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Pyrazolone ligation-mediated versatile sequential bioconjugations†

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Bioconjugation chemistries are critical tools in biotherapeutics discovery. The past efforts have been exclusively focused on two-segment conjugations. However, emerging research directions, such as polypharmacy biotherapeutics, desire multiple-component bioconjugations where more than two pharmacologically related biomolecules can be assembled into a single construct in high efficiency. We present here a set of sequential bioconjugation chemistries centered on a pyrazolone structural motif. It starts with a clickable “pyrazolone ligation” between a hydrazine group and a β -ketoester moiety followed by the conjugation between the newly formed pyrazolone core and an aldehyde-bearing biomolecule through a Knoevenagel reaction forming a Michael addition acceptor that can effectively capture a thiol-bearing biomolecule. When utilized intermolecularly, it quickly assembles four segments together forming a quadruple functional construct. When applied intramolecularly, it offers a set of highly diverse biomolecule scaffolds including stapled peptides and poly-macrocyclic peptides. We envision broad utilities of such sequential ligation chemistries.

Introduction

Bioconjugation chemistries are being routinely employed in the discovery of biotherapeutics. They play crucial roles in constructing dual functional single molecule constructs by joining two bioactive entities through a covalent linkage.¹ One notable success is the regulatory approval of 8 antibody–drug conjugate (ADC) drugs by the end of 2019, such as Adcetris and Kadcyla.² In addition, these methods offer powerful tools enabling fundamental research at the interface of chemistry and biology.

A significant number of bioconjugation reactions have been established,^{1,3} a frequently-utilized short list consists of alkyne–azide cycloaddition,⁴ oxime/hydrazone ligation,⁵ Staudinger ligation,⁶ tetrazine ligation,⁷ alkylation of cysteine thiol,^{1a,3} and acylation of the ϵ -amine of lysine.^{1a,3} The past efforts have been exclusively focused on improving the conversion efficiency, reaction kinetics, selectivity, and the drug-likeness of the newly-formed junctions. Currently these criteria can often be satisfactorily met.⁸ However, new directions constantly emerge in pharmaceutical and biotechnology field that raise the bar for our capabilities in generating new chemical classes. For instance, the integration of multiple pharmacological elements such as multiple agonists and/or antagonists, tissue targeting motif, and pharmacokinetic modulators into a single therapeutic construct represents a highly promising direction.⁹ In

addition, multiple modalities, such as small molecule,¹⁰ antibody,¹¹ protein,¹² peptide,¹³ oligonucleotide,¹⁴ gene¹⁴ and cell therapies,¹⁵ presently have all been well-embraced by the community. The desire of efficiently integrating multiple pharmacological components, possibly in heterogeneous molecular formats, requires the development of new bioconjugation chemistries as the existing methods are designed for connecting two molecules. Driven by such needs, we have been embarking on a new field that focuses on developing chemistries which can covalently joint multiple components together with high efficiency (Fig. 1).

Here we describe a set of sequential bioconjugation chemistries centered on a pyrazolone structural motif. It begins with a clickable “pyrazolone ligation” between a hydrazine group and a β -ketoester moiety generating a 5-membered cyclic junction site. Subsequently the resulting pyrazolone can be readily conjugated to an aldehyde through a Knoevenagel reaction forming a Michael addition acceptor that can

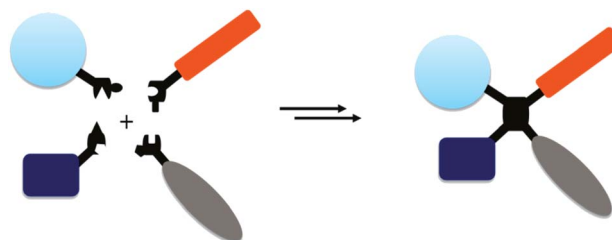


Fig. 1 Cartoon representation of a 4-component sequential bioconjugation.

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effectively capture a free thiol group. When such chemistry is utilized intermolecularly, it can quickly assemble four conjugation partners together forming a quadruple functional construct. When engaging intramolecular reactions, it offers a set of highly diverse biomolecule scaffolds including further derivatized stapled peptides and bicyclic peptides, and potentially tricyclic peptides.

