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Segmental isotopic labeling by asparaginyl endopeptidase-mediated protein ligation

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Received: 9 January 2018 / Accepted: 7 March 2018 / Published online: 13 March 2018 © Springer Science+Business Media B.V., part of Springer Nature 2018

Abstract

Segmental isotopic labeling can facilitate NMR studies of large proteins, multi-domain proteins, and proteins with repetitive sequences by alleviating NMR signal overlaps. Segmental isotopic labeling also allows us to investigate an individual domain in the context of a full-length protein by NMR. Several established methods are available for segmental isotopic labeling such as intein-mediated ligation, but each has specific requirements and limitations. Here, we report an enzymatic approach using bacterially produced asparagine endopeptidase from *Oldenlandia affinis* for segmental isotopic labeling of a protein with repetitive sequences, a designed armadillo repeat protein, by overcoming some of the shortcomings of enzymatic ligation for segmental isotopic labeling.

Keywords Asparaginyl endopeptidase \cdot Designed armadillo repeat protein \cdot Segmental isotopic labeling \cdot Protein *trans*ligation \cdot Protein NMR

Abbreviations

PTS	Protein-trans splicing
CTS	Conditional protein trans-splicing
EPL	Expressed protein ligation
IPL	Intein-mediated protein ligation
NCL	Native chemical ligation
SML	Sortase-mediated ligation
AEP	Asparaginyl endopeptidase
AML	AEP-mediated ligation
SrtA	Staphylococcus aureus sortase A
HSQC	Heteronuclear single-quantum correlation
	spectroscopy
OaAEP1	Oldenlandia affinis asparaginyl endo-
	peptidase 1 _b
HINT domain	Hedgehog/intein domain
dArmRP	Designed armadillo repeat protein

Electronic supplementary material The online version of this article (https://doi.org/10.1007/s10858-018-0175-4) contains supplementary material, which is available to authorized users.

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GFP	Green fluorescent protein
MBP	Maltose-binding protein
Ulp1	Ubiquitin-like-specific protease 1
IMAC	Immobilized metal ion affinity
	chromatography
IPTG	Isopropyl β -D-1-thiogalactopyranoside
CBD	Chitin-binding domain

Introduction

Segmental isotopic labeling is a powerful labeling technique in protein NMR spectroscopy to facilitate NMR studies of larger proteins by not only alleviating signal overlaps in larger proteins but also preserving features of uniformly isotope-labeled samples for triple resonance experiments (Skrisovska et al. 2010; Volkmann and Iwaï 2010). Several approaches have been successfully applied to produce segmentally isotope-labeled proteins. Segmental isotopic labeling methods can be divided into three categories: (1) chemoselective ligation, (2) protein trans-splicing (PTS), and (3) protease-mediated ligation. Each method has some specific requirements and limitations. Chemoselective ligation such as the expressed protein ligation (EPL)/intein-mediated protein ligation (IPL) requires an α -thioester group at the C-terminus and an N-terminal cysteine residue (Fig. 1a) (Dawson et al. 1994; Evans et al. 1998; Muir et al. 1998; Fig. 1 Different approaches for segmental isotopic labeling. a Expressed protein ligation (EPL)/Intein-mediated protein ligation (IPL) using native chemical ligation (NCL). b Protein *trans*-splicing (PTS) by split inteins. c Salt-dependent conditional protein *trans*splicing by split inteins (CTS). d Sortase-mediated ligation (SML). e AEP-mediated ligation (AML)



Xu et al. 1999). PTS requires fusions with a split intein or HINT (hedgehog/intein) domains, which could reduce the solubility of split fragments and might require refolding steps (Fig. 1b) (Yamazaki et al. 1998; Otomo et al. 1999a, b; Muona et al. 2010; Aranko et al. 2013, 2014). However, PTS can also be used for segmental isotopic labeling of a central fragment by multiple-fragment ligation steps using two orthogonal split inteins (Otomo et al. 1999b; Busche et al. 2009). Enzymatic ligation for segmental labeling typically requires specific recognition sequences in the substrates and an enzyme (Mao et al. 2004; Kobashigawa et al. 2009; Freiburger et al. 2015).

