn99, Thr135, Asp144 and His178 that are located on the interface between domains A and B, as it is shown in figure 5A. The second calcium ion is located at the protein surface, and it involves interactions with conserved residues Gly167 and D169. The third site presents an E87K substitution that is detrimental to calcium binding.

The most interesting feature of our model is that it suggests the presence of two additional starch- binding domains (shown in figure 5B in orange and red). Starch binding domains have a variable size ranging from 30 to 200 residues, and they are present in approximately 10% of the starch degrading enzymes. It has been proposed that starch-binding domains improve the affinity with granular starch, and that they allow the hydrolysis of otherwise inaccessible glycosidic bonds (33, 34). This approximation effect increases the effective concentration of the enzyme and, therefore, the hydrolysis rate (35). These domains are of great interest in biotechnology as they can be used for the production of fusion proteins, and also as a model for the study of sugar-protein interactions (36, 37). Novamyl[®], a maltogenic amylase from B. stearothermophilus, is the only domain-five a-amylase which 3D structure has been solved so far (30). The effect of these extra domains and of the calcium ions on substrate binding and catalysis will be further investigated in a following research work on the recombinant enzyme.

CONCLUSION

 α -amylase BBM1 is an inducible enzyme controlled by the CcpA protein, and it is transcribed as monocistronic mRNA. This protein is secreted as a mature enzyme of 619 residues, and its structure is stabilized by the calcium ions located between domains A and B. The sequence analysis suggests that α -amylase BBM1 might have two additional starch-binding domains in contrast to most conventional amylases.

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REFERENCES

- Van derMaarel M, Van derVeen B, Uitdehaag H, Leemhuis H, Dijkhuizen L. Properties and applications of starch converting enzymes of the α-amylase family. J. Biotechnol. 2002 Mar 28; 94 (2): 137-155.
- 2. Turner P, Mamo G, Karlsson EN. Potential and utilization of thermophiles and thermostable enzymes in biorefining. Microb Cell Fact. 2007 Mar 15; 6: 9.
- Kirk O, Borchert TV, Fuglsang CC. Industrial enzyme applications. Curr Opin Biotechnol. 2002 Aug; 13 (4): 345-351.
- Prakash O, Jaiswal N. alpha-Amylase: an ideal representative of thermostable enzymes. Appl Biochem Biotechnol. 2010 Apr; 160 (8): 2401-2414.
- Quintero M, Montoya OI, Gutiérrez P. Purificación y caracterización de una alfa-amilasa producida por la cepa nativa *Bacillus*sp. BBM1. Dyna. 2010 Jun; 162: 31-38.
- Hall TA. BioEdit: a user-friendly biological sequence alignment editor and analysis program for Windows 95/98/NT. Nucleic Acids Symp Ser. 1999, 41: 95-98.
- Altschul SF, Gish W, Miller W, Myers EW, Lipman DJ. Basic local alignment search tool. J Mol Biol. 1990 Oct 5; 215 (3): 403-410.
- Gruber AR, Lorenz R, Bernhart SH, Neuböck R, Hofacker IL. The Vienna RNA websuite. Nucleic Acids Res. 2008 Jul 1; 36: W70-W74.
- Durbin R, Eddy S, Krogh A, Mitchinson G. Biological sequence analysis: probabilistic models of proteins and nucleic acids. Cambridge: Cambridge UniversityPress; 1998.
- Boeckmann B, Bairoch A, Apweiler R, Blatter MC, Estreicher A, Gasteiger E, *et al*. The SWISS-PROT protein knowledgebase and its supplement TrEMBL in 2003. Nucleic Acids Res. 2003 Jan 1; 31 (1): 365-370.
- Larkin MA, Blackshields G, Brown NP, Chenna R, McGettigan PA, McWilliam H, *et al.* Clustal W and Clustal X version 2.0. Bioinformatics. 2007 Nov 1; 23 (21): 2947-2948.
- Kumar S, Nei M, Dudley J, Tamura K. MEGA: a biologist-centric software for evolutionary analysis of DNA and protein sequences. Brief Bioinform. 2008 Jul; 9 (4): 299-306.
- Fujimoto Z, Takase K, Doui N, Momma M, Matsumoto T, Mizuno H. Crystal structure of a catalytic-site mutant alphaamylase from *Bacillus subtilis*complexed with maltopentaose. J Mol Biol. 1998 Mar 27; 277 (2): 393-407.
- Altschul SF, Madden TL, SchäfferAA, Zhang J, Zhang Z, Miller W, et al. Gapped BLAST and PSI-BLAST: a new generation of protein database search programs. Nucleic Acids Res. 1997 Sep 1; 25 (17): 3389-3402.
- Penninga D, van derVeen BA, Knegtel RM, van Hijum SA, Rozeboom HJ, Kalk KH, *et al.* The raw starch binding domain of cyclodextringlycosyltransferase from *Bacillus circulans* strain 251. J Biol Chem. 1996 Dec 20; 271 (51): 32777-32784.
- Kobe B, Guss M, Huber T (editors). Methods in Molecular Biology, Vol. 426: Structural Proteomics: High-throughput Methods. Totowa, NJ: Humana Press; 2008. Eswar N, Eramian D, Webb B, Shen MY, Sali A. Protein structure modeling with MODELLER; p. 145-159.
- Madhaiyan M, Poonguzhali S, Kwon SW, Sa TM. Bacillus methylotrophicussp. nov, a methanol-utilizing, plant-growthpromotingbacteriumisolatedfrom rice rhizospheresoil. Int J Sys Evol Microbiol. 2010 Oct; 60 (10): 2490-2495.
- Chen XH, Koumoutsi A, Scholz R, Eisenreich A, Schneider K, Heinemeyer I, *et al.* Comparative analysis of the complete genome sequence of the plant growth-promoting bacterium *Bacillus amyloliquefaciens* FZB42. Nat Biotechnol. 2007 Sep; 25 (9): 1007-1014.
- Movva NR, Nakamura K, Inouye M. Regulatory region of the gene for the ompA protein, a major outer membrane protein of

