

REVIEW



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1,4-Dihydropyridine: synthetic advances, medicinal and insecticidal properties

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1,4-Dihydropyridine (1,4-DHP) is one of the foremost notable organic scaffolds with diverse pharmaceutical applications. This study will highlight recent accomplishments in the construction of 1,4-DHP with structural and functional modifications using multi-component one-pot and green synthetic methodologies. The various intrinsic therapeutic applications, ranging from calcium channel blocker, anti-oxidative, anticancer, anti-inflammatory, anti-microbial, anti-hypertensive, anti-diabetic, anticoagulants, anti-cholinesterase, neuro-protective, and other miscellaneous activities, have been

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summarized with a focus on their structure–activity relationship (SAR) investigations. In addition, the insecticidal properties have been collated and discussed. Researchers in the fields of medicinal chemistry and drug development will find the summarized conclusions of this study incredibly informative, instructional, and valuable.

1. Introduction

Pyridine has remained one of the most remarkable scaffolds in chemistry since its discovery owing to its broad range of uses in fields ranging from medicine to materials.^{1,2} The reduction of the unsaturated double bond(s) forms major scaffolds, which facilitates their wide variety of applications, among the several successful pyridine chemical transformations. Different scaffolds have been created based on the number of double bonds that are decreased. Piperidine (three double bonds), tetrahydropyridines (two double bonds), and dihydropyridines (DHPs) are examples of these compounds (one double bond). These scaffolds have been found in a variety of natural and synthetic substances, with a proclivity for inducing molecules with striking and unusual pharmacological effects. The semi-saturation of one pyridine double bond with two substituents can theoretically generate five isomeric structures of nitrogen heterocyclic DHPs, although the 1,2-DHPs and the 1,4-DHPs are the most prevalent and significant.³ Scheme 1 depicts the reductive chemical transformation of pyridine, and the structural characteristics of the 1,2-DHP and 1,4-DHP scaffolds.

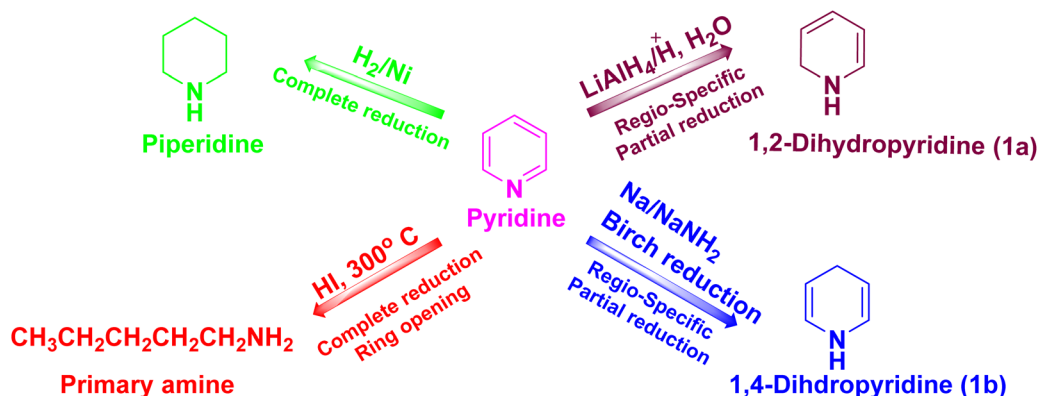
The 1,4-DHP scaffold has gained traction as a preferred choice of medicinal chemistry researchers for developing novel molecules with therapeutic significance, and it is regarded as one of the privileged structures in the drug discovery process due to its ability to import valuable medicinal properties into molecules.^{4–10} Privileged structures, according to the IUPAC definition, exhibit “substructural diversity, which reveal required (often drug-like) activities on molecules comprising that feature. Mostly include a semi-rigid skeleton, on which can exist numerous hydrophobic residues without undertaking hydrophobic collapse”.¹¹ Research data on 1,4-DHP derivatives have been collected from PubMed, an open-access database, and the statistics of the data are depicted in Fig. 1. The data indicate that researchers have been working on the

development of newer 1,4-DHP derivatives since 1969. From 1970 to the 1990s, publications grew dramatically; after that, they began to decline. The largest number of research publications published in a year in the previous ten years was 60, which is somewhat disappointing. Having accessible, thoroughly evaluated information on this research area is one approach to enhancing the research in this field. There have been few attempts at a complete study of the 1,4-DHPs,¹² and only a handful have been discovered with a specific focus on therapeutic regions.^{4,13} There is no study that we are aware of that focuses on the insecticidal agrochemical applications of 1,4-DHP derivatives. To draw researchers' attention to this field, a complete evaluation of the literature on synthetic advances in producing different 1,4-DHP derivatives and their medicinal and agrochemical applications are necessary. This review provides a summary of the current advances in the development of synthetic techniques for 1,4-DHP derivatives and the numerous intrinsic medicinal applications, with an emphasis on their structure–activity relationship (SAR) studies. Additionally, the insecticidal applications as agrochemicals have been compiled and reviewed.

2. Synthetic advances

2.1. The classic reaction: the Hantzsch reaction

A multi-component reaction (MCR) is used in this process, in which two or more starting elements react together to produce pure results.^{14–16} The key benefits of MCRs are that they are environmentally benign, green, and long-lasting reactions for the synthesis of biomolecules from readily available starting materials. Bond breaking and making occur concurrently in a single reaction in MCRs, resulting in the formation of multiple bonds.^{17,18} That reaction produces selected products with a high yield and purity according to MCR's technique. MCRs are becoming increasingly attractive in the production of



Scheme 1 The possible reductions of pyridine and generation of the 1,2-DHP (1a) and 1,4-DHP (1b) scaffolds.

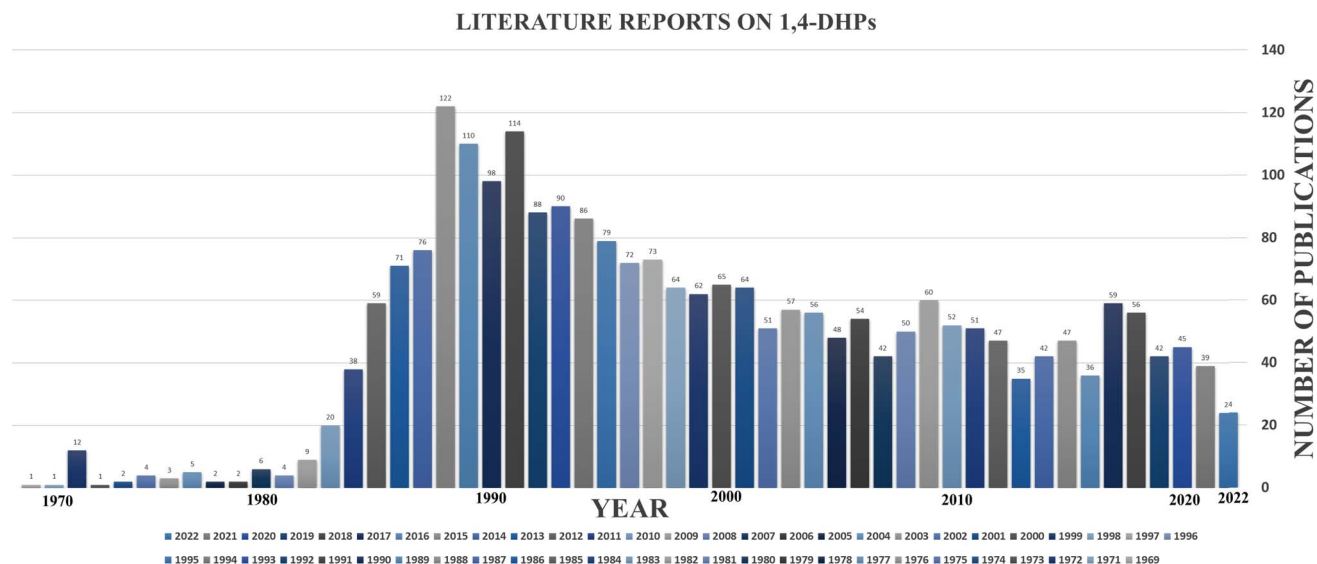
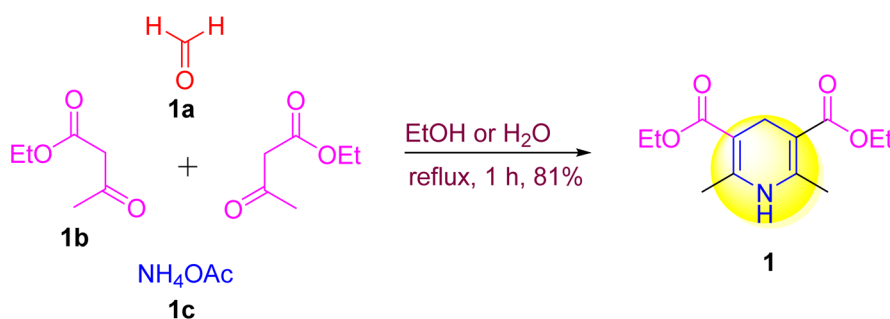


Fig. 1 The number of publications published on 1,4-DHP scaffolds since 1969 (<https://pubmed.ncbi.nlm.nih.gov/?term=1%2C4-dihydropyridine>).



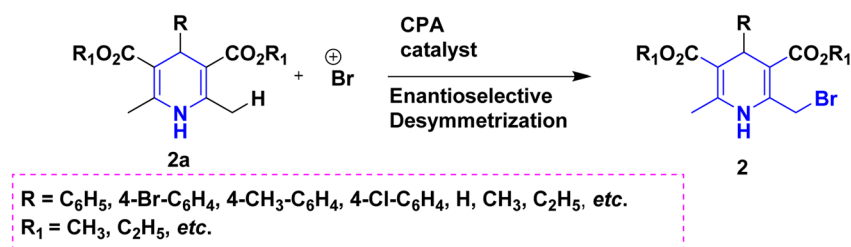
Scheme 2 The Hantzsch synthetic method, a classical 1,4-DHP synthesis.

bioactive chemicals. The Hantzsch reaction, a classic and iconic MCR established for the synthesis of physiologically active heterocyclic (N, S, and O) compounds,¹⁹ was first used to make 1,4-DHP (**1**) (Scheme 2), in which one mole of an aldehyde **1a** reacts with two moles of acetoacetic ester **1b** and an amine **1c** to undergo a four-component, one-pot reaction.

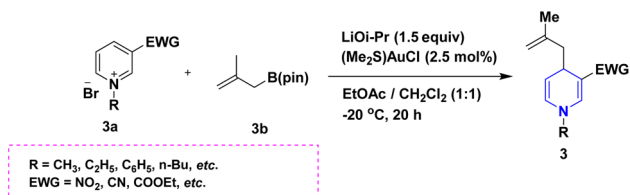
2.2. A decade of advancements in 1,4-DHP synthetic protocols

Owing to the broad range of possible options in synthetic methods development and the construction of structurally

varied 1,4-DHP compounds, the Hantzsch method for the synthesis of 1,4-DHPs has swiftly acquired popularity. To maximize its value, several researchers have been working on future advancements. By desymmetrizing enantio-selective bromination, Han *et al.* (2022) discovered an effective approach for the synthesis of enantio-enrich Hantzsch type 1,4-DHPs **2** starting with symmetrical 1,4-DHPs **2a** with the methyl group at the second or sixth position. The inert C–H bond was changed into the adaptable C–Br bond, ensuring high efficiency in the alteration of the chiral 1,4-DHP derivatives (Scheme 3).²⁰ According to the location of the azinium ion that is attacked,



Scheme 3 Hypothesis for preparation of chiral Hantzsch type 1,4-DHPs.

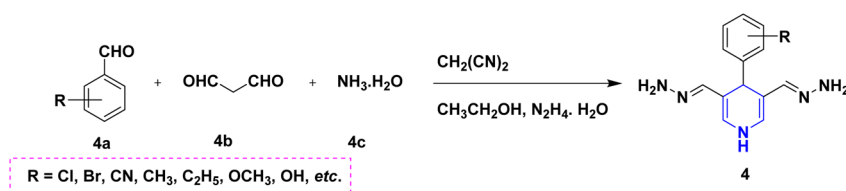


Scheme 4 Nucleophilic allylation of azinium ions.

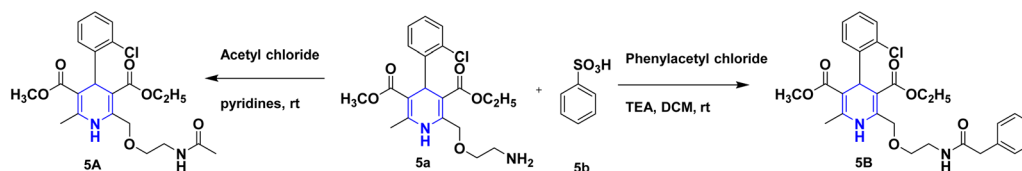
Brien *et al.* (2022) detailed the gold(i)-catalyzed nucleophilic allylations of pyridinium and quinolinium ions **3a** with various allyl pinacol boronates **3b** to yield diverse functionalized 1,4-DHPs **3**. Allyl gold(i) intermediates created by transmetalation from allyl boronates are entirely selective for the reactions. The reactions have unique regioselectivity for attacking the substrates' fourth position, and do not require any additional safety measures to keep out air or moisture (Scheme 4).²¹ A fluorescent hydrazine hydrate probe (DMA) was created by Xue *et al.* (2021) using 1,4-DHP derivatives **4** that were produced from a variety of aldehydes **4a**, malonaldehyde **4b** in the presence of hydrazine hydrate **4c**. This probe's fluorescence emission peak is in the near-infrared (667 nm) range, which exhibits strong hydrazine hydrate selectivity. The probe demonstrates aggregation-induced emission (AIE) properties, which is

significant. To identify hydrazine hydrate in the solution and achieve the quantitative detection of hydrazine hydrate gas, the probe is also equipped with a portable test paper. This probe is capable of quickly and accurately detecting hydrazine hydrate on both a qualitative and quantitative level (Scheme 5).²² In the presence of benzene sulphonic acid **5b** and pyridines, Schiller *et al.* (2022) reported the amlodipine-based acetylated DHP **5A** using acetyl chloride and benzylated 1,4-DHP **5B** using phenyl acetyl chloride. Amlodipine **5a**, at a high dose of 300 μM , showed an inhibitory impact on the four newly synthesized 1,4-DHPs evaluated at hP2X5FL. Low micromolar ATP-induced currents were increased by isradipine and even more nimodipine. The 1,4-DHP derivatives of amlodipine are not effective hP2X5 antagonists, but amlodipine itself may be used as a starting material for future synthesis to improve its affinity. A stimulatory effect on inflammatory processes may also be a side effect of nimodipine therapy (Scheme 6).²³

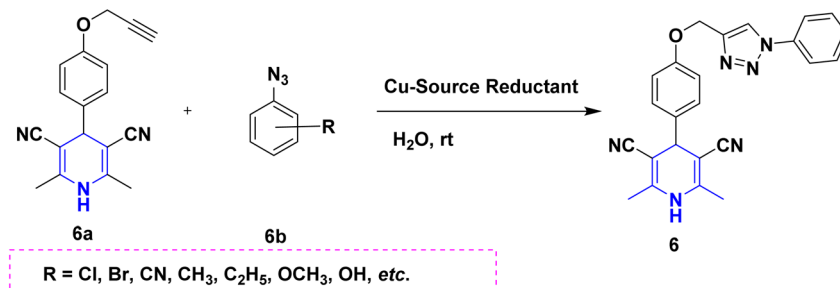
Using alkyne 1,4-DHPs **6a** and substituted azides **6b** for the synthesis of functionalized triazoles, Bijani *et al.* (2022) described a copper-catalyzed azide-alkyne cyclization (CuAAC) reaction for the synthesis of a library of 1,4-DHP-based 1,2,3-triazole derivatives **6** and assessed their cytotoxic potential on colorectal adenocarcinoma. The findings suggested that the majority of these drugs exhibit substantial antiproliferative action, with IC_{50} values ranging from 0.63 ± 0.05 to 5.68 ± 0.14



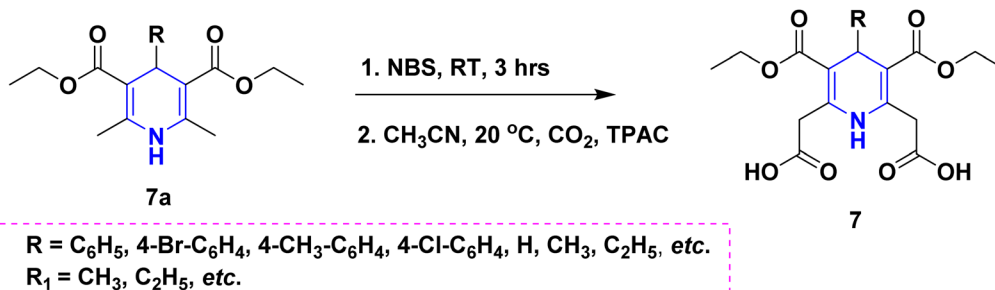
Scheme 5 Synthesis of 1,4-DHP using the fluorescent probe.



Scheme 6 Synthesis of amlodipine 1,4-DHP derivatives.



Scheme 7 Synthesis of 1,4-DHPs based triazole derivatives.



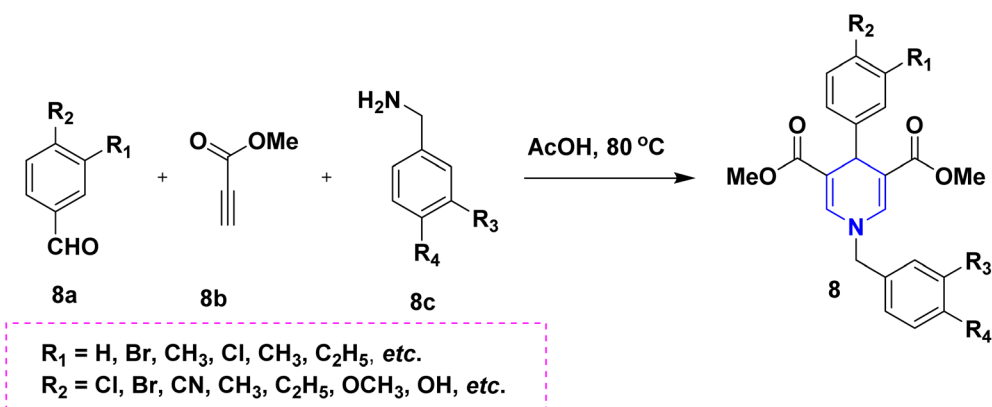
Scheme 8 Synthesis of dicarboxylic-1,4-DHP derivatives.

