A Novel Neoagarotriose-Producing Agarase from the Marine Bacterium Gilvimarinus Agarilyticus JEA5

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Abstract

Background The degradation of agar by bacterial agarases has many commercial and academic applications. We recently identified a novel neoagarotriose-producing β-agarase, Gaa16B, in the marine bacterium *Gilvimarinus agarilyticus* JEA5. This is the first report to describe neoagarotriose production from β-agarase.

Results The Gaa16B agarase, which belongs to the glycoside hydrolase 16 (GH16) family of β-agarases, shows less than 70.9% amino acid similarity with previously characterized agarases. The coding region of Gaa16B is 1800 bp long, encoding 600 amino acids, and exhibits features typical of agarases belonging to the GH16 family. A recombinant Gaa16B lacking the carbohydrate binding region (rGaa16Bc) was overexpressed in *Escherichia coli* and purified as a maltose-binding protein (MBP) fusion protein. Activity assays revealed the optimal temperature and pH of rGaa16Bc to be 55 °C and pH 6–7, respectively, and the protein was highly stable at 55 °C for 90 min. Additionally, rGaa16Bc activity was strongly enhanced (2.3-fold) in the presence of 2.5 mM MnCl₂. The $K_m$ and $V_{max}$ of rGaa16Bc for agarose were 6.4 mg/ml and 953 U/mg, respectively. Thin layer chromatography analysis revealed that rGaa16Bc can hydrolyze agarose into neoagarotetraose, neoagarotriose, and neoagarobiose, and the production of neoagarotriose by rGaa16Bc was successfully validated by high-resolution electrospray ionization mass spectrometry.

Conclusion The biochemical properties of Gaa16B and the results of the hydrolytic pattern analysis suggest that Gaa16B could be useful to produce functional neoagaro-oligosaccharides for industrial applications.

Background

Agar is the main component of the cell wall in several species of red seaweed and consists of a mixture of two polysaccharides: agarose and agarpectin. Agarose consists of a linear chain of alternating 3-O-linked β-D-galactopyranose and 4-O-linked 3,6-anhydro-α-L-galactose units [1]. Agar is widely used as a gelling substance in the pharmaceutical, cosmetic, and food industries, as well as in the life sciences, where it is used in a range of techniques including microbiome culturing media, electrophoresis gel, and chromatography resin. In addition to these classical applications, recent studies have reported that many agar-derived oligosaccharides exhibit various biological and therapeutic properties, and hence have the potential for diverse applications in the cosmetic, medicinal, and pharmaceutical industries [2, 3].

Agarases hydrolyze both agar and agarose, and based on the mechanism of their actions, they are categorized into two main groups, namely α-agarases and β-agarases. The α-agarases hydrolyze α-1,3-linkages between the composite sugars in agarose to generate agaro-oligosaccharides, while β-agarases hydrolyze only β-1,4-linkages in agarose to generate neoagaro-oligosaccharides. The β-agarases are classified into four families on the basis of their amino acid sequence homology: glycoside hydrolase 16 (GH16), GH50, GH86, and GH118 [4]. To date, many β-agarases have been found from many different genera including *Agarivorans* [5], *Alteromonas* [6], *Bacillus* [7], *Catenovulum* [8], *Pseudoalteromonas* [9], *Pseudomonas* [10], *Vibrio* [11], *Streptomyces* [12], *Saccharophagus* [13] and *Gilvimarinus* [14].

