

REMOTE CONTROL OF STEREOCHEMISTRY: COMMUNICATING INFORMATION VIA CONFORMATION

JONATHAN CLAYDEN

School of Chemistry, University of Manchester, Oxford Road, Manchester M13 9PL, U.K.

E-Mail: j.p.clayden@manchester.ac.uk

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ABSTRACT

Stereochemistry is information, and stereoselective reactions are the means by which that information may be communicated within and between molecules. The control of remote stereogenic centres can be achieved by stereochemical relay, and the use of thermodynamic control over conformational preference is turning out to be a very powerful method for long-range transmission of stereochemical information.

Introduction

Stereoselective synthesis, which aims to control relative and absolute configuration at new stereogenic centres, has been a remarkable success story [1]. The award, in 2001, of a Nobel Prize to three of the founding thinkers in the area of asymmetric catalysis [2] confirmed as one of the great achievements of late 20th century chemistry the science of stereoselectivity. Stereochemistry is information – binary information – and stereochemical control, mediated by stereoselective reactions, can be seen as a chemical means of information transfer – of communication of information. The reliability of the communication depends on a handful of interrelated factors, but it is clear that stereocontrol is best when proximity and rigidity are features of the reaction sites and molecules involved: it has long been a stereochemical tenet that flexible acyclic molecules provide generally poor substrates for stereoselective reactions.

The need for stereoselective reactions to take place close to the stereocontrolling influence in general limits the distance through which stereocontrol can take place to some four or five bond lengths. The communication of stereochemistry over greater distances than this is generally described as *remote stereocontrol*, or *remote asymmetric induction* [3]. Stereocontrol apparently reaching over distances of up to 12 or 13 bond lengths has been achieved by using a metal ion to grab a remote reaction site and bring it into temporary proximity with a stereocontrolling influence [4]. But stereocontrol at a reaction site truly distant in space generally requires specific structural features to allow stereochemical information to be communicated – "unrelayed" 1,5- stereocontrol with a selectivity of 90:10 has been described as "remarkable" [5]. These structural features "relay" the stereochemistry to the new site.

Stereochemical relays may take the form of temporary stereogenic centres, which bridged the gap between the two reaction sites. For example, Paterson's intermediate 3 en route to ebelactone [6] contains a 1,5-relationship between an isolated stereogenic centre and its nearest neighbour. After first setting up a proximal 1,2-relationship in 2 using a stereoselective aldol reaction of 1, a stereospecific Ireland-Claisen {3,3}-sigmatropic rearrangement expands the 1,2-relationship to a 1,5-relationship (Scheme 1).

Scheme 1. Stereospecific rearrangement as a means of relaying stereochemistry.

The "temporary" stereogenic centre need not be present in an isolable synthetic intermediate, but instead might be formed transiently at some point along the overall reaction pathway. Thomas [7] has achieved up to 1,7 stereocontrol in alkenes such as **6** using this strategy: the tin-bearing centre in the intermediate **5** is under the control of the hydroxyl group by virtue of tin-oxygen coordination within a six membered ring (Scheme 2).

Scheme 2. Relaying stereochemistry through a transient stereogenic centre.

Scheme 3 shows both strategies combined: the strategy of Scheme 2 controls the 1,7-relationship in 7 which is extended, by the strategy of Scheme 1, to a 1,9-relationship in 8 [7].

Scheme 3. Extending a remote stereochemical relationship.

A more subtle means of achieving stereocontrol from a distance makes use of a controlling centre to orientate not temporary *configuration* but simply to bias the *conformation* of that part of the molecule lying between the reaction site and controlling influence. Still very elegantly demonstrated the power of such methods for stereocontrol where the conformation of a macrocyclic ring relays the stereochemistry [8]. A conformational relay effect accounts for the protecting group dependence of the stereoselectivity exhibited by alkylation reactions of the phenylmorpholinones 9 (Scheme 4) [9]. Changing the planar *N*-Boc protecting group of 10a to the conformationally mobile *N*-benzyl group of 10b *inverts* the 1,3-diastereoselectivity in the alkylation of 12, the sodium enolate of 9. The Boc group is

stereochemically inert, and an "inherent" *anti* stereoselectivity is observed in reactions of **12a**. However, in **12b** the steric interaction between the phenyl-bearing centre and the *N*-benzyl group forces the benzyl group onto the top face of the molecule, and favours alkylation from the bottom face. The *N*-benzyl group of **12b** facilitates the relay of stereochemical information from the old centre to the new.

