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# Mechanistic insight of TiCl<sub>4</sub> catalyzed formal [3 + 3] cyclization of 1,3-bis(silyl enol ethers) with 1,3-dielectrophiles†

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The mechanism of TiCl<sub>4</sub> mediated formal [3 + 3] cyclization of 1,3-bis(silyl enol ethers) with 1,3-dielectrophiles is studied at the B3LYP level of density functional theory (DFT) to rationalize the experimental regioselectivity. Methyl and trifluoromethyl substituted 1,3 dielectrophiles are studied theoretically since they show different regioselectivities. Two different mechanisms involving 1,2 and 1,4 addition of 1,3-bis(silyl enol ethers) on 1,3-dielectrophiles are studied for each dienophile. The intramolecular transition metal catalyzed and non-catalyzed dynamic shift of the silyl moiety is also studied. The structure of the 1,3 dienophile and the associated Mulliken charges are the driving forces for different regioselectivities in methyl and trifluoromethyl dienophiles.

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#### Introduction

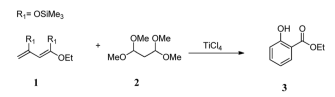
Functionalized arenes are interesting class of organic molecules, and find applications in natural products, polymers and medicinal chemistry.3 The traditional approaches for the synthesis of substituted arene involve direct substitution of already existing benzene scaffolds. The widely used strategies based on this classical approach involve electrophilic4 and nucleophilic substitution5 (in the case of aryl halides) and transition metal catalyzed coupling reactions.6 However, these strategies suffer from some serious drawbacks such as activating/deactivating effects of electron-donating/ withdrawing groups,7 sequences in multistep reactions, low yields and decreased availability of functionalized arenes as starting materials.7,8

A significant alternative is the "acyclic approach" for the synthesis of highly functionalized arenes, which are based on acyclic precursors.<sup>8</sup> These cyclocondensation reactions include [3 + 2 + 1] Dötz reaction,<sup>9</sup> transition metal catalyzed [2 + 2 + 2] cyclisation, [4 + 2] cycloadditions,<sup>10</sup> [4 + 2] Danheiser alkynecyclobutenone cyclization,<sup>11</sup> 1,6-electrocyclization reactions,<sup>12</sup> cyclocondensations of dielectrophiles with dinucleophiles and

The [3 + 3] addition of 1,3-bis(silyloxy)-1,3-butadienes with 3-silyloxy-2-en-1-ones is an important TiCl<sub>4</sub> mediated one pot cyclization reaction based on the "acyclic approach". The significance of the reaction includes its high regioselectivity, we excellent yields, moderate reaction conditions, and commercial availability of the catalyst. TiCl<sub>4</sub> is an efficient Lewis acid and shows significant affinity for oxygen containing organic compounds. The catalyst is effective in many organic transformations, Particularly in Pinacol coupling reaction, Pyrrolidine synthesis, Claisen rearrangements, and asymmetric aldol reaction. The alkene production from molecular reductive coupling of carbonyl compounds such as aldehydes, ketones, acylsilanes, ketoesters, and oxoamides are also prominent applications of titanium in organic chemistry.

Chan and coworkers (in 1979) reported the synthesis of arenes by formal [3 + 3] cyclization of 1,3-bis(silyl enol ethers) with 1,3-dielectrophiles, using titanium(n) as a Lewis acid

<sup>†</sup> Electronic supplementary information (ESI) available: Total electronic, zero-point and Gibbs free energies along with the Cartesian coordinates of the optimized geometries are shown in the ESI. See DOI: 10.1039/c5ra19238b



Scheme 1  $TiCl_4$  catalyzed cyclization of 1,3 bis(silyl enol ether) with 1,1,3,3-tetramethoxypropane.

<sup>[5 + 1]</sup> benzannulations between alkenoyl keteneacetals and nitroalkanes.<sup>13</sup> However, these annulations proceeds in harsh reaction conditions, require highly expensive catalysts, suffers from regiochemical ambiguities and lack of substrate generality.<sup>8</sup>

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(Scheme 1).<sup>22</sup> The reaction is widely used for synthesis of valuable functionalized arenes.<sup>7</sup> For example, Lewis acid promoted reaction of 1,3-bis(silyl enol ether) 1 with 2 delivers functionalized arene 3.

Similarly, a biphenyl derivative 6 can be generated (Scheme 2) by the reaction of 1,3-bis(silyl enol ether) 4 and 3-silyloxy-2-en-1-one 5.

Despite much synthetic advancements, mechanistic details of formal [3+3] cyclization are very limited. A proposed mechanism by Langer and coworkers<sup>7</sup> is presented in Scheme 3.  $TiCl_4$  selectively activates one of the two electrophilic centers of the unsymmetrical 1,3-dielectrophile by forming six membered chelation ring, which leads to the formation of exclusively one isomer and results in regioselective cyclization. Nature of the Lewis acid also affects the reaction pathway because specific functional groups are activated by specific Lewis acid.

