1 SpyCombinator Assembly of Bispecific Binders

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22 ABSTRACT

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24 Bispecific antibodies are a successful and expanding therapeutic class, bridging two cell-types 25 or engaging two different molecules on the same cell. Bispecifics unlock avenues towards synergy, resistance evasion, and signaling bias. Standard approaches to generate bispecifics are 26 27 complicated by the need for disulfide reduction/oxidation or cloning of each binder molecule 28 in two different formats. Here we present a modular approach to bispecifics using 29 SpyTag/SpyCatcher spontaneous amidation, where all binders are cloned in the same format, 30 bearing a SpyTag. Two SpyTag-fused antigen-binding modules can be precisely conjugated 31 onto DoubleCatcher, a tandem SpyCatcher where the second Catcher is unreactive until unveiling of reactivity using site-specific protease. Assembly on DoubleCatcher is efficient in 32 33 phosphate-buffered saline at 37 °C, with half-times less than 5 min for both SpyCatcher arms 34 and over 97% bispecific homogeneity. We engineer a panel of DoubleCatchers, locked through disulfide bonds to direct binders to project in different directions from the hub. We establish a 35 generalized methodology for one-pot assembly and purification of bispecifics in 96-well plate 36 37 format. A panel of Fab, affibody or nanobody binders recognizing different sites on HER2 38 were coupled to DoubleCatcher, revealing unexpected combinations with anti-proliferative or pro-proliferative activity on HER2-addicted cancer cells. Bispecific activity depended 39 40 sensitively on both the order of the binders within the assembly and the geometry of 41 DoubleCatcher scaffolds. These findings support the need for straightforward assembly in 42 different formats. SpyCombinator provides a simple and scalable tool to discover synergy in 43 bispecific activity, through modulating receptor organization and geometry.

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46 Introduction

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48 Bispecific antibodies are biological matchmakers, bringing together two different target 49 molecules or two different parts of the same target (Hofmann et al., 2020). Bispecifics have 50 had clinical success in cancer therapy by directing cytotoxic T cells to cancer-specific antigens (Hofmann et al., 2020). There are also approved bispecific therapeutics that bridge proteins on 51 the surface of the same cell (Labrijn et al., 2019; Vijavaraghavan et al., 2020). Cell signaling 52 53 often naturally depends on co-localization of particular ligands at the cell-surface (Batada et al., 2006). Synthetic control of such signaling has been successful using bifunctional small 54 molecules (Zheng et al., 2022), DNA nanostructures (Shaw et al., 2014), and non-antibody 55 56 protein-based binders (Mohan et al., 2019). The response of the cell is often exquisitely 57 sensitive to the timing and spatial orientation of this bridging between surface targets (Moraga et al., 2015). Therefore, despite the numerous successes from bispecifics, there remains a huge 58 59 range of unexplored combinations of bispecifics that could illuminate the principles of cell 60 signaling and provide valuable therapeutic leads. It is an exciting challenge in protein 61 engineering to facilitate the screening of bispecific combinations.

The most common routes to generate bispecifics depend upon rearrangement of IgG 62 architecture, particularly through knob-into-hole generation of heterodimeric Fc domains 63 (Spiess et al., 2013), or disulfide bond shuffling in the Fabs (Dengl et al., 2020; Labrijn et al., 64 65 2013). Alternative bispecific pairing has been achieved by conjugating the two binders using sortase (Andres et al., 2020), transglutaminase (Plagmann et al., 2009), click chemistry (Szijj 66 and Chudasama, 2021) or split inteins (Han et al., 2017). However, these routes require each 67 68 binder to be fused to the respective conjugation domain, such that each binder must be recloned, re-expressed, and re-purified in two formats to explore all possible bispecific 69 70 combinations. Such a requirement is a major problem as the field moves towards exploring 71 thousands of binders across the proteome (Cao et al., 2022; Colwill et al., 2011; Dübel et al., 72 2010).

Our group previously developed an efficient way to ligate genetically-encoded building blocks, 73 74 through spontaneous isopeptide bond formation between a peptide Tag and a complementary 75 Catcher protein (Keeble and Howarth, 2020; Keeble et al., 2022). SpyTag/SpyCatcher has 76 been widely used for modular assembly of binders, enzymes and vaccines (Keeble and 77 Howarth, 2020; Pardee et al., 2014). SpyTag003/SpyCatcher003 is a pair that we subsequently 78 developed for reaction at a rate close to the diffusion limit, which maintains compatibility with 79 all previous Spy versions (Keeble et al., 2019). SpyTag/SpyCatcher has been previously 80 employed for bispecific assembly but faces the above limitation of having to clone each binder 81 in two different formats, with one version fused to SpyTag and the other version fused to 82 SpyCatcher (Chen et al., 2022; Keeble et al., 2022; Sutherland et al., 2019). In this work we 83 establish a protease-activatable SpyCatcher003, to enable the generation of bispecific 84 combinations where each binder only needs to be cloned into one common format - the 85 SpyCombinator strategy. We show that binders can be assembled onto a panel of scaffolds, in order for the binders to project in different directions and change the orientation of the target 86 87 receptor. We evaluate the SpyCombinator system against human epidermal growth factor 88 receptor 2 (HER2), which has a central role in breast, gastric, and ovarian cancers through proliferative and anti-apoptotic signaling (Scholl et al., 2001). Employing SpyCombinator on 89 90 a panel of 9 binders (affibody, nanobody or antibodies) allowed simple generation of 81 binder 91 pairs against HER2 (9 with bivalent and 72 with bispecific binding activity). In order to demonstrate the mutability of our modular bispecific assembly approach, DoubleCatcher 92 93 scaffolds that vary in dimensional architecture and rigidity were designed by modifying the 94 linker separating Catcher domains, or by introducing an intramolecular disulfide bond. Again, SpyCombinator was applied to dimerize a subset of 9 pairs of anti-HER2 binders using the 95 panel of 7 DoubleCatcher variants, to generate 63 bivalent binders spanning a diverse format 96 97 space. These pairs of binders show major differences in their effect on cell survival and 98 division, illustrating the potential of the SpyCombinator approach to accelerate combinatorial 99 discovery and fine-tune cellular responses to designed receptor modulators.

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101 **Results**

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103 Masked SpyCatcher003 is a protease-activatable variant of SpyCatcher003

Our approach uses a tandem SpyCatcher003 for specific irreversible heterodimerization of any 104 105 two antigen-binding units fused to SpyTag or its variants (i.e. SpyTag002 or SpyTag003) (Fig. 106 1A). The N-terminal SpyCatcher003 is accessible for reaction. The C-terminal SpyCatcher003 107 is masked. The non-reactive SpyTag003 D117A (SpyTag003DA) mutant (Keeble et al., 2019) is fused to the C-terminus of the second SpyCatcher003 via a flexible linker containing a 108 tobacco etch virus (TEV) protease cleavage site (ENLYFQ/G) (Fig. 1A). Before cleavage, 109 110 SpyTag003DA should bind tightly to SpyCatcher003, with $K_d = 21$ nM for the non-linked pair (Keeble et al., 2019), thus masking the reactive Lys of SpyCatcher003. The intramolecular 111 SpyCatcher003/SpyTag003DA interaction should be entropically favored over a competing 112 113 intermolecular interaction with SpyTag003 in solution. Following cleavage at the TEV site, the SpyTag003DA peptide will be free to dissociate, unmasking the reactive Lys of 114 SpyCatcher003 to enable reaction with a second supplied SpyTag-linked binder. We performed 115 116 all cleavages with superTEV protease, because of its efficient activity without the need for reductant (Cabrita et al., 2007). 117

We piloted the SpyCombinator strategy on a single SpyCatcher003 moiety. Masked 118 119 SpyCatcher003 was cleaved with superTEV protease and incubated with a model fusion protein of SpyTag003 linked to maltose-binding protein (MBP) (Fig. 1B). This reaction 120 121 proceeded efficiently to > 99% completion, with a half-life of reaction ($t_{1/2}$) of 2.7 ± 0.7 min 122 (Fig. 1B, C, D). To reduce the extent to which SpyTag003DA dissociation limits the rate of reaction between TEV-cleaved Masked SpyCatcher003 with SpyTag003-MBP, the reaction 123 124 temperature was elevated to 37 °C and the molar excess of SpyTag003-MBP relative to Masked 125 SpyCatcher003 was increased to 5-fold. The muted reactivity of Masked SpyCatcher003 before cleavage was probed by incubating equimolar amounts of the construct with SpyTag003-MBP 126 and monitoring the disappearance of Masked SpyCatcher003 over time by SDS-PAGE and 127 128 densitometry (Fig. 1D, E). Under these conditions, we observed 3.7% reaction of Masked 129 SpyCatcher003 with SpyTag003-MBP after 60 min, supporting effective silencing of 130 SpyCatcher003 reactivity.

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132 Establishing DoubleCatcher as a tool for controlled combination of distinct proteins

133 To design a platform that facilitates the conjugation of two Tagged proteins, Masked SpyCatcher003 was genetically fused to the C-terminus of a native SpyCatcher003 protein via 134 a flexible Gly/Ser linker, to give DoubleCatcher (Fig. 2A, with sequence shown in 135 Supplementary Fig. 1). The reactivity of each SpyCatcher003 moiety within the DoubleCatcher 136 137 construct was determined by monitoring the reaction of DoubleCatcher with SpyTag003linked model proteins before and after cleavage by MBP-superTEV protease. First, 138 DoubleCatcher was incubated with 2-fold excess of SpyTag003 linked to the green fluorescent 139 140 protein mClover3 in the absence of TEV protease (Fig. 2A, B). 91% reactivity of the N-141 terminal SpyCatcher003 was observed after 1 h, with a $t_{1/2}$ of 0.58 ± 0.05 min (Fig. 2C).

