Foldamer-Mediated Remote Stereocontrol: > 1,60 Asymmetric Induction**

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Abstract: An N-terminal l-α-methylvaline dimer induces complete conformational control over the screw sense of an otherwise achiral helical peptide foldamer formed from the achiral quaternary amino acids Aib and AcC. The persistent right-handed screw-sense preference of the helix enables remote reactive sites to fall under the influence of the terminal chiral residues, and permits diastereoselective reactions such as alkene hydrogenation or iminium ion addition to take place with 1,16-, 1,31-, 1,46- and even 1,61-asymmetric induction. Stereochemical information may be communicated in this way over distances of up to 4 nm.

In a typical stereoselective reaction, existing stereochemistry governs the formation of a new stereogenic center by controlling the direction of attack of a reagent or a catalyst at a reactive site. Close spatial contact between the controlling center and the reaction site is typically required, and 1,2- or 1,3-asymmetric induction (where the two sites are separated by one or two bonds) can routinely be expected to give high levels of stereoselectivity. Asymmetric induction over longer distances (“remote asymmetric induction” usually refers to 1,4-asymmetric induction and beyond) is possible, but only if the flexibility of the molecule is limited, usually by the (sometimes temporary) formation of a cyclic structure. Asymmetric induction can be achieved over more than 20 bonds in non-cyclic molecules by using semi-rigid structures in which relayed interactions between a series of polarized groups allow a controlling stereogenic center to influence the local environment of a remote reactive site.

Here we present a way to achieve asymmetric induction over much greater distances by using molecules that have inherent helicity, or foldamers. The conformational properties of foldamers as structural analogues of biomolecules have been investigated widely, but their ability to control reactivity remains largely unexplored. By linking an appropriate controller and a reactive site through a molecular fragment strongly disposed to adopt a helical structure, we show that it becomes possible for the controlling center to govern the stereochemical environment at a remote reactive site located several nanometers away.

The helical foldameric parts of our molecules were made from oligomers of aminoisobutyric acid (Aib, 1) and 1-amino-cyclohexanecarboxylic acid (AcC, 2) (Figure 1a). Peptidyl-like oligomers of these achiral quaternary amino acids typically adopt well-defined helical structures in solution, but because the individual amino acids each possess a plane of symmetry, their oligomers exist as a rapidly interconverting equal mixture of left- and right-handed helices. A bias towards a single screw sense was induced by ligating to the N-terminus of these oligomers one or more residues of the chiral quaternary amino acid L-α-methylvaline (αMv, 3). The structure of L-α-methylvaline is compatible with the helical structure of Aib oligomers and we reasoned that incorporating a sufficient number of chiral quaternary L-amino acid residues would maximize the chances of inducing a high degree of preference for the right-handed screw sense in the helical chain. Two oligomers were synthesized, 4 and 5, containing one and two αMv residues respectively (Figure 1b). Circular dichroism (CD) spectra of 4 and 5 in MeOH, and titration of 5 in THF with DMSO, were consistent with the adoption of a characteristic helical structure in these solvents (see the Supporting Information).

Before carrying out the remote stereoselective reactions, we quantified the screw-sense preference induced in the helical chains of the oligomers 4 and 5 by incorporating a 13C label asymmetrically into the two methyl groups of the fifth Aib residue of the chain. The anisochronicity of the two 13C labeled Me groups in the 13C NMR spectra in MeOH and in THF of both 4 and 5 confirmed that the population of left- and right-handed helical conformers is unequal, and the location of the majority of the label in the same (upfield) member of the pair of anisochronous signals, in conjunction with the CD spectra in MeOH, indicates that both 4 and 5 prefer a right-handed preferred screw sense. Analysis of the spectra acquired at a range of temperatures (Figure 1c and the Supporting Information) showed that while the oligomer 5 displays good conformational selectivity for the right-handed screw sense in MeOH, in THF (and especially at low temperature) the preference for a single, right-handed, screw sense becomes almost quantitative. At −50°C, for example, we calculate that the 13C-labelled residue of 5 finds itself in a right-handed helical environment more than 98% of the time. Data for both 4 and 5, in MeOH and in THF at +40°C (°C)

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and at −50°C, are shown in Table 1. Two chiral residues are evidently necessary to induce this almost complete screw-sense preference: studies of related compounds containing three chiral residues (see the Supporting Information) showed that the third offered no improvement in the degree of conformational control.

Support for the conclusion that an N-terminal l-αMv dimer provides close to quantitative screw-sense control was provided by conformational analysis of two related compounds, 6a Cbz(αMv)2AibGlyNH2 and 6b Ac-(αMv)2AibGlyNH2 (Figure 1e). The X-ray crystal structure of 6a shows a well-formed right-handed 310 helix, and density functional theory calculations on 6b indicate that the lowest energy left-handed helical conformation is more than 10 kJ mol⁻¹ higher in energy than the lowest energy right-handed helical conformation.

Our next challenge was to utilize the helical screw-sense preference to induce, at a distance, the formation of a new stereogenic center.

Scheme 1: Asymmetric induction of an l-Phe residue by hydrogenation of a didehydrophenylalanine residue embedded in an induced right-handed helix.

Table 1: Conformational preferences in 4 and 5 derived by analysis of their 13C NMR spectra.

