Prediction of the efficiency of diastereoisomer separation on the basis of the behaviour of enantiomeric mixtures

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The driving force of homo- and heterochiral complex formation in mixtures of chiral compounds is, probably, the effort of the system to separate the most symmetric associates from the less symmetric ones. A possible way to achieve separation of these associates is distribution between two phases. Therefore, during the separation of (a certain part) diastereoisomers similar trends can be observed as in the course of the distribution of enantiomeric mixtures between two phases, although in the first case a third chiral compound (namely the resolving agent) is present. Of course in this case the pursuit of symmetry is not so obvious as in case of enantiomeric mixtures. It should be noted that the outcome may thus be modified by the intervention of kinetic control. One can conclude that the structure of chiral compounds encodes the result of the (optical) resolution.

Introduction

In many cases, living organisms contain only one of the two enantiomers of chiral molecules, but often racemic compounds (1 : 1 mixture of the two enantiomers) are obtained in the chemical syntheses. The biological activity of enantiomers may be different or even opposite, so enantiomeric separations are necessary and inevitable. Many of the methods described in the literature for the separation of enantiomers involve the formation of diastereoisomers followed by liberation of the separated enantiomers. These enantiomeric separation methods are discussed and systematized in several articles.1-10

In the course of the resolution processes, racemic compounds are reacted with another chiral reagent (resolving agent). The diastereoisomers so obtained are separated, and their decomposition affords the corresponding enantiomeric mixtures. Usually, pure enantiomers can only be obtained by further purification of these enantiomeric mixtures (Scheme 1).

The composition of the salt crystallized during the fractionated precipitation of the diastereoisomeric salts and enantiomeric mixtures is determined by the SDE11 capacity (self disproportion of enantiomers) of the involved chiral compounds.

The most commonly used methods for the separation of enantiomeric mixtures are based on the exploitation of the distribution of hetero- and homochiral associates between two phases (most often between solid and liquid or vapour phases).12 According to the most recent research enantiomeric enrichment can occur using achiral chromatography13-16 and by separation of racemates using a chiral selector on an achiral17,18 stationary phase or chiral phase chromatography (HPLC),19 respectively.

A correlation between the eutectic composition of binary melting point diagrams of diastereoisomeric mixtures (eeDiast) and efficiency of resolution (F) was established by us (Scheme 2).20

Furthermore, it was found by analyzing the results of 45 resolutions, that the average enantiomeric purity (eeRes = 78%) of enantiomeric mixtures isolated from crystalline diastereoisomers correlates to the average value of the measured eutectic composition of the starting racemic compounds (eeDiast = 73%). At the same time, when the eutectic composition of the resolving agent is higher than the eutectic composition of the racemic compound (in 29 cases), a better correlation was observed between the average value of enantiomeric purities (eeDiast = 80%) of enantiomeric mixtures isolated from the crystalline diastereoisomers and the average value of eutectic compositions of enantiomeric mixtures of the resolving agent (eeRes = 78%) (Table 1).21

Based on these observations, we suppose that the composition of a crystalline diastereoisomer is determined either by the eutectic composition of the racemic compound or that of the resolving agent and the higher ee value has a more dominant effect (Scheme 3).

Consequently, a correlation can be found between the binary melting point/composition phase diagram of the diastereoisomeric mixtures and the phase diagrams of the enantiomers which are the constituents of the diastereoisomers.
If we wish to separate the enantiomers of a racemic mixture using a structurally similar resolving agent (equivalent or half equivalent amount), the isomers of the racemic compound are transformed into “quasi enantiomeric mixtures”. In the course of separation of these “quasi enantiomeric mixtures” (of diastereoisomers) the same methods based on the distribution between two phases can be used which are suitable methods for the separation of enantiomeric mixtures (also diastereoisomeric related supramolecular enantiomeric associates).\(^7\)

### Separation of diastereoisomers from the melt

The mixtures of racemic compounds and resolving agents can be considered as mixtures of diastereoisomeric supramolecular structures. These diastereomeric supramolecular structures exist in solutions and in melts, therefore these diastereoisomeric associates can be separated by crystallization from the melt. In this case the mixture of the racemic compound and the resolving agent is melted, then the crystalline phase – obtained by controlled cooling – can be separated by filtration (if it is possible).