Next we incubated DoubleCatcher with two-fold excess SpyTag003-MBP for 2 h to 142 saturate the N-terminal SpyCatcher003 moiety. Unreacted SpyTag003-MBP was removed 143 144 from the reaction mixture by overnight incubation with bead-immobilized SpyCatcher003. Addition of superTEV protease to the crude mixture (4 h, 34 °C) induced cleavage of the 145 SpyTag003DA mask from the C-terminal SpyCatcher003 (Fig. 2D). 2 µM of the cleaved 146 147 construct was then reacted with a 5-fold molar excess of SpyTagged Small Ubiquitin-like 148 Modifier (SUMO-SpyTag003) and the reaction was monitored over time, by SDS-PAGE and 149 densitometry (Fig. 2E, F). Reaction to fill the second site of DoubleCatcher was rapid with $t_{1/2}$

150 of 3.6 ± 0.7 min, showing that the cleaved SpyTag003DA can be efficiently replaced by a 151 reactive Tag.



Figure 1. Masked SpyCatcher003 has protease-activatable reactivity. (A) SpyCombinator 152 approach. DoubleCatcher consists of an accessible N-terminal SpyCatcher and a C-terminal 153 154 SpyCatcher masked by the unreactive SpyTag003DA peptide. The first SpyTag-linked binder reacts with the N-terminal Catcher but is blocked from reacting at the second Catcher by the 155 tethered unreactive SpyTag003DA peptide. Addition of TEV protease induces cleavage of 156 SpyTag003DA, so that the second Catcher can be loaded with a different SpyTag-linked 157 binder. SpyTag003 in cyan with D117A indicated in yellow, TEV protease site in orange, Fabs 158 in pink, and isopeptide bond in red. A colon indicates covalently linked products. (B) Cartoon 159 following protease-controlled unmasking of SpyCatcher003. Incubation of Masked Catcher 160 with TEV protease enables cleavage and dissociation of SpyTag003DA from SpyCatcher003, 161 allowing SpyTag003-linked proteins to react with SpyCatcher003 ('Unmasked Catcher'). (C) 162 Efficient coupling after TEV protease unmasking. After incubation of Masked Catcher with 163 TEV protease, the unmasked Catcher (5 µM) is incubated with equimolar SpyTag003-MBP 164 for the indicated time, before SDS-PAGE with Coomassie staining. (D) Quantification of 165 SpyCatcher003 reactivity with or without masking (individual replicates with means 166 167 connected, n = 3), indicating the mean half-time of reaction, $t_{1/2}$. (E) Minimal reactivity of Masked SpyCatcher003 prior to TEV protease cleavage. Masked SpyCatcher003 was 168 incubated with SpyTag003-MBP (both partners at 5 µM) for the indicated time at 37 °C, before 169 170 SDS-PAGE/Coomassie.



171 Figure 2. Dual conjugation of Tagged proteins to DoubleCatcher is fast and site-specific. (A) Schematic of covalent conjugation of SpyTag003-mClover3 to the reactive N-terminal 172 Catcher of DoubleCatcher. (B) Time-course of reaction between DoubleCatcher (DC; 2.5 µM) 173 and SpyTag003-mClover3 (5 µM), analyzed by SDS-PAGE/Coomassie. (C) Depletion of 174 175 unreacted DoubleCatcher over the course of reaction was quantified by densitometry, 176 following (B). (D) Schematic depicting unmasking and conjugation of the C-terminal Catcher 177 of DoubleCatcher. (E) Reaction at the unmasked Catcher of DoubleCatcher. Following saturation of the N-terminal Catcher of DoubleCatcher with SpyTag003-MBP ('A'), the 178 179 sample was incubated with TEV protease for 2 h at 34 °C. The resulting unmasked DoubleCatcher:SpyTag003-MBP (2 µM) was then reacted with SUMO-SpyTag003 ('B', 10 180 μ M) over the time indicated, before SDS-PAGE/Coomassie. (F) Quantification of reaction at 181 the unmasked Catcher of DoubleCatcher, based on (E). All triplicate data points are shown, 182 with the line connecting the mean. 183

184 Establishment of scalable heterodimerization of proteins using SpyCombinator

A universal bispecific assembly platform for high-throughput screening requires a simple and 185 scalable methodology. In particular, procedures involving multiple rounds of column 186 187 chromatography will provide a bottleneck to the combinatorial complexity of bispecifics that can be interrogated. Therefore, we next sought to optimize a one-pot reaction in which any two 188 Tagged binders can be conjugated to DoubleCatcher (Fig. 3A), and the resulting bispecific 189 190 molecule can be purified (Fig. 3B). For optimization, model ligands were assembled onto DoubleCatcher: these were a SpyTag construct linked to three tandemly-repeated affibodies 191 against HER2 (SpyTag-AfiiHER2₃, ligand A) and SpyTag003 linked to a high stability MBP 192 193 tandem fusion ((MBPx)₂-SpyTag003, ligand B) (Veggiani et al., 2016). These ligands lack free 194 Cys (avoiding impurity from inter-ligand disulfides) and have a difference in mass that permits 195 unambiguous detection of reactant and product bands by SDS-PAGE (Fig. 3A). The intact masses of these purified reactants in PBS pH 7.4 were validated by electrospray-ionization 196 197 mass spectrometry (Supplementary Fig. 2).

In step 1 of our one-pot assembly route, DoubleCatcher reacts with a small molar excess 198 199 (1.25-fold) of SpyTag-AffiHER23, to ensure negligible unreacted DoubleCatcher in solution 200 (Fig. 3A). SDS-PAGE revealed 97% of the reaction contained the desired singly-occupied 201 DoubleCatcher:SpyTag-AffiHER2₃ species (Fig. 3C, lane DC + A). As observed for Masked 202 SpyCatcher003 reactivity tests (Fig. 1D), 3% contained DoubleCatcher conjugated to two 203 3C). confirm where the homodimeric SpyTag-AffiHER2₃ molecules (Fig. То 204 DoubleCatcher:(SpyTag-AffiHER2₃)2 would run, we added superTEV protease (1/12th molar 205 equivalent) to the crude mixture; the homodimer in the reaction mixture increased to 30% ('DC + A + TEV' lane in Fig. 3C). The identity of this by-product was corroborated by the increase 206 207 in the density of the band corresponding to AffiHER23 homodimer in a control reaction 208 containing twice the concentration of SpyTag003-AffiHER2₃ (DC + A + TEV + A' lane in 209 Fig. 3C).

For subsequent unmasking and saturation of the C-terminal SpyCatcher003 moiety, 210 superTEV protease (1/12th molar equivalent) and (MBP_X)₂-SpyTag003 (5-fold molar excess, 211 'B') were added simultaneously to the singly saturated DoubleCatcher (Step 2, Fig. 3A). Thus, 212 immediately upon cleavage of SpyTag003DA, the concentration of (MBP_X)₂-SpyTag003 in 213 214 solution should be 20 times that of unreacted SpyTag-AffiHER23 and so homodimer formation 215 is minimized. Indeed, AffiHER23 homodimer accounted for only 3% of the total species in the 216 reaction mixture ('DC + A + TEV + B' in Fig. 3C). Trace (3%) formation of DoubleCatcher conjugated to two (MBPx)2-SpyTag003 molecules ('MBP homodimer') was observed by SDS-217 218 PAGE, validated by the formation of a band of equivalent mobility in a reaction with 219 DoubleCatcher, superTEV protease and (MBP_X)₂-SpyTag003 alone ('DC + B + TEV + B' lane 220 in Fig. 3C).

221 For depletion of excess SpyTag- or SpyTag003-linked binder from the desired product, the crude reaction mixture was incubated with bead-immobilized SpyCatcher003, by adding 222 223 beads in a 1:1 suspension with PBS directly to the reaction mixture (Step 3, Fig. 3A and Fig. 224 3B). The purity of DoubleCatcher:SpyTag-AffiHER2₃:(MBP_X)₂-SpyTag003 following purification was assessed by SDS-PAGE densitometry as 95% (excluding superTEV protease) 225 (Purification lane in Fig. 3C). Based on SDS-PAGE, superTEV protease (70 kDa, Fig. 3D, E) 226 227 was the major contaminant. Since TEV protease has exceptional specificity for its target 228 sequence, which is not found in the human proteome, and has been expressed before without 229 effect on cells (Gray et al., 2010), we did not consider it necessary to remove. However, it should also be possible to add amylose-agarose to the well, for depletion of the MBP-linked 230 superTEV protease, if considered helpful. 231

We also validated the bispecific assembly by mass photometry, allowing single molecule-based assessment of each species in solution (Asor and Kukura, 2022). The peak

corresponding to DoubleCatcher:SpyTag-AffiHER2₃:(MBP_X)₂-SpyTag003 was observed with
 a mass of 144 kDa, matching well to the 144.6 kDa predicted mass (Fig. 3D). The second peak
 of 70 kDa, fitted well to MBP-superTEV protease (Fig. 3D), as validated by running MBP-

superTEV protease on its own (Fig. 3E).



Figure 3. One-pot conjugation of DoubleCatcher with Tagged model proteins to yield a 238 bispecific with high purity. (A) Schematic depicting assembly of SpyTag-AffiHER2₃ (pink) 239 240 and (MBP_X)₂-SpyTag003 (purple) onto DoubleCatcher (DC, blue). N represents the 241 concentration of DoubleCatcher and then the relative molarity of each Tag-binder is marked. 242 Immobilized SpyCatcher003 removes free Tagged proteins, leaving the bispecific in solution. (B) One-pot bispecific assembly as in (A), monitored by SDS-PAGE/Coomassie. (C) 243 244 Schematic of purification of excess Tag-fusion by immobilized SpyCatcher003 S49C in Step 3. (D) Mass photometry of SpyCombinator reaction. Peaks are shown for the product of 245 246 coupling, as in (B), with observed and expected M_w indicated, confirming the identity of the target bispecific. The other species in the reaction was TEV protease. (E) Mass photometry of 247 TEV protease alone, annotated as in (D). 248

Anti-HER2 bispecifics assembled by SpyCombinator induce various anti-proliferative or proliferative effects in HER2-addicted cancer cells