Scheme 4. Relay via an N-benzyl group.

In the alkylations of diketopiperazines 13, a switch of protecting group from conformationally inconsequential N-Me to conformationally powerful N-p-methoxybenzyl by contrast enhances selectivity (Scheme 5) [10]. While 13a (R = H) is alkylated with 2:1 diastereoselectivity, 13b (R =p-MeOPh) is alkylated with diastereoselectivity of 30:1. Modelling suggests that the enolate 16 derived from 13b adopts a conformation (Scheme 5) in which steric repulsion between the adjacent isopropyl and p-methoxybenzyl group favours conformations with this group lying anti to the isopropyl group. In turn, this N-substituent pushes the second N-substituent onto the bottom face, directing the electrophile to approach the enolate from above. Effectively, stereochemical information is relayed via both benzyl groups to the reacting centre.

Scheme 5. Stereochemical relay via two N-PMB groups.

Renaud has developed a related concept [11] in which stereochemistry is relayed from a chiral *catalyst* to a relatively distant reaction centre through the conformation of an intervening bond (Scheme 6) [12]. The agent of the relay takes the form of an "achiral auxiliary", an oxazolidinone derivative of 2-aminophenol. Complexation of chiral Lewis acid MgBr₂.19 to acryloyl oxazolidinone 17 is proposed to favour one of the two diastereoisomeric conformations 20 about the exocyclic N-CO bond, and indeed increasing the size of R from H through Me, Et, Bn and PMB gave a corresponding increase in product enantiomeric ratio (er) [13] from 57:43 to 94:6 as the differentiation between the faces of the acrylamide is increased.

Scheme 6. Stereochemical relay through a transient N-CO axis.

In related reactions of 21, a temporary, labile (i.e. kinetically unstable) stereogenic nitrogen centre relays stereochemistry from the chiral catalyst to a reaction centre (Scheme 7) [11, 14]. The "achiral auxiliary" in this case is a pyrazolidinone, readily available by the conjugate addition of hydrazine to 3,3-dimethylacrylate. The synthesis allows for easy variation of the substituent R, and the results suggest that this group R has an important role to play in relaying the effect of the chiral Lewis acid to the reaction site: the bigger the group R the higher the selectivity. The tetrahedral N atom inverts rapidly, but in the presence of a chiral Lewis acid it prefers the configuration shown as 24a or 24b, converting the achiral (or *prochiral*) auxiliary into a chiral auxiliary. The chiral ligand biases the stereochemistry at N which in turn, being relatively close to the reaction centre, controls the face-selectivity of the Diels-Alder reaction.

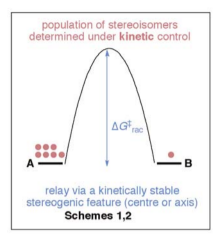
Scheme 7. Stereochemical relay through a transient stereogenic centre at N.

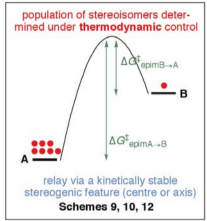
A similar effect is evident in the conjugate addition of nucleophiles to **21** in the presence of chiral magnesium Lewis acids [15]. Introducing larger *N*-substituents R apparently turns on the relay effect.

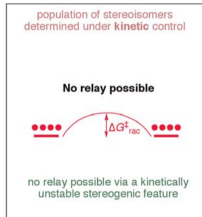
Relays of stereochemical information from chiral catalysts have an inbuilt drawback: the catalyst must activate the reaction centre and therefore interact with it in some way: the source of asymmetry can never be truly removed spatially from its effect.