Despite its potential usefulness, the applications of segmental isotopic labeling have been relatively limited. This is probably mostly because segmental isotopic labeling is often seen as technically demanding and labor-intensive, as well as requiring undesired amino-acid changes for ligation. For example, multiple-step protein purifications of at least two fragments are required together with optimizations for the ligation conditions to generate the ligated protein (Otomo et al. 1999a; Skrisovska and Allain 2008; Minato et al. 2012).

There have been several improvements reported for each of these methods. PTS can also be performed in vivo without multiple protein purification steps (Züger and Iwai 2005; Muona et al. 2010). Ligation by EPL can also be improved by different thiol reagents or on-column ligation (Johnson and Kent 2006; Skrisovska and Allain 2008; Michel et al. 2013; Gallagher et al. 2017). Conditional splicing using a salt-inducible intein has been recently exploited to alleviate the solubility issue of split inteins (Fig. 1c) (Ciragan et al. 2016). Enzymatic ligation using sortase A (SrtA) from *Staphylococcus aureus* is an attractive approach because it requires only short tags instead of larger split intein

fragments (Mao et al. 2004). Sortase-mediated ligation (SML) has increasingly become popular in protein modification (Fig. 1d) (Mao et al. 2004; Antos et al. 2016). However, SML has strict sequence requirements and thus does not permit arbitrary ligation sites (see below).

For segmental isotopic labeling of proteins, the major challenge is to achieve both high ligation efficiency and very high purity, which is required for NMR studies (Muona et al. 2010; Minato et al. 2012). One of the drawbacks of any enzymatic ligation, including SML, could be the reverse reaction catalyzed by the enzyme because the ligated product still contains the recognition sequence. The reverse reaction could be suppressed by constantly removing one of the products (Freiburger et al. 2015). Moreover, SrtA is a relatively inefficient enzyme with the low turnover rate. Therefore, a large amount (stoichiometric amount) of the enzyme and/or longer reaction time is often used to produce a large quantity of ligated proteins required for NMR investigations (Kobayashigawa et al. 2009; Freiburger et al. 2015). Variants of SrtA have been developed to improve the transpeptidase kinetics of SrtA (Dorr et al. 2014). However, another shortcoming of SrtA is an insertion of "Leu-Pro-Xxx-Thr-Gly-Gly" (LPXTGG) sequence at the ligation site, which is necessary for SrtA to ligate. The recognition sequence introduces a non-natural sequence and could thus affect biological functions of ligated proteins. Therefore, it might not be the best choice for NMR studies of biologically active proteins.

In contrast, the PTS approach using an intein bearing Ser at the +1 position has been successfully demonstrated to be valuable for producing a segmentally isotope-labeled protein with the native sequence, which can be accommodated owing to the sequence diversity of naturally occurring inteins and HINT domains (Ciragan et al. 2016).

Recently, we demonstrated segmental isotopic labeling of a single-domain globular protein without any refolding step using asparaginyl endopeptidase (AEP) from Oldenlandia affinis (OaAEP1) (Mikula et al. 2017). AEPs from plants catalyze an efficient transpeptidase reaction in backbone cyclization of cyclotides (Fig. 2) (Nguyen et al. 2014; Harris et al. 2015). Importantly, OaAEP1 was produced from an E. coli bacterial expression system similar to SrtA, making it an attractive alternative enzyme for segmental isotopic labeling (Harris et al. 2015; Mikula et al. 2017). The natural recognition sequences of OaAEP1 are minimal, "Gly-Leu"(GL) and "Thr-Arg-Asn/Gly-Leu" (TRN/GL), in the natural substrate of Kalata B1 precursor (a cyclotide) for the N- and C-termini respectively, in which "/" indicates the cleavage and ligation site (Fig. 1e) (Harris et al. 2015). Additionally, OaAEP1 seems to be tolerant of variations in the substrate sequence, even though the full range of tolerable mutations is still unknown. This suggests that the practical sequence requirement can even be shorter in length with the

(a) Enzymatic ligation



Fig. 2 Reaction types catalyzed by AEP. a Enzymatic bi-molecular *trans*-ligation. b Backbone cyclization. c The proximity effect in enzymatic ligation

theoretical requirement of only one residue of "Asn" (Harris et al. 2015; Mikula et al. 2017), but the scope of possible sequence changes remains to be investigated. We previously reported that proteins with the sequences of "TRN/AL" or "TRN/CL" in the C-terminal propeptide sequence were much more efficient substrates than the original "TRN/GL" for recombinant *Oa*AEP1 at least for the backbone cyclization of a model protein of green fluorescent protein (GFP) (Mikula et al. 2017).