μM. The available findings suggest that powerful anticancer compounds can be made from 1,2,3-triazole conjugates coupled with DHP (Scheme 7).²⁴ Malhi *et al.* (2022) described a series of 1,4-DHPs with a carboxylic moiety to synthesize 2,2'-[3,5-bis(ethoxycarbonyl)-4-phenyl-1,4-dihydropyridine-2,6-diyl] diacetic acid **7**, the unique dicarboxylic derivatives of 1,4-DHP obtained through the electrocarboxylation of tetrasubstituted-1,4-DHP derivatives **7a**. Additionally, the synthesized compounds exhibit superb activity in a variety of bacteria. Their activity against *E. coli*, *S. aureus*, *B. subtilis*, *A. niger*, and *P. glabrum* was found to be very effective at 4 μg mL⁻¹ in comparison to conventional doses of amoxicillin and fluconazole (Scheme 8).²⁵ In a straightforward one-pot synthesis involving three components of aromatic aldehydes **8a**, methyl propiolate **8b**, and a benzylamine **8c**, Döring *et al.* (2021) reported that the novel 1,4-DHPs **8** are accessible. While those generated with the dicarbonyl molecule have alkyl substituents at the corresponding 1,4-DHP positions, their target was to generate compounds that have no substituents in the second and sixth positions. In an MRP4-overexpressing cell line experiment, all compounds displayed superior activities to the most well-known MRP4 inhibitor MK571, and the activities may be attributed to the varied aromatic residue substitution patterns within the symmetric molecular framework (Scheme 9).²⁶ In a further reaction involving 2-chlorobenzaldehyde **9a** and cyanothioacetamide **9b** in the presence of an excess of *N*-methylmorpholine, Kurskova *et al.* (2022) reported on the production

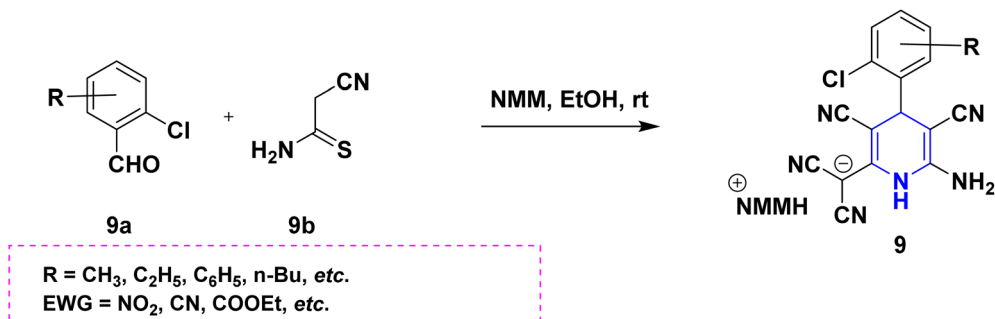
of 2-amino-4-(2-chlorophenyl)-6-(dicyanomethyl)-1,4-dihydropyridine-3,5-dicarbonitrile **9** (Scheme 10).²⁷

In a condensation procedure involving certain aldehyde derivatives **10a**, ethyl acetoacetate **10b**, and ammonium acetate **10c** in the presence of superparamagnetic manganese ferrite nanoparticles at 80 °C, Moradi *et al.* (2022) described the effective one-pot synthesis of several 1,4-DHPs **10**. The advantages of this method for synthesizing 1,4-DHPs include selectivity, high product purity, outstanding yields, quick reaction times, ease of processing, and environmentally favorable settings. Furthermore, the product yield can be slightly decreased by recovering and reusing the catalyst across several runs (Scheme 11).²⁸

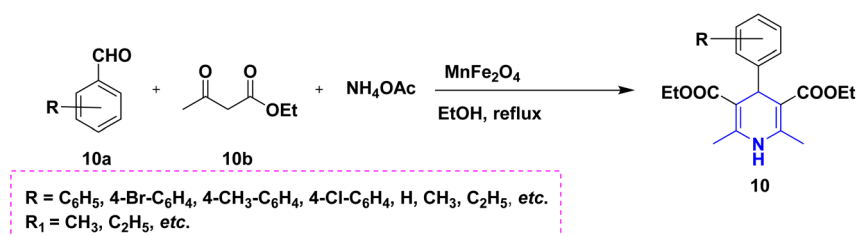
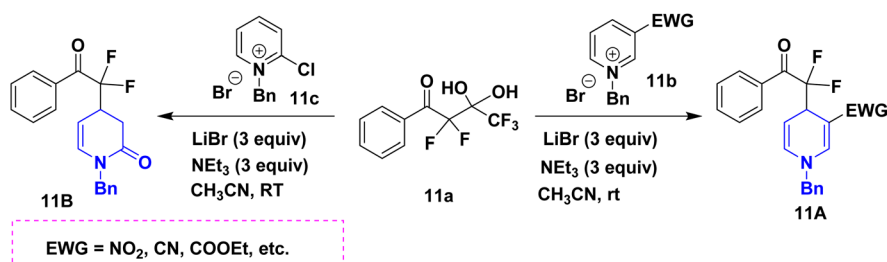
For the first time, Patra *et al.* (2022) took advantage of the activated pyridinium salts with, -difluorinated gem-diols **11a**, creating a highly regioselective and simple synthesis of 1,4-DHPs ornamented with a high-value *gem*-difluoro methylene motif **11** in high yields. The protocol has a wide substrate scope, is scalable, and is compatible with many different functional groups. The DHP and pyridone compound **11B** that were thus produced were particularly noteworthy because they could be used to create densely functionalized oxazabicyclo[3.3.1]nonane scaffolds (Scheme 12).²⁹ The Hantzsch-like process was used to synthesize a novel series of diethyl 3,5-dicarboxylate derivatives of 1,4-DHPs **12A**. As raw materials, aromatic aldehydes **12a**, aromatic amines **12c**, and ethyl propiolate **12b** were used to construct and refine the synthetic methodology shown



Scheme 9 Synthesis of symmetrical 1,4-DHPs.

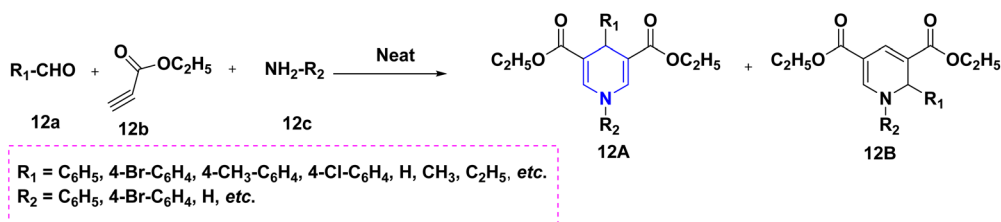


Scheme 10 Synthesis of 1,4-DHPs using NMO.

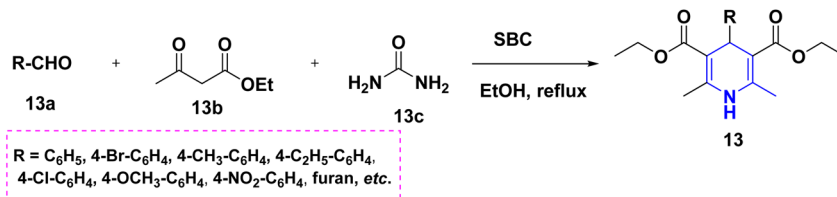
Scheme 11 Synthesis of 1,4-DHPs by MnFe₂O₄ nanoparticles.Scheme 12 Synthesis of 1,4-DHPs adorned with *gem*-difluoromethylene motif.

here. The diethyl-3,5-dicarboxylate of 1,2-DHP, **12B**, was discovered as an unanticipated by-product in this synthesis process (Scheme 13).³⁰ The novel compounds, diethyl 3,5-dicarboxylate derivatives of 1,4-DHPs **13**, have been reported using an array of aromatic aldehydes **13a**, ethylaceto acetate **13b**, and urea **13c** in the presence of sub-critical (SBC) and ethanol as a solvent medium (Scheme 14). This study is significant since it takes less time, uses a traditional work-up, and does not use column chromatography.³¹

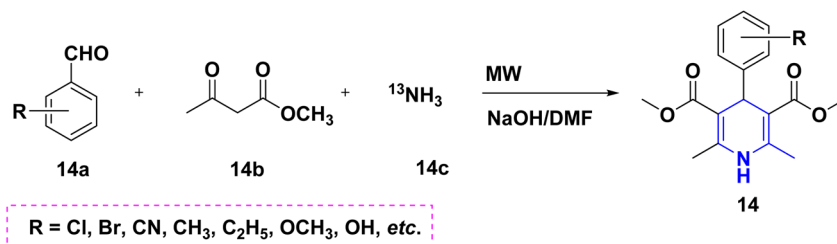
Radioactive ¹³N-labeling experiments were carried out by Blower *et al.* (2021). The findings were interesting and are quite useful for the radiochemical synthesis of [¹³N] 1,4-DHP **14** derivatives using several benzaldehydes **14a** and β-keto ester **14b** in the presence of radiolabeling ammonia **14c** (Scheme 15). The 1,2,4-triazole-tagged 1,4-DHP derivatives **15** were developed with a highly efficient, green MCR protocol by the multicomponent reaction, *viz.*, aldehydes **15a**, 1*H*-1,2,4-triazol-3-amine **15d**, diethyl acetylene dicarboxylate **15b** and active methylene



Scheme 13 Synthesis of 1,4-DHPs using neat conditions.



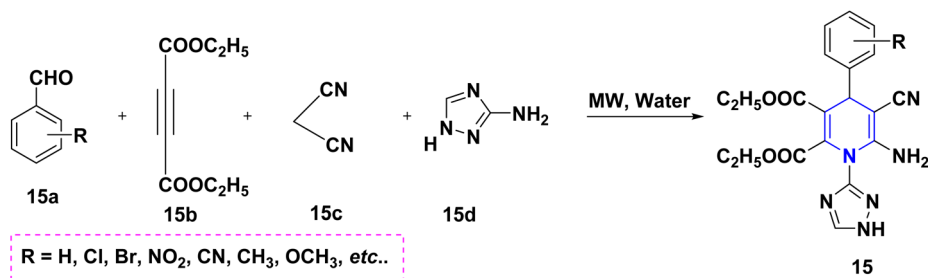
Scheme 14 Synthesis of 1,4-DHPs using SBC conditions.



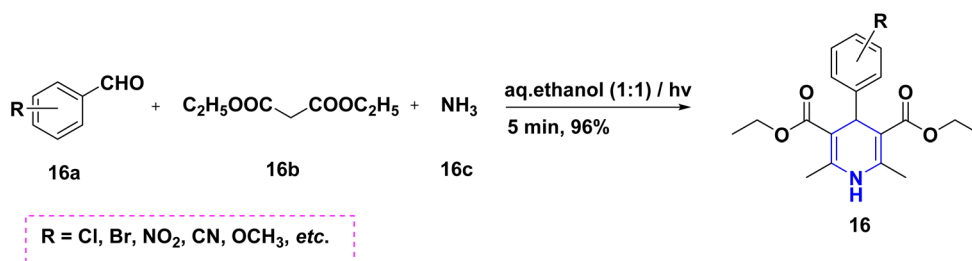
Scheme 15 Synthesis of 1,4-DHP using microwave conditions.

compounds **15c**. Interestingly, the reactions took only 12 minutes to complete in a green water solvent under microwave (MW) conditions (Scheme 16). Excellent yields (94–97%), functional group tolerance, and selectivity were all achieved throughout the reaction.³² Similarly, the condensation of benzaldehyde **16a**, ethyl acetoacetate **16b**, and ammonia **16c** in the presence of aqueous ethanol at room temperature was described as a visible light-mediated technique for the production of poly substituted 1,4-DHPs, **16**. When compared to thermal procedures, this approach has the benefit of minimizing byproduct generation and having quicker reaction times

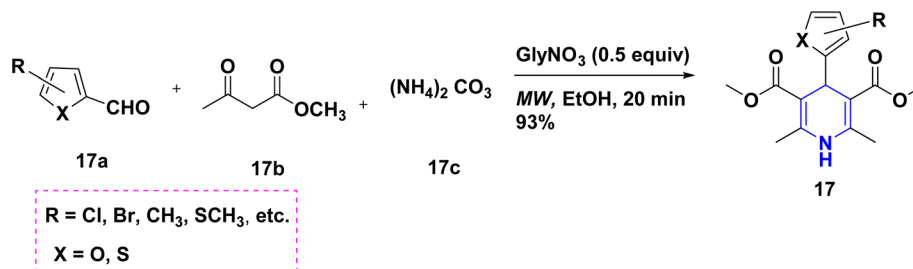
(Scheme 17).³³ By using the same starting ingredients, such as benzaldehyde and ethyl acetoacetate, 1,4-DHPs, **16** are synthesized in the presence of Fe nanoparticles³⁴ and magnesium nitride.³⁵ By using ionic liquid and MW mediation, the catalytic efficiency of glycine nitrate (GlyNO₃) was investigated and developed into a multicomponent reaction of 1,4-DHPs **17** starting from aromatic or heteroaromatic symmetrical or unsymmetrical aldehydes **17a**, methyl acetoacetate **17b**, and ammonium carbonate **17c** (Scheme 18). The use of ionic liquid glycine nitrate to boost product yields was a key feature of this research.³⁶



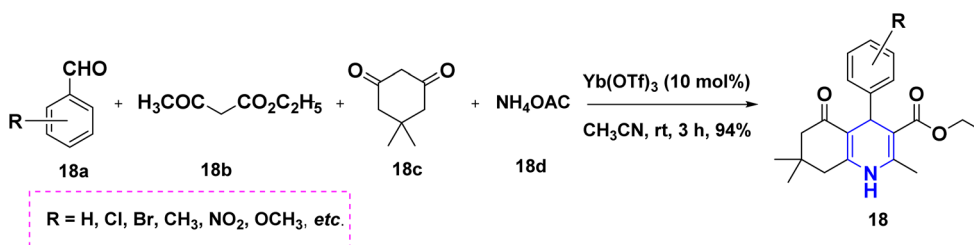
Scheme 16 Synthesis of 1,4-DHP using microwave in water conditions.



Scheme 17 Synthesis of 1,4-dihydropyridine in visible light.

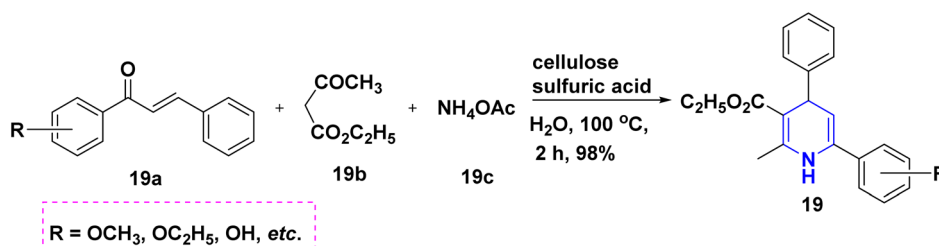


Scheme 18 Glycine nitrate ionic liquid catalyzed synthesis of 1,4-dihydropyridines.

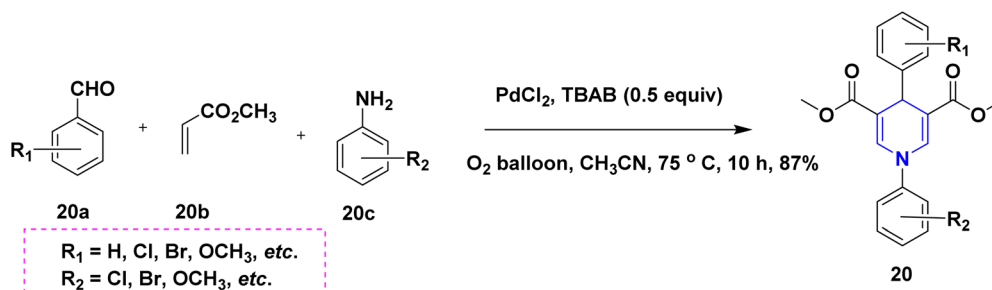
Scheme 19 Yb(OTf)₃ Lewis acid-catalyzed synthesis of 1,4-DHPs.

In 2011, Gestwicki and coworkers devised a synthetic technique for highly functionalized 1,4-DHPs. In the MCR of benzaldehyde **18a**, ethyl acetoacetate **18b**, dimedone **18c**, and ammonium acetate **18d**, Yb(OTf)₃ is utilized as a Lewis acid catalyst. With good yields, this approach has been tested with a variety of aldehydes, including aliphatic, aromatic, and bulky aldehydes. This research is significant since it confirms the use of Yb(OTf)₃, a stable rare-earth metal, as a catalyst (Scheme 19).³⁷ A solid support biodegradable cellulose sulphuric acid

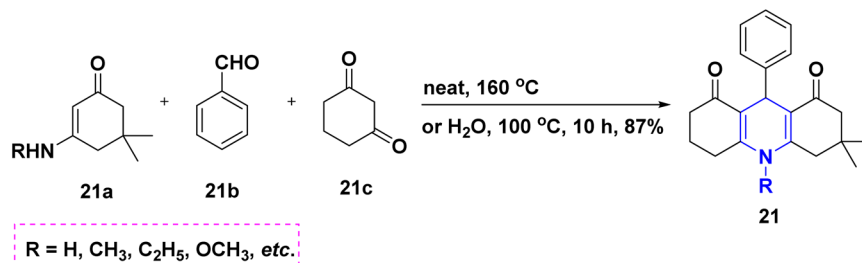
(CSA) was applied in the non-toxic and reusable catalyzed water-mediated one-pot synthesis of C5-unsubstituted 1,4-DHPs, **19** with excellent yields using chalcones **19a**, ethyl acetoacetate **19b** and ammonium acetate **19c** (Scheme 20).³⁸ The development of highly economical cellulose sulphuric acid as a catalyst and C5-unsubstituted 1,4-DHPs are the high points of this work. The palladium(II) catalyzed cascade reaction of 1,4-DHPs **20** started from substituted benzaldehyde, **20a**, aniline **20b** and electron-deficient alkenes **20c** in the presence of O₂ atmosphere



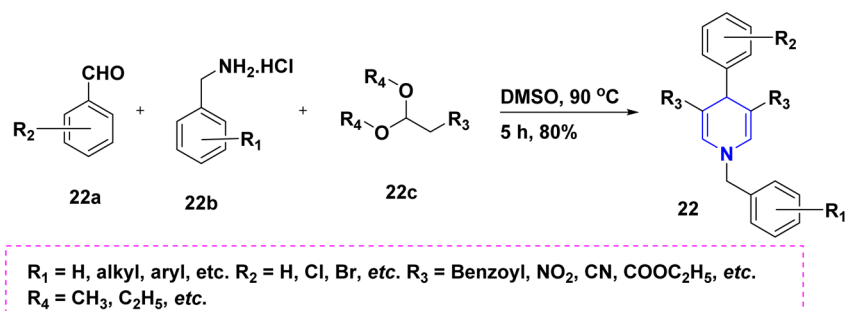
Scheme 20 The cellulose sulphuric acid mediated synthesis of C5-unsubstituted 1,4-DHPs.



Scheme 21 Palladium-assisted synthesis of 1,4-DHPs.



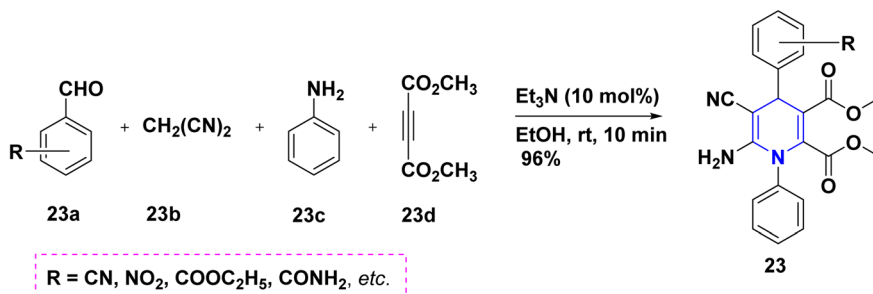
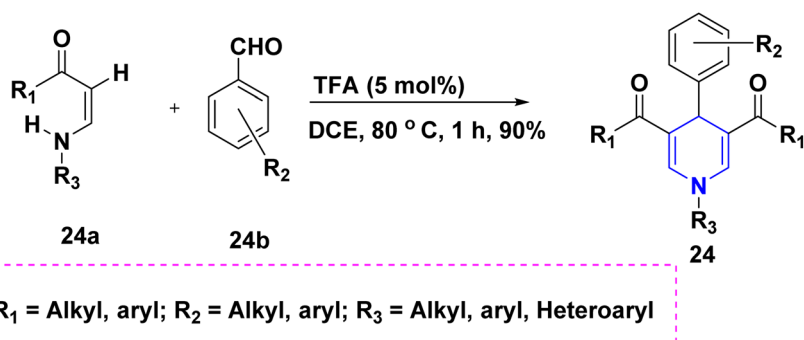
Scheme 22 Synthesis of unsymmetrical 4-arylacridinediones under neat condition.



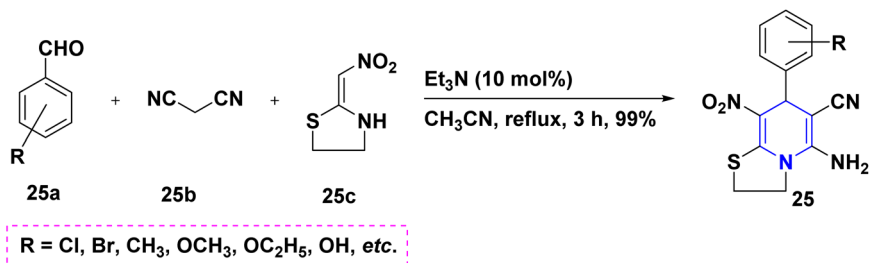
Scheme 23 Three-component reaction for the generation of 1,4-DHPs.

