Synthesis of Tetrasubstituted Alkenes via Metathesis

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Abstract: Fully substituted olefin generation via metathesis is presented. Catalyst development, optimization of reaction conditions and substrate screening are included. In addition, asymmetric alkene metathesis, the cross metathesis reaction for this transformation and its application in natural products will be discussed.

Keywords: tetrasubstituted alkene; metathesis; catalyst

1. Introduction

As an alternative technology for olefin generation, the metathesis reaction has been focused on since its recent and rapid development [1,2]. This transformation allows the efficient production of otherwise readily unavailable olefins including medium-sized heterocycles, carbocycles, trisubstituted olefins and heteroatom-substituted olefins (Scheme 1) using structurally diverse catalysts (Figure 1) [3,4]. Nowadays, the protocol is widely utilized to synthesize valuable polymers, natural products and medicinal compounds for humans [5–7].

Scheme 1. Metathesis reaction and its basic transformation.
Although metathesis is a fascinating technology for alkene synthesis, this reaction still needs improvement in terms of reactivity and selectivity [1]. The general pattern that the more substituted a substrate is, the less reactive it is, shows that steric bulkiness hampers the incorporation of metal carbenoids in the substrate. For the selectivity issue of alkene geometry, various methodologies have been developed to control the $E$ or $Z$ geometry of the resulting olefins [8–12]. However, the general problems of $E$- or $Z$-selective alkene metathesis haven’t yet been solved when it comes to the case of tetrasubstituted olefins (Figure 2).

For the reasons mentioned above, a great effort has already been devoted to the efficient generation of tetrasubstituted alkenes via metathesis. A variety of catalysts, reaction conditions including solvents, substrate screening and applications have been tried. This article presents an overview of these endeavors.

2. Results and Discussion

2.1. Early Studies with a Molybdenum Catalyst

Actually, tetrasubstituted alkene generation via metathesis was first reported two decades ago, when the rapid ring closing metathesis (RCM) of the substrate to form the sterically encumbered medium-sized ring skeletons $1–4$ using a highly reactive molybdenum catalyst (A) was described (Scheme 2) [13–15]. These results indicate that the molybdenum carbenoid can be incorporated into
the sterically hindered substrates easily. However, this molybdenum metal species was soon replaced by the ruthenium catalysts B–D, because of their air and moisture stability and wide utility. Consequently, tetrasubstituted alkene generation via ruthenium catalysts was also studied.

Scheme 2. Early utilization of a Mo catalyst A in tetrasubstituted olefins.

2.2. Early Studies with a Ruthenium Catalyst

The most challenging obstacle to the utilization of ruthenium carbenoids to produce tetrasubstituted alkenes was the reactivity of the ruthenium species. Unlike the molybdenum species A which catalyzed the transformation of the diene 3’ into the corresponding product 3 in high yield, the ruthenium catalyst E couldn’t perform the same role (Scheme 2 and Equation 3 in Scheme 3) [16].

Scheme 3. Unsuccessful RCM and successful relay RCM.

Equation 3

Equation 4
Considering that the relay-RCM of dienyne 5' to the corresponding diene product 5 was reported using the same catalyst E, the incorporation of the ruthenium carbenoid into the substrate was regarded as a rate determining step in this process (Scheme 3). Definitely, more reactive ruthenium catalysts were necessary.

2.3. New Catalyst Design and Its Application

As is well known, dramatic improvement of ruthenium catalyst reactivity was achieved by replacing the tricyclohexylphosphine (PCy₃) ligand with an N-heterocyclic carbine (NHC) ligand [17]. Actually, it was reported that a similar ruthenium catalyst F with an imidazolylidene ligand could produce tetrastubstituted cycloalkenes or pyrroline products (Entry 1, Scheme 4) [18]. Furthermore, the same imidazolylidene with a more sterically demanding dimesityl group was introduced to catalyze the medium-sized RCM of tetrastubstituted alkenes (Entry 2, Scheme 4) [19]. The Grubbs-Hoveyda catalyst D also showed a high performance in producing furans 10 and 11 (Entry 3, Scheme 4) [20].

