Review

Synthesis of Imidazole-Based Medicinal Molecules Utilizing the van Leusen Imidazole Synthesis

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Received: 10 February 2020; Accepted: 1 March 2020; Published: 3 March 2020

Abstract: Imidazole and its derivatives are one of the most vital and universal heterocycles in medicinal chemistry. Owing to their special structural features, these compounds exhibit a widespread spectrum of significant pharmacological or biological activities, and are widely researched and applied by pharmaceutical companies for drug discovery. The van Leusen reaction based on tosylmethylisocyanides (TosMICs) is one of the most appropriate strategies to synthetize imidazole-based medicinal molecules, which has been increasingly developed on account of its advantages. In this review, we summarize the recent developments of the chemical synthesis and bioactivity of imidazole-containing medicinal small molecules, utilizing the van Leusen imidazole synthesis from 1977.

Keywords: van Leusen; TosMICs; imidazole; synthesis

1. Introduction

Imidazole ring, which is widely found in natural products and medical molecules, is one of the most prominent, five-membered, nitrogen-containing, heterocyclic scaffolds. Furthermore, imidazole-based heterocyclic compounds, which possess a vital position in medicinal chemistry, have been playing a central role in the treatment of numerous types of diseases, and new derivatives for medicinal use are being energetically developed worldwide [1–5]. Due to the peculiar structural characteristic of imidazole scaffold with a worthy electron-rich feature, it is advantageous for imidazole groups to combine with various receptors and enzymes in biological systems, through diverse weak interactions, thereby showing a variety of biological activities. At present, a legion of imidazole-containing compounds with high a medical potential as a clinical drug have been widely used to treat diverse types of illnesses, such as antibacterial [6,7], antifungal [8,9], anti-inflammatory [10,11], antiviral [12,13], anti-parasitic [14,15], anticancer [16,17], antihistaminic [18,19], and enzyme inhibition [20,21]. Imidazole and its derivatives encompass a vast range of medical activities, as shown in the following Table 1.
the van Leusen imidazole synthesis based on TosMICs, which is the cycloaddition reaction, is one of
decades, there have been numerous classical strategies for synthesizing this ring compound in the
cycloaddition reaction in a proton solvent, meanwhile, the effects of R1 and R2 on the formation of
in organic synthesis by the Dutch professor van Leusen in 1972, this reagent is also known as van
2. General van Leusen Imidazole Synthesis
due to its excellent advantages like simple manipulation, easily obtained raw materials and a wide
paid attention to by pharmaceutical chemists and organic synthesis researchers. However, there is

Table 1. Various pharmacological activities and chemical structures of imidazole-based molecules.

| Pharmacological Activities | Chemical Structures |
|----------------------------|---------------------|
| Antibacterial              | ![Chemical Structure 1](image1) ![Chemical Structure 2](image2) |
| Antifungal                 | ![Chemical Structure 3](image3) ![Chemical Structure 4](image4) |
| Anti-inflammatory          | ![Chemical Structure 5](image5) ![Chemical Structure 6](image6) |
| Antiviral                  | ![Chemical Structure 7](image7) ![Chemical Structure 8](image8) |
| Antiparasitic              | ![Chemical Structure 9](image9) ![Chemical Structure 10](image10) |
| Anticancer                 | ![Chemical Structure 11](image11) ![Chemical Structure 12](image12) |
| Antihistaminic             | ![Chemical Structure 13](image13) ![Chemical Structure 14](image14) |
| Enzyme inhibitor           | ![Chemical Structure 15](image15) ![Chemical Structure 16](image16) |
Owing to the significant pharmacological or biological activities and the enormous medicinal value of imidazole-based molecules, the synthesis of the imidazole-skeleton small molecule has been paid attention to by pharmaceutical chemists and organic synthesis researchers. However, there is still a need for a simple and efficient way to construct the imidazole heterocyclic skeleton. In recent decades, there have been numerous classical strategies for synthesizing this ring compound in the laboratory, including van Leusen imidazole synthesis [22], Debus-Radziszewski imidazole synthesis [23], Wallach imidazole synthesis [24], etc. Among these synthetic strategies, it is well-known that the van Leusen imidazole synthesis based on TosMICs, which is the cycloaddition reaction, is one of the most convenient and attractive protocols for the preparation of imidazole-based small molecules, due to its excellent advantages like simple manipulation, easily obtained raw materials and a wide range of substrates, which has been developed rapidly in the past decades (Scheme 1).

\[
\text{R}^1\text{CH}=\text{NR}^2 + \text{Tos}^+\text{NC} \xrightarrow{\text{base}} \text{R}^1\text{N}^-\text{R}^2(\text{H})^\text{+}
\]

Scheme 1. General van Leusen imidazole synthesis.