Results and discussion

Pyrazolone ligation

We envisioned the 5-membered cyclic functional group pyrazolone could serve as a core structural motif that may enable an efficient covalent assembly of multiple biomolecules into a single construct. Such vision was based on our hypothesis that the pyrazolone core could be formed through the conjugation between a β -ketoester-bearing biomolecule and a hydrazine-bearing biomolecule, and the fact that it has been reported as reactive toward aldehyde-bearing biomolecules providing the possibility of subsequent conjugations.¹⁶

The synthesis of pyrazolone compounds through a tandem reaction between β -ketoester and hydrazine has been practiced in classical medicinal chemistry. Although it was typically conducted under conditions that are too harsh to biomolecules.¹⁷ To the best of our knowledge, there have been no reported attempt of employing such chemistry to bioconjugation. On the other hand, despite that hydrazone ligation is an attractive procedure attributed to the specificity and simplicity; its usefulness is significantly limited by the reversibility.^{5,18} Our proposed “pyrazolone ligation”, involving a tandem process of hydrazone ligation-lactam cyclization which locks the metastable hydrazone bond into an irreversible structure moiety, should offer an improved alternative to the hydrazone ligation.

We started our investigation by examining the reactivity of different β -ketoesters **1** with hydrazine (Fig. 2a). Among the initially assessed compounds, it was found that β -oxo-benzenepentanoic methyl ester possessed significantly faster reaction kinetics than β -oxo-benzenepropanoic methyl ester and its derivatives. In addition, the electron-withdrawing substituents on the latter were favorable for the ligation rate while the electron-donating groups were detrimental. In all cases, there was no non-cyclized hydrazone intermediate observed in the reaction mixture, suggesting this thermodynamically favorable 5-membered lactam cyclization is fast while the hydrazone formation represents the rate-limiting step. In addition, it was confirmed that such tandem reaction between alkyl hydrazine and β -oxo-benzenepentanoic methyl ester was highly regioselective, 96 : 4 for methylhydrazine and single product for bulkier α -methylbenzylhydrazine as determined by LC-MS and NMR (Fig. S1†). Such data suggests that the conjugation between β -oxo-benzenepentanoic methyl ester- and hydrazine-containing peptides should be highly regioselective.

In the next step, we studied the structure-reaction kinetics relationship of the pyrazolone ligation by derivatizing both β -ketoester and hydrazine segments. It was found that the contraction by one methylene or expansion up to two methylene groups between the phenyl group and β -ketoester unit had

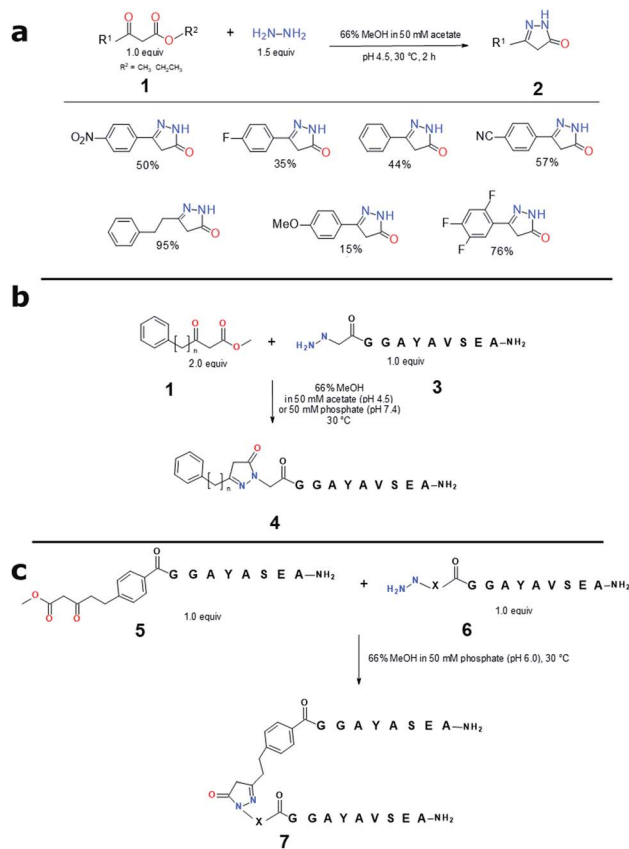


Fig. 2 The screening and optimization of pyrazolone ligation substrates. (a) Screening of β -ketoesters **1** with hydrazine. The reactions were performed at 5.00 μmol scale, and the yield was determined by LC-MS of the crude reaction mixture at 2 h. (b) Optimization of β -ketoester structure with a hydrazine-bearing peptide **3** ($n = 1, 2, 3, 4$). (c) Optimization of the hydrazine structure with a β -ketoester-bearing peptide **5**; the x groups in peptides **6** were $-\text{CH}_2-$ (peptide **3**), $-(\text{CH}_2)_5-$, or $-\text{CH}_2-\text{C}_6\text{H}_4-$.

negligible impact on the ligation rate with the hydrazine peptide **3** (Fig. 2b, S2 and Table S1†). In contrast, among the hydrazine groups examined by ligating peptides **6** to the β -ketoester-bearing peptide **5**, 2-hydrazinyl acetyl group outperformed others (Fig. 2c, S3 and Table S2†). In addition, the 2-hydrazinyl acetyl-peptides can be readily prepared and possess a favorable chemical stability profile. Furthermore, for all these ligations involving short peptides, only a single product was observed on UPLC-MS (Fig. S2 and S3†). The collective data led us to choose β -oxo-benzenepentanoic methyl ester and 2-hydrazinyl acetyl as our preferred pyrazolone ligation precursors, and they were used in the following studies where the scope of the ligation was evaluated.

