Kinetic Controlled Tag-Catcher Interactions for Directed Covalent Protein Assembly

Lee Ling Tan, Shawn S. Hoon, Fong T. Wong*

Molecular Engineering Lab, Biomedical Sciences Institutes, Biopolis Drive, Singapore, Singapore

* wongft@bmsi.a-star.edu.sg

Abstract

Over the last few years, a number of different protein assembly strategies have been developed, greatly expanding the toolbox for controlling macromolecular assembly. One of the most promising developments is a rapid protein ligation approach using a short polypeptide SpyTag and its partner, SpyCatcher derived from Streptococcus pyogenes fibronectin-binding protein, FbaB. To extend this technology, we have engineered and characterized a new Tag-Catcher pair from a related fibronectin-binding protein in Streptococcus dysgalactiae. The polypeptide Tag, named SdyTag, was constructed based on the native Cna protein B-type (CnaB) domain and was found to be highly unreactive to SpyCatcher. SpyCatcher has 320-fold specificity for its native SpyTag compared to SdyTag. Similarly, SdyTag has a 75-fold specificity for its optimized Catcher, named SdyCatcherDANG short, compared to SpyCatcher. These Tag-Catcher pairs were used in combination to demonstrate specific sequential assembly of tagged proteins in vitro. We also demonstrated that the in vivo generation of circularized proteins in a Tag-Catcher specific manner where specific Tags can be left unreacted for use in subsequent ligation reactions. From the success of these experiments, we foresee the application of SdyTags and SpyTags, not only, for multiplexed control of protein assembly but also for the construction of novel protein architectures.

Introduction

Protein engineering is an important tool for many industrial, therapeutic and research applications [1]. One important aspect is fusion protein construction for applications, such as protein therapeutics (antibody fusion proteins [2, 3]), biomaterial engineering (scaffold matrices and functional groups [4]) and genetic regulation (DNA binding repressors or activators [5]). Most fusion proteins are constructed as genetically encoded proteins connected by flexible, rigid or cleavable linkers. This strategy is highly dependable on soluble expression and proper folding of the fused multi-domains constructs. To overcome solubility bottlenecks, another strategy is to assemble independently expressed and properly folded proteins in vitro. This strategy employs stable non-covalent interactions, such as dimerizing linkers [6], streptavidin-biotin [7], and covalent linkages such as inteins [8] and sortases [9]. While covalent in vitro assembly
strategies bypass solubility and folding issues to form permanent linkages between proteins, some constraints of these ligation partners limit the molecular topologies that can be achieved. For example, sortases and inteins-mediated ligations can only occur between the N and C-terminus of target proteins.

Recently, a rapid covalent ligation strategy was developed by Zakeri et al [10]. This system, consisting of a peptide tag (SpyTag) and a reactive domain (SpyCatcher), was engineered by deconstructing the isopeptide forming CnaB2 domain from *S. pyogenes* [10]. SpyTag-SpyCatcher is an important addition to the protein engineering toolbox as it is robust, rapid and flexible in its placement within the protein. Applications for this protein ligation system have been numerous; these include controlled macromolecular assembly [11], protein cyclisation [12], cell labelling [13], antibody engineering [14] and hydrogel synthesis [15]. To realize the goal of assembling higher order macroassemblies from modular protein parts, multiple orthogonal interaction domains are needed. Using SpyTag and SpyCatcher along with other engineering strategies, this can be performed to a certain extent [11–15]. However, an orthogonal ligation partner, with SpyTag-like robustness, kinetics and flexibility, promises to significantly advance macromolecular assembly possibilities. In this study, we report the construction and optimization of a *S. dysgalactiae* derived Tag-Catcher pair that can be used in concert with SpyTag-SpyCatcher. The new Tag-Catcher pair will be used alongside SpyTag-SpyCatcher partners, to demonstrate kinetically controlled directed protein ligation, domain specific protein circularization and macromolecular assembly. During our preparation of this manuscript, Dr. M. Howarth at University of Oxford have also published an alternative Tag-Catcher system, SnoopTag-Catcher, that can also be utilized with SpyTag-Catcher [16].

**Materials and Methods**

**Protein expression**

*E. coli* codon-optimized DNA sequences for Tag-Catcher constructs were synthesized using gBlocks (Integrated DNA Technologies). These were cloned into pET28 vectors using restriction enzyme digestion and ligation. The final hexahistidine-tagged constructs were transformed into T7 express *E. coli* (NEB) for overnight protein expression at 16°C, 100 μM isopropyl β-D-1-thiogalactopyranoside (IPTG). The cultures were centrifuged at 10000 g for 10 minutes at 4°C. The resulting pellets were resuspended in 25 mL of 100 mM 4-(2-hydroxyethyl)-1-piperazineethanesulfonic acid (HEPES) pH 7.4, 10 mM imidazole, 150 mM sodium chloride before sonication. The resulting lysate was then centrifuged at 19000 g for 1 hour at 4°C. The decanted supernatant was incubated with Ni-NTA agarose for 1 hour at 4°C. The resin was washed with 15 mL 100 mM HEPES pH 7.4, 60 mM imidazole, 150 mM sodium chloride and the bound protein was eluted with 5 mL 100 mM HEPES pH 7.4, 150 mM imidazole, 50 mM sodium chloride. The elute was diluted two fold with 100 mM HEPES pH 7.4 before loading onto a HiTrap-Q anion exchange column for FPLC (AKTA start, GE Healthcare life sciences, buffer A: 100 mM HEPES, pH 7.4, buffer B: 100 mM HEPES, pH 7.4, 1 M sodium chloride). A linear gradient elution was used. The eluted fractions were combined and buffer exchanged into 50 mM HEPES pH 7.4, 10% glycerol for storage. Catchers were expressed at ~1 mg/L yields. Tag-EGFPs were expressed at 20–30 mg/L yields. Amino acid sequences of the constructs can be found in the S1 Table.