TosMIC, one of the most significant reactants, has many good features at room temperature, including being a stable solid and being odorless and colorless. Since it was introduced and applied in organic synthesis by the Dutch professor van Leusen in 1972, this reagent is also known as van Leusen’s reagent. Up to now, TosMIC and its derivatives have been recognized as one of the most significant building blocks in nitrogen heterocyclic synthesis, which have been fruitfully employed especially in the preparation of imidazole-based heterocycles [25–29].

Therefore, this review will summarize the developments of the synthesis of imidazole-based molecules, utilizing the van Leusen imidazole synthesis, based on TosMICs from 1977. It is expected that this review article will be beneficial for new opportunities to search for a reasonable design for less toxic and higher bioactive imidazole-containing drugs.

2. General van Leusen Imidazole Synthesis

In 1977, van Leusen et al. first published that TosMIC and aldime undergo a base-induced cycloaddition reaction in a proton solvent, meanwhile, the effects of R1 and R2 on the formation of 17 were qualitatively analyzed. They discovered that α-tosylbenzyl isocyanate and α-tosylethyl isocyanate 18 could form 1,4,5-trisubstituted imidazoles 19 (Scheme 2). Based on the various advantages of this reaction, it is widely named as the van Leusen imidazole synthesis [22].

\[
\text{R}^1\text{CH}=\text{NR}^2 + \text{Tos}^+\text{NC} \xrightarrow{\text{base \ ToH}} \text{R}^1\text{N}^-\text{R}^2(\text{H})^\text{+}
\]

Scheme 2. The first example of van Leusen imidazole synthesis.

As shown in Scheme 3, the van Leusen imidazole synthesis allows the preparation of imidazole through [3 + 2] cycloaddition reaction from aldimes and a reaction with TosMICs, which contain reactive isocyanide carbons, active methylene, and leaving groups such as C2N1 “3-atom synthon”. The cyano moiety can be a gradual cycloaddition to polarize a double bond under a base condition. The elimination of p-TosOH forms the intermediate 4-tosyl-2-imidazoline to produce the target 1,4,5-trisubstituted imidazoles 19, accompanied by the elimination of p-TosOH, which is negative to the obtained 1,5-disubstituted imidazoles.
3. Developments of the van Leusen Imidazole Synthesis

In 1995, Frankowski and co-workers were reported to lead the known D-xylo-pentodialdose to form imidazo-L-xylo-piperidino derivatives, based on a sequential eight-step reaction. In this process, the imidazole-base molecule was obtained through a van Leusen reaction as a key step. As shown in Scheme 4, applying this van Leusen methodology, 20 and TosMIC 18 were transformed into the imidazole derivative 21. At last, removal of the protecting group gave to the target product 22, which is a kind of bicyclic azasugar and a glycosidase inhibitor [30].

In 1998, the Sisko group synthesized the 1,4,5-trisubstituted imidazole 25, which displayed potent binding with p38 MAP kinase, a recently discovered protein kinase that participates in an inflammation regulatory mechanism. As shown in Scheme 5, 1,4,5-trisubstituted imidazole 24 was accessed by a novel and facile protocol, based on the reaction of an α-ketoaldimine 23 with an aryl-substituted TosMIC reagents 18 as the key step [31].

