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A feasible zeolite-assisted ortho C-H nitration of nitrostilbenes has been developed for the first time, which can be used in situ acetyl nitrate as a mild, easily handled and commercially available nitrating reagent, leading to the synthesis of polynitrostilbenes with double selectivities and in good yields.

Nitrostilbenes, such as 2,2',4,4',6,6'-hexanitrostilbene (HNS), are well known high-energy materials with excellent thermally stability and detonation properties. Furthermore, nitrostilbenes based derivatives can be found in various natural products, which play an increasingly prominent role in the area of non-linear optical (NLO) application. In energetic materials chemistry, the presence of nitro groups tends to decrease the heat of formation but contributes markedly to the overall energetic performance. There is a lot of interests and challenges in the syntheses and chemistry of energetic materials containing C=C unsaturated double bonds together with more nitro groups attached in the molecule. However, traditional nitration conditions are limited in scope due to poor regioselectivity, poor chemoselectivity arising from overnitration, a lack of compatibility with easily oxidized functionalities, result in the complexity of the mixture of products formed. Acetyl nitrate in acetic anhydride solution as typical electrophilic reagent reacted rapidly with many alkenes to give mixtures of nitro-alkenes, β-nitro nitrates, and β-nitro acetates. For example, nitration of styrenes and stilbenes with acetyl nitrate gave 40-70% yields of β-nitro acetates. [Eq. (1), Scheme 1] Recently, Maiti and coworkers reported the preparation of (E)-nitroolefins from nitration of olefins using AgNO₃/TEMPO, t-BuONO/TEMPO, Fe(NO₃)₃/TEMPO. These methods while offering significant improvements in direct nitration of olefins, problematic issues include metal reagent employed as well as the absence of recovery and reuse of the catalyst.

The use of solid catalysts is potentially attractive because of the ease of recovery and reuse of the catalyst, with addition possibilities that the solid might enhance the selectivity, particularly if it is a zeolite. Our initial successes in regioselective nitration of simple aromatics or naphthalene over zeolite by Kyodai nitration promote us towards the direction. Improved para-regioselective for single ring aromatics has also been achieved with zeolites as the catalyst, particularly when nitric acid is used in combination with acid anhydride. [Eq. (2), Scheme 1] Although this method is effective, a situation may arise where one needs to introduce a nitro group through nitration of substrates with -C=C- unsaturated double bonds as well as aromatic rings. Further, nitration of nitrostilbenes exhibits a high degree of selectivity including regioselective and chemoselectivity towards aryl rings and -C=C- double bond. Herein we report a zeolite-assisted double selective nitration method by which nitrostilbenes can be converted to their ortho-nitrated derivatives in high yield using in situ acetyl nitrate as a mild, easily handled and commercially available nitrating reagent. [Eq. (3), Scheme 1]

Our initial nitration of trans,trans-1,3-distyryl-2,4,6-trinitrobenzene using HNO₃/H₂SO₄, and a complex mixture of products was observed, which are mainly consisted of oxidation products. (Table S1, entry 1) However, when treated with HNO₃/Ac₂O, an isolated ortho-nitrated product 1 along with a wide product distributions were observed. (Table S1, entries 2-4) With the presence of zeolites (SiO₂/Al₂O₃ = 25) in HNO₃/Ac₂O, an exciting result was obtained. By further optimizing the amount of nitration reagent, catalyst species and loading amount, we discovered that 1 equivs of HNO₃, 0.20 g of zeolite in excess acetic anhydride at room temperature in 5
min could provide 95% isolated yield of 1. (Table S1, entry 4) Remarkably, running the nitration with NO₂O₂ or NO₃O₃ had no positive effect on the outcome of the reaction. (Table S1, entries 12-13) The –C=C– double bonds in nitration products disappeared, which might be attributed to oxidation in the double bonds.