**In vitro reactions**

In our assay to analyze Catcher constructs, 50 μM Catcher constructs were incubated with 65 μM SdyTag-EGFP in 100 μM sodium phosphate buffer, pH 7, at 25°C. The reaction was quenched with SDS-PAGE loading buffer after 40 minutes. The samples were boiled at 95°C
for 7 minutes before loading onto a 4–12% bis-tris Invitrogen precast gel. The resulting gel was stained with SimplyBlue SafeStain (Invitrogen), then imaged and analyzed using Gel Doc EZ system (Bio-Rad). Triplicates were used to measure % yield.

To measure rates, 10 μM SpyTag-EGFP was incubated with 10 μM Catcher in 100 μM sodium phosphate buffer, pH 7, at 25°C. A portion was taken out at intervals and quenched with SDS-PAGE loading buffer. The samples were boiled at 95°C for 7 minutes before loading onto a 4–12% bis-tris Invitrogen precast gel. The resulting gel was stained with SimplyBlue SafeStain, then imaged and analyzed using Gel Doc EZ. Triplicates were performed to determine rates.

For triSdyTag and bi-SdyCatcher reaction, proteins were reacted at different concentrations and different ratios for overnight in 100 μM phosphate buffer, pH 7, at 25°C. The reactions were quenched with SDS-PAGE loading buffer and boiled at 95°C for 7 minutes before loading onto a 3–8% tris-acetate Invitrogen precast gel. Further details for the specific reactions can be found in Supporting information (S1 File).

Homology modelling

I-TASSER [17] was used to obtain a set of 5 homology models. These models were then re-assessed using the ERRAT program [18]. The model, which was chosen, had an overall quality value of 93 (percentage of the model which falls between 95% confidence level).

Mass analysis

Mass spectrometry were performed on purified SdyCatcher\textsubscript{DANG short} and purified recombinant CnaB protein (Table 1).

To determine if isopeptide bonds were formed, excess SdyTag-EGFP (100 μM) and SdyCatcher\textsubscript{DANG short} (20 μM) was incubated overnight, in 100 mM Tris buffer, pH 6.8, at 25°C, before being subjected directly to mass spectrometry (Biopolis Shared Facilities, Singapore). Mass spectra can be found in S1 Fig. ExPASy ProtParam was accessed to calculate average mass of the proteins without N terminal methionine and using subtraction of 18 Da to account for isopeptide formation reaction. N-glucosylation is accounted for by addition of 178 Da [19] and GFP maturation by subtraction of 20 Da [20].

Western blot

To examine lysate fractions for TriSdyTag after slow (16°C, 100 μM IPTG, overnight) and fast expression (37°C, 100 μM IPTG, 4 hours), the cultures were centrifuged at 10000 g for 10

| Sample | Proteins | Expected [M +H]+, Da | Observed [M +H]+, Da | Delta mass, Da |
|--------|----------|---------------------|---------------------|---------------|
| Purified recombinant CnaB domain | Wild type self-ligated CnaB domain | 16596.4 | 16596.8 | 0.4 |
| Purified SdyCatcher\textsubscript{DANG short} | SdyCatcher\textsubscript{DANG short} | 16774.4 | 16774.8 | 0.4 |
| | Glucosylated self-ligated wild type CnaB domain | 14233.6 | 14233.9 | 0.2 |
| Excess SdyTag-EGFP incubated with SdyCatcher\textsubscript{DANG short} | SdyTag—maturated EGFP | 14411.6 | 14412 | 0.3 |
| | SdyCatcher\textsubscript{DANG short} | 30633.5 | 30633.7 | 0.2 |
| | Glucosylated SdyCatcher\textsubscript{DANG short} (covalently ligated) | 44849.1 | 44849.5 | 0.4 |
| | SdyCatcher\textsubscript{DANG short} : N-acetylated SdyTag-EGFP (covalently ligated) | 45027.1 | 45026.6 | -0.5 |

doi:10.1371/journal.pone.0165074.t001
minutes at 4°C. The resulting pellets were resuspended in 25 mL of 100 mM HEPES, pH 7.4, before sonication. The lysate is then centrifuged at 19000 g for 1 hour at 4°C. The supernatant is then loaded onto SDS-PAGE. The proteins were separated by SDS-PAGE, were then transferred from the gel to nitrocellulose membrane and probed with Anti-6X His tag antibody (HRP, Abcam). The western blot analyses were visualized, according to manufacturer's protocol (SuperSignal West Pico Chemiluminescent Substrate, Cat. No. 34080, Thermo Scientific).