In 2000, they also described an active and gentle procedure for preparing multisubstituted imidazoles in one-pot, from an aryl-substituted TosMIC and a generated imine, in situ. One of the reactions is shown in Scheme 6, cycloaddition of imine 28, prepared imine in situ from a 40% aqueous solution of pyruvaldehyde 26 and amine 27, then, ketone 29 was prepared in DMF, using aryl-substituted TosMIC 18 and K₂CO₃, with a 75% yield. Surprisingly, when the reaction was carried out at 0 °C, the product 29 was blended with 15–20% of the 1,4-disubstituted imidazole 30 [32].
In the same year, the Vanelle group converted 6-nitropiperonal into the methylimin 31 through a condensation reaction with methylamine in ethanol. The molecule with imidazole ring 32 was formed by reacting equimolar quantities of TosMIC 18 and aldime 31 with K$_2$CO$_3$ in methanol solvent, under a refluxing condition. Then, through the classical Knoevenagel reaction, compound 33 were obtained, which displayed an effective in vitro leishmanicidal bioactivity (Scheme 7). Compound 33a, higher in vitro leishmanicidal bioactivity, might lead to a promising significant therapeutic agent (Scheme 8) [33].

From 2005 to 2006, in order to access unusual imidazole-based heterocyclic structures, a series of experimental results were reported by the Gracias and Djuric group.

Initially, they reported a new protocol employing the van Leusen three-component reaction and the ring-closing metathetical reaction in a sequence method, to form a fused bicyclic imidazole ring. The general method is shown in Scheme 9 (top), the reaction underwent smoothly with the condensation reaction of 4-pentenal 34 and allylamine 35, in DMF at room temperature, to give the imine in situ, which was followed by the addition of the TosMIC reagent 18 and a base, to access the van Leusen imidazole product 36 in high yield. Next, a ring-closing metathesis (RCM) reaction was performed through the second-generation Grubbs catalyst to get imidazole 37 and their derivatives (Scheme 9, top) [34].

Next, they developed a concise route of fused imidazo azepine analogs via the stepwise van Leusen/intramolecular enyne metathesis synthesis. The condensation reaction of 4-pentenal 34 with but-2-yn-1-amine 38 in DMF at room temperature gives the imine in situ, and then adds phenyl TosMIC 18 and K$_2$CO$_3$ to access the van Leusen imidazole product 39 in high yield. Subsequent cyclization via the intramolecular enyne metathesis reaction results in the formation of the cyclized product 40 containing the diene functional group and their derivatives (Scheme 9, bottom) [35].
Firstly, they synthesized fused imidazo-pyridine and imidazo-azepine derivatives by using a sequential van Leusen/intramolecular Heck route. The imidazole 47 and 49 were accessed via the condensation reaction between an appropriate aldehyde-containing vinylogous bromide 45 and an amine-containing double bond 46, respectively. Then, preformation of the imine, the desired TosMIC reagent 18 and K2CO3 were added, and the cyclization was allowed to proceed at room temperature to get imidazo-[1,5-a]pyridine 48 or imidazo[1,5-a]azepine 50 (Scheme 11) [37].
Next, they described an elegant method to obtain a fused imidazole ring, employing the van Leusen three-component reaction, followed by Pd/Cu catalyzed intramolecular C-arylation. Meanwhile, another available protocol to form 5,6-dihydroimidazol[2,1-a]-isoquinoline system 53, via a radical cyclization reaction of tethered aryl halides onto the imidazole ring, was also published. In their article, the imidazole precursors were accessed through an alkylation chemistry, and the van Leusen reaction was used to assemble imidazole followed by a transition metal catalyzed C-C bond framework (Scheme 12) [38].

![Scheme 12. Synthesis of fused imidazole rings 53.](image)

In 2006, Dömling’s group reported that substituting pyrroloimidazoles 56 were assembled by van Leusen multicomponent reaction (MCR) of TosMIC 18, indole carbaldehydes 54, and primary amines 55. They were interested in bioactivity of the products and these imidazole-containing skeletons were screened in a phenotypic assay for neurite outgrowth. The test results indicated that these small molecules would serve as useful chemical probes to research neurite growth and might be used as a therapeutic application for axon regeneration of lesions of the human spinal cord and brain (Scheme 13) [39].

![Scheme 13. Synthesis of chemical probes 56.](image)

Based on their previous research, they also reported a novel synthesis route of potential aspartyl-protease inhibitors in 2007. In their method, 1,4,5-trisubstituted 1-(4-piperidyl)-imidazoles 59 could be obtained through an isocyanide-based van Leusen MCR of \( \alpha \)-substituted TosMICs 18, aldehydes 57, and 4-aminopiperidine 58 (Scheme 14) [40].