Here, we report the use of asparagine endopeptidase (AEP) for production of segmental isotope-labeled proteins by overcoming the shortcoming of AEP-mediated ligation for complementing existing segmental isotopic labeling techniques and discuss the possible applications and limitations.

Materials and methods

GFP substrates for trans-ligation

The N-terminal substrate GSH_6 -(GFP)-TRNCL was produced from the vector pEMRSF2, derived from pJTRSF50 using two oligonucleotides, I940: 5'-ACA TAT GGG CAG CCA TCA TCA CCA TCA CCA C and J052: 5'-AGG AAG CTT ACA GAC AAT TTC GGG TAC TAC CGC G (Muona et al. 2010; Mikula et al. 2017). The C-terminal substrate of GLPH₆-(GFP(N159S))-GLT was produced as a SMT3 fusion protein (the yeast SUMO protein) from the plasmid pBHRSF277 bearing an N159S mutation in GFP introduced by using the two oligonucleotides, J166: 5'-GAA GTT AGC TTT GAT CCC ACT CTT TTG TTT GTC TGC and J167: 5'-GCA GAC AAA CAA AAG AGT GGG ATC AAA GCT AAC TTC and pBHRSF208 as the template. The protein was expressed as a SUMO fusion and purified as previously described (Guerrero et al. 2015; Mikula et al. 2017).

Trans-ligation using model globular proteins

Purified substrate proteins were mixed at an equimolar ratio at a final concentration of 5 μ M each in the reaction buffer of 50 mM HEPES, pH 7.0, 50 mM NaCl, 1 mM EDTA, and 0.5 mM TCEP. Pre-activated recombinant OaAEP1, of which self-proteolysis was induced for the activation by lowering pH of the buffer to 4.0 for 5 h, was added at a final concentration of 0.2 µM to GFP variants. Reactions were incubated at room temperature in 1.5 mL Eppendorf tubes. The samples for SDS-PAGE analysis were taken every 10 min for 100 min for GFP variants. The time course of the trans-ligation of two GFP variants was analyzed by SDS-PAGE. The samples taken at 10-min intervals for 100 min were diluted 1:1 with 2-times SDS loading buffer, and loaded on 12% SDS polyacrylamide gels. The SDS-gels were stained with Coomassie Blue R (GE Healthcare) and quantified using NIH ImageJ software with the assumption that the dye binds equally to the proteins (Schneider et al. 2012).

Segmental isotopic labeling of designed armadillo repeat protein (dArmRP)

The full-length dArmRP (YM₃A) consists of the N-terminal capping repeat (Y), three consensus repeats (M) and a C-terminal artificially designed capping repeat (A). The gene of YM₃A was split into two fragments of YM and MMA with the N- and C-terminal additions as below. The N-terminal fragment was designed to have the C-terminal sequence of "NCL" for ligation by OaAEP1. The N-terminal fragment was cloned into pHYRSF53 as H₆-SUMO fusion at the N-terminus, using the two oligonucleotides, HK542: 5'-AAG GAT CCG AAC TGC CGC AGA TGA C and J153: 5'-GGA AGT GTT GTT TCA GGG TCC TGA TGG CAA CGC GCT GAC CC, resulting in pEMRSF9 bearing H₆-(SMT3)-(YM) with "NCL" at the C-terminus. The gene of the C-terminal fragment of MMA with the N-terminal sequence of "GL" for ligation was amplified by PCR using the two oligonucleotides, J157: 5'-ACA TAT GGG TCT TAA CGA ACA AAT CCA AG and J223: 5'-TTA TGA ATT CGT GGG ACT GCA GCT TCT C. The PCR product was cloned into pMHBAD14, resulting in pEMBAD26 bearing GL-(M₂A)-H₆ (Parmeggiani et al. 2008; Oeemig et al. 2009; Alfarano et al. 2012).