The proposed mechanism involves coordination of 7 with TiCl<sub>4</sub> to give intermediate **Int1**, which is then attacked by the terminal carbon atom of 4 to give intermediate **Int2**. The **Int2** loses OSiMe<sub>3</sub> moiety to deliver **Int3**. Cyclization of **Int3** followed by aromatization in **Int4** delivers a functionalized arene 8 (Scheme 3). The above mechanism allows to rationalize the regioselectivity of a particular formal [3 + 3] reaction, however, it is relatively silent regarding the different regioselectivities starting from two structurally similar substrates (Scheme 4, *vide infra*). Moreover, the mechanism is based on a 1,4 addition of

Scheme 2 TiCl<sub>4</sub> catalysed regioselective [3 + 3] addition of 1,3-bis-(silyloxy)-1,3-butadienes with 3-silyloxy-2-en-1-ones.

Scheme 3 Proposed mechanism of TiCl $_4$  catalysed 3 + 3 addition of 1,3-bis(silyloxy)-1,3-butadienes with 3-silyloxy-2-en-1-ones.

Scheme 4 Illustration of observed regional regional regional regional regional forms of 1,3-bis(silyloxy)-1,3-butadienes with 3-silyloxy-2-en-1-ones.

1,3-bis(silyloxy)-1,3-butadienes on 1,3-dienophile. A competitive 1,2 addition is not invoked in the mechanism.

With the computational tools in hand, we became interested to investigate the mechanism of the formal [3+3] addition and the results are presented here. There appears no theoretical reports on the mechanism of the titanium catalyzed [3+3] addition of 1,3-bis(silyloxy)-1,3-butadienes with 3-silyloxy-2-en-1-ones.

## Computational methods

All calculations were performed with Gaussian 09. Geometries of the structures were optimized without any symmetry constraints at hybrid B3LYP method using 6-31G\* basis set. The B3LYP method consists of three parameter hybrid functional of Becke in conjunction with the correlation functional of Lee, Yang, and Parr. The B3LYP method provides a nice balance between cost and accuracy, and it is known to perform reasonably well for the prediction of geometries and behaviour of neutral and charged species23 including transition metals. The B3LYP method has been used in the theoretical simulations of properties and reactions catalysed by Titanium.24 Each optimized structure was confirmed by frequency analysis at the same level (B3LYP/6-31G\*) as a true minimum (no imaginary frequency) or a transition state (with one imaginary frequency). Imaginary frequencies of transition states were also evaluated to confirm that their associated eigenvector correspond to the motion along the reaction coordinates. Moreover intrinsic reaction coordinates (IRC) calculations were performed to confirm that the transition states connect to the concerned starting materials and products. IRC was performed until the stationary point was reached with RMS gradient less than 1  $\times$ 10<sup>-4</sup>. Stationary points located through IRC were then fully optimized at the above mentioned method. The reported energies for all structures are in kcal mol<sup>-1</sup> and include unscaled zero point energy corrections.

Density functional theory (DFT) calculations were performed to gain mechanistic insight for the regioselectivity observed for the [3+3] addition of 1,3-bis(silyloxy)-1,3-butadienes with 3-silyloxy-2-en-1-ones (Scheme 4). An *ortho* phenyl substituted benzoate ester 6 is the dominant product from 5 (a methyl ketone) whereas *para* phenyl substituted benzoate ester 10 is the major product from 9. The regioselectivity is altered when

methyl ketone is replaced with a trifluoromethyl ketone, 9. A number of questions needs to be answered in order to logically

investigate the mechanism; is the silyl group in dielectrophiles 5 and 9 dynamic between the two oxygen atoms  $(5 \rightarrow 5')$  and 9  $\rightarrow$  9')? If yes, then which isomer is more stable? What is the stability order for these two isomers when they are bound to titanium? Is it a direct addition of 4 on 5, or a 1,4 addition?

Therefore, in this study, we have attempted to address all these questions and the results are described below. However, to reduce the computational cost, SiMe<sub>3</sub> group is replaced with SiH<sub>3</sub>.

# Shift of the silyl group

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 $\alpha$ ,  $\beta$  unsaturated ketones 5 and 9 can exist in isomeric species 5' and 9', respectively by the shift of the silyl group between two oxygen atoms (Scheme 5). The dynamic shift is studied both in the presence and absence of the transition metal (Ti) and it is observed that the shift of the silyl group both in fluorinated and non-fluorinated substrates, 9 and 5, respectively, is a kinetically favourable process even in the absence of the transition metal (titanium). A transition state for the silyl shift in 5 is located at a barrier of 2.91 kcal mol<sup>-1</sup> from 5 (titanium free conditions), and the reaction is exothermic by 0.36 kcal mol<sup>-1</sup>. The low activation barrier for the silyl shift may be attributed to the close proximity of the silyl group to the keto oxygen in 5. The O1-Si bond (see Fig. 1 for numbering) in the transition state slightly increases to 1.89 Å compared to 1.73 Å in 5 (Fig. 2). The O1-Si bond in the product 5' is 2.27 Å. On the other hand, the O2-Si bond decreases to 1.88 Å in TS1 from 2.22 Å in 5, and finally to 1.74 Å in 5'. The stronger bond of silicon with both oxygen atoms may be another reason for low activation barrier.