<table>
<thead>
<tr>
<th>Compound</th>
<th>Solvent</th>
<th>T [°C]</th>
<th>(K^\circ)</th>
<th>(h.e.^\circ) [%]</th>
<th>(P:M^\circ)</th>
</tr>
</thead>
<tbody>
<tr>
<td>4</td>
<td>MeOH</td>
<td>+40</td>
<td>50</td>
<td>75:25</td>
<td></td>
</tr>
<tr>
<td>4</td>
<td>MeOH</td>
<td>−50</td>
<td>4.9</td>
<td>83:17</td>
<td></td>
</tr>
<tr>
<td>5</td>
<td>MeOH</td>
<td>+40</td>
<td>64</td>
<td>83:17</td>
<td></td>
</tr>
<tr>
<td>5</td>
<td>MeOH</td>
<td>−50</td>
<td>9.5</td>
<td>91:9</td>
<td></td>
</tr>
<tr>
<td>5</td>
<td>THF</td>
<td>+40</td>
<td>92.9</td>
<td>96:4</td>
<td></td>
</tr>
<tr>
<td>5</td>
<td>THF</td>
<td>−50</td>
<td>60</td>
<td>98.5:1.5</td>
<td></td>
</tr>
</tbody>
</table>

[a] Equilibrium constant \(K\) for interconversion of \(P\) and \(M\) conformers at −50°C, calculated by line shape analysis of the variation of the 13C NMR spectrum with temperature. [b] The value \(\Delta h.e. = \Delta h.e.\) (see Figure 1c,d) interpreted as “helical excess” (that is, \([P]−[M]/[P]+[M]\)). [c] Calculated ratio of \(P\) and \(M\) conformers derived from either \(K\) or h.e. [d] Value is identical upon two-fold dilution.
This unsaturated residue was converted into phenylalanine by chemoselective hydrogenation in the presence of Crabtree’s catalyst,\textsuperscript{20} [Ir(cod)(PCy\textsubscript{3})(ppy)]PF\textsubscript{6} -, in ethanol or in dichloromethane. Two diastereoisomeric products were formed in each case, and the synthesis of authentic samples of 10\textsubscript{a} and 10\textsubscript{b} from L- and D-Phe showed that the major compound was the L-Phe containing oligomer. The ratios of 9\textsubscript{a}:9\textsubscript{b} and of 10\textsubscript{a}:10\textsubscript{b} are shown in Scheme 1. Selectivities were higher in the reactions of 8 than of 7, and were higher in dichloromethane than in ethanol. Hydrogenation of 8 in dichloromethane gave more than 95:5 selectivity for the formation of 10\textsubscript{a}, which arises from attack of hydrogen on the Re-face of the double bond of 7 (the back face as shown in Scheme 1). Given the remoteness of the nearest stereogenic center, we propose that this selectivity arises purely from the preferred screw sense of the helical foldamer structure in which the reactive site is embedded. Lower selectivities with hydroxyl solvents and with a single controlling chiral residue presumably result from the lower levels of screw-sense control attained in these cases (see Table 1). Next, we sought reactions of terminal prochiral groups that would allow asymmetric induction to be relayed from one end of a foldamer chain to the other. A survey of the reactions of C-terminal reactive C-C, C=O, and C=N groups revealed that trapping of a C-terminal N-acyliminium ion by a nucleophilic arene takes place with excellent diastereoselectivity.

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With this evidence that the remote asymmetric reactions at the terminus of a helical foldamer may be mediated solely by the screw sense of the foldamer, we proceeded to lengthen the chain by successive homologation of 16 with achiral helical fragments 20 comprising Ac\textsubscript{3}cAlb\textsubscript{5} pentamers, as shown in Scheme 3a. A series of carboxylic acids 21–23 were coupled with allylamine to make the amides 24–26, and then isomerized to their enamide isomers 27–29. The enamides 27 and 28 were treated with trifluoromethanesulfonic acid in the presence of 1,3,5-trimethoxybenzene to give amide products 30 and 31 as mixtures of diastereoisomers (Scheme 3b,c). Authentic samples of both diastereoisomeric products were synthesized from 21 or 22 and the enantiomers of 15, and comparison by \textsuperscript{1}H NMR (800 MHz, [D\textsubscript{3}],[D\textsubscript{6}]THF) of these compounds with the product mixtures 30 and 31 by \textsuperscript{1}H NMR showed that selectivities remained close to 10:1 in these reactions, even as the remote relationship between the controlling center and the newly formed asymmetric center was extended from 1,16 in 14 to 1,31 in 30 to 1,46 in 31.

Finally, under the same conditions, enamide 29 yielded two diastereoisomers of the product 32 in a ratio of 88:12 (Scheme 3d). This reaction displays 1,61 asymmetric induction, a value almost three times the greatest through-bond distance previously reported,\textsuperscript{23} representing the communication of stereochemical information over a distance of about 4 nm,\textsuperscript{24} through more than six induced right-handed turns of the helical structure.

To summarize, we have shown that the screw sense of a molecular helix, induced under thermodynamic control, can act as the mechanism by which information may be transmitted over long distances (over 60 bonds) on the molecular scale. By inducing complete preference for right-handed helicity in a chain having no inherent screw-sense preference of its own, an appropriate controller enables distant reactive sites to fall under its influence, resulting in a reaction displaying 1,61 asymmetric induction. Such remote control of chemical reactivity is characteristic of allosteric enzymes and receptors, and conceptually similar designs could be used.
to make artificial analogues of such biological signal transduction mechanisms.

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Scheme 3. Remote asymmetric induction through achiral helices. a) Synthesis of the three oligomers 21–23 and their conversion to the enamides 27–29. b) 1,3Remote asymmetric induction through nine achiral residues to yield 30. c) 1,4Remote asymmetric induction through fourteen achiral residues to yield 31. d) 1,6Remote asymmetric induction through nineteen achiral residues to yield 32.