(Scheme 21).³⁹ The highlight of the method is an exploration of an effective Pd-catalyst to provide anti-Markovnikov products using primary anilines and alkenes. The unsymmetrical 1,4-DHPs **21** has been investigated from enaminones **21a**,

benzaldehyde **21b**, and 1,3-cyclohexanedione **21c** in the presence of neat conditions (Scheme 22).⁴⁰ The access to a complex product, green solvent and no column chromatography method are some of the advantages of this technique.

Scheme 24 Et_3N catalysed synthesis of polysubstituted 1,4-DHPs.

Scheme 25 TFA catalyzed synthesis of 1,4-DHPs.



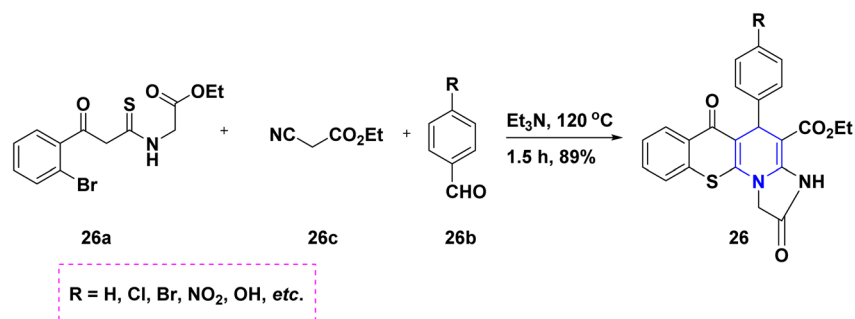
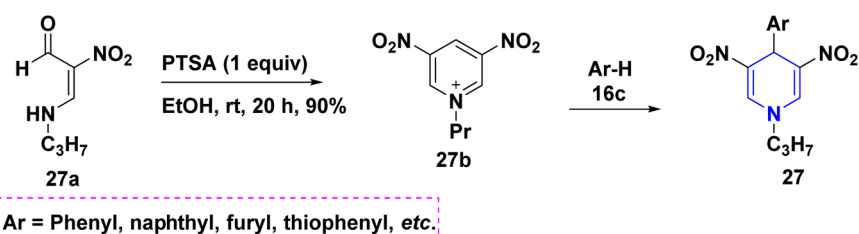
Scheme 26 Synthesis of 3-cyano-5-nitro-substituted 1,4-DHPs.

Benzaldehyde **22a**, quaternary ammonium hydrochloride salt **22b**, and acetals **22c** react to form 1,4-DHP **22** with an 80% yield without the aid of a catalyst. The synthesized 1,4-DHPs also displayed a diversity of fluorescence wavelengths as a result of various derivatives at the third and fifth positions (Scheme 23).⁴¹ The one-pot cascade reaction of benzaldehyde **23a**, malononitrile **23b**, arylamine **23c**, and acetylene dicarboxylate **23d** provide highly substituted 1,4-DHPs **23** with excellent yields, as shown in (Scheme 24).⁴² Trifluoroacetic acid (TFA) catalyzed the synthesis of 1,4-DHPs **24** *via* domino reaction of *p*-nitrobenzaldehyde **24b**, enamines **24a** (Scheme 25).⁴³

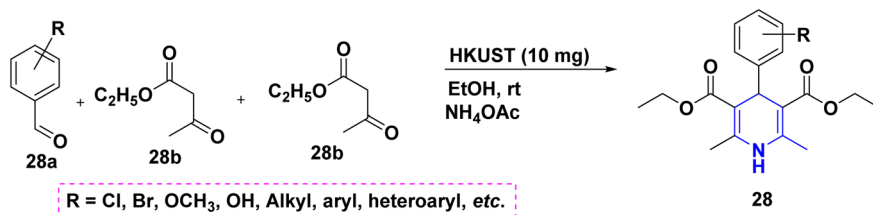
By reacting different benzaldehydes **25a**, malononitrile **25b**, and cyclic nitroketene *N,S*-acetals **25c** with a triethylamine (Et_3N) base, it is possible to create the highly substituted thiazolo[3,2-*a*]pyridines **25**. The focus of this work is the investigation of *N,S*-acetals and their derivatives (Scheme 26).⁴⁴ An Et_3N -mediated one-pot cascade ring formation reaction towards the synthesis of imidazo[1,2-*a*]thiochromeno[3,2-*e*]pyridines **26** started from substituted *N,S*-acetals **26a**, ethyl cyanoacetate **26b** and benzaldehydes **26c** under neat conditions (Scheme 27).⁴⁵ *p*-

Toluenesulfonic acid (PTSA) catalyzed the condensation reaction of two moles of 2-formyl-2-nitroenamine **27a** and one mole of electron-rich aromatic compound **27b** towards the synthesis of C4-substituted 3,5-dinitro-1,4-DHPs **27** (Scheme 28).⁴⁶

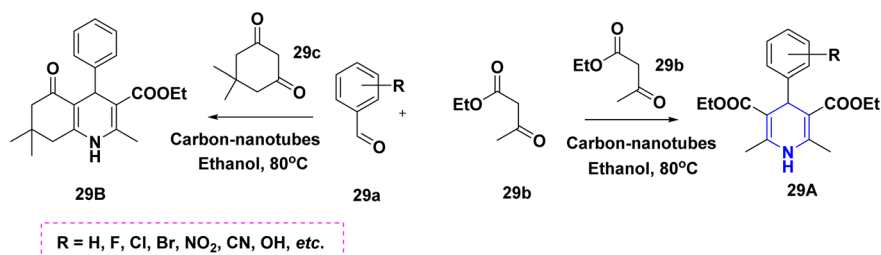
The one-pot multicomponent reaction of benzaldehyde **28a** with ethyl acetoacetate **28b** in the presence of metal-organic catalyst HKUST-1 (a type of MOFs with a molecular formula of $[\text{Cu}_3(\text{BTC})_2(\text{H}_2\text{O})_n]$) towards the synthesis of 1,4-DHPs **28** has been carried out in neat conditions. This protocol affords numerous advantages such as good yields with high purity, no column chromatography, short reaction time and recyclability of the catalyst (Scheme 29).⁴⁷ A similar synthesis of 1,4-DHPs was obtained by using a recyclable biocatalyst, nicotinic acid⁴⁸ and alginic acid.⁴⁹ Similar single pot cascade reactions for the synthesis of 1,4-DHPs have been reported by applying other conditions such as tetrabutylammonium hydrogen sulfate (TBAHS),⁵⁰ Brønsted acid MIL-101- SO_3H ⁵¹ and MW irradiation.⁵² A new one-pot cascade reaction of aldehydes **29a**, ethyl acetoacetate **29b**, and dimedone **29c** was carried out to give 1,4-DHPs **29A** and **29B** using aminated multi-walled carbon nanotubes

Scheme 27 Synthesis of imidazo[1,2-*a*]thiochromeno[3,2-*e*]pyridine.

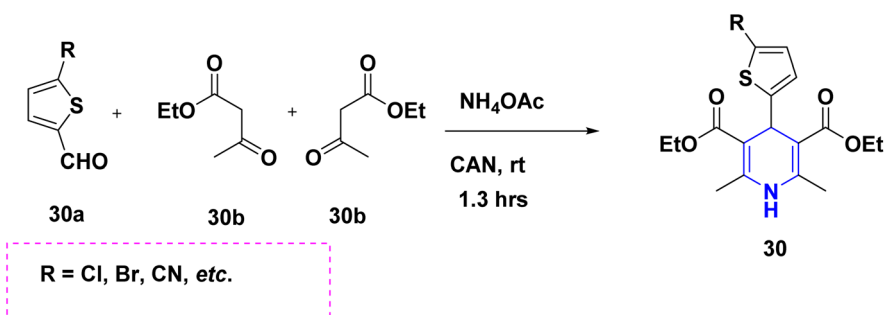
Scheme 28 Synthesis of 3,5-dinitrosubstituted 1,4-DHPs.



Scheme 29 HKUST-1 catalyzed synthesis of 1,4-DHPs under solvent-free conditions.



Scheme 30 Preparation of 1,4-DHPs in the presence of aminated MWCNTs.

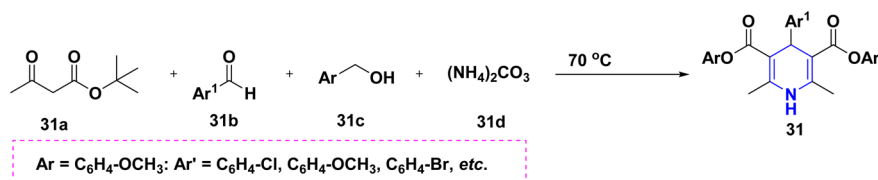


Scheme 31 Synthesis of 1,4-DHP using CAN catalyst.

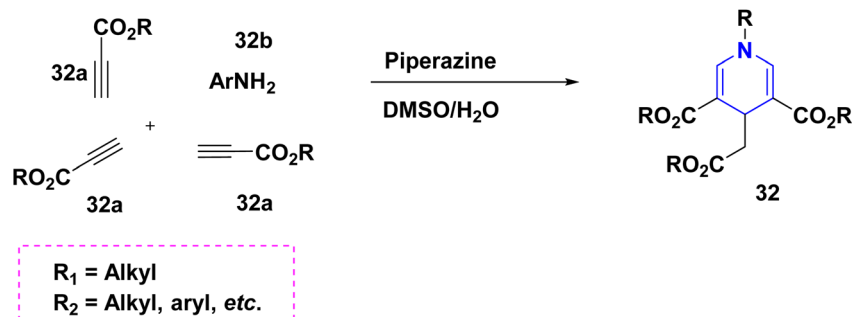
under heat conditions. The benefits of this method include the recyclable catalyst, easy workup and good yield of products (Scheme 30).⁵³ Ceric ammonium nitrate (CAN) catalyzed the tandem multicomponent reaction of various 1,3-diones and 5-bromothiophene-2-carboxaldehyde **30a**, ethyl acetoacetate **30b**, for the synthesis of 1,4-DHPs **30** with significant yields (Scheme 31).⁵⁴

According to Dharma Rao *et al.* (2018), an effective synthesis of symmetrical 1,4-DHP derivatives **31** was initiated from *tert*-butyl- β -ketoester **31a**, aryl aldehyde **31b**, benzyl alcohol **31c**, and ammonium carbonate **31d** at 70 °C in the presence of a green

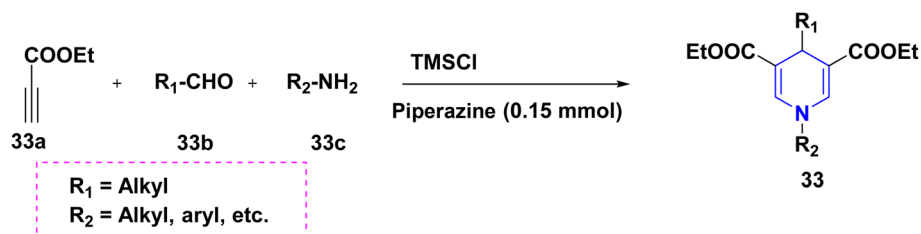
solvent and under neat conditions (Scheme 32).⁵⁵ The one-pot domino methodology was established for the synthesis of 4-DHPs carrying a C4 acetate moiety **32**. They were made from alkyl propiolates **32a** and primary amines **32b** in DMSO solvent with piperazine as a base. The application has been realized with ease, and allows for the efficient and high-yield production of 1,4-DHPs (Scheme 33).⁵⁶ Jieping Wan *et al.* (2014) described the symmetrical 1,4-DHPs **33** produced in the presence of TMSCl from electron-deficient alkynes **33a**, aldehydes **33b**, and amines **33c** by utilizing a secondary amine for the preparation of *N*-substituted 1,4-DHPs started from aldehyde **33b** with



Scheme 32 Synthesis of 1,4-DHP under catalyst-free conditions.



Scheme 33 Synthesis of 1,4-DHP under piperazine catalyst.

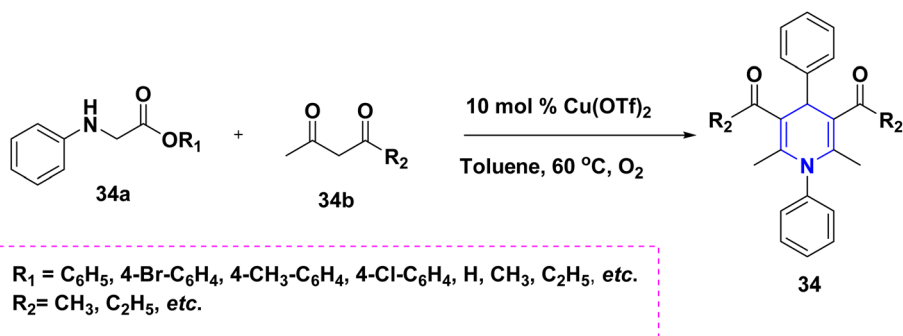
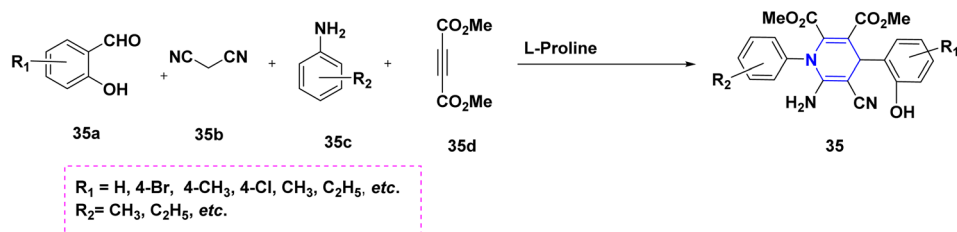


Scheme 34 Synthesis of 1,4-DHP using piperazine (0.15 mmol) and TMSCl catalyst.

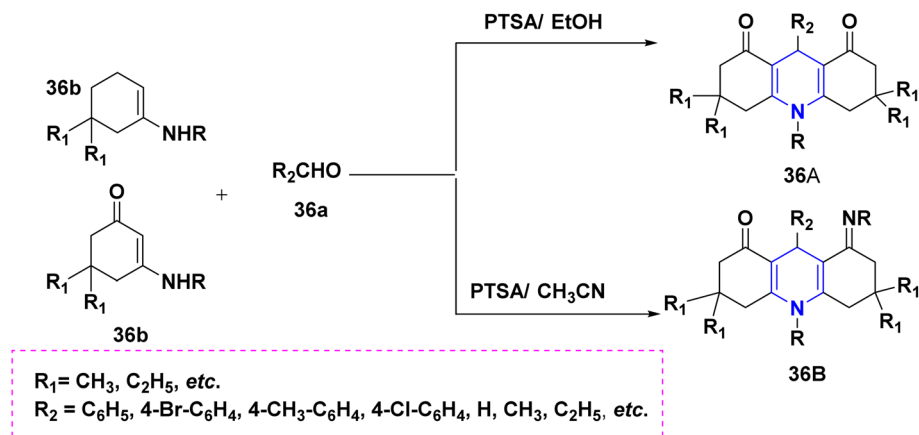
acetylene derivative **33a** via enaminoester intermediates (Scheme 34).⁵⁷

Zhu *et al.* (2016) used copper as a less hazardous catalyst and molecular oxygen as the terminal oxidant to synthesize highly substituted 1,4-DHPs **34** generated from *N*-arylglycine esters **34a** and 1,3-dicarbonyl compounds **34b**. The key advantage of this one-pot tandem reaction was the excellent product yield

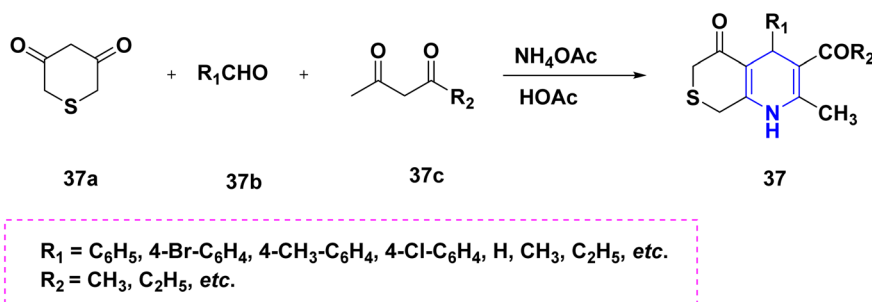
(Scheme 35).⁵⁸ Perumal *et al.* (2014) described a cascade process towards the synthesis of DHPs **35** generated from aldehydes **35a**, malononitrile **35b**, amines **35c**, and activated alkynes **35d** under neat conditions. Huisgen dipolar additions are used in this cascade technique to produce 1,4-DHPs as a single diastereomer (Scheme 36).⁵⁹ Using PTSA in ethanol and acetonitrile as solvents, Liu *et al.* (2018) reported an atom economy reaction of

Scheme 35 Synthesis of 1,4-DHPs using $\text{Cu}(\text{OTf})_2$ catalyst.

Scheme 36 Synthesis of 1,4-DHP using L-proline catalyst.



Scheme 37 Synthesis of 1,4-DHP using PTSA.

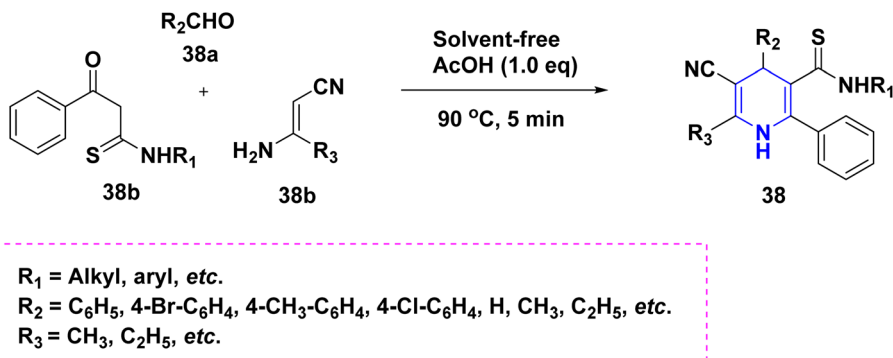
Scheme 38 Synthesis of 1,4-DHP using NH_4OAc .

enaminones **36b** with aldehydes **36a** to give symmetrical 1,4-DHPs **36A** and **36B**. The production of four carbon–carbon bonds and two carbon–nitrogen bonds in a single pot procedure are the highlights of this reaction (Scheme 37).⁶⁰ Chang-Sheng Yao *et al.* (2010) developed a one-pot multicomponent process using active methylene **37a**, aldehyde **37b**, and ketoester **37c** for the synthesis of libraries of thiopyrano[3,4-*b*]pyridine derivatives **37** with shortened reaction times with excellent yields (Scheme 38).⁶¹ Li *et al.* (2013) described a study towards the preparation of a chemo selectivity product of 1,4-DHPs in higher yields **38** by using the simple starting materials such as β -ketothioamides (KTAs), aldehydes **38a** and β -enaminonitriles

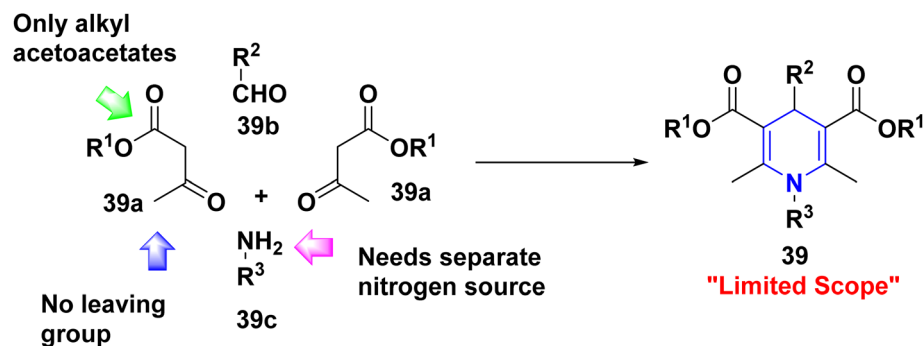
38b in the presence of AcOH under a neat protocol (Scheme 39).⁶²

2.3. Limitations of Hantzsch 1,4-DHPs synthesis

Even though the Hantzsch 1,4-DHP synthesis is a well-known reaction, it has a few disadvantages, including the use of only alkyl acetoacetates as substituents, the difficulty of adding an aryl group in the -keto ester, and the need for a distinct nitrogen supply (Scheme 40). Exploration of innovative techniques toward the synthesis of highly functionalized 1,4-DHPs by overcoming these challenges will now receive a lot of attention.