A similar behaviour is also observed for the fluorinated substrate **9**. A transition state (Fig. 3) for silyl shift is observed at a barrier of 4.06 kcal  $\mathrm{mol}^{-1}$ , and the reaction is exothermic by 0.9 kcal  $\mathrm{mol}^{-1}$ . A relatively higher activation barrier from **9** (compare  $5 \rightarrow 5'$ ) may be attributed to the electron withdrawing effect of the CF<sub>3</sub> functional group which decreases the electron density on the keto oxygen. Therefore, the nucleophilic attack of the keto oxygen on the silyl group is relatively difficult. The shift of the silyl group in **5** and **9** is kinetically a favourable process even in the absence of transition metal.

O OSiMe<sub>3</sub>
Ph 5'

O OSiMe<sub>3</sub>
Ph 5'

O OSiMe<sub>3</sub>
Ph 
$$CF_3$$

**Scheme 5** Reversible silyl shifts between oxygens of 3-silyloxy-2-en-1-ones.

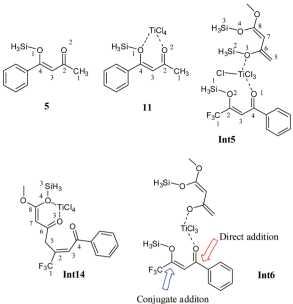


Fig. 1 Numbering scheme for discussion, and description of 1,2 and 1,4 addition.

## Complexation with titanium

Phenyl ketones 5' and 9' are more stable compared to their alkyl/trifluoroalkyl ketones 5 and 9, respectively, therefore, compounds 5' and 9' are anticipated as active species in the reaction. However, the situation is somewhat different after complexation with titanium. Although 5' is more stable than 5; but, its titanium tetrachloride complex 11' is relatively unstable by 2 kcal mol<sup>-1</sup> compared to 11, a complex from 5. The greater stability of 11 relative to 11' can be explained on the basis of attractive interaction between the hydrogens of CH<sub>3</sub> with the chlorides of TiCl<sub>4</sub>. H–Cl bond distances in 11 are 2.89 Å and 2.85 Å (Fig. 4).

For 9, this reversal of stability on complexation with titanium is not observed (Fig. 5). The complex of 9' with titanium (12') is

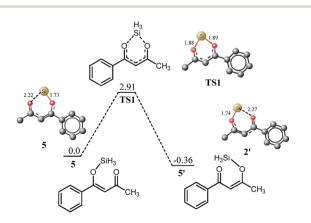


Fig. 2 Potential energy diagram for silyl shift between 5 and 5'. All values are relative to 5 at 0.0 kcal mol<sup>-1</sup>. All bond distances are in Angstroms. Unnecessary hydrogen atoms are removed for clarity.

Paper

Fig. 3 Potential energy diagram for silyl shift between 9 and 9'. All values are relative to 9 at  $0.0 \text{ kcal mol}^{-1}$ .

Fig. 4 Potential energy diagram for complexation of 5 and 5' with titanium to deliver 11 and 11', respectively. All values are in kcal mol<sup>-1</sup> with respect to 5 at 0 kcal mol<sup>-1</sup>. All bond lengths are in Angstroms, and unnecessary hydrogen atoms are removed for clarity.

still more stable than **12** by 0.38 kcal mol<sup>-1</sup>. The higher stability of **12**' over **12** is supportive to our hypothesis above that interaction of hydrogen of  $CH_3$  with the chlorides of  $TiCl_4$  is the main driving force for the higher stability of **11** over **11**'. The attractive interactions ( $CH_3 \cdots Cl$ ) in **11** are replaced by repulsive interactions in **12** ( $CF_3$  with chlorides).

In the next step of our mechanistic investigation we analyzed the distribution of charges in **11**, **11**′, **12** and **12**′ (shown in Fig. 6) in order to predict the reactivity based on charge densities. In both **11** and **11**′, the position of highest positive charge density is the carbon bearing the methyl group regardless of the fact whether this carbon is a keto carbon or silyloxy bearing carbon. In **11**, C2 and C4 (see Fig. 1 for labelling) have 0.5 and 0.389 positive charges, respectively whereas in **11**′, C2 and C4 bear 0.46 and 0.45 positive charges respectively. Since **11** is

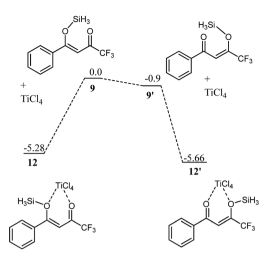


Fig. 5 Potential energy diagram for complexation of  $\bf 9$  and  $\bf 9'$  with titanium to deliver  $\bf 12$  and  $\bf 12'$ , respectively. All values are in kcal  $\rm mol^{-1}$  with respect to  $\bf 9$  at 0 kcal  $\rm mol^{-1}$ .

more stable and has higher positive charge on methyl ketone carbon therefore we believe that this isomer will be the active participating species in the reaction. For the fluorinated arene 12, the situation is quite opposite; a high positive charge is observed on carbon bearing the phenyl ring (C4). In 12′, carbons 2 and 4 have 0.327 and 0.439 positive charges, respectively whereas in 12, carbons 2 and 4 bear 0.394 and 0.414 positive charges, respectively. It is interesting to note that both isomers (of any system) have similar sequence of charge distribution.

Discussion in the subsequent section has been divided into fluorinated and methyl systems, and in each section 1,4 and 1,2 addition are discussed in detail.

