Mechanism and stereoselectivity in NHC-catalyzed β-functionalization of saturated carboxylic ester†

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To understand the mechanism and origin of the stereoselectivity of the [3 + 2] annulation reaction between a carboxylic ester and an isatin generating spirooxindole lactone catalyzed by N-heterocyclic carbene (NHC), density functional theory (DFT) calculations have been carried out. DFT results indicate that the catalytic cycle begins with the coupling of the catalyst with benzotriazole ester, followed by α-deprotonation to produce the enolate intermediate. The subsequent 1,4-proton transfer affords the homoenolate intermediate. The next crucial step is the stereoselective C–C bond formation. Then proton transfer takes place leading to the formation of the lactone intermediate. Finally, the elimination of the catalyst furnishes the final product. The presence of 1-hydroxybenzotriazole (HOBt) dramatically accelerates the proton transfer step. More importantly, HOBt has a non-negligible impact on stereoselective C–C bond formation, and the SR-configured product is the major stereoisomer of the annulation product, which is in good agreement with the experimental observations. The differential π–π stacking, C–H–π, lone pair (LP)–π and repulsion interactions are found to be responsible for the stereoselectivity. The obtained mechanistic insights should provide valuable information for understanding the important roles of the NHC catalyst and HOBt additive and be helpful for designing better NHC catalysts for this kind of reaction.

1. Introduction

As one of the most versatile and useful building blocks, saturated carboxylic compounds (e.g., esters and aldehydes) have been widely used in fields as diverse as pharmacology, chemistry, and materials science. Because of their great synthetic value, the functionalization of saturated carboxylic compounds has attracted considerable attention. However, the majority of the research work deals with either the carbonyl carbon atom or the α-carbon atom functionalization of saturated carboxylic compounds. Developing direct and efficient methods for the β-functionalization of these compounds is highly desirable and is still a challenge for chemists.

During the past few years, N-heterocyclic carbene (NHCs) have become a hot topic of research due to their widely applications as ligands in organometallic catalysis4 and in Lewis base catalysis. More importantly, NHCs have received considerable attention because of their diverse applications in organocatalytic reactions with excellent efficiency, such as umpolung reaction, benzoin, Stetter, Mannich, cycloadditions, redox reactions, CO₂ fixation, cooperative catalysis and cascade reactions. Indeed, many cycloaddition reactions using NHC catalysts affording various heterocycles have been reported, including the [2 + 2], [2 + 4], [3 + n] (n = 2, 3, 4), and [4 + n] (n = 2, 3) cycloadditions.

Recently, Xu et al. developed a catalytic protocol for the synthesis of spirooxindole lactones via β-functionalization of saturated carboxylic esters in tetrahydrofuran (THF) solvent at room temperature. It was found that the reaction of carboxylic esters with isatins could be effectively catalyzed by using NHC as catalyst and 1-hydroxybenzotriazole (HOBt) as additive. Additional experiments showed that HOBt is important in the current reaction because the yields and enanto- and diastereo-selectivities were higher in the presence of HOBt as compared with that in the absence of HOBt.

However, the mechanism of the NHC-catalyzed [3 + 2] annihilation reaction between carboxylic esters and isatins (Scheme 1) has not been studied theoretically until now. How does the catalytic reaction proceed? Which step is the stereocontrolling step? What role does the additive HOBt play? In this work, we will examine the NHC-catalyzed [3 + 2] annulation reaction between a carboxylic ester and an isatin using density functional theory calculations. We concentrate on the reaction mechanism as well as the origin of stereoselectivity. The objective of this work is to provide mechanistic insights for the catalyzed reaction shown in Scheme 1 and explain the enhanced
stereoselectivity in the presence of HOBt in comparison to that in its absence.

