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A chiral pentanidium and pyridinyl-sulphonamide ion pair as an enantioselective organocatalyst for Steglich rearrangement†

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Enantioselective ion pair catalysis has gained significant attention due to its ability to exert selectivity control in various reactions. Achiral counterions have been found to play crucial roles in modulating reactivity and selectivity. The modular nature of an ion pair catalyst allows rapid alterations of the achiral counterion to achieve optimal outcomes, without the need to modify the more onerous chiral component. In this study, we report the successful development of a stable chiral pentanidium pyridinyl-sulphonamide ion pair as a nucleophilic organocatalyst for asymmetric Steglich rearrangement. The ion pair catalyst demonstrated excellent performance, leading to enantioenriched products with up to 99% ee through simple alterations of the achiral anions. We conducted extensive ROESY experiments and concluded that the reactivity and enantioselectivity were correlated to the formation of a tight ion pair in solution. Further computational analyses provided greater clarity to the structure of the ion pair catalyst in solution. Our findings reveal the critical roles of NMR experiments and computational analyses in the design and optimisation of ion pair catalysts.

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Introduction

Asymmetric ion pair catalysis is a powerful strategy to achieve an enantioselective outcome in a reaction.^{1,2} The chiral cation or anion can transmit stereochemical information to the reaction centres on their respective counterions, thus enabling enantioselectivity to be achieved. The use of chiral cations such as quaternary ammonium salts in phase transfer catalysis (PTC) has been extensively reported in the literature.^{2–5} Some prominent strategies that evolved from chiral cation PTC include asymmetric counteranion-directed catalysis,⁶ chiral anion PTC,⁷ and cation- or anion-binding catalysis.^{8,9}

The use of a chiral cation ion pair catalyst with transition metal binding to a ligand on the chiral cation (Ooi¹⁰ and Peters¹¹) or a ligand on the achiral anion (Phipps^{12,13}) has been previously reported (Fig. 1A). An alternative strategy will be chiral cation-directed transition metal catalysis, where the

active metal centre is achiral and chiral information from the cation is transmitted by virtue of proximity through tight ion pairing. In this aspect, we have contributed to the use of chiral guanidinium cations such as bisguanidinium (BG) and pentanidium (PN) for various enantioselective oxidation reactions.^{14–17} Recently, the group of Jacobsen has reported an asymmetric anion-binding transition metal catalysis *via* hydrogen bond based organocatalysts.¹⁸

Ion pair catalysis allows modular design of an asymmetric catalytic system. For instance, a privileged chiral scaffold can be deployed as the cation or anion for optimal stereochemical induction, while the counterion can be an achiral catalyst that is responsible for activation of the reactants. This modularity enables a variety of cooperative catalytic concepts to be possible. Chiral cation-directed transition metal catalysis¹⁹ and asymmetric photocatalysis *via* ion-pairing²⁰ are two examples of such concepts.

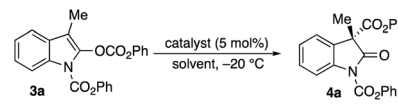
Building on our prior research on chiral guanidinium-directed transition metal catalysis,^{14–17,19,21} we now introduce the utilization of an anionic nucleophilic catalyst in chiral cation-directed nucleophilic organocatalysis. The DMAP scaffold, known for its versatility in catalysing an array of reactions including esterification, acylation, and halogenation, has been extensively investigated as a nucleophilic organocatalyst.²² Over the past few years, chiral DMAP catalysts have also been developed. For example, Ruble and Fu have designed ferrocene-based planar DMAP derivatives,^{23,24} while Suga and his colleagues have developed BINOL-based chiral DMAP derivatives.^{25,26} The research groups of Vedejs,^{27,28} Conon,²⁹ Sibi,³⁰ and Spivey³¹ have each prepared