For segmental isotopic labeling, the N-terminal fragment (YM) was expressed in 20% ¹³C, 100% ¹⁵N-labeled M9-medium containing ¹⁵NH₄Cl (1 g/L) and a mixture of $^{13}C_6$ D-glucose (0.2 g/L) and unlabeled D-glucose (0.8 g/L) as nitrogen and carbon sources. T7 Express Competent E. coli cells (New England Biolabs) were transformed with the plasmid (pEMRSF9) for protein expression. The cells were grown at 37 °C in 2 L of M9-medium supplemented with 25 µg/mL kanamycin and induced with a final concentration of 1 mM of isopropyl β -D-1-thiogalactopyranoside (IPTG) when the OD_{600} reached 0.6. The cells were further incubated for 3 h at 200 rpm before harvesting. The protein was purified by immobilized metal ion affinity chromatography (IMAC) using a HisTrap 5 mL column (GE Healthcare) as described previously and dialyzed against 2 L of 20 mM Tris buffer, pH 8.0 and 150 mM NaCl overnight (Guerrero et al. 2015). The C-terminal fragment of GL-(MMA)-H₆ was expressed in the E. coli strain ER2566 at 37 °C using pEMBAD26 in 2 L of LB medium supplemented with 100 µg/mL ampicillin and induced for 3 h with a final concentration of 0.02% arabinose. The protein was purified by IMAC using a 5 mL HisTrap column and dialyzed against 2 L of 20 mM Tris buffer, pH 8.0 and 150 mM NaCl, overnight, at 8 °C.

The labeled N-terminal fragment (YM)-NCL (133 µM) and the unlabeled C-terminal fragment GL-(MMA)-H₆ (67 μ M) was mixed at a 2:1 molar ratio in a volume of 1.5 mL. The ligation reaction was initiated by addition of 420 µL of the pre-activated recombinant OaAEP1, of which proteolytic activity was activated by lowering pH to 4.0 for 5 h, at a final concentration of 2.7 µM. The preparation of activated recombinant OaAEP1 using the plasmid pBHRSF184 (Addgene ID #89482) was previously reported (Mikula et al. 2017). The reaction mixture was immediately transferred into a dialysis tube (3.5 kDa MWCO) and dialyzed at room temperature against 0.5 L of 50 mM HEPES buffer, pH 7.0, 50 mM NaCl, 1 mM EDTA, and 0.5 mM TCEP overnight. The reaction mixture was further purified by anion exchange chromatography using a MonoQTM 5/50 GL column (GE Healthcare Life Sciences). The fractions containing the ligated product were collected and dialyzed overnight at 8 °C against 1 L of 20 mM sodium phosphate buffer, pH 6.0 and concentrated to 0.58 mM using an ultracentrifugation device.

For the comparison, a full-length protein was also produced in the labeled M9 medium using the plasmid (pADHRSF57) following the same purification protocol and concentrated to 0.45 mM in 20 mM sodium phosphate buffer, pH 6.0.

Amino acid tolerance of *Oa*AEP1 at the N-terminal P1" and P2" sites

To test the promiscuity of OaAEP1 at the N-terminus (P1" site in Fig. 6a), we prepared GFPs with three different

amino acids (Gly, Ala, and Ser) at the N-terminus. GLP-(GFP)-RNALPH₆, ALP-(GFP)-RNALPH₆, SLP-(GFP)-RNALPH₆ were produced as the N-terminal SUMO-fusion proteins from the plasmids of pJTRSF82, pKERSF10, and pKERSF11, respectively. The removal of the N-terminal SUMO domain in these fusion proteins by Ulp1 protease created the N-terminal Gly, Ala, or Ser. pJTRSF82 was derived from pJTRSF55 using the following oligonucleotides, #78GFP_N: 5'-TGG GAT CCA AAG GAG AAG AAC NTT NC and J164: 5'-GAT GAT GAT GAT GAT GAC CCA GAG CAT TTC GGG TAC T, and HK122: 5'-CTA AAG CTT AAT GAT GAT GAT GAT GAT G to add the C-terminal His₆ tag. pKERSF10 and pKERSF11 were constructed using the following oligonucleotides, J136: 5'-TAG GTA CCC GGC AGT GCT CCA CCA ATC TGT TCT C and J138: 5'-TAG GTA CCC GGC AGT GAT CCA CCA ATC TGT TCT C, respectively and cloned between the SpeI and KpnI sites of pJTRSF82. For testing the mutations at the P2" position, GVP-(GFP)-RNALPH₆, GIP-(GFP)-RNALPH₆, GFP-(GFP)-RNALPH₆, GMP-(GFP)-RNALPH₆, were produced as the N-terminal SUMO-fusion proteins from the plasmids of pJTRSF125, pJTRSF127, pJTRSF128, and pJTRSF129, respectively. JTRSF125, pJTRSF127, pJTRSF128, and pJTRSF129 were created by PCR using the following oligonucleotides containing the mutations, J290: 5'-TAG GTA CCC GGA ACT CCT CCA CCA ATC TGT, J292: 5'-TAG GTA CCC GGG ATT CCT CCA CCA ATC TGT, J293: 5'-TAG GTA CCC GGC ATT CCT CCA CCA ATC TGT and JT010: 5'-TAG GTA CCC GGG AAT CCT CCA CCA ATC TGT, respectively.