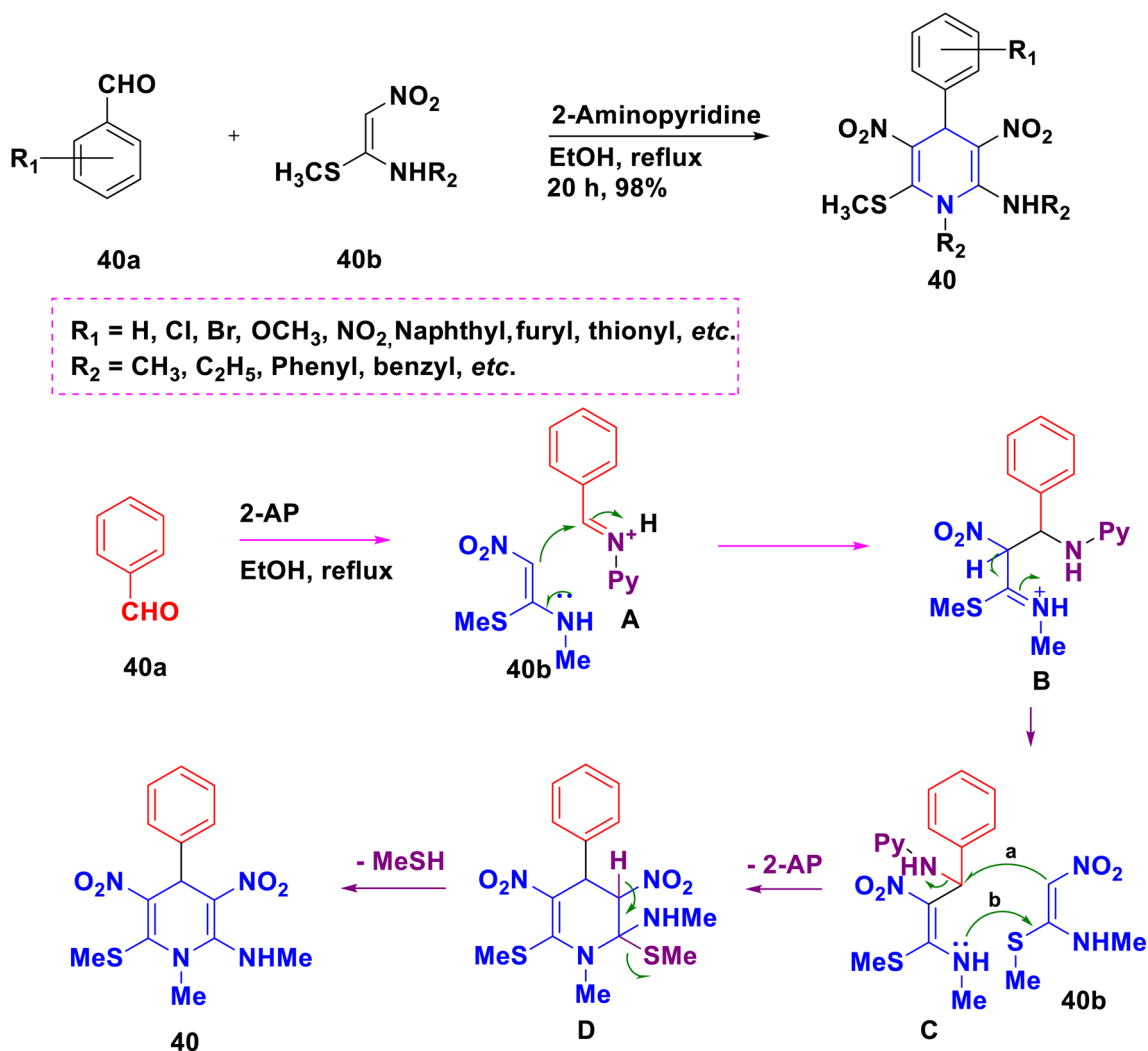
Scheme 39 Synthesis of 1,4-DHP neat conditions.



Scheme 40 1,4-DHPs Hantzsch synthesis and limitations.

Surya Prakash Rao and Parthiban (2014) investigated the chemistry of *N*-methyl-1-(methylthio)-2-nitroethenamine (NMSM) **40b** as a nitrogen source, and used one-pot MCR to synthesize poly substituted 1,4-DHPs **40** from aldehydes **40a** and NMSM **40b** with 2-aminopyridine (2-AP) as a base (Scheme 41).⁶³ As illustrated in Scheme 41, a plausible mechanism has

been presented. The iminium ion **A** is formed in the first step when benzaldehyde **40a** combines with 2-AP. The iminium ion which interacts with NMSM **40b** provides the intermediate **B**, and swiftly changes to give the unstable intermediate **C**, which has a more stable nitroketene-*N,S*-acetal substructure. Intermediate **D** and 2-AP are formed when intermediate **C** interacts



Scheme 41 The pseudo three components for the synthesis of 1,4-DHP and plausible mechanism.

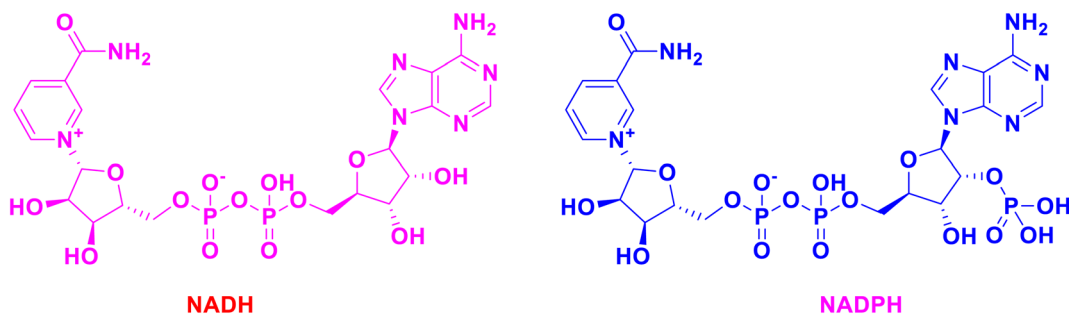


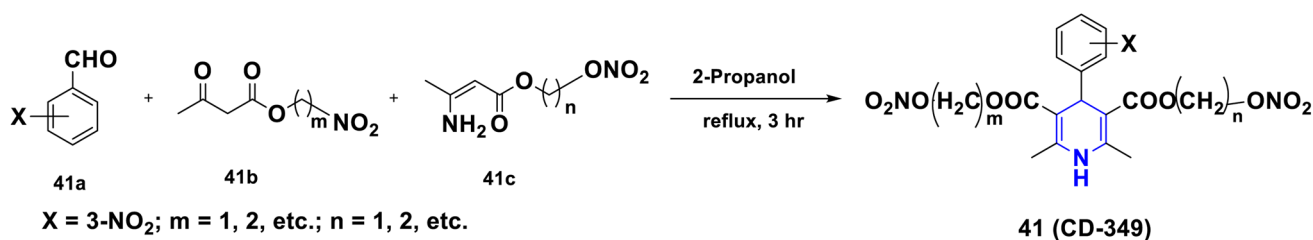
Fig. 2 Structure of NADH and NADPH.

with one additional unit of NMSM **40b**. Finally, methanethiol is eliminated intramolecularly, yielding 1,4-DHP **40**. The fact that primary amines stimulate the MCR suggests that iminium ion **A** is formed as the first intermediate.

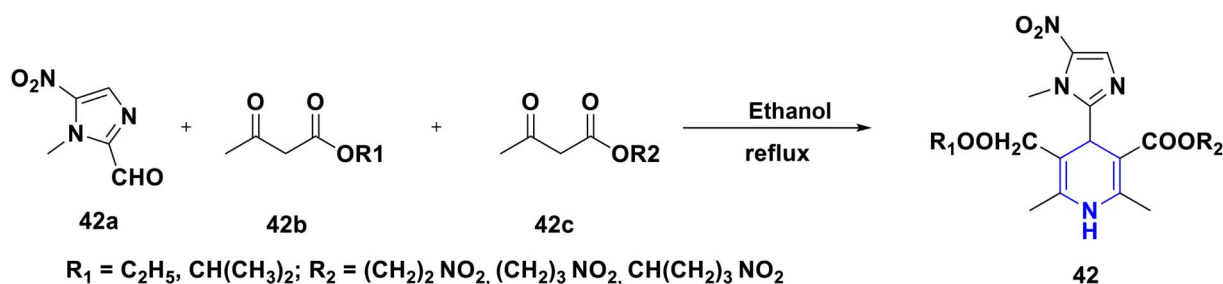
2.4. Medicinal importance of 1,4-DHPs

For nearly a century, the 1,4-DHPs have been known for their enormous biological value and uses. Both NADH and NADPH have reactive portions that function as cofactors in enzymes that perform oxidation–reduction processes, while keeping 1,4-DHP in their structures (Fig. 2).^{64,65} Synthetic 1,4-DHPs have a wide range of therapeutic uses.^{66,67} As a result of their wide range of applications and vast structural variety, 1,4-DHPs have become a favored scaffold among synthetic and medicinal chemists (Fig. 2).^{68–72} To aid medicinal chemists and biologists in identifying and rationalizing the molecule's structural and functional importance in inducing the particular activity, some of the most notable medicinally important molecules of 1,4-DHPs, as well as the synthetic protocols used to achieve them, are discussed herein.

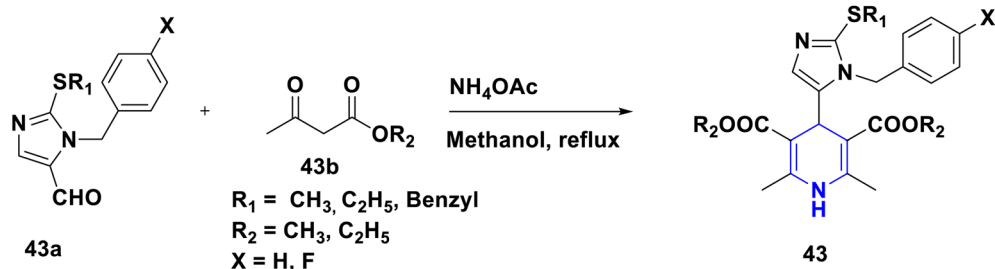
2.4.1. Antihypertensive activity. The role of nitrooxyalkylester insertion at the 3- and 5-positions of 1,4-DHPs **41** in exerting antihypertensive action was discovered by Ogawa *et al.* (1993). As indicated in the scheme, the synthesis involves an MCR reaction between the corresponding aldehyde **41a** and the nitrooxyalkylester **41b** and amine source **41c**. The study discovered **CD-349**, a chemical compound that is more powerful than the antihypertensive medication, nifedipine (Scheme 42).⁷³ Nekoeian *et al.* (2009) reported on novel nitroxyalkyl-containing 1,4-DHP derivatives **42** in a similar way. The compounds were tested in two-kidney and one-clip hypertension rat models. In comparison to nifedipine, a commonly used medicine, the two nitroxy group-containing compounds have shown a weak lowering impact on the mean arterial pressure. Because of its putative interaction with Ca channels, the substituent at the fifth position of 1,4-DHP had a significant influence on inducing the antihypertensive ability, according to the SAR. Furthermore, it is established that the presence of the nitroxy group is not required for the action. When compared to nifedipine, a medication with no nitroxy group, there is no



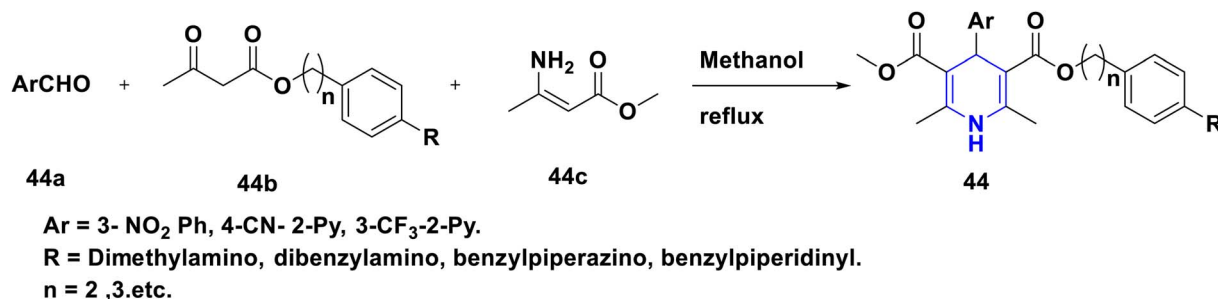
Scheme 42 The antihypertensive activity of CD-349.



Scheme 43 Antihypertensive activity of nitroxyalkyl 1,4-DHP derivatives.



Scheme 44 Antihypertensive activity of 5-imidazolyl DHP derivatives.

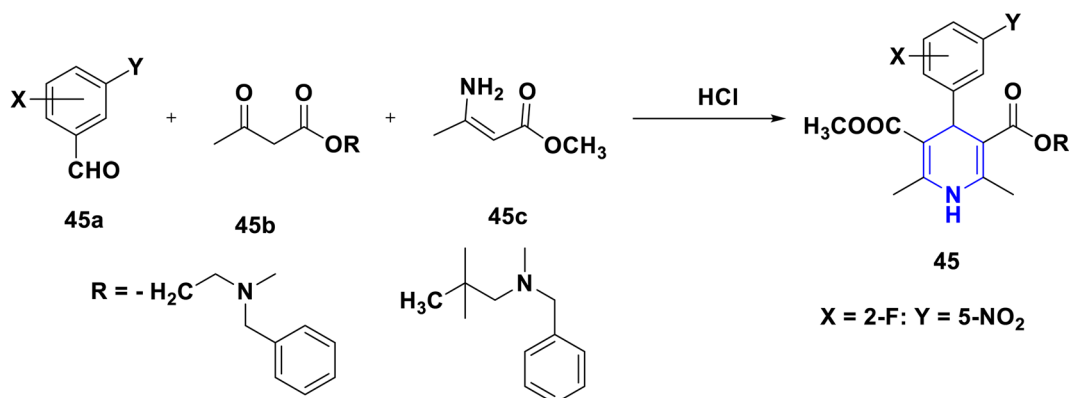


Scheme 45 Antihypertensive activity of 1,4-DHPs bearing 3-[4-(substituted amino)phenyl]alkyl ester derivatives.

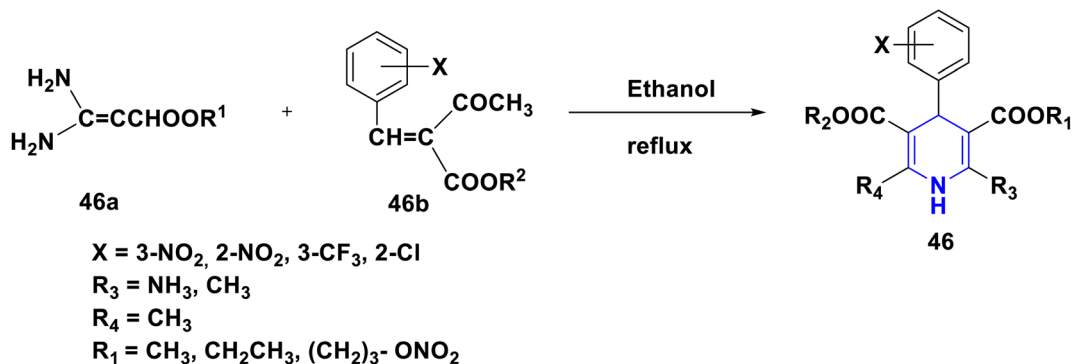
indication of a drop in MAP (Scheme 43).⁷⁴ Hadizadeh *et al.* (2009) developed and synthesized a new class of imidazolyl comprising 1,4-DHPs 43. DOCA-induced hypertension in rats was used to test the drug's antihypertensive efficacy. The influence of the benzyl ring, particularly the *para* position replacement with electron-withdrawing groups, was shown in SAR analyses. All of the examined substances improved the antihypertensive properties. The molecule with the fluorine substituent ($\text{X} = \text{F}$) was shown to be the most active, and even more effective than the common medication amlodipine (Scheme 44).⁷⁵ Novel 3-[4-(substituted amino)phenyl]esters containing 1,4-DHP derivatives 44 were developed, produced, and evaluated for antihypertensive efficacy by Ashimori *et al.* (1991). The results are encouraging, with a longer duration and stronger impact than nifedipine, the usual medicine.

Researchers were encouraged by the positive results to pursue the compounds for further pharmacological exploration (Scheme 45).⁷⁶

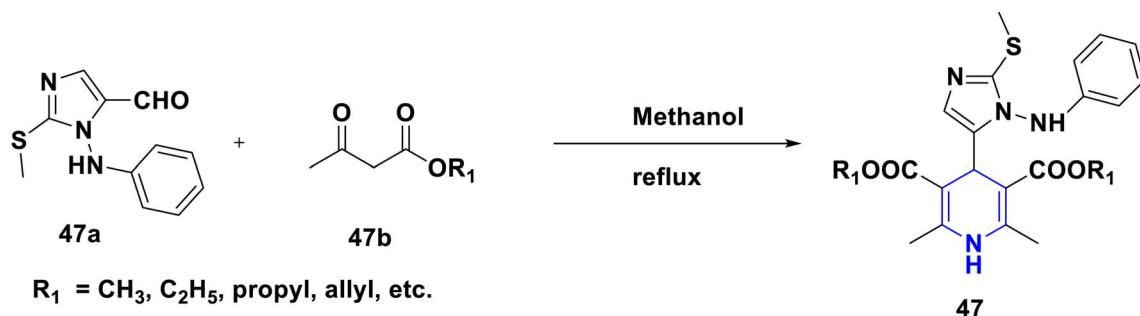
Kanno *et al.* (1992) developed and synthesized a variety of novel 1,4-DHPs with amino-2,2-dimethyl-propyl ester groups or 4-disubstituted phenyl and an aminoethyl ester 45, and tested their antihypertensive efficacy in normotensive and spontaneously hypertensive rats. In comparison to nifedipine and nicardipine, several have shown strong activity and a longer duration of their effect. The 2-fluoro-5-nitrophenyl at the fourth position and 2,2-dimethylaminopropylester at the third position in 45 make it the most active compound in this class (Scheme 46).⁷⁷ For the screening of the antihypertensive activity of their synthesized novel 2-amino-1,4-DHP compounds with nitroxy-alkoxycarbonyl group substitutions at the third or fifth



Scheme 46 Antihypertensive activity of 1,4-DHPs bearing 3-aminoethylester derivatives.



Scheme 47 Antihypertensive activity of 1,4-DHPs bearing nitroxy-alkoxycarbonyl derivatives.



