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We report herein a highly \textit{exo}-selective intramolecular Diels-Alder reaction of alkenyl boronates which employs an N-B dative bond-involved bicyclic rigid tether. Complex C(\textit{sp}^3)-rich polycyclic molecules containing up to 8 stereocenters can be readily formed via an operationally simple two-step procedure.

The Diels-Alder reaction, constructing six-membered rings by stereospecific \([4+2]\) cycloadditions, has shown unparalleled efficiency in the field of chemical synthesis, especially of natural products.\(^1,2\) Still, however, many complex biologically important molecules, such as the ones shown in Figure 1,\(^3\) contain highly functionalized six-membered rings that may not be directly disconnected by conventional \([4+2]\) mode due to the presence of multiple oxygen and/or nitrogen substituents. One possible solution to this problem would be using a convenient precursor that can be readily transformed to the desired functional groups. Boron-substituted building blocks therefore provide an attractive strategy because C-B bonds may be reliably installed before and converted into C-O, C-N or C-C bonds after the Diels-Alder reaction. Alkenyl boronates were first used in \([4+2]\) cycloadditions more than five decades ago by Matteson\(^4\) and Woods.\(^5\) Later on, various alkenylboron reagents including dialkyl boranes,\(^6\) boron dihalides,\(^7\) boronic acids and boronates\(^8\) have been studied and a review has summarized these important progresses.\(^9\) Recently, this strategy has been successfully applied in the total synthesis of pancratistatin and lycorine.\(^10\) Despite the mentioned significant advancement, in many cases, the reactions suffered from low reactivity, low \textit{endo}/\textit{exo} selectivity and/or low functional group tolerance. During our studies on new organoboron reagents,\(^11\) we have been interested in the stereoselective cycloadditions of alkenyl boronic acids taking advantage of new masking structures on boron atoms. Herein, we wish to report an exclusively \textit{exo}-selective intramolecular Diels-Alder reaction with broad substrate scope.

\begin{figure}[h]
\centering
\includegraphics[width=\textwidth]{figure1.png}
\caption{Selected examples of six-membered carbon rings in natural products.}
\end{figure}

MIDA (N-methyliminodiacetic acid) boronates\(^12\) and diethanolamineboronates\(^13\) are relatively stable surrogates of boronic acids and the former, particularly, have shown great potential in chemoselective Suzuki-Miyaura coupling.\(^11a,12,14\) Structurally, these compounds include a rigid [3.3.0] bicyclic moiety where the bridgehead nitrogen and boron atoms are connected by an N to B dative bond.\(^15\) Thus, the rest two substituents on the N and B atoms (R and Y, Scheme 1a) were placed as cis relationship with little conformational flexibility. We therefore wondered if such a structural preregionization might be employed in intramolecular Diels-Alder reaction to promote the reaction of electronically unactivated substrates and to induce not only high regioselectivity but also high stereoselectivity. To forge this well-defined framework in a highly modular manner, we planned to condense an alkenyl boronic acid and an aminodiacetic or an aminodiethanol on which a diene is preinstalled (Scheme 1b).
We firstly chose to use an iminodiacetic acid tether, i.e. where X is O in Scheme 1b. The diene-incorporated iminodiacetic acid 1 and the alkenyl boronic acids 2 were readily prepared by routine methods (for experimental details, see ESI). We then tested the design by a two-step process involving intermolecular dehydrative condensation and intramolecular cycloaddition. Thus, heating a mixture of (E)-styril boronic acid (2a, 1.0 equiv.) and N-dienyliminodiacetic acid (1, 1.1 equiv.) in toluene and DMSO (10:1 v/v) containing 4 Å molecular sieves at 80 °C cleanly gave the desired MIDA boronate-type compound 3a, as evidenced by the characteristic AB-type -CH₂- signals in ¹H NMR spectrum of the crude product (chemical shift 4.06, 4.16 ppm, coupling constant 16.8 Hz). This crude compound can be used without further purification for the key [4+2] cycloaddition reaction. Thus, simply heating a solution of 3a in 1,2-dichlorobenzene at 160 °C for 8 hours led to full conversion, affording a single observable isomer according to ¹H NMR analysis of the crude product. After purification by flash chromatography on silica gel, extensive NMR experiments including ¹H-¹H COSY, HSQC and HMBC supported an exo-[4+2] cycloaddition product 4a (64% yield over two steps based on (E)-styril boronic acid). The structure was further confirmed by X-ray diffraction analysis of a single crystal of 4a (Figure 2). Notably, this compact cage-like hetero-propellane structure contains five rings and four of them are annulated non-aromatic rings.

