1
Introduction: A Survey of How and Why to Separate Enantiomers

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This book is about the separation of enantiomers by synthetic methods, which is to say methods involving some chemical transformation as part of the separation process. We do not in this book cover chromatographic methods for the separation of enantiomers [1]. Nor do we focus on methods based on crystallizations as these have been amply reviewed elsewhere (see below). We are concerned mainly therefore with resolutions that involve a synthetic component, so mostly with the various flavours of kinetic resolutions through to more modern methods such as divergent reactions of a racemic mixture (DRRM). This introduction briefly clarifies the scope of the book.

The reasons such methods are of continued importance are threefold:

1) Society: the need for enantiopure compounds. New molecules as single enantiomers are important to our continued well-being because they are the feedstocks of new medicines, agrochemicals, fragrances and other features of modern society in a chiral world. Of the 205 new molecular entities approved as drugs between 2001 and 2010, 63% were single enantiomers [2]. Nature provides an abundance of enantiopure compounds, but we seek, and need, to exceed this by obtaining useful unnatural molecules as single enantiomers, and we may reasonably want to access both enantiomers of some compounds.

2) Academia: the basic science involved in the behaviour of chiral compounds. If we seek the state of the art in our discipline, we cannot help but think that rapid and selective chemical distinction between enantiomers, which results in their facile separation, is something beautiful in itself. There have been many successful methods developed for the synthetic separation of enantiomers, as we shall see, and these are both de facto interesting and instructive to consider for the design of future examples of such processes. The relationship between kinetic resolution and asymmetric catalysis is strong, and one can inform the design of the other. It is hoped that the diverse examples described in this book stimulate thoughts in the reader of what is possible next.

3) Industry: the need for new methods. There remain many classes of compounds that still cannot be resolved, or where efficiencies are too low for widespread

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adoption. It is still the case that classical resolution techniques are overwhelm-
ingly used over other more complex methods. Of the 128 drug candidate
molecules assessed in a recent industry survey, half were being developed as
single enantiomers, and the sources of the stereocentres were mainly the chiral
pool (55%) with resolution (28%) and asymmetric synthesis (10%) responsible
for fewer examples [3]. This is, predictably, a feature of economics as much
as science and one must not be too quick to judge new fields like asymmetric
catalysis versus older ones like classical resolution. Pasteur added something
enantiopure to a racemate in 1853 [4], whereas the catalytic prowess of a metal
centre surrounded by chiral ligands was first demonstrated only in 1968 [5].
In addition, many chiral acids and bases have proven to be useful in classical
resolutions, while Nature does not seem to be so generous in its supply of
molecules that can effect catalytic, asymmetric transformations. The great
progress made in synthetic chemistry has not (yet) brought us to the posi-
tion that allows us to make any enantiopure substance in quantity given that
resources are always limited. That leaves us with the synthesis of a racemate
from which we pick one enantiomer out. Such a process can be remarkably
efficient and cost-effective, if such tools are available, but the great successes
described in this book should not hide the fact that we require better separation
methods with wide applicability if we are to avoid an overreliance on just using
whatever Nature provides.

The various methods considered in this book may be classified as follows.

1.1
Classical Methods

A racemate can be resolved with ease if it happens that the enantiomers form
separate crystals – a so-called conglomerate. It becomes possible to separate,
physically, the enantiomorphous crystals – a process sometimes referred to as *triage*.
This is what Pasteur famously achieved in 1848 with a sample of ammonium
sodium tartrate [6]. Such good fortune is quite rare and is in any case not a
’synthetic method’ in the strictest sense (nor is it practical on a large scale).
Other physical processes (alone, without any attendant synthetic process) such as
evaporation or sublimation can be used to increase the enantiomeric excess of
organic compounds, including amino acids [7].

Another important but non-synthetic method involves harvesting crops of enan-
tioenriched crystals by seeding a supersaturated racemate and is known as *resolution
by entrainment*, or *preferential crystallization*. These approaches have been well
reviewed and will not be covered here [8]. Rare (but spectacular) examples where
crystallizations of conglomerates are observed combined with racemization events
in solution, leading to *total spontaneous resolutions* [9], are briefly mentioned in
Chapter 7 as there have been interesting recent developments on that front.
Included therein are special cases where diastereomeric interactions in solution
combined with racemization may yield more than a 50% yield of one enantiomer,
1.2 Kinetic Resolution (‘KR’)

A conceptually simple and well-known method for the separation of enantiomers is the kinetic resolution. A racemate is subjected to some reaction using a chiral agent and one of the enantiomers of the racemate reacts more quickly than the other (Scheme 1.2). The ‘selectivity factor’, $s$, is an expression of the difference in the rates of the two reactions.