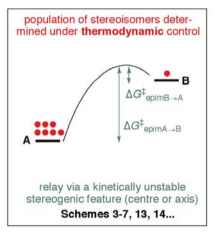
Unlike in Schemes 1 and 2, the stereochemical features proposed to give rise to the observed relay effects in Schemes 3-7 have a lifetime which is short on the laboratory timescale (though presumably long relative to the half-life of the reaction itself [16,17]). For example, although they have features of a chiral axis, the N-CH₂Ar bonds of **12b** and **16** and the N-CO bonds of **20** are not atropisomeric at the temperature of the reaction – in other words their chirality relies solely on the thermodynamic influence of a nearby centre, not on a kinetic barrier to bond rotation. The evidence that a true relay effect is involved is

therefore necessarily circumstantial – the dependence of the level of stereoselectivity in the reactions on the presence or absence of certain proposed stereochemical features. While the result of these reactions is not in doubt, the detailed mechanism by which stereochemical control is achieved is by no means clear, since it has not yet proved possible in these systems to study spectroscopically the conformation and stereodynamics of the bonds concerned. Scheme 8 illustrates schematically the various possibilities for achieving stereochemical relay.









Scheme 8. Kinetic and thermodynamic control in strategies for stereochemical relay. A and B represent two alternative stereo- or conformational isomers of the centre or bond facilitating the relay, in which one is more highly populated than the other due to either kinetic or thermodynamic control. A and B may or may not interconvert on the laboratory timescale.

The idea of temporarily storing information in an *atropisomeric* bond (i.e. one with kinetically stable stereochemistry which is not labile on the timescale of the reaction, and which for example can be studied by low temperature NMR techniques) was expanded by Curran [18] into the concept of "prochiral auxiliaries". Curran envisaged that atropisomeric stereochemistry generated under kinetic control would subsequently relay stereocontrol to further centres in a substrate. The idea has never been fully realized in its original form, and the most significant recent developments in the use of temporary atropisomeric chirality to relay stereochemical information have made use of thermodynamic rather than kinetic control over the temporarily stereogenic bond.

Kawabata and Fuji have shown that appropriately protected amino acids **25** may be alkylated apparently with retention of configuration via the hindered enolate **26** (Scheme 9) [19]. Extensive kinetic investigations have shown the origin of this "through time" relay effect (generally termed "chiral memory" [20], and to be contrasted with the "through space" relays above) is the conformation of the starting amide, coupled with the kinetic stability of the atropisomeric intermediate enolate. As the enolate **26** is formed from what semi-empirical calculations suggest to be the major conformer of **25**, the preferred orientation of its NR₂ group persists. Because C-N bond rotation is slow, and alkylation of the atropisomeric enolate is diastereoselective, the product **27** is formed largely retentively (90:10 er). Thus thermodynamic control over conformation is trapped and given kinetic stability by the formation of the atropisomeric (at low temperature) enolate **26**, the axial configuration of which persists even once the original controlling centre has disappeared [19, 21].

Scheme 9. Chiral memory – stereochemical relay through time.

Being very fast, radical reactions can often display chiral memory effects, and Curran has reported a reaction in which comparison of the temperature dependence of the stereose-lectivity and of the starting conformer distribution suggests that both chiral memory and chiral relay effects are in operation (Scheme 10) [17]. The stereogenic centre of **28** biases

the population of the two amide conformers in such a way that radical cyclization gives principally 30a rather than 30b. At low temperature the radicals 29 cyclize faster than they can interconvert, and while there is some stereochemical leakage from 29b to 30a the product ratio is largely the result of the ratio of conformers of 28. The importance of the axis in relaying stereochemistry is evident in related reactions of resolved *M*-31 which gives 32 with 90% stereospecificity [22].

Scheme 10. Chiral memory in radical cyclizsations.