2. Computational methods

Geometries of all the structures involved in the reaction were optimized using the M06-2X functional\textsuperscript{13} with the 6-31G(d) basis set. Previous studies have confirmed the reliability of the chosen M06-2X functional.\textsuperscript{13} Frequency calculations were performed in order to check the minimum has all real frequencies while the transition state has one and only one imaginary frequency and to obtain thermodynamic corrections at 298.15 K and 1 atm pressure. In order to confirm the connection of the transition states and the desired minima, intrinsic reaction coordinate (IRC) calculations were also performed.\textsuperscript{14} To account for solvent effect, single-point energy calculations were carried out at the M06-2X/6-311++G(d,p) level in the solvent phase modeled by the Cramer–Truhlar continuum solvation model SMD\textsuperscript{15} in tetrahydrofuran (THF, \(\varepsilon = 7.43\)). To assign the atomic charges, natural bond orbital (NBO) analyses were also carried out.\textsuperscript{16} All calculations were carried out with the Gaussian 09 software package.\textsuperscript{17}

3. Results and discussion

3.1 Reaction mechanism

In this section, we present and discuss the detailed results for the [3 + 2] cycloaddition reaction between 4-nitrophenyl 3-phenylpropanoate and isatin depicted in Scheme 1. On the basis of our calculation results, the detailed mechanism for the whole catalytic cycle is presented in Scheme 2. Before considering the possible reaction pathways of the current reaction, we explore first the actual catalyst in the current reaction. As shown in Scheme 2, in the presence of cesium carbonate Cs\(_2\)CO\(_3\), the catalyst precursor Pre_Cat will undergo deprotonation to form the active NHC catalyst and release bicarbonate anion HCO\(_3^-\).\textsuperscript{18} Experimentally, 4-nitrophenyl 3-phenylpropanoate was used as one substrate (isatin as the other substrate) and 1-hydroxybenzotriazole (HOBt) as an additive. Computationally, we use the benzotriazole ester, depicted in Scheme 2 as one starting reactant because benzotriazole ester can be generated through the reaction of 4-nitrophenyl 3-phenylpropanoate with HOBt. As presented in Scheme 2, the catalytic reaction proceeds via six steps: (1) the addition of the active catalyst NHC to benzotriazole ester R1 to form the acyl azolium intermediate IM1 and release OBt\(^-\); (2) the deprotonation of the \(\alpha\)-C–H in IM1 to generate the enolate intermediate IM2; (3) the proton transfer of IM2 leads to the homoenolate intermediate IM3; (4) the stereoselective carbon–carbon bond formation process which can be achieved from the addition of isatin R2 to IM3 leading to intermediate IM4; (5) the proton transfer followed by the intramolecular lactonization will eventually provide access to the five-membered ring intermediate IM6, and (6) the elimination of the NHC catalyst yields the spiroxindole lactone product P with two chiral carbon atoms. The corresponding free energy profiles and optimized structures are shown in Fig. 1 (steps I–III) and 2 (steps IV–VI). In the following part, the reaction mechanism will be examined in greater detail.

**Step I: addition of catalyst NHC to benzotriazole ester R1.** As shown in Fig. 1, the reaction is initiated by the coupling of the NHC catalyst with benzotriazole ester R1 leading to the formation of intermediates IM01 and IM01-1 via the respective transition states TS1 and TS1-1. The calculated free energy barriers of TS1 and TS1-1 are 15.6 kcal mol\(^{-1}\) and 18.3 kcal mol\(^{-1}\), respectively. The relative free energy of IM01 and IM01-1 are 6.3 and 9.7 kcal mol\(^{-1}\), respectively. Clearly, formation of intermediate IM01 is both kinetically and thermodynamically much more favorable than IM01-1. Therefore, the description herein focuses on the reaction pathways associated with intermediate IM01. As shown in Fig. 1, for the transformation of TS1 \(\rightarrow\) IM01, the distance of C1–C2 bond shortens from 2.239 Å to 1.560 Å, while that of C2–O bond increases from 1.447 Å to 1.658 Å. These results demonstrate that the formation of C1–C2 bond is occurred together with the cleavage of C2–O bond. Subsequent dissociation of OBt\(^-\) from IM01 leads to the acyl azolium intermediate IM1.

Scheme 3 shows the NBO charge analysis results. As can be seen from Scheme 3, for Cat + R1 \(\rightarrow\) IM01 process, the positive charge value on the C1 atom increases from 0.148e to 0.414e, while that on the C2 atom decreases from 0.840e to –0.041e. Furthermore, the negative charge value on the N atom decreases from –0.489e to –0.415e, while that of the carbonyl oxygen O3 atom increases from –0.543e to –0.909e. These results indicate...
that the electron density is transferred from Cat to R1 during the process of Cat + R1 → IM01.