The work presented above demonstrates the efficient assembly of bispecific ligands through 251 252 DoubleCatcher. We then aimed to complement the SpyCombinator platform with a screening approach of equal throughput on binders against the important cell-surface target of HER2. 253 The relative activity of each signaling pathway and downstream physiological outcome has 254 255 been attributed to different ligands stabilizing receptor dimers in distinct conformations, 256 inducing distinct phosphorylation of the 17 sites in HER2's intracellular domain (Freed et al., 257 2017). Although the functional selectivity of native Receptor Tyrosine Kinase ligands has been 258 extensively studied, this bias is yet to be exploited for personalized cancer therapies. Thus, we 259 sought to assemble a large matrix of anti-HER2 bispecifics which differ in format and screen for their ability to alter HER2 activity (Brack et al., 2014; Stüber et al., 2021). Binders specific 260 to each subdomain of the HER2 extracellular region were identified (Fig. 4A, B), including 261 Fabs based on the clinically used Pertuzumab ('Pert') and Trastuzumab ('Tras', known 262 commercially as Herceptin). Tras is a Trastuzumab variant with a point mutation to enhance 263 264 binding stability (Jain et al., 2013). Each binder was expressed as a SpyTag003-fusion in Expi293F cells, except the nanobody (nanoHER2) (Verheije et al., 2013) and affibody 265 (AffiHER2) (Feldwisch et al., 2010) which were expressed in E. coli. We confirmed the 266 267 identity of each binder using electrospray-ionization mass spectrometry (Supplementary Fig. 3). Fusions to SpyTag are conventionally expressed with a spacer to favor high reactivity with 268 269 SpyCatcher and optimal activity of the fused domain (Keeble and Howarth, 2019). We 270 expressed a variant of Tras with no linker to SpyTag003, to test the importance of linker flexibility on activity, termed Tras NoLink. The NoLink variant was confirmed to react to 97% 271 in 60 min in PBS pH 7.4 at 37 °C with DoubleCatcher, an efficiency comparable to the linker-272 273 containing parental construct (99% in 60 min under equivalent reaction conditions; Supplementary Fig. 4), confirming that reactivity is not impaired by removing the linker 274 275 between the Fab and the reactive peptide. Each binder was dimerized with itself or another binder using SpyCombinator methodology (Fig. 3A). We studied the effects of the resulting 276 277 matrix of anti-HER2 bispecifics and homodimers on metabolic activity in HER2-addicted 278 BT474 cells (a human ductal breast carcinoma cell-line) or SKBR3 cells (a human 279 adenocarcinoma cell-line), quantifying using the resazurin assay (Stüber et al., 2021).

Following incubation of BT474 or SKBR3 cells with anti-HER2 bispecific assemblies (100 nM) for 96 h, the change in metabolic activity was calculated relative to untreated cells (Fig. 4C, D). We discovered that the different assemblies exhibited a range of antagonistic and agonistic activities. We found that bispecific constructs displaying one or more arm binding to domain II (39S, HFS2, or Pert) are consistently anti-proliferative.

285 Anti-HER2 bispecifics also showed different potencies in SKBR3 cells relative to 286 BT474 cells. Firstly, a higher number of bispecifics had significant proliferative activity in BT474 (20 constructs) than in SKBR3 (10 constructs). Secondly, constructs with significant 287 anti-proliferative activity were more potent in BT474 (average metabolic activity of significant 288 289 antagonists = $58.4 \pm 20.4\%$) than in SKBR3 (average metabolic activity of significant 290 antagonists = $64.2 \pm 7.0\%$). In BT474 cells, the potent antagonist Tras NoLink-Tras (74.1 \pm 6% (BT474) metabolic activity after 96 h treatment) showed no activity when assembled in the 291 292 opposite orientation: Tras-Tras NoLink ($103 \pm 1\%$ for BT474 metabolic activity). Importantly, 293 the same orientation-dependence was observed in SKBR3 cells for the significantly active 294 bispecifics nanoHER2-HFS2, Tras-AffiHER2, nanoHER2-AffiHER2, MF3958-Tras NoLink, 295 Tras-39S, 39-Tras NoLink, Tras-MF3958; where assembly of these binders in the opposite 296 orientation removes their significant agonistic or antagonistic activity.

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298 Figure 4. SpyCombinator bispecific panel exhibits a range of pro- or anti-proliferative 299 activities on cancer cells. (A) Binding sites on HER2 for the panel of anti-HER2 binders. 300 For each subdomain (I-IV) of HER2's extracellular domain at least one binding molecule was identified and the cartoon illustrates their binding site on HER2 (gray), with each binder in a 301 302 different color. (B) Subdomains of HER2's extracellular domain are colored, with anti-HER2 binders from (A) overlaid in gray. (C) SpyCombinator anti-HER2 panel effect on SKBR3 303 304 cell proliferation. The vertical column indicates the binder at the N-terminal site of 305 DoubleCatcher. The horizontal column shows the binder at the C-terminal site of DoubleCatcher. The effect on proliferation of SKBR3 cells with 100 nM bispecific after 4 306 days from resazurin assay is shown as a heat map, with red showing reduced proliferation 307 308 and blue showing enhanced proliferation, compared to the DoubleCatcher alone control at 309 100%. (D) Bispecific panel effect on proliferation of BT474, as in (B). (E) Titration curves 310 for the top 5 bispecifics with significant anti-proliferative activity in SKBR3 cells, assayed as in (C). (F) Titration curves for the top 5 bispecifics with significant pro-proliferative activity 311 312 in SKBR3 cells, assayed as in (C).

Examining the effect of removing the linker between binder and SpyTag003, the average difference between bispecifics with at least one Fab0.11 versus at least one Tras NoLink was only 2%. The similarity between the activities of Tras and Tras NoLink supports the usage of linker-free binder-SpyTag fusions in future screens, where it will be beneficial to maximize rigidity in the bispecific molecule and impart control over target geometries. Together, these results highlight the importance of screening epitope-diverse bispecific binders in multiple formats and orientations.

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321 DoubleCatcher architecture can be customized

As with most bispecific antibodies, the format of a bispecific binder built using SpyCombinator 322 323 may be influenced by the identity of the SpyTag003-linked binder, and by the order in which 324 the binders are coupled. However, the overall shape, size, and flexibility of the bispecific 325 molecule may also be determined by the DoubleCatcher module itself. To amplify the format 326 space to be screened, matrices of bispecific antibodies could therefore be developed in three 327 dimensions: varying two binder arms against the targets of interest and, in parallel, varying the 328 module. The highest-scoring AlphaFold2-predicted DoubleCatcher structure for DoubleCatcher is displayed in Fig. 5A (left), with the position and orientation (N to C) of the 329 330 two SpyTag binding sites represented by cyan vectors. Modifications to the DoubleCatcher architecture that change the relative spacing and orientation of the cyan vectors will reflect 331 332 changes in the structural relationship between the two target-binding arms in the final 333 bispecific.

334 Modifications to the architecture of DoubleCatcher may be achieved by varying the 335 linker connecting the two SpyCatcher003 moieties. The amenability of DoubleCatcher to linker modification was tested by substituting the Gly/Ser linker with the central helix from the 336 Bacillus stearothermophilus ribosomal protein L9 (Kuhlman et al., 1997). The DoubleCatcher 337 338 variant with the helical linker is termed 'DoubleCatcher H-lock' (Fig. 5A, right, and sequence 339 in Supplementary Fig. 1). Size-exclusion chromatography analysis of Ni-NTA-purified 340 DoubleCatcher H-lock revealed its apparent molecular mass was 17 kDa larger than DoubleCatcher (Fig. 5D, E), while the predicted increase in mass is only 3.5 kDa, as validated 341 by electrospray-ionization mass spectrometry (Supplementary Fig. 2). 342

343 One approach to increasing the rigidity of the DoubleCatcher module that overcomes 344 the flexibility of the SpyCatcher003 termini is to engineer a covalent linkage between the two Catcher domains. A pair of residues (one residue from each SpyCatcher003) that reside in loop 345 346 regions and close proximity (1.2 nm) within each of the the five highest-confidence predicted 347 structures of DoubleCatcher was chosen as the site for cysteine substitution to engineer a SpyCatcher003-SpyCatcher003 intramolecular disulfide bond. The AlphaFold2-predicted 348 349 structure of the double-cysteine variant designed from the top-predicted DoubleCatcher 350 structure is presented in Fig. 5B and is termed 'DoubleCatcher α -lock'. The substantially less 351 variable average distance between N_E of the reactive Lys residues of each SpyCatcher003 in 352 the top 5 AlphaFold2-predicted structures of DoubleCatcher α -lock (4.7 ± 0.04 nm) suggests the two SpyCatcher003s may have low relative mobility. The DoubleCatcher double-cysteine 353 variants designed from the four remaining predicted structures of DoubleCatcher are presented 354 355 in Fig. 5C. Therefore, the combination of SpyTagged antigen binders using these 356 DoubleCatcher variants should provide conformationally distinct bispecific binders that bind 357 targets in different orientations.

358 Size-exclusion chromatography profiles of Ni-NTA-purified double-cysteine DoubleCatcher 359 variants reveal that the construct exists as two species: one main peak corresponding to 360 monomer and one minor peak that corresponds to covalent dimer (Fig. 5D).