RESULTS

Our own work [23, 24] has established the surprising ease with which stereogenic centres placed adjacent to certain functional groups – in particular tertiary aromatic amides such as 33 – are able to govern their orientation, often with very high degrees of conformational

control. Scheme 11 summarizes some of the substituents which display this ability, notably sulfoxides such as **36**, which give rise to conformational ratios about Ar-CO of up to 200:1 [25].

Scheme 11. Typical conformational ratios in tertiary amides bearing adjacent stereogenic centres.

The relatively slow rate of rotation about bonds in aromatic amides [26] meant that we have been able to quantify conformational ratios by NMR, and we have used the ability of a stereogenic centre to control the orientation of a functional groups as a means of synthesizing new classes of non-biaryl atropisomers enantioselectively [27, 28]. However, given that we were also well aware that atropisomeric amide axes are able to control the formation of new stereogenic centres [29, 30], it occurred to us that by placing these two aspects of stereoselectivity in series we could use a rotationally restricted amide as a means of relaying stereochemical information (Scheme 12). The sulfoxide group of 36 is introduced straightforwardly by quenching the lithiated amide (±)-34 with Andersen's menthyl sulfinate 35. The sulfoxide is then removed by sulfoxide-lithium exchange, and like the enolate 26, organolithium *M*-34 retains its absolute stereochemistry at least over a period of minutes at low temperature, long enough to relay its stereochemistry to the new stereogenic centre of 37. Lactonization gave *ent*-O-methylisoochracein 38 in 92:8 er [31].

Scheme 12. Chiral memory via the Ar-CO axis of a tertiary aromatic amide.

Amide conformation can be used as a means of relaying stereochemical information through space as well as time. Our first spatial stereochemical relay [24, 32] used the racemic silane $\bf 39$, in which steric effects impose an $\bf 88:12$ conformational preference on the amide (quantified by NMR in CDCl₃ at room temperature), to synthesize the meso amide $\bf 41$ (Scheme 13). The lithio derivative of $\bf 40$ has low kinetic stability on the laboratory timescale, even at low temperature [33], but the persisting conformational influence of the adjacent silyl-bearing centre ensures the success of the relay.

Scheme 13. 1,5-Stereochemical relay through an amide axis.

A reliable source of conformational control over aromatic amides, and one which is easily introduced in enantiomerically pure form, is the (-)-ephedrine-derived oxazolidine featured in 44. It was known that condensation of an aromatic aldehyde with (-)-ephedrine 43 leads preferentially to one of the two possible diastereoisomeric oxazolidines [34]. Intriguingly, though, an adjacent amide will adjust its conformation to ensure that the oxygen atoms of oxazolidine and carbonyl group, and likewise their bulky substituted nitrogen atoms, lie on opposing faces of the ring which bears them [24, 35]. Thus condensation of 42 with 43 gives 44, which adopts preferentially the conformation shown and retains that conformation through the formylation and addition reactions leading to 46. Stereoselectivity of Grignard additions is governed by amide conformation [30], and reaction of 45 with PhMgBr is fully diastereoselective (Scheme 14) [24].

Scheme 14. 1,5-Stereocontrol by stereochemical relay from an oxazolidine [36].

In all of the examples above, stereochemical relay induces asymmetry at an otherwise locally achiral remote site in a molecule. That otherwise achiral site could alternatively be a point of coordination for a metal. Scheme 15 and Table 1 show how the phosphines 47 [35] and 48 [36], whose chirality in the vicinity of phosphorus is the consequence of stereochemical relay from their chiral centres via the amide, can be moderately effective chiral ligands in the palladium-catalysed asymmetric allylic substitution of acetate 49 by dimethyl malonate (Scheme 15).

local Ar–CO conformation generates chiral environment at P

Ph₂P

Centre controls Ar–CO conformation

Ph₂P

Centre controls Ar–CO conformation

AT

Conformation

OAC ligand (1 mol%)

Ph

MeO₂C

CO₂Me

[PdCl(
$$\pi$$
-C₃H₅)]₂ 0.5 mol%

MeC(OSiMe₃)=NSiMe₃, CH₂Cl₂, KOAc, 25°C

Scheme 15. Enantioselective allylic substitution catalysed by palladium in the presence of amidophosphine ligands.

entry	ligand	Time	yield	product	er
1	47	3 days	60	(-)-50	95:5
2	48	24 h	93	(-)-50	91:9
3	52	24 h	85	(+)-50	23:77

Table 1. Ligands for the enantioselective substitution of 45.