The enantiomer ratio is shown using a formalism borrowed from Ed Vedejs in Chapter 6. As can be seen, the composition varies dynamically as the reaction
proceeds. At the outset, the system displays its best synthetic selectivity between the enantiomers because an equal amount of each enantiomer is present. As the reaction proceeds, statistics begins to interfere as the fast-reacting enantiomer is depleted. Success is a curse – as the fast-reacting enantiomer concentration reduces it becomes difficult to stop the slow-reacting enantiomer from participating – a ‘mass action’ problem. This explains the two best-known features of kinetic resolutions – that the maximum yield for the process is necessarily 50%, but that high enantiomeric excesses are found at the extremes of conversion – that is, that the enantioenrichment of the product (whatever that may be) is largest at the start of the reaction while the enantiomeric excess of the starting material left behind is largest just before the end of the reaction. In a kinetic resolution, one cannot usually have one's cake (high enantiomeric excess) and eat it (high yield).

A simple approach to kinetic resolution is to use a stoichiometric reagent. In the example shown in Scheme 1.3, an enantioenriched alcohol is obtained from a racemate through the addition of an enantiopure-acylating agent [13]. If 1 equivalent of the reagent were used, a 1 : 1 mixture of diastereomers would result with no kinetic differentiation. Stoichiometric kinetic resolutions thus necessarily employ a sub-stoichiometric amount of the reagent – in the example shown only 0.1 equivalents is used. This method for the separation of enantiomers is described in detail by Maddani, Fiaud and Kagan in Chapter 2.
The obvious alternative is to use a chiral catalyst to effect some reaction on the racemate. Frequently, the reaction does not introduce a new stereocentre, and a typical example is shown in Scheme 1.4. This process, developed by Birman, involves an acylation of an alcohol effected by a catalytic quantity of an enantiopure nucleophilic catalyst [14]. Catalytic methods are covered by Hélène Pellissier in Chapter 3.

\[
\begin{align*}
\text{OH} & \quad \text{Ph} \\
\text{rac} & \quad \text{CO}_2 \text{H} \\
\text{MeO} & \quad \text{O} \\
\text{O} & \quad \text{OMe} \\
(0.9 \text{ equiv}) & \quad (0.75 \text{ equiv}) \\
i-\text{Pr}_2\text{NET} (1.8 \text{ equiv}), \text{CH}_2\text{Cl}_2, \text{rt}, 12 \text{ h} \\
\end{align*}
\]

**Scheme 1.4** Example of a catalytic kinetic resolution of a racemic alcohol.

The catalyst can be an enzyme, and some of the most impressive examples have been of this type. In the example shown in Scheme 1.5, a lipase acts on a kilogram of the racemic starting material to generate enantioenriched products that may be separated by distillation after filtration of the enzyme from the reaction mixture [15]. The use of enzymes in kinetic resolutions is covered by Humphrey, Ghanem and Turner in Chapter 4.

\[
\begin{align*}
\text{OH} & \quad \text{NMe}_2 \\
\text{rac} & \quad \text{O} \\
\text{Candida antarctica lipase} & \quad \text{O} \\
(36\%) & \quad (45\%, >95\% \text{ ee}) \\
\end{align*}
\]

**Scheme 1.5** Example of catalytic kinetic resolution effected by an enzyme.

1.3 Dynamic Kinetic Resolution (‘DKR’)

Kinetic and classical resolutions both suffer from the weakness that the maximum yield of a product is 50%. This limit is removed if the enantiomers of the starting material can be interconverted (Scheme 1.6). The challenge is significant – to run a kinetic resolution in the forward direction and a simultaneous racemization of the
starting materials that leaves any compound derived from the starting materials unaffected. The racemization needs to be rapid with respect to the asymmetric transformation, to prevent build-up of the slow-reacting enantiomer.

\[
\begin{align*}
\text{Product from } R & \quad \text{Product from } S
\end{align*}
\]