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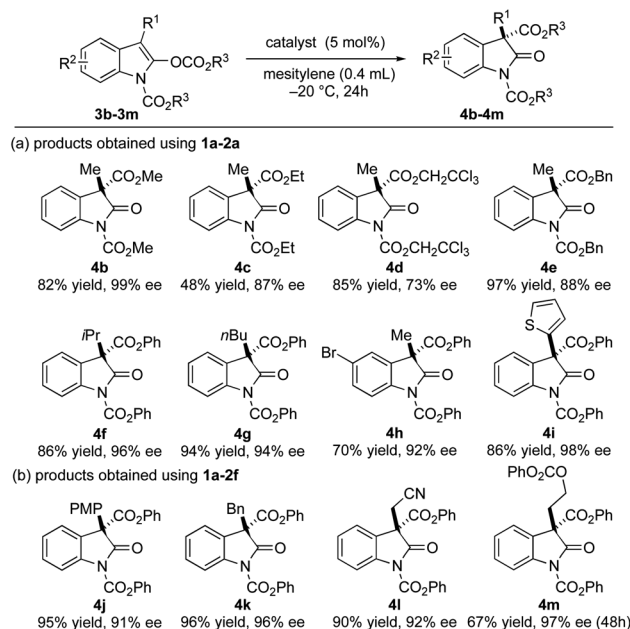
Table 1 Optimization of reaction conditions in enantioselective Steglich rearrangement of oxindole derivatives^a

				
Entry	Catalyst	Solvent	Yield ^b	ee ^c
1 ^d	1a-Cl, DMAP	Toluene	39%	0%
2 ^d	1a-Cl, Na-2a	Toluene	8%	7%
3 ^d	1a-2a	Toluene	99%	0%
4	1a-2a	Toluene	74%	87%
5	1b-2a	Toluene	82%	73%
6	1c-2a	Toluene	63%	55%
7	1a-2b	Toluene	90%	70%
8	1a-2c	Toluene	68%	84%
9	1a-2d	Toluene	—	—
10	1a-2e	Toluene	66%	77%
11	1a-2f	Toluene	97%	79%
12	1a-2g	Toluene	63%	83%
13	1a-2a	MeCN	<5%	20%
14	1a-2a	DCM	63%	25%
15	1a-2a	THF	65%	30%
16	1a-2a	Et ₂ O	94%	75%
17	1a-2a	Mesitylene	99%	92%
18 ^e	1a-2a	Mesitylene	81%	86%
19 ^f	1a-2a	Mesitylene	91%	86%

^a Reaction conditions: **3a** (0.04 mmol, 1.0 equiv.), catalyst (5 mol%), and solvent (0.4 mL) at $-20\text{ }^{\circ}\text{C}$ for 24 h. ^b Isolated yield. ^c Determined by chiral HPLC analysis. ^d Reaction was carried out at room temperature. ^e Reaction was carried out at $0\text{ }^{\circ}\text{C}$. ^f The catalyst loading was 3 mol%.

was observed using solvents that have a higher dielectric constant than toluene or mesitylene³⁹ (such as acetonitrile or dichloromethane), giving products with severely degraded ee values and varying yields, whereas the products afforded in less polar solvents had higher ee values (entries 16–17). Meanwhile, increasing the temperature or decreasing the catalyst loading both decreased the yields and ee values of products (entries 17–19). In summary, the product **4a** can be afforded with the best yield and ee value using 5 mol% chiral ion pair catalyst **1a-2a** in mesitylene at $-20\text{ }^{\circ}\text{C}$.

To gain a better understanding on the nature of our ion pair catalyst, an investigation into the substrate scope was carried out (Fig. 2). We first probed the migrating acyl group on the substrates. Alkyl and benzyl substituents on the migrating acyl group (**3b-3e**) showed good compatibility with the ion pair catalyst **1a-2a** to achieve moderate to good ee values, albeit with poorer yield for the ethyl ester derivative (**4c**). It is partially due to the purification process as the substrate **3c** and the product **4c** share similar polarity. The substitution groups on the oxindole scaffold were also examined. For C-3 substitution, excellent ee values of the corresponding products were achieved with the catalyst **1a-2a** when using a larger alkyl (**3f-3g**) or a 2-thienyl substituent (**3i**). The bromine substitution on the C-6 position had a minimal effect on the enantioselectivity but had a negative impact on the yield (**3h**). The ion pair catalyst **1a-2f** was also able to affect the rearrangement for the substrates bearing C-3