# Fluorinated 1,3 dielectrophile

#### 1,2 Addition

The bis silyl ether 4 contains two oxygen atoms which are available for binding with titanium therefore, it is believed that the bis silyl ether binds to 12 before the reaction takes place. The titanium is penta coordinated in 12' and the binding of 4

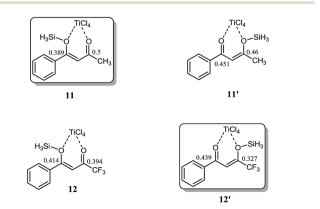


Fig. 6 Mulliken charges analyses on carbon 2 and 4 in 11, 11', 12 and 12'.

with titanium of **12** would generate an octahedral complex, **Int5**. Coordination of oxygen with titanium is followed by loss of the silyl chloride (Scheme 4). The literature reveals that octahedral complexes of titanium are well known, and are quite stable.<sup>25,26</sup>

Complexation of 4 with 12′ generates an intermediate octahedral complex Int5 where both oxygen atoms are in *cis* orientation. O1–Ti and O3–Ti bond lengths are 2.12 and 2.38 Å, respectively. The large bond distance of siloxy-Ti may be due to steric reasons (Fig. 7). The chlorides–Ti bonds *trans* to oxy ligands are relatively short (2.24 Å) whereas other Ti–Cl bonds are somewhat elongated (2.29 and 2.32 Å, shown in Fig. 8). The *cis* orientation of the oxy ligands is necessary for subsequent addition reaction. Any *trans* orientation of these group will not be favourable for C–C bond formation. A chloride ligand on titanium in Int5 is parallel to the silyl group at a distance of 3.29 Å.

A transition state for the concomitant elimination of the silyl and chloride groups is located at a barrier of 19.36 kcal  $\mathrm{mol}^{-1}$ . The transition state (TS3) has the geometry very similar to Int5 except the bond lengths change at the reaction site. The Ti–Cl bond elongates to 2.45 Å in TS3 from 2.24 Å in Int5. Cl–Si and O3–Si2 bond distances are 2.35 and 1.96 Å, respectively in the transition state. The reaction has low activation barrier, and

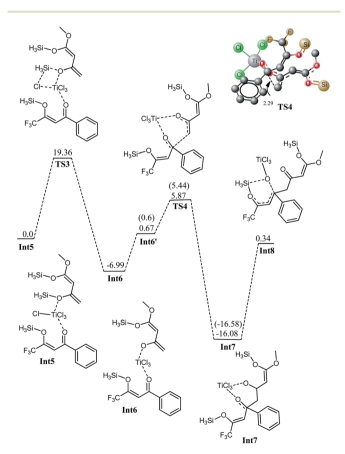


Fig. 7 Potential energy diagram for elimination of  $H_3Si-Cl$  followed by C–C bond formation (Int5  $\rightarrow$  Int8); all energies are relative to Int5 at 0 kcal mol $^{-1}$ . All bond lengths are in Angstroms. Unnecessary hydrogen atoms are removed for clarity. Values in parenthesis correspond to trimethylsilyl substituted derivatives.



Fig. 8 Optimized geometries of Int5, TS3 and Int6. All bond lengths are in Angstroms, and unnecessary hydrogen atoms are removed for clarity.

easily accessible at room temperature. Moreover this step is also driven thermodynamically. The product of the reaction, Int6 is  $6.99 \text{ kcal mol}^{-1} \text{ more stable than the starting material (Int5)}.$ The loss the silvl group generates an oxy ligands where both organic ligands are not well suited for C-C bond formation. Conformation changes are required to bring both organic fragments in proper orientation for C-C bond formation. This conformational change costs 7.66 kcal mol<sup>-1</sup>. A transition state is located at a barrier of 5.2 kcal mol<sup>-1</sup> from Int6'. This low activation energy may be attributed to relatively higher positive and negative charge densities on C4 and C6 carbons, respectively. The C-C bond being formed has a bond length of 2.29 Å in the transition state. Some other bond lengths also change considerably during the C-C coupling reaction. The O3-Ti bond length increases from 1.78 Å in Int6' to 1.90 Å in the TS4 whereas the O1-Ti bond length decreases from 2.06 to 1.94 Å.

Since we have replaced the trimethylsilyl group with  $SiH_3$  in order to reduce the computational cost, it was of interest whether this simplification affects the activation barriers, and may change the regioselectivities. Towards this end, activation barriers for the key step (1,2 or 1,4 (*vide infra*)) are studied with  $Si(CH_3)_3$  groups as well, and the activation barriers are given in the parenthesis (see Fig. 7 and 10). It is very obvious from Fig. 7 that the activation barrier is not affected considerably by replacing trimethylsilyl (4.84 kcal mol<sup>-1</sup>) group with  $SiH_3$  (5.20 kcal mol<sup>-1</sup>).