Results and Discussions

Identification of a novel Tag-Catcher pair

A BLAST search using *S. pyogenes* CnaB2 domain (accession code: AFD50637.1) as a query against the non-redundant protein sequences database identified a homologous CnaB protein domain from *S. dysgalactiae* with 63% percent sequence identity. This domain is located in a fibronectin-binding protein (accession code: CAA80122.1), consisting of an N-terminal collagen binding domain, repeat units of collagen binding protein domain B, a CnaB domain, fibronectin-binding repeats and a C-terminal cell anchoring motif (Fig 1A). Comparison between the *S. dysgalactiae* protein and the fibronectin-binding protein, from which SpyTag-Spy-Catcher was engineered (PrtF2, accession code: AAT38844), revealed that sequence conservation is restricted to the C-terminal region containing collagen binding protein domain B, CnaB, fibronectin binding repeats and anchor domains. Sequence alignment between the *S. dysgalactiae* CnaB domain and its *S. pyogenes* counterpart showed significant conservation in the N-terminal portion (residues 1–102, Fig 1B). Sequence identity is 68% for the latter.

![Diagram](https://example.com/figure1.png)

**Fig 1.** *S. dysgalactiae* CnaB domain analysis. (a) Graphical view of domains in *S. dysgalactiae* fibronectin-binding protein, accession code: CAA80122.1. This was not drawn to scale. (b) Sequence alignment of *S. pyogenes* and *S. dysgalactiae* CnaB domains using Clustal Omega [21] and ESPript 3.0 [22]. Structural beta-sheet elements (β1-β10) of the *S. pyogenes* protein are also indicated based on resolved *S. pyogenes* domain (PDB ID: 2X5P [23]). Residues involved in isopeptide formation are indicated with stars. The *S. dysgalactiae* CnaB domain used for alignment is located at amino acids 828–950 of the *S. dysgalactiae* fibronectin-binding protein (1117 amino acids long).

doi:10.1371/journal.pone.0165074.g001
compared to 38% for the C-terminal portion (residue 103–123). Key residues involved in iso-epitope formation were also conserved (Fig 1B, indicated with stars). Following the annotated boundaries of PrtF2 (accession code: CAA80122.1), a deconstructed standalone \textit{S. dysgalactiae} CnaB domain was expressed in T7 Express \textit{Escherichia coli} (New England Biolabs, NEB) with an N-terminal his-tag. Using electrospay ionization-mass spectrometry (ESI-MS) for mass determination of the purified construct, we observed masses of 16596.8 and 16774.8 Da (Table 1). These correspond, within error, to the expected average masses of a self-ligated CnaB domain with and without spontaneous $\alpha$-N-6 gluconoylation of the N-terminal His-tag [19], 16596.4 Da and 16774.4 Da respectively. Although this demonstrates \textit{in vivo} intra-ligation of the recombinant CnaB domain, we cannot conclude it will do the same in the native full length fibronectin binding protein. However, the intra-ligated recombinant protein indicates that a Tag-Catcher system could be engineered from the \textit{S. dysgalactiae} CnaB domain.

**Catcher construction**

Drawing from previous SpyCatcher structure function studies [10, 24], we dissected \textit{S. dysgalactiae} CnaB domain into a Tag-Catcher pair, henceforth termed SdyTag (encoding DPIV-MIDNDKPT) and SdyCatcher respectively. To examine and optimize SdyTag-SdyCatcher interactions, we constructed and analyzed a panel of SdyCatcher variants with SdyTag. I-TASSER [17] homology modeling of the \textit{S. dysgalactiae} domain yielded a model with root mean square deviation of 0.57 Å (405 atoms) to the resolved SpyTag-SpyCatcher structure (PDB: 4MLI [24]) (Fig 2A). Structural and sequence alignments of SdyCatcher with SpyCatcher were then utilized to design SdyCatcher variants (Figs 1B, 2A and 2B).

From \textit{in silico} analyses, two main variants of SdyCatcher were designed (Fig 2B). First, we hypothesized that Ile34 (on native SdyCatcher) could interact with Glu119 (SdyTag) and thus mutations at residue position 34 (Catcher domain) could affect ligation efficiencies (Fig 2A inset). To investigate this, Catcher constructs with mutations Ile34Glu (DENG constructs) and Ile34Ala (DANG constructs) were generated, together with wild type constructs (DING constructs). DENG SdyCatcher constructs were found least efficient compared to DING and DANG constructs. DANG constructs were the most efficient (Fig 2A–2C). Yields of the short versions of DENG, DING (wild type) and DANG SdyCatcher constructs were 23, 30 and 73% respectively. These observations are likely due to the predicted electrostatic repulsion between Glu34 on DENG Catchers and Glu119 on SdyTag (Fig 2A). Isoleucine has also been suggested to inhibit Tag docking efficiencies due to its bulkier size, compared to alanine [10]. The Sdy-Catcher observations, where Glu substitution produced the least efficient Catcher, also differ from the previous SpyCatcher construction where an Ile34Glu mutation was shown to improve efficiencies with the Tyr119 containing SpyTag [10].