With the fully optimized reaction conditions, we explored the scope and limitations of the method. Generally, all the nitration products were obtained in a good isolated yield. (Table 1) (E)-1,3,5-trinitrostilbene gave excellent yields of the desired nitro product (2j, 93%) under the optimized conditions. High regioselectivity for the less hindered ortho C-H bond was observed for substrates bearing a para chloro (2b) and bromo (product 2i) or meta chloro group (2e and 2k). In contrast, when another chloro group was introduced in ortho position (2d), the major product isolated with the same regioselectivity but with low yields. However, if the sterically challenging ortho substituted nitrostilbenes derivatives as substrates often provided complex mixtures under usual conditions. Para-substituents with –CH₃ or –OCH₃ reacted smoothly to give the expected nitro products (2e and 2f). While, when (E)-2-(4-methylstyryl)-1,3,5-trinitrobenzene as substrate, small amounts of Z-isomer was detected (2n). Notably, if the -OCH₃ group appeared in meta-position, di-nitration products were isolated in good yields (2g and 2m). Strong electron-withdrawing substituent with 4-nitro also gave a nitro product (2h), albeit in lower yield. Unfortunately, 3-nitro nitrostilbenes (e.g. (E)-1,3,5-trinitro-2-(3-nitrostyryl) benzene) did not form the desired nitration product under the reaction conditions. Attempting the reaction on another type of nitrostilbenes, which were found to be problematic under traditional conditions, resulted in successful nitration in excellent yields (2o and 2p). The regio-chemical designation of the products (2e and 2e) was unequivocally proven by single crystal X-ray diffraction.

In order to check the possibility of reuse of the zeolite, it was recovered following extraction of the products and was regenerated by heating overnight in air at 450 °C. Nitration reactions were then conducted under identical conditions using the recovered zeolite. The results showed that under the standard conditions, there was only a slight decrease (to 93%) in the yield of 2a even after using the same zeolite six times under identical conditions. To demonstrate the scalability of the selective nitration method, 3,3′-(1,1′-E)-(2,4,6-trinitro-1,3-phenylene)bis-(ethene-2,1-diyli)bis(chlorobenzene) was reacted in a 15 mmol scale. In this experiment, the desired nitrostilbene product (2e) was separated in excellent yield (90%).

Nitrination by acetyl nitrate with high ortho-para ratios result from “ortho effect” on steric ground or coordination mechanism. Prins applied liquid and solid state MAS NMR to Ac₂O/HNO₃ liquid and in beta zeolite and found acetyl nitrate and acetic acid which were coordinated to the framework aluminium of the zeolite. The Bronsted acid site in the zeolite wall are responsible for the generation of NO₃⁻ ion from acetyl nitrate during electrophilic attack on substrates. It is further supported by the results that when increasing the Si/Al ratio in beta zeolite, a decrease in ortho selectivity is observed. (Table S1, entries 6, 8 and 9) Also, the higher selectivity for the zeolite may be ascribed to the 3D structure of zeolite in comparison to the 2D structure of K10. (Table S1, entries 6 and 11)

Molecular electrostatic potential (ESP) was employed in the explanation of chemoselectivity, which reveals the regions of the molecule to which an electrophile would initially be attracted.

**Table 1** Substrate scope of nitrostilbenes nitration

**Table 2** Nitration of other nitrostilbenes

Next, we investigated the nitration of nitrostilbenes analogues by decreasing the number of nitro groups under identical conditions (Table 2). The presence of (E)-2,4-dinitro-1-styrylsilene did not alter the expected outcome of the reaction (3a, 85%). However, the principal (E)-2,4-trinitrostilbenes as well as ortho-nitrated products were found, when the hydrogen atom in para-position was substituted by -CH₃ or -OCH₃ group (3b-e). With the presence of another methoxy group, two nitro groups were introduced in aryl ring and one nitro group in -C=C- double bond (3f). If strong electron-withdrawing nitro group was introduced in meta position, 3g was afforded in acceptable yields. Also, isomerization of the trans- geometry of the reactant 2, 4-trinitrostilbenes to cis- geometry in the products occurred, which was similar to the report by Srinivasan’s nitration with fuming nitric acid. This isomerization may be due to the favorable π-π interaction between the two aryl rings during nitration. Further investigation on nitrating trans-stilbene and trans-para-nitrostilbene under usual conditions revealed that this protocol was ortho C-H nitration enhanced (3h-j).