Of the many agarases that have been reported, most of them are β-agarases, except for some α-agarases isolated from *Alteromonas agarilyticus* GJ1B [15], *Thalassomonas agarivorans* JAMB-A33, *Catenovulum agarivorans* [16] and *Catenovulum sediminis* WS1-A [17]. Several reports have described that β-agarases hydrolyze agarose to produce various neoagaro-oligosaccharides such as neoagarobiose, neoagarotetraose, and neoagarohexaose [1]. However, to our knowledge, an agarase that produces neoagarotriose has not yet been reported. In this study, we identified that the gene *gaa16b*, isolated from the marine bacterium *Gilvimarinus agarilyticus* JEA5, encodes a novel neoagarotriose-producing β-agarase. We cloned the *gaa16b* gene and overexpressed it as a recombinant protein in *E. coli* BL21 (DE3). Maltose-binding protein (MBP)-tagged Gaa16B was then purified and analyzed to investigate its biochemical properties and hydrolytic patterns.
Molecular characteristics

We have previously published a draft genome of the marine bacterium *Gilvamarinus agarilyticus* JEA5, which exhibits agarolytic activity [18], and identified Gaa16B as a predicted agarase. The nucleotide sequence of *gaa16b* has an open reading frame (ORF) of 1800 bp, encoding a putative agarase of 600 amino acid residues. The molecular mass and isoelectric point (pl) of Gaa16B were predicted to be 65 kDa and 4.2, respectively. An analysis of the amino acid sequence predicted a signal peptide in the N-terminal region, a GH16 domain in the middle of the protein, and two carbohydrate binding module 6 (CBM6) motifs in the C-terminus. The deduced amino acid sequence of Gaa16B showed the highest identity (90.7%) and similarity (93.8%) with *Gilvamarinus polysaccharolyticus* (WP_049721028.1), whereas it exhibited less than 55.9% identity and 70.9% similarity with functional characterized agarases (Table 1).

| Gaa16B Protein                                      | Identity (%) | Similarity (%) | Gap (%) | Accession No.   | Remark           |
|-----------------------------------------------------|--------------|----------------|---------|-----------------|------------------|
| *Gilvamarinus polysaccharolyticus*                  | 90.7         | 93.8           | 2.7     | WP_049721028.1  | Uncharacterized  |
| *Gilvamarinus chinensis*                            | 89.7         | 93.5           | 2.2     | WP_020208740.1  | Uncharacterized  |
| *Microbulbifer thermotolerans*                       | 75.1         | 85.8           | 5.4     | WP_067155675.1  | Uncharacterized  |
| *Saccharophagus degradans*                           | 55.9         | 70.9           | 4.2     | ABD80437.1      | Characterized    |
| *Pseudomonas sp. ND137*                              | 43.5         | 55.5           | 29.0    | BAD88713.1      | Characterized    |
| *Microbulbifer elongatus*                            | 37.8         | 50.5           | 28.8    | BAC99022.1      | Characterized    |

Overexpression of recombinant agarase Gaa16B

To characterize the Gaa16B protein, the catalytic region (the GH16 domain) of the *gaa16b* gene was amplified by PCR and cloned into the pMal-c2x expression vector. The recombinant Gaa16B catalytic domain (rGaa16Bc) fused to an MBP tag was expressed in *E. coli* BL21 (DE3) and purified using the pMal fusion protein purification system. The purified rGaa16Bc protein was electrophoresed on an SDS-PAGE gel and after staining, exhibited a strong single band with an approximate molecular mass of 75 kDa, which corresponded with the predicted molecular mass (33 kDa for rGaa16Bc plus 42 kDa for the MBP tag) (Fig. 1.).