The second set of modifications consisted of removing 6 C-terminal amino acids from the Catcher construct (β9, Fig 1B). Our initial dissection point of \textit{S. dysgalactiae} CnaB domain to generate the SdyTag-SdyCatcher partners, was between two aspartic acids; Asp110 (C terminal residue of SdyCatcher) and Asp111 (N terminal residue of SdyTag). To determine if this Asp (Catcher)-Asp (Tag) interaction would affect reaction rates unfavorably, C-terminal truncated constructs were made to remove the C-terminal Asp110 (long and short constructs, Fig 2B). Structural analyses of the SpyCatcher-SpyTag complex and experimental studies of C-terminus truncated SpyCatchers suggest that C-terminal portion of SpyCatcher can be removed without significant change in SpyTag ligation activity [24]. In contrast to SpyCatcher [10, 24], removal of SdyCatcher’s C-terminal portion increased ligation yields significantly (Fig 2B and 2C). Yields of DANG short and long SdyCatcher constructs were 73 and 26% respectively. Whilst the C-terminal portion of SpyCatcher was predicted to have minimal effect on Tag-Catcher
ligation efficiencies [24], significant differences in yields between long and short SdyCatcher constructs indicate that the C-terminal portion of the SdyCatcher is important in Tag-Catcher ligation kinetics, possibly via inhibition of SdyTag placement onto the Catcher domain.

Overall, SdyCatcher_{DANG short} construct was considered the most efficient SdyCatcher construct (73% yield after 40 minutes reaction at 25˚C, pH 7, Fig 2C, Accession number: TBC) and was used for subsequent SdyCatcher experiments in this study. Covalent linkage of the ligated proteins was confirmed by ESI-MS of ligation reaction of SdyCatcher_{DANG short} with SdyTag-EGFP incubated with SdyCatcher_{DANG short}.
EGFP (Table 1). Alanine mutation of Asp117 on the SdyTag did not yield any ligated products thus confirming the critical role of this conserved residue for isopeptide bond formation (Lane 6, Fig 2D). Reaction of SdyTag-EGFP with SpyCatcher also yielded minimal products under the same conditions (7%, Fig 2C).

Tag-Catcher interactions

Based on our observations of SdyCatcher variants, we decided to investigate further Tag-Catcher interactions. Sequence alignments showed that *S. pyogenes* and *S. dysgalactiae* Catcher motifs that interact with the Tag (β2, Figs 1B and 2A) were highly similar. In contrast, despite conservation of isopeptide-forming Asp117 and hydrophobic region predicted to be necessary for Catcher interaction [24], Tags were predicted to be highly different in overall charge at pH 7 (Fig 3A). Isoelectric points (pIs) of Tags derived from *S. pyogenes* (SpyTag) and *S. dysgalactiae* (SdyTag) are predicted to be 9.7 and 3.4 respectively [26]. This difference is due to substitution of neutral residues on SpyTag with negatively charged ones on SdyTag (Ala and Tyr to Asp) and positively charged residues on SpyTag with neutral residues on SdyTag (His and Lys to Pro and Tyr respectively). To probe this further, we examined ligation of SpyTag-EGFP with SpyCatcher or with SdyCatcher_{DANG short} under various pH conditions, ranging from pH 5.2–9 (Fig 3B). Similar experiments was also performed with SdyTag-EGFP with both Catchers (Fig 3C). There are minimal changes in specificities of SpyTag for SpyCatcher versus SdyCatcher_{DANG short} with pH. SpyTag ligations were faster with lower pH (Fig 3B), similar to observations in previous SpyTag-Maltose binding protein experiments [10]. Likewise, Sdy-Tag ligation reactions were faster at lower pH. From pH 7 to 6.8, an observed doubling of yield for SdyTag-EGFP with SdyCatcher_{DANG short} was estimated based on densitometry (40% to 80% with respect to SdyTag-EGFP, Fig 3C). Catcher specificities for SdyTag-EGFP, where the Sdy-Catcher ligation is significantly faster compared to SpyCatcher, were also maintained up to pH 6.3. At pH 5.2, differences in yields of SdyTag-EGFP for SdyCatcher_{DANG short} and SpyCatcher were minimal (Fig 3C). This suggest that at sufficiently low pHs, interactions of SdyTag with Catchers has minimal effect on the overall ligation reaction and ligation becomes dependent on rate of isopeptide formation between the highly conserved catalytic residues in the Tag-Catcher combinations.

Specificities of Tag-Catchers

From our observations of SdyTag with SpyCatcher (Figs 2C and 3C), we predicted that the two Tags can be utilized in combination under kinetic control. To quantify the kinetic preferences of the various Tags and Catchers, we calculated and compared second order rate constants of both SpyCatcher and SdyCatcher_{DANG short} with their non-native and native Tags fused to EGFP (Table 2). Rate constants were calculated from Catcher depletion measurements during equimolar ligation of the respective Catcher and Tag at 25°C, pH 7 (See S2–S4 Figs). The most significant observation is the 320-fold preference of SpyCatcher for its native SpyTag over SdyTag. SdyTag also has a 75-fold specificity for SdyCatcher versus SpyCatcher while SpyTag has minimal preference for SpyCatcher and SdyCatcher. Our observations of the reactivity of Catcher variants (Fig 2) in conjunction with previous studies on C-terminal truncated SpyCatcher [24], explains why SdyCatcher_{DANG short} ligate almost as efficiently with SpyTag compared to SpyCatcher. SpyTag ligation has been shown earlier to be independent of C-terminal truncations of its native Catcher [24], thus SpyTag interactions is not inhibited by both Catchers. Glu34 on the SpyCatcher would also be able to accommodate Tyr119 (SpyTag) but not Asp119 (SdyTag), while Ala34 (SdyCatcher) would be able to accommodate both Tags.
Kinetic controlled Tag-Catcher ligations for directed assembly

Comparison of rate constants between Catcher-Tag partners (Table 2) suggests that directed ligation schemes can be generated under two conditions. First, the 320-fold preference of Spy-Catcher for its native SpyTag over SdyTag can be used to select for SpyTag in a mixture of Tags.
(Fig 4A and 4B). Second, 75-fold preference of SdyTag with SdyCatcherDANG short over SpyCatcher can be utilized to select for SdyCatcherDANG short in a mixture of Catchers (Fig 4C and 4D). Directed protein assembly is also performed with the considerations that SdyCatcher does not demonstrate preference for SdyTag over SpyTag and that SpyTag can also react almost as efficiently with both Catchers.

