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Enantiomerically enriched tetrahydropyridine allyl chlorides†

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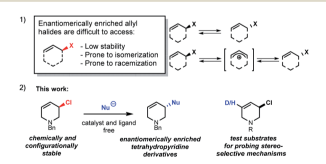
Enantiomerically enriched allyl halides are rare due to their configurational lability. Here we report stable piperidine-based allyl chloride enantiomers. These allyl chlorides can be produced via kinetic resolution, and undergo highly enantiospecific catalyst-free substitution reactions with C, N, O and S-based nucleophiles. DFT calculations and experiments with deuterium-labelled chloro-tetrahydropyridine, selectively prepared using H/D primary kinetic isotope effect, were used to investigate the mechanisms of resolution and substitution reactions. The allyl chlorides may also serve as valuable mechanistic tools for probing stereoselective reaction pathways.

Introduction

The stereoselective synthesis of molecules with chlorosubstituted stereogenic centres is challenging. Despite considerable effort, few asymmetric chlorination methods have been reported.¹⁻³ Synthesis of enantiomerically enriched allyl halides has proven to be especially difficult as they are generally prone to fast isomerization and racemization (Scheme 1(1)).4

Enantioenriched allyl chlorides are virtually unknown, except as a feature of diastereomeric compounds, where other stereogenic elements exert control over the observed stable allyl halide configuration.2d,5 The selective preparation of enantiomerically enriched allyl chlorides^{5a} could be important as stereogenic centres containing C-Cl bonds are found in

Results and discussion



Scheme 1 Properties of allyl halide enantiomers and this work.

pharmaceuticals, 56,6 and the allyl halides may undergo stereospecific reactions.1,6,7

Tetrahydropyridines (THPs) are a subgroup of nitrogen heterocycles found in biologically active molecules.8 THPs are also important precursors to functionalized piperidines. Piperidine is among the most common motifs found in licensed pharmaceuticals,8a and is frequently found in best-selling brand-name medicines. 9,10 3-substituted piperidine derivatives are at the core of many potent therapeutic agents (Fig. 1), 9,11,12 and methods for their preparation have attracted growing attention.13 However, asymmetric syntheses of 3-substituted piperidines via direct functionalization is scarce. 14

Due to the importance of THPs, racemic 3-chloro-1,2,3,6tetrahydropyridines were examined in asymmetric allylic additions (AAAs) with Zr-nucleophiles and copper catalysts. To our surprise, we were able to isolate enantiomerically enriched allyl chlorides from the reaction mixture. The kinetic resolution of

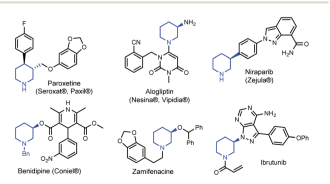


Fig. 1 Examples of biologically active 3-substituted piperidines.

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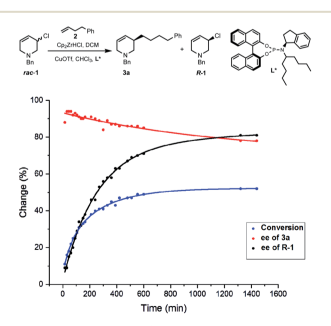
halides has received little attention,^{2e,15} with resolution of allyl halides being limited to a single report of activated allyl fluorides.¹⁶ Here we report the synthesis of enantioenriched allylic 3-chloro-THPs *via* kinetic resolution and investigate alternative preparation methods. The chemically and configurationally stable allyl chlorides can be used to prepare a wide range of THP derivatives that may be useful in synthesis and medicinal

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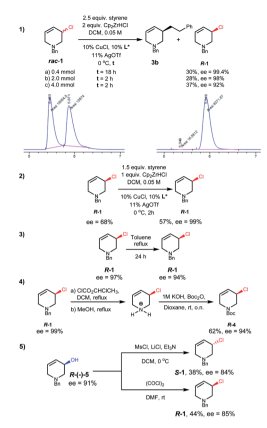
chemistry (Scheme 1(2)).