The protein expression was induced at 22 °C and proteins were purified as described previously (Mikula et al. 2017). Cyclization reactions of GFP variants were carried out, of which three varied the N-terminal residue ('GLP', 'ALP', and 'SLP') and four varied the second residue (the P2" position in Fig. 6a; 'GVP', 'GIP', 'GMP, and 'GFP') together with Ala at the P1' position (Fig. 6). These were performed using 5 μ M of the substrate and 0.2 μ M of activated recombinant *Oa*AEP1 and analyzed as previously reported (Mikula et al. 2017).

NMR measurements

 $[{}^{1}\text{H}, {}^{15}\text{N}]$ -HSQC spectra were recorded on a Bruker Avance III HD spectrometer equipped with a cryogenically cooled probe at a ${}^{1}\text{H}$ frequency of 850 MHz. The segmentally labeled sample was concentrated to 0.58 mM, 625 µL. The uniformly labeled dArmRP was concentrated to 0.45 mM, 3.8 mL. 450 µL solution containing 5% D₂O of each of the samples was transferred into NMR tubes for the measurements.

Results

Trans-ligation of two globular domains without any affinity for each other by *Oa*AEP1

Efficient enzymatic ligation of two globular domains with a peptide bond would make it very simple to introduce segmental isotopic labeling for multi-domain proteins. Robust enzymatic activities of backbone cyclizing AEPs from plants are very attractive for protein ligation, compared to the widely used SrtA. AEPs would require a smaller amount of enzyme and the sequence modification can be a half or fewer residues in length of what is used with SrtA. While the reaction is much faster than SrtA with smaller enzyme-to-substrate ratios, the ligation efficiency of two globular proteins (GFPs) without any affinity for each other to create a tandem fusion was not impressive (Fig. 3). The low efficiency is presumably due to the occurence of the reverse reaction (Fig. 2a). Therefore, it might be unlikely to achieve a high ligation yield (>90%) (Fig. 3c). This could restrict the application of AML for trans-ligation of multiple domains when the ligation efficiency is critical, e.g., segmental isotopic labeling. The shorter recognition sequence of *Oa*AEP1 might also lead to non-specific cleavages because possible non-specific degradation was previously observed when a small domain of the B1 domain of IgG binding protein G (GB1) was ligated to GFP (Mikula et al. 2017). Thus, smaller less stable proteins and flexible peptide linkers might not be suitable as the substrate of OaAEP1. This promiscuity of OaAEP1 might thus limit the application of AML and needs to be further investigated.

Ligation of a designed armadillo repeat protein

Similar to SML, a ligated product of interest by AML remains the substrate for AEP, resulting in the reverse reaction (Fig. 2a). Therefore, the ligation yield might not be very high unless the ligated product is energetically or entropically favorable (Fig. 3). For example, the ligation yield has been improved by continuously removing one of the products during the reaction (Freiburger et al. 2015). As OaAEP1 is very efficient having > 200 times faster ligation kinetics than SrtA, such an approach to remove a product might not be very practical. To overcome this problem, the proximity effect can be exploited for improving the ligation efficiency (Fig. 2c). Backbone cyclization of the natural substrate (Kalata B1) of OaAEP1 is very efficient as efficiency close to 100% was easily achieved (Harris et al. 2015). This is presumably due to the proximity effect by N- and C-termini being closely neighbored in the structure (Fig. 2b).