**Step II: α-C–H deprotonation.** From intermediate IM1, α-C–H deprotonation occurs to generate the enolate intermediate IM2. Many computational studies have shown that the mediators assisted deprotonation favors the direct deprotonation. In this work, three possibilities including direct, OBt−/C0 and HCO3−/C0 assisted deprotonation have been investigated.

1. **Direct deprotonation.** For direct deprotonation, we cannot locate the corresponding transition state despite lots of attempts. We performed relax potential energy scan for the C4–Ha bond. The corresponding pointwise potential curve was provided in Fig. S1 in the ESI† Fig. S1† clearly shows that the energy keeps increasing when the Ha atom getting farther from the C4 atom of IM1. The high energy barrier estimated for direct deprotonation process (~70 kcal mol⁻¹), indicates that the direct deprotonation pathway is kinetically unfeasible. In view of this, a mediator assisted deprotonation, facilitated by the previously formed OBt− (or HCO3−) has been examined.

2. **OBt− assisted deprotonation.** The anion OBt− generated in the first step can remove the hydrogen atom (Ha) in IM1 via the transition state TS2 to generate intermediate IM2. The calculated free energy barrier is 2.4 kcal mol⁻¹ for TS2 with respect to IM1, demonstrating that the OBt− mediated deprotonation of α-C–H is a fast process under the experimental conditions. During this process (IM1 → TS2 → IM2), the distance of C4–Ha bond is elongated from 1.099 Å to 1.372 Å (IM1 → TS2), whereas the distance of Ha–O bond is shortened from 1.245 Å, to 0.973 Å (TS2 → IM2 + OBt). These results indicate that the deprotonation process is completed. The transformation from IM1 to IM2...
Fig. 1  Free energy profile obtained at SMD(THF)/M06-2X/6-311++G(d,p) level (kcal mol\(^{-1}\)) and optimized structures for steps I–III. Bond lengths are given in Å.

Scheme 3  The NBO charges of selected atoms in NHC, R1 and IM01 (units of e).
is found to be slightly exergonic with a reaction energy of −0.9 kcal mol⁻¹.

(3) HCO₃⁻-assisted deprotonation. In another possibility, the previously formed HCO₃⁻ was used as a promoter. The energy barrier calculated for the HCO₃⁻-assisted deprotonation is 7.6 kcal mol⁻¹ via B_TS2. As shown in Fig. 1, the distance of C4–Hb bond increases from 1.099 Å in IM1 to 1.283 Å in B_TS2, while that of Ha–O bond decreases from 1.359 Å in IM1 to 0.968 Å in H₂CO₃.

Taken together, the mediators (BtO⁻ and HCO₃⁻) promote the deprotonation process. The most favored proton transfer pathway is found to be that from IM1 to IM2 via TS2 with the assistance of OBT⁻.

Step III: 1,4-proton transfer. The 1,4-proton transfer of IM2 forms intermediate IM3. Similar to the discussion of step II, both direct and mediator (HCO₃⁻, BtO⁻ and HOBt) promoted proton transfer processes have been investigated.

For direct 1,4-proton transfer, migration of the Hb atom connected to C5 atom to O3 atom via the corresponding five-membered ring transition state D_TS3 with a free energy barrier of 31.1 kcal mol⁻¹ (43.8 kcal mol⁻¹ with respect to reactants) (Fig. 1). Therefore, direct 1,4-proton transfer in intermediate IM2 is a kinetically inaccessible process under the experimental conditions.

We also calculated the HCO₃⁻ and OBT⁻ assisted proton transfer. The corresponding energy profiles are also presented in Fig. 1. However, the high activation energies of B_TS3 (39.8 kcal mol⁻¹ relative to reactants, associated with the HCO₃⁻ mediated proton transfer) and TS3 (40.3 kcal mol⁻¹ relative to reactants, associated with the OBT⁻ mediated proton transfer) indicate that the mediator (HCO₃⁻ and BtO⁻) has minimal effects on the proton transfer event. Thus these pathways are also kinetically unfavorable, and can be ruled out.

