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A Broad-Spectrum Synthesis of Tetravinylethylenes

Kelsey L. Horvath,^[a] Christopher G. Newton,^[a] Kimberley A. Roper,^[a] Jas S. Ward^[a] and Michael S. Sherburn*^[a]

Abstract: The first general synthesis of compounds of the tetravinylethylene (TVE) family is reported. Ramirez-type dibromoolefination of readily accessible penta-1,4-dien-3-ones generates 3,3-dibromo[3]dendralenes, which undergo twofold Negishi, Suzuki-Miyaura or Mizoroki-Heck reactions with a wide variety of olefinic coupling partners. This route delivers a broad range of unsymmetrically-substituted tetravinylethylenes with up to three different alkenyl substituents attached to the central C=C bond. The extensive scope of the approach is demonstrated by the preparation of the first higher order oligo-alkenic through-conjugated/cross-conjugated hybrid compounds. An unsymmetrically-substituted TVE is shown to undergo a domino electrocyclization-cycloaddition with high site-selectivity and diastereoselectivity, thereby demonstrating the substantial synthetic potential of substituted TVEs for controlled, rapid structural complexity generation.

Tetravinylethylene (TVE) is one of the smallest structures exhibiting both through-conjugation and cross-conjugation (Scheme 1). Specifically, the structure comprises intersecting pairs of (E)-1,3,5-hexatrienes, (Z)-1,3,5-hexatrienes and 2methylene-1,4-pentadienes ([3]dendralenes). Nothing is known about the impact of substitution upon the behavior of TVEs and how this might be exploited. Furthermore, related structures containing more than five C=C units have not been reported in the literature. The first synthesis of TVE, a landmark contribution from the classical era of unsaturated hydrocarbon chemistry, was reported by Skattebøl and co-workers in the 1960s.¹⁻³ We recently disclosed the first preparatively useful approach to TVE, a one-step multi-gram scale synthesis, and demonstrated that the hydrocarbon is bench-stable when stored as a neat liquid.^{4,5} The one-step synthesis of TVE (Scheme 1, 1a) utilized a Pd(0)catalyzed fourfold Stille cross-coupling sequence between tetrachloro-ethylene 2 and vinyltributylstannane 3a. In addition to the parent TVE, five symmetrically-substituted TVEs 1b were similarly prepared using substituted alkenyltributylstannanes 3b in the exhaustive Stille cross-coupling process.4,5

As a starting point for future synthetic applications, the parent TVE has been shown to rapidly generate multi-cyclic systems.^{4,5} Additionally, the hybrid through/cross-conjugated structure of the TVE unit represents the smallest organic four-directional branch point in a possible conducting molecular wire.⁶ To realize the potential of the TVE family in target synthesis and materials applications, synthetic access to unsymmetrically-substituted systems must be granted. Until now, no unsymmetrically-substituted TVEs have been reported. Our previous synthetic approach (Scheme 1), in common with all other substituted TVE

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syntheses reported in the literature,⁷ is only suited to the attachment of four identical substituted alkenyl-groups to the central C=C core.



Scheme 1. The existing synthetic route to TVEs with its limitations, and the new, broad scope synthesis.

Herein we report a broad-reaching solution, delivering the first 29 unsymmetrically substituted TVEs carrying up to three different substituents (Scheme 1, bottom). We also report the first compounds exhibiting extended π -systems based upon TVE subunits. Moreover, we establish the synthetic potential of these unsymmetrically-substituted TVEs, by way of a one-pot, siteselective and diastereoselective domino pericyclic sequence that generates three C-C bonds and six contiguous stereocenters. The new synthetic approach (Scheme 1, bottom) involves the conversion of penta-1,4-dien-3-ones 4 (readily accessed through into twofold aldol-type condensations) 3.3-dibromo-[3]dendralenes 5, which in turn undergo a range of twofold cross-couplings to furnish diversely-substituted tetravinylethylenes 6.

