

A Fluorous-Tagged “Safety Catch” Linker for Preparing Heterocycles by Ring-Closing Metathesis

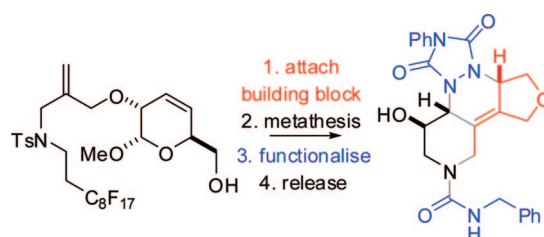
Catherine O’Leary-Steele,[†] Christopher Cordier,[†] Jerome Hayes,[‡]
Stuart Warriner,[†] and Adam Nelson^{*†}

School of Chemistry and Astbury Centre for Structural Molecular Biology, University
of Leeds, Leeds, LS2 9JT, United Kingdom, and Chemical Development,
GlaxoSmithKline, Old Powder Mill, Leigh, Tonbridge,
Kent, TN11 9AN, United Kingdom

a.s.nelson@leeds.ac.uk

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ABSTRACT



A fluorous-tagged “safety catch” linker is described for the synthesis of heterocycles with use of ring-closing metathesis. The linker facilitates the purification of metathesis substrates, the removal of the catalyst, the functionalization of the products, and the release of only metathesis products. The synthesis of a range of heterocycles is described.

Ring-closing metathesis has revolutionized organic synthesis.¹ Ruthenium complexes are particularly functional group tolerant,² but the catalyst residues often need to be scavenged.³ Recently, we developed a fluorous-tagged linker for synthesizing heterocycles by metathesis but a fluorous-tagged catalyst was needed to allow easy product purification.⁴ We now describe a fluorous-tagged “safety catch”⁵ linker that facilitates the synthesis, purification, and func-

tionization of metathesis products without the use a fluorous-tagged catalyst (Scheme 1). We use the term “linker” to describe compounds (e.g., **1**) which are functionalized to yield metathesis substrates (e.g., **2**).

It was envisaged that functionalization of **1** (\rightarrow **2**) would be followed by removal of excess reagents by fluorous-solid phase extraction⁶ (F-SPE). Initiation of a metathesis cascade would be expected at the terminal alkene⁷ of **2** (\rightarrow **3**). Cyclization (\rightarrow **4**) would be followed by a second ring-closing metathesis (\rightarrow **5**) in which a catalytically active methylene complex was regenerated.⁸ Crucially, the product **5** would still be fluorous-tagged; F-SPE would thus allow

[†] University of Leeds.

[‡] GlaxoSmithKline.

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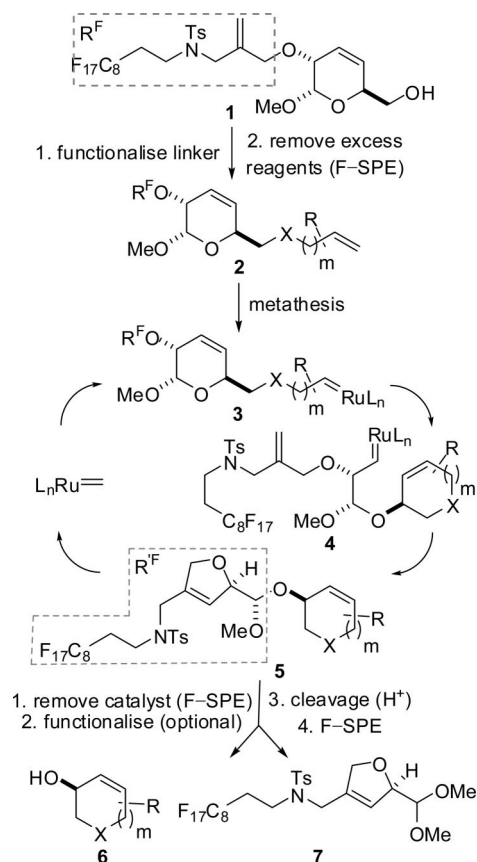
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Scheme 1. Design of the Fluorous-Tagged “Safety Catch” Linker **1**



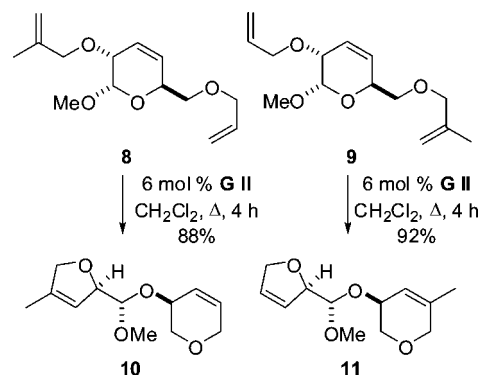
removal of the metathesis catalyst and removal of the excess reagents in subsequent functionalization steps. Finally, acetal cleavage would release only metathesis products (e.g., **6**) (and not unreacted substrates such as **2**) from the fluorous tag. The fluorous-tagged linker **1** was, therefore, designed to be a “safety catch”¹⁵ linker since the cleavage step should release only metathesis products.