To apply the Tags for directed protein ligation, dual Tagged or dual Catcher proteins were constructed. A dual SpyTag-EGFP-SdyTag (Fig 4A), along with its SpyTag only equivalent (SpyTag-EGFP-SpyTag, Fig 4B) were designed and expressed. Similarly, a dual Catcher, consisting of SpyCatcher-SdyCatcherDANG short (Fig 4C), and its SpyCatcher only equivalent (SpyCatcher-SpyCatcher, Fig 4D) were constructed and expressed.

To evaluate selectivity of each Catcher, the dual-Tag construct was used to ligate specific Catcher constructs in a step-wise in vitro reaction (Fig 4E). We first incubate the dual-Tag with SpyCatcher to obtain the specifically ligated SpyCatcher:SpyTag-EGFP-SdyTag. As predicted, with the high preference of SpyCatcher for SpyTag over SdyTag, we observed only one major product with the incubation of SpyCatcher with dual-Tag (lanes 1–3, Fig 4E). This product is predicted to be SpyCatcher:SpyTag-EGFP-SdyTag, where SdyTag is unreacted. Under these conditions, there is also little cross-reactivity between Tag-Catcher partners as we observe minimal double SpyCatcher ligated product (SpyCatcher:SpyTag-EGFP-SdyTag:SpyCatcher). Based on densitometry, less than 10% of the products were ligated non-specifically twice (with a protein product of 60 kDa, lanes 1–2, Fig 4E). With the further addition of SdyCatcherDANG short, the double-ligated product (SpyCatcher:SpyTag-EGFP-SdyTag:SdyCatcher) is observed (lane 4, Fig 4E). In contrast, addition of SpyCatcher did not result in any change in protein profile (lane 3, Fig 4E). In the SpyTag-only equivalent, the sequential ligation reaction was not possible, where both singly and doubly ligated SpyTag-EGFP-SpyTag products were observed under the same conditions (lanes 5 and 6, Fig 4E).

Next, we demonstrate directed ligation using SpyCatcher-SdyCatcherDANG short construct (Fig 4F). Upon incubation of SpyCatcher-SdyCatcherDANG short with SdyTag-EGFP, we observed the appearance of a major protein band corresponding to a singly ligated product (60 kDa, lanes 5 and 6, Fig 4F). From the rate constants, this is predicted to be SpyCatcher-SdyCatcherDANG short-SdyTag-EGFP. Based on densitometry, <10% of the products were ligated non-specifically twice (90 kDa, lanes 5 and 6, Fig 4F). Only the further addition of SpyTag-EGFP, produced the predicted SpyTag-EGFP:SdyCatcherDANG short:SdyTag-EGFP (90 kDa, lane 7, Fig 4F) after 10 minutes. In comparison, minimal product was observed when SpyCatcher-SpyCatcher was reacted with SdyTag-EGFP after 40 minutes under the same conditions (Lane 8, Fig 4F). Together, these results demonstrate that SdyTag-EGFP can be directed onto SpyCatcher- SdyCatcherDANG short to produce SpyCatcher- SdyCatcherDANG short-SdyTag-EGFP, which can be, in turn, ligated further with SpyTag-EGFP.

Table 2. Reaction rate constants between different ligation partners.

| Catcher Tag | SdyTag | SpyTag |
|-------------|--------|--------|
| Reaction with SpyCatcher | | |
| SpyCatcher  | 0.6    | 193    |
| SdyCatcherDANG short | 45     | 4      |

Triplicate measurements were used to calculate average and standard deviations, (STD).

doi:10.1371/journal.pone.0165074.t002
Utilizing the selectivities of SpyCatcher and SdyTag, we have demonstrated that under kinetic control, one-pot directed protein ligation is possible. Although, our current assembly was accomplished with EGFP fusions, we do not foresee any difficulty fusing other proteins onto the Tag-Catchers as shown from previous studies [11, 12, 15]. To further enhance yields, solid phase addition, as demonstrated by SnoopTag [16], or step-wise purifications could also be utilized.

Construction and polymerization of circular proteins

Next, we also demonstrated utility of the Tag-Catcher pairs for macromolecular control. Although, circular proteins have been constructed previously using SpyTag-SpyCatcher [11, 12], development
of the SdyTag presents us with an opportunity to generate specific internally-circularized proteins with specific free Tags to be used for further protein assembly possibilities.