Fig. 3 AEP-mediated protein ligation in *trans*. a Schematic illustration of *trans*-ligation of two GFP variants catalyzed by *Oa*AEP1. b SDS-PAGE analysis of the time course of the ligation reaction of two GFP substrates. M stands for the molecular marker. Arrows indicate the bands of substrates and the ligated product. c The estimated ligation yield versus time



Previously, we produced a nicked maltose-binding protein (MBP) with AEP-tag by using a dual co-expression system of two fragments that can reassemble, in which the nicked site has N- and C-termini in spatial proximity (Mikula et al. 2017). It was possible to ligate > 90% of the nicked MBP by the recombinant OaAEP1 with two residue mutations in the loop by exploiting the natural sequence of "NG" in MBP, supporting the notion that the proximity effect can improve the ligation efficiency, similar to the effect seen in backbone cyclization (Mikula et al. 2017). For avoiding subsequent in vitro refolding steps of MBP, time-delayed dual co-expression was used by co-expressing the two split fragments, first in one medium, then in the other (differently labeled), to produce the nicked MBP with only one of the fragments labeled (Züger and Iwai 2005; Mikula et al. 2017).

However, not all proteins can be produced as nicked proteins by co-expression of two split fragments. This requirement might constrain the wider applications of this approach. Therefore, we were interested in other protein fragments with inherent affinities in vitro. Armadillo repeats are found in many proteins with a repetitive amino acid sequence of about 40 residues in length. Designed armadillo repeat proteins (dArmRP) have been developed as a promising modular scaffold protein for the engineering of binding molecules that recognize extended polypeptide chains (Parmeggiani et al. 2008; Reichen et al. 2014). Interestingly, dArmRP fragments that have been split between the consensus repeats can spontaneously assemble into one globular protein by non-covalent association (Watson et al. 2014). Thus, this re-associated complex would be an ideal substrate for AEP. Moreover, segmental isotopic labeling is particularly useful for proteins with repeating sequences because NMR signals tend to overlap due to the similar chemical environments within the repeating sequences (Busche et al. 2009). As a model system, we chose a consensus dArmRP (YM_3A) protein, consisting of three identical internal repeats (M), flanked by N- and C-terminal capping repeats (Y and A, respectively). YM₃A was split into two fragments within a loop after the first internal repeat (M_1) , named YM, and M₂A fragments, respectively (Figs. 4a, 5a). We added four residues at the C-terminus of the YM fragment as an AEP-tag, i.e., "KNCL", which is the same sequence we previously used (Fig. 5b) (Mikula et al. 2017). For the C-terminal fragment of M₂A, we created a "GL" sequence at the N-terminus. The ligated product would have a threeresidue insertion because we utilized a Gly residue in the native sequence (Fig. 5b). We ligated these two fragments in vitro by *Oa*AEP1 at a 1:25 enzyme-to-substrate ratio. The ligation was not as fast as the two-GFP ligation, but the ligated product yield exceeded 50% (Fig. 4b). Increasing the amount of YM fragment by twofold improved the ligated product yield, presumably because the equilibrium





Fig.4 Ligation of the designed armadillo repeat fragments by OaAEP1. **a** Schematic representation of the production of segmentally labeled dArmRP by protein-ligation catalyzed by OaAEP1. **b** SDS-PAGE analysis of *trans*-ligation between the N-terminal YM and C-terminal M₂A from dArmRP. 0 and 3 h indicate hours after

was shifted towards the complex. The ligated product yield was >90% after overnight incubation (Fig. 4b).

Segmental isotopic labeling by OaAEP1

Next, we proceeded to produce a segmentally isotopelabeled sample by preparing the N-terminal fragment of YM in the ¹⁵N-labeled medium. The solution mixture containing the two purified labeled YM and unlabeled M_2A fragments was simply incubated with the activated recombinant *Oa*AEP1 at an enzyme-substrate-ratio of 1:25 and the product was further purified by anion exchange chromatography to remove unreacted substrates and the enzyme (Fig. 4c, Supplementary Fig. 1). Figure 5c shows the HSQC spectrum of the segmentally ¹⁵N-labeled YM₃A with a three-residue insertion in the YM region (red in Fig. 5). The spectrum