Int7 is believed to undergo an interesting rearrangement of OTiCl<sub>3</sub> to the silyl group which subsequently delivers Int9 by elimination of TiCl<sub>3</sub>(OSiH<sub>3</sub>). Although transition state for this oxy-titanium shift is located for methyl system (vide infra); however, no such transition state is located for fluorinated analogue. Rather, a true minimum Int8 is observed in which oxytitanium is bound to both, silyl moiety and carbon 5. The O3-C5 and O3-Si1 bond distances are 2.66 Å and 3.04 Å, respectively. Thermodynamic cost for this shift is 16.42 kcal mol<sup>-1</sup>. Int9 undergoes coordination of the 2<sup>nd</sup> silyloxy oxygen (O4) which is followed by elimination of another Si and Cl, very similar to the one observed in Int5. The transition state for the elimination of silyl chloride (TS5) is not modelled here. The activation barrier for the elimination of silyl chloride, is believed similar to the one observed in Int5. The product of the silyl chloride elimination, Int10 undergoes nucleophilic attack on the CF<sub>3</sub>-CO group. Kinetic barrier for the step is 26.56 kcal mol<sup>-1</sup>. The reaction is also favourable thermodynamically; the product (Int11) lies 18.7 kcal mol<sup>-1</sup> lower in energy than Int10

(Fig. 9). The cyclized product is believed to undergo loss of a water molecule and finally tautomerization to deliver the product. The possibility of elimination of silyl chloride with concomitant cyclization to deliver the final product cannot be excluded safely.

#### 1,4 Addition

The competitive reaction for Int6' is 1,4 addition on enone fragment instead of a 1,2 addition. The activation barrier for C-C bond formation through 1,4 addition was found higher compared to the 1,2 attack; 13.10 kcal mol<sup>-1</sup> for 1,4 addition compared with 5.2 kcal mol<sup>-1</sup> for the 1,2 attack. The higher activation barrier for Int6 → Int12 (Fig. 10) compared to Int6-Int6  $\rightarrow$  Int7 may be due to steric crowding at the carbon bearing the siloxy and CF<sub>3</sub> moieties, or may be due to electronic factors because of reduced charge density at keto carbon C2. The C6-C2 bond distance is 2.39 Å in the transition state (TS7), compared to 2.29 Å in the transition state for the 1,2 attack, TS4. The titanium has trigonal bipyramidal geometry in TS7 where fluorine atoms of CF<sub>3</sub> do not coordinate to the titanium. The reaction is thermodynamically favourable ( $E_R = 16.12$  kcal mol<sup>-1</sup>). The activation barrier for the similar reaction but involving Si(CH<sub>3</sub>)<sub>3</sub> has a very similar activation barrier (12.37 kcal mol<sup>-1</sup>). These findings are consistent with those in Fig. 7 that replacing a trimethylsilyl group with SiH<sub>3</sub> does not change the activation barrier to alter the regioselectivities.

**Int12** undergoes elimination of O-SiH<sub>3</sub> (next to CF<sub>3</sub>) and TiCl<sub>3</sub> moiety to generate the species for the next step, **Int13**. This process may be a direct one step process for the methyl ketone (*vide infra*) or may be a multistep process as in the case of trifluoro species. For the trifluoro species, first the silyl oxygen coordinates to titanium and a transition state **TS8** has been located for this coordination. This is just an addition reaction because it does not involve elimination of any other

Fig. 9 Potential energy diagram for cyclization in Int10 to generate Int11. All energies are in kcal  $mol^{-1}$  relative to Int10 at 0.0 kcal  $mol^{-1}$ .

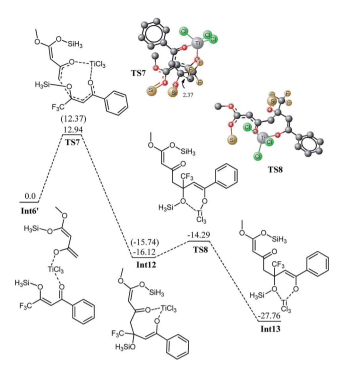


Fig. 10 Potential energy diagram for 1,4 addition in Int6 to generated Int12, followed by change in coordination to generate Int13. Hydrogen atoms are removed for clarity. All bond lengths are in Angstroms. Values in parenthesis correspond to trimethylsilyl substituted derivatives.

group. The cyclic species undergoes cleavage of OSiH3 and TiCl<sub>3</sub> to generate Int14 (see ESI†). In Int14, a silyl group is parallel to a chloride on titanium therefore; elimination of silyl chloride generates the enolxy species Int15. The geometry of Int15 has some interesting features; C7 is coordinating with titanium at a distance of 2.34 Å, the carboxylate oxygen (O4) along with the ortho carbon C7 behave as bidentate ligand for titanium, the titanium in Int15 has distorted octahedral geometry. Nucleophilic attack of C7 on C4 in Int15 generates the cyclic species Int16. The reaction has kinetic demand of 33.20 kcal mol<sup>-1</sup> (Fig. 11). In the transition state, C7-C4 bond distance is 2.68 Å whereas the C-Ti distance is increased to 2.75 Å. Although kinetically unfavourable, the reaction is highly favourable thermodynamically. The reaction is exothermic by 19.32 kcal mol<sup>-1</sup>. Thermal energy available at room temperature is not sufficient to surpass this barrier which clearly illustrates that this is not the pathway which should be observed experimentally. The regioselectivity observed experimentally is in marked contradiction with the product obtained from 1,4 addition.