In the first application case study, we aimed to conjugate a known albumin binding domain (ABDcon)¹⁹ to the C-terminus of a glucagon-like peptide 1 (GLP-1) analog (Fig. 3a). The resulting construct was expected to be fully active based on the well-established GLP-1 structure-activity relationship (SAR)²⁰ while possessing prolonged *in vivo* half-life attributed to the albumin binding. The synthesis of both GLP-1 analog **8** and the



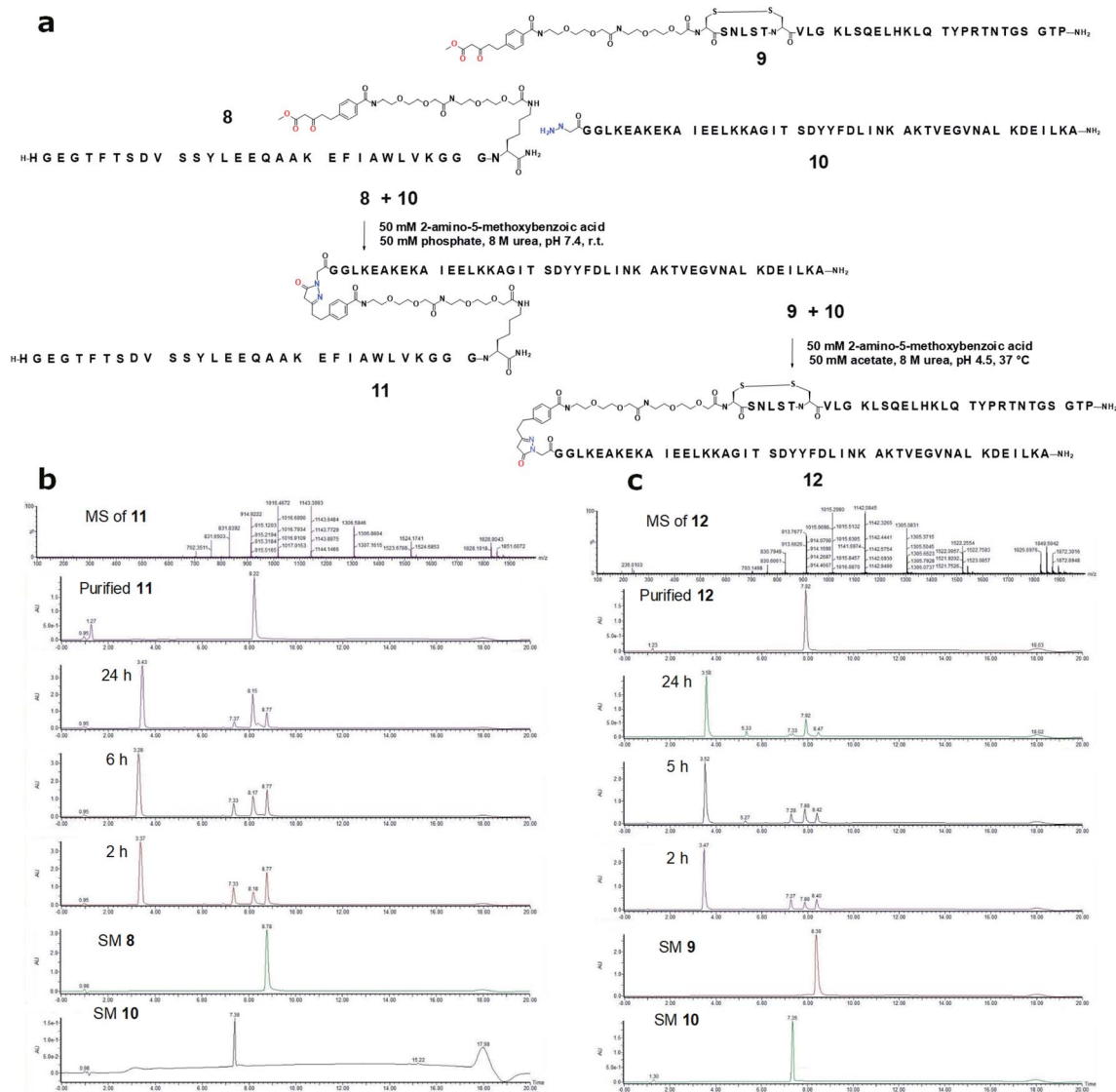


Fig. 3 Pyrazolone ligation. (a) Structures of GLP-1 analog **8**, sCT analog **9**, and ABDcon **10**. (b) UPLC trace and MS of the pyrazolone ligation between GLP-1 **8** and ABDcon **10**. (c) UPLC trace and MS of the pyrazolone ligation between sCT analog **9** and ABDcon **10**.