![Scheme 14. Synthesis of potential aspartyl-protease inhibitors 59.](image)

In 2009, a range of 1,5-disubstituted-4-methylimidazole preparations were reported by Fodili and co-workers. Compounds 63 were prepared from acetimine analogs 62, which were obtained from dehydroacetic acid 60 and primary amine 61. These were enabled to react with TosMIC 18 in the presence of catalytic amounts of bismuth triflate. A plausible mechanism was put forward that
involved the first formation of an imidazoline intermediate, followed by methyl migration, and then subsequent aromatization provided access to the target imidazole (Scheme 15) [41].

![Scheme 15. Synthesis of 1,5-disubstituted-4-methyl imidazoles 63.](image)

In 2012, Hulme and Moliner reported a novel synthesis of two pharmacological relevant classes of molecules forming the imidazoquinoxaline scaffold 66 and 68, respectively. This method involves the use of 1,2-phenylenediamines 64 and glyoxylic acid derivatives, namely ethyl glyoxylate 65 or benzylglyoxamide 67, along with TosMIC 18 in a microwave-assisted van Leusen three-component condition. Then there was a deprotection-cyclization step to form two biological-activity-enticing imidazoquinoxaline families (Scheme 16) [42].

![Scheme 16. Synthesis of biologically enticing imidazoquinoxaline 66 and 68.](image)

In 2016, the Pirali’s group reported a method for synthesizing 4-PI analogues via the van Leusen MCR, which is by far the most direct method for obtaining functionally rich imidazole. This transformation can be used to synthesize four different series of compounds—1,4,5-trisubstituted and 1,5-, 1,4- and 4,5-disubstituted imidazoles 71. Meanwhile, 4,5-diaryl imidazoles were analyzed by a 3D quantitative structure-activity relationship (SAR). Based on their docking score and synthetic feasibility, the compounds were selected, synthesized, and biologically evaluated. Compared with 4-PI, the IDO1 inhibitor products through this experimental method have enhanced the potency. Both in enzymatic and cellular assays, the most active compounds showed lower micromolar potency, but no detectable cellular toxicity. The analysis displayed that a putative hydrogen bond between the Ser167 of the protein and the NH of the imidazole ring might be responsible for the improvement of the potency, together with favorable interactions with the partially occupied pocket B. The results above suggested that the 4,5-disubstituted imidazole framework might give a new orientation for the design of IDO1 inhibitors, in future (Scheme 17) [43].

![Scheme 17. Synthesis of potential IDO1 inhibitors 71.](image)
At the same year, the Suresh group developed a direct, sequential, copper-catalyzed N-arylation-condensation reaction utilizing chiral cyclic 1,2-diamines 73 and ortho-haloaryl aldehydes or ketones 72. The corresponding chiral tricyclic 1,4-benzodiazepines 74 were synthesized in high yield. Subsequently, 1,4-diazapine 75 was converted into a novel tetracyclic N-fused ImBDs through van Lusen imidazole synthesis (Scheme 18) [44].

A range of 5-aryl-1-alkylimidazole derivantes 79 were synthesized by utilizing the van Leusen MCR by Bojarski and co-workers, in 2017. The van Leusen reaction was performed by the stepwise cycloaddition of TosMIC 18 to the polarization double bond of a preformed imine 78. Imidazole is synthetized by the elimination of p-TosOH, from the cyclic intermediate. Then, aromatic aldehydes bind to the appropriate amine. All compounds were highly selective to 5-HT5A. These compounds were metabolically stable in human liver microsome, showed less toxicity in HEK-293 and HepG2 cells and were water-soluble (Scheme 19) [45].