To validate the design, we prepared the trienes **8** and **9** from a known glucose derivative (see the Supporting Information). Treatment of **8** and **9** (4 mM in CH₂Cl₂) with 6 mol % Grubbs’s second generation catalyst gave the expected metathesis products **10** and **11** (Scheme 2). Thus, irrespective of the initiation site,⁷ the metathesis cascade proceeded smoothly, cleaving the central dihydropyran ring. The study validated the “safety catch” linker design since hydrolysis of the resulting acyclic acetals would yield the required dihydropyran products.

Scheme 3 describes the synthesis of the linkers **1** and **18**. Reaction of the anion of **12** with ethyl α -bromomethyl acrylate,⁹ and reduction, gave the allylic alcohol **13**. A Fukuyama–Mitsunobu reaction¹⁰ between **13** and the sulfonamide⁴ **14**, and deprotection, gave the fluorous-tagged

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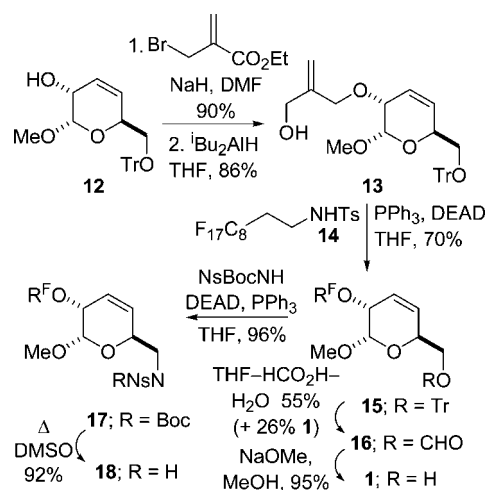
Scheme 2. Validation of the Design of the Linker **1**^a



^a **G II** is Grubbs’s second generation catalyst.

linker **1**. Finally, Fukuyama–Mitsunobu reaction with Ns-BocNH, and deprotection, gave the fluorous-tagged sulfonamide **18**.

Scheme 3. Preparation of the Fluorous-Tagged Linkers **1** and **18**^a



^a For the definition of R^F, see Scheme 1.

The linkers **1** and **18** were functionalized with a range of reactants (see Figure 1, Table 1, and the Supporting Information). Thus, the substrates were prepared by using the Fukuyama–Mitsunobu reaction,¹⁰ allylation, silaketal formation,¹¹ or esterification. In general, the fluorous-tagged products were purified by F-SPE alone, and the purities were determined by HPLC.

The cascade reactions of a range of the metathesis substrates were successful (Table 1). Six- and seven-

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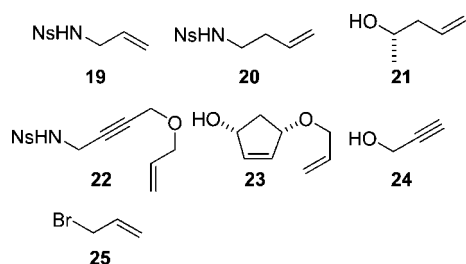
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Table 1. Heterocycle Synthesis by Functionalization of the Linker, Metathesis, and Release (See Scheme 1 for the Definitions of R^F and R^{F'})

entry	linker (reactant, method ^b)	functionalisation product ^b	yield ^c /%	method ^a (catalyst mol %)	yield ^c /%	metathesis product ^b	cleavage method ^a	product [yield ^c / %]
1	1 (19, A)		87 ^d (93 ^e)	B (3 × 5)	90 ^d (94 ^e)		C	26 (R=H) [70]
2	1 (20, A)		98 ^d (92 ^e)	B (3 × 5)	55 (80, ^d 69 ^e)		C	27 (R=H) [77]
3	1 (21, A ^f)		26	B (2 × 5)	98 ^d (85 ^e)		C	28 (R=H) [62] (93, ^d 72 ^e)
4	1 (22, A)		98	B (2 × 5)	41		C	29 (R=H) [67]
5	18 (23, A)		>98 ^d (83 ^e)	B (6 × 5)	87		C	30 (R=H) [35]
6	18 (24, A)		>98 ^d (84 ^e)	B ^g (5)	53 (83, ^d 65 ^e)		C	31 (R=H) [23]
7	1 (25, D)		72 ^d (93 ^e)	B (2 × 5)	51		<i>h</i>	–

^a Method A: reactant (4 equiv), PPh₃ (4 equiv), DEAD (4 equiv), THF, rt then F-SPE. Method B: (i) Hoveyda–Grubbs second generation catalyst, CH₂Cl₂, reflux; (ii) P(CH₂OH)₃, Et₃N then silica; (iii) F-SPE. Method C: 3% TFA in CH₂Cl₂, rt then F-SPE. Method D: (i) NaH, THF, 0 °C; (ii) allyl bromide, rt; (iii) MeOH then F-SPE; ^b See Scheme 1 for the definitions of R^F and R^{F'}. ^c Unless otherwise stated, isolated yield of product. ^d Mass of product after F-SPE. ^e Purity (%) determined by HPLC after F-SPE. ^f 10 equiv of the sulfonamide, PPh₃, and DEAD were used. ^g In the presence of an ethylene atmosphere. ^h Not undertaken.

