Basics of Retrosynthesis Analysis

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Abstract: Retrosynthesis is a crucial strategy in organic chemistry, allowing chemists to deconstruct complex molecules into simpler starting materials by working backward from a target structure. This paper introduces fundamental concepts of retrosynthesis, including various bond-breaking techniques such as heterolysis and homolysis, and the use of synthetic equivalents. By examining single-functional and two-functional-group disconnections, we explore methods for identifying efficient synthesis pathways. Additional retrosynthetic strategies, like functional group interconversion and electrocyclic disconnection, are discussed to address unique challenges in synthesis planning. Through guiding principles and real-world examples, this paper demonstrates how retrosynthesis facilitates efficient and creative molecule construction, ultimately enhancing our ability to design complex chemical syntheses.

Keywords: Chemistry, Organic chemistry, Synthesis, Retrosynthesis

1. Background and Introduction

Before getting into the topic of retrosynthesis, we have to have a clear understanding of: what exactly is retrosynthesis? Retrosynthesis, in essence, is a method used in chemistry that involves deconstructing a target molecule to determine the starting materials needed for its synthesis. To illustrate, consider the task of constructing a triangle. One might suggest using sticks and glue, or perhaps assembling smaller triangles to form a larger one of equivalent area. There are countless methods to achieve the same outcome. Similarly, in retrosynthesis, chemists approach the problem by working backward from the desired molecule, exploring various pathways to construct it from available starting materials. This backward-thinking approach not only fosters creativity but also aids in the efficient synthesis of complex molecules.

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Figure 1&2: Methods of how to construct a triangle

2. Methods of Retrosynthesis

Let us now turn to one of the most critical aspects of retrosynthesis: the methodologies employed in this process. Drawing on the earlier example of constructing a triangle, we approached the task by breaking it down into smaller components. In a similar manner, a fundamental strategy in retrosynthesis is the selective breaking of chemical bonds. The goal is to identify and cleave the appropriate bond to simplify the target molecule into more manageable fragments. Consider, for instance, the mortise and tenon structure in woodworking. The most effective way to separate the two connected pieces is to cut at the joint; attempting to do so elsewhere would prove significantly more challenging. Thus, the careful selection of which bond to break is crucial in retrosynthesis. The question then arises: how do we determine the correct bond to break? The following methods provide some guidance in this regard.



Figure 3: Breaking a mortise and tenon structure from the joint

2.1. Bond-breaking

2.1.1. Classification: Heterolysis and homolysis

To begin, bond-breaking strategies can be categorized based on the transfer of electrons, namely into heterolysis and homolysis. Heterolysis involves the complete transfer of an electron pair, resulting in the formation of ions, whereas homolysis entails the equal splitting of an electron pair, leading to the creation of two radicals. In the context of retrosynthesis, heterolysis is the more prevalent method, as it produces synthons—molecular fragments that are either anionic or cationic. However, because these charged synthons cannot exist independently in their ionic forms, they must be paired with neutral counterparts in actual total synthesis. This is where the concept of synthetic equivalence comes into play. Identifying the appropriate synthetic equivalents for each type of synthon is a fundamental aspect of organic synthesis. Below are some examples that illustrate this concept.

Target Molecule	Synthons	Synthetic Equivalent
R-R'	$R^{\Theta} + R'$	RMgX, RLi, R ₂ Cd R'X
RCH ₂ CH ₂ OH	R^{Θ} + CH_2CH_2OH	RMgX, RLi, R ₂ Cd
RCOOH	R^{Θ} + COOH	RMgX, RLi CO ₂
RCOR'	R [⊖] + [⊕] COR'	RMgX, RLi, R ₂ Cd R'COCl, R'COOH
RC:CR'	$R^{\Theta} + CECR'$	RMgX, RLi HC=CR', R'X

Figure 4: Examples of synthons [1]

2.1.2. One-functional-group disconnection for heterolysis

To further elaborate on heterolysis, it can primarily be divided into two types: one-functional-group disconnection and two-functional-group disconnection (which are also known as one-group-disconnection, and two-group-disconnection for abbreviation). In the case of one functional-group disconnection, it is helpful to consider the resonance forms of the molecule. Referring to Figure 6, we observe the ketone and its resonance form, particularly the one on the left, where the oxygen carries a negative charge and the carbon carries a positive charge. In this configuration, the oxygen can be identified as a hydroxyl functional group (with an additional hydrogen). Thus, when breaking bonds, through a process known as Functional Group Interconversion (FGI), which will be discussed later, the ketone can be transformed into a hydroxyl group with a positively charged connected carbon, consistent with the resonance form of the ketone.