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Figure 5. Panel of DoubleCatcher architectures. (A) Schematic of DoubleCatcher or 362 DoubleCatcher with helical spacer. AlphaFold2-predicted structures show DoubleCatcher dark 363 blue and the Tag as a cyan vector, indicating the direction that a Fab-SpyTag fusion would 364 project. The flexible linker in DoubleCatcher (left) was replaced with a stable α -helical linker 365 in DoubleCatcher H-lock (right). (B) Schematic of DoubleCatcher locking through an 366 engineered disulfide bond. AlphaFold2-predicted structure, as in (A). The residues selected for 367 Cys substitution (left) and the disulfide bond in the resulting double-cysteine mutant 368 (DoubleCatcher α -Lock) are labeled as red spheres with sulfur in yellow. (C) Schematic of the 369 other DoubleCatcher Lock variants, labeled with their Cys substitutions. (D) Size-exclusion 370 371 chromatography of DoubleCatcher variants normalized by A₂₈₀. Chromatograms are staggered by 1 from adjacent curves in the y-axis. The apparent molecular weights of each species were 372 calculated following calibration with molecular weight standard proteins (Bio-Rad) (grey 373 arrows). (E) Dynamic light scattering of DoubleCatcher variants in PBS. Traces are staggered 374 by 1 from adjacent curves in the y-axis. (F) Quantification of dynamic light scattering, showing 375 mean hydrodynamic radius $(R_h) \pm 1$ s.d. (n = 10) for each DoubleCatcher. 376

377 DoubleCatcher double-cysteine monomer was incubated with CuSO₄ to drive disulfide bond formation prior to downstream analyses. The identity and disulfide bond formation of each 378 DoubleCatcher was validated by electrospray-ionization mass spectrometry (Supplementary 379 380 Fig. 2). The DoubleCatcher variants were analyzed by dynamic light scattering (DLS) (Fig. 5F), with quantification shown in Fig. 5G. As expected, both the mean hydrodynamic radius 381 (R_h) and the mean s.d. of the double-cysteine mutants $(3.5 \pm 1.0 \text{ nm}, \text{mean} \pm 1 \text{ s.d.}, \text{n} = 10)$ 382 383 were less than those for the cysteine-free variants (4.5 \pm 1.7 nm, mean \pm 1 s.d., n = 10), consistent with reduced flexibility within the architecture. 384

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SpyCombinator-assembled anti-HER2 bispecific binders are sensitive to DoubleCatcher architecture

To investigate the sensitivity of HER2 signaling to dimer geometry, a small-scale matrix of 388 anti-HER2 bispecific binders was assembled by using each of the 7 designed DoubleCatcher 389 variants to dimerize Tras NoLink, nanoHER2, and AffiHER2 in all 9 possible permutations. 390 The assembled bispecific binders were added to HER2-addicted SKBR3 cells (100 nM) for 96 391 392 h, after which the change in metabolic activity was calculated (Fig. 6A, B). For each pair of binders, cells were treated with a mixture of the purified binders (100 nM) without 393 394 DoubleCatcher, to test for monovalent binder activity- 'No DC' matrix in Fig. 6A. 395 Simultaneously, purified DoubleCatcher variants (100 nM) were added to cells, to test for any activity of scaffold alone (Fig. 6A). 396

397 In a similar way to the orientation-dependent activity of bispecifics assembled in Figure 398 4, we repeatedly found that the potency of specific combinations of binders were enhanced when assembled onto a different DoubleCatcher variant. The most pronounced example of this 399 is for the pro-proliferative DoubleCatcher β -lock bispecifics containing nanoHER2: relative to 400 401 the DoubleCatcher-assembled bispecifics, those assembled with DoubleCatcher \beta-Lock 402 induced +25.6 % greater proliferation in SKBR3 cells. Similarly, the anti-proliferative activity of Tras NL-Tras NL ($84.8 \pm 6.8\%$ metabolic activity for the DoubleCatcher-assembled 403 bispecific) was enhanced when dimerized with DoubleCatcher H-Lock (75.9 \pm 2.5%), 404 405 DoubleCatcher α -Lock (79.5 ± 3.9%), DoubleCatcher γ -Lock (80.4 ± 3.4%), DoubleCatcher δ -Lock (73.6 ± 4.9%), and DoubleCatcher ε -Lock (72.5 ± 4.3%). In addition to increased 406 potency, it was also observed that the ability of some pairs of binders to either agonize or 407 antagonize HER2 signaling was switched when assembled into a different architecture. 408 409 Specifically, the bispecific AffiHER2-Tras NoLink showed mild agonistic activity on the 410 DoubleCatcher scaffold (112.5 \pm 6.6% metabolic activity), yet the same bispecific assembled using DoubleCatcher H-Lock exhibited strong antagonistic activity under the same conditions 411 412 $(82.5 \pm 11.1\%)$. The same switch in activity was even more marked for Tras NoLink-AffiHER2 413 assembled in DoubleCatcher γ -Lock (78.6 ± 3.1 %) relative to DoubleCatcher (127.7 ± 4.1%), 414 accentuating the value of screening multiple formats in the assembly of bispecifics.

415

416 **Discussion**

Here we have demonstrated SpyCombinator assembly, allowing controlled and precise 417 construction of bispecific binders through sequential SpyTag/SpyCatcher conjugation. The 418 419 SpyCombinator platform offers a plug-and-play approach in which the antigen-binding building blocks exist in a common format, fused to the commonly-employed SpyTag peptide 420 or its subsequent generations (Keeble and Howarth, 2020; Keeble et al., 2019). Unlike standard 421 approaches to bispecific assembly (Hofmann et al., 2020), SpyCombinator eliminates the need 422 423 to clone and express each pair of binding proteins into more than one format. Neither SpyTag nor SpyCatcher require any cysteines and the overall assembly is ligated through amide bonds 424 425 (Keeble and Howarth, 2020), so SpyCombinator avoids the careful optimization and instability 426 that may arise from bispecific assembly relying on disulfide bond reduction and oxidation.



DoubleCatcher variant

427 Figure 6. Architectural modifications to anti-HER2 bispecifics change the signaling effects. (A) Different DoubleCatcher scaffolds change HER2 signaling. Matrices of anti-HER2 428 429 bispecifics were assembled by SpyCombinator using distinct DoubleCatcher variants. Treatment of SKBR3 cells with the bispecifics (100 nM, 4 days) revealed DoubleCatcher-430 dependence on their pro- or anti-proliferative potency, as determined by resazurin assay. Mean 431 metabolic activity, relative to untreated control, are presented as stacked heat maps, with red 432 showing reduced proliferation and blue enhanced proliferation. Activity following 100 nM 433 DoubleCatcher variant alone was measured in parallel. Below each variant, vectors depict the 434 position and orientation of C-N SpyTag003-binding groove within each SpyCatcher003 435 moiety, such that the binder projects in the direction of the vector arrowhead. (B) Graphical 436 437 representation of DoubleCatcher-dependence for cell proliferation, following (A). Each data-438 point is shown (n = 3), with the bar indicating the mean, ± 1 s.d..

439

440 Optimization of SpyCombinator conditions produced a generalized pipeline that generates 441 bispecifics with up to 97% purity relative to homodimeric by-products. The modularity of the 442 platform invites customization of each constituent, exemplified here through DoubleCatcher 443 variants with different linkers or intramolecular disulfide bonds. There is potential in future 444 work for substantial diversification of the DoubleCatcher module, to generate additional 445 bispecific architectures (Baker, 2019).

In this study, anti-HER2 binders varying in epitope and size were permuted into a 446 447 matrix of bispecifics. We identified anti-HER2 bispecifics with major differences in their pro-448 or anti-proliferative activity. Interestingly, this proliferative activity depended not only on the 449 identity of the anti-HER2 monomers, but also on the order of coupling to DoubleCatcher, affirming the importance of screening paratope orientation (Scheer et al., 2012; Stüber et al., 450 2021). Furthermore, in assembling a small panel of anti-HER2 binders into bispecifics using 451 452 DoubleCatchers of varying geometries and flexibilities, we found that the activity or potency of particular pairs of binders is dependent on their scaffold. Cellular effects of bispecific 453 454 bridging depend upon an intricate balance of allosteric interactions between intracellular kinase 455 domains (Zhang et al., 2006), altered heterodimer formation with other receptors (Gu et al., 456 2014), and receptor dynamics at the plasma membrane or endolysosomal compartments 457 (Scheer et al., 2012; Stüber et al., 2021). Therefore, being able to screen large numbers of bispecific combinations in multiple formats is crucial to the discovery of the most potent 458 459 effector activity. HER2 dimerization is facilitated predominantly by interactions between 460 domain II residues (Arkhipov et al., 2013), so these bispecifics are likely to block HER2 dimerization and activation. Conversely, bispecifics harboring the small, non-domain II 461 binders nanoHER2 (16 kDa) or AffiHER2 (15 kDa) display potent proliferative activity. 462 463 Indeed, the bispecific constructs with significant agonistic activity in both SKBR3 and BT474 464 cells all comprised at least one of nanoHER2 or AffiHER2. Where agonistic activity of 465 biparatopic anti-HER2 molecules has been observed, it has been suggested to result from favouring HER2 homodimerization over heterodimerization, or from a shorter distance 466 between antigen-binding sites (Gu et al., 2014; Kast et al., 2021; Scheer et al., 2012). 467 Considering their small size, DoubleCatchers coupled to either nanoHER2 or AffiHER2 may 468 469 bridge HER2 molecules close enough for trans-phosphorylation by HER2 kinase domains. 470 This result may be explained by the differential total expression of HER2, as well as the 471 expression of Epidermal Growth Factor Receptor (EGFR) and HER3 relative to HER2. BT474 expresses HER2 to a greater extent (~ 2.8×10^6 copies per cell) (Kamashev et al., 2022) relative 472 473 to SKBR3 (~1.2 \times 10⁶ copies) (Hendriks et al., 2013). Both express HER3 to a similar level $(1-1.2 \times 10^6 \text{ copies})$ but SKBR3 expresses ~3-fold more EGFR $(2.2 \times 10^5 \text{ copies})$ than BT474 474 $(7 \times 10^4 \text{ copies})$ (Brockhoff et al., 2001). HER2-HER3 dimerization mediates activation of 475 PI3K/Akt and proliferation axis (Junttila et al., 2009), while HER2-EGFR dimers favor 476 477 MAPK/ERK and pro-survival signaling (Kirouac et al., 2016). Therefore, for bispecifics preventing HER2 homodimerization but not occluding domain II of HER2 (for example, 478 479 domain I-binding H2-18 or MF3958), heterodimerization may occurs more favorably. This could result in a higher frequency of proliferative HER2-HER3 dimers in BT474 than SKBR3. 480

481 Differences in activities of anti-HER2 bispecifics have been attributed to the 482 architecture of the molecule, in addition to binder identity. Specifically, this includes the binder's ability to oligomerize HER2 monomers, resulting in arrested receptor diffusion 483 484 compartmentalization into membrane nanodomains (Stüber et al., 2021); the binder-defined 485 distance between cross-linked HER2 molecules (Kast et al., 2021); the distortion of HER2 486 extracellular domain upon binding, such that its ability to homo/hetero-dimerize is affected 487 (Gu et al., 2014); and the amenability of the binder's format to increased bispecific-HER2 complex stoichiometries (Brack et al., 2014). Our DoubleCatcher panel introduces variation in 488

489 relative paratope orientation, distance between paratopes, and scaffold rigidity. We hypothesize that these parameters and their combinations each influence HER2 dynamics by 490 one or more of the mechanisms highlighted. However, further experiments are required to 491 492 decipher their biochemical behaviours, such as employing BRET to study homo- vs. heterodimer formation (Siddiqui et al., 2013), single-particle tracking analysis to quantify changes in 493 receptor diffusion kinetics (Freed et al., 2017), or nanoparticle-aided cryo-electron microscopy 494 495 to gain insight into the conformational homogeneity, and thus rigidity, of different 496 SpyCombinator-assembled bispecific molecules (Ki et al., 2021).