Initial exploration¹⁷ of Cu-catalyzed AAA with chlorotetrahydropyridines showed recovery of scalemic allyl chloride, indicating slow (or indeed no) interconversion between starting material enantiomers during the reaction. The reaction of 1-benzyl-3-chloro-1,2,3,6-tetrahydropyridine (*rac-1*) with 4-phenyl-1-butene (2) was followed in time (Scheme 2). After 30 minutes at –10 °C the reaction reached 16% conversion, giving product 3a in 94% ee, with the ee of 3a then decreasing ~15% due to slow consumption of the less reactive starting enantiomer over time. The ee of 1 increased from 0 to 80% over 22 hours, where it remained unchanged (within experimental error).¹⁷ Allyl chlorides bearing different protecting groups (Boc, Cbz, Ts and Ms) were tested in Cu-catalyzed AAAs with 4-phenyl-1-butene,¹⁷ however; these reactions were low yielding and not stereoselective.

We then investigated conditions to optimize the ee of 1.¹⁷ With styrene, diluting the reaction conditions, and using DCM as solvent, *R*-1 was obtained in useful yield with excellent selectivity (Scheme 3(1a)). Kinetic resolution was found to be much faster on larger scales (Scheme 3(1b and c)). These reactions also afforded alkylation product 3b in 49–65% yield and 84–88% ee. The mass balance in these reactions is good in that neither the small nor large scale reactions of *rac*-1 with styrene or 2 gave detectable amounts of side products. Conveniently, scalemic allylic chloride samples can be recycled, with



Scheme 2 Kinetic resolution of *rac-*1 using a copper–ligand catalyst and an alkylzirconium reagent.



Scheme 3 (1) Kinetic resolution of rac-1 and HPLC traces of racemic and enantiomerically enriched 1 using a chiral non-racemic stationary phase. (2) Kinetic resolution of scalemic 1. (3) Configurational stability of R-1 to heating. (4) Removal and addition of a protecting group to give N-Boc protected R-1 with little loss of enantiomeric excess (5) alternative synthesis of R- and S-1 from an enantiomerically enriched alcohol R-5 via selective inversion (to give S-1) or selective retention (to give R-1).

resolution on such material giving up to 99% ee using lower equiv. of alkyl-Zr reagent (Scheme 3(2)).

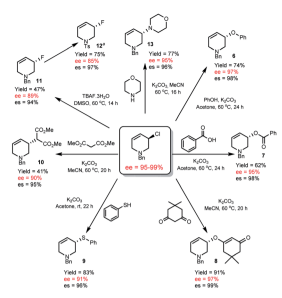
R-1 shows remarkable thermal stability; in toluene heated to reflux for 24 hours only 3% ee erosion was observed without any detectable side product formation (Scheme 3(3)). If desired the *N*-protecting group of *R*-1 can be exchanged, for example to *N*-Boc, through a pyridinium salt intermediate with only a small decrease in ee (Scheme 3(4)).¹⁷

Alternative routes to enantioenriched **1** were tested (see ESI†). Enantiomerically enriched (Scheme 3(5), 91% ee) alcohol *R*-5, after optimization,¹⁷ could be chlorinated with either inversion or retention, and reasonable stereospecificity, to give either enantiomer of **1**.

Though thermally stable, *R*-1 is a versatile chiral nonracemic building block and undergoes a variety of highly stereospecific substitution reactions. Using mild conditions we were able to access a variety of THP derivatives including ethers (6), esters (7), vinylogous ethers (8), thioethers (9), malonates (10), fluorides (11) and amines (13) with very high stereospecificity (94–99% es, Scheme 4).

The absolute configuration of *R*-1 was assigned¹⁷ by converting allyl fluoride 11 to 12 (Scheme 4), which has been

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Scheme 4 Enantiospecific substitution reactions of R-1. es = % enantiospecificity; es = [(ee of product/ee of R-1) \times 100]. ^aThe absolute configuration of 12 is known, see ESI† for the conditions to prepare 12.

determined by X-ray crystallographic analysis, 18 and knowledge (vida infra) that R-1 to 11 occurs via an S_N2 pathway.