From the observed rate constants (Table 2) and kinetic controlled Tag-Catcher assembly results (Fig 4), SpyCatcher has demonstrated preference for its native SpyTag in the presence of both SpyTag and SdyTag in vitro. We hypothesize that the same selection should occur in vivo. To demonstrate this, we designed a multi-Tag construct. This multi-Tag construct consist of 3 SdyTags, 1 SpyTag and 1 SpyCatcher, arranged as SdyTag-SpyTag-SdyTag-SdyTag-gtobacco etch virus (TEV) protease site-SpyCatcher (triSdyTag, Fig 5A). We predicted that Spy-Tag-SpyCatcher ligation would proceed rapidly compared to SdyTag-SpyCatcher ligation, subsequently, in situ intra-polypeptide ligation, under slow expression conditions, is expected to mainly produce the SpyTag-SpyCatcher ligated circularized protein (Fig 5A).

The triSdyTag construct were expressed in T7 Express E. coli at 100 μM IPTG under 16°C overnight (slow expression) or 37°C for 4 hours (fast expression). The cell pellets were lysed and their protein profiles were examined by Western Blot analysis with an anti-6xHistag antibody (HRP, Abcam) (Fig 5B). Not surprisingly, multi-mer proteins were observed at 37°C whilst the 16°C expression yielded a primary protein band at ~20 kDa (77% of the lysate based on densitometry). TriSdyTag is predicted to have a mass of 22 kDa.

Using Ni-NTA purification and followed by fast protein liquid chromatography (FPLC), ~20 kDa sized proteins were purified from 16°C expression of triSdyTag (lane 1, Fig 5C). Circularization of the triSdyTag product was confirmed with TEV protease digestion, where a single band of lower mobility was observed after overnight digestion (lanes 1 and 2, Fig 5C). We further verified presence of three accessible Tags on the triSdyTag with a maximum addition of three SdyCatchers onto tri-SdyTag (lanes 3–8, Fig 5C). To determine if the circularization was specific (SpyTag to SpyCatcher), we used SpyCatcher ligation as a means to determine the percentage of free SpyTag present in the triSdyTag. TriSdyTag had no observable product with SpyCatcher (lane 3, Fig 5D). This was similar to the reaction of SdyTag-EGFP with SpyCatcher, which had no product under the same conditions (lane 2, Fig 5D). Thus, this suggests that there are minimal free SpyTag in the circular protein and subsequently, also verifies that the in situ protein circularization took place specifically via SpyTag and SpyCatcher. In the presence of a 3:1 SdyTag: SpyTag population and under slow expression conditions, we have demonstrated in vivo selectivity of the SpyCatcher for its native Tag and the construction of a circularized protein with accessible internal and N-terminal SdyTags.

Besides ligating the circular triSdyTag with SdyCatcher fusions, a triSdyTag could be utilized to form a protein network, similar to the Spy hydrogel network created using tetraSpyTag with double SpyCatcher containing constructs [15]. In this assembly, the network would be formed between circular proteins instead of linear Tag-containing proteins. A bi-SdyCatcher “connector” was constructed (SdyCatcherDANG short—SdyCatcherDANG short Fig 5A) and added to the triSdyTag in vitro (Fig 5E). With excess TriSdyTag, an additional set of N-mers were observed compared to excess or equimolar of bi-SdyCatchers (Fig 5E). To determine the maximum size of the protein assembly, we also ran the 2:1 Catcher:Tag reaction (lane 1, Fig 5E) on a size exclusion chromatography column, Tosoh TSKgel G3000SWxl, which has a calibrated range of 10–500 kDa for globular proteins (S5 Fig). The maximum size observed was > 500 kDa.

Conclusions

In this study, we engineered and characterized a new protein ligation Tag-Catcher pair, SdyTag-SdyCatcher. Like SpyTag-Catcher, protein ligation occurs via the formation of an isopeptide bond. We showed that SdyTag-SdyCatcher ligations can also occur between N, C-
Fig 5. Construction and polymerization of circular proteins. (a) Schematics for in vivo and in vitro ligations of triSdyTag constructs. (b) Western blot showing E. coli lysates for the production of triSdyTag with 100 μM IPTG, 16°C (lane 2) and 37°C (lane 1). (c) Characterization of triSdyTag. Lane 1: Purified triSdyTag, lane 2: purified triSdyTag after TEV digestion overnight, lanes 3–7: reaction of SdyCat with triSdyTag at 1, 5, 20, 90 minutes and overnight at 25°C, pH 7 and lane 8: Novex Sharp ladder. (d) Presence of free SpyTag was examined by reaction of SpyCatcher with either SpyTag-EGFP (lane 1), SdyTag-EGFP (lane 2) or circularized triSdyTag (lane 3) at 5 minutes, pH 7, 25°C. Only SpyTag-EGFP (lane 1) has a product (SpyTag-EGFP:SpyCatcher) with SpyCatcher under these conditions. (e) Polymerization of tri-SdyTag. Lanes 1–3: Reaction of bi-SdyCatcher with triSdyTag at 30 μM in 2:1, 1:1, 1:4 Catter:Tag molar ratios, lane 4: Novex Sharp ladder and lanes 5–8: reaction of bi-SdyCatcher with triSdyTag at 10 μM in 4:1, 2:1, 1:1, 1:4 Catter:Tag molar ratios. The reactions were left overnight at 25°C, pH 7.

doi:10.1371/journal.pone.0165074.g005
terminus and internal sites under various pH conditions. We took advantage of the unreactive nature of SdyTag for SpyCatcher to perform kinetic controlled directed protein assembly in vivo and in vitro. Based on analyses of EGFP-fused Tags, SpyCatcher has a 320-fold preference for its native SpyTag compared to SdyTag. SdyTag also has a 75-fold preference for SdyCatcher DANG short compared to SpyCatcher. Although our kinetics were performed using EGFP-fused Tags and might not be applicable for all fusion proteins, selectivity between Tags have also been demonstrated using a standalone multiple Tag-SpyCatcher system (Fig 5).