The 1,2 addition is a favourable pathway energetically, and it explains the right regioselectivity. Although the calculations above reveal that 1,2 attack of enol on the keto oxygen delivers the product which is observed experimentally yet there is another set of similar calculations but with CF<sub>3</sub> ketone bound to titanium in the first step rather than a phenyl ketone. We have also considered this as a possible reaction in our study.

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Fig. 11 Potential energy diagram for intramolecular cyclization in Int15 to generate Int16, all energy values are relative to Int15 at 0.0 kcal  $mol^{-1}$ . Hydrogen atoms are removed for clarity all bond lengths are in

#### 1,2 Attack on CF<sub>3</sub> ketone

Calculations have been performed on a titanium bound complex starting from CF<sub>3</sub> ketone, Int17. Typically the nature and the number of step are almost comparable to those for phenyl ketone regarding both 1,2 and 1,4 attack. We have repeated the same set of calculations for CF3 ketone to explore the associated kinetics and thermodynamics. The first step is elimination of silvl chloride from Int17 for which a transition state has been located at a barrier of 19.67 kcal mol<sup>-1</sup> from starting complex. The activation barrier is very comparable to the elimination of silyl chloride in **Int5** (19.36 kcal mol<sup>-1</sup>). The reaction is thermodynamically favourable by 8.52 kcal mol<sup>-1</sup>. A few important changes in bond distances on going to transition states are given in the table below (Table 1).

Nucleophilic attack of enol on the C2 in Int18' is a kinetically favourable ( $E_{\rm act}$  6.22 kcal mol<sup>-1</sup>, see Fig. 12) but this activation energy is slightly higher (1 kcal mol<sup>-1</sup>) compared to the activation barrier for nucleophilic attack on phenyl ketone Int6 ( $E_{act}$  $= 5.2 \text{ kcal mol}^{-1}$ ). This difference in activation barrier may be attributed to the density of the positive charge of 12' and 12

Table 1 Comparison of some important geometric parameters in Int17 and TS10

Bond	Int17	TS10
TiCl	2.24	2.45
Ti-O	2.36	2.00
O-Si	1.72	1.95
Si-Cl	3.31	2.35

Fig. 12 Potential energy diagram of Ti catalyzed 1,2 addition of enol on  $CF_3$  ketone, all values are in kcal  $mol^{-1}$  relative to Int17 at 0.0 kcal  $mol^{-1}$ 

(vide supra). The important C6-C2 bond (being formed) has bond length of 2.37 Å in the transition state TS11 compared to 3.61 Å and 1.54 Å in Int19 and Int18, respectively. The similar reaction using Si(CH<sub>3</sub>)<sub>3</sub> group instead of SiH<sub>3</sub> has the activation barrier of 6.63 kcal mol<sup>-1</sup> (see ESI, Fig. SI1†).

The next key step in this pathway is the cyclization of Int20 to Int21, a reaction which generates the cyclic intermediate (Int21) from enol by nucleophilic attack (Fig. 13). The calculated activation barrier for the cyclization is 20.60 kcal mol<sup>-1</sup>. This activation barrier is although high but still accessible under the reaction conditions. Although energetics associated with this pathway are not very high (may proceed under reaction conditions); however, its comparison with 1,2 attack on phenyl ketone (1,2 attack starting from Int6) shows that this pathway is energetically less favourable. Moreover, the theoretical findings are consistent with the experimental regioselectivities (compare the structure of Int21 and Int11 with Int10).

The 1,4 attack on CF<sub>3</sub> ketone is expected to be kinetically demanding, although it delivers the desired product. The 1,4 addition on CF3 ketone is also modelled, and found to have high activation barriers (see Fig. SI2 and SI3†). The first 1,4 attack has an activation barrier of 15.99 kcal mol $^{-1}$  (much higher than any of the above three mechanisms). This high activation barrier is expected because this reaction not only involves 1,4 attack which is kinetically demanding but it Paper

F<sub>3</sub>C Cl<sub>3</sub> Cl<sub>3</sub>

Fig. 13 Potential energy diagram for cyclization in Int20. All values are in kcal  $mol^{-1}$  relative to Int20 at 0.0 kcal  $mol^{-1}$ .

involves intermediate 18' on which attack is even more demanding kinetically. Analysis of the pathways, shown above, clearly indicates that the charge density shown in Fig. 6 is a good tool to predict the regioselectivity of the reaction. Moreover, pathways with 1,2 attack on ketone are energetically more favourable whereas conjugate addition is thermodynamically less favourable.

#### Methyl system

From Fig. 6, it is apparent that the methyl ketone **11** has higher Mulliken charge density on C2; therefore, this structure is only considered for mechanistic studies. The phenyl ketone **11**′ is not considered for both **1**,2 and **1**,4 attack because higher activation barriers are expected than **11** (*vide supra*). Moreover, the regioselectivity expected through a **1**,2 attack is in contradiction to what is observed experimentally.