Escherichia coli. Proc Natl Acad Sci USA. 1980 Jul; 77 (7): 3845-3849.

- Takkinen K, Pettersson RF, Kalkkinen N, Palva I, Söderlund H, Kääriäinen L. Amino acid sequence of alpha-amylase from *Bacillus amyloliquefaciens* deduced from the nucleotide sequence of the cloned gene. J Biol Chem. 1983 Jan 25; 258 (2): 1007-1013.
- Nakajima R, Imanaka T, Aiba S. Nucleotide sequence of the Bacillus stearothermophilusalpha-amylase gene. J Bacteriol. 1985 Jul; 163 (1): 401-406.
- 22. Yamazaki H, Ohmura K, Nakayama A, Takeichi Y, Otozai K, Yamasaki M, *et al.* Alpha-amylase genes (amyR2 and amyE+) from an alpha-amylase-hyperproducing*Bacillus subtilis* strain: molecular cloning and nucleotide sequences. J Bacteriol. 1983 Oct; 156 (1): 327-337.
- 23. Fickett JW. The gene identification problem: an overview for developers. Comput Chem. 1996 Mar; 20 (1): 103-118.
- 24. Stephens MA, Ortlepp SA, Ollington JF, McConnell DJ. Nucleotide sequence of the 5' region of the *Bacillus licheniformis* alpha-amylase gene: comparison with the B. amyloliquefaciens gene. J Bacteriol. 1984 Apr; 158 (1): 369-372.
- Sibakov M, Palva I. Isolation and the 5'-end nucleotide sequence of *Bacillus licheniformis* alpha-amylase gene. Eur J Biochem. 1984 Dec 17; 145 (3): 567-572.
- Kim JH, Guvener ZT, Cho JY, Chung KC, Chambliss GH. Specificity of DNA binding activity of the *Bacillus subtilis*catabolite control protein CcpA. J Bacteriol. 1995 Sep; 177 (17): 5129-5134.
- Weickert MJ, Chambliss GH. Genetic analysis of the promoter region of the *Bacillus subtilis*alpha-amylase gene. J Bacteriol. 1989 Jul; 171 (7): 3656-3666
- Gordon JJ, Towsey MW, Hogan JM, Mathews SA, Timms P. Improved prediction of bacterial transcription starts sites. Bioinformatics. 2006 Jan, 22 (2): 142-148.
- Nielsen JE, Borchert TV. Protein engineering of bacterial alphaamylases. Biochim Biophys Acta. 2000 Dec 29; 1543 (2): 253-274.
- Machius M, Declerck N, Huber R, Wiegand G. Activation of Bacillus licheniformis alpha-amylase through a disorder-->order transition of the substrate-binding site mediated by a calciumsodium-calcium metal triad. Structure. 1998 Mar 15; 6 (3): 281-292.
- Fujimoto Z, Takase K, Doui N, Momma M, Matsumoto T, Mizuno H. Crystal structure of a catalytic-site mutant alphaamylase from *Bacillus subtilis*complexed with maltopentaose. J Mol Biol. 1998 Mar 27; 277 (2): 393-407.
- 32. Liu Y, Shen W, Shi GY, Wang ZX. Role of the calcium-binding residues Asp231, Asp233, and Asp438 in alpha-amylase of *Bacillus amyloliquefaciens* as revealed by mutational analysis. CurrMicrobiol. 2010 Mar; 60 (3): 162-166.
- 33. Dauter Z, Dauter M, Brzozowski AM, Christensen S, Borchert TV, Beier L, et al. X-ray structure of Novamyl, the five-domain "maltogenic" alpha-amylase from *Bacillus stearothermophilus*: maltose and acarbose complexes at 1.7A resolution. Biochemistry. 1999 Jun 29; 38 (26): 8385-8392.
- 34. Guillén D, Santiago M, Linares L, Pérez R, Morlon J, Ruiz B, et al. Alpha-amylase starch binding domains: cooperative effects of binding to starch granules of multiple tandemly arranged domains. Appl Environ Microbiol. 2007 Jun; 73 (12): 3833-3837.
- Rodríguez-Sanoja R, Oviedo N, Sánchez S. Microbial starchbinding domain. Curr Opin Microbiol. 2005 Jun; 8 (3): 260-267.
- Boraston AB, Bolam DN, Gilbert HJ, Davies GJ. Carbohydratebinding modules: fine-tuning polysaccharide recognition. Biochem J. 2004 Sep 15; 382 (Pt 3): 769-781.
- Shoseyov O, Shani Z, Levy I. Carbohydrate binding modules: biochemical properties and novel applications. Microbiol Mol Biol Rev. 2006 Jun; 70 (2): 283-295.