To clarify the mechanism of stereospecific substitution we prepared isotopically labelled rac-1-d. Reduction of benzylpyridinium bromide in CD₃OD selectively adds D to the C3 position of 14-d, which was epoxidized to 15-d. We used the primary kinetic isotope effect of H/D deprotonation19 as a strategy to prepare deuterium-labelled allyl alcohol 5-d. This sequence gave a 2.4:1 D/H ratio at C5 after optimization.17 Rac-1-d was obtained with 61-65% D saturation as judged by ¹H and ²H NMR spectroscopic experiments after chlorination (Scheme 5(1)).20 Acyclic allyl chlorides generally undergo S_N2'

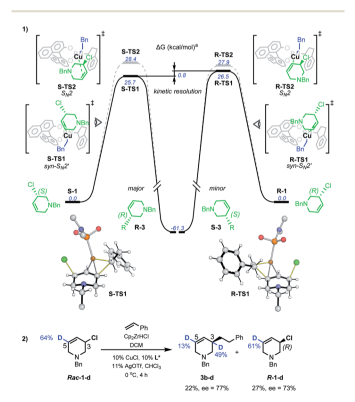
13%

Scheme 5 (1) Preparation of *rac-*1-d. $^{a}\%D = [n_{5-d}/(n_5 + n_{5-d})] \times 100$. (2) Mechanistic studies on substitution reactions. ^bConditions for substitution reactions are the same as shown in Scheme 4

substitutions as the carbon with the leaving group can freely rotate about the vicinal olefin,21 but mechanistic studies with cyclic allyl chlorides suggests S_N2 substitutions are generally favoured followed by anti-S_N2' and syn-S_N2' pathways. 4e-h,22

Our results (Scheme 5(2)) show that the %D saturation at C5 of rac-1-d is usually conserved throughout these substitutions, strongly suggesting S_N2 reactions. An exception to this trend is seen with thioether 9-d which shows some D transfer to C3. As the es of R-1 to 9 is high (96% es), it suggests that the reaction occurs by a mixture of pathways, in favour of S_N2, with both S_N2 and syn-S_N2' leading to the same enantiomer. Density Functional Theory (DFT) calculations excluded transannular aziridinium ion formation due to the large free energy difference between an azabicyclic intermediate and 1 (37.8 kcal mol⁻¹).¹⁷ Such levels of energy necessarily imply unfavourable transition structures to access it (TS \geq 37.8 kcal mol⁻¹), thus eliminating the possibilities of an intra-nucleophilic reaction pathway with double inversion.

DFT studies were also used to probe the kinetic resolution of rac-1 (Scheme 6(1)). Geometry optimisations were carried out with M062X functional and the 6-31G(d) basis set for C, H, N, O, P, Cl atoms and the LANL2DZ effective core potential/valence double zeta basis set for Cu. Single point energy corrections were obtained at the M062X/def2-TZVPP level of theory with chloroform solvation described by an implicit Solvation Model based on Density (SMD), then corrected using D3-dispersion energy as developed by Grimme.17



Scheme 6 Mechanistic study of Cu-catalysed AAAs. aSMD-M062X/ def2-TZVPP//M062X/6-31G(d)/LANL2DZ computed Gibbs energies at $298.15 \text{ K and 1 mol L}^{-1}$