**Fig. 2** Substrate scope of enantioselective Steglich rearrangement of oxindole derivatives.

benzyl or *para*-methoxyphenyl groups (**3j-3k**) in excellent yields and good to excellent ee values. Meanwhile, sensitive substituents such as the acidic cyanomethyl group could be tolerated (**3l**). The acyloxyethyl pendant was also particularly noteworthy as it demonstrated insensitivity to β -elimination or loss of the acyloxy group even after 48 h (**3m**). Both **4l** and **4m** were obtained in excellent yields and ee values. The absolute configuration of the major enantiomer of **4e** was determined to be *S* by comparing the HPLC trace of a known compound in the literature.²⁵

The influence of temperature and the properties of organic solvent on the viability of ion pair formation and the strength of the association between ions is well-documented.^{41,42} NMR spectroscopy serves as a powerful method to examine the behaviour of the chiral ion pair catalyst in the solution state. This information can then be correlated to the results observed in our optimization process (Table 1).

In our NMR investigation, we varied both temperatures and solvents using the ion pair catalyst **1a-2f** as a model. The catalyst **1a-2f** was chosen for the study as it demonstrated slightly better and more consistent solubility across all the temperatures during NMR studies.

Two-dimensional ROESY (Rotating-frame Overhauser Effect Spectroscopy) experiments can reveal spatial correlations between two groups. A negative off-diagonal cross-peak, represented by a blue signal that maps to two different signals, suggests that the associated protons are generally less than 5 Å apart. ROESY experiments were conducted on the ion pair catalyst **1a-2f** in two solvents: chloroform-*d* and toluene-*d*₈. The experiments were performed at two different temperatures: $25\text{ }^{\circ}\text{C}$ and $-20\text{ }^{\circ}\text{C}$. At $25\text{ }^{\circ}\text{C}$, the reaction produced a racemic product. However, at $-20\text{ }^{\circ}\text{C}$, a high degree of enantioselectivity



was observed (Table 1, entries 3 and 4). No intermolecular spatial correlations were found in the 2D ROESY spectrum of **1a–2f** in chloroform-*d* at either temperature (refer to Fig. S1–S4†). Instead, only intramolecular contacts were present. The 2D ROESY spectra for **1a–2f** in toluene-*d*₈, a less polar solvent, showed distinct spatial correlations between two of the aryl protons in **2f** and the “arms” of the **1a** cation, specifically the *tert*-butyl protons. These cross-peaks were absent at 25 °C (see Fig. 3A) but appeared at –20 °C (refer to Fig. 3B).

Molecular Dynamics (MD) simulations of **1a–2f** using a multiscale approach⁴³ with ORCA^{44–46} provided a distribution of distances that contribute to the observed ROE signals (as seen in Fig. 3C). The relative areas derived from the MD simulations align qualitatively with the experimental 1D ROESY results (calculation details can be found in the ESI†). The pyridyl *C*-3 protons (depicted in red in Fig. 3) and the aryl protons on *C*-6 (depicted in green in Fig. 3) are, on average, in closer proximity to the *tert*-butyl protons of the cation **1a** than the

pyridyl *C*-2 protons (depicted in blue in Fig. 3). These findings suggest that aromatic solvents with low polarity (such as toluene) coupled with lower temperatures result in a tighter association of the ion pair. This increased association enhances the ability of the chiral cation to exert its enantio-discriminating influence on substrates activated by the nucleophilic anion.