Scheme 48 Antihypertensive activity of 1,4-DHPs bearing 2-methylthio-1-phenylamino-5-imidazolyl derivatives.

position in **46**, Koyabhashi *et al.* (1995) utilized spontaneously hypertensive rats. Compounds having *tert*-amino groups on each side of an ester chain have been shown to have a powerful and long-lasting effect, and have been discovered to be more powerful than nifedipine (Scheme 47). Fassihi *et al.* (2004) described the analogues of nifedipine alkyl ester **47**. The 2-methylthio-1-phenylamino-5-imidazolyl group has been substituted for the *o*-nitrophenyl group at the fourth position in the design. The calcium-channel antagonist properties of the synthesized derivatives were tested utilizing a higher-potassium contraction of guinea-pig ileal longitudinal smooth muscle. More than two methylene chains in the position of the third and fifth ester substituents lowered the activity of the symmetrical

aliphatic esters and phenyl alkyl ester series, according to SAR analyses. Overall, the compounds with an isopropyl ester substitution at the third and fifth positions exhibited promising activity in comparison to the standard drug, nifedipine. Amazingly, **47** derivatives were shown to be more effective than nifedipine (Scheme 48).⁷⁸

SAR has been established, and is graphically depicted in Fig. 3 based on the interpretation of the results. 1,4-DHP's central core is a necessary prerequisite for performing the activity. The common core requirement to execute the activity was discovered to be the methyl group substitutions at the second and sixth positions, and the ester substitutions at the third and fifth positions. It was found that 1,4-DHPs with

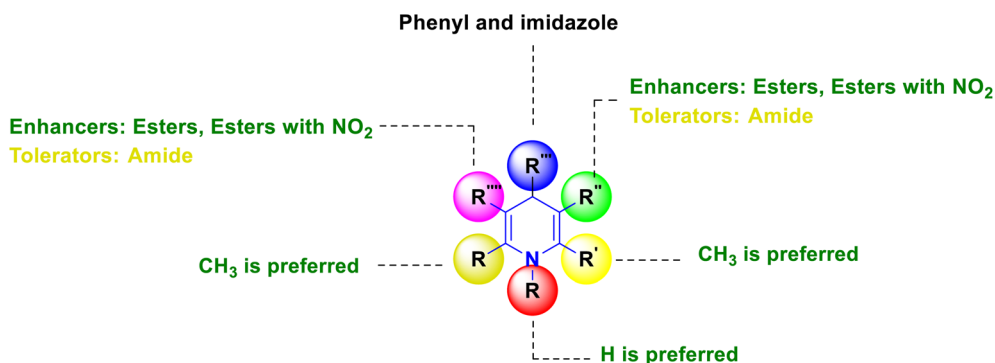
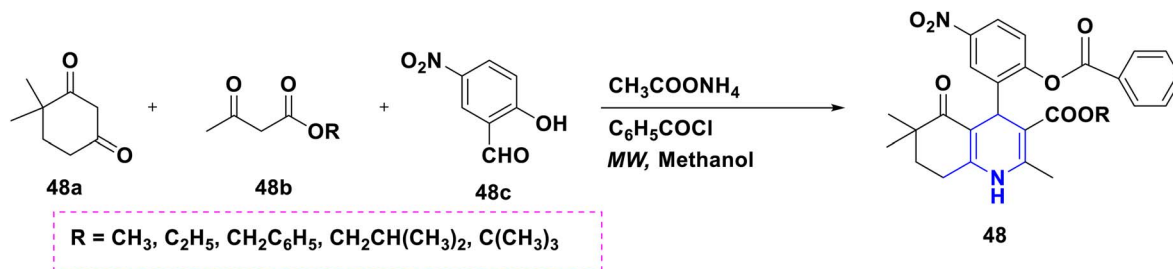
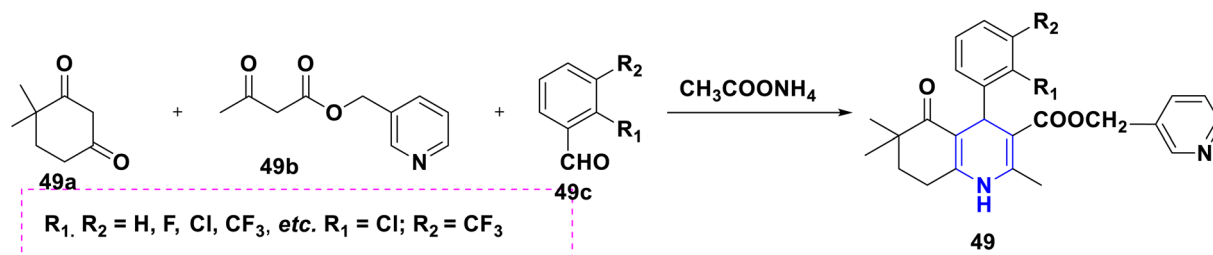


Fig. 3 SAR studies of antihypertensive activity of 1,4-DHPs.



Scheme 49 Cyclohexane ring fused 1,4-DHPs as potential calcium channel blockers.



Scheme 50 The condensed 1,4-DHP ring system act as potential calcium channel blockers.

a nitroxy ester in the third position are more potent than nifedipine. When compared to other compounds that induce activity by electron-withdrawing groups, compounds having a fluoro group at the fourth position of the phenyl ring demonstrated superior activity. Similarly, the substitution of phenyl, 4-(substituted pyridyl) and imidazole at the fourth position have significantly improved activity.

2.4.2. Calcium channel blockers. Ozer *et al.* (2018) synthesized cyclohexane ring fused 1,4-DHPs **48**, and investigated the effects on the calcium channel blocking activities by substituents with different ester moieties and the benzoyl inserted in the 4-phenyl ring. The observed pharmacological data revealed that, while all compounds are powerful relaxants on the isolated rat aortic smooth muscle, adding a benzyloxy substituent to the phenyl ring reduced their relaxant activity (Scheme 49).⁷⁹ Bladen

et al. (2014) synthesized libraries of a new 1,4-DHP ring structure (hexahydroquinoline) **49**, and showed that they can block both L- and T-type calcium channels. Modification of a crucial ester group within this class of compounds not only adjusts the blocking affinity for both L- and T-type channels, but also permits the synthesis of DHPs with 30-fold selectivity for T-type channels over L-type channels. These results indicate that a pharmacophore based on a condensed DHP-based scaffold might be used to develop a novel class of T-type selective inhibitors (Scheme 50) (Fig. 4).⁸⁰ The SAR investigation on the fourth, fifth, and sixth positions of the series of 1,4-DHPs APJ2708 (**50A**) published by Yamamoto *et al.* (2006) identified a new class of N-type calcium channel blockers with fewer cardiovascular side effects than the standards, cilnidipine and APJ2708. The compounds, 4-(3-chloro-phenyl)-2-methyl-6-

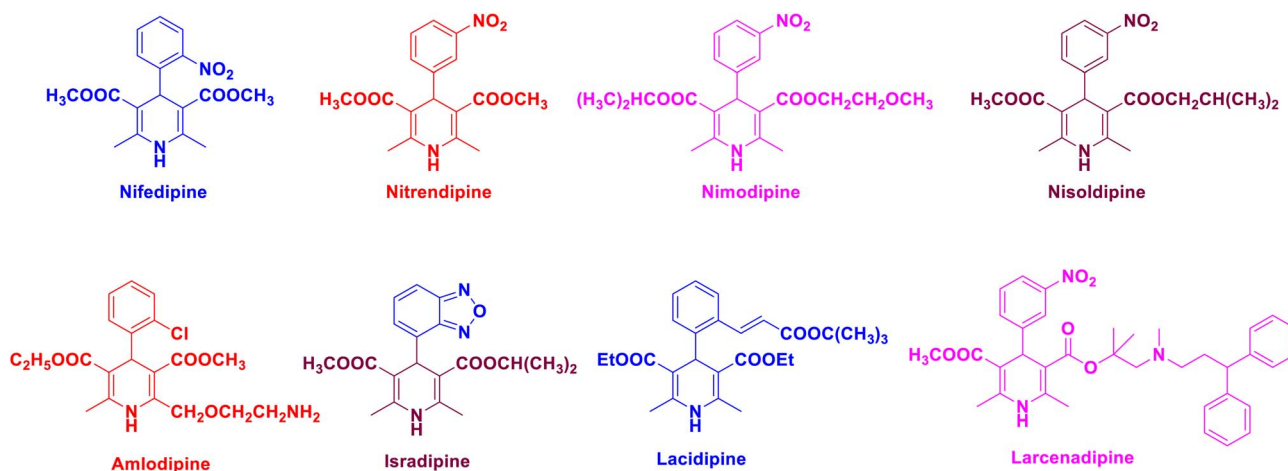
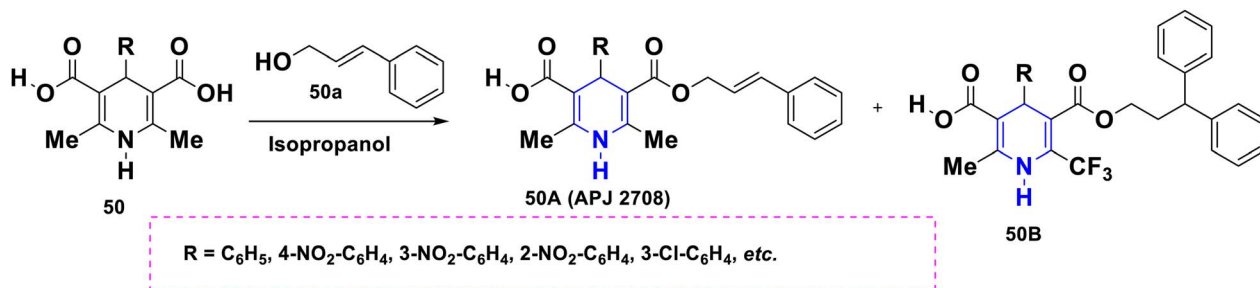
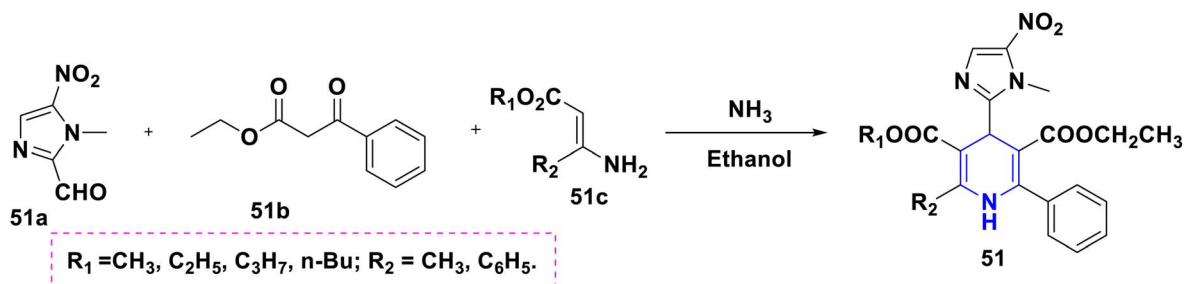


Fig. 4 Structure of 1,4-DHP based channel blockers.



Scheme 51 1,4-DHPs act as potential calcium channel blockers.

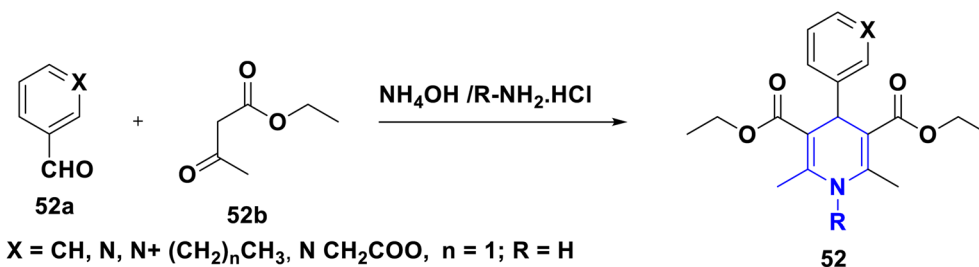


Scheme 52 The nitroimidazolyl-1,4-DHPs as potential calcium channel blockers.

trifluoromethyl-1,4-dihydropyridine-3,5-dicarboxylic acid 5-(3,3-diphenylpropyl)ester **50B**, which exhibited almost identical activity against N-type calcium channels but 1600-fold less activity for L-type than cilnidipine, was among them. The relevance of the inhibitory action against N-type channels in the action of neuropathic pain was reemphasized by this finding (Scheme 51).⁸¹ Lipophilic nitroimidazolyl-1,4-DHPs **51** were explored by Miri *et al.* (2006). It claims that a lipophilic aromatic substitution at the C-6 position of the DHP core improves penetration into the organs, but that replacing both methyl groups in the aromatic ring at both the C-2 and C-6 positions of DHP results in negligible calcium antagonistic activity owing to increased steric hindrance. Due to an extended alkyl chain at the C-3 position, the product 5-ethyl-3-*n*-propyl-1,4-dihydro-2-methyl-6-phenyl-4-(1-methyl-5-nitro-2-imidazolyl)-3-pyridine dicarboxylate **51** demonstrated better activity (Scheme 52).⁸² Rucins *et al.* (2014) developed, synthesized, and tested new 1,4-DHP derivatives featuring the cationic pyridine group at the fourth portion and the *N*-propargyl moiety as a substituent at the first position of the 1,4-DHP **52**. The 4-(*N*-dodecyl)

pyridinium group bearing compounds (without the *N*-propargyl group) and (with the *N*-propargyl group) had the strongest calcium antagonistic activities with an IC₅₀ value of almost 5–14 μM against neuroblastoma SH-SY5Y and vascular smooth muscle A₇T₅ cell with an IC₅₀ of 0.6–0.7 μM. This indicated that they primarily targeted L-type calcium channels, and also showed a little free radical scavenging activity overall. 4-(*N*-Dodecyl)pyridinium-moiety containing molecules might act as prototype molecules for cardio-protective or neuro-protective drugs (Scheme 53).⁸³

Based on the interpretation of the findings, the SAR has been constructed and is graphically represented in Fig. 5. The central core of 1,4-DHP is a necessity for carrying out the activity. The methyl group substitutions at the second and sixth positions were found to be the collective core requirement to exert the activity. The activity has been reduced as a result of replacing the methyl group with a trifluoromethyl group and phenyl systems. Therefore, the recommended functional group at the second and sixth positions is the methyl group. The required functional group is the ester functional group at the third and



Scheme 53 1,4-DHPs cationic pyridine moiety act as calcium channel blockers.

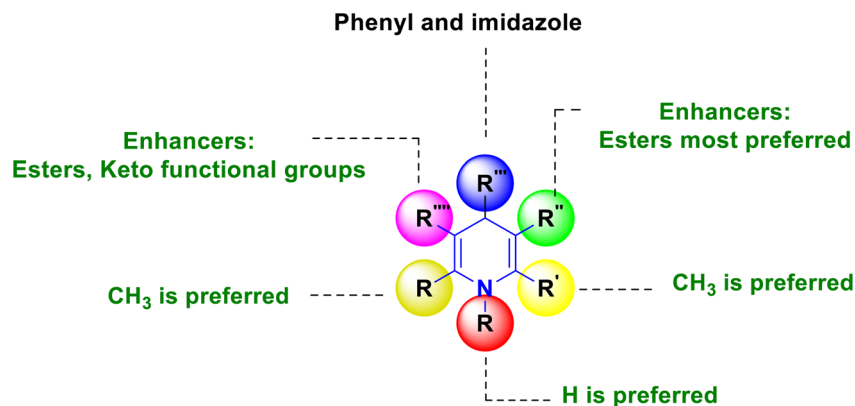


Fig. 5 SAR studies of calcium channel blockers activity of 1,4-DHPs.

fifth positions. The activity has been decreased by using an ester instead of a keto functional group. As a result, the ester functional group is preferred. A considerable impact on the activity has been shown when the phenyl group, a six-membered aromatic system, is favored over imidazole, a five-membered aromatic heterocyclic system, at the fourth position. The relaxant activity of 1,4-DHP was decreased by adding a benzyloxy group to the phenyl ring found in the fourth position. This

implies that the molecules become significantly larger when a large ring is added to the phenyl ring, which may reduce their capacity to block calcium channels. The results for the asymmetrical esters showed that lengthening the substituent in the three carbons of the ester substituent increased activity in the alkyl ester analogues of the new derivatives of nifedipine, in which the *o*-nitrophenyl group at the fourth position is replaced by a 1-methyl-5-nitro-2-imidazolyl substituent and the methyl

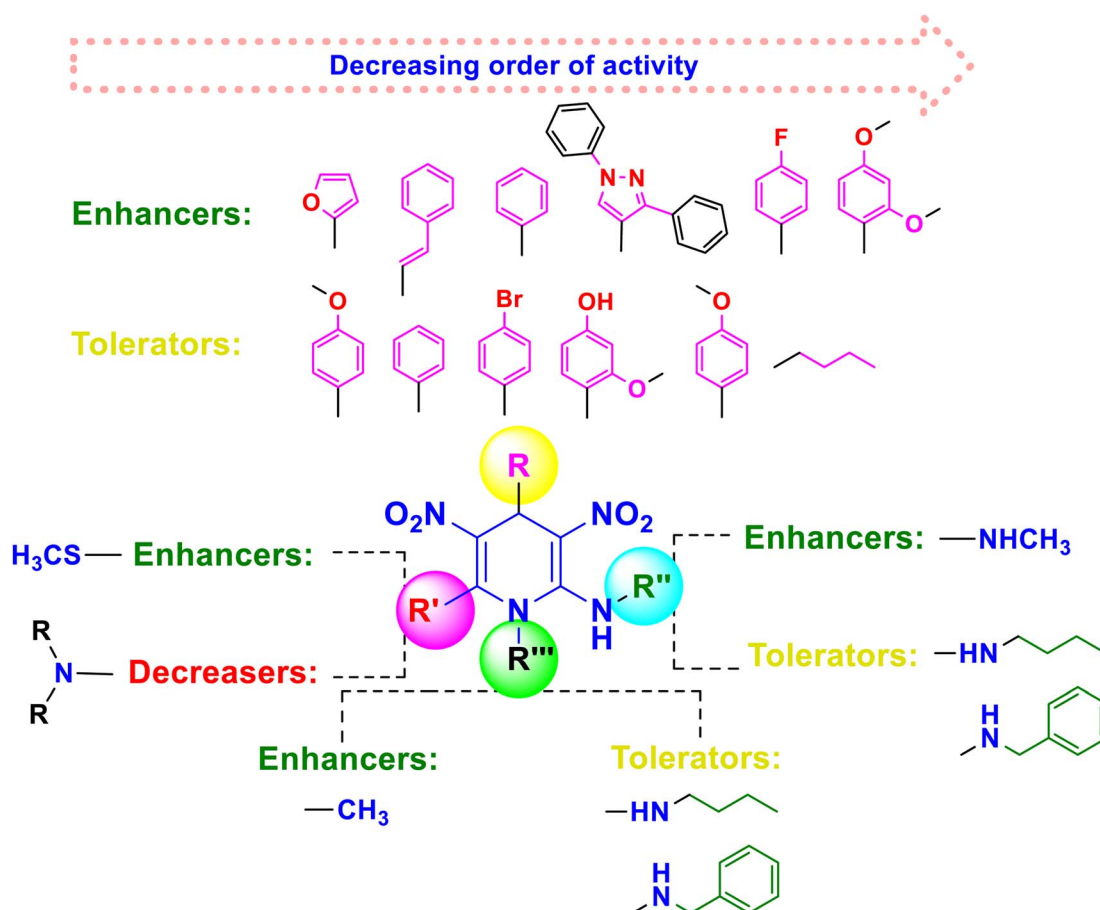
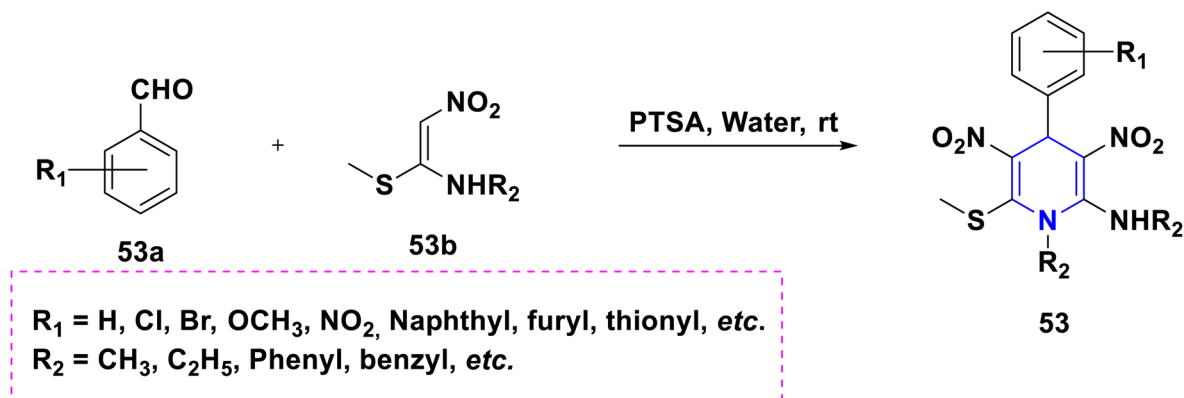


Fig. 6 SAR studies of nitro group containing 1,4-DHPs.