the addition of *Oa*AEP1, O/N and M stand for overnight incubation and molecular marker, respectively. Arrows indicate C- and N-terminal fragments and full-length dArmRP as the ligated product. **c** SDS-PAGE analysis of the segmentally labeled dArmRP after anionexchange chromatography

with the reduced number of dispersed peaks indicates the well-folded fragment as previously reported, suggesting the successful segmental isotopic labeling (Watson et al. 2014). We compared the HSQC spectra between the original YM₃A without any ligation and the ligated YM₃A (Fig. 5c and Supplementary Fig. 2) and observed small shifts of several peaks and additional peaks, presumably originating from the newly inserted residues. Even though the amino acid changes are smaller than ones required for SrtA, the insertion in the loop has some influences on the NMR spectrum, and possibly on the three-dimensional structure as well, inducing some chemical shift changes (Supplementary Fig. 2). This observation implies that a small insertion might still influence the functionality of the ligated protein, and it might be necessary to be further minimized for producing biologically active proteins with native primary structures.





(b)

N-terminal fragment

- Y: SELPOMTOOLNSDDMQEQLSATRKFSQILSD
- M: GNEQIQAVIDAGALPALVQLLSSPNEQILQEALWALSNIASGKNCL

C-terminal fragment

- M: GLNEQIQAVIDAGALPALVQLLSSPNEQILQEALWALSNIASG
- M: G-NEQIQAVIDAGALPALVQLLSSPNEQILQEALWALSNIASG
- A: G-NEQKQAVKEAGALEKLEQLQSHENEKIQKEAQEALEKLQSH EFHHHHHH

Ligated product

- Y: SELPOMTOQLNSDDMQEQLSATRKFSQILSD
- M: G-NEQIQAVIDAGALPALVQLLSSPNEQILQEALWALSNIASGKN
- M: GLNEQIQAVIDAGALPALVQLLSSPNEQILQEALWALSNIASG
- M: G-NEQIQAVIDAGALPALVQLLSSPNEQILQEALWALSNIASG
- A: G-NEQKQAVKEAGALEKLEQLQSHENEKIQKEAQEALEKLQSH EFHHHHHH



ω₁ (¹⁵N) [ppm]



Fig. 5 Segmental labeling of dArmRP. **a** A cartoon model of the structure of the segmentally labeled dArmRP (YM₃A) in which [¹³C, ¹⁵N]-labeled YM region and M₂A unlabeled regions are colored in red and dark blue, respectively. **b** Sequences of N- and C-terminal fragments in red and black, respectively; [¹³C, ¹⁵N]-labeled N-terminal fragment for YM domains containing a C-terminal "KNCL" rec-

ognition tag for AEP (underlined and in italics); C-terminal fragment for M_2A domains with the N-terminal "GL" recognition tag for AEP (underlined); the expected sequence of the ligated product. **c** [¹H, ¹⁵N]-HSQC spectra of uniformly labeled (left) and segmentally [¹³C, ¹⁵N]-labeled dArmRP produced by AML (right)



Fig. 6 The promiscuity of *Oa*AEP1 at the N-terminal recognition sequence. **a** Schematic representation of the substrate (GFP) and reaction catalyzed by *Oa*AEP1. The C-terminal P1–P3 and P1'–P3' sites of the C-terminal propeptide are indicated. P1" and P2" denote the N-terminal residues that replace the P1' and P2' residues after cleavage of the C-terminal propeptide. The amino acid sequence

corresponding to the natural substrate of KalataB1 are shown on the

top. The amino acid types reported in this work and previous reports

are shown below (*a*Harris et al. 2015 and *b*Mikula et al. 2017). **b** SDS-PAGE analysis of backbone cyclization of GFP substrates by OaAEP1. Variants of the N-terminal recognition sequence at the P1" or P2" site are indicated above the gels; 0, 0.5, 1, and 2 h indicate the number of hours after addition of the enzyme; M stands for molecular weight marker, and arrows indicate the bands of linear and circular GFPs