Twelve substituted penta-1,4-dien-3-ones **4** were prepared, by way of sequential Claisen-Schmidt condensations⁸ (see SI for details). There are only three examples of Ramirez dibromoolefinations of penta-1,4-dien-3-ones **4** in the literature⁹⁻¹¹ and, in our hands, low yields were obtained under standard Ramirez reaction conditions.¹² Gratifyingly, in the majority of cases, the modified protocol introduced by Lautens and co-workers¹³

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furnished workable yields, and twelve substituted 3,3-dibromo[3]dendralenes¹⁴ **5** were prepared in this manner (Table 1).





 Table 2. Twofold Negishi cross-coupling of 3,3-dibromo[3]dendralenes 5 with

 vinylzinc bromide furnishes the first unsymmetrical tetravinylethylenes 6.

3,3-Dibromo[3]dendralenes comprising the same (**5a-g**) or different (**5h-I**) alkenyl substituents performed equally well, to generate the first examples of TVEs possessing two or three different alkenyl groups about the central C=C bond. TVEs bearing carbocyclic aromatic substituents with different electronic characteristics (**6j-I**) and the first TVEs carrying heterocyclic aromatic substituents (**6f-i**) are included in this selection.

Twofold cross-couplings of 3,3-dibromo[3]dendralenes **5** are by no means limited to the introduction of a pair of unsubstituted vinyl groups: in fact, alkenyl groups bearing substituents at any position are tolerated. The syntheses of eighteen of these more highly substituted TVEs are summarized in Table 3. Double Negishi couplings of 3,3-dibromo[3]dendralene **5a** with 2propenyl-, (*E*)-styrenyl- and (2-methylprop-1-en-1-yl)zinc bromides furnished six novel tetra- and hexa-substituted TVEs **6n-s**. Cross-couplings other than Negishi reactions are also successful, as demonstrated by the products of double Suzuki-Miyaura¹⁷ and Mizoroki-Heck¹⁸ processes, **6t-v** (3 examples)

 Table 1. Lautens-modified Ramirez dibromo-olefination of penta-1,4-dien-3-ones 4 permits the first general synthesis of 3,3-dibromo[3]dendralenes 5.

Unlike some other dendralenic structures, which exhibit a propensity to dimerize and polymerize,¹⁵ 3,3-dibromo[3]dendralenes **5** are generally white or yellow crystalline solids that can be stored neat without appreciable decomposition, and are easily handled using standard laboratory techniques.

While there are many examples of cross-coupling reactions of 1,1-dibromoalkenes in the literature,¹⁶ there is only one example of a cross-coupling of a 3,3-dibromo[3]dendralene, involving the Sonogashira sp^2 -sp coupling of a cyclic system with trimethylsilylacetylene.¹⁰ In light of the lack of precedent, both for sp^2 - sp^2 cross-couplings and also for reactions of acyclic systems, we were delighted to find that twofold Negishi reactions of 3,3-dibromo[3]dendralenes **5** with vinylzinc bromide proceed well using [Pd(dppf)Cl₂] as pre-catalyst to form the first unsymmetrical TVEs **6** (Table 2).

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and **6w-ae** (9 examples), respectively. These results establish the wide scope of the 3,3-dibromo[3]dendralene cross-coupling approach to TVE synthesis and its broad tolerance of substitution upon the incoming alkenyl-groups.



 Table 3. Twofold Negishi, Suzuki-Miyaura or Mizoroki-Heck reactions grant access to unsymmetrical, multi-substituted tetravinylethylenes. ^aX-ray crystal structures of these four TVEs²² are in the SI.

Following the trend previously seen,^{4,5} the 25 new TVEs depicted in Tables 2 and 3 are bench-stable substances. No special equipment or methods are required when manipulating these compounds: they survive storage neat at ambient temperature for several weeks. The lack of susceptibility of this new family of π -bond rich systems towards uncatalyzed Diels-Alder dimerization and autoxidation (typical of related cross-conjugated¹⁵ and through-conjugated¹⁹ polyenic systems) bodes well for future applications.