Experimentally, HOBt was used as an additive, promoting us to investigate the HOBt assisted proton transfer process. According to our calculation results, the 1,4-proton transfer proceeds via the seven-membered ring transition state A_TS3 leading to the formation of the homoenoate intermediate IM3 in the presence of HOBt (Fig. 1). Computational results show that the transition state for this process (A_TS3 (25.1 kcal mol⁻¹)) is much lower in energy than those of the unassisted and the BtO⁻ and HCO₃⁻ assisted proton transfer processes (i.e., D_TS3 (43.8 kcal mol⁻¹), B_TS3 (39.8 kcal mol⁻¹) and TS3 (40.3 kcal mol⁻¹)). As can be seen from Fig. 1, the Hb atom migrates from C5 to O, at the same time, the H atom transfers from O to O3 via the seven-membered ring transition state A_TS3. The bond distances of C5–Hb, Hb–O, O–H and O3–H in A_TS3 are 1.363, 1.257, 1.993 and 0.982 Å, respectively, indicating the HOBt mediated 1,4-proton transfer occurs in a concerted manner but not synchronous.

Step IV: the stereoselective C–C bond formation. The fourth step is the stereoselective determining step, which involves the addition of the C5 atom of IM3 to the C6 atom of R2 leading to the formation of C–C bond. As depicted in Scheme 4, four stereocchemically distinct attack modes i.e. (Re, Re), (Re, Si), (Si, Re) and (Si, Si) were located resulting into the formation of intermediates IM4(RR&RS&SR&SS), which have two chiral carbon atoms (i.e., C5 and C6). It should be noted that the formation of the C5–C6 bond is concerted with a proton transfer of the enolate proton to the isatin moiety leading to the formation of intermediate IM4. According to experiments, improved enantioselectivity could be achieved when HOBt was used as an additive, both the non- and HOBt-assisted C–C bond formation pathways have thus been explored.

In the absence of HOBt, the activation energy barriers of this step via transition states D_TS4(RR&RS&SR&SS) are 13.2/20.2/14.0/17.9 kcal mol⁻¹ (29.6/36.6/30.4/34.3 kcal mol⁻¹ with respect to reactants), as shown in Fig. S2. Considering that the activation energy barrier of D_TS4(RR) is lower than those of D_TS4(RS&RS&SS), the reaction pathway associated with D_TS4(RR) should be more favorable than the other pathways. Therefore, the dominant product in the absence of HOBt should be the RR configuration product. This is inconsistent with the experimental results. The experimentally observed major product without the use of HOBt was the SR-stereoisomer of the cycloaddition product. Considering this inconsistency, more experimental studies are expected to be necessary to better understand the stereoselectivity associated with the reaction, depicted in Scheme 1 in the absence of HOBt.

Since HOBt was used as an additive in experiment, it might assist in this C–C bond formation. We thus have included an explicit molecule of HOBt in this step to form intermediate IM4. Fig. 2 shows that the free energy barriers of this step in the presence of HOBt via transition states TS4(RR&RS&SR&SS) are 9.8/10.2/7.9/12.1 kcal mol⁻¹ (26.2/26.6/24.3/28.5 kcal mol⁻¹ relative to reactants), which is found to be lower than the transition states in the absence of HOBt (for details see the ESI, Fig. S2†). To understand this result, we examined the non and HOBt assisted transition state D_TS4 and TS4 structures (shown in Fig. S3†). As depicted in Fig. S3,† additional interactions such as O–H···O and O–H···N are present in the HOBt-assisted transition state compared to the unassisted transition state. These interactions are depicted as follows: O–H···O(a, b, c and d) and O–H···N(e). Moreover, Fig. 2 clearly shows that the energy barrier of TS4(SR) is remarkably lower than those of other configuration transition states (i.e., TS4(RR), TS4(RS) and TS4(SS)), for simplicity of presentation, we discuss the reaction pathways starting from IM4(SR) here (as shown in Fig. 2). Results for the less favorable pathways which feature RR/RS/SS-configuration are provided in the ESI (Fig. S4†). Intrinsic reaction coordinate (IRC) results for transition state TS4(SR) reveal that the C5–C6 bond formation occurs along with the Hb atom migration from O3 to O7, leading to intermediate IM4(SR). From TS4(SR) to IM4(SR), the distance of C5–C6 bond changes from 2.502 Å to 1.570 Å, at the same time, the distance of O3–Hb bond increases from 0.975 Å to 1.426 Å, while the Hb–O7 bond shortens from 2.136 Å to 1.051 Å. The above structural analysis indicates that the C5–C6 bond formation is accompanied by the transformation of the Hb atom (from O3 to O7).