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Starting from a Cu-L* complex in which the ligand *n*-alkyl groups were abbreviated in computational models, ¹⁷ rac-1 can bind to the metal centre at the N lone-pair. Two diastereomeric complexes can form with similar stabilities: of which S-C1 $(0.8 \text{ kcal mol}^{-1})$ is slightly less favourable than **R-C1** (-0.2 kcal mol⁻¹). Such a qualitative difference could potentially suggest a role in stereoselectivity and explain the specificity observed with N-benzyl-protected allyl chlorides. 17 For displacement of chloride by the catalyst, five possible pathways were investigated: syn- and anti- oxidative addition, anti-S_N2', S_N2 and syn-S_N2'. Alternative conformations were considered for each possibility.17 The syn-S_N2' transition structure (TS) S-TS1 was the most favourable overall, proceeding from the reaction of S-1 with a barrier of 25.7 kcal mol^{-1} to give the (R)-product of alkylation. For the (S)-enantiomer, this was followed by the $S_N 2$ TS S-TS2 (28.4 kcal mol^{-1}). Copper-catalyzed allylic alkylations are often described as occurring through complexation of a Cu(1) complex to the allylic olefin followed by oxidative addition to generate an allyl-Cu(III)²³ complex, but here anti-S_N2', syn- and anti-oxidative addition were found to be comparatively unfavourable, except with the anti-S_N2'-oxidative addition pathway (S-TS3) that is only 0.1 kcal mol⁻¹ higher than S-TS2.¹⁷ Our computational model is nevertheless qualitatively correct to disfavour S-TS3 over S-TS2 as it would otherwise give the opposite enantiomer than observed experimentally. Similarly to S-1, R-1 was found to preferentially react via a syn-S_N2' pathway, with the barrier for R-TS1 at 26.5 kcal mol⁻¹. As found for S-1, the next most stable was $S_N 2$ TS *R*-TS2 at 27.9 kcal mol⁻¹. Based on these computations, the alkylation of R-1 is kinetically disfavoured vs. S-1 ($\Delta \Delta G_{\text{syn-S}_N}^{\frac{1}{2}} = 0.8 \text{ kcal mol}^{-1}$) via dominant syn-S_N2' pathways for both enantiomers. This is consistent with the absolute sense of enantioenrichment observed experimentally, in which the (R)-enantiomer of both substrate and product accumulates. In addition to the major syn-S_N2' pathway, our

D-labelled rac-1-**d** was subjected to Cu-catalyzed resolution (Scheme 6(2)), and in accordance with computation, both S_N2 and S_N2 pathways are operative: product 3b-**d** shows D-incorporation at the C3 (49%) and C5 (13%) positions consistent with syn- S_N2 ' and then S_N2 being the most favourable pathways. No deuterium-isomerization of starting 1-**d** was observed and R-1-**d** was isolated from the reaction mixture with 73% ee.

result possibly implicate involvement of a minor S_N2 mecha-

nism to give the same product stereochemistry.

Conclusions

We have found allyl chlorides that are thermodynamically stable and can be prepared in highly enantioenriched form. A rare kinetic resolution of allylic chlorides formed piperidine-based allyl chlorides with high ee. The allyl chloride enantiomers can be separated by chromatography using a chiral non-racemic stationary phase, or prepared by stereospecific (with retention or inversion) chlorination of the corresponding alcohol. The allyl chloride undergoes highly enantiospecific substitution reactions with nucleophiles to give a wide range of tetrahydropyridine products which are important in biology

and medicine. Experiments with D-labelled chlorotetrahydropyridine and DFT calculations were used to investigate the mechanistic pathways of nucleophilic substitution and kinetic resolution reactions. This work provides new routes for the formation of enantiomerically enriched tetrahydropyridine derivatives. Further, *rac-*1, *R-* or *S-*1, *R-*1-d and other configurationally stable allyl halides may serve as probe substrates for investigating reaction pathways in mechanistic studies of reactions.²⁴

Conflicts of interest

Oxford University Innovation has filed a patent application (GB1815018.5) with S. P. F. and S. K. named as inventors. The remaining authors declare no competing financial interests. A version of this manuscript has been shared and posted on ChemRxiv.

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relocation of D by S_N2 ' transformations. However, the latter pathway likely contributes only at minute levels as D atoms at C3 position of **rac-1-d** could only be detected by sensitive 2 H-NMR spectrscopic analyses. See ESI.†

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