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**Table 1** Substrate scope of nitrostilbenes nitration

<table>
<thead>
<tr>
<th>Substrate</th>
<th>Product</th>
<th>Yield (%)</th>
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<tbody>
<tr>
<td>1</td>
<td>2a</td>
<td>90%</td>
</tr>
<tr>
<td>2</td>
<td>2b</td>
<td>85%</td>
</tr>
<tr>
<td>3</td>
<td>2c</td>
<td>66%</td>
</tr>
</tbody>
</table>

**Table 2** Nitration of other nitrostilbenes

<table>
<thead>
<tr>
<th>Substrate</th>
<th>Product</th>
<th>Yield (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>4</td>
<td>3a</td>
<td>85%</td>
</tr>
<tr>
<td>5</td>
<td>3b</td>
<td>65%</td>
</tr>
<tr>
<td>6</td>
<td>3c</td>
<td>60%</td>
</tr>
</tbody>
</table>

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Molecular electrostatic potential (ESP) was employed in the explanation of chemoselectivity, which reveals the regions of the molecule to which an electrophile would initially be attracted. The
structure of trans-stilbene was shown to be nearly planar, while with the number of nitro group increased, the structures of nitrostilbenes were twisted. (Fig. S1, Fig. S2, Fig. S4, Fig S6 and Fig. S8). Also, the ESP surface minima of trans-stilbene was about -14.35 kcal/mol and the electrons were well dispersed over the molecule. (Fig.1 (a) and Table S7) However, the ESP surface minima of nitrostilbenes were concentrated on the nitro-group for its lone pair electrons. (Fig.1 (b) and (c)) The ESP quantitative analysis of trans-C=C- double bond and aryl rings, trans-stilbene (-14.23 kcal/mol, -14.35 kcal/mol), trans-1-styryl-2,4-dinitrobenzene (-1.47 kcal/mol, -6.35 kcal/mol), trans, trans-1,3-distyryl-2,4,6-trinitrobenzene (-1.35 kcal/mol, -7.26 kcal/mol), well explained the decreases in chemical activity of trans-C=C- double bonds. (Table S2, Table S4 and Table S6) Also the ESP surface minima values of nitrostilbenes in aryl rings were higher than trans-stilbene’s but lower than their corresponding trans-C=C- double bond’s.

Figure 1 Electrostatic potential distribution (left) and average negative values around corresponding atoms in kcal/mol (right).

The average negative values around corresponding atoms were also presented in Fig. 1 (right) and the meta position of the aryl ring owned the minimum values for nitrostilbenes. However, it was found that the existing nitro group decreased the electrostatic interaction of trans-C=C- double bond with NO$_3^-$ sharply, which would favor substitution at the ortho position of the benzene ring. The high regioselectivity for nitration of nitrostilbenes by acetyl nitrate was activated by zeolite and gave NO$_3^-$ from a linear coordination mechanism.

The high regioselectivity for nitration of nitrostilbenes by acetyl nitrate and from a linear coordination mechanism. Efforts in further expanding the scope of the reaction and elucidating the mechanism are ongoing and will be reported in due course.

Notes and references

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† Electronic Supplementary Information (ESI) available: Detailed experimental procedures, spectral data, electrostatic potential calculations and crystallographic data. See DOI: 10.1039/c000000x/


[13] Nitration of (E)-2-(2-chlorostyryl)-1,3,5-trinitrobenzene and 2,2‘-((1(E),1’E)-(2,4,6-trinitro-1,3-phenylene)bis(ethene-2,1 diyl))bis(chlorobenzene) under the typical conditions.

[14] Experimental details of the structure determination can be found in the Supporting Information. CCDC-965295 and CCDC-999375 contain the supplementary crystallographic data for this paper. These data can be obtained free of charge from The Cambridge Crystallographic Data Centre via http://www.ccdc.cam.ac.uk/data_request/cif.