In 1989, the crystal structure of **51** was reported [37]. The benzene-1,2,3-tricarboxamide adopts the conformation shown in Scheme 16: each amide carbonyl group points in a direction opposing its neighbours – presumably controlled by steric or electronic (dipole) interactions or both. On the assumption that the preference for nearby tertiary amide groups to adopt mutually opposing conformations persists in solution, we made the amide **52**, a homologue of the ligand **48** [36]. The phosphine was included in an allylic substitution reaction and gave moderate enantiomeric excess (Table 1, entry 3), but importantly the major enantiomer of the product **50** was opposite to that generated in the presence of the ligand **48**. This is exactly what is to be expected if the amides lie opposed to one another: the local environment of the phosphorus atom in **52** is enantiomeric with the local environment of the phosphorus centre of **48**, despite the stereochemistry originating from the same enantiomer of ephedrine.

Scheme 16. Conformational interactions between adjacent amides.

The *anti* alignment preferred in aromatic dicarboxamides turns out to be quite general, even with more remote relationships between the amide substituents [38]. The NMR spectrum of xanthene-1,8-dicarboxamide 53, for example, contains a single 6 H singlet corresponding to the *gem*-dimethyl group, and double lithiation and electrophilic quench yields a compound 54 which in principle contains two stereogenic axes and may therefore exist as a pair of diastereoisomers. However, only a single diastereoisomer is obtained, which NMR showed to be C_2 symmetric, HPLC on a chiral stationary phase showed to be chiral and racemic, and X-ray crystallography showed to have the stereochemistry shown in Scheme 17 [39].

Scheme 17. Anti-preference in a xanthene-1,8-dicarboxamide.

Even amides borne on separate non-rigidly interconnected aromatic rings have a strongly preferred conformation. Amide **56** contains two stereogenic axes, but double ortholithiation of **55** gave only a single diastereoisomer syn-**56** (Scheme 18). We were able to prove its stereochemistry by HPLC on a chiral stationary phase: a separation into two enantiomers was just visible, indicating formation of racemic, C_2 -symmetric syn-**56** rather than achiral, centrosymmetric anti-**56** [40]. However, this C_2 -symmetric diastereoisomer turned out to be the less stable of the two, because heating in toluene gave a mixture of compounds in which the second, achiral diastereoisomer prevailed. The amides have a clear thermodynamic preference for the S_2 -relationship and biaryl 2,2'-dicarboxamides fall into the class of molecules which are "flexible with a defined shape" [41].

Scheme 18. Kinetic and thermodynamic stereocontrol in biphenyl-1,1'-dicarboxamides.

The question that posed itself at this point was how far stereochemistry could be communicated using conformational preference alone as the means of stereochemical relay. In 1988, Noe [42] had reported some chemistry which, though characterized by only poor stereoselectivity, gave the first indication that conformational control allows communication of stereochemical information over long distances. The anomeric effect forces paraformaldehyde oligomers to adopt a helical conformation in solution, and by tethering short chains of paraformaldehyde to a source of chirality it appears to be possible to induce the helix in 57 to prefer one of two diastereoisomeric conformations. Stereochemical information is transmitted to the new chiral centre in 58 by Grignard attack on a ketone. Stereoselectivity drops with chain length but still reaching 2:1 for the 1,10-relationship formed in the reaction shown in Scheme 19. The importance of the polyether helix in mediating an interaction between the origin and terminus of the stereochemical relay was confirmed by control experiments showing that a hydrocarbon chain of the same length failed to induce selectivity.