**Scheme 1.6** Generic dynamic kinetic resolution showing the ratio of enantiomers with conversion.

This seemingly insurmountable task has been solved, with several impressive systems having been demonstrated to date, although the number of cases is far lower than those for regular kinetic resolutions. In the example shown in Scheme 1.7, the kinetic resolution of secondary alcohols is achieved with a lipase while the racemization is effected by a ruthenium complex [16]. This example illustrates well the striking effectiveness of two orthogonal chemical processes that might a priori be expected to interfere with each other. Dynamic kinetic resolutions (DKRs) are reviewed by Nakano and Masato Kitamura in Chapter 5. The chapter also includes a description of rarer processes that are frequently confused with DKR but which are mechanistically distinct [17], such as the dynamic kinetic asymmetric transformation (DYKAT); the similarity is that DYKAT and DKR achieve complete conversion of a racemate to one enantioenriched product but the DYKAT product is typically not convertible back to starting material (needed for a separation of

\[
\begin{align*}
\text{Candida antarctica lipase} \\
\begin{array}{c}
\text{Ph} \\
\text{Ph} \\
\text{Ph} \\
\text{Ph}
\end{array} & \quad \begin{array}{c}
\text{Ph} \\
\text{Ph} \\
\text{Ph} \\
\text{Ph}
\end{array} \\
\text{OC} & \quad \text{OC}
\end{align*}
\]

**Scheme 1.7** Example of dynamic kinetic resolution.
1.4 Divergent Reactions of a Racemic Mixture (‘DRRM’)

The essential weakness of a simple kinetic resolution is the build-up of the slow-reacting enantiomer of the starting material, a process that may be solved by DKR. There is another solution – to have the slow-reacting enantiomer be consumed by another reaction, to give a second product that may be separated from the first. In other words, to have the enantiomers give distinct (enantioenriched but not enantiomeric) products at similar rates (Scheme 1.8) – known as a divergent reaction of a racemic mixture. As for the DKR process, the starting material ideally remains as a racemate throughout, in which case the maximum yield of any given product is 50%. Strictly speaking for the process to be called a resolution, there must be a means of separating the products and converting them back to the enantiopure starting materials. The term DRRM encompasses processes that involve the addition of one or two chiral reagents to a racemate. The concept is well illustrated by the latter, involving addition of two chiral reagents to a racemate as shown; this subclass of DRRM is frequently referred to as parallel kinetic resolution (PKR).

The specific example shown (Scheme 1.9) involves two pseudo-enantiomeric reagents [18]. One of the reagents reacts rapidly with the (R)-enantiomer of the starting alcohol while the other reacts preferentially with the (S)-enantiomer. Although a kinetic resolution of the racemate would be possible with either reagent alone, the mass action problem would mean that the reagent’s selectivity would need to be very high to match the yield and enantiomeric excess values obtained from the DRRM. This process was discovered by Ed Vedejs, who reviews the field in Chapter 6.
Scheme 1.9 Example of a divergent reaction of a racemic mixture.

1.5 Other Methods

At the end of this book, I summarize a small number of ‘neglected’ cases not covered elsewhere, such as the use of circularly polarized light, polymerizations, ‘ripening’ processes, dynamic combinatorial chemistry and even several thermodynamic processes. In some of these cases, the potential for future significance in the synthetic separation of enantiomers is clear but the fields are not yet mature because the methods are young, exhibit recurrent issues, or have been developed for other reasons – in the case of polymerizations the eye of the polymer chemist is typically on the polymer, not on any enantiomeric excess that might be contained in the residual solution. This chapter is a little unusual – it appears that these disparate methods have never previously been collected together; the separate fields typically do not cite each other despite their ultimately sharing a common theme.

One striking example mentioned in this final chapter requires us to bend the term racemate to include ‘very near racemates’ that contain a very small enantiomeric excess. Enrichment of such samples by direct crystallization-based methods would typically only be attempted by committed optimists. In such a situation, we could synthesize more of the excess enantiomer preferentially if we had an appropriately asymmetric autocatalytic reaction – our initial excess enantiomer could replicate at the expense of the other. Preparatively, this is the effective separation of the enantiomers we used at the outset. Such a system has its physical realization in the Soai autocatalysis in which a very small enantiomeric excess of a pyrimidyl alcohol is amplified over several cycles to give an almost enantiopure sample of the alcohol (Scheme 1.10) [19].
Nearly racemic

\[
\text{Zn}(i-	ext{Pr})_2
\]

High enantiomeric excess
(after several cycles)

Scheme 1.10 The Soai reaction that achieves a form of synthetic separation of enantiomers through selective autocatalysis.

These and other processes are not by any means widespread or generalizable, and they have not yet had an impact on the industrial preparation of enantiopure compounds, but are included as suggestions of where the vibrant and important field of the synthetic separation of enantiomers might go next. Some of these methods also remind us of the ‘prototypical’ synthetic separation of enantiomers that may have played a role in the origin of life.

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References

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