Step V: the proton transfer and intramolecular lactamization. As shown in Fig. 2, intermediate IM4(SR) undergoes consecutive proton transfer and intramolecular lactonization to reach intermediate IM6(SR). First, the proton Hc transfers to the C4 atom generates intermediate IM5(SR). This proton
transfer step via transition state TS5(SR) has an activation energy of 10.2 kcal mol\(^{-1}\). For IM4(SR) \(\rightarrow\) TS5(SR) \(\rightarrow\) IM5(SR) reaction course, the bond distance of O’–Hc is elongated from 1.300 Å in TS5(SR) to 2.058 Å in IM5(SR), while that of Hc–C4 is shortened from 1.330 Å in TS5(SR) to 1.111 Å in IM5(SR).

Subsequently, lactonization of IM5(SR) generates the slightly more stable intermediate IM6(SR). Note that we can not locate the corresponding transition state despite lots of attempts. From IM5(SR) to IM6(SR), the distance of C2–O7 bond shortens from 2.804 Å to 1.523 Å.

**Step VI: formation of the cycloaddition product.** In the final step, NHC catalyst releases from intermediate IM6(SR) affording the final cycloaddition product P(SR) through the transition state TS6(SR) with an activation energy of 6.0 kcal mol\(^{-1}\). As shown in Fig. 2, the distance of C1–C2 bond is elongated from 1.546 Å in IM6(SR), to 2.031 Å in TS6(SR).

### 3.2 Origin of selectivity

As shown in Fig. 2 and discussed above in step IV, the addition of the homoenoalate intermediate IM3 to isatin R2 leading to the formation of C–C bond is the stereoselectivity-determining step. Consequently, transition states TS4(RR), TS4(RS), TS4(SR) and TS4(SS) are found to be responsible for determining the stereoselectivity of the entire reaction. The activation energy barriers for generating the RR/RS/SR/SS configuration products via TS4(RR/RS/SR/SS) are calculated to be 26.2/26.6/24.3/28.5 kcal mol\(^{-1}\) (the energy difference between TS4(RR/RS/SR/SS) and reactants). Obviously, the SR-configurational product P(SR) is the major product, agrees well with the experimental observations. Based on the Curtin–Hammett Principle, the product distribution is determined by the standard free energy difference of the stereoselectivity-determining transition states. The energy difference between
TS4(SR) and TS4(RS), which are responsible for enantioselectivity, have an energy difference of 2.3 kcal mol\(^{-1}\) corresponding to an enantiomeric excess of 96%. The computed enantioselectivity is in good agreement with the experimental reported value (93%). Similarly, the energy difference between TS4(SR) and TS4(RR), which is related to diastereoselectivity, is 1.9 kcal mol\(^{-1}\). This energy difference corresponds to a diastereomeric ratio of 92%, which is in good agreement with the experimentally observed de of 91% (diastereomeric ratio > 20 : 1).

Furthermore, in order to better understand the origin of stereoselectivity of the NHC-catalyzed [3 + 2] annihilation reaction, we performed the noncovalent interaction (NCI) analysis\(^{21}\) which can distinguish the strong interactions as well as the repulsion interactions. NCI analysis results for the stereoselectivity-determining transition states TS3(RR), TS3(RS), TS3(SR), and TS3(SS) are presented in Fig. 3. As depicted in Fig. 3, it is clear that a large green cloud between the \(\pi\)-systems of the NHC-bounded homoenolate (IM3-part) and HOBt is observed in TS3(SR), but absent in the other transition states. In addition, the repulsion interactions exist in TS3(RR), TS3(RS), and TS3(SS). Moreover, a number of noncovalent interactions including \(\pi\cdots\pi\) interactions, lone pair (LP)\cdots\pi interactions, and C-H\cdots\pi interactions are identified in TS3(SR). Therefore, TS3(SR) is more stable than the other transition states (TS3(RR), TS3(RS) and TS3(SS)), which is in good agreement with the experimentally observed enantioselectivity.