497 Control over Tag/Catcher activity has previously been achieved using pH (Vester et al., 498 2022), redox (Matsunaga et al., 2013; Wu et al., 2018), temperature (Cao et al., 2017), or light 499 (Hartzell et al., 2021; Ruskowitz et al., 2023). While these switches have an array of valuable 500 applications, the simple generation and large change in reactivity of the protease-uncaging 501 system is advantageous for bispecific assembly. In particular, redox changes on the Catcher would be undesirable for assembling building blocks that themselves contain disulfides, 502 notably nanobodies and various peptide hormones. A potential limitation of SpyCombinator is 503 504 that SpyTag/SpyCatcher is derived from Streptococcus pyogenes, so immunogenicity may affect the use of DoubleCatcher bispecifics as clinical candidates (Rahikainen et al., 2021). 505 506 Nonetheless, Catcher bispecifics can be important leads to inform the design of biologics and 507 identify novel targets/epitopes for next generation therapeutics. Also, many factors other than 508 being non-self contribute to the immunogenicity of therapeutics (Yachnin et al., 2021), e.g. 509 bacterially-derived non-immunoglobulin scaffolds such as affibodies have shown promising 510 results for sustained therapy in clinical trials (Klint et al., 2023).

A major effort is under way to generate recombinant binders to the entire human 511 proteome (Cao et al., 2022; Colwill et al., 2011; Dübel et al., 2010). The data here and 512 513 elsewhere clearly illustrate how one binder is not enough and multiple binders to different 514 regions of a target are valuable (Kast et al., 2021). Therefore, with an arsenal of tens of 515 thousands of binders, methods such as SpyCombinator to screen the bispecific possibilities will accelerate the understanding and application of synergy for modulating cell signaling. 516 Increasing numbers of functional proteins have been SpyTagged (Hentrich et al., 2021; Keeble 517 and Howarth, 2019; Kellmann et al., 2023), allowing purification (Khairil Anuar et al., 2019; 518 519 Vester et al., 2022), immobilization (Wang et al., 2020), multimerization (Brune and Howarth, 520 2018) and networking to other molecular functionalities (Keeble and Howarth, 2020). Beyond proteins, SpyTag/SpyCatcher has been functionalized with nucleic acids (Kröll et al., 2022), 521 522 oligosaccharides (Li et al., 2022) or small molecules (Geissinger et al., 2020), which will allow 523 modular assembly of bispecifics with more diverse activities. We anticipate applications of 524 SpyCombinator beyond cell signaling, such as enhancing the recruitment of stem cells for 525 tissue repair (Cousens et al., 2009) or optimizing enzyme networks for catalysis (Wei et al., 526 2020).

527

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538 AUTHOR CONTRIBUTIONS

539 C.L.D. performed all experiments. A.H.K. designed and purified initial constructs. C.L.D. and 540 M.H. designed the project. C.L.D. and M.H. wrote the manuscript. All authors read and

541 approved the manuscript.

542

543 DECLARATION OF INTERESTS

544 M.H. is an inventor on a patent on spontaneous amide bond formation (EP2534484) and a 545 SpyBiotech co-founder and shareholder. M.H. and A.H.K. are inventors on a patent on 546 SpyTag003/SpyCatcher003 (UK Intellectual Property Office 1706430.4). C.L.D., A.H.K. and 547 M.H. are inventors on a patent application on the bispecific approach here (United Kingdom 548 Patent Application No. 2313175.8).

549

550 Materials and Methods

551

552 Plasmids and Cloning

553 Constructs were cloned by PCR methods using Q5 High-Fidelity Polymerase (New England
554 Biolabs) and fragments were assembled by Gibson assembly. All constructs were validated by
555 Sanger Sequencing.

pET28a-SpyTag003-MBP (GenBank Accession no. MN433888, Addgene plasmid ID 556 133450), pET28-SpyTag003-mClover3 (Addgene plasmid ID 133453) (Keeble et al., 2019), 557 558 pDEST14-SpyCatcher003 S49C (Addgene plasmid ID 133448) (Vester et al., 2022), pET28-559 MBP-super TEV protease (Addgene plasmid ID 171782) (Keeble et al., 2022), pET28a-560 SnoopTag-AffiHER2-SpyTag (N-terminal His6-SnoopTag-anti-HER2 Affibody-SpyTag) (GenBank accession no. KU296975) (Addgene deposition in progress) (Brune et al., 2017), 561 562 and pET28a-SnoopTag-SpyTag-(AffiHER2)3 (GenBank accession no. KU296976) (Addgene deposition in progress) (Veggiani et al., 2016) have been described. pET28a-SUMO-563 564 SpyTag003 (N-terminal His₆ tag-SUMO protein-SpyTag003) was cloned previously by Irsyad Khairil Anuar (University of Oxford) (GenBank and Addgene deposition in progress). 565 pDEST14-SpyCatcher003-TEVs-SpyTag003DA ('Masked SpyCatcher0003'; N-terminal His6 566 tag-SpyCatcher003-TEV protease cleavage site-SpyTag003 D117A, GenBank and Addgene 567 568 deposition in progress) was derived from pDEST14-SpyCatcher003 (GenBank Accession no. 569 MN433887, Addgene plasmid ID 133447) by incorporating the TEV protease cleavage site 570 (ENLYFQ/G) and SpyTag003 D117A (RGVPHIVMVAAYKRYK) (Keeble et al., 2019) by 571 Gibson assembly. pDEST14-SpyCatcher003-(GSG)₃-SpyCatcher003-TEVs-SpyTag003DA 572 (DoubleCatcher; N-terminal His6 tag-TEV protease cleavage site-two SpyCatcher003 moieties connected by a (GSG)3 spacer-TEV protease cleavage site-SpyTag003 DA, GenBank and 573 574 Addgene deposition in progress) was synthesized as a gene fragment (Integrated DNA 575 Technologies) and inserted into pDEST14. The Gly/Ser spacer in this construct was then replaced 576 with α-helical spacer (sequence an 577 PANLKALEAQKQKEQRQAAEELANAKKLKEQLEK) (Kuhlman et al., 1997), and this 578 construct was termed DoubleCatcher H-Lock (GenBank and Addgene deposition in progress). 579 Double-cysteine substitution variants of DoubleCatcher were derived from pDEST14-580 SpyCatcher003-(GSG)₃-SpyCatcher003-TEVs-SpyTag003DA by Gibson assembly: 581 DoubleCatcher α -Lock (GenBank and Addgene deposition in progress), DoubleCatcher β -582 Lock (GenBank and Addgene deposition in progress), DoubleCatcher y-Lock (GenBank and 583 Addgene deposition in progress), DoubleCatcher δ-Lock (GenBank and Addgene deposition 584 in progress), and DoubleCatcher ɛ-Lock (GenBank and Addgene deposition in progress). 585 pET28a-MBP_X-MBP_X-SpyTag003 (His₆-MBP_X-SSSGGSGGGSG linker-MBP_X-SpyTag003; GenBank and Addgene deposition in progress), where MBP_X is a variant of MBP with stronger 586 maltose binding (Veggiani et al., 2016), was cloned by Dr Henry Wood (University of Oxford). 587 588 pcDNA3.1-Tras heavy chain-GSG-SpyTag003 ('Tras'), pcDNA3.1-Tras heavy chain-

589 SpyTag003 ('Tras NoLink') and pcDNA3.1-Tras light chain (GenBank and Addgene 590 deposition in progress) were cloned previously by Jamie Lam (University of Oxford) from 591 previous constructs in pOPIN, which have been described (Vester et al., 2022).

592 Sequences for the heavy and light chains of anti-HER2 Fabs were synthesized (Twist 593 Bioscience) and cloned into pcDNA3.1 plasmid: 39S (Oganesyan et al., 2018), MF3958 (Geuijen et al., 2018), H2-18 (Hu et al., 2015), HFS2 (Salimi et al., 2018) and Pert (Gennaro 594 595 et al., 2014). Each Fab heavy chain had the final assembly: N-terminal signal peptide-Heavy 596 variable domain-Human IgG1 C_H1 domain-EPKSC IgG1 upper hinge region residues-597 SpyTag003-His₆ tag, with the exception of HSF2 whose C_H1 domain was of mouse origin, and 598 Pert for which the hinge region was omitted (GenBank and Addgene deposition in progress). 599 Each Fab light chain had the final assembly: N-terminal signal peptide-Light variable domain-600 Human kappa light chain, with the exception of HSF2 whose kappa light chain was of mouse origin. pET28a-nanoHER2-SpyTag003 (GenBank and Addgene deposition in progress), based 601 602 on the 11A4 nanobody against HER2 (Verheije et al., 2013), was cloned with a C-terminal His₆ 603 tag.