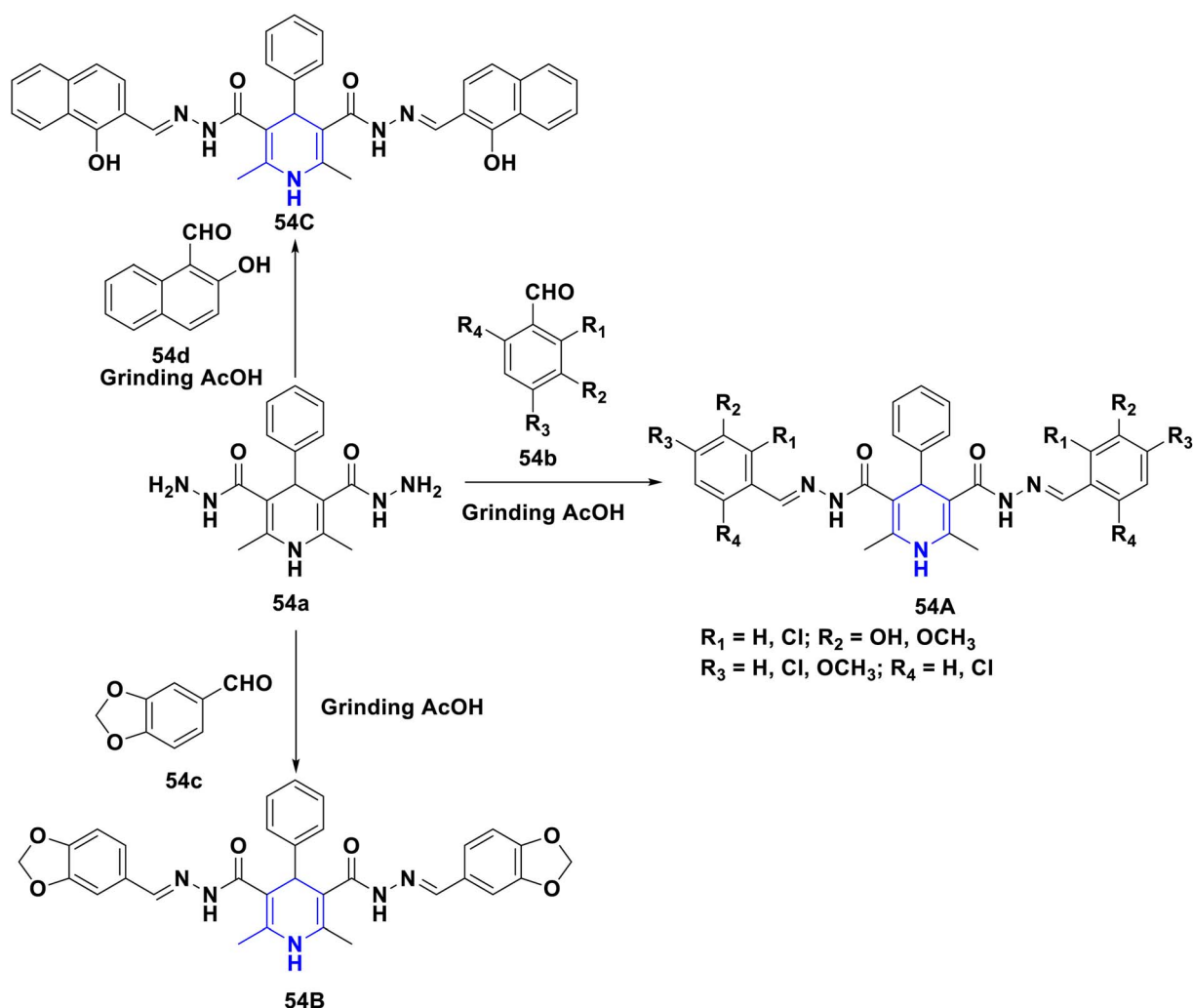


Scheme 54 The nitro substituted 1,4-DHPs as anticancer agents.

group at the sixth position is replaced by a phenyl substituent. The activity dropped when the length increased, and while the steric hindrance also increased.

2.4.3. Anticancer agents. Parthiban *et al.* (2020) designed and synthesized 1,4-DHPs with various structural and

functional group changes. SWISSADME has been used to evaluate the physicochemical characteristics and drug-like molecular nature of the compounds. The PTSA (*para*-toluene sulfonic acid) catalyzed a tandem one-pot synthesis approach from NMSM **53b** and related aldehydes **53a** using a simple,



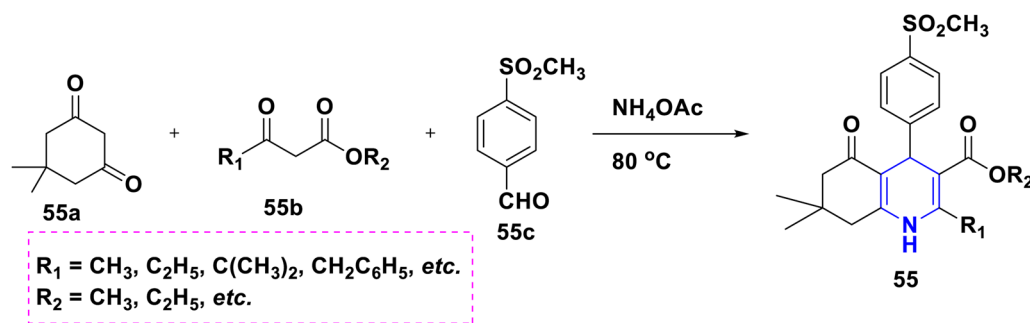
Scheme 55 The 1,4-DHPs-3,5-dicarbohydrazide derivatives as anticancer agents.

inexpensive, and environmentally friendly water-mediated process. All of the substances were tested *in vitro* against lung adenocarcinoma (A549) and laryngeal carcinoma (Hep2) cancer cell lines. The antioxidant capabilities were assessed using the DPPH decrease level (%). According to the SAR studies (Fig. 6), the furan substituted 1,4-DHPs displayed the most potent anticancer activity and was almost equal to doxorubicin, a commonly used cancer medication (Scheme 54).⁸⁴

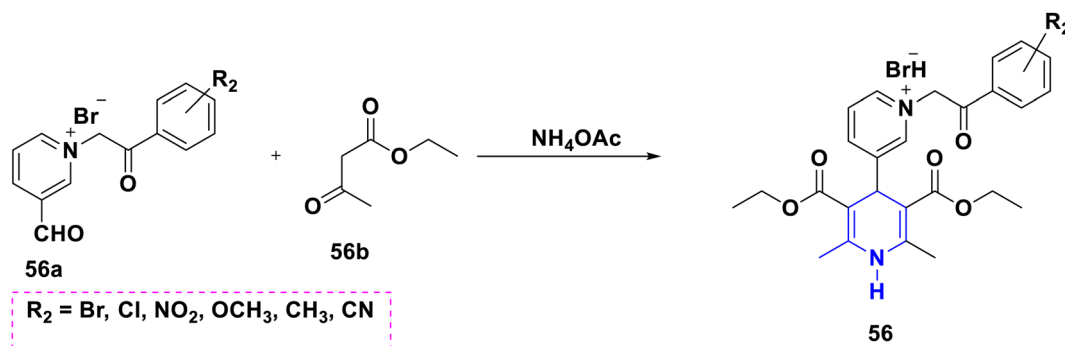
Gomha *et al.* (2020) reported a new series of 2,6-dimethyl-4-phenyl-1,4-dihydropyridine-3,5-dicarbohydrazone derivatives (54A–C) started from 2,6-dimethyl-4-phenyl-1,4-dihydropyridine-3,5-dicarbohydrazide (54a) with various benzaldehydes (54b–d) in the presence of catalytic drops of acetic acid without any catalyst and solvent at ambient temperature, which resulted in moderate to higher yields. The majority of the newly synthesized compounds were screened for anticancer activity against the HepG2 cancer cell line, with several showing good IC₅₀ values. Additionally, using Autodock Vina, *in silico* molecular docking of the synthesized molecules revealed their binding mechanisms inside the active site of DYRK1A (Scheme 55).⁸⁵

Sabakhi *et al.* (2015) conducted research into novel COX-2 inhibitors, a novel series of 1,4-DHPs that were synthesized and studied *in vitro* and *in silico* to understand their biological properties. Novel synthesized compounds at the *para* (*p*) position of the C-4 phenyl ring with a COX-2 SO₂Me pharmacophore, distinct alkoxy carbonyl groups (COOR₂) at the C-3 position and hydrophobic groups (R₁) at the C-2 position of

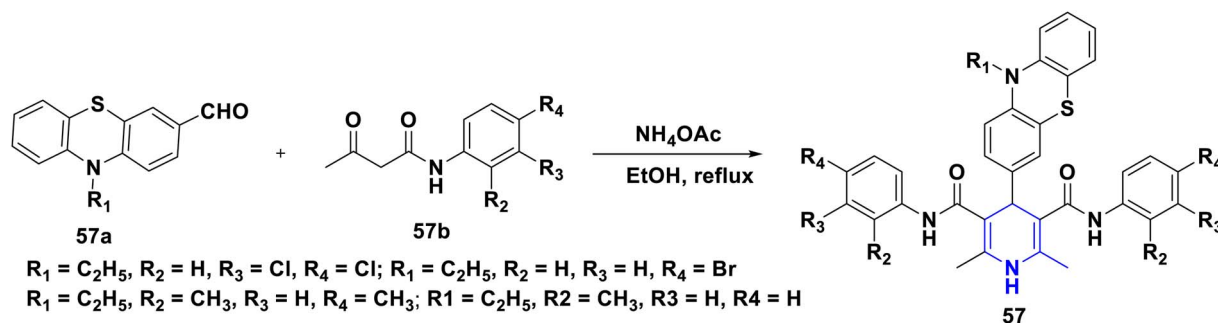
1,4-DHP showed specific inhibition of the COX-2 isozyme. Compound 55 exhibited the most powerful and selective COX-2 inhibitor with an IC₅₀ of 0.30 μM and a COX-2 selectivity index of 92. To establish possible binding models for chemical 55, molecular docking research was conducted. The *p*-SO₂Me-phenyl fragment of 55 was found to be placed into the secondary COX-2 binding sites Phe (518), Arg (513), His (90) and Gly (519) in the research study. The SAR discovery shows that compound 55, ethoxycarbonyl and methyl as R₁ and COOR₂ replacements, has the required molecular geometry as a result of selective inhibition of the COX-2 isozyme, and might be a viable starting point for discovering novel drugs (Scheme 56).⁸⁶ Denish *et al.* (2017) have reported on the 1,4-DHP derivatives having the benzylpyridinium moiety 56, and evaluated its anticancer properties using three human cancer lines: lung cancer A549, colorectal adenocarcinoma Caco-2 and glioblastoma U87MG. Among them, compound 56 had strong anticancer properties. The anticancer activity of the synthesized compounds was similar to that of common pharmaceuticals (carboplatin, gemcitabine, and daunorubicin). As a result, 1,4-DHPs that have been synthesized might be regarded as promising compounds for further drug development as anticancer medicines (Scheme 57).⁸⁷ Sivaramakarthikeyan *et al.* (2019) reported the synthesis of a series of new phenothiazinyldihydropyridine dicarboxamides 57 using a multi-step synthetic method. The molecules containing *o*-bromo/fluoro, *p*-fluoro, *o*/*p*-methyl, and *m/p*-methoxy substituents have anti-inflammatory efficacy that was equivalent to or superior to



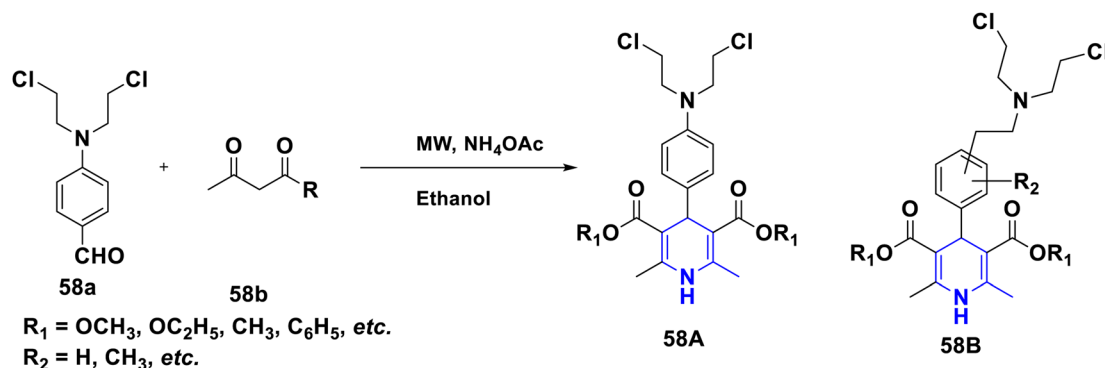
Scheme 56 The sulfoxide 1,4-DHPs as anticancer agents.



Scheme 57 The benzylpyridinium moiety 1,4-DHPs as anticancer agents.



Scheme 58 The phenothiazine-1,4-DHPs-dicarboxamides as anticancer agents.



Scheme 59 The nitrogen mustard pharmacophore hybrid as anticancer agents.

that of diclofenac sodium. Furthermore, as compared to conventional ascorbic acid, the molecules having *o*-bromo, *m*-nitro and fluoro substituents demonstrated higher radical scavenging capabilities. Anticancer research also found that *p* and *m*-chloro-derivatives had the highest anticancer efficacy against all pancreatic cancer cells examined. Molecular dynamics studies were conducted on B-cell lymphoma and B-cell lymphoma, and revealed a promising binding affinity in docking (8.10 kcal mol) (Scheme 58).⁸⁸ Singh *et al.* (2014) developed and synthesized novel nitrogen mustard pharmacophore hybrids such as DHP-M **58A** without a spacer and DHP-L-

M **58B** with an ethyl spacer. They were evaluated *in vitro* for their cytotoxicity properties against the COLO 205 (colon), A 549 (lung), IMR-32 (neuroblastoma) and U 87 (glioblastoma) cancer cell lines by MTT assay with chlorambucil and docetaxel standard drugs. Most of the derivatives tested and exhibited cytotoxic activity that ranged from mild to substantial. DHP-M had the greatest anticancer properties in the four cancer cells studied (**58A**). This might be because **58B** provides less steric impediment (Scheme 59).⁸⁹

The SAR has been constructed, and is visually depicted in Fig. 7 based on the interpretation of the findings. Performing

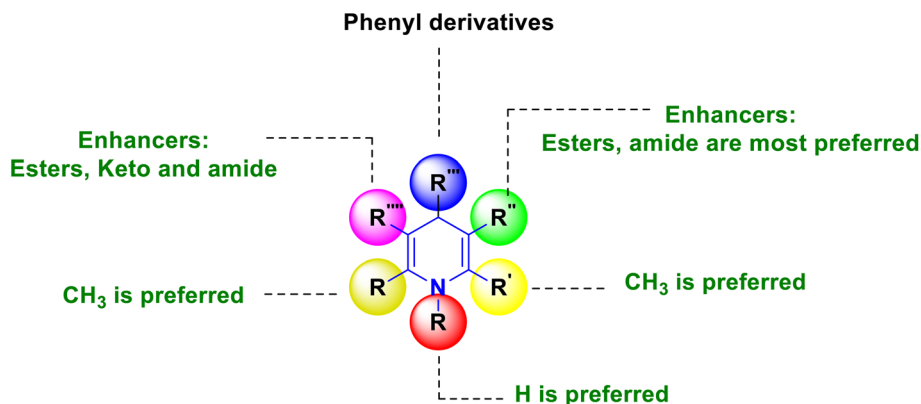
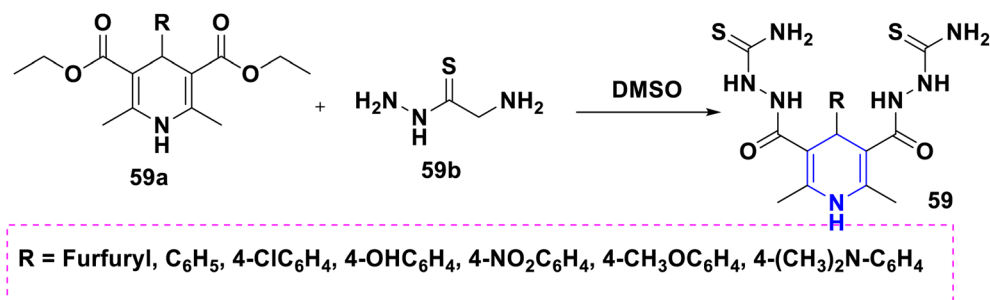
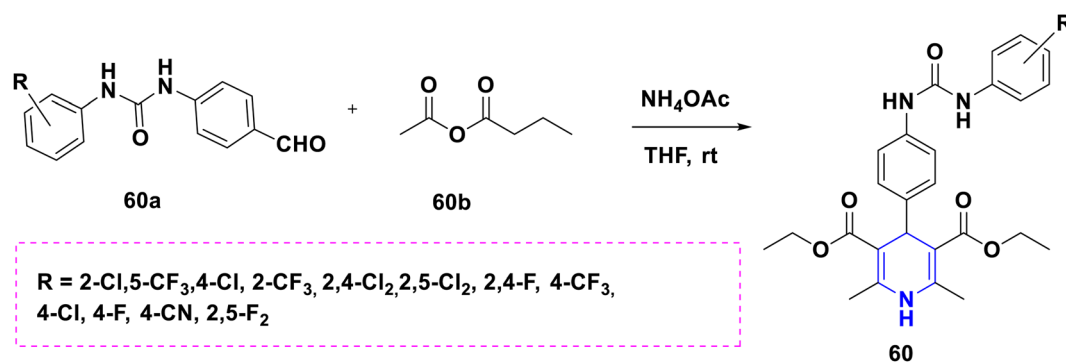


Fig. 7 SAR studies of anti-cancer activity of 1,4-DHPs.



Scheme 60 The dihydrazine carbothioamides 1,4-DHPs as anti-inflammatory agents.



Scheme 61 The 4-(3-aryleureido)phenyl-1,4-DHPs urea derivatives as anti-inflammatory agents.

the activity requires the 1,4-DHP central core. It was discovered that the methyl group substitutions at the second and sixth positions were the essential component needed to exert the activity above the other alkyl systems. Considerable functional group variations, such as amide, ester, and acyl hydrazine, have been studied at the third and fifth positions. More activity was elicited by the dicarbohydrazone and amide functional group than the ester functional group. At the fourth position substituted phenyl derivatives, a six-membered aromatic system showed better activity than the six-membered aromatic heterocyclic system, pyridine. Better anticancer activity was demonstrated by compounds having alkoxycarbonyl groups (COOR₂) in the third position and a SO₂Me and nitrogen

mustard pharmacophore at the *para* position of the phenyl ring, which is in the fourth position.