**Figure 1.** Reactants used to derivatize the linkers **1** and **18**.

membered nitrogen and oxygen heterocycles were formed in good to excellent yield. In the case of the terminal alkyne substrate (entry 6), the reaction was performed under an ethylene atmosphere,¹² and a 53% yield of the fluorine-tagged product **31** (R = R^F) was obtained. More

complex cascade reactions in which two new heterocyclic rings were formed were also successful (entries 4 and 5). Unlike with our previous linker,⁴ it was not possible to prepare eight- or nine-membered heterocycles (see the Supporting Information for the substrates studied); instead, dimerization was competitive with cyclization and, hence, release from the linker. Six metathesis products [**26–31** (R = H)] were released directly from the linker by treatment of the corresponding metathesis products with 3% TFA in CH₂Cl₂ (entries 1–6, Table 1).

The metathesis products could also be functionalized before release from the fluorine tag (see Table 2 and Figure 2). In each case, the excess reagents were removed by F-SPE only. Thus, removal of the *o*-nitrophenylsulfonamide group from **26** (R = R^F), derivatization, and release from

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Table 2. Functionalisation of the Metathesis Products and Release from the Fluorous Tag^a

entry	starting material	purity/%	functionalization method ^b	product	mass recovery ^c /% (purity ^d /%)	cleavage method ^b	product	yield ^e /%
1	26 (R = R ^F)	94	A	33 (R = R ^F)	87 (>90)	B	33 (R = H)	82
2	26 (R = R ^F)	94	C	34 (R = R ^F)		B	34 (R = H)	67 ^f
3	26 (R = R ^F)	>99	E	35 (R = R ^F)		B	35 (R = H)	57 ^f
4	29 (R = R ^F)	>99	D	36 (R = R ^F)	86 (87)	B	36 (R = H)	59
5	36 (R = R ^F)	87	A	37 (R = R ^F)	79 (>95)	B	37 (R = H)	67

^a See Scheme 1 for the definition of R^F. ^b Method A: (i) PhSH, DBU, MeCN; (ii) BnNCO; (iii) F-SPE. Method B: (i) 3% TFA in CH₂Cl₂; (ii) F-SPE. Method C: (i) PhSH, DBU, MeCN; (ii) Ac₂O, pyridine; (iii) F-SPE. Method D: (i) 4-phenyl-[1,2,4]-triazole-3,5-dione, CH₂Cl₂; (ii) F-SPE. Method E: (i) PhSH, DBU, MeCN; (ii) DMAP and isoxazole-5-carbonyl chloride; (iii) F-SPE. ^c Mass of product after F-SPE only. ^d Purity (%) determined by HPLC after F-SPE only. ^e Isolated yield of purified product. ^f Isolated yield of product over 2 steps.

the fluorous tag yielded the tetrahydropyridines **33** (R = H), **34** (R = H), and **35** (R = H) (entries 1–3). Alternatively, the diene **29** (R = R^F) underwent efficient Diels–Alder reaction with 4-phenyl-[1,2,4]-triazole-3,5-

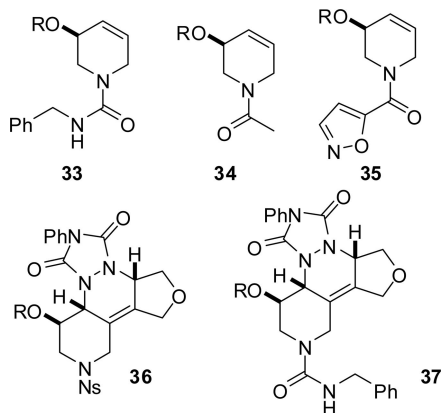
dione to yield **36** (R = R^F): the resulting adduct could either be released directly from the fluorous tag [\rightarrow **36** (R = H), entry 4] or after deprotection and derivatization [\rightarrow **37** (R = H), entry 5].

In summary, we have developed a linker for the synthesis of arrays of heterocyclic products using metathesis cascade reactions. The design of the fluorous-tagged linker allowed (a) easy purification of metathesis substrates; (b) easy removal of the catalyst from the metathesis products; (c) functionalization of the products before release; and (d) the release of only metathesis products.

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Supporting Information Available: Details of all experimental procedures, including unsuccessful metathesis substrates, and NMR spectra for all novel compounds. This material is available free of charge via the Internet at <http://pubs.acs.org>.

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**Figure 2.** Derivatized metathesis products after release from the fluorous tag, R = H.