Using dual-Tag (SpyTag-EGFP-SdyTag) and dual-Catcher (SpyCatcher-SdyCatcher) constructs, we have demonstrated methods for directed protein ligation in vitro. Although our assembly was accomplished "linearly", we expect that SdyTag can also be used for macromolecular assembly of proteins in the many other unique architectures demonstrated previously with the SpyTag-SpyCatcher ligation strategy (e.g. star-shaped, H-arms [11]). The ability to control the order of proteins assembled would also be particularly advantageous towards assembling a specific sequence of protein domains, in particular for biosynthetic pathways [27].

Addition of the SdyTag-SdyCatcher to the protein ligation toolbox also presented an opportunity to re-examine protein circularization. Circularized proteins are advantageous for their thermal stability and resistance to degradation [28]. Previous strategies for end-to-end circularization include chemical ligation and sortases [28]. With the SpyTag-SpyCatcher alone, domain specific circularization has been demonstrated by Arnold et al [11]. Using both SdyTag and SpyTag in this study, specifically circularized proteins with free SdyTags, which can be further used for in vitro macromolecular assembly, can be expressed in vivo. In addition to the circular construct in this study, we anticipate the in vivo specificity of the Tag-Catchers would be an attractive ability in generating another dimension to un-natural protein scaffolds.

In our characterization and engineering of the SdyTag-SdyCatcher, we managed to demonstrate directed protein assembly using SdyTag and SpyTag, however, we recognize that the current capabilities of SdyTag-SdyCatcher is limited due to SdyTag’s eventual cross-reactivity to SpyCatcher. Despite this, our optimisation experiments have highlighted importance of different Tag-Catcher interactions that could be potentially used in multiplexing protein ligations.

Supporting Information

S1 Fig. Deconvoluted electrospray ionization- mass spectrometry (ESI-MS) spectra. Spectra of (a) the standalone S. dysgalactiae CnaB domain, (b) SdyCatcher DANG short alone and of (c, d) incubation of excess SdyTag-EGFP with SdyCatcher DANG short. (TIFF)

S2 Fig. Yield (%) of ligated SpyTag product with time (minutes). SpyTag-EGFP, 10 μM, is incubated with with SdyCatcher (10 μM, diamond) and SpyCatcher (10 μM, square). (TIFF)

S3 Fig. Yield (%) of ligated SdyTag product with time (minutes). SdyTag-EGFP, 50 μM, is incubated with (a) SdyCatcher (50 μM) and (b) SpyCatcher (50 μM). (TIFF)

S4 Fig. Rate constant for reactions. SpyTag-EGFP with (a) SpyCatcher and (b) SdyCatcher at 10 μM concentrations. SdyTag-EGFP with (c) SpyCatcher and (d) SdyCatcher at 50 μM concentrations. Trend line equations and coefficients are shown. (TIFF)
S5 Fig. Size exclusion column chromatography of 2:1 bi-SdyCatcher to trioSdyTag ligation reaction. The elution time for the different mass standards are also annotated. Indicated peaks have the masses: (1) >669 kDa, (2) ~300 kDa, (3) ~210 kDa, (4) ~150 kDa, (5) ~100 kDa and (6) <30 kDa.

S1 File. Methods for in vitro reactions and size exclusion chromatography.

S1 Table. Protein sequences of constructs used in the study.

Acknowledgments
We are grateful to Sydney Brenner for discussion and Pei Yu Lim and Hui Ling Yeo for their help with the western blot.

Author Contributions
Conceptualization: FTW SSH LLT.
Investigation: FTW LLT.
Methodology: FTW SSH LLT.
Project administration: FTW.
Visualization: FTW SSH LLT.
Writing – original draft: FTW SSH LLT.
Writing – review & editing: FTW SSH LLT.

References
1. Foo JL, Ching CB, Chang MW, Leong SSJ. The imminent role of protein engineering in synthetic biology. Biotech Adv. 2012; 30:541–549.
2. Carter PJ. Introduction to current and future protein therapeutics: a protein engineering perspective. Exp Cell Res. 2011; 317:1261–1269. doi: 10.1016/j.yexcr.2011.02.013 PMID: 21371474
3. Chen X, Zaro JL, Shen WC. Fusion protein linkers: property, design and functionality. Adv Drug Delivery Rev. 2013; 65:1357–1369.
4. DiMarco RL, Heilshorn SC. Multifunctional Materials through Modular Protein Engineering. Adv Mat. 2012; 24:3923–3940.
5. Bogdanove AJ, Daniel FV. TAL effectors: customizable proteins for DNA targeting. Science. 2011; 333:1843–1846. doi: 10.1126/science.1204094 PMID: 21960622
6. Ghosh I, Hamilton AD, Regan L. Antiparallel leucine zipper-directed protein reassembly: application to the green fluorescent protein. J Am Chem Soc. 2000; 122:5658–5659.
7. Laitinen OH, Nordlund HR, Hytönen VP, Kulomaa MS. Brave new (strept) avidins in biotechnology. Trends Biotech. 2007; 25:269–277.
8. Lue RY, Chen GY, Hu Y, Zhu Q, Yao SQ. Versatile protein biotinylation strategies for potential high-throughput proteomics. J Am Chem Soc. 2004; 126:055–1062.
9. Levary DA, Parthasarathy R, Boder ET, Ackerman ME. Protein-protein fusion catalyzed by sortase A. PLoS One 2011; 6(4):e18342. doi: 10.1371/journal.pone.0018342 PMID: 21494692
10. Zakeri B, Fierer JO, Celik E, Chittock EC, Schwarz-Linek U, Moy VT, et al. Peptide tag forming a rapid covalent bond to a protein, through engineering a bacterial adhesin. Proc Nat Acad Sci. 2012; 109: E690–E697. doi: 10.1073/pnas.1115485109 PMID: 22366317
1. Zhang WB, Sun F, Tirrell DA, Arnold FH. Controlling macromolecular topology with genetically encoded spytag–spycatcher chemistry. J Am Chem Soc. 2013; 135:13988–13997. doi: 10.1021/ja4076452 PMID: 23964715