ABDcon peptide **10** were completed smoothly by applying the standard Fmoc/*t*Bu-based solid phase peptide synthesis (SPPS). The pre-made β -ketoester building block 4-(5-methoxy-3,5-dioxopentyl)benzoic acid **8** (Fig. S12 and S13†) was incorporated to the GLP-1 sequence as the last on-resin coupling step following the Mtt removal from C-terminal Lys side chain and the subsequent couplings of two mini-PEG linker (8-amino-3,6-dioxaoctanoic acid, or "OEG"). Similarly, the hydrazine was also introduced as the final step during the synthesis of ABDcon **10** by treating the resin-bound N-terminal bromoacetylated peptide with hydrazine in DMSO. The standard resin cleavage condition (TFA/H₂O for **8** and TFA/TIS/H₂O for **10**) and reverse-phase (RP)-HPLC purification afforded both peptides in good yield and excellent purity (Fig. S17 and S22†).

The pyrazolone ligation condition between peptides **8** and **10** was screened against various parameters including pH (4.5, 7.4),⁵ temperature (r.t., 30, 40 °C),⁵ catalyst (2-amino-5-

methoxybenzoic acid, aniline, 3,5-diaminobenzoic acid, at 10 eq. and 50 mM),²¹ urea (0–8 M), NaCl (0, 0.15, 3 M),²² and buffer salts (acetate, phosphate, *C*-(1*H*-imidazol-2-yl)-methylamine, *N,N*-dimethylethylenediamine, *p*-phenylenediamine) (Table S3†).²³ It was found that both 50 mM catalyst 2-amino-5-methoxybenzoic acid and 8 M urea were critical for high conversion efficiency. Under such conditions, the impact of pH on reaction rate was not significant, and the ligation achieved around 80% completion after 6 h at 40 °C, or 1 day at 30 °C, or 2 days at r.t. (Table S4†). In a modest scale-up ligation reaction, peptides **8** and **10** were mixed at 1 : 1 molar ratio with a final concentration of 800 μ M of each in the pH 7.4 buffer with 8 M urea and 50 mM catalyst 2-amino-5-methoxybenzoic acid. 20 mg of the ligated product GLP-1-ABDcon **11** was obtained with an isolation yield of 55% after 48 h agitation at r.t. and subsequent RP-HPLC purification (Fig. 3a and b).



The pyrazolone ligation was next assessed on constructing an ABDcon–salmon calcitonin (sCT) conjugate (Fig. 3a and c). Differently from GLP-1, sCT bears one disulfide and the conjugation site needs to be at its N-terminus where the modification doesn't compromise its biological activities.²⁴ The ligation of peptides **9** and **10** was conducted by mixing both peptides at 500 μ M final concentration in pH 4.5 buffer (50 mM acetate) in the presence of 8 M urea and 50 mM 2-amino-5-methoxybenzoic acid overnight at 37 °C. 11.8 mg of ABDcon–sCT conjugate **12** was obtained after RP-HPLC purification with an isolation yield of 64%.

With two successful case studies, we chemically stressed the ligation products **11** and **12** to evaluate the stability of the newly-formed pyrazolone junction. We monitored the UPLC traces of both GLP-1-ABDcon and ABDcon–sCT conjugates in pH 4.5 and 7.5 buffers at 4, 25 and 37 °C over 7 days (Fig. S4 and S5†). ABDcon–sCT **12** demonstrated highly favorable stability profiles at all conditions with a negligible amount of degraded products detected. In contrast, despite being stable at 4 °C, a significant amount of degradation was observed for GLP-1-ABDcon **11**, in particular at pH 4.5, 37 °C. The strikingly different stability suggested the observed degradation of GLP-1-ABDcon took place primarily on the peptide sequence rather than at the pyrazolone junction. These case studies collectively established the pyrazolone ligation as an attractive new bioconjugation tool.