<table>
<thead>
<tr>
<th>Number of resolutions</th>
<th>(ee_{eeu \text{ Rac}}) (average) %</th>
<th>(ee_{eeu \text{ Res}}) (average) %</th>
<th>(ee_{e\text{Dia}}) (average) %</th>
<th>(F) (average)</th>
</tr>
</thead>
<tbody>
<tr>
<td>45</td>
<td>73</td>
<td>78</td>
<td>78</td>
<td>0.57</td>
</tr>
<tr>
<td>29</td>
<td>78</td>
<td>80</td>
<td>0.62</td>
<td></td>
</tr>
</tbody>
</table>
An example of such a separation is the crystallization of a diastereoisomeric mixture of MEN–DBTA molecular complex from a melt incorporating menthol (MEN) and (R,R)-dibenzoyl-tartaric acid ((R,R)-DBTA) (Scheme 4). It can be seen from the above example that the behavior of mixed chiral compounds is valid not only for diastereoismeric salts but for diastereoisomers in general. During resolution of racemic menthol with (R,R)-DBTA, the parts of the diastereoisomeric complexes are kept together by weak second order interactions, only. Only the more stable molecular complex that crystallizes more quickly makes the separation possible.

Separation of diastereoisomers by sublimation of enantiomeric mixtures

Enantiomeric separations can be effectuated even if the mixture of diastereoisomers is obtained in a solid–solid reaction. An example is the reaction of solid 2-iodo-trans-cyclohexanol (ICH) and solid (R,R)-DBTA for three months (Scheme 5). It was observed in the course of the fractionated vacuum sublimation23 of the above mentioned mixture of compounds, that one of the enantiomers of ICH sublimated as the first fraction at a relatively low temperature (T1, Scheme 5), and the other ICH enantiomer sublimated at a higher temperature (T2), after thermal decomposition of the molecular complex formed previously in the solid–solid reaction.

Separation of diastereoisomers by distillation of enantiomeric mixtures

In the case of resolutions using half an equivalent of resolving agent it is expected, that the enantiomeric proportion of racemic compound may be separated from the corresponding diastereoisomer formed (distributed between two phases). In the reaction of methylana (2-methylamino-1-phenylpropane, MA) and (R,R)-DBTA, after the precipitation of the diastereoisomeric salt, the residual free amine, namely (S)-MA could be obtained by distillation under vacuum,24 while the other enantiomer was obtained by the separation of the solid diastereoisomer residue (Scheme 6).

Separation of diastereoisomers by fractionated distillation of enantiomeric mixtures

This method is also suitable for fractionated separations if the resolving agent forms a salt that can decompose without any damage. An adequate example for this is the resolution of racemic anara (2-amino-1-phenylpropane, AN) by half an equivalent of the structurally related (S)-N-phthaloyl-α-phenyl-ethylamine (PPEA) (Scheme 7). Again, the free, optically active base ((R)-AN) could be distilled off at T1, then the solid diastereoisomeric salt could be decomposed at a higher temperature (T2), by ring closure of the phthaloyl derivative, so the other amine enantiomer ((S)-AN) could be distilled off in the second stage.

Separation of diastereoisomers by extraction of enantiomeric mixtures with a supercritical fluid (carbon dioxide)

In the course of a half equivalent resolution, the remaining free enantiomer may also be removed by extraction from the
reaction mixture after the crystallization of the diastereoisomeric salt. This extraction can be accomplished using a supercritical fluid, most often supercritical carbon dioxide. In case of resolution of trans-cyclohexane-1,2-diol (trans-CHD) by (R,R)-tartaric acid ([(R,R)-TA]) the free enantiomeric portion was separated by extraction with supercritical CO2 from the mixture of the excess of trans-CHD and the crystalline diastereoisomeric complex (Scheme 8). Of course, the other enantiomer can be recovered from the diastereoisomeric complex.

