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Stereoselective synthesis of vinyl nitriles through a Ramberg–Bäcklund approach†

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The applicability of vinyl nitriles in the preparation of pharmaceuticals, polymers, and other valuable materials benefits from robust preparative methodologies. In this work, we present a novel approach for the synthesis of vinyl nitriles based on the Ramberg–Bäcklund olefination reaction (RBR). While there are few examples for accessing functionalized olefins using the RBR, we believe that this methodology embodies useful means for installing privileged vinyl nitrile building blocks, such as the acrylonitrile fragment of the US FDA-approved antiviral rilpivirine.

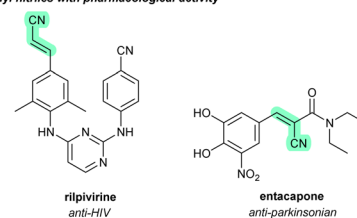
Classic olefination reactions, such as the Wittig,¹ Julia–Kocienski,² Horner–Wadsworth–Emmons,³ and Peterson reactions,⁴ require two complementary reactants with reaction-specific functional groups: a nucleophile (such as phosphorus, sulfonyl, phosphonate, or silicon ylides) and a carbonyl electrophile. In contrast, the Meyer modification⁵ of the Ramberg–Bäcklund reaction (RBR)⁶ only requires a sulfonyl group that is flanked by two activated C–H bonds (such as methyl, methylene, or methine). This is significant as the sulfonyl group (either as such or masked as a less oxidized *S*-functional group) can be brought through a synthetic campaign up to a late-stage.⁷ The stereochemistry of the RBR has been widely discussed and the consensus is that it depends on the strength of the base used. Strong bases lead to the formation of *E* isomers, while weak bases lead to *Z* olefins.⁸ Although there is a relative scarcity of RBRs for the preparation of functionalized olefins (vinyl groups bearing a functional group),⁹ some have recently been utilized for synthesizing valuable species that are difficult to access by other methods.¹⁰ To the best of our knowledge, there are no reports on the preparation of vinyl nitriles using the Ramberg–Bäcklund reaction.

The vinyl nitrile group is a privileged moiety in many pharmaceuticals (Fig. 1a) and features varied synthetic

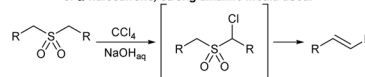
applications.^{11,12} In materials science, polymers based on cyanoacrylates are widely used as adhesives¹³ and support materials.^{14,15} Vinyl nitriles have also been used as building blocks for synthesizing aryl piperidine carbinols, valuable intermediates in the synthesis of paroxetine-like compounds.¹⁶

In recent times, significant attention has been focused on alcohol ammoxidation,¹⁷ the dehydration of oximes¹⁸ and amides,¹⁹ and the Heck-type coupling between acrylonitriles and aryl halides,²⁰ for the synthesis of vinyl nitriles. However, many of these methods, albeit catalytic, rely on transition metal (TM)-based catalysts. While some of these efforts are an improvement over more traditional approaches, as a result of

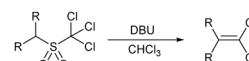
a) Vinyl nitriles with pharmacological activity



b) Meyer, 1969: Ramberg–Bäcklund reaction (RBR) with in situ formation of α -halosulfone, strong alkaline media used.



c) Braverman, 1998: Milder RBR conditions by using DBU



d) This work: Stereoselective synthesis of vinyl nitriles through a RBR under mild conditions.

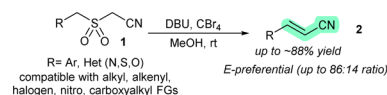


Fig. 1 (a) Vinyl nitriles with pharmacological activity. (b) Meyer modification of the Ramberg–Bäcklund reaction (RBR). (c) Use of DBU for milder conditions in the RBR. (d) This work: vinyl nitrile synthesis from α -benzyl- α' -cyano sulfones using a RBR under mild conditions.