Pyrazolone-mediated sequential conjugations

It was reported decades ago that pyrazolone can react with the aldehyde group through Knoevenagel condensation forming a Michael addition acceptor that however can react with another pyrazolone at a significantly faster rate than the first step, forming a single product consisting of one aldehyde and two pyrazolone units (Fig. S6†).²⁵ By treating the carbohydrates with phenylpyrazolone, this chemistry has provided a useful method for determining the carbohydrate composition of natural polysaccharides or glycoproteins.²⁵ More recently, the application of such reaction has been extended to the preparation of antibody drug conjugates.¹⁶ In one case, a pre-made pyrazolone-tagged cytotoxic agent was loaded to a Her2 antibody, bearing two aldehyde groups providing a drug to antibody ratio of 4 to 1.^{16b} To realize our goal of assembling multiple biomolecules into a single construct, the reactivity between pyrazolone and aldehyde offers promise. However, the resulting construct bearing two pyrazolone segments is often undesired for constructing multiple functional therapeutic biomolecules. To address this, we proposed to trap the highly reactive Knoevenagel condensation product by an external thiol-containing biomolecule through an extremely fast thiol 1,4-addition. Such strategy not only prevents further reaction with a second pyrazolone segment post initial Knoevenagel condensation, but also allows us to incorporate a fourth biomolecule segment into the product, realizing a true sequential bioconjugation process.

We first chose formaldehyde, and a short peptide Cholecystokinin 26-33 (CCK-8)²⁶ which was extended by two OEG and one acetylated Cys residue at the N-terminus (peptide **14**), to test the sequential conjugation. A slightly molar excess amount of

formaldehyde (final concentration: 414 μ M) and CCK-8 thiol **14** (290 μ M) were mixed with GLP-1-ABDcon **11** (276 μ M) in a phosphate buffer at neutral pH and r.t. It was found the Knoevenagel condensation between GLP-1-ABDcon **11** and formaldehyde and the subsequent Michael addition with CCK8 thiol **14** proceeded cleanly with fast kinetics. The tandem process was completed within 60 min and pyrazolone peptide **11** was quantitatively converted to a trifunctional construct **15** consisting of albumin binding domain, sCT and CCK8 with an isolation yield of 65% (9.5 mg) (Fig. 4a and b).

We next substituted formaldehyde with an aldehyde-containing GLP-1 derivative **13** and re-assessed the sequential conjugation by using ABDcon–sCT conjugate **12** and CCK8 thiol **14**. The final concentration of peptides **12**, **13**, and **14** was 130, 136 and 136 μ M respectively. It was found that the transformation of peptides **12**, **13** and **14** took place equally efficiently to the one of peptides **11**, **14** and formaldehyde. The reaction was completed within 60 min providing a quadri-functional construct **16** consisting of albumin binding domain, GLP-1, sCT and CCK8 with an isolation yield of 60% (5.8 mg) (Fig. 4a and c).

These two applications established an unprecedented sequential conjugation procedure involving pyrazolone ligation, Knoevenagel condensation and Michael addition. It demonstrated high efficiency in assembling multiple building segments, each possessing a unique pharmacological property, into a trifunctional or quadri-functional construct.

Pyrazolone-mediated versatile macrocyclization

The high efficiency of pyrazolone-centered intermolecular sequential conjugation inspired us to further explore its application in intramolecular reactions. Such intramolecular processes could offer versatile methods to quickly access novel macrocyclic peptide scaffolds, a molecular format that is often highlighted by its antibody-mimicking potency and significantly improved chemical, biophysical and enzymatic stabilities.

We started testing the intramolecular pyrazolone ligation by placing the β -ketoester and hydrazine groups at the side chain of residues Lys17 and Lys24 of a GLP-1 analog. They were incorporated to the resin-bound peptide through Lys(ivDde) and Lys(Mtt) chemistries respectively. Immediately after resin-cleavage by TFA/H₂O, it was found that the major crude product was the desired GLP-1 analog with Lys17–Lys24 side chain crosslinked by a pyrazolone motif, peptide **19** (Fig. 5a and S25†). Such result suggested the intramolecular pyrazolone ligation took place concurrently during the resin-cleavage step. The kinetics were extremely fast, which was in stark contrast to the intermolecular ligation where the reaction rate was significantly slower. This GLP-1 case study establishes the intramolecular pyrazolone ligation as an effective alternative method for peptide crosslinking, including α -helix stabilization through side-chain stapling.