Notably, this compact cage-like hetero-propellane structure contains five rings and four of them are annulated non-aromatic rings.

We next explored the scope of this tethered DA reaction. The results are shown in Scheme 2. Remarkably, in all cases, only one stereoisomer was observed, in sharp contrast to previous DA reactions of alkenyl boronates where a mixture of endo and exo products were usually formed. Various substituents on the phenyl ring (Me, OMe, SMe, F, Cl and Br) were well tolerated, affording the respective cycloaddition products (4b-4i) in two steps in good to excellent yields. Thio-phenyl was also compatible in this D-A approach (4j), giving the product in 80% yield. In addition, aliphatic alkenyl boronic acids could be successfully used in this process and the products were isolated in high yields and high exo-selectivity (4k and 4l). These results suggest that the generally excellent stereocontrol is governed by the N-B dative bond-induced rigid [3.3.0] bicyclic moiety. Alkenyl boronic acids bearing electron-withdrawing groups are often prone to decomposition due to protodeboronation, thus posing challenges for further transformations. To our satisfaction, β-carbonyl alkenyl boronic acids were also viable in our system, although only moderate yields were obtained (4m-4o). Interestingly, in the case of 4m, D-A reaction occurred during the dehydrative condensation process. Therefore, in a single step, the final products were readily prepared.

Encouraged by the above results, we extended the process to DABO boronate-like substrates. Literature works have indicated that the DABO structures are less rigid than the MIDA counterparts. In this work, nevertheless, we were able to employ the DABO-type N-B coordination tether for successful intramolecular [4+2] reaction. In event, the dehydration step could be smoothly done under mild conditions. Thus, the two components (5 and 2, Scheme 3) were stirred with anhydrous magnesium sulphate in dichloromethane at ambient temperature to complete the condensation step. After removal of the solids by filtration, the crude product was immediately used in the intramolecular [4+2] reaction with conditions similar to the MIDA-like cases. Again, only the exo-stereoisomer was obtained. The
substrate scope was briefly surveyed. In all cases, very good exo-selectivity was achieved. Substituents such as methyl, methoxy, methymercapto, nitro and chlorine groups on the phenyl rings (7b–7f), and a phthalamide group (7g) were all tolerated and the corresponding cyloaddition products were isolated in moderate to good yields. It is worth noting that cyclohexene-3-boronic acid (2a), which had led to low yield of the product (4a) in the MIDA-type case, now served as an efficient dienophile, giving D-A product 7h in good yield. This outcome might be attributable to the mild condensation conditions that prevent decomposition of 2a.

was thus identified as 7l (shown in Scheme 4) by X-ray diffraction analysis. The other isomer was therefore deduced to be 7l'. Remarkably, in this operationally simple two-step procedure, complex sp²-rich polycyclic molecules containing up to 8 stereocenters were readily formed.

In order to understand the origin of the exceptionally high exo/endo-selectivity in this tethered Intramolecular [4+2] reaction, density functional theory (DFT) calculations were performed at the M06-2X/6-31G* level of theory (Scheme 5). The Intramolecular [4+2] reaction of 3a was selected as the model reaction. TSex and TSend represent the transition states of exo-selective and endo-selective reaction channel respectively. Compared with the lowest-lying conformation of 3a, the gas phase activation free energy barrier for TSend is 26.1 kcal/mol, and TSex is located 3.2 kcal/mol higher than TSend. This energy difference indicates the Intramolecular [4+2] reaction to be highly exo-selective, in consistence with the experimental results.

Scheme 5. Computational study of the endo/exo selectivity of 3a.

Conclusions

In summary, a highly exo-selective intramolecular Ueis-Alder reaction has been developed which utilizes an N-B dative bond induced (See, u.) bicyclic boronate unit as the structurally well-defined tether. In an operationally simple two-step process, various readily available alkanyl boronic acids were successfully used as the dienophile, affording the [4+2] cycladdition products in high yields and stereoselectivity. We are currently studying to extend this reaction for asymmetric synthesis of complex biologically molecules.

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References

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