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their catalytic nature and their avoidance of highly toxic species (e.g., cyanide) or multistep sequences, there is a pressing need for reliable methods to access vinyl nitriles with high stereoselectivity. We believe that the RBR has not been extensively explored for synthesizing electron-withdrawing group (EWG)-substituted olefinic products, such as vinyl nitriles. Most RBRs are performed in aqueous, highly alkaline media (Fig. 1b), which limits their applicability in late-stage olefinations due to poor chemoselectivity. This can lead to undesired anionic polymerization on EWG-substituted olefinic products.²¹ A milder approach involves the use of guanidine-type bases in RBRs, enabling higher tolerance for the preparation of functionalized olefins, as exemplified by the syntheses of 1,1-dichloroalkenes by an RBR under anhydrous conditions, achieving yields of up to 70–90% (Fig. 1c).²²

As part of our interest in synthetic methodologies based on sulfur functional groups for advanced synthetic applications,²³ including the generation of olefinic blocks for materials,²⁴ we aimed to develop a robust method for the synthesis of vinyl nitriles using the RBR (Fig. 1d). We used 2-(benzylsulfonyl) acetonitrile (**1a**) as an α -aryl- α' -cyano sulfone probe to investigate RBR-based olefinations. This compound was easily prepared from benzyl mercaptan using an alkylation–oxidation sequence (see the ESI†) and proved to be a convenient choice for our research. The treatment of **1a** with tetrabromomethane (CBr₄) and 1,8-diazabicyclo(5.4.0)undec-7-ene (DBU) delivered, with complete conversion, olefinic products **2a-E** and **2a-Z** (ratio 85 : 15, respectively) in 69% isolated yield after 2.5 h (entry 1, Table 1).

A screening of nitrogenated bases was performed on the same substrate (**1a**), with diisopropylamine (iPr₂NH, DIPa) and diisopropylethylamine (iPr₂NEt, DIPEA) proving productive (entries 3 and 4, Table 1) but not as high yielding as DBU (entry 2, Table 1). We observed no correlation between the screened nitrogenated bases and the olefin stereochemistry in any of the cases studied, as the **2e-Z** product was the prevailing

stereochemical outcome. No clear relationship between base strength and reaction yields was established, although bases with higher pK_a values generally showed better results. As anticipated, neither episulfones nor α -halogenated sulfones were detected.²⁵

Besides CBr₄, additional halogen sources were also screened including tetrachloromethane (CCl₄) (entry 10, Table 1),⁵ *N*-bromosuccinimide (NBS) (entry 11),²⁶ dibromotetrachloroethane (DBTCE) (entry 12),²⁷ and *p*-toluenesulfonyl bromide (*p*-TsBr) (entry 13). While DBTCE, NBS and *p*-TsBr delivered olefinic products (entries 11–13), CBr₄ yielded the best results. Interestingly, no conversion was observed when CCl₄ was used (entry 10).

Under the optimized conditions, we next explored the substrate scope to test functional group tolerance and performance (Table 2). All substrates were easily prepared through the *S*-alkylation of the corresponding thiols or thioacetates with chloroacetonitrile, followed by oxidation using either *m*-chloroperoxybenzoic (mCPBA) acid or Oxone® (see the ESI†). Substrates bearing electron-donating (e.g., *t*-Bu and OMe; **1c** and **1d**) or electron-withdrawing (e.g., NO₂ and CO₂Me; **1e** and **1g**) groups underwent complete olefination, with the latter substrates delivering lower yields, presumably because of an increased propensity to react through undesired anionic polymerization pathways as a result of the increased electrophilicity of the olefinic products (**2e** and **2g**; entries 5 and 7, Table 2). In most cases, a strong stereochemical preference for *E* over *Z* products was observed,²⁸ except for 2-((2-methylbenzyl)sulfonyl)acetonitrile (**1f**, entry 8), which gave poorer stereoselectivity (**2h-E** : **2h-Z** = 73 : 27) and inferior yield (22%). In our case, this result can be rationalized through the plausible conformations of diastereoisomeric adducts of **1** with DBU that precede the corresponding W-type transition states (Scheme 1). The presence of an *o*-substituent such as Me in **1h** leads to an increase of destabilizing repulsive interactions on kinetically preferred conformers **A** and **B** (especially on **B**),²⁹ thus diminishing the *E* : *Z* ratio. An indole-based substrate (**1m**, entry 13) showed a similar outcome, conceivably caused by the *N*-mesyl group albeit more intense. An α,β -unsaturated sulfone (**1i**, entry 9) led to the corresponding olefin **2i**, retaining the original *E*-stereochemistry, but exhibiting a modest *E* preference on the second unsaturation. In the case of sulfone **1j** (entry 10), we were able to couple a β -dehydrobromination³⁰ for the preparation of a 2-en-4-yne nitrile derivative, although this process seems to have impacted the stereoselectivity as the *Z* isomer (**2j-Z**) was obtained preferentially and the product featured partial Br substitution by OMe (see the ESI†). Interestingly, both tertiary α' -ethyl-substituted sulfone (**1n**, entry 14) and an α -benzyl-substituted substrate (**1o**, entry 15) exhibited similar stereoselectivity as in the previous cases. It is important to note that the α -tertiary benzylic substrate **1o** was prepared through dianion-type alkylation, a novel approach when applied to switchable regioselective alkylations of α -benzyl- α' -cyano sulfones (see the ESI†). Substrates with heterocyclic rings such as furyl (**1k**), thiophenyl (**1l**) and indolyl (**1m**) (entries 11–13, Table 2) were suitable for this