In 2019, they also synthesized a range of fluorinated indole-imidazoles to find more effective 5-HT7 receptor selective agonists as molecular probes. As shown in Scheme 20, the target compounds 82 were obtained from the relevant indoles undergoing the formylation process and the stepwise van Leusen imidazole synthesis with TosMIC 18. Meanwhile, they indicated that compound 82a might be a potential analgesic or a long-sought tool for studying the receptor effect of 5-HT7, based on the anti-nociceptive function on a murine model of neuropathic pain (Scheme 21) [46].
In 2017, Sharma and co-workers developed a brief, novel, and simple operational van Leusen method. They presented the first van Leusen process towards synthesis of high functionalized dihydrodibenzo[b,f]imidazo[1,2-d][1,4]thia/oxazepine 86, by reacting TosMIC 18 and dibenzo[b,f][1,4]thia/oxazepines 85, which were obtained from compound 83 and 84, under a basic condition (Scheme 22) [47].

Scheme 22. Synthesis of highly functionalized molecules 86.

In 2018, the Weaver group reported that two corresponding series of molecules based on the SAR design—N1-substituted 5-indoleimidazoles and N1-substituted 5-phenylimidazoles were accessed. The latter (and more efficient) series were obtained by an accidental rearrangement of imines that are intermediate in the van Leusen imidazole synthesis reaction. The one-pot synthesis of compounds 89 in two steps utilizing the van Leusen imidazole synthesis process is shown in Scheme 23. First, imine formation between 2-indolecarboxaldehyde 87 and the appropriate benzyl amine 88 took place, which then reacted with TosMIC 18 and K2CO3 to give indoleimidazoles. They have also conducted SAR studies on their novel and more potent compounds. Compounds 92 were synthesized starting from known amine 91 and aromatic aldehyde 90 (Scheme 24). These inhibitors represent a promising future for the development of IDO1 inhibitors with specific physical and chemical properties that are easy to synthesize [48].

Scheme 23. One-pot synthesis of potential IDO1 inhibitors 89.

Scheme 24. One-pot synthesis of potential IDO1 inhibitors 92.

In the same year, Kondaparla et al. synthetized a range of short chain 4-aminoquinoline-imidazole derivatives 95 using two steps, in one-pot, via the van Leusen MCR standard method. The detailed synthetic protocol for the synthesis of target compounds 95 is shown in Scheme 25, compound 94 as amine was utilized to react with TosMIC 18 and acetaldehyde 93 through a multicomponent cyclisation reaction. In the end, all synthesized compounds were screened against the chloroquine sensitive and chloroquine-resistant strains of Plasmodium falciparum. In this study, substitution on the N-(2-(1H-imidazol-1-yl)ethyl)-7-chloroquinolin-4-amine moiety had greatly influenced the antiplasmodial bioactivity [49].
In the same year, Laali and co-workers reported that an available route provided access to various C5-substituted imidazoles 97, 98, and 99, through one-pot tandem via the van Leusen-Suzuki, van Leusen-Heck, and van Leusen-Sonogashira methods, respectively. Imidazolium-ILs as solvents, along with piperidine-appended imidazolium [PAIM][NTf2] as task-specific basic IL and facile aldimines 96, and TosMIC 18 in the mild condition, were employed in their article (Scheme 26) [50].

In the late 2018, Guan and co-workers developed a novel and highly efficient synthesis of polysubstituted 1H-imidazo-[4,5-c] quinoline derivatives through the stepwise van Leusen/Staudinger/aza-Wittig/carbodiimide-mediated cyclization method. Azides 102 were accessed by the van Leusen reaction of 2-azidobenzaldehyde 100, amine 101, and TosMIC 18. Then, 1H-imidazole-[4,5-c] quinoline 103 was formed by a tandem aza-Wittig reaction with isocyanate, in a moderate to good yield (Scheme 27) [51].

In 2019, Lammi et al. reported an imidazolyl peptidomimetic, which has shown proprotein convertase subtilisin/Kexin 9 (PCSK9) inhibitor bioactivity in the micromolar dosage range. As shown in Scheme 28, the first imidazole derivative 105 was accessed in good yield, starting from \( p \)-anisaldehyde 104, methylvamine 77, and TosMIC 18. Then, compound 105 was treated with \( n \)-butyllithium and...
DMF as a formylating reagent at low temperatures, to obtain the aldehyde derivative 106 in high yield. The target compounds 107a RIm13 and 107b RIm14 were accessed via a quasi-iterative protocol, with the emphasis on the alternating van Leusen three-component reactions with two formylation steps. In the end, they found that RIm13 represents currently one of the most potent proprotein convertase subtilisin/kexin 9/low-density lipoprotein receptor, (PCSK9/LDLR) protein–protein interaction of small molecules (Scheme 28) [52].