Expanding possible ligation sites by OaAEP1

We further asked if we could use OaAEP1 for other sequences because another AEP, butelase 1, was found to be promiscuous at the N-terminal sequence (Nguyen et al. 2014). We mutated the N-terminal Gly to Ala or Ser of the GFP model substrate, which was created after removing the N-terminal SUMO domain by Ulp1 digestion, and tested the backbone cyclization of the GFPs by OaAEP1 (Fig. 6) (Mikula et al. 2017). The circular form of GFP migrates faster than the linear form in the SDS-PAGE after successful backbone cyclization and can thus be used to monitor the transpeptidase activity (Iwai et al. 2001). Both Ala and Ser at the N-terminus (the P1" position) in the GFP were cyclized as efficiently as GFP with the N-terminal Gly when the P1' position is Ala (Fig. 6b). Previously, Gln and Lys was successfully used as the N-terminal residue at the P1" position for backbone cyclization of peptides by OaAEP1 (Harris et al. 2015). Additionally, we tested a few hydrophobic amino-acid types at the P2" position following the N-terminal residue for backbone cyclization of the model GFP (Fig. 6b). These data suggest that the sequence requirement by OaAEP1 is not strictly limited to "NGL" but could be used for other sequences, thereby widening the application of AML using OaAEP1. However, the full scope of permissible amino acids has not been fully elucidated because the number of amino-acid combinations is 3.2×10^6 for the five positions in the recognition sequence.

Discussion

The ability to efficiently ligate peptide fragments and thereby introduce segmental labeling in proteins at any desired region without extensive optimizations and sequence alterations is a challenge of great importance in many areas, including protein NMR as well as protein engineering. Transpeptidases with high turnover rates can be an attractive approach because enzyme-mediated ligation would then require only a small amount of the enzyme for ligation.

In this study, we demonstrated that bacterially produced asparaginyl endopeptidase from *Oldenlandia affinis* (OaAEP1) could be used to efficiently produce a segmentally isotope-labeled dArmRP. The high turnover and shorter recognition sequence of OaAEP1 make AML an attractive alternative to SML. The minimal length required for the ligation site using OaAEP1 can be as short as three residues in total, which is shorter than the required sequence of "LPXTGG" for SrtA-mediated ligation. This feature is advantageous when ligated products need to have native sequences. However, for the higher ligation efficiency, the ligation reaction must be preferred by the enzyme to the reverse reaction, such as, e.g. by the proximity effect.

The ligation site we used in this model system of dArmRP contained the sequence "KNGL", after completion of the reaction. However, the N-terminal residue of the C-terminal fragment can also be Ala or Ser instead of Gly for a transpeptide reaction, thereby expanding the possible sequence combinations by OaAEP1. Our experiments with a few variants at the P2" position suggest that "Leu" at the P2" position could also be replaced at least by a few hydrophobic residues such as "Val", "Ile", "Met", and "Phe" together with Ala at the P1' position (Fig. 6b). Further characterization of the specificity of OaAEP1 in detail will widen the application of OaAEP1 for segmental isotopic labeling as well as other protein engineering applications. AML does not require any fusions with large protein fragments like split inteins but currently requires only three residues at the C-terminus as the AEP-tag. Compared with AML, the PTS approach was successfully used to produce a native sequence after splicing, because of the high diversity of intein sequences and high specificity of the splicing reaction (Ciragan et al. 2016). Although the promiscuity of OaAEP1 could be beneficial for finding an appropriate ligation site in the native sequence, the tradeoff can be non-specific cleavages, and this might be necessary to be suppressed in, if not all, some cases.

There are now several approaches for segmental isotopic labeling. Each method still has specific requirements and restrictions. It is unlikely to have only one robust method that could be used for every situation of all proteins. Further characterization and engineering of inteins, AEPs, and SrtA could establish a general toolbox for segmental isotopic labeling of proteins and thereby segmental isotopic labeling can be used more widely for NMR studies of larger proteins, proteins with a repetitive sequence, and multi-domain proteins. Particularly transient interactions, present within large multi-domain proteins, might be difficult to observe without segmental isotopic labeling even though they may play critical roles in biological functions (Minato et al. 2017; Shiraishi et al. 2018). Acknowledgements This work is supported by grants from the Academy of Finland (137995, 277335). The NMR facility at the Institute of Biotechnology, University of Helsinki is supported by Biocenter Finland and Helsinki Institute of Life Science (HiLIFE). The authors thank K. Elsner, A. Hietikko, S. Jäskeläinen, J. Tommila, and E. Maschke for their technical assistance.

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