Scheme 4. Development of new catalysts and their application.

| entry | catalyst | conditions | products (yield) | references |
|-------|----------|------------|-----------------|------------|
| 1     | F        | 5 mol%, CH₂Cl₂ 40 °C, 13-24h | 6 Ts 80%, 3 EtO₂C₂CO₂Et 64% | 18 |
|       |          |            | 4 99%, 7 83%    |            |
| 2     | G        | 5-14 mol%, CH₂Cl₂ 40 °C, 60-72h | 8 Boc 76%, 9 BnO₂P 81% | 19 |
| 3     | D        | CH₂Cl₂ rt 24h then 40 °C 20h for 10 | 10 38%, 11 66% for 11 | 20 |

2.4. Optimization of Reaction Conditions

In addition to catalyst development, other reaction conditions such as solvents and substrate were optimized (Scheme 5). In 2001, Furstner et al. utilized supercritical CO₂ to improve the RCM
procedure. This unique condition showed improved results for the fully substituted alkenes 6 and 3 [21]. Another research carried out studied the substrate. Using the established Grubbs 2nd generation catalyst, Haufe et al. achieved efficient RCM of a fluoride-substituted alkene 12. Due to the biological importance of fluoride-containing molecules, this paper was highlighted and later applied to the synthesis of other natural products (Scheme 10, *vide infra*) [22]. A polymer-supported metathesis catalyst H was also utilized for this type of conversion. An azepine structure 13 could be constructed using the polymer-supported catalyst H, which could be reused [23]. More dramatic advances were achieved when Blecher et al. changed solvents from normal benzene to hexafluorobenzene. This solvent change allowed the highly efficient RCM of medium-sized heterocycles in high yield [24].

After optimizing reaction conditions, a large improvement in the tetrasubstituted olefin metathesis reaction was achieved. However, this type of RCM still needed generality for various substrates.

**Scheme 5.** Optimization of the metathesis reaction in tetrasubstituted alkenes.

| entry | catalyst | conditions | products (yield) | references |
|-------|----------|------------|----------------|------------|
| 1     | ![Catalyst G](image) | 5 mol%, 40 °C, supercritical CO₂, 24-120h | Ts 6 83%, EtO₂C 3 51% | 21 |
| 2     | ![Catalyst C](image) | 2 mol%, toluene, 80 °C, 8h | ![Product 12](image) 45% | 22 |
| 3     | ![Catalyst H](image) | 5 mol%, PhCF₃, 50 °C | ![Product 13](image) 93% | 23 |
| 4     | ![Catalyst I](image) | 1-3 mol%, C₆F₆, 80 °C, 1h | ![Products 14](image) 99%, ![Products 15](image) 99%, ![Products 16](image) 99%, ![Products 17](image) 83%, ![Products 18](image) 99% | 24 |

2.5. Further Catalyst Development

More advanced catalyst development research began with the Grubbs-Hoveyda catalyst D. Grela et al. reported that the NO₂-substituted aromatic ligand activates the catalyst J in the tetrasubstituted olefin
metathesis reaction (Entry 1, Scheme 6) [25]. It seems that the electronic effect facilitates the insertion of olefin in the ruthenium carbenoid. However, a more outstanding advance was achieved with the sterically less-demanding liganded catalyst K.

Scheme 6. Development of more advanced catalysts J–M.

| entry | catalyst | conditions | products (yield) | references |
|-------|----------|------------|----------------|------------|
| 1     | J        | 5 mol% CH₂Cl₂, 40 °C | 3 (0-14%), 6 (22-82%), 19 (20-29%) | 25 |
| 2     | K        | 5 mol% C₆D₆, 60 °C | 6 (>95%), 14 (>95%), 3 (>95%), 4 (>95%), 15 (88%), 18 (87%) | 26 |
| 3     | K        | 5000 ppm MTBE, 50 °C, 8h | 19 (90%), 20 (97%) | 27 |
| 5     | L        | 0.2-0.5 mol% toluene, 80 °C, 20h | 3 (56-84%), 21 (91-99%), 15 (53-95%), 18 (89-98%), 13 (63-85%) | 29 |
| 6     | M        | 5 mol% toluene, 100 °C, 20h | 3 (29%) | 30 |

*MTBE = methyl tert-butyl ether.
Replacement of the mesityl substituent of the toluene moiety made the catalyst \( K \) much more reactive than the original catalyst \( D \) [26]. Actually, the catalyst \( K \) was so reactive that it was sufficient to produce azacycles 19 and 20 at low catalyst loading (Entry 3) [27]. Due to this high performance, the catalyst \( K \) was used by Stoltz et al. to synthesize elatol, a bioactive natural product [28].