The optimum conditions for rGaa16Bc activity

We next investigated the effect of temperature, pH, thermostability, and metal ions on rGaa16Bc activity. The recombinant protein was exposed to temperatures ranging from 45 °C to 70 °C, as shown in Fig. 2A. rGaa16Bc
activity gradually increased from 45 °C to 55 °C, exhibiting its highest agarolytic activity at 55 °C. Its activity dramatically decreased at temperatures over 60 °C and it lost almost all activity at 70 °C. The effect of pH on rGaa16Bc activity was tested using pH values ranging from 4.0 to 10.0, in three different buffers, as shown in Fig. 2B. rGaa16Bc showed over than 80% activity in the pH range 5–8, with its highest activity observed at pH 6. No activity was detected at pH 4 or pH 10. Although the maximum activity of rGaa16Bc was observed at 55°, the protein showed low thermostability at this temperature (Fig. 2C). It showed less than 40% activity at 55 °C after 30 min and it lost almost activity after 60 min. However, rGaa16Bc activity was stable at 45 °C for up to 90 min and maintained approximately 80% of its activity until 120 min. The rGaa16Bc protein was also very stable at 50 °C for 60 min and maintained approximately 70% of its activity until 90 min. To determine the effect of metal ions and chelators, the enzyme assay was carried out in the presence of 2.5 mM KCl, CaCl$_2$, MnCl$_2$, NaCl, MgCl$_2$, CuSO$_4$, ZnSO$_4$, FeSO$_4$, or EDTA (Fig. 2D). The relative activity of rGaa16Bc was inhibited by CuSO$_4$ and ZnSO$_4$, but was enhanced in the presence of 2.5 mM MnCl$_2$, CaCl$_2$, MgCl$_2$, and FeSO$_4$. In particular, MnCl$_2$ more than doubled the activity of rGaa16Bc.

The enzymatic reactions for calculating $K_m$ and $V_{max}$ values were performed under optimal conditions and the results showed that the $K_m$, $V_{max}$ and $K_{cat}$ of rGaa16Bc for agarose were 6.4 mg/mL, 953 U/mg, and 201.2 s$^{-1}$, respectively.

**Hydrolytic pattern of rGaa16B**

To determine the hydrolysis pattern and products of rGaa16Bc on agarose, thin layer chromatography (TLC) was used to investigate the hydrolysates at different reaction time points. The TLC results (Fig. 3) show that rGaa16Bc hydrolyzes agarose, generating various neoagaro-oligosaccharides during the initial stages of the reaction. With increasing incubation times, the ratio of the larger oligosaccharides gradually decreased while neoagarotetraose increased (Fig. 3A). After an overnight incubation, neoagarotetraose was observed to be the main product, although neoagarobiose and another spot, which we hypothesized was neoagarotriose, were also observed on the TLC plate (Fig. 3B). After hydrolyzing both neoagarotetraose and neoagarohexaose with rGaa16Bc, the predicted neoagarotriose was observed on the TLC plate (Fig. 3C).

A high-performance liquid chromatography (HPLC) with evaporative light scattering detector (ELSD) mass spectrometry (MS) (HPLC-ELSD-MS) analysis confirmed that rGaa16Bc hydrolyzes agarose into neoagarotetraose (m/z 629.1 [M·H]$^{-}$), neoagarotriose (m/z 469.2 [M + H]$^{+}$), and neoagarobiose (m/z 347.0 [M + Na]$^{+}$) (Fig. 4A). The presence of neoagarotriose was further confirmed using high-resolution ESI mass spectrometry, which identified a peak with the molecular formula C$_{18}$H$_{28}$O$_{14}$ (m/z 467.1397 [M·H]$^{-}$ (calculated for C$_{18}$H$_{27}$O$_{14}$ 467.1401)) (Fig. 4B).

**Discussion**

We have previously reported a draft genome sequence for the agar degrading bacterium *G. agarilyticus* JEA5 [18], and were the first to describe the molecular characteristics and biochemical properties of a β-agarase (Gaa16A) isolated from *G. agarilyticus* JEA5 [19]. Here, we describe Gaa16B, a novel neoagarotriose-producing β-agarase. This newly identified agarase possesses the typical functional domains characteristic of β-agarases belonging to the GH16 family. Almost all GH16 β-agarases contain a GH16 domain at the N-terminus and carbohydrate binding modules in the C-terminus of the protein [20]. Gaa16B exhibits both the GH16 domain and two carbohydrate VI modules at the N- and C-termini of the protein, respectively.

The Gaa16B amino acid sequence showed the highest identity and similarity with a carbohydrate binding protein from *G. polysacchariticus* (WP_049721028.1), which has not yet been characterized. The NCBI database contains three genomes from the *Gilvimarinus* genus; however, only one agarase (Gaa16A from *G. agarilyticus* JEA5) has been characterized thus far [19]. Compared to the amino acid sequences of characterized agarases, Gaa16B displayed the highest similarity to an agarase from *Saccharophilus degradans* 2–40 (with a sequence identity and similarity of only 55.9% and 70.9%, respectively).
To determine the optimum reaction conditions for this enzyme, we investigated the effects of temperature, pH, thermostability, and metal ions on rGaa16Bc function. It showed the highest agarolytic activity at 55 °C and pH 6, although it also exhibited relatively high agarolytic activity (80% or more) between pH 5–8. This high activity in a relatively wide pH range may be an advantage for the industrial use of rGaa16Bc. Furthermore, since agar hardens at temperatures below 40 °C, it is important to have high activity above 40 °C. rGaa16B showed optimum activities at these higher temperatures, as well as under neutral ionic conditions that do not require neutralization. These properties could be particularly advantageous for industrial use.

The kinetic characterization assays revealed that the \( K_m \) value of rGaa16Bc (6.4 mg/mL) is only slightly higher than that of other reported GH16 \( \beta \)-agarases, which is probably due to the absence of the CBMs. To assess the agarolytic activity of Gaa16B, the \textit{gaa16b} gene was amplified without the carbohydrate binding region and cloned into a pMal-c2x vector. Most of the expressed protein was insoluble when the full length recombinant Gaa16B (including the carbohydrate binding region) was cloned and expressed in \textit{E. coli}, and the soluble fraction also showed very low agarolytic activity (data not shown). In contrast, the recombinant rGaa16B without the CBMs showed very high activity compared to full length Gaa16B. Other studies have also reported kinetic characterizations of GH16 \( \beta \)-agarases lacking the CBMs and reported similar findings. A recombinant agarase containing only the GH16 catalytic region from \textit{M. thermostolerans} JAMB-A94 exhibited a higher \( K_m \) value than the full length fusion protein [21]. In addition, Wang et al. reported that a recombinant Aga0917 lacking a CBM from \textit{Pseudoalteromonas fuligina} YT-15-1 showed a remarkably high (39.6 mg/ml) \( K_m \) value compare to other known \( \beta \)-agarases [22].

The TLC results showed that agarose was rapidly fragmented by rGaa16Bc. In the early stages of the reaction, rGaa16Bc hydrolyzed the agarose to generate neoagarobiose, neoagarotetraose, neoagarohexaose, and various larger oligosaccharides. The amount of neoagarosaccharides smaller than neoagarohexaose increased in a time-dependent manner. This hydrolytic pattern suggested that rGaa16Bc functions as an endo-type \( \beta \)-agarase. It has been reported that endo-type agarases randomly degrade agarose and gradually lower the viscosity of agarose solution, while exo-type agarases tend to produce single major products and gradually decrease the viscosity of agarose solutions [23].