2.4.4. Anti-inflammatory molecules. By condensation of compounds 59a with thiosemicarbazide 59b, Idhayadhulla *et al.* (2015) created a novel family of dihydrazine carbothioamides 59. The synthesized compounds were evaluated for anti-inflammatory effects on Swiss albino rats (Scheme 60).⁹⁰ Novel derivatives of 4-(3-aryleureido)phenyl-1,4-dihydropyridine 60 were synthesized using a straightforward cascade reaction of oxidation, reduction, as well as nucleophilic addition sequences, in moderate to excellent yields. All of the compounds were subjected to anti-inflammatory ability against pro-inflammatory cytokines, such as tumor necrosis factor-

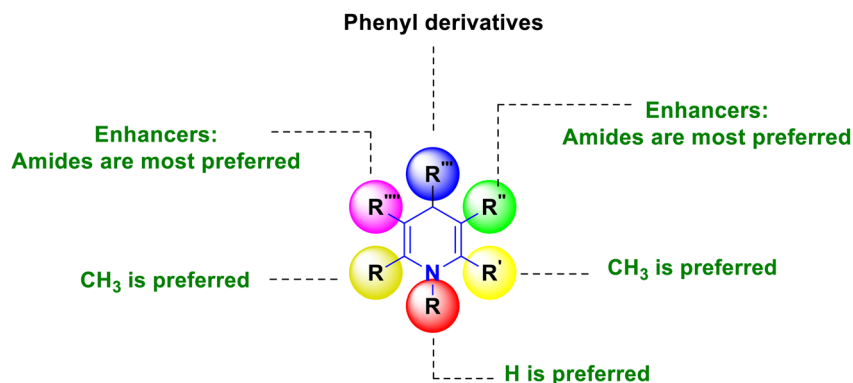
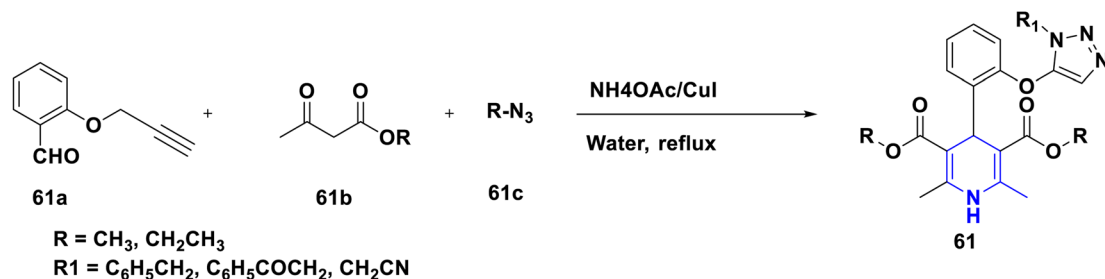


Fig. 8 SAR studies of anti-inflammatory agents of 1,4-DHPs.



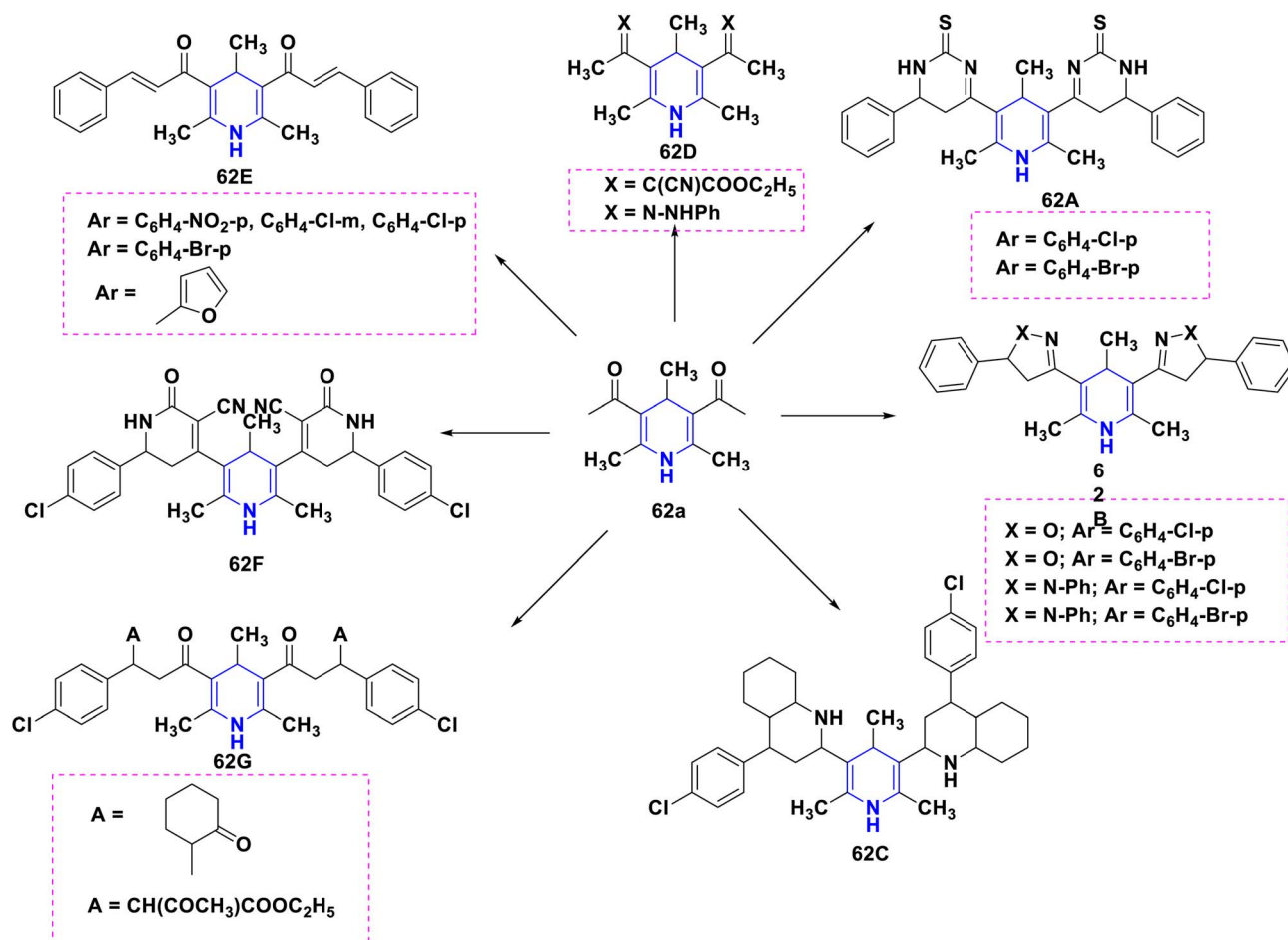
Scheme 62 The 1,2,3-triazoly1,4-DHPs hybrids as anti-microbial agents.

alpha (α), TNF- α , and interleukin-6, (IL-6), and also studied anti-microbial activity (anti-bacterial and anti-fungal). Compound **60** exhibited potential anti-inflammatory action, inhibiting TNF- α and IL-6 by 74–83% and 91–96%, correspondingly, with the standard conventional dexamethasone drug (71% and 86%) at a MIC of $10 \mu\text{M mL}^{-1}$, respectively. This is the major study on the anti-inflammatory efficacy of new urea containing 1,4-DHP compounds, which are important molecular objectives (Scheme 61).⁹¹

The SAR analyses suggest that 1,4-DHP's capacity to reduce inflammation was improved by the addition of dihydrazine

carbothioamides or amide groups at the third and fifth positions, as well as 4-(3-aryleureido) or amide groups at the fourth position of the phenyl rings (Fig. 8).

2.4.5. Anti-microbial agents. Archana *et al.* (2017) described a novel family of 1,2,3-triazoly1,4-dihydropyridine hybrids **61** through the cascade reaction of propargyl salicylaldehyde/*o*-propargyl naphthaldehyde **61a**, β -keto compounds **61b**, and organic azides **61c**. The newly synthesized hybrids were tested *in vitro* against four distinct human infections, including *Proteus mirabilis*, *S. aureus*, *K. pneumonia* and *E. coli*. These outcomes were matched to those of tetracycline,

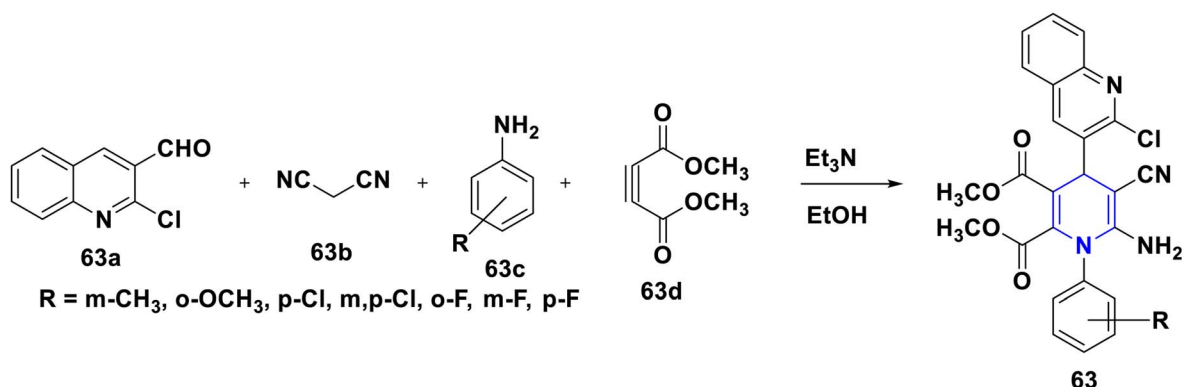


Scheme 63 The 3,5-diacetyl1,4-dihydro-2,4,6-trimethylpyridine as anti-microbial agents.

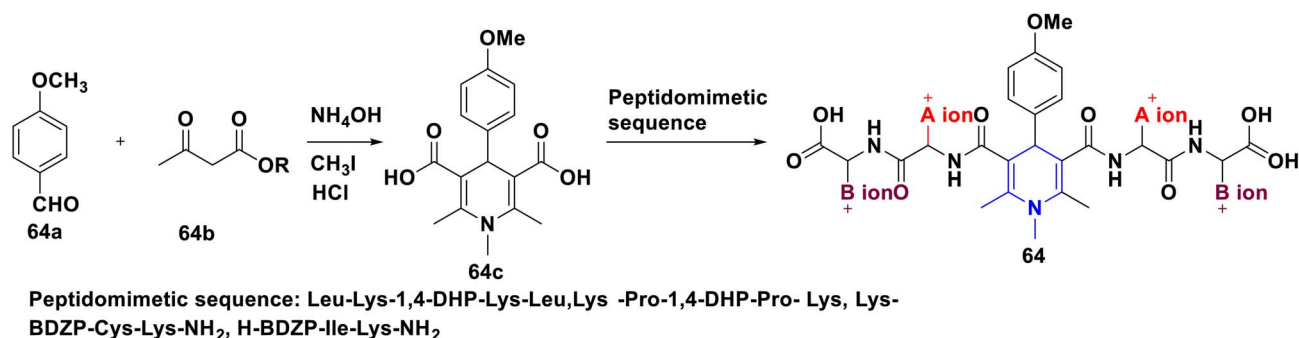
a common antibiotic. These compounds were also tested for their anti-inflammatory effect against bovine serum albumin (BSA). They were then compared to the standard medication diclofenac, and a few hybrids were found to have antibacterial and anti-inflammatory properties that are equivalent to those of reference medicines (Scheme 62).⁹² Melha *et al.* (2013) reported that aromatic aldehydes, thiourea, hydroxylamine hydrochloride, and hydrazine hydrate were condensed with 3,5-diacetyl-1,4-dihydro-2,4,6-trimethylpyridine (**62a**) to generate the corresponding 1,4-DHP derivatives **62A–G**. All of the newly synthesized compounds were subjected to their antimicrobial activity, and compounds **62A–C** were shown to have stronger action against Gram-positive bacteria than Gram-negative bacteria out of all compounds examined (Scheme 63).⁹³

Nkosi *et al.* (2016) developed a new multi-component method for the synthesis of quinoline-bearing DHP derivatives (**63**) involving 2-chloro-3-formyl quinolone **63a**, malononitrile **63b**, arylamines **63c**, and dimethyl acetylenedicarboxylate **63d** by using Et_3N as a base. These molecules were selected for their anticancer activity in an A549 lung cancer cell line, and showed significant anti-proliferative properties. The antibacterial activity of **63** against *Pseudomonas aeruginosa* ATCC 27853, *Escherichia coli* ATCC 25922, and *Staphylococcus aureus* ATCC showed promising activity at the minimum inhibitory concentration (MIC) values of $16\text{ }\mu\text{g mL}^{-1}$ against *Staphylococcus aureus* and $32\text{ }\mu\text{g mL}^{-1}$ against *Pseudomonas aeruginosa* and *Escherichia coli* (Scheme 64).⁹⁴ Lapidot

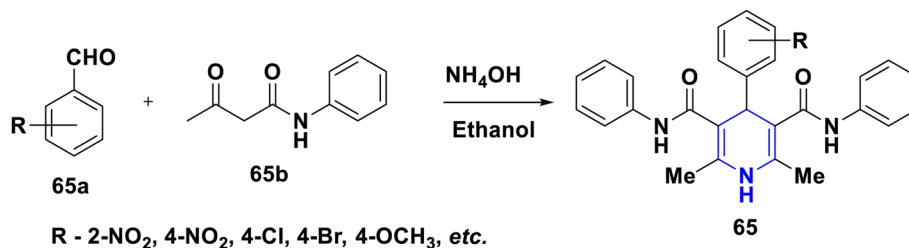
et al. (2015) developed a novel 1,4-DHP **64** as anti-microbial broad-spectrum cationic peptidomimetics. This method involves two steps: the first one is the Hantzsch reaction, which is then followed by the coupling reaction for the synthesis of 1,4-DHP skeleton involving two dipeptides attaching cationic side chains. The synthesized peptidomimetics tested its anti-bacterial activity, and showed noteworthy MICs in the range of $35\text{--}100\text{ }\mu\text{g mL}^{-1}$ against Gram-(+) and Gram-(−) bacterial pathogens (Scheme 65).⁹⁵ Narsinghani *et al.* (2017) synthesized the 1,4-DHP derivative **65**, and evaluated their anti-bacterial and anti-fungal properties. They revealed that a bromo group located at the *para* position of the phenyl ring and a chloro atom attached to the *para* position of the carbamoyl phenyl ring showed excellent anti-bacterial activities against Gram-negative strains (81.76% and 75.94%, respectively). The hydroxy group at the second position and the fifth position of the bromo group attached to the phenyl moiety of the 1,4-DHP compounds showed good antifungal properties (86.85%). If the chloro group was attached to the *para* position of the carbamoyl phenyl ring, these molecules exhibited significant antibacterial, as well as antifungal activities. Furthermore, this activity needed electron-withdrawing groups such as fluoro or chloro at the third and fifth positions of the DHP ring (Scheme 66).⁹⁶ Gomha *et al.* (2020) reported the reaction of (2,6-dimethyl-4-phenyl-1,4-dihydropyridine-3,5-diyl)-bis(4-amino-4*H*-1,2,4-triazole-3-thiol) (**66a**) with several aldehydes in the presence of bromine/acetic acid towards the synthesis of 1,4-DHP hybrid with 1,2,4-



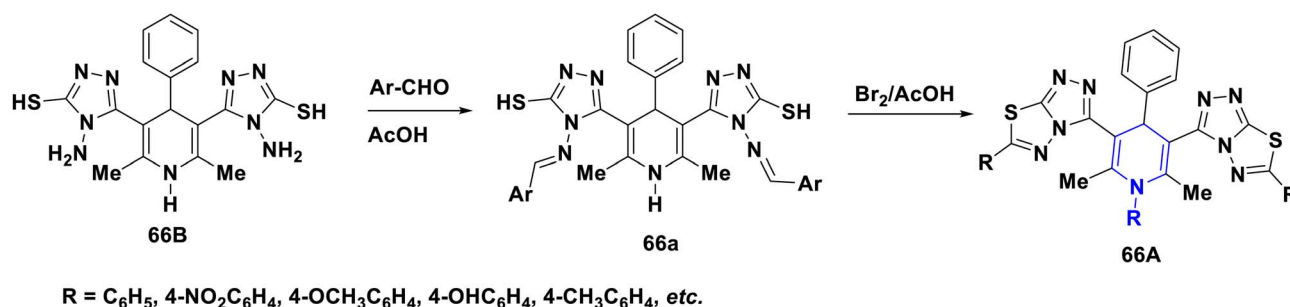
Scheme 64 The quinoline bearing 1,4-DHP derivatives as anti-microbial agents.



Scheme 65 The cationic peptidomimetics centred on a hydrophobic 1,4-DHP as anti-microbial agents.



Scheme 66 The carbamoyl 1,4-DHPs as anti-microbial agents.



Scheme 67 The triazole 1,4-DHPs as anti-microbial agents.

triazole **66A** and 1,4-dihydropyridine-bis-triazolothiadiazoles hybrid derivatives (**66B**). The antimicrobial activity of the prepared hybrids was screened and showed excellent activity compared to standards like ampicillin, gentamicin and amphotericin B through the agar diffusion well method (Scheme 67).⁹⁷

Based on the interpretation of the results, the SAR has been created and is represented visually in Fig. 9. The 1,4-DHP central core is necessary for carrying out the activity. It was found that the methyl group substitutions at the second and sixth positions were crucial for exerting the activity that was superior to other alkyl systems. Numerous functional group changes have been researched at the third and fifth positions, including ester, amide or dipeptides, anhydride, cyano, keto, and five-membered heterocyclics such as triazole-3-thiol. Ester and keto functional groups produced more activity than amide, anhydride, cyano, and five-membered heterocyclics. Therefore, the activity has been improved by the ester and keto functional

groups. Six-membered aromatic systems with electron-withdrawing group such as fluoro or chloro at the third and fifth positions, 1,2,3-triazolyl of phenyl ring, 2-chloro quinoline and alkyl derivatives, have demonstrated improved activity at the fourth position.

2.4.6. Anti-oxidant activity. Sun *et al.* (2012) reported a reaction of dimethyl or diethyl 2,6-dimethyl-4-phenyl-1,4-DHP-3,5-dicarboxylate **67a**. Various chalcone **67b** afforded a novel chalcone substituted at the third position of 1,4-DHPs **67**. The synthesized molecules were also studied for their radical scavenging activities. Among them, thienyl substituted 1,4-DHPs have shown promising anti-oxidant activities (Scheme 68).⁹⁸ Kruk *et al.* (2020) reported the chemiluminescence properties of 4-flavonil-1,4-DHP derivatives **68** by spectrophotometry methods, which showed better light emission from the structure. From this study, all observed luminescence molecules showed superoxide dismutase activity due to the transformation of superoxide radicals (Scheme 69).⁹⁹ Dhinakaran

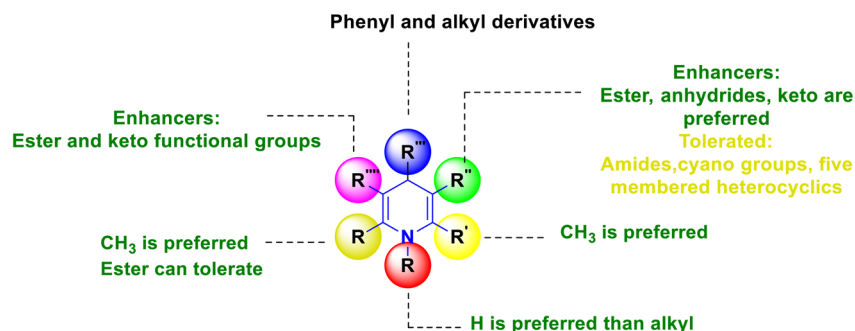
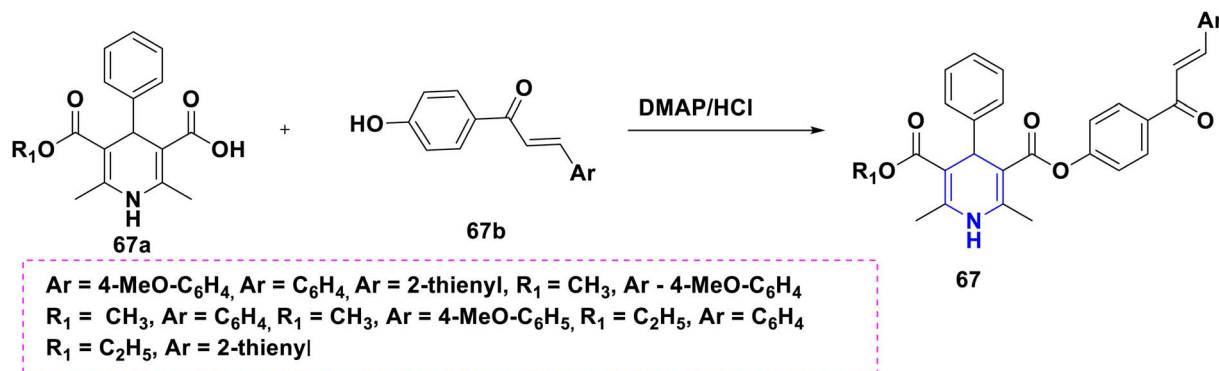
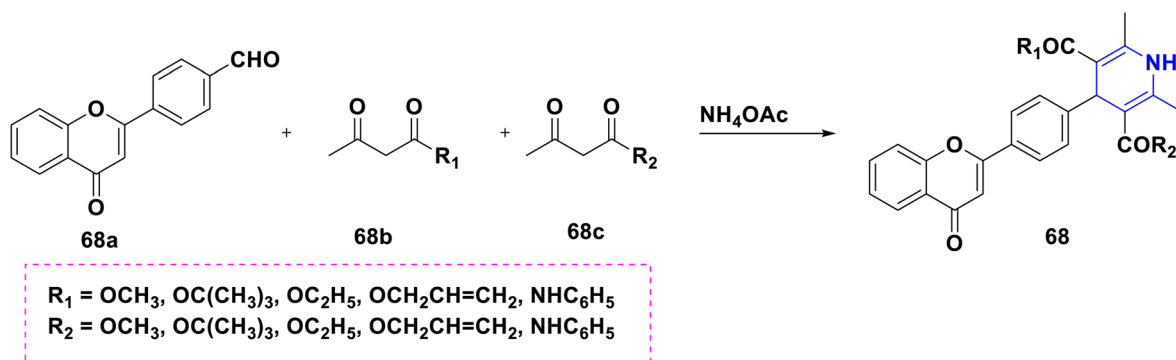


Fig. 9 SAR studies of anti-microbial agents of 1,4-DHPs.