Research on catalyst optimization is still ongoing. In 2010, Plenio et al. reported the bis-NHC liganded catalyst \( L \). In particular, this electron-deficient ligand helped the catalyst to produce various tetrasubstituted olefins by RCM (Entry 5, Scheme 6) [29]. A 6-membered ring ligand was also reported recently. Substitution of imidazoline on the pyridine ligand also showed potential RCM in fully substituted olefins 3 (Entry 6) [30]. Although its chemical yield is relatively low, it is an impressive result as it provides possibilities for ligand modification.

### 2.6. Asymmetric RCM in Tetrasubstituted Alkenes

As seen in Scheme 7, an experiment on asymmetric RCM was also investigated. Collins et al. tried to utilize the metathesis reaction for the introduction of tetrasubstituted asymmetric alkenes. After a set of ligand screens, desymmetrization of the triene 23′ to the optically active dihydropyran 23 could be carried out successfully. It was also reported that the optically active dihydrofuran skeleton could be produced through the same desymmetrization reaction [31].

**Scheme 7.** Asymmetric RCM of 23′ to 23.

![Scheme 7](image)

2.7. Cross Metathesis in Tetrasubstituted Alkenes

As research on RCM in tetrasubstituted alkenes proceeds, experiments on utilizing this reaction for cross metathesis were also attempted. Howell et al. reported cross metathesis of the terminal alkene 24′ with \( \beta \)-lactam 25 to obtain the tetrasubstituted unsaturated \( \beta \)-lactam 24 in high yield using the Grubbs-Hoveyda catalyst \( D \) (Scheme 8). The reaction was quite rapid when utilized for the generation of biologically important structures with various substituents, hydroxyl, aliphatic, silyl and aromatic groups, although the olefin geometry of the product was unselective. Further results showed that the electron-rich \( \beta \)-lactam didn’t produce the same conversion under the same conditions, indicating that the electronic factors are important for this conversion [32].
Scheme 8. Cross metathesis in tetrasubstituted alkenes 24.

Due to little success of metathesis in tetrasubstituted alkenes, its application has been also limited. However, some results show that this conversion can be applied to the synthesis of interesting molecules when it is utilized appropriately. Actually, Yoshida et al. showed an example. A dimethylene compound 26' could be converted into 26'' via the alkene metathesis reaction under the catalysis of Grubbs 2nd catalyst. Once 26'' formed, it was aromatized into the phenol 26 by spontaneous tautomerization in situ (Scheme 9) [33]. This reaction could be carried out in both CH₂Cl₂ and benzene, although the later showed better yield despite the slow reaction rate.

Scheme 9. RCM in tetrasubstituted alkenes and its aromatization.
Another application of this RCM reaction in the total synthesis of elatol, a bioactive natural product, was achieved by Stoltz et al. [28]. A fused bicycle 27 was produced from bis-terminal methylene 27' through catalysis of the less sterically demanding ruthenium carbenoid K, described above (Scheme 10). The chloroalkene 27 could be obtained at a 97% yield and 1.3 g scale, because this procedure was so practical. It is interesting that 27 could be produced using the 2nd generation Grubbs catalyst B at 85% after 24 h. This intermediate 27 could be transformed into (+)-elatol via simple functional group interconversion.

**Scheme 10.** RCM in tetrasubstituted alkenes and its application to natural product synthesis.

These two examples show that RCM in tetrasubstituent alkenes is a highly efficient technology for the synthesis of complex molecules. More research to expand its generality is still ongoing.

### 3. Conclusions

Since the development of molybdenum and ruthenium catalysts, the metathesis reaction has served as a key carbon-carbon bond transformation. It has also allowed us to obtain otherwise unavailable tetrasubstituted alkene skeletons in an effective manner, although its substrate scope and low chemical yield hampers its wide utilization. Nowadays, however, challenges to overcome these limitations and expand its efficiency into the enantioselective reaction and the synthesis of natural products are actively ongoing. It is envisioned that this effort will result in outstanding advances in the synthesis of tetrasubstituted alkenes.

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