Uniquely different from the well-established alpha-helix stapling protocols, the pyrazolone ligation-mediated stapling offers the opportunity of post-stapling modification of the



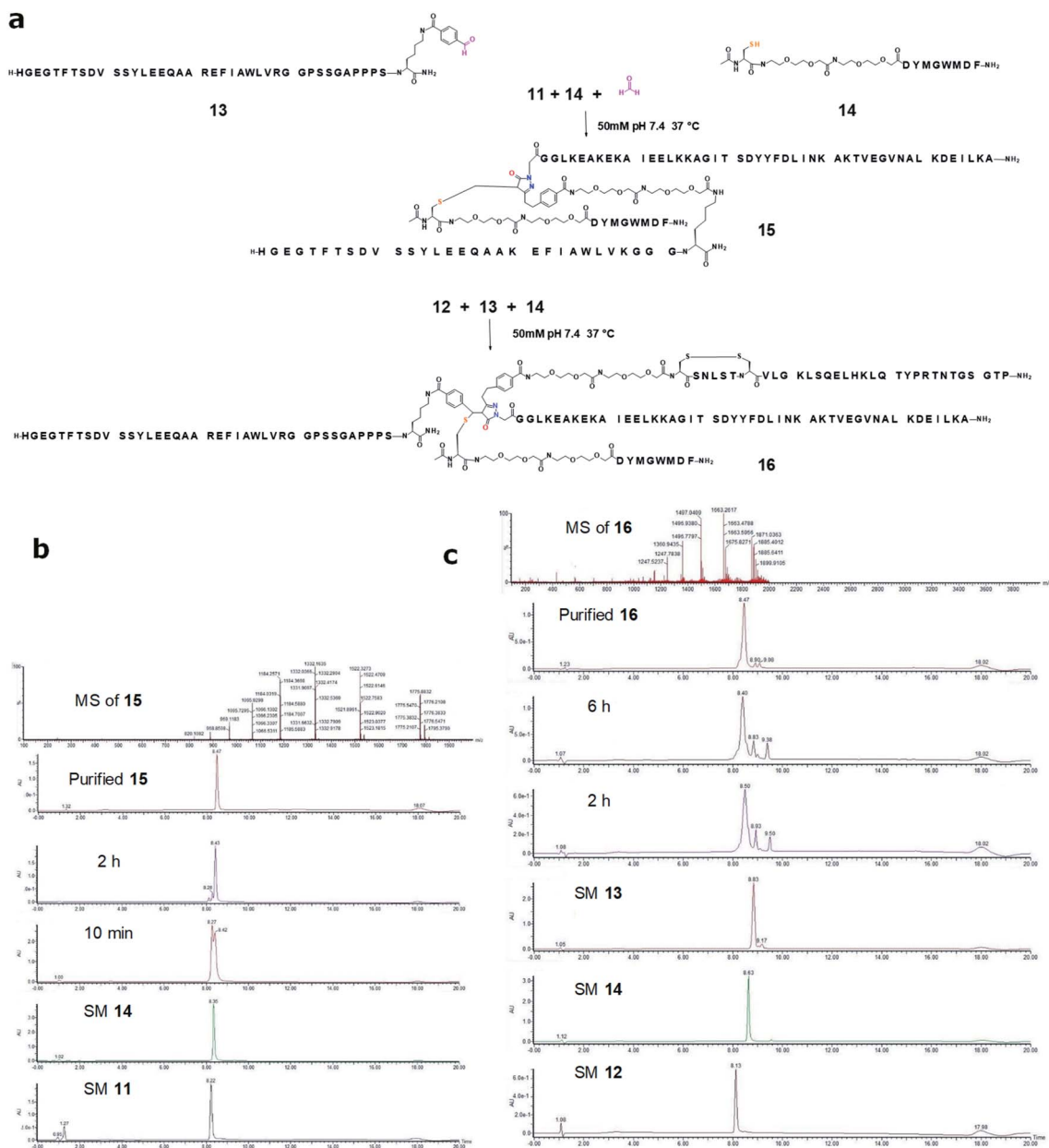


Fig. 4 Pyrazolone-mediated sequential ligation. (a) Reaction schemes. (b) UPLC trace and MS of the reaction among GLP-1-ABDcon **11**, formaldehyde and a Cys-bearing CCK8 **14**. (c) UPLC trace and MS of the reaction among ABDcon-sCT **12**, an aldehyde-bearing GLP-1 analog **13**, and a Cys-bearing CCK8 **14**.