2. Schoene C, Fierer JO, Bennett SP, Howarth M. SpyTag/SpyCatcher cyclization confers resilience to boiling on a mesophilic enzyme. Angew Chem Int Ed. 2014; 53:6101–6104.

3. Chen AY, Deng Z, Bilings AN, Seker UOS, Lu MY, Citorik RJ, et al. Synthesis and patterning of tunable multiscale materials with engineered cells. Nat Mat. 2014; 13:515–523.

4. Fierer JO, Veggianni C, Howarth M. SpyLigase peptide–peptide ligation polymerizes affibodies to enhance magnetic cancer cell capture. Proc Nat Acad Sci. 2014; 111:E1176–E1181. doi: 10.1073/pnas.1315776111 PMID: 24639550

5. Sun F, Zhang WB, Mahdavi A, Arnold FH, Tirrell DA. Synthesis of bioactive protein hydrogels by genetically encoded SpyTag-SpyCatcher chemistry. Proc Nat Acad Sci. 2014; 111:11269–11274. doi: 10.1073/pnas.1401291111 PMID: 25049400

6. Veggianni G, Nakamura T, Brenner MD, Gayet RV, Yan J, Robinson CV, et al. Programmable polyproteins built using twin peptide superglues. Proc Nat Acad Sci. 2016; 113:1202–1207. doi: 10.1073/pnas.1519214113 PMID: 26787909

7. Yang J, Zhang Y. I-TASSER server: new development for protein structure and function predictions; 2015. Database: I-TASSER [Internet]. Accessed: http://zhanglab.ccmb.med.umich.edu/I-TASSER/

8. Colovos C, Yeates TO. Verification of protein structures: patterns of nonbonded atomic interactions. Prot Sci. 1993; 2:1511–1519.

9. Geoghegan KF, Dixon HB, Rosner PJ, Hoth LR, Lanzetti AJ, Borzilleri KA, et al. Spontaneous o-N-6-Phosphogluconoylation of a “His Tag” in Escherichia coli: The Cause of Extra Mass of 258 or 178 Da in Fusion Proteins. Anal biochem. 1999; 267:169–184. doi: 10.1006/abio.1998.2990 PMID: 9918669

10. Niwa H, Inouye S, Hirano T, Matsuno T, Kojima S, Kubota M, et al. Chemical nature of the light emitter of the Aequorea grebe fluorescent protein. Proc Nat Acad Sci. 1996; 93:13617–13622 PMID: 8942983

11. Li W, Cowley A, Uludag M, Gur T, McWilliam H, Squizzato S, et al. The EMBL-EBI bioinformatics web and programmatic tools framework. Nucleic Acids Res. 2015; 43:W580–W584. doi: 10.1093/nar/gkv279 PMID: 25845596

12. Robert X, Gouet P. Deciphering key features in protein structures with the new ENDscript server. Nuc Acids Res. 2014; 42:W320–W324.

13. Oke M, Carter LG, Johnson KA, Liu H, McMahon SA, Kerou M, et al. The Scottish Structural Proteomics Facility: targets, methods and outputs. J Struct Funct Genomics. 2010; 11:167–180. doi: 10.1007/s10969-010-9090-y PMID: 20419351

14. Li L, Fierer JO, Rapoport TA, Howarth M. Structural analysis and optimization of the covalent association between SpyCatcher and a peptide Tag. J Mol Biol. 2014; 426:309–317. doi: 10.1016/j.jmb.2013.10.021 PMID: 24161952

15. DeLano WL. The PyMOL Molecular Graphics System, Version 1.8 Schrödinger, LLC.

16. Peptide calculator. Innovagen. 2015. Available: http://pepcalc.com/. Accessed October 5 2015.

17. Dueber JE, Wu GC, Malmirchegini GR, Moon TS, Petzold CJ, Ullal AV, et al. Synthetic protein scaffolds provide modular control over metabolic flux. Nat Biotech. 2009; 27:753–759.

18. Antos JM, Popp MW, Ernst R, Chew GL, Spooner E Ploegh HL. A straight path to circular proteins. J Biol Chem. 2009; 284:16028–16036. doi: 10.1074/jbc.M901752200 PMID: 19359246