604

605 Bacterial Protein Expression

pDEST14 constructs were transformed into chemically competent E. coli C41 (DE3), a gift 606 from Anthony Watts (University of Oxford), except for the pDEST14-SpyCatcher003-607 608 SpyCatcher003-TEV site-SpyTag003DA disulfide-containing variants, which were 609 transformed into chemically competent E. coli T7SHuffle (NEB). pET28a and pETDuet 610 constructs were transformed into chemically competent E. coli BL21 (DE3) (Agilent 611 Technologies), with the exception of pET28a-SnoopTag-AffiHER2-SpyTag, which was 612 transformed into chemically competent E. coli T7 Express (NEB). Colonies were picked into 10 mL LB containing either 100 µg/mL ampicillin (pDEST14/pETDuet) or 25-50 µg/mL 613 614 kanamycin (pET28a), and grown at 37 °C at 200 rpm for 4-16 h. Starter cultures were diluted 100-fold into either 1 L LB supplemented with appropriate antibiotic and 0.8% (w/v) glucose, 615 1 L LB supplemented with antibiotic alone (T7 Express only), or 2 × TY supplemented with 616 antibiotic and 0.5% (v/v) glycerol (T7SHuffle only), and grown at 37 °C with shaking at 200 617 rpm. At OD₆₀₀ 0.5-0.6, expression was induced with 0.42 mM isopropyl β-D-1-618 619 thiogalactopyranoside (IPTG; Fluorochem). pDEST14-transformed cultures were then 620 incubated for 4 h at 30 °C and 200 rpm, while all other cultures were incubated for 16 h at 18 °C and 200 rpm post-induction. Cells were harvested by centrifugation at 4,000 g for 15 min 621 at 4 °C and pellets were stored at -80 °C until purification. 622

623

624 Mammalian Protein Expression

Anti-HER2 Fabs were expressed in Expi293F cells (Thermo Fisher), which were maintained 625 in Expi293 Expression Medium (Thermo Fisher) supplemented with 50 U/mL penicillin and 626 627 50 µg/mL streptomycin and grown in a humidified Multitron Cell incubator (Infors HT) at 37 628 °C with 8% (v/v) CO₂ and shaking at either 125 rpm (125 mL flasks; Corning) or 240 rpm (50 629 mL mini bioreactor tubes; Corning). After > 3 passages, cells were diluted to 3×10^{6} cells/mL in antibiotic-free Expi293 Expression Medium and transfected using an ExpiFectamine 293 630 Transfection Kit (Thermo Scientific). ExpiFectamine 293 Transfection Enhancers 1 and 2 were 631 632 added 20 h post transfection and cultures were incubated for 5 days post-transfection, when 633 cell viability fell below 50%. Cell supernatants were supplemented with cOmplete, Mini, EDTA-free Protease Inhibitor Cocktail (Roche) and clarified by centrifugation at 4,000 g at 4 634 635 °C, followed by filtration through a 0.22 µm syringe filter (Thermo Fisher). One tenth 636 supernatant volume of Ni-NTA wash buffer (50 mM Tris-HCl pH 7.8, 300 mM NaCl, 10 mM imidazole) was added to filtered supernatant, which was then pH-adjusted to pH 7.6 using 1 M 637

Tris-HCl pH 8.0. Supernatants were stored at -80 °C until purification. The typical yield for
SpyTag003-fused Fabs ranged from 6 to 156 mg per L of culture.

640

641 Ni-NTA purification of His6-tagged proteins

After thawing, bacterial cell pellets were resuspended in 1× Ni-NTA buffer (50 mM Tris-HCl 642 pH 7.8, 300 mM NaCl) supplemented with cOmplete, Mini, EDTA-free Protease Inhibitor 643 644 Cocktail (Roche) and 1 mM phenylmethylsulfonyl fluoride (PMSF), and lysed by sonication 645 on ice. SpyCatcher003 S49C variants and DoubleCatcher double-cysteine variants were further supplemented with 10 mM 2-mercaptoethanol to break disulfide bonds to other proteins. 646 647 Clarified cell lysates were centrifuged at 30,000 g for 30 min at 4 °C and adjusted to pH 7.6 648 using 1 M Tris-HCl pH 8.0, before purification by Ni-NTA affinity chromatography (Qiagen) as previously described (Fierer et al., 2014). Elution of all proteins was performed in the 649 absence of 2-mercaptoethanol. For anti-HER2 binding proteins to be heterodimerized by 650 651 DoubleCatcher for treatment of cells, the following washes were performed directly prior to elution: 50 column volumes (CV) wash buffer containing 0.1% (v/v) Triton X-114 (50 mM 652 Tris-HCl pH 7.8, 300 mM, 10 mM imidazole), followed by 10 CV wash buffer (50 mM Tris-653 654 HCl pH 7.8, 300 mM). Clarified mammalian supernatants were purified identically. Eluted proteins were dialyzed three times against PBS pH 7.4 (137 mM NaCl, 2.7 mM KCl, 10 mM 655 Na₂HPO₄, 1.8 mM KH₂PO₄) in 3.5 kDa molecular weight cut-off Spectra/Por tubing (Spectrum 656 Labs) at 4 °C. SpyCatcher003 S49C variants to be coupled to resin were instead dialyzed into 657 658 coupling buffer (50 mM Tris-HCl pH 8.5, 5 mM EDTA). Protein concentrations were 659 determined from A₂₈₀ using their predicted extinction coefficient from ExPASy ProtParam. 660 Typical yield for SpyCatcher003-containing variants was 10-20 mg per L of culture. A typical 661 yield for affibody expression was ~3 mg per L of culture, and 30 mg per L for nanobody 662 expressions. Expression of SpyTag003-fused, non-antigen binding proteins yielded 5-15 mg 663 protein per L of culture.

Prior to assembly of bispecific molecules with DoubleCatcher for mammalian cell 664 treatment, endotoxin was removed from DoubleCatcher scaffolds by phase separation with 665 666 Triton X-114 (Aida and Pabst, 1990). Purified protein samples were transferred to 1.5 mL endotoxin-free microcentrifuge tubes (StarLab, cat. E1415-1510), to which 1% (v/v) Triton 667 668 X-114 was added. Samples were incubated on ice for 5 min, until all Triton X-114 was 669 dissolved. Samples were then incubated at 37 °C for 5 min and centrifuged for 1 min at 16,000 g at 37 °C. The supernatant was pipetted off, and the entire procedure was repeated twice. 670 Endotoxin concentration of any bispecific constituent to be added to cells was determined using 671 the Limulus amebocyte lysate (LAL) Chromogenic Endotoxin Quantitation Kit (Thermo 672 673 Fisher) according to the manufacturer's instructions, which confirmed the endotoxin level to 674 be below 1 endotoxin unit/mL.

675

676 SpySwitch purification of SpyTag003-tagged proteins

677 MBP_X-MBP_X-SpyTag003 cell pellets were lysed and clarified as outlined above, except 678 thawed pellets were resuspended in 1 × SpySwitch buffer (50 mM Tris-HCl pH 7.5, 300 mM NaCl). SpySwitch purification was performed at 4 °C, as described previously (Vester et al., 679 2022). Briefly, SpySwitch resin (0.75 mL per 1 L bacterial culture) was equilibrated with 2×10 680 681 CV SpySwitch buffer, prior to incubation with clarified cell lysate on an end-over-end rotator 682 for 1 h at 4 °C, after which resin was washed with 4 × 10 CV SpySwitch buffer. pH-dependent elution was performed by incubating the resin with 6×1.5 CV SpySwitch elution buffer (50 683 684 mM acetic acid, 150 mM NaCl, pH 5.0) for 5 min each, which was immediately neutralized by 685 mixing the flow-through with 6×0.3 CV of 1 M Tris-HCl pH 8.0. Proteins were subsequently dialyzed into PBS pH 7.4. 686 687

688 Polyacrylamide gel electrophoresis

SDS-PAGE was performed using 8-16% polyacrylamide gels in an XCell SureLock system (Thermo Scientific). SDS-PAGE was run at 200 V in 25 mM Tris-HCl pH 8.5, 192 mM glycine, 0.1% (w/v) SDS. Gels were stained with Brilliant Blue G-250 and destained with Milli-Q H₂O prior to imaging on a ChemiDoc XRS+ imager. ImageLab 6.1.0 software (Bio-Rad) was used for densitometric quantification of bands. For imaging on an iBright FL1500 imaging system (Thermo Fisher), analysis was performed using iBright Analysis Software Versions 5.0.1 and 5.2.0 (Thermo Fisher).

696

697 Size-Exclusion Chromatography

Ni-NTA- or SpySwitch-purified protein samples (SpyCatcher003-containing variants, 698 699 SpyTag003-linked non-antigen binding proteins) were injected onto a pre-equilibrated HiLoad 700 16/600 Superdex 200 pg column (GE Healthcare), or a HiLoad 16/600 Superdex 75 pg column 701 (SUMO-SpyTag003 and affibodies; GE Healthcare). Samples were run on an ÄKTA Pure 25 (GE Healthcare) fast protein liquid chromatography machine at 1 mL/min in PBS pH 7.4 at 4 702 703 °C. The absorbance profile of column elutions was recorded at 230 nm, 260 nm, and 280 nm. 704 Each column was calibrated using 660 kDa to 1.35 kDa molecular weight protein standards 705 (thyroglobulin, IgG, ovalbumin, myoglobin, and vitamin B12) (Bio-Rad). Following peak 706 analysis by SDS-PAGE, fractions containing purified protein at the expected molecular weight were concentrated using Vivaspin 20 centrifugal filters (Cytiva) at 4 °C and then stored at -80 707 708 °C.

709

710 **Resin coupling**

SpyCatcher003 S49C was attached to SulfoLink Coupling Resin (Thermo Fisher) according to 711 the manufacturer's instructions, with the following changes: SpyCatcher003 S49C in coupling 712 713 buffer was concentrated to 20 mg/mL using a Vivaspin 20 5 kDa molecular weight cut-off centrifugal filter at 4 °C, reduced with 1 mM tris(2-carboxyethyl)phosphine (TCEP) for 30 min 714 715 at 25 °C, and coupled to SulfoLink Coupling Resin at 20 mg reduced protein per mL of packed 716 resin. Following washes with 1 M NaCl, the coupled resin was washed with TP buffer (25 mM 717 orthophosphoric acid adjusted to pH 7.0 with Tris base) to remove SpyCatcher003 S49C that 718 had bound non-covalently to the resin. The coupled resin was stored in 20% (v/v) ethanol in 719 PBS pH 7.4 at 4 °C for \leq 3 months.

720

721 Isopeptide bond reconstitution reactions

722 All reactions were carried out in triplicate in PBS pH 7.4. 2.5 µM SpyCatcher003 (at 25 °C), 723 Masked SpyCatcher003, or DoubleCatcher variants (each at 37 °C) was reacted with 5 µM SpyTag003-linked protein. At each time-point or end-point indicated, reactions were quenched 724 725 by adding 6 × SDS loading buffer [0.23 M Tris-HCl pH 6.8, 24% (v/v) glycerol, 120 µM 726 bromophenol blue, 0.23 M SDS], with subsequent heating at 95 °C for 5 min in a Bio-Rad 727 C1000 thermal cycler. Reactions were analyzed by SDS-PAGE and the depletion of SpyCatcher003-derived reactant over time was quantified by densitometry. The extent of 728 729 reaction between binding partners at each time point is defined as $100 \times [1 - (SpyCatcher003 -$ 730 derived reactant)/(average density of SpyCatcher003-derived reactant in control lanes)].