The solution-state structure of the ion pair catalyst **1a–2f** was determined *via* theoretical calculations. The top 26 conformations located⁴⁷ account for 95% of the total Boltzmann population. The top seven conformations, which represent 70% of the population, show **1a** binding to **2f** in a similar orientation. The primary differences between these conformations involve the *tert*-butyl groups' arrangement (Fig. 3D). Electron density-based independent gradient model (IGM) analysis^{48,49} indicates that the main intermolecular interaction consists of non-classical hydrogen bonds (NCHBs) between the benzylic C–H or aryl C–H and the sulphonamide group's oxygens (Fig. 3E). The intermolecular interaction between **1a** and **2f** is estimated to be

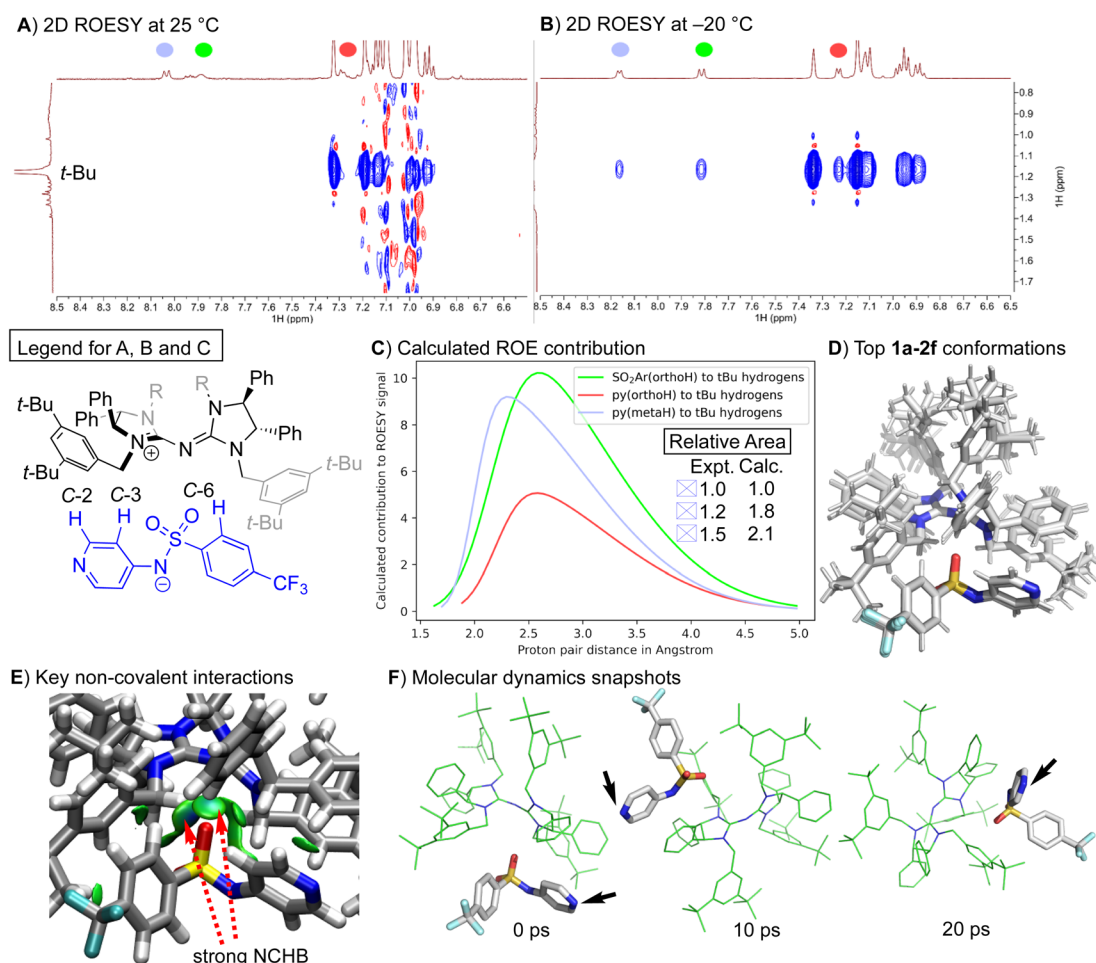
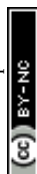