Scheme 68 The chalcone-substituted 1,4-DHPs as anti-oxidant agents.

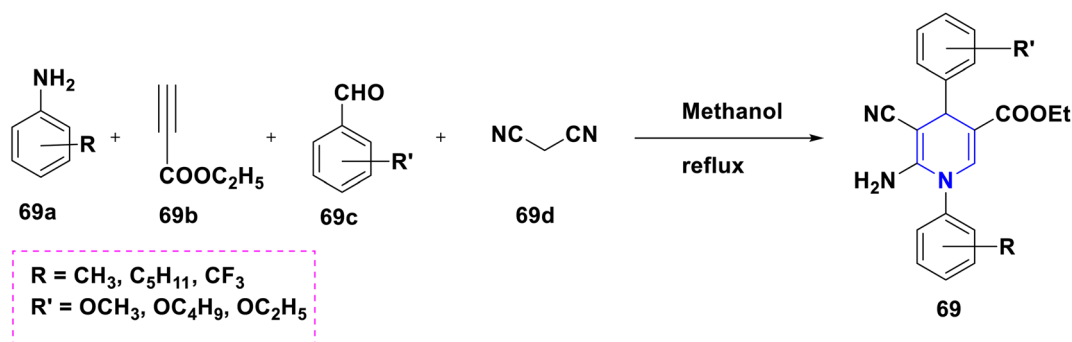


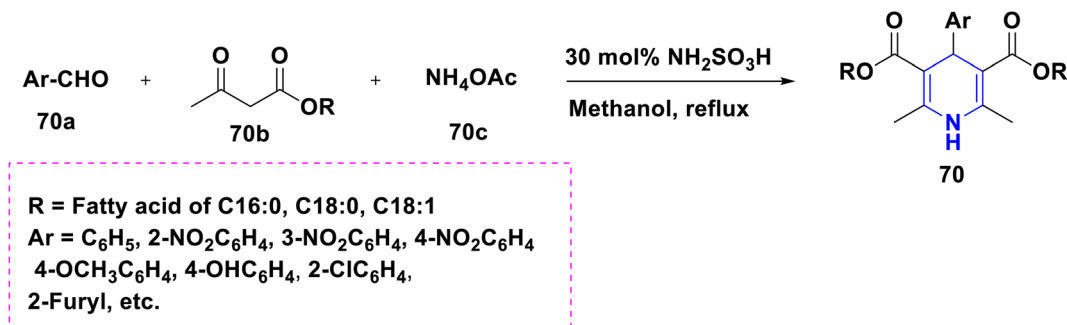
Scheme 69 The flavonil-1,4-DHPs as anti-oxidant agents.

et al. (2015) reported a cascade method towards the synthesis of *N*-aryl, 1,4-DHPs **69**. They evaluated their cytotoxicity properties using the A549 cell line and antioxidant activities by DPPH methods, and also found better activities (Scheme 70).¹⁰⁰ Cabrera *et al.* (2019) developed the Hantzsch cascade reactions for the synthesis of 1,4-DHPs **70**, starting with various aldehydes **70a**, β-ketoester **70b** and ammonium acetate **70c** using a reusable sulfamic acid catalyst. These 1,4-DHP derivatives were studied for their antioxidant activity by using two substituents groups. The results revealed that the nitro group and long chain fatty derivatives embedded with 1,4-DHP scaffolds showed

better antioxidant properties, and were almost similar to the standard drug vitamin E (Scheme 71).¹⁰¹

Based on the interpretation of the results, the SAR has been created and is represented visually in Fig. 10. The 1,4-DHP central core is necessary for carrying out the activity. It was found that the methyl group substitutions at the second and sixth positions were crucial for exerting the activity that was superior to other alkyl systems. The ester functional group has enhanced activity compared to the cyano functional group at the third position. The chalcone and long chain fatty derivatives embedded at the third and fifth positions show enhanced activity. Phenyl, six-membered aromatic systems have

Scheme 70 The *N*-aryl-1,4-DHPs as anti-oxidant agents.



Scheme 71 The long-chain fatty DHPs as anti-oxident agents.

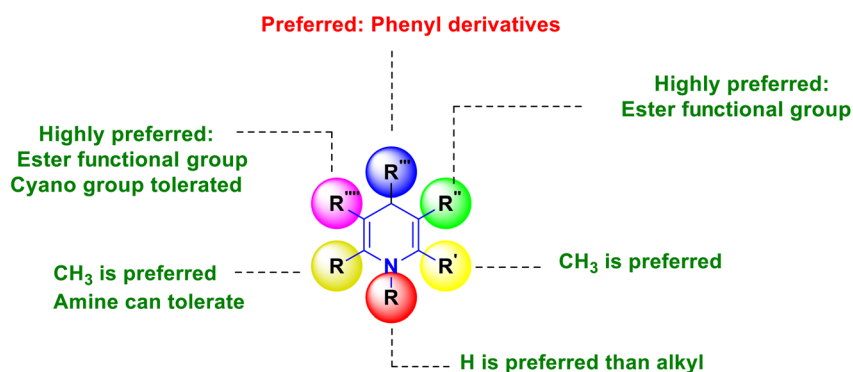
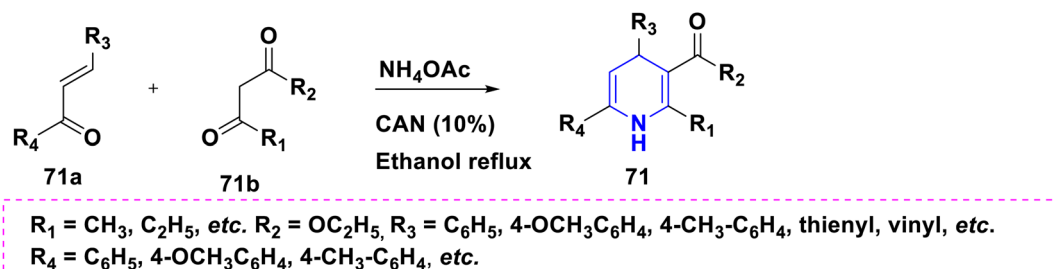


Fig. 10 SAR studies of the antioxidant activity of 1,4-DHPs.

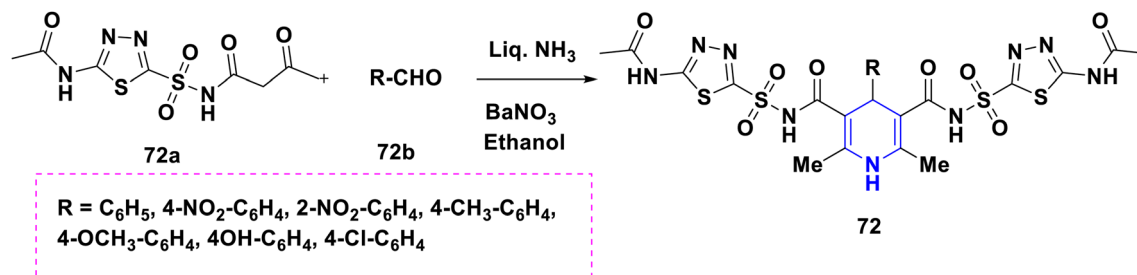
demonstrated improved activity at the fourth position. The 4-flavonil or flavonoids-substituted phenyl system at the fourth position have increased the activity.

2.4.7. Miscellaneous activities. Tenti *et al.* (2013) discovered the ceric ammonium nitrate-catalyzed one-pot three-component reaction from various α,β -unsaturated ketones **71a**, β -dicarbonyl compounds **71b** and ammonium acetate towards the synthesis of 4,6-diaryl-1,4-DHPs **71**. These compounds were evaluated for their neuroprotective activity. Among them, 1,4-DHP having a 2-thienyl substituent exhibited reasonable neuroprotective properties (Scheme 72).¹⁰² Babu *et al.* (2019) reported the barium nitrate catalyzed reaction of thiadiazolsulfonylamide **72a**, different aldehydes **72b** using liquid ammonia medium for the preparation of novel thiadiazolsulfonylamide 1,4-DHPs **72**, and estimated the

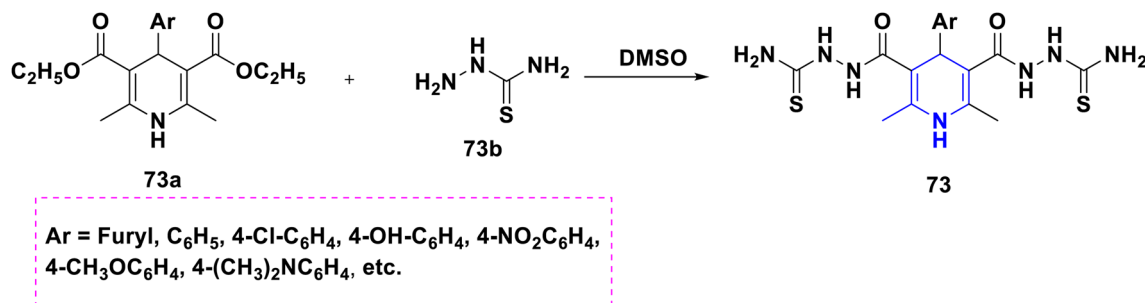
anticholinesterase properties by the Ellman method. All of the molecules showed potent inhibition action when compared with the present drug, donepezil HCl, against acetylcholinesterase and butyl cholinesterase enzymes. This outcome indicates the significance of thiadiazolsulfonylamide compounds for anticholinesterase properties, and showed their efficacy and reliability (Scheme 73).¹⁰³ Kumar *et al.* (2011) described the condensation reaction of 1,4-DHP ester **73a** with thiosemicarbazide **73b** towards the synthesis of thiosemicarbazide 1,4-DHPs **73** and studied their anticoagulant properties (Scheme 74).¹⁰⁴ Ahamed *et al.* (2018) developed an ultrasound irradiation method for the synthesis of a novel series of 1,4-DHPs (**74A–C**) with better yields. The newly synthesized derivatives have been studied for both antimicrobial and anticoagulant properties. Compound **74A** (MIC: 0.25 $\mu\text{g mL}^{-1}$) shows



Scheme 72 The 4,6-diaryl-1,4-DHPs as neuroprotective agents.



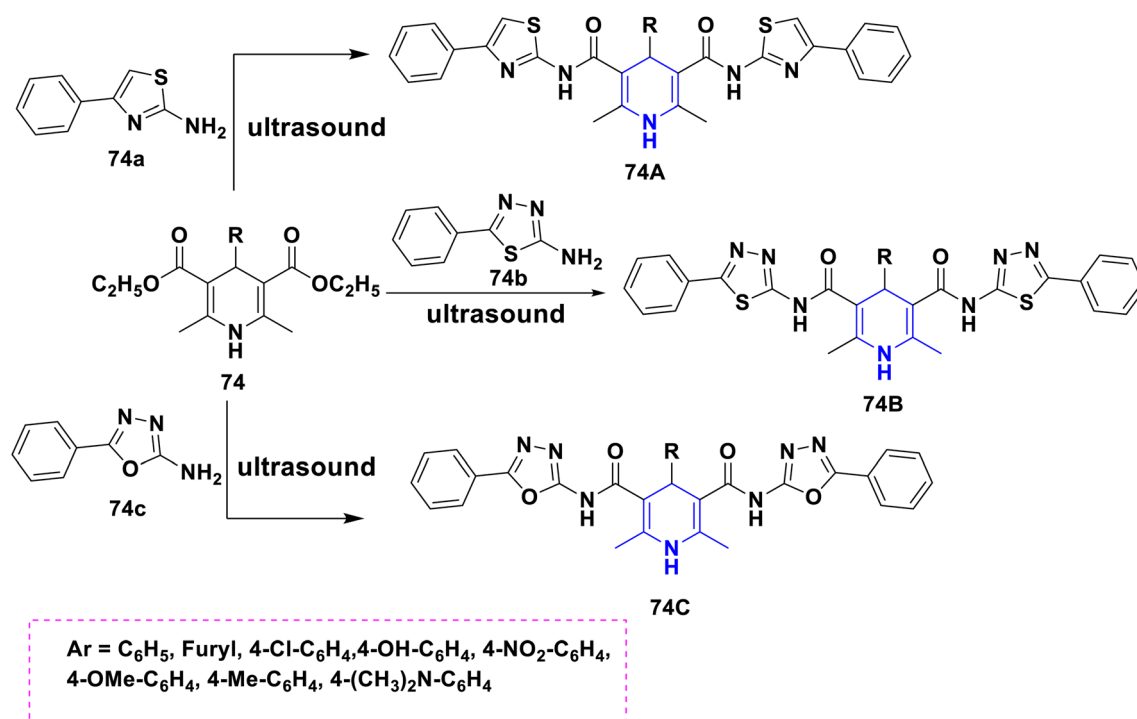
Scheme 73 The thiadiazol sulfonfyl anamide 1,4-DHPs as anticholinesterase agents.



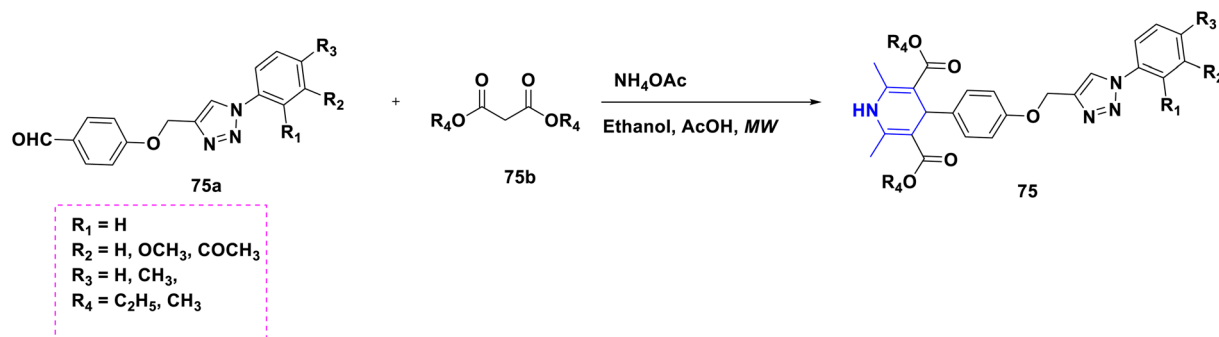
Scheme 74 The thiosemicarbazide 1,4-DHPs as anticoagulant agents.

potential active against *Escherichia coli*, and compound **74B** (MIC: 0.5 $\mu\text{g mL}^{-1}$) also exhibited promising activity against *Pseudomonas aeruginosa* equal to that of standard ciprofloxacin (MIC: 1 $\mu\text{g mL}^{-1}$). The antifungal property of **74C** (MIC: 0.5 $\mu\text{g mL}^{-1}$) was more active than that for the clotrimazole (MIC: 1 $\mu\text{g mL}^{-1}$)

standard against *Candida albicans*. The anticoagulant activity was determined. Compound **74B** exhibited good potential in anticoagulant activity, which was almost comparable with the standard of heparin by activated partial thromboplastin time (APTT) and prothrombin time (PT) coagulation



Scheme 75 The phenylthiadiazol-2-amine 1,4-DHPs as anticoagulant agents.



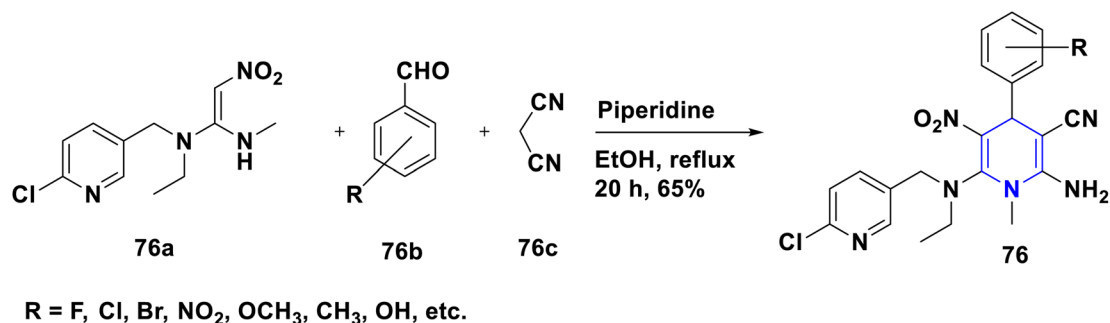
Scheme 76 The triazol 1,4-DHP derivatives as antidiabetic agents.

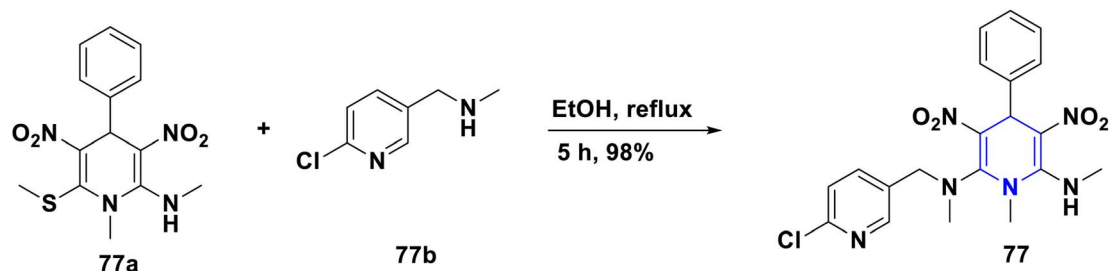
assays. The prepared molecule **74C** was screened for its anti-cancer properties using HeLa (cervical), HepG2 (liver) and MCF-7 (breast) cancer cell lines, and displayed greater toxicities ($\text{GI}_{50} = 0.02 \text{ lm}$). The compound **74B** ($\text{GI}_{50} = 0.03 \text{ lm}$) also found equipotent activity against the MCF-7 cell line. Henceforth, the molecules (**74A–C**) might be potent compounds for the development of novel groups of antimicrobial and anticoagulant drugs (Scheme 75).¹⁰⁵ E. Preveen Kumar *et al.* (2019) described the MW mediated one-pot reaction using triazole derivatives **75a**, β -keto ester **75b** and ammonium acetate towards the synthesis of triazole 1,4-DHP derivatives **75** and assessed their anti-diabetic properties. Comparing all derivatives, compound **75** