Peptide oligomers from ultra-short peptides using sortase

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ABSTRACT

Sortase A catalyzed ligation of ultra-short peptides leads to inter/intra-molecular transpeptidation to form either linear or cyclic oligomers dependent upon the peptide length. Cyclic peptides were the main products for peptides with more than 15aa. However, for ultra-short (< 15aa) peptides, cyclic oligomers became predominant in prolonged reactions. Peptides with 1-3 aminoglycines were equally active but peptide oligomers from peptide containing more than one aminoglycine were prone to hydrolysis.

1. Introduction

Bioactive peptides have many unique and unbeatable features in comparison with proteins and other synthesized polymers. Bioactive peptides regulate many physiological processes, acting at some sites as endocrine or paracrine signals and at others as neurotransmitters or growth factors [1], and show useful properties for human health, including antimicrobial, antifungal, antiviral, and antitumor activities [2,3]. In addition, peptides can interact specifically with toxic metal ions to act as effective chelating agents [4,5], and peptides with specific sequences can self-assemble to form materials with regular nanostructures [6-8]. Although most peptides can be produced using the molecular biology approach, they are produced in limited quantities at high costs and associated with some difficulties in purification and isolation. In addition, some peptides with special structures, such as cyclic peptides, can hardly be produced by the biological method. This is especially true for the biomedical applications in which long peptides with functional sequences are usually needed [8].

Staphylococcus aureus Sortase A (SrtA_{Staph}), a cysteine transpeptidase, recognizes an LPXTG motif near C-terminus of a protein substrate and cleaves the peptide bond between threonine and glycine residues to form a thioacyl enzyme intermediate between the catalytic cysteine and the substrate threonine [9,10]. The intermediate reacts with the N-terminal of an oligoglycine in nucleophilic substitution with formation of amide bond between the substrate threonine and the incoming glycine [11]. This SrtA_{Staph}-catalyzed transpeptidation reaction has been used for protein/peptide ligation and labeling [12,13-17,26,27]. Recently, Wu et al. found that Sortase A mediated transpeptidation seemed to be length dependent for peptides with 16–19 amino acids [25]: a cyclic dimer was the major product for peptides with 16 and 17 amino acids while the main product of 19aa peptide was a cyclic monomer. However, SrtA_{Staph}-mediated peptide ligation and thus peptide oligomer production using ultra-short (< 15 amino acids) peptides has not been studied.

2. Experimental details

2.1. Peptides

All peptides for this study (> 90% in purity) were synthesized by Genescript Corp. Methyl ester modified peptides (peptide-OMe) were produced according to method reported previously [18]. The replacement of the glycine in the minimal SrtA_{Staph} recognition sequence, LPRTG, by the threonine methyl ester (-OMe) was shown to minimize product hydrolysis [18].

2.2. Sortase a expression and purification

Wild type Staphylococcus aureus Sortase A containing residues 60–206 (St-SrtA_{Staph}), named as SrtA_{Staph}, was produced in Escherichia coli (E. coli). Plasmid pET15bSt-SrtA_{Staph} (a gift from Dr. Robert Clubb, UCLA) was used for expression. pET15bSt-SrtA_{Staph} was transformed into BL21-Gold(DE3) competent cells (Stratagene, CA). Cultures were grown in 1.0 L of Luria-Bertani (LB) media supplemented with ampicillin (200 μg/mL) at 37 °C with shaking at 250 rpm. After cells were collected by centrifugation at 4,000 rpm, cell pellets were resuspended in the buffer B (20 mM Tris-HCL, pH 7.5, 150 mM NaCl, 15 mM 2-mercaptoethanol)
and lysed by sonication. SrtA<sub>Staph</sub> protein bearing N-terminal His tag was purified by affinity chromatography using a His-Select Nickel Affinity gel (Sigma). Purified protein was eluted with buffer B containing 200 mM imidazole and then concentrated using Amicon Ultra Centrifugal filters (Millipore) with MWCO of 10 kDa. SrtA<sub>Staph</sub> with good reactivity was obtained.

2.3. Sortase FRET activity assay

Sortase A substrate Dabcyl-LPQGT-Edans (Anaspec, CA) was dissolved in the reaction buffer (300 mM Tris-HCl, pH = 7.5, 150 mM NaCl, 5 mM CaCl<sub>2</sub>) and added at a final concentration of 10 µM. Peptide cleavage was monitored as an increase in fluorescence intensity over time at 460 nm (λ<sub>ex</sub> = 360 nm) with a SynergyMX spectrometer (BioTek Instrument, VT). Reaction was carried out in sortase activity buffer in a volume of 100 µL at 25 °C with 12 µM SrtA<sub>Staph</sub> and 1 mM triglycine.

2.4. Sortase-mediated peptide oligomer synthesis

Reactions were carried out by combining 250 µM methyl ester peptides and 12 µM SrtA<sub>Staph</sub> in sortase buffer (300 mM Tris-HCl, pH = 7.5, 150 mM NaCl, 5 mM CaCl<sub>2</sub>) and incubated at 37 °C for times indicated. At each time point, 20 µL aliquots were withdrawn, and reaction was quenched by adding 5 µL of 0.1% trifluoroacetic acid (TFA). The reaction aliquots were purified through ZipTip<sub>C-18</sub> column (Millipore) and loaded on the target plate for MALDI-TOF/TOF analysis.

2.5. MALDI – TOF/TOF MS

Peptide products in the reaction mixtures were analyzed via Matrix-Assisted Laser Desorption Ion-Time of Flight/Time of flight (MALDI-TOF/TOF) mass spectrometry. All MALDI mass spectra were acquired using a Bruker UltraFlextreme MALDI-TOF/TOF mass spectrometer (Bruker Daltonics). The instrument was used in reflector mode and acquisition was optimized for a mass range from 600 to 3000 Da. All measurements were done with constant laser power of 40. The matrix used was α-cyano-4-hydrocinamic acid (10 mg/mL in 50% acetonitrile with 0.05% trifluoroacetic acid). Experiments were carried with fixed amount of exogenous internal standard (neurotensin, monoiso-GLPR T- ---7 6 25 2 5 - - GLPR T - - - -)

3. Results and discussions

We started our study on SrtA<sub>Staph</sub>-mediated ultra-short peptide ligation using the minimal SrtA<sub>Staph</sub>-recognition sequence based on publications, GGGLPRT-OMe, with a methyl ester at the C-termini. SrtA<sub>Staph</sub>-catalyzed small peptide ligation was very efficient and 97% of GGGLPRT-OMe was converted into peptide oligomers at the end of 24 h reaction (Fig. 1). Resulting peptides were a mixture of dimers, trimers, tetramers, and pentamers in linear and cyclic forms, with dimer and trimer peptides as the main products.

Kinetic studies revealed that linear peptide oligomers formed early and rapidly (Fig. 2). Significant amounts of linear dimer, trimer, and tetramer were found after one hour of reaction when very low concentrations of corresponding cyclic oligomers were detected. However, the amount of cyclic oligomers increased as the reaction time was extended. There was a good dynamic correlation between linear oligomer disappearance and cyclic oligomer formation over the reaction period of 0–5 h. These data show that SrtA<sub>Staph</sub>-catalyzed small peptide ligation includes two reactions: 1) formation of linear oligomers through the head-to-tail peptide ligation, and 2) cyclization of linear oligomers to form cyclic products by followed intramolecular transpeptidation. Interestingly, despite the quick accumulation of linear oligomers at the beginning of the reaction, linear oligomers in hydrolyzed forms (without methyl ester at the C-termini) were maintained at relatively stable levels within the first five hours of reaction. Increase of linear oligomers in hydrolyzed form was observed after a long time (> 5 h) reaction. There was good correlations between the formation of cyclic peptides and the production of corresponded linear peptides in the hydrolyzed forms, indicating the presence of SrtA<sub>Staph</sub>-mediated ring opening reactions in cyclic oligomers. Oligomers containing more peptide units were especially prone to SrtA<sub>Staph</sub>-mediated hydrolysis, and all GGGLPRT tetramers (both cyclic and linear) formed at the beginning of reactions disappeared at the end of 24 h reaction (Fig. 1).

It is known that the polyglycine sequence at the N-terminus of the
peptide can greatly affect SrtA-Staph-mediated protein ligation. Good reactive activity was usually observed for proteins/peptides with three or more glycine residues at N-terminus [22]. SrtA-Staph mediated transpeptidation would not happen for peptides or proteins with only one N-terminal glycine [22,23]. To test the reactivity of SrtA-Staph to small peptides with less than three N-terminal glycines, we extended our study to peptide GLPRT-OMe and GGLPRT-OMe. Both GLPRT-OMe and GGLPRT-OMe gave oligomer products (Fig. 2), indicating that a single aminoglycine was sufficient for nucleophilic attack at the small LPXT containing substrates. These results were in consistent with findings from recent studies [26,27]. However, significant differences in reaction kinetics and product distribution were found for GGGGLPRT-OMe, GGLPRT-OMe, and GLPRT-OMe (Fig. 2): first, reactions of GLPRT-OMe and GGLPRT-OMe were associated with fewer amounts of linear oligomers. The sum of cyclic oligomers in the final products of GLPRT-OMe and GGLPRT-OMe reached 85%, which was higher than the 65% in GGGGLPRT-OMe; second, SrtA-Staph-mediated ring-opening (hydrolysis) reaction was only observed for the cyclic products from GG GLPRT-OMe but not these from GLPRT-OMe or GGLPRT-OMe. Cyclic oligomers from GLPRT-OMe and GGLPRT-OMe increased steadily during the course of reaction (Fig. 3A & B). Obviously, LPRT-G or LPRT-GG linkages in cyclic oligomers from GLPRT-OMe and GGLPRT-OMe were not good substrates for SrtA-Staph, because the longer glycine linker limited the steric accessibility of the substrate to the active site, supporting the finding that LPXTG sequences containing three or more N-terminal glycines are good nucleophiles of SrtA-Staph [22,23]; third, the main products for GLPRT-OMe and GGLPRT-OMe at the end of 24 h reaction were cyclic trimers and tetramers. Neither GLPRT-OMe nor G GLPRT-OMe formed a cyclic dimer, a main cyclic product of GGGGLPRT-OMe. In contrast, cyclic pentamer was only found for LPRT-OMe. Because GGGGLPRT-OMe, GGLPRT-OMe, and GLPRT-OMe existed as random coils in solutions, potential contributions of peptide secondary structures to SrtA-Staph-mediated small peptide ligation could be excluded. Kinetic results suggested that linear dimers did exist at the early phase of GLPRT-OMe and GGLPRT-OMe reactions (Fig. 3C & D) but they failed to proceed intramolecular transpeptidation to form cyclic products. We know that side chain interactions among amino acid residues in cyclic peptides will be much stronger than those in linear ones. Therefore, the strong ring stain might explain why dimers were the main cyclic oligomer for GGGGLPRT-OMe but not for GLPRT-OMe and GGLPRT-OMe. Data from GLPRT-OMe, GGLPRT-OMe and GG LPRT-OMe (Fig. 1) suggest that peptide lengths had dramatic effects on the final products of SrtA-Staph-mediated small peptide ligations.