The rGaa16Bc enzyme produced mainly neoagarotetraose and neoagarobiose. Interestingly, the predicted neoagarotriose spot was observed to lie between those of neoagarobiose and neoagarotetraose, but it was not in line with the spot of the agarotriose which was used as a standard. To verify that rGaa16Bc does indeed produce neoagarotriose, the molecular mass of the hydrolytic products was measured by LC/MS. Electrospray ionization mass spectrometry of the reaction product showed peaks at 347.0 m/z [neoagarobiose + Na]+, 469.2 m/z [neoagarotriose + H]+ and 629.1 m/z [neoagarotetraose - H]-. The molecular weight of agarotriose (C\textsubscript{18}H\textsubscript{30}O\textsubscript{15}) and neoagarotriose (C\textsubscript{18}H\textsubscript{28}O\textsubscript{14}) are 486.4 and 468.4 g/mol, respectively. The sequence of agarotriose is G-A-G (G, D-galactose; A, 3-6-anhydro-a-L-galactose), whereas that of neoagarotetraose is A-G-A. This is the first report to describe a neoagarotriose-producing agarase. Some \( \alpha \)-agarases have been reported to produce agarotriose, but to our knowledge, there are no previous studies that have noted an agarase producing neoagarotriose. The \( \alpha \)-agarase AgaD from \textit{Thalassomonas} sp. LD5 was shown to hydrolyze agarotetraose and generated agarotriose with a molecular weight of 486 g/mol and the G-A-G structural arrangement [24]. In the current study, rGaa16Bc produced neoagarotriose after hydrolyzing agarose, neoagarohexaose, and neoagarotetraose. Furthermore, it generated neoagarotriose with a molecular weight of 468 g/mol and an A-G-A structure.

### Conclusions

In conclusion, a recombinant \( \beta \)-agarase (rGaa16B) from \textit{Gilvamarinus agarilyticus} JEA5 was cloned, overexpressed, and purified, and its biochemical properties were analyzed. This is the first report of an agarase that produces neoagarotriose. Gaa16B has a GH16 family domain as well as a carbohydrate binding domain, both of which are typical features of GH16 \( \beta \)-agarases. The Gaa16B catalytic domain showed high activity at elevated temperatures and within a relatively wide pH range. It also has good thermostability. These
characteristics suggest that Gaa16B is a good candidate for industrial applications, such as in the cosmetic, pharmaceutical, and food industries. Further research on this agarase would require the determination of its crystal structure and a three-dimensional structural analysis, in order to understand the details of the catalytic mechanism by which Gaa16B produces neoagarotriose.

Methods

Molecular characterization of Gaa16B

In our previous study, we isolated the marine bacterium *Gilvamarinus agarilyticus* JEA5, which exhibited agarolytic activity, and we sequenced its genome using next-generation sequencing (NGS) [18]. The predicted agarase was identified by searching the draft genome sequence of *G. agarilyticus* JEA5 (NCBI accession No. WP_041522726) using the Basic Local Algorithm Search Tool (BLAST), and was then named Gaa16B and submitted to NCBI GenBank (NCBI accession No. KP716980). Signal peptide analysis was performed using the SignalP 4.1 server [25] and conserved domains were predicted by SMART [26] and ScanProsite [27]. The identities and similarities of the Gaa16B amino acid sequence were investigated using the EMBL Pairwise Sequence Alignment Tool [28].

Cloning of the gaa16b agarase gene

We isolated genomic DNA from *G. agarilyticus* JEA5 (KCCM43129) using an AccuPrep® Genomic DNA Extraction Kit (Bioneer, Daejeon, Korea Rep) following the manufacturer’s protocol. Primer pairs were designed to amplify only the catalytic region (GH16 domain) of Gaa16B and the resulting protein fragment was designated as Gaa16Bc. Gaa16B-F (5´- TTC AGA ATT CGG ATC GCC GAC TGG GAC GGC TTA - 3´) and Gaa16B-R (5´- TTG CCT GCA GGT CGA GCT CTA GGT AAT GTC GGT ATT GT-3’) primers were designed so that the 5’ region shares 15 bp with the both ends of the *BamH*I and *Sal*I digested vector sequence. The PCR mixture consisted of 1 µl of genomic DNA template (200 ng/µl), 35.5 µl sterile deionized water, 5 µl 10X Ex Taq buffer, forward and reverse primers (20 pmol each), 4 µl dNTPs (2.5 mM), and Ex Taq DNA polymerase (3 U). PCR amplification conditions were as follows: initial denaturation at 94 °C for 5 min; 30 cycles of denaturation at 94 °C for 30 s, annealing at 58 °C for 30 s, extension at 72 °C for 1 min 30 s; and a final extension at 72 °C for 5 min. PCR reactions were carried out using a TaKaRa PCR Thermal Cycler Dice® Gradient (Takara Bio Inc, Japan). The PCR products were purified using an AccuPrep® Gel Purification Kit (Bioneer, Daejeon, Korea Rep), after which they were cloned into a *BamH*I and *Sal*I digested pMal-c2x expression vector (New England Biolabs, UK) using a Ez-Fusion™ Cloning Kit (Enzymomics, Korea Rep) following the manufacturer’s protocol. The recombinant plasmids were transformed into *E. coli* DH5α using standard protocols. The clones were purified using an AccuPrep® Nano-Plus Plasmid Mini Extraction Kit (Bioneer, Daejeon, Korea Rep) and then again transformed into *E. coli* BL21 (DE3) cells for protein expression.