Figure 5: Example of one group disconnection

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Figure 6: Resonance form of ketone [2]

2.1.3. Two-functional-group disconnection for heterolysis

For two functional-group disconnection, we can analyze it from the perspective of the consonant pattern. In this scenario, the two functional groups involved are hydroxyl and ketone (Figure 7). Given that oxygen is an electronegative atom, it tends to attract the shared electron pair towards itself, creating a partially positive charge on the carbon to which it is connected. Consequently, this partially positive carbon draws electrons from the adjacent carbon, imparting a partially negative charge to it. This results in a positive-negative-positive pattern, referred to as a consonant pattern, which facilitates the bond-breaking process. It is important to note that bonds corresponding to a dissonant pattern are more difficult to break than those in a consonant pattern.



Figure 7: Example of two-group disconnection [3]

2.2. Other strategies besides bond-breaking

However, bond-breaking alone is not sufficient to address all challenges in retrosynthesis, prompting the introduction of additional strategies.

2.2.1. Fine-tuning

One such strategy is fine-tuning, which involves making small adjustments during the retrosynthesis process. While fine-tuning alters the problem at hand, it does not necessarily simplify it. Within this approach, we have Functional Group Interconversion (FGI), which transforms one functional group into another; Functional Group Addition (FGA), which introduces a new functional group; and Functional Group Removal (FGR), which eliminates a functional group, although it is crucial to understand how this group will be reintroduced during total synthesis.

2.2.2. Electrocyclic disconnection

Electrocyclic disconnection is another method, which involves the reverse process of cycloaddition by reorganizing and breaking bonds through the movement of electrons within a ring. An example of this is the reverse process of Diels-Alder reaction [4].





Figure 8: Reverse process of Diels-Alder reaction

2.2.3. Illogical disconnection

Illogical disconnection is a method that paradoxically simplifies the molecule by forming new bonds.

2.2.4. Rearrangement

Rearrangement strategies are employed to further refine the retrosynthetic pathway by rearranging the molecule with certain conditions.

3. Guiding Principles

3.1. Focus on Simplification

The first guiding principle is that we should always focus on simplification of the molecule. Since methods such as fine-tuning can only shift the problem instead of simplifying them, we should focus on simplification by reducing reliance on fine-tuning and taking advantage of symmetry to streamline the process.

3.2. Most Reactive Functional Group Needs to be Cut Off the First

The second guiding principle is that the most reactive functional group should be cut off first, as these groups are typically the last to be introduced in a forward synthesis. To use a house-building analogy, if we consider the carbon-carbon skeleton as the framework of the house, then the bonds to heteroatoms can be likened to the furniture. So frame first, funiture last. And in backwards process, which is retrosynthesis, furniture first, then framework.

3.3. Two Group Disconnection is Usually Better than One Group Disconnection

Last but not the least, the third guiding principle is that two-functional-group disconnections are generally preferred over one functional-group disconnections. Due to the presence of consonant patterns, it is often easier to break two functional groups apart.

4. Real world example

With reference to a real-life example, we can analyze how we can do retrosynthesis with the methods mentioned above.



Figure 9: Real-life total synthesis process [5]

Firstly, we can start our retrosynthesis by analyzing which molecule is the target molecule, which is the starting material, and what are the similarities between them to therefore think of a process to retrosynthesize it.



Figure 10: Starting material (left) and target molecule (right)

The figure shown above are the starting material (left) and target molecule (right). The two parts circled are the similarity between the two molecules since them have similar ring structures and consist similar elements.

So our target is to remain this structure in the target molecule, not cutting it, and try to convert it to the ring structure in the starting material.

Since this functional group, MeO, does not appear in the starting material, we have to cut this off. With reference to guiding principle 2, most reactive cut off first. Since MeO is not that reactive for us to directly cut off, we use FGI to convert this ether into hydroxyl and then into CO2, a much more reactive functional group, which is a lot easier to cut off.



Figure 11: Methodology of how to convert MeO into CO2

After removing the functional group, CO₂, by FGR, via FGA, we introduce a bromine (third molecule) for the construction of a double bond in the next step (second molecule) and an OTBS (third molecule) to change into hydroxyl functional group (second molecule). So next, we use electrocyclic and bond breaking to from this double bond (second molecule) and break the ring structure in the third molecule. Finally, by breaking the bond in the second molecule and with a FGI, we got the starting material.



Figure 12: Methodology after removing MeO into the starting material

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