731

732 Assembly of dimeric molecules using DoubleCatcher

733 With the exception of reconstitution assays, the assembly of bispecific molecules from two 224 Sector (002) bished metains of interest (Partoin A) and (Partoin B) are a referred in 100 mL

SpyTag(003)-linked proteins of interest, 'Protein A' and 'Protein B', was performed in $100 \,\mu$ L

- total volume per well in 96-well cell culture plates (Thermo Scientific, cat. 167008): 1)
- 736 DoubleCatcher variant with concentration N (here, $N = 1-75 \mu$ M) was first mixed with $1.25 \times N$ Protein A-SpyTag003 for 2 h at 25 °C with shaking at 550 rpm; 2) $5 \times N$ Protein B-

738 SpyTag003 and $1/12 \times N$ MBP-superTEV was added directly to the reaction mixture and 739 incubated for 4 h at 34 °C with shaking at 550 rpm; 3) reaction mixture was incubated with 740 SulfoLink resin-immobilized SpyCatcher003 S49C for 16 h at 4 °C with shaking at 550 rpm, 741 to remove excess unreacted SpyTag003-linked binders from solution. All coupling reactions 742 were carried out in PBS pH 7.4.

743

744 Dynamic Light Scattering

745 Size-exclusion chromatography-purified protein samples were centrifuged for 30 min at 16,900 746 g at 4 °C to remove any aggregates. Samples were filtered through 0.33 mm diameter sterile 747 syringe filters with a 0.22 µm hydrophilic polyvinylidene difluoride (PVDF) membrane 748 (Sigma-Aldrich) and diluted to a final concentration of 25 μ M for protein samples > 35 kDa, or 75 μ M for samples < 35 kDa into sterile filtered PBS pH 7.4. 20 μ L diluted sample was 749 750 loaded into a reusable quartz cuvette and measurements (10 scans of 10 s each) were recorded 751 at 20 °C using an Omnisizer (Viscotek). Data were analyzed in duplicate using OmniSIZE 3.0. 752 For each sample, the radius of hydration (R_h) was plotted with error bars ± 1 s.d.

753

754 Cell Culture

755 SKBR3 cells were from ATCC. BT474 cells were from Cancer Research UK, Lincoln's Inn Fields. SKBR3 cells were grown in complete DMEM: Dulbecco's Modified Eagle Medium-756 757 high glucose (DMEM) supplemented with 10% (v/v) fetal bovine serum (FBS), 100 U/mL 758 penicillin and 100 µg/mL streptomycin (1 × pen/strep; Sigma-Aldrich), 1% (v/v) GlutaMAX 759 (Thermo Fisher Scientific) at 37 °C and 5% (v/v) CO₂. BT474 cells were grown in complete 760 RPMI: RPMI-1640 supplemented with 10% fetal bovine serum (FBS), 100 U/mL penicillin 761 and 100 μ g/mL streptomycin (1 × pen/strep; Sigma-Aldrich), and insulin (5 μ g/mL; Sigma) at 37 °C and 5% (v/v) CO₂. Cells were passaged at 70-80% confluency and were sub-cultured for 762 763 fewer than 3 months. Cell-lines were validated as mycoplasma-negative by PCR.

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765 Metabolic Activity Assay

BT474 or SKBR3 cells were seeded into 96-well plates at 1.25×10^4 cells per well and grown 766 in DMEM supplemented with 0.5% (v/v) FBS, 1% (v/v) GlutaMAX and 50 U/mL penicillin 767 768 and 50 µg/mL streptomycin (0.5 × pen/strep) at 37 °C for 24 h. 100 nM and subsequent serial 769 dilutions of anti-HER2 binders coupled to DoubleCatcher were prepared in 125 µL DMEM 770 with 0.1% (v/v) FBS, 1% (v/v) GlutaMAX and $0.5 \times \text{pen/strep}$. Spent media was removed and protein solutions were added to cells. DoubleCatcher or MBP-superTEV alone were applied to 771 772 cells at the same concentrations as the anti-HER2 binders as a negative control for cell killing. Cells were treated for 96 h at 37 °C with 5% (v/v) CO₂, after which 40 µL 0.15 mg/mL 773 774 Resazurin (Alamar Blue; Sigma-Aldrich) prepared in PBS pH 7.4 and dissolved in PBS and 775 sterile-filtered through a 0.22 µm pore-sized membrane filter was added directly to each well, 776 and plates were incubated for 4 h at 37 °C with 5% (v/v) CO₂. Fluorescence (λ_{ex} 544 nm, λ_{em} 777 590 nm) was measured using a FLUOstar Omega plate reader (BMG Labtech). The 778 fluorescence intensity of media, without cells, containing Resazurin ('background') was 779 subtracted from each measurement. The percent metabolic activity relative to untreated cells 780 was then defined as $100 \times (background-subtracted fluorescence of cells)/(background-$ 781 subtracted fluorescence of untreated cells). Untreated cells are defined as cells grown in 782 DMEM + 1% (v/v) FBS, $0.5 \times \text{pen/strep}$ alone.

783

784 Mass Spectrometry

Intact protein mass spectrometry in positive ion mode was performed using a RapidFire 365 jet-stream electrospray ion source (Agilent) coupled to a 6550 Accurate-Mass Quadrupole Time-of-Flight (Q-TOF) (Agilent) mass spectrometer. 50 μ L protein samples at 10 μ M in PBS

788 pH 7.4 were prepared on a 384-well polypropylene plate (Greiner). Samples were acidified to 1% (v/v) formic acid, before aspiration under vacuum for 0.4 s and loading onto a C4 solid-789 phase extraction cartridge. Following washes with 0.1% (v/v) formic acid in water (1.5 mL/min 790 791 flow rate for 5.5 s), samples were eluted to the Q-TOF detector with deionized water containing 792 85% (v/v) acetonitrile and 0.1% (v/v) formic acid (1.25 mL/min flow rate for 5.5 s). Data were analyzed using Mass Hunter Qualitative Analysis software B.07.00 (Agilent). Expected 793 794 molecular weights for full-length proteins were calculated using the ExPASy ProtParam tool, 795 with the N-terminal fMet (bacterial expression) or signal peptide sequence (mammalian 796 expression) removed. Signal peptide cleavage was predicted by SignalP 6.0 (Nielsen et al., 797 2019). -2 Da was calculated for each predicted disulfide bond. The small amount of +178 Da 798 peak relates to gluconoylation, a common spontaneous post-translational modification for 799 proteins overexpressed in E. coli BL21 (Geoghegan et al., 1999). 800

801 Mass Photometry

Microscope coverslips (24×50 mm, Menzel Gläser) were cleaned by sequential sonication in 802 803 50% (v/v) isopropanol and Milli-Q H₂O (5 min each), dried under a clean nitrogen stream, and 804 assembled into flow chambers with isopropanol/Milli-Q H₂O-rinsed and nitrogen-dried silicone gaskets (6 mm × 1 mm, GBL103280, Grace Bio-Labs) as outlined (Young et al., 2018). 805 Data were acquired on a Refeyn TwoMP mass photometer. Immediately prior to mass 806 807 photometry, protein samples were diluted into freshly prepared and degassed MP sample buffer 808 [PBS pH 7.4, 300 mM NaCl, 0.5% (v/v) glycerol, 1 mM NaN₃, sterile-filtered through a 0.22 809 um pore-size syringe filter (Thermo Fisher)]. A dynamin protein mass standard (MS1000 20-810 40×), diluted into MP sample buffer, was used to calibrate the mass photometer each time a 811 new cover slip was used (Foley et al., 2021). The focal position of the microscope was found 812 in flow chambers containing 20 µL sample, after which 0.5-10 µL of MP sample buffer was 813 replaced with an equivalent volume protein sample, to a final working concentration of 2-50 nM, to account for differences in the dissociation characteristics of different proteins. After \leq 814 815 10 s, images were acquired for 60 s at 331 Hz. Each sample was measured at least twice 816 independently. All data were processed using DiscoverMP v1.2.3 software (Refeyn Ltd) (Foley et al., 2021). Masses were plotted as mass histogram (bin width = 2.5 kDa) and fitted to 817 818 Gaussian non-linear regression curves to identify the mean masses of each protein 819 subpopulation within the heterogeneous sample in GraphPad Prism 9 (GraphPad Software). 820 Expected molecular weights for protein constituents were calculated using the ExPASy ProtParam tool, with the N-terminal fMet (bacterial expression) or signal peptide sequence 821 822 (mammalian expression) removed.

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824 Data Analysis and Graphics Visualization

825 Data visualization and statistical tests were performed using GraphPad Prism 9 (GraphPad 826 Software) and MATLAB R2023a (Mathworks). Protein structures were visualized using 827 PyMOL 2.5.4 (Schrödinger). The structure of HER2 was assembled in PyMOL from PDB IDs 828 1N8Z (Extracellular Domain) (Cho et al., 2003), 2KS1 (transmembrane domain) (Mineev et 829 al., 2010), and 3PP0 (kinase domain) (Aertgeerts et al., 2011) as previously (Goodsell, 2022). 830 Binder structures are based on the following: Tras by PDB ID 1N8Z (Cho et al., 2003); Pert 831 by PDB ID 1S78 (Franklin et al., 2004); 39S by PDB ID 6ATT (Oganesyan et al., 2018); MF3958 by PDB ID 504G (Geuijen et al., 2018); H2-18 by PDB ID 3WLW (Hu et al., 2015); 832 and AffiHER2 by PDB ID 3MZW (Eigenbrot et al., 2010). Protein structures for 833 834 DoubleCatcher variants were predicted using ColabFold versions 1.3 and 1.5.2 (Mirdita et al., 835 2022), and structures for HER2 extracellular domain in complex with HFS2 or nanoHER2 were predicted using ColabFold version 1.4 (Mirdita et al., 2022). Structural predictions were 836 performed using the full-length protein sequence, except for HFS2 where the signal peptide 837

was removed. For all ColabFold-simulated structures, the structure predicted with the highestconfidence is presented.