Overexpression and purification of recombinant Gaa16Bcr

The *E. coli* cells carrying Gaa16Bc-pMal-c2x were cultured overnight at 37 °C in 5 mL of Luria broth (LB) containing 100 mg/mL of ampicillin. From the overnight culture, 3 mL was re-inoculated into 250 mL of fresh media and incubated until mid-logarithmic phase (OD$_{600nm}$ = 0.6-0.7). To overexpress recombinant proteins, isopropyl-β-d-thiogalactopyranoside (IPTG) was added to a final concentration of 0.05 mM and the cultures were incubated at 20 °C for 20 hours. The cells were harvested by centrifugation at 8000 × g for 15 min. The collected cells were re-suspended in a column buffer (200 mM NaCl, 20 mM Tris-HCl) and frozen at -20 °C overnight.

The frozen cells were thawed on ice and then disrupted by sonication. The supernatants were separated by centrifugation at 11,000 rpm for 20 min at 4 °C. The recombinant Gaa16Bc (rGaa16Bc) was purified from the soluble fraction using the pMal™ Protein Fusion & Purification System (New England Biolab, UK) according to the manufacturer's instructions and analyzed by SDS-PAGE. Gels were stained with Coomassie brilliant blue to visualize proteins. The concentrations of the purified recombinant proteins were determined using a BCA Protein Assay Reagent Kit (Thermo Fisher Scientific Inc, USA).
Enzyme activity assays

To determine the biochemical properties of rGaa16Bc, we performed enzyme assays to assess the optimum temperature and pH for enzyme activity, as well as protein thermostability, and the effects of metal ions and chelators. The enzyme activities were measured using modified DNS methods [29]. The reaction conditions were as follows: 100 µL of 1% agarose (Lonza, Switzerland), 95 µL of proper buffer and 5 µL of diluted enzyme, incubated for 5 min at 55 °C. The optimum temperature assay was carried out at 45 to 70 °C with 5 °C intervals. To assess the optimum pH, enzyme activity was measured from pH 4.0 to pH 10.0 with pH 1.0 intervals. To analyze thermostability, rGaa16Bc was pre-incubated at 45, 50, or 55 °C for 30, 60, 90 and 120 min. and enzyme activity was measured every 30 min until 120 min. The relative activity was calculated in comparison to the maximum agarolytic activity, which was set at 100%. To determine the kinetic parameters of the recombinant protein, the agarolytic activities of rGaa16Bc were measured using various concentration of agarose ranging from 0.25 to 10 mg/mL under optimal condition. The \( K_m \), \( V_{max} \), and \( K_{cat} \) values were calculated using GraphPad Prism, version 8.3.1 (GraphPad Software, Inc, USA).