Separation of diastereoisomeric molecular complexes by fractionated crystallization

The above demonstrated methods can be applied for the separation of enantiomers having an asymmetric center on a phosphorous atom. For example, the resolution of several racemic alkyl-, alkoxy-, and aryl-substituted 3-methyl-3-phospholene oxides were accomplished via molecular complex formation with (R,R)-TADDOL (α,α′,α′′,α′′′-tetraphenyl-1,3-dioxolan-4,5-dimethanol) as the resolving agent (Scheme 9). If half an equivalent of (R,R)-TADDOL was used, the more stable diastereoisomer crystallized which could be isolated by conventional methods, such as filtration.

Separation of diastereoisomeric coordination complexes by fractionated crystallization

In other cases, the cyclic P-chiral 3-phospholene oxides were separated into their enantiomeric mixtures by resolution with the Ca2+ or Mg2+ salts of DBTA (DBTC). As is shown in Scheme 10, when half an equivalent of resolving agent was used, the favourable diastereoisomer was precipitated. After filtration and decomposition of the diastereoisomeric complex, an enantiomeric mixture of the 3-phospholene oxide (MPO) was obtained. The antipode of the first isolated MPO enantiomer was recovered from the mother liquors of the resolutions.

Separation of diastereoisomers from enantiomeric mixtures using mixtures of two immiscible solvents

Based on the above examples, formation of solid phase, solid–liquid phases or solid–gas phases systems is always necessary for the separation of diastereoisomers. Now the question is: could one achieve chiral separation between two immiscible liquid phases? The answer yes, of course. Diastereoisomers may be separated by distribution between two liquid phases (Scheme 11).
If the racemic methylanara (MA) is reacted with half an equivalent of sodium salt of \((R,R)\)-tartaric acid (\((R,R)\)-TAN) in a mixture of water and benzene, the enantiomer and diastereoisomer are distributed between the two liquid phases.\(^\text{29}\)

The aqueous phase contains the neutral salt of TAN-MA, while the other enantiomer can be found in the organic layer. So the separation of diastereoisomers or enantiomers can be accomplished without crystallization, using two immiscible solvents. The two solvent phases can also provide particularly good separation if the diastereoisomer can crystallize due to its insolubility in the applied two solvents.

![Scheme 8](image)

**Scheme 8** Separation of the trans-CHD enantiomers via molecular complex formation followed with supercritical fluid extraction.

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**Separation of diastereoisomers by crystallization involving the formation of solvates**

It is very common during the separation of diastereoisomers, that the enantiomeric purity and yield of separation is increased when solvates were formed during the crystallization of the diastereoisomers. For example, if trans-chrysanthemic acid (CRS) is resolved with \(N,N\)-dimethyl-aminodiol (obtained by the transformation of an intermediate of chloramphenicol) in methanol, a methanol solvate of the diastereoisomer could be isolated in very good diastereoisomeric purity, but the yield was 52%, only (Scheme 13). Whereas the resolution was accomplished in diisopropyl ether, or methyl isobutyl ether in the presence of methanol, the methanol solvate of the diastereoisomer was also crystallized, but both the purity and the yield increased significantly.\(^\text{31}\)

**Separation of diastereoisomers by crystallization in the presence of a structurally related achiral reagent**

Frequently, the resolution can be accomplished only via solvation. In the previous example the resolution could be carried out using another solvent. When the racemic \(\alpha\)-phenylethylamine (PEA) was resolved using half an equivalent of a derivative of a PEA enantiomer, a diastereoisomer was isolated which contained \((S)\)-PEA in 60% excess in acetone (Scheme 14).\(^\text{32}\) When an achiral compound (such as urea) with a related structure to one

![Scheme 11](image)

**Scheme 11** Resolution of MA using two immiscible solvents.

![Scheme 9](image)

**Scheme 9** Separation of diastereoisomeric molecular complexes of MPO by fractionated crystallization.

![Scheme 10](image)

**Scheme 10** Resolution of MPO via diastereoisomeric coordination complex formation and fractionated crystallization.
part of the resolving agent was added to the solution before crystallization, much purer diastereoisomer was isolated with an (S)-PEA enantiomeric excess of 90%.

So the achiral solvate forming reagent – which is structurally related to either the resolving agent or the racemic compound – promoted the crystallization of the diastereomer, and increased enantiomeric excess of the diastereomer.