linker, a feature that has not been reported previously. We first treated the pyrazolone stapled GLP-1 analog **19** with formaldehyde and an albumin binder palmitic acid motif **17** in phosphate buffer at neutral pH and 37 °C. Palmitic acid peptide **17** was decorated by a free thiol *via* a solubility-enhancing spacer. The final concentration of **17**, **19** and formaldehyde were 316 μM, 300 μM and 450 μM respectively. The reaction proceeded smoothly and completed within 30 min, offering a palmitoylated protracted stapled GLP-analog **S-3** with an isolation yield of 70% (5.6 mg) (Fig. S7†). We secondly attempted to introduce a second therapeutic segment in addition to the time-action modulating palmitoyl segment. The conjugation was

conducted by mixing the pyrazolone stapled GLP-1 **19** (final concentration: 500 μM), a CCK-8 peptide **18** (550 μM) bearing an aldehyde group and the palmitoyl segment **17** (550 μM) under the same condition. Similarly, the reaction proceeded cleanly and was completed within 60 min providing a palmitoylated dual agonist **20** consisting of CCK-8 and stapled GLP-1 with an isolation yield of 62% (6.3 mg) (Fig. 5a and d). The bis-pyrazolone product **S-4** was observed as a minor side product in this reaction (Fig. S8†).

We then explored the pyrazolone-mediated poly-macrocyclization. We first introduced a Cys residue to the C-terminus of pyrazolone-stapled GLP-1 **19**. It was confirmed