#### 1,2 Attack

The first step is typically cleavage of a chloride and a silyl group in **Int22**. The titanium in **Int22** has an octahedral geometry where both oxygen ligands are *cis* to each other. The O3–Ti–O2 bond angle is 81.77 degrees. The O3–Ti and O2–Ti bond distances are 2.40 and 2.06 Å, respectively. The Ti–Cl (*trans* to oxy ligands) bond lengths are in the range of 2.24–2.25 Å whereas the other chloride–titanium bond lengths are 2.30–2.31 Å. One of the chlorides is bent towards silyl group primarily because of the dipole dipole interactions (Fig. 14). A transition state for the cleavage of silyl chloride in **Int22** is located at a barrier of 18.74 kcal mol<sup>-1</sup> (from **Int22**, see Fig. 15). In the transition state **TS13**, Cl–Ti bond distance increases to 2.46 Å. The alkoxide–titanium bond distance decreases to 2.01 Å in

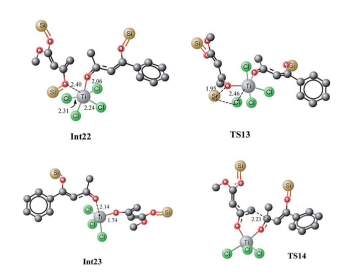


Fig. 14 Optimized geometries of Int22, TS13, Int23 and TS14. Unnecessary hydrogen atoms are removed for clarity. All bond lengths are in Angstroms.

**TS13**. The geometry around titanium also changes. The Cl–Ti–O3 (silyl) angle drops to 159.0 degrees in **TS13** from 176.18° in **Int22**. This activation barrier for the cleavage of a silyl and a chloride is very comparable to the one observed for the trifluoro system (*vide supra*). These findings clearly illustrate that

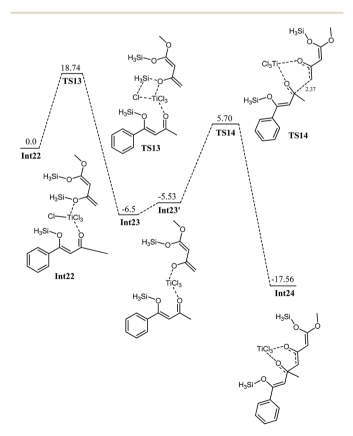


Fig. 15 Potential energy diagram for elimination of silyl chloride followed by 1,2 addition of enol to methyl ketone to generate Int24. All values are in kcal  $\text{mol}^{-1}$  relative to Int22 at 0.0 kcal  $\text{mol}^{-1}$ .

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the cleavage is not significantly influenced by the structure of the complex. The reaction is exothermic by 6.5 kcal mol<sup>-1</sup>. The complex Int23 has trigonal bipyramidal geometry around titanium. The two chlorides and alkoxide ligands do not make a perfect trigonal planar structure. O-Ti-Cl bond angle is 114-115 degrees, whereas Cl-Ti-Cl bond angle is 127.65 degrees. Two chlorides and the alkoxide ligands are in one plane (trigonal). The third chloride and keto oxygen are in the perpendicular plane. The alkoxide-Ti and keto-titanium bond distances are 1.74 and 2.14 Å, respectively. The subsequent C-C bond formation in Int23 has kinetic demand of 11.23 kcal mol<sup>-1</sup>, whereas the reaction is thermodynamically favourable by 12.03 kcal mol<sup>-1</sup>. In the transition state TS14, the C2-C6 bond distance is 2.23 Å. Moreover, the O3-Ti and O3-C5 bond distances are 1.88 and 1.31 Å, respectively. The newly formed O2-Ti and C2-O2 bond distances are 1.94 and 1.30 Å, respectively. The Int23, generated as a result of C-C bond formation, undergoes an interesting cleavage/shift of titanium oxy group on the silyl group. This shift is predicted to have a kinetic demand of 19.12 kcal mol<sup>-1</sup>. In the transition state, O1-C2 bond distance is 2.15 Å whereas the O2-Si1 bond distance is 3.17 Å.

The titanium oxy group interacts very loosely with the silyl groups, as it is reflected with bond separation in the TS. This behaviour of the methyl system is remarkably different than what is observed for the trifluoromethyl analogue. In the case of trifluoromethyl system, this type of transition state is not observed rather an intermediate is observed in which titanium oxy species is located between carbon and silyl group (*vide supra*).

It is believed that the Int25 generated in the above step (Fig. 16) soon loses  $Cl_3Ti$ –Osi $H_3$  species and binds to  $TiCl_4$  to generate Int26, as a precursor for the next step. Cleavage of a silyl and chloride in Int26 generates an intermediate Int27 ready for cyclization. Cyclization in Int27 is a kinetically and thermodynamically favorable process (Fig. 17). The titanium in TS16 has a trigonal bipyramidal geometry where two chlorides and an oxo ligand lie in the plane of trigon; however, the CH and chloride ligands are in the vertical axis. The CH–metal bond distance increases to 3.02 Å in the transition state from 2.22 Å in Int27. Moreover, the CH and keto carbon bond distance is 2.87 Å which indicates a very early transition state. The calculated activation barrier for the cyclization is 19.54 kcal  $mol^{-1}$ . Dehydration and tautomerization in Int28 may deliver the final aromatic product.

#### 1,4 Addition

**Int23** generated by the loss of a silyl chloride molecule can also undergo a 1,4 addition on the  $\alpha$ ,  $\beta$  unsaturated ketone to deliver intermediate **Int29** which is thermodynamically more stable than the intermediate **Int23** by 3.03 kcal mol<sup>-1</sup>. The reaction has kinetic demand of 19.57 kcal mol<sup>-1</sup> which is higher than the kinetic demand for the 1,2 addition in **Int23**′ (11.23 kcal mol<sup>-1</sup>). These observations are also consistent with the theoretical calculations for the trifluoromethyl system (*vide supra*) where direct attack is more favourable than the 1,4 addition.