Scheme 19. Relay of stereochemistry via a helix.

Given that amides even with rather remote relationships are also capable of communicating with one another, we hoped to be able to use pairs of conformationally communicating groups to mediate remote stereocontrol by a form of staged stereochemical relay. The concept is illustrated in Fig. 1. A set of otherwise freely rotating groups u-z adopt, under the influence of a stereocontrolling influence at A, a single orientation which results in the transmission of information about the shape of A through u-z to B. Cowburn and Welland had demonstrated a similar concept using quantum dot magnets spaced 135 nm apart [43]: stereochemical relay using amide dipoles could reduce the scale of the system by a factor of about 300.

 $^{^1\,}$ The 1 and 8 positions of a xanthene ring are about 4.6 Å apart.

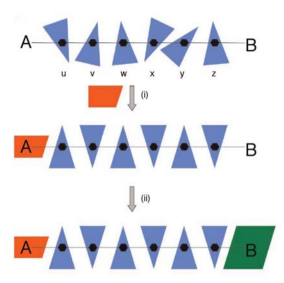


Figure 1. A strategy for the conformational relay of information from A to B. Groups u-z are freely rotating but on incorporation (i) of a stereochemical influence at A adopt an "all-anti" alignment, allowing information about the shape of A to be relayed to B via u-z. (ii) Formation of new stereochemistry at B can take place diastereoselectively despite the remoteness of A.

Stages in the development of this idea are shown in the Schemes 20 [32], 21 [39, 44] and 22 [40]. In Scheme 20, the conformation of the axis adjacent to the stereogenic centre of **39** is relayed round the ring by introduction of a second axis into **59** adjacent to the first. When the second stereogenic centre of **60** is constructed adjacent to this axis, its stereochemistry is under the ultimate control of the centre lying *para* across the ring [32]. In Scheme 21, the idea is taken a stage further: the two amides of **53** are already related by the *anti* conformational preference of such systems; introduction of an ephedrine-derived oxazolidine into **61** forces both axes to adopt a single absolute conformation. The stereochemistry of the oxazolidine is relayed through both amides and allows the subsequent addition of a nucleophile to the carbonyl group of **62** to proceed with complete (1,9)-stereochemical control, and **63** is formed as a single diastereoisomer [39, 44]. In Scheme 22, the amides also adopt conformations which allow the molecule as a whole to minimize its dipole moment. Thus the oxazolidine of **65** controls both amide groups and thus also ultimately directs nucleophilic attack on the remote carbonyl group of **66**, leading to complete 1,8-stereocontrol in **67** [40].

Scheme 20. Remote (1,6)-stereocontrol relayed through a benzene-1,2-dicarboxamide.

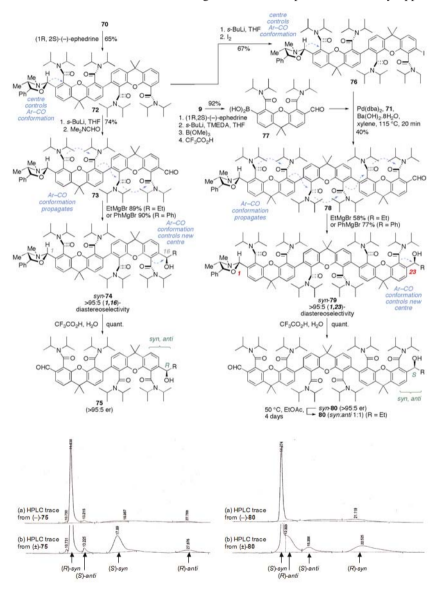
Scheme 21. Remote (1,9)-stereocontrol relayed through a xanthene-1,8-dicarboxamide.

Scheme 22. Remote (1,8)-stereocontrol relayed via a biphenyl-1,1'-dicarboxamide.

The next logical step was to make some molecules containing both the features of both 1,8-xanthenedicarboxamides and biphenyl-2,2'-dicarboxamides. Extensive work to establish optimum conditions for Suzuki couplings allowed us to synthesize the conformationally uniform (by NMR) bis-xanthene **70** via the route shown in Scheme 23 [44].