![Diagram](image-url)

**Scheme 28.** Synthesis of inhibitory activity molecules 107a and 107b.

In 2020, Rashamuse and co-workers described a microwave-assisted cycloaddition of TosMIC 18 with imines and aldehydes to form 1-substituted 5-aryl-1H-imidazoles. Imidazoles 110 were also obtained in a one-pot, two-steps reaction with a yield comparable to that obtained through step-by-step irradiation of aldehyde 108 and aliphatic amine 109, with a neat microwave at 60 °C for 4 min, followed by the addition of TosMIC 18, K2CO3, and CH3CN, and the reaction mixture was placed under microwave conditions (Scheme 29) [53].

![Diagram](image-url)

**Scheme 29.** Synthesis of a novel class of potential inhibitors of microbial pathogen 110.

Moreover, the antibacterial properties of these fragments in vitro were tested by the minimum inhibitory concentration (MIC) bioassay. The results showed that the MIC value of compound 110a against *Staphylococcus aureus* was 15.6 μg/mL, while 110b displayed a similar MIC value against *Bacillus cereus*, and indicated that these compounds might be further developed to specifically target microbial pathogens (Scheme 30).
In 2014, the Bunev group reported a novel process for the synthesis of 1,4,5-trisubstituted imidazole-containing trifluoromethyl group 112, which contained two-component condensation reaction, N-aryltrifluoroacetimidoyl chlorides 111 reacted with TosMIC 18, as well as sodium hydride in dry THF at room temperature, under argon atmosphere (Scheme 31, top) [54].

![Scheme 31. Synthesis of imidazoles 112 and 114 containing the trifluoromethyl group.](image)

Then, in 2019, they also previously described a procedure, in which 1-imidoylbenzotriazoles [N-aryl-1-(1H-benzotriazol-1-yl)-2,2,2-trifluoroethan-1-imines] 113 reacted with TosMIC 18, according to the van Leusen reaction to obtain a good yield of 1-aryl-4-(4-methylbenzenesulfonyl)-5-(trifluoromethyl)-1H-imidazoles 114, which is difficult to access. The yield of 114 almost did not depend on the substituent in the N-aryl fragment of initial imidoylbenzotriazole 113 (Scheme 31, bottom) [55].

A possible mechanism for synthesis of the imidazole-containing trifluoromethyl group is shown in Scheme 32. Initially, deprotonation of TosMIC with sodium hydride forms stabilized carbanion 18, which attacks the carbon–nitrogen bond’s carbon atom of 115, to give the intermediate adduct 116. Elimination of the R ion from the latter generates intermediate 117, which undergoes intramolecular cyclization and leads to imidazole 118.

![Scheme 32. A possible mechanism for the synthesis of imidazole-containing trifluoromethyl group.](image)

4. Other van Leusen Imidazole Synthesis

In 2015, Fodili and collaborators described the synthesis of a 1,4-disubstituted 5-methylimidazole 121. As shown in Scheme 33, compound 121 was prepared by reacting enamine 119 with TosMIC 18, under the presence of tert-butylamine and a catalytic amount of bismuth (III) triflate in methanol. In this research, it was the first example of a usual rearrangement in the van Leusen imidazole synthesis and showed that the imidazole ring system can be prepared through reaction with TosMIC and a tautomeric enamine, to form a secondary ketamine. The possible mechanism involves the formation of the van Leusen imidazoline intermediate, followed by a C-C bond cleavage and then subsequent tosylation substitution [56].
In 2019, the Suresh group demonstrated the formation of imidazoles in the presence of water as a solvent and a base-free condition. The reaction of dihydro $\beta$-carboline imines $122$ and $p$-toluenesulfonylmethyl isocyanides $18$ formed the corresponding substituted $N$-fused imidazo 6,11-dihydro $\beta$-carboline derivatives $123$, with good yields under mild and green condition (Scheme 34) [57].