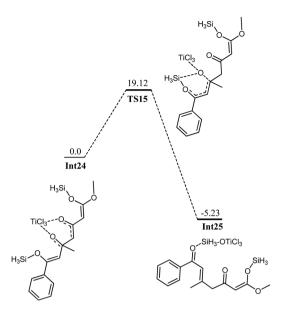


Fig. 16 Potential energy diagram for migration of  $OTiCl_3$  to silyl moiety in Int24 to generate Int25. All values are in kcal  $mol^{-1}$  relative to Int24 at 0.0 kcal  $mol^{-1}$ .

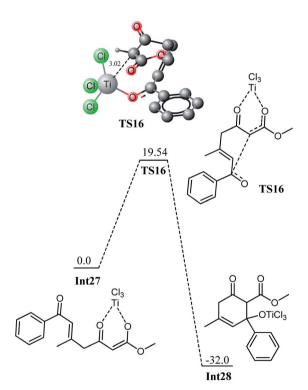


Fig. 17 Potential energy diagram for cyclization in Int27. All values are in kcal  $mol^{-1}$  with respect to Int27 at 0.0 kcal  $mol^{-1}$ .

From these calculations, one can infer that 1,2-addition is a more feasible pathway compared to 1,4-addition for these formal [3 + 3] addition reactions (1,4 addition). The difference in activation barrier for 1,2 and 1,4 attack is 8.34 kcal mol<sup>-1</sup> for the methyl system whereas this difference is about 7.9 kcal mol<sup>-1</sup> for the trifluoromethyl analogue.

Paper

Fig. 18 Potential energy diagram for Ti catalyzed 1,4 addition of enol to enone in Int23 All values in kcal mol<sup>-1</sup> with respect to Int23 at 0.0 kcal mol<sup>-1</sup>

The subsequent cyclization has a calculated activation barrier of 19.72 kcal mol<sup>-1</sup> which is not significantly different than the one calculated for the cyclization in Int27 which clearly illustrates that the selectivity is governed at the first addition step of the formal [3 + 3] addition reaction. The 1,2 addition (1,2 addition), shown in Fig. 16 and 17, delivers the product with the experimentally observed regioselectivity. The 1,4 addition (shown in Fig. 18) does not deliver the product with the correct regioselectivity. These findings are consistent with our the calculations that 1,2 addition is a favourable process with lower activation barriers compared to the 1,4 addition.

In summary, we have shown that the experimentally observed different regioselectivities for CH3 and CF3 enones can be explained by a common mechanism where 1,2 addition of 1,3-bis(silyl enol ethers) on 1,3-dielectrophiles is a more favorable pathway than the competing 1,4 addition. The differences in regioselectivities originate from different isomeric structures of enones entering in the catalytic cycle.

## Conclusions

Density functional theory calculations have been performed to gain mechanistic insight of the formal [3 + 3] addition of 1,3bis(silyl enol ethers) and 1,3-dielectrophiles to explain the experimentally observed regioselectivities. For this purpose, reaction of 1,3-bis(silyl enol ethers) with two different 1,3-dielectrophiles (5 and 9) is studied. The silyl moieties in 1,3-dielectrophiles are dynamic even in the absence of a metal catalyst  $(E_{\rm act} < 5 \text{ kcal mol}^{-1})$ . The isomeric dielectrophiles (for example 5

and 5') differ in their reactivity in formal [3 + 3] addition, and it depends on the strength of the positive charge (Mulliken charges) on the electrophilic carbon atom. Calculations reveal that a single mechanism can justify the experimentally observed different regioselectivities for 5 and 9 depending on the isomeric species entering in the catalytic cycle (11 and 12'). 1,2 and 1,4 addition mechanisms have been studied for both isomeric forms of each dielectrophile. The calculations reveal that 1,2 addition of 1,3-bis(silyl enol ethers) on 1,3-dielectrophile is a favorable process over the conjugate (1,4) addition. A typical mechanism involves coordination of 1,3-bis(silyl enol ethers) and 1,3-dielectrophiles with TiCl4 followed by elimination of silyl chloride. The elimination of silyl chlorides has an activation barrier of about 19-20 kcal mol<sup>-1</sup>. The elimination of the silvl chloride is followed by a nucleophilic attack on the dielectrophile. This nucleophilic addition governs the selectivity of the reaction. For trifluoromethyl, 1,2 addition (1,2 addition) is kinetically much favorable ( $E_{act} = 5.2 \text{ kcal mol}^{-1}$ ) than the 1,4 addition ( $E_{\text{act}} = 13.1 \text{ kcal mol}^{-1}$ , Fig. 18). For the methyl analogue, the difference in the activation barriers for the 1,2 and 1,4 addition is even more pronounced in the favor of 1,2 addition (about 8.34 kcal mol<sup>-1</sup>). The trends in the activation barrier for 1,2 and 1,4 addition for both dielectrophiles can be correlated to Mulliken charges at the electrophilic centers.

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