**Crystallization of diastereoisomers based on kinetic and thermodynamic control**

The decisive role of kinetic and thermodynamic control was observed in the separation of enantiomeric mixtures. This phenomenon can also be found for both the separation of quasi-enantiomeric mixtures and conventional resolutions. For example, the effect of kinetic control was observed in the resolution of pregabalin (PRE) by mandelic acid (S)-MA (Scheme 15). When the crystalline diastereoisomeric salt was isolated after 15 minutes crystallization, the enantiomeric excess of (S)-PRE isolated from the salt was 98%, while the ee decreased significantly if the crystallization was carried out over 48 hours.

This means that the thermodynamic control has a disadvantageous effect on this process. The same phenomenon was observed in the course of the reciprocal process, when racemic MA was resolved by (S)-PRE. Namely, crystallization of the diastereoisomeric salt was controlled kinetically.

In other cases it is necessary to wait until thermodynamic equilibration, because in these cases the process is controlled thermodynamically. An adequate example is the resolution of an intermediate of tamsulosin (TAM) with (R,R)-DBTA (Scheme 16). In this case the diastereoisomeric salt contained the (R)-TAM enantiomer in excess. However, after an hour of crystallization, practically racemic TAM was found in the salt, but excellent enantiomeric excess could be achieved when the diastereoisomer was crystallized for 48 hours.

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**Scheme 12** The separation of diastereoisomers by crystallization from immiscible solvents.

**Scheme 13** Separation of diastereoisomers by crystallization involving the formation of solvates.

**Scheme 14** Separation of diastereoisomers by crystallization involving the formation of solvates.

**Scheme 15** The effect of kinetic control on the separation of diastereoisomers.
compounds (racemate and resolving agent) con-
average eutectic compositions of the involved chiral
enantiomeric mixtures from series of resolutions with the
similar. Comparison of the average ee values of the obtained
diastereoisomer forming chiral compounds are not structurally
tioration of the formed (crystalline) diastereoisomers even if the
racemate and/or the resolving agent determines the composi-
tion of the eutectic composition of the diastereoisomeric salt.

Table 2 The average ee data of the examined resolutions

<table>
<thead>
<tr>
<th>No. of experiments</th>
<th>Average value of ee_{EuRac}</th>
<th>Average value of ee_{EuRes}</th>
<th>Average value of ee_{Dia}</th>
<th>Average value of F</th>
</tr>
</thead>
<tbody>
<tr>
<td>13 (10^3)</td>
<td>80%</td>
<td>78%</td>
<td>0.62</td>
<td></td>
</tr>
</tbody>
</table>

* The used compounds are not structurally related. ee_{EuRac}: ee of an enantiomer at the eutectic composition of the racemate; ee_{EuRes}: ee of a resolving agent enantiomer (R or S) at the eutectic composition; ee_{Dia}: ee of an enantiomer of the original racemate at the eutectic composition of the diastereoisomeric salt.

Conclusion

The above mentioned examples demonstrate that the properties of the involved enantiomeric mixtures determine the enan-
tiomer (diastereoisomer) distribution between two phases and in
this way the behaviour of the chiral compounds used deter-
mines the efficiency of the process during enantiomer or dia-
tereoisomer separations. There are numerous methods of
choice (solid–solid, solid–liquid, solid–gas, liquid–liquid
distributions) and, of course, we should choose the most
favourable method (if there is more than one possibility).

We have recognized that in the resolution processes the
diastereoisomers behave similarly to their constituent enan-
tiomeric mixtures if the resolving agent was structurally related
to the racemic compound.

We demonstrated that the eutectic composition of the
racemate and/or the resolving agent determines the composition of the formed (crystalline) diastereoisomers even if the
diastereoisomer forming chiral compounds are not structurally
similar. Comparison of the average ee values of the obtained
enantiomeric mixtures from series of resolutions with the
average eutectic compositions of the involved chiral
compounds (racemate and resolving agent) confirmed the
above observation, namely the higher eutectic composition
governs the enantiomer separation (Table 2).

We also think that the eutectic composition of the dia-
astereoisomer forming enantiomers determines the efficiency of the
resolutions in cases of crystalline diastereoisomeric salt,
molecular- or coordination complex formations and these
governing effects are valid when the separation is based on the
distribution between two liquid phases.

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