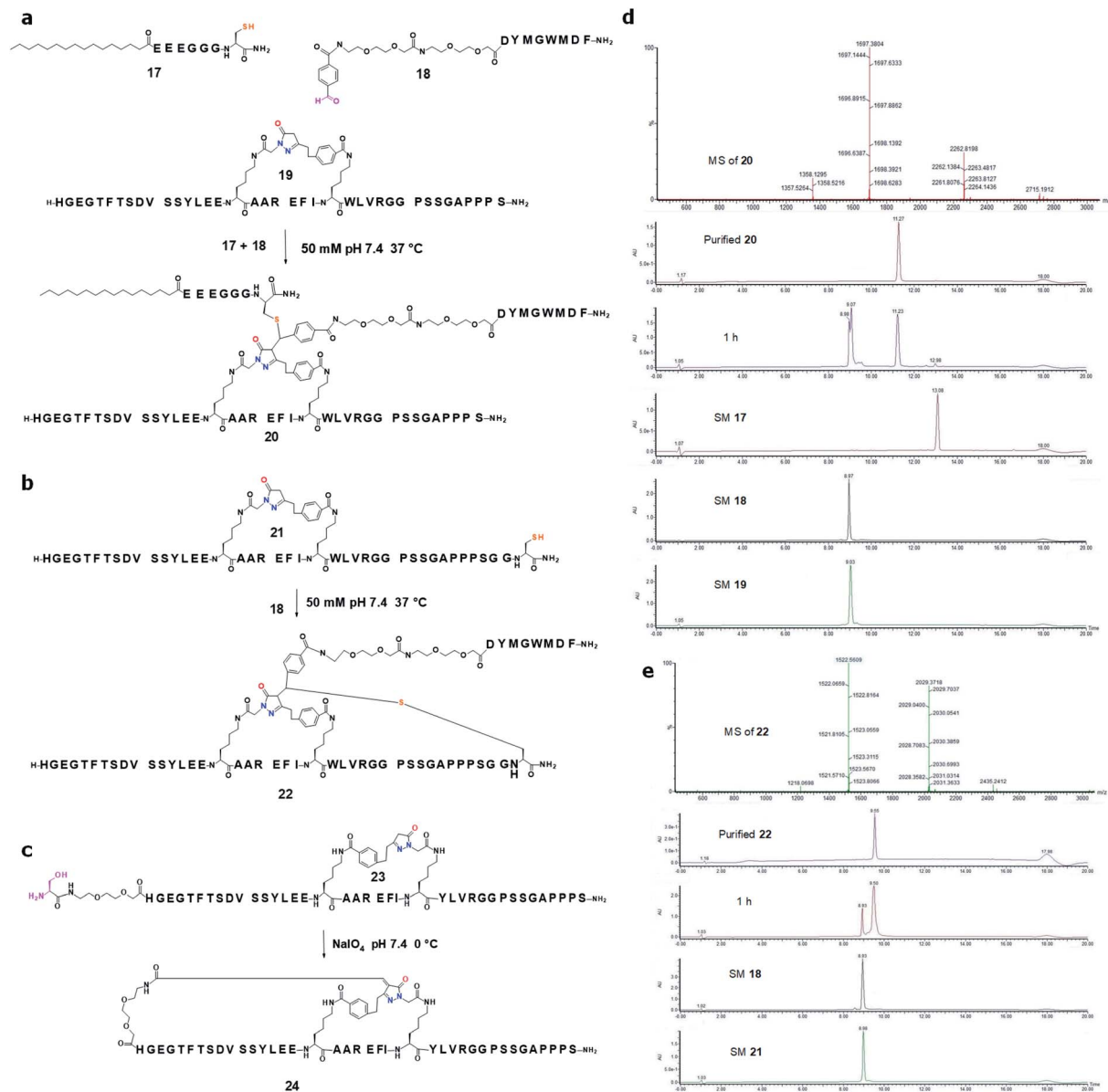


Fig. 5 Pyrazolone-mediated macrocyclization. (a) Reaction scheme of modification of pyrazolone-stapled GLP-1 peptides **19**. (b) Reaction scheme of constructing bicyclic peptides through pyrazolone and an internal Cys residue. (c) Reaction scheme of constructing bicyclic peptides through pyrazolone and an internal aldehyde group. (d) UPLC traces and MS of reaction scheme (a). (e) UPLC traces and MS of reaction scheme (b).

that the free thiol was fully compatible with the intramolecular pyrazolone ligation process affording the desired product **21** (Fig. S26[†]). Upon treatment with a formaldehyde (final concentration: 44 μ M) in neutral phosphate buffer at 37 °C, the C-terminal region of **21** (29 μ M) was cyclized to the pyrazolone core through a tandem process involving intermolecular Knoevenagel condensation–intramolecular Michael addition producing a bicyclic peptide **5-S** with an isolation yield of 72% (5 mg) (Fig. S9[†]). The bis-pyrazolone product was observed when this reaction was run at the peptide concentration of 300 μ M. When the aldehyde-bearing CCK8 **18** was used instead of formaldehyde, a similar condition offered a CCK8-conjugated bicyclic GLP1 analog **22** with an isolation yield of 66% (6.2

mg) (Fig. 5b and e). Interestingly, this ligation was set up with both peptides at 330 μ M while not observing any formation of the bis-pyrazolone product. In both bioconjugation reactions, the cyclization proceeded cleanly and achieved full conversion within 60 min.

We next introduced an aldehyde group to the N-terminus of a pyrazolone-stapled GLP-1 analog. A Ser residue was installed at the N-terminus of the GLP analog *via* one OEG linker resulting in peptide **23** (Fig. S27[†]), which was found to be smoothly converted to 2-oxoacetamide by the treatment with 1.5 eq. sodium periodate (NaIO₄) at 0 °C.²⁷ Within one pot, the newly generated aldehyde group immediately cyclized onto the pyrazolone core through an intramolecular Knoevenagel reaction generating



bicyclic peptide **24** with an isolation yield of 54% (2.7 mg) (Fig. 5c and S10†). We initially tested this conversion on a peptide with the Ser residue attached directly to the N-terminal His of GLP-1 analog **19**, despite the NaIO₄ treatment proceeded efficiently, the cyclization was very slow, presumably due to the unfavorable conformational constraint during this macrocyclization. We also introduced a Cys residue to the C-terminus of peptide **23** (peptide **S-6**) (Fig. S28†) aiming to construct a tricyclic peptide. We expect that once the N-terminal Ser is converted to the 2-oxoacetamide group, the formation of the second macrocycle will take place *via* an intramolecular Knoevenagel reaction followed by the third cyclization through an intramolecular thiol Michael addition. After the treatment of NaIO₄, the product with mass corresponding to the sulfone form of the desired tricyclic peptide product was observed in the reaction mixture; however, the majority of the products were found to be associated with the direct oxidation of the C-terminal Cys thiol group (Fig. S11†). Alternative design of a synthetic scheme to introduce both aldehyde and thiol groups into the pyrazolone-stapled peptide are required to achieve higher quality synthesis of tricyclic peptides.

Conclusion

We have reported an unprecedented sequential bioconjugation strategy that can quickly assemble four individual segments into a quadri-functional single construct. When applied intramolecularly, it serves as a unique and versatile peptide polymacrocyclization technology. Furthermore, the combined employment of intermolecular conjugation and intramolecular macrocyclization offers a set of diverse biomolecule scaffolds. The process starts with the pyrazolone ligation between a β -ketoester and a hydrazine, followed by Knoevenagel condensation with an aldehyde and subsequent Michael addition with a thiol. Although the intermolecular pyrazolone ligation requires catalysts to enhance the reaction rate, the intramolecular pyrazolone macrocyclization can be completed instantly. The following Knoevenagel condensation and Michael addition both possess fast kinetics producing the respective products within 60 min. We envision such new bioconjugation strategy will provide quick accesses to polyfunctional biomolecule constructs enabling quick proof of concept assessment of various polypharmacy hypothesis. We hope the pyrazolone-mediated sequential conjugation chemistry presented here will serve the first demonstration and spark more innovative method development in this emerging multiple-component bioconjugation field.

Conflicts of interest

F. L. is currently a full-time employee and shareholder of Novo Nordisk.

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