Fig. 3 Assignment of peaks to the structure is indicated by the colour code in the legend. (A) Selected region of the 2D ROESY spectrum of **1a–2f** in toluene-*d*₈ at 25 °C. (B) Selected region of the 2D ROESY spectrum of **1a–2f** in toluene-*d*₈ at –20 °C. (C) 1D ROESY contribution estimated from molecular dynamics simulation (refer to the ESI† for details) and experimental integrals from 1D ROESY of selected peaks. (D) Overlay of the top seven conformations for **1a–2f** optimized at ALPB(toluene)/GFN2-XTB:r²SCAN-3c. (E) NCHB: Non-classical hydrogen bond. Independent Gradient Model (IGM) isosurface (isovalue at 0.01 a.u.) visualized by VMD⁴⁰ to reveal key intermolecular interactions between **1a** and **2f**. (F) MD snapshot from a 20 ps trajectory of the most stable conformation at the same level of theory. Black arrow indicates the position of the nucleophilic nitrogen of the anion **2f**.



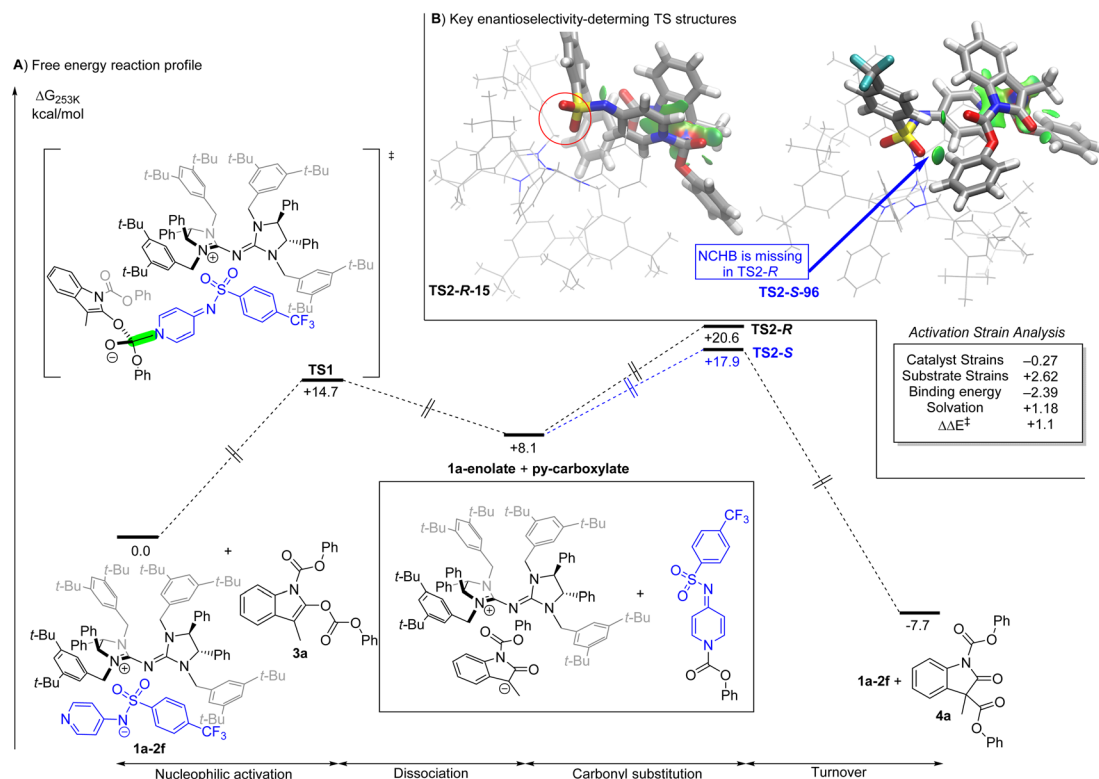


Fig. 4 A) Free energy profile of the **1a–2f** catalyzed enantioselective Steglich reaction. Values of free energy profiles are calculated at SMD(toluene)/M06-2X⁵⁴/def2-TZVPP//ALPB(toluene)/GFN2-XTB:r²SCAN-3c. (B) Activation strain analysis on the best TS structures from each set of TS: negative values favor TS2-R-15, while positive values favor TS2-S-96 (leads to the experimentally observed enantiomer). IGM isosurface (0.01 a.u.) based on promolecular density to highlight the origin of the substrate strain.