Access to unsymmetrically-substituted TVEs promotes new opportunities in rapid structure complexity generation, and an interesting question of diene site selection in [4+2]cycloadditions of polyenes (Scheme 2). Heating diphenyl-di-(1-naphthyl)-TVE 6aa in benzene brings about a disrotatory electrocyclization to generate cyclic [4]dendralene¹⁴ 7, with a *cis*-disposition between the two cyclohexadiene substituents. When performed in the presence of dienophile *N-tert*-butylmaleimide **8**, a subsequent endo-mode cycloaddition occurs, with approach to the less sterically encumbered π -diastereoface of [4]dendralene 7. The two non-equivalent semi-cyclic dienes of [4]dendralene 7 (highlighted in different colors in Scheme 2) are expected to be more reactive than the cyclohexadiene site,⁵ and the phenylsubstituted diene is preferred over the 1-naphthyl-substituted site. Overall, this operationally trivial domino sequence generates densely-functionalized decalin 9, with three new C-C bonds, six contiguous stereocenters, and a cyclic [3]dendralene subunit that invites additional synthetic manipulation.²⁰



Scheme 2. Stereoselective domino $[6\pi]$ electrocyclization/[4+2]cycloaddition of TVE **6ad** proceeds with diene site selectivity to generate multicyclic product **9**. H atoms not attached to stereocenters are omitted from the X-ray crystal structure^{22,23} for clarity.

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Scheme 3. First preparation of higher order acyclic C=C-based π -frameworks related to TVE. The new π -systems are highlighted in green. Ethyl groups are truncated in the X-ray crystal structure^{22,23} for clarity.

The robust nature of the new pentaene-based TVE compounds, combined with their ease of synthesis, prompted the question of whether larger structures with unprecedented acyclic π frameworks were within reach. The successful extension of these methods to the syntheses of hexaenes 14 and 15, and heptaenes 13, 16 and 17 are depicted in Scheme 3. Dibromotetraene 10, dibromo-pentaene 11 and tetrabromo-triene 12, prepared through Lautens-modified Ramirez dibromo-olefination reactions of commercially available ketonic precursors (see SI), underwent twofold or fourfold Negishi or Mizoroki-Heck couplings to furnish the bench-stable polyenic products. The UV absorption spectrum of heptaene 17 exhibits features attributable to the longest through-conjugated subunit (see SI). The solid state molecular structure of tetra-ester 17, obtained from single crystal X-ray analysis, is consistent with this observation, revealing an essentially in-plane 2,4,6,8,10dodecapentaenoate unit (Scheme 3). Acyclic, hybrid crossconjugated/through-conjugated C=C structures are very poorly represented in the literature.²¹ The structures depicted in Scheme 3 represent the first examples of substances of this type.

In summary, we have described the first general synthetic approach to TVEs and related through/cross-conjugated acyclic C=C-based systems. This work reports 34 new TVEs and related structures, including the first examples exhibiting unsymmetrical substitution patterns, extended throughconjugation and/or cross-conjugation. The synthetic approach is robust, involving twofold Negishi, Suzuki-Miyaura or Mizoroki-Heck couplings of acyclic 3,3-dibromo[3]dendralenes. The new structures are bench stable compounds that undergo operationally simple, selective, complexity-generating domino sequences. The acyclic C=C-bond-rich structures described herein are representatives of a sizeable region of structural space that was previously inaccessible but is now available for exploration and investigation. Applications in target synthesis and in conducting materials will follow.

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Keywords: hydrocarbons • polyenes • cross-coupling • electrocyclic reactions • cycloaddition

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Master key for TVE. The first general synthesis of tetravinylethylenes (TVEs) involves the preparation and cross-coupling of dibromo-[3]dendralenes. Their value in controlled, rapid structural complexity generation is demonstrated. The first higher order acyclic oligo-alkenic throughconjugated/cross-conjugated hybrid compounds are also reported.



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