Table 1 Screening of nitrogenated bases and electrophilic halogen sources for the RBR synthesis of vinyl nitriles

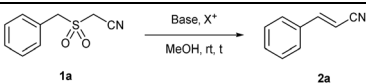
						
Entry	Base	Equiv.	X ⁺ source	Equiv.	Yield %	dr (<i>E</i>) : (<i>Z</i>)
1	DBU	2	CBr ₄	1.1	69	85 : 15
2	DBU	3	CBr ₄	1.1	85	86 : 14
3	iPr ₂ NH	3	CBr ₄	1.1	28	80 : 20
4	iPr ₂ NEt	3	CBr ₄	1.1	24	83 : 17
5	DABCO	3	CBr ₄	1.1	—	—
6	TMP	3	CBr ₄	1.1	—	—
7	Et ₃ N	3	CBr ₄	1.1	—	—
8	Piperidine	3	CBr ₄	1.1	—	—
9	DMAP	3	CBr ₄	1.1	—	—
10	DBU	3	CCl ₄	1.1	—	—
11	DBU	3	NBS	1.1	12	75 : 25
12	DBU	3	DBTCE	1.1	65	85 : 15
13	DBU	3	<i>p</i> -TsBr	1.1	41	83 : 17



Table 2 Substrate scope of the RBR with α -aryl- α' -cyanosulfones **1**. Reagents and conditions: DBU (3.0 eq.), CBr₄ (1.1 eq.), MeOH, rt, 2.5–72 h

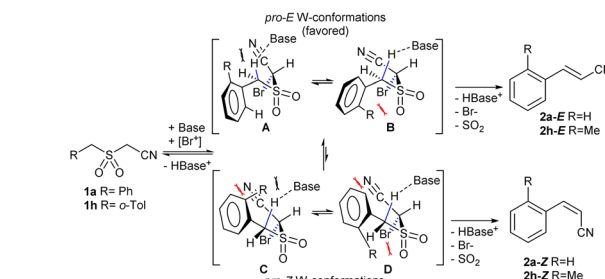


Entry	Sulfone (1)	Vinyl nitrile (2)	Yield %	E/Z ratio
1	1a		85	86 : 14
2	1b		76	83 : 17
3	1c		88	82 : 18
4	1d		61	83 : 17
5	1e		49	82 : 18
6	1f		76	77 : 23
7	1g		49	83 : 17
8	1h		22	73 : 27
9	1i		60	58 : 42
10	1j		48 ^a	31 : 69
11	1k		55	80 : 20
12	1l		58	75 : 25
13	1m		60	54 : 46
14	1n		70	72 : 28
15	1o		88	72 : 28
16	1p		Trace	—
17	1q		—	—

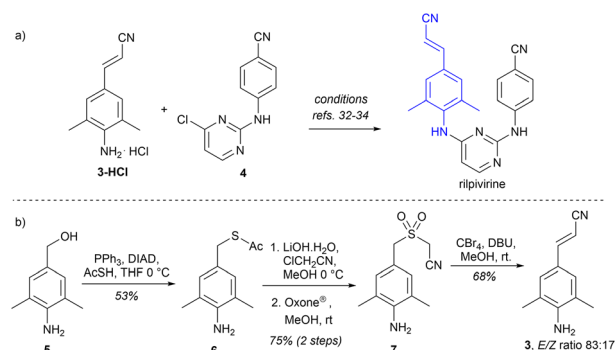
^a ca. 7% of (2E,4E)-4-methoxy-5-phenylpenta-2,4-dienitrile was also detected, see the ESI.†

method and the stereopreference remained *E* selective, except for **1m** (*vide supra*). Unfortunately, α -alkyl substrates (**1p** and **1q**) seemed unreactive under our conditions which can be understood by the relatively low α -acidity (entries 16 and 17).