–36 kcal mol^{–1} from promolecular density with IGM analysis. The nitrogen atom on the pyridine ring faces outward, positioned ideally for substrate activation during the reaction. This orientation is maintained despite the facile anion's movement around the cation (black arrows in Fig. 3F). This can be attributed to the kinetically stable NCHB between **1a** and **2f** (Fig. 3E).

The chiral ion pair catalysed Steglich rearrangement⁵⁰ is calculated to proceed *via* a stepwise mechanism (Fig. 4). The pyridinyl-sulphonamide anion **1f** activates the reactant **3a** *via* TS1 (nucleophilic activation in Fig. 4). The dissociation of the intermediate to the **1a**-enolate and py-carboxylate allows the subsequent product formation step to occur. The nucleophilic C-attack on the carbonyl group of the py-carboxylate furnishes the oxindole product **4a**, together with regeneration of the catalyst. The turnover frequency is calculated to be 6.8 h^{–1} with TS2 being the turnover-determining TS.^{51,52}

The calculated ee value of 93% from the ensemble of 69 R- and S- carbonyl substitution transition state (TS) structures is higher than the experimental ee value of 79% (Table 1, entry 11). The % ee calculated from $\Delta\Delta H^\ddagger$ is 71%, which is in good agreement with the experimental value. The discrepancy arises mainly from the entropy contribution that is likely to have significant numerical noise due to the numerical nature of the hessian. Full DFT on the selected optimized TS reduces the positive contribution of entropy to the calculated level of enantioselectivity (see the ESI[†]). Given the considerable

reduction in computational effort by using a multiscale method, the agreement is reasonable.

An activation strain analysis⁵³ of the two most stable conformations from the ensemble of TS structures reveals that the key contributions to enantioselectivity arise from the more stable substrate geometry in the TS and solvation. As revealed by the IGM analysis, the substrates in TS2-S-96 have essentially one additional NCHB between the SO₂ of **1f** and aryl C–H, which leads to a more favourable substrate geometry and thus less strain. From the less favourable binding energy of TS2-S-96 relative to TS2-R-15, it can be concluded that the geometry in TS2-S-96 was adopted to maximize substrate stability and solvent interaction.

Conclusions

In conclusion, we have extended the capability of chiral cation-directed catalysis by pairing with anionic nucleophiles. In this work, the nucleophilic pyridinyl-sulphonamide works synergistically with pentanidium to form an ion pair capable of inducing a high level of enantioselectivity. This chiral ion pair catalyst has been successfully utilized in the enantioselective Steglich rearrangement of oxindole derivatives, achieving up to 99% ee. Through ROE integrals obtained under experimentally relevant conditions, we conclude that a tightly bound ion pair existence is solvent and temperature dependent. Its existence is

important for the high level of enantioselectivities observed in the Steglich rearrangement. Molecular dynamics simulation provided microscopic details on the structural preferences of the ion pair catalyst. A stepwise mechanism and the origin of enantioselectivity were postulated *via* theoretical calculations. This study demonstrates the potential for pairing chiral cations with achiral anionic organocatalysts through hydrogen bonding. We anticipate that this concept could also be extended to organocatalysts that are activated *via* halogen bonding or carbene formation.

Data availability

The data that support the findings of this study are available in the ESI† of this article. Data for this paper, including computational results, total energies, thermochemical corrections, and cartesian coordinates of all optimized structures are available at FigShare at <https://doi.org/10.6084/m9.figshare.23294474>.

Author contributions

Z. Y. and C.-H. T conceived the project and contributed to the methodology development. Z. Y. performed the experiments (catalysts synthesis, reaction testing, and characterization). C. X. and X. Z. contributed to part of experimental data. Z. Y., C. W. K. and C.-H. T. wrote the manuscript. C. W. K. performed the theoretical calculations. C. B. C. supervised the NMR experiments performed by Z. Y. All authors contributed to revising the manuscript.

Conflicts of interest

The authors declare no conflict of interest.

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