The important role of peptide length in SrtA-Staph-catalyzed small peptide cyclization was confirmed by using small peptides with increased peptide lengths (7–10 amino acids) but containing only one N-terminal glycine (GHKLPT-OMe, GHHHPPT-OMe, and GVPGVGLPRT-OMe) (Fig. 4). Like other small peptides, all three selected peptides do not have specific secondary structures and exist as random coils in solutions (Fig. 4, top). Cyclic peptides were also the main products (yield > 81%) for all three peptides. Peptide lengths had the same effect on SrtA-Staph-catalyzed peptide ligation: cyclic dimers were major products of GHHHPPT-OMe and GVPGVGLPRT-OMe but were not found for GHKLPT-OMe (Fig. 4, bottom).

SrtA-Staph-mediated small peptide ligations are summarized in Fig. 5. SrtA-Staph catalyzed small peptides to form linear peptide oligomers efficiently (yield > 85%) and rapidly (< 60 min). However, SrtA-Staph was unable to position both ends of small peptide oligomers (less than 14 amino acids) in a productive binding arrangement for the intramolecular amide bond to form cyclic oligomers because of steric hindrance. However, once bigger peptide oligomers were formed, they would undergo intramolecular transpeptidation to form cyclic products. SrtA-Staph catalyzed cyclization was slow in comparison with linear oligomerization, and might take hours to complete. Because adding extra peptide units to large oligomers would become more difficult and oligomers containing multiple copies of G-LPRT sequences became prone to SrtA-Staph hydrolysis, peptide oligomers (both linear and cyclic) with more than 30 amino acids were hardly synthesized from small (< 10 amino acid in length) peptides. On the contrary, once formed, the small ring formed by shorter peptides will have strong steric hindrance and thus cannot readily adopt a conformation to rebind to SrtA for hydrolysis. Therefore, only middle sizes (15–30 amino acid residues) of cyclic peptides formed in SrtA catalyzed small peptide ligation.

An interesting question was what would be the products of peptides with more than 15 amino acids. Such peptides met the minimal length requirement (> 14 amino acids) for cyclization but even dimers of these peptides would exceed the length limit (< 30 amino acids) to generate stable ligation products. We tested SrtA-Staph catalyzed peptide ligation on a 19-aa peptide (GGGWLGALFKALSKLLPRT-OMe). As we had expected, oligomers from this peptide were not found. However, high yield (~83%) and single cyclic product from the direct cyclization of GGGWLGALFKALSKLLPRT-OMe was produced (Fig. 4, Bottom).

There has been an increased interest and rapid expansion in the study of cyclic peptides over the last decade. Strategies for synthesizing cyclic peptides include side chain to side chain, side chain to terminal group, and terminal group to terminal group (head-to-tail) cyclization. These methods, however, are inefficient for cyclization of large peptides (> 10 residues) due to the large entropic barriers for such reactions and competing intermolecular oligomerization. Findings from this study provide useful information to the synthesis of linear and cyclic oligomers from ultra-short peptides for wide pharmaceutical and biomedical applications. We know that cyclic peptides may have other advantages over their linear counterparts due to their unusual biological activity, improved thermodynamic stability, and increased resis-
tance to protease digestion [24, 28, 29]. In addition, nano-structured surfaces and scaffolds are essential for assisting bone and neuron cell growth and have a wide range of biomedical applications [30, 31]. Peptides or peptide oligomers with specific sequences can also self-assemble to form materials with regular nanostructures and demonstrate unusual stability [7, 8]. Because of good biocompatibility, biomaterials using self-assembled peptides instead of synthetic polymers...
have been generated recently [32,33]. Taking peptide GGWLGALFKALSKLLPRT-OME as an example: although both linear and cyclic peptide self-assembled in water, they behaved differently (Fig. 6A) and formed peptide aggregates with distinct super-molecular structures (Fig. 6B).

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Appendix A. Transparency document

Supplementary data associated with this article can be found in the online version at doi:10.1016/j.bbrep.2017.02.005.

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