To simulate vectors that reflect the directionality of SpyTag003, the PyMOL modevectors tool was used to connect Ser57 to His53 and Ser179 to His175 by their carbonyl main-chain oxygens (Law, 2020) (DoubleCatcher numbering, according to sequence listed in Supplementary Fig. 1). The arrowhead represents the N-terminus of the SpyTag003 peptide, such that a binder with a C-terminal SpyTag003 would project from the arrowhead of the vector when it is coupled to DoubleCatcher.

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847 Statistics and reproducibility

848 Statistical significance for metabolic activity assays were calculated by two-way analysis of 849 variance (ANOVA) with Dunnett's correction for multiple comparison. For representative SDS-PAGE (Fig. 1C, E; Fig. 2B, E; Supplementary Fig. 4A, B), observations were confirmed 850 at least once with similar or identical conditions. For Fig. 3 D, E, heterodimer assembly and 851 852 subsequent analyses by mass photometry and SDS-PAGE were repeated at least twice with similar results. For Fig. 4C, D and Fig. 6D, the bispecific assemblies and subsequent metabolic 853 854 activity assays were repeated at least twice independently with similar results. No statistical 855 method was used to predetermine sample size. No data were excluded from the analyses. The 856 experiments were not randomized. The Investigators were not blinded to allocation during 857 experiments and outcome assessment.

858 859 Data availability

Amino acid sequences of Masked SpyCatcher003-containing variants are available in Supplementary Fig. 1. Sequences of other constructs are available in GenBank as described in the section Plasmids and cloning. Plasmids encoding DoubleCatchers and related binders will be deposited in the Addgene repository (https://www.addgene.org/Mark_Howarth/). Further information and request for resources and reagents should be directed to and will be fulfilled by the lead contact, M.H..

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867 868 **References**

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Supplementary Figures

SpyCombinator Assembly of Bispecific Binders

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SpyTag003

RGVPHIVMVDAYKRYK

Masked SpyCatcher003

MSYYHHHHHHHDYDIPTTGAMVTTLSGLSGEQGPSGDMTTEEDSATHIKFSKRDEDGRELAGATMELRDSSGKTISTWIS DGHVKDFYLYPGKYTFVETAAPDGYEVATPIEFTVNEDGQVTVDGEATEGDAGSSGS<mark>ENLYFQG</mark>GGSG**RGVPHIVMVAA** YKRYK**

DoubleCatcher

MSYY<mark>HHHHHH</mark>DYDIPTT<mark>ENLYFQG</mark>GAMVTTLSGLSGEQGPSGDMTTEEDSATHIKFSKRDEDGRELAGATMELRDSSGK TISTWISDGHVKDFYLYPGKYTFVETAAPDGYEVATPIEFTVNEDGQVTVDGEATEGDAHT<mark>GSGGSGGSG</mark>VTTLSGLSG EQGPSGDMTTEEDSATHIKFSKRDEDGRELAGATMELRDSSGKTISTWISDGHVKDFYLYPGKYTFVETAAPDGYEVAT PIEFTVNEDGQVTVDGEATEGDAGSSGS<mark>ENLYFQG</mark>GGSG**RGVPHIVMVAAYKRYK***

DoubleCatcher H-Lock

MSYYHHHHHHDYDIPTT<mark>ENLYFQG</mark>GAMVTTLSGLSGEQGPSGDMTTEEDSATHIKFSKRDEDGRELAGATMELRDSSGK TISTWISDGHVKDFYLYPGKYTFVETAAPDGYEVATPIEFTVNEDGQVTVDGEATEGDAHT<mark>GSPANLKALEAQKQKEQR QAAEELANAKKLKEQLEKGS</mark>VTTLSGLSGEQGPSGDMTTEEDSATHIKFSKRDEDGRELAGATMELRDSSGKTISTWIS DGHVKDFYLYPGKYTFVETAAPDGYEVATPIEFTVNEDGQVTVDGEATEGDAGSSGS<mark>ENLYFQG</mark>GGSG**RGVPHIVMVAA YKRYK***

DoubleCatcher α -Lock

MSYYHHHHHHHDYDIPTT<mark>ENLYFQG</mark>GAMVTTLSGLSGEQGPSGDMTTEEDSATHIKFSKRDEDGRELAGATMELRDS**C**GK TISTWISDGHVKDFYLYPGKYTFVETAAPDGYEVATPIEFTVNEDGQVTVDGEATEGDAHT<mark>GSGGSGGSG</mark>VTTLSGLSG EQGPSGDMTTEEDSATHIKFSKRDEDGRELAGATMELRDS**C**GKTISTWISDGHVKDFYLYPGKYTFVETAAPDGYEVAT PIEFTVNEDGQVTVDGEATEGDAGSSGS<mark>ENLYFQG</mark>GGSG**RGVPHIVMVAAYKRYK***

DoubleCatcher β-Lock

MSYYHHHHHHDYDIPTT<mark>ENLYFQG</mark>GAMVTTCSGLSGEQGPSGDMTTEEDSATHIKFSKRDEDGRELAGATMELRDSSGK TISTWISDGHVKDFYLYPGKYTFVETAAPDGYEVATPIEFTVNEDGQVTVDGEATEGDAHT<mark>GSGGSGGSG</mark>VTTLSGLSG EQGPCGDMTTEEDSATHIKFSKRDEDGRELAGATMELRDSSGKTISTWISDGHVKDFYLYPGKYTFVETAAPDGYEVAT PIEFTVNEDGQVTVDGEATEGDAGSSGS<mark>ENLYFQG</mark>GGSG**RGVPHIVMVAAYKRYK***

DoubleCatcher y-Lock

MSYY<mark>HHHHHH</mark>DYDIPTT<mark>ENLYFQG</mark>GAMVTTLSGLSGEQ**C**PSGDMTTEEDSATHIKFSKRDEDGRELAGATMELRDSSGK TISTWISDGHVKDFYLYPGKYTFVETAAPDGYEVATPIEFTVNEDGQVTVDGEATEGDAHT<mark>GSGGSGGSG</mark>VTTLSGLSG EQGPSGDMTTEEDSATHIKFSKRDEDGRELAGATMELRDSS**C**KTISTWISDGHVKDFYLYPGKYTFVETAAPDGYEVAT PIEFTVNEDGQVTVDGEATEGDAGSSGS<mark>ENLYFQG</mark>GGSG**RGVPHIVMVAAYKRYK***

DoubleCatcher δ -Lock

MSYYHHHHHHHDYDIPTT<mark>ENLYFQG</mark>GAMVTTLSGLSGEQGPSGDMTTEEDSATHIKFSKRDEDGRELAGATMELRDSSGK TISTWISDGHVKDFYLCPGKYTFVETAAPDGYEVATPIEFTVNEDGQVTVDGEATEGDAHT<mark>GSGGSGGSG</mark>VTTLSGLSG EQGPSGDMTTEEDSATHIKFSKRDEDGRELAGATMELRDSSGKTISTWISDGHVKDFYLCPGKYTFVETAAPDGYEVAT PIEFTVNEDGQVTVDGEATEGDAGSSGS<mark>ENLYFQG</mark>GGSG**RGVPHIVMVAAYKRYK***

DoubleCatcher *ɛ*-Lock

MSYYHHHHHHHDYDIPTT<mark>ENLYFQG</mark>GAMVTTLSGLSGEQGPSGDMTTEEDSATHIKFSKRDEDGRELAGATMELRDSSG**C** TISTWISDGHVKDFYLYPGKYTFVETAAPDGYEVATPIEFTVNEDGQVTVDGEATEGDAHT<mark>GSGGSGGSG</mark>VTTLSGLSG EQGPSGDMTTEEDSATHIKFSKRDEDGRELAGATMELRDSSGKTISTWISDG**C**VKDFYLYPGKYTFVETAAPDGYEVAT PIEFTVNEDGQVTVDGEATEGDAGSSGS<mark>ENLYFQG</mark>GGSG**RGVPHIVMVAAYKRYK***

Supplementary Figure 1. Amino acid sequences of SpyTag003, Masked SpyCatcher003, and DoubleCatcher variants. The His₆-tag is shown with gray shading, TEV protease cleavage site with green shading, GSG spacer with cyan shading, cysteines in red, SpyTag003DA mask in bold, and the stop codon with *.



Supplementary Figure 2. Mass spectrometry of building blocks. RapidFire Electrospray Ionization Mass Spectrometry on Masked SpyCatcher, DoubleCatcher variants and each Tagged binder. The observed mass is indicated above the main peak. The expected mass was calculated from ExPASy ProtParam, based on disulfide bonds being formed. The minor +178 Da peak relates to gluconoylation, which is a common post-translational modification for proteins overexpressed in *E. coli* BL21 (Geoghegan et al., 1999).



Supplementary Figure 3. Mass spectrometry of anti-HER2 binders. RapidFire Electrospray Ionization Mass Spectrometry of SpyTag-linked anti-HER2 Fabs, an affibody, and a nanobody, as used to assemble the matrix of bispecifics. The observed mass is indicated above the main peak. Since the hinge region is omitted from the Pert-SpyTag003 heavy chain sequence, there is no interchain disulfide bond in the Pert Fab. The expected mass was calculated from ExPASy ProtParam. The minor +178 Da peak relates to gluconoylation.



Supplementary Figure 4. Removing the linker between Tras Fab and SpyTag003 does not affect reactivity with DoubleCatcher. The tripeptide linker connecting Tras Fab and SpyTag003 was removed to generate Tras NoLink (Tras NL), to reduce the flexibility between the DoubleCatcher and the binder. (A) Time-course of reaction between 2.5 μ M DoubleCatcher and 5 μ M Tras in PBS pH 7.4 at 37 °C for the indicated time, analyzed by SDS-PAGE/Commassie. (B) As in (A) with Tras NL. (C) Quantification of reactivity of Tras or Tras NoLink for DoubleCatcher. Each datapoint is shown (n = 3), with the line connecting the mean.