Fig. 2 Free energy profile obtained at SMD(THF)/M06-2X/6-311++G(d,p) level (kcal mol\(^{-1}\)) and optimized structures for steps IV–VI. Bond lengths are given in Å.
3.3 The nature of catalyst

In this section, global reactivity indexes (GRIs) analysis was carried out in order to understand the role of the NHC catalyst. The results are presented in Table 1. In GRI analysis, the molecular global electrophilicity character represented by the electrophilicity index $\omega$, can be derived from the following expression,

$$\omega = (\mu^2/2\eta) \, (\text{eV}),$$

where $\mu$ and $\eta$ are chemical potential and chemical hardness, respectively. $\mu$ and $\eta$ can be calculated from the one-electron energies of the highest occupied molecular orbital (HOMO) and the lower unoccupied molecular orbital (LUMO), $\varepsilon_H$ and $\varepsilon_L$ with the expressions

$$\mu = (\varepsilon_H - \varepsilon_L)/2$$

and

$$\eta = (\varepsilon_L - \varepsilon_H),$$

respectively.\[23\] Within the Kohn-Sham scheme,\[24\] the empirical (relative) nucleophilicity index $N$ introduced by Domingo et al. was written as

$$N = \varepsilon_{H(\text{Nu})} - \varepsilon_{H(\text{TCE})}. $$

The nucleophilicity scale was referenced to tetracyanoethylene (TCE). Table 1 shows that from R1 to IM01, the electronic chemical potential increases ($-4.10 \, \text{eV}$ vs. $-3.46 \, \text{eV}$), which is consistent with what we found in NBO charge analysis, i.e., the charge goes from NHC to R1. Table 1 also shows that the coupling of NHC catalyst with carboxylic esters R1 strengthens its nucleophilicity (the nucleophilicities of IM2 (4.96 eV) and IM3 (5.40 eV) are larger than that of R1, $N = 2.67 \, \text{eV}$), but weakens its electrophilicity (the electrophilicity of IM2 (IM3) is smaller than that of R1, 0.67 (0.79) eV vs. 1.10 eV). The GRI results suggest that the coordination of NHC to R1 makes the nucleophilic addition to R2 easier.

| Species | $\eta$ (eV) | $\mu$ (eV) | $\omega$ (eV) | $N$ (eV) |
|---------|-------------|------------|--------------|----------|
| R1      | 7.66        | -4.10      | 1.10         | 2.67     |
| IM01    | 7.05        | -3.46      | 0.85         | 3.61     |
| IM2     | 5.71        | -2.77      | 0.67         | 4.96     |
| IM3     | 4.83        | -2.77      | 0.79         | 5.40     |

4. Conclusion

The reaction mechanism of the NHC-catalyzed cycloaddition reaction between carboxylic esters and isatins generating spirooxindole lactones with high enantio- and diastereo-selectivities has been investigated by using density functional theory (DFT) calculations at SMD(THF)/M06-2X/6-311++G(d,p) level. The role of the additive HOBt is also examined in detail.
The current computations reveal that the catalytic cycle comprises six steps: the NHC catalyst first attacks carboxylic ester, followed by $\alpha$–C–H deprotonation gives the enolate intermediate. Subsequently, proton transfer occurs leading to the homoenolate intermediate and the next step is the stereoselective C–C bond formation. The fifth step is the generation of lactone intermediate and finally liberation of catalyst gives the cycloaddition product. DFT results reveal that the additive stereoselectivity is traced to di with the experimentally reported enantioselectivity computed stereoselectivity in the presence of HOBt agrees very well with the homoenolate intermediate and the next step is the stereo- 

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