A possible mechanism for the present metal- and base-free imidazole framework is shown in Scheme 35. Initially, the precursor dihydro $\beta$-carboline imine $122$ acts as a base that captures proton from TosMIC $18$, to provide a C-nucleophile, which would add to the dihydro $\beta$-carboline imine $122$, and then be cyclized to form the intermediate $125$ through the intermediate $124$. Another molecule of the starting dihydro $\beta$-carboline imine $122$ captures a proton from intermediate $125$, then removes the tosyl group, which might result in the construction of imidazole derivative $123$. Subsequently, the product imidazole $123$ might also act as a base following the rational reaction mechanism, as described in Scheme 35.
At the same year, Necardo et al. found an unusual multicomponent synthesis of 4-tosyl-1-arylimidazoles 127, by considering aryl azides as the electrophilic partners for the TosMIC 18-mediated van Leusen cycloaddition. In this transformation, it is the first example of the reaction of two TosMIC molecules participating in van Leusen imidazole synthesis (Scheme 36) [58].

![Scheme 36. Synthesis of 127 by van Leusen imidazole synthesis with two TosMIC.](image)

A plausible scenario for the MCR is shown in Scheme 37. In the initiation step, the TosMIC anion attacks N-3 of the azide 128 to produce intermediate 129. Then, N-1 intercepts the isocyanide in a 6-endo-trig cyclization to form anion 130, which is quenched by a proton source to give 131. Owing to its instability, compound 131 processes a [4 + 2] cycloreversion to formation 132, with a loss of nitrogen. Subsequently, the imine of 132 passes through an attack by a second molecule of the TosMIC anion, followed by ring closure, to produce 133. At this point, after protonation, intermediate 134 regains aromaticity via a base-assisted mechanism, with the expulsion of the most acidic proton and loss of hydrogen cyanide and sulfinate. Under strong basic condition, excess t-BuOK deprotonates the newly formed hydrogen cyanide, avoiding the release of toxic HCN. It is worth noting that the two molecules of TosMIC partake in the reaction mechanism in two different paths, with the second molecule in Scheme 37 undergoing a fragmentation resulting in the incorporation of a C–H into the target molecule.

![Scheme 37. A plausible scenario for the van Leusen imidazole synthesis with two TosMIC molecules.](image)

In 2019, the one-pot three-component van Leusen chemistry was used for the DNA-encoded libraries (DELS) synthesis by the Staz group, which provided the first published DNA-compatible approach to form unusual highly functionalized imidazoles. The target DEL productions 138 were obtained from the aldehyde functionalized DNA molecule 136, amine 137, and TosMIC derivatives 18 by the van Leusen reaction. Moreover, a wide variety of amines, commercial TosMIC molecules and aldehyde as the variety element in the three ingredients heterocyclization were investigated under an optimized condition, respectively. This transformation meaningfully expanded the application of van Leusen imidazole synthesis and DNA compatible chemistries (Scheme 38) [59].
In summary, under the in-depth research and application in imidazole-based medicinal chemistry and the progress in other disciplines—such as cell biology, molecular biology, pharmacology, and organic chemistry—an increasing number of imidazole-containing drugs with lower toxic, better efficacy, superior pharmacokinetic characteristics, effective pathologic probes and diagnostic agents would be used in clinics. This could make remarkable contributions for the protection of mankind’s health. Therefore, the van Leusen imidazole synthesis based on TosMICs will play an increasingly central part in the synthesis of bioactive compounds as clinic drugs in drug design and synthesis. We could focus on the changing of the various aldimine groups and TosMIC derivatives in the van Leusen reaction to modify the imidazole derivatives in future. Above all these have clearly indicated the infinite potentiality of van Leusen imidazole synthesis in medicinal chemistry. Additionally, we hope this review would build a full foundation and reference source which would open up new thoughts for researchers to focus on in imidazole-based medicinal molecule design and synthesis chemistry.

Author Contributions: All authors wrote the paper. All authors have read and agreed to the published version of the manuscript.

Funding: Financial support of this research provided by the Science and Technology Planning Project of the Jilin Province (20160414015GH) is greatly acknowledged.

Conflicts of Interest: The authors declare no conflict of interest.

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