Analysis of hydrolytic action and substrate specificity

TLC was used to analyze the hydrolytic pattern of rGaa16Bc. D-galactose (Sigma-Aldrich, USA), neoagarobiose, agarotriose, neoagarotetraose, and neoagarohexaose (all from Carbosynth, UK) were used as standards. Agarose, neoagarohexaose, neoagarotetraose, and neoagarobiose were hydrolyzed by rGaa16Bc under standard conditions. The hydrolytic products were applied to a silica gel 60 TLC plate (Merck, Germany), which was developed using a solvent system of n-butanol: acetic acid: dH2O (2:1:1 [v/v]). Spots were visualized by spraying orcinol dip reagent (80 mg of orcie monohydrate was dissolved in 160 mL of acetone, to which 8 mL of sulfuric acid was added), followed by heating at 110 °C in a drying oven for 10 min.

Product detection by HPLC-ELSD-MS

The agarose degradation products obtained from the treatment with rGaa16Bc were analyzed by HPLC-ELSD-MS using a C18 reversed phase column, YMC-Pack Pro C18 (250 mm x 4.6 mm, 32 µm particle size) from YMC (Kyoto, Japan). The HPLC-ELSD-MS system comprised an Agilent 1260 Infinity module coupled to both an evaporative light scattering detector 1260 Infinity and a single quadrupole mass spectrometer 6120B (Agilent, Santa Clara, CA, USA) equipped with an electrospray ionization source. For the mobile phase, a linear solvent gradient of 100% water / 0% acetonitrile to 100% acetonitrile was used for 5 min followed by the use of 100% acetonitrile for an additional 5 min. The flow rate of the mobile phase was a constant 0.5 mL/min, and the injection volume was 20 µL. The signals corresponding to neoagarobiose and neoagarotetraose in the HPLC-ELSD chromatogram were identified by comparing the retention time and pseudomolecular ions with those of the analytical standards. The presence of neoagarotriose in the chromatogram was confirmed by the molecular formula obtained using high-resolution ESI mass spectrometry using a SYNAPT-G2 system.

Declarations

Ethics approval and consent to participate

Not applicable

Consent for publication

Not applicable

Availability of data and materials

All data generated or analyzed during this study are included in this published article.

Competing interests
The authors declare that they have no competing interests.

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**Authors’ contributions**

YL, EJ, YJL and HC performed the majority of the laboratory works. TYE and YG contributed to the interpretation of the results. YHK, SDM and SAH also performed the laboratory experiments. DHK helped to revise the manuscript. YL wrote the manuscript and overall work was supervised by CO. All authors read and approved the final manuscript.

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30. Declarations.
SDS-PAGE of the rGaa16Bc protein. Samples of rGaa16Bc were separated on a 12% SDS-PAGE gel and stained with Coomassie brilliant blue. M: Molecular mass marker (Thermo scientific, USA). Lane 1: whole cell lysates from E. coli BL21 (DE3) before induction; lane 2: whole cell lysates after IPTG induction; lane 3: total soluble cellular extract after induction; lane 4: total insoluble cellular extract after induction; lane 5: purified rGaa16Bc.
Figure 2
Characterization of biochemical properties of purified rGaa16Bc. The effect of temperature (A), pH (B), thermostability (C) and metal ions and chelators (D) on the rGaa16Bc activity.
Thin layer chromatography of hydrolysis products of rGaa16Bc on agarose, neoagarotetraose (NA4), and neoagarohexaose (NA6). D-galactose (G), neoagarobiose (NA2), NA4, NA6 and agarotriose (A3) are used as standard. The assays were conducted under standard conditions. Lane 1-5, agarose hydrolyzed by rGaa16Bc for 10, 30, 60, 120 and 180 min; lane 6, agarose hydrolyzed by rGaa16Bc overnight; lane 7-8, neoagarotetraose and neoagarohexaose hydrolyzed by rGaa16Bc for 1 hour.
Identification of agarose degradation products. HPLC-ELSD chromatogram of the agarose degradation products (A) and positive-ion ESI-MS spectrum of b (B). a, neoagarobiose ([M+Na]+ 347.0); b, neoagarotriose ([M+H]+ 469.2); c, neoagarotetraose ([M-H]- 629.1).