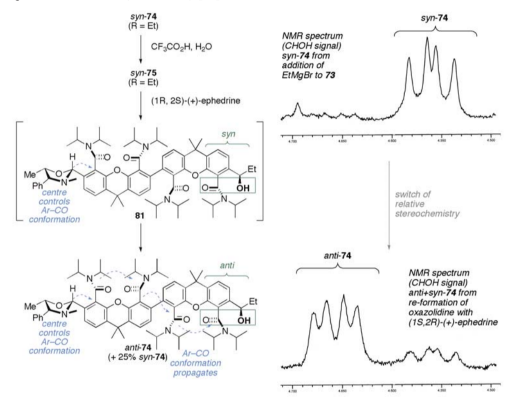
Scheme 23. Synthesis of a bis-xanthene.

From **70**, a further series of lithiations, functionalizations and coupling gave two aldehydes, **73** and **78**, in which the electrophilic centre is separated from a terminal (-)-ephedrine-derived oxazolidine by a chain of four or six amide groups respectively. The X-ray crystal structure of **78** shows all six amides arranged with their dipoles successively opposed.



Scheme 24. Stereochemical relay in a bis- and tris-xanthene.

Nucleophilic addition of EtMgBr and PhMgBr to each aldehyde generated a single stereo-isomer of the product **74** or **79**, as judged by NMR and by HPLC (Scheme 24), in reactions displaying relayed unprecedented 1,16- and 1,23-remote stereocontrol. Stereochemical homogeneity at the newly formed centre was confirmed by hydrolysis of the oxazolidines to yield aldehydes **75** and **80**. Comparison of the HPLC traces of these aldehydes with those of authentic mixtures of stereoisomers showed some degree of thermal epimerization (from the deprotection step), but allowed the degree of stereocontrol at the new centre to be quantified as > 95:5 (Scheme 24) [44].



Scheme 25. Switching relative stereochemistry from a distance.

Alternative explanations for the stereocontrol achieved in **74** and **79**, such as association between the oxazolidine ring of one molecule with the Grignard reagent as it adds to another, were ruled out by showing for example that diastereoselectivity is unchanged in reactions of racemic oxazolidines. Conclusive evidence that stereochemical information may be relayed through the amides was obtained simply by re-condensing **74** (R = Et) with ephedrine enantiomeric with that first used to form the oxazolidine (Scheme 25). Initially, presumably, **81** is formed, but under the conditions of the oxazolidine formation, the amides of **81** relax into their preferred orientation. Necessarily, the relative stereochemistry

between the last amide in the sequence and the stereogenic centre adjacent to it becomes inverted, and this local inversion of relative stereochemistry is clearly evident in the chemical shifts of the CHOH signals of the two samples of 74 (Scheme 25).

Conclusion

The chemistry in Schemes 24 and 25 shows conformational information being relayed through a molecule in a manner which parallels the allosteric conformational changes in enzymes and receptors which relay information in biological molecules. So far the "output" of the conformational change has been a stereoselective reaction, detectable chromatographically or spectroscopically. Other inputs and outputs can be envisaged, though: for example, Krauss and Koert [45] have developed molecules which use conformational information to mediate signal transduction between a metal binding site and a fluorophore (Scheme 26). Tricyclic 82 adopts the preferred conformation shown as 82a, but on treatment with zinc a switch occurs to the ring-flipped 82b. Evidence for the ring flip is provided by NMR or by a change in the fluorescence spectrum as the pyrene fluorophores move apart [46].

The possibility that the synthetic methodology of remote stereocontrol might be applied to a challenge in nanotechnology – signal transduction and processing – is intriguing, and it is clear that the future in this area is very bright. There are huge opportunities for the combination of various types of conformational control elements to lead not just to systems capable of relaying information but of processing and storing that information as well.

Scheme 26. Conformational relay of binding information to a fluorophore.

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