Rilpivirine (EDURANTTM, Fig 1a, Scheme 2a) is an antiviral agent against human immunodeficiency virus 1 (HIV-1), approved by the US FDA in 2011.³¹ Strategically, rilpivirine synthesis relies on the availability of *E*-3-(4-amino-3,5-dimethylphenyl)acrylonitrile (**3**), which is coupled to 4-((4-chloropyrimidin-2-yl)amino)benzonitrile (**4**) (Scheme 2a). A patent study describes the synthesis of **3** starting with a Pd(OAc)₂/P(*o*Tol)₃-



Scheme 1 Plausible conformers preceding W-planar transition states leading to stereoisomers **2-E** and **2-Z**; if *o*-substituent R = Me (**1h**), repulsive interactions increase for A–B conformers leading to a decrease of the *E* : *Z* ratio. Red marks denote significant steric strain.

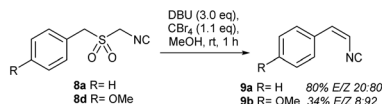


Scheme 2 (a) Assembly of rilpivirine by coupling acrylonitrile **3** and chloropyrimidine **4**. (b) A transition metal-free synthesis of *E*-3-(4-amino-3,5-dimethylphenyl)acrylonitrile **3**, a strategic intermediary for the synthesis of rilpivirine.

catalyzed coupling,³² while another, more recent, case uses a Heck reaction catalyzed by Pd/C.³³ Both routes require the use of catalysts based on transition metals (*e.g.*, Pd(OAc)₂, P(*o*Tol)₃, and Pd/C), which are not ideal for industrial setups.³⁴ Our approach to access arylacrylonitriles through the RBR seemed ideal for the development of a transition metal-free formal synthesis of rilpivirine through the preparation of **3** (Scheme 2b). Masked benzylmercaptane **6** was obtained from aminoalcohol **5** by a *thio*-Mitsunobu reaction, followed by tandem acetyl cleavage/S-alkylation using chloroacetonitrile under basic media, and oxidation with Oxone® to deliver sulfone precursor **7**. Finally, the RBR method applied on **7** cleanly delivered acrylonitrile **3** with 68% olefination yield in an 83 : 17 *E* : *Z* ratio. Remarkably, no *N*-protecting group was needed in the sequence. Efforts on the optimization of the efficiency and stereoselectivity enhancement towards *E*-**3** are currently underway in our laboratory.

Submission of an isocyanide analog of **1a** ((isocyanomethyl)sulfonyl)methyl benzene (**8a**)^{23a} delivered olefinic products in comparable yields; however, the stereoselectivity was completely reversed, as the **9a-E** : **9a-Z** ratio was 20 : 80 (Scheme 3). This stereoselectivity “switch” was also revealed for 4-methoxy isocyanide **9d**, supporting a stereodivergency phenomenon triggered by the isocyanide group.³⁵





Scheme 3 Z-Preferential RBR-based olefination of α -benzyl- α' -isocyano sulfones **8a** and **8d**.

Conclusions

We have developed a method for the RBR-based olefination of α -benzyl- α' -cyano sulfones (**1**) with a preference towards *E* stereoisomeric acrylonitriles (*E*-**2**). Our approach is tolerant to substrates bearing both electron-donating and electron-withdrawing groups but limited towards 2-substituted (aryl) substrates that delivered stereoselectivity loss, plausibly caused by steric strain. Heterocycles in general were also suitable for olefination, whereas fully aliphatic sulfones could not be transformed satisfactorily. The method was applied to a transition metal-free synthesis of *E*-3-(4-amino-3,5-dimethylphenyl) acrylonitrile (**3**), a strategic block used for the synthesis of the antiviral rilpivirine. The extrapolation of the olefination method to analogous selected α -benzyl- α' -isocyano sulfones (**8a** and **8d**) featured a stereochemical reversal as *Z* isomers were preferred. We are currently optimizing the route to acrylonitrile **3**, and a comprehensive study regarding the synthetic development of the RBR-based olefination of α -benzyl- α' -isocyano sulfones will be released in the near future.

Conflicts of interest

There are no conflicts to declare.

Acknowledgements

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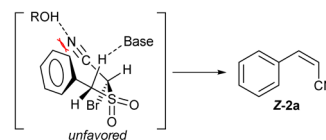
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"weak" H—NC (cyano) bonding $\rightleftharpoons$ Ph - CN steric strain prevails



vs.

"strong" H—NC (isocyano) bonding $\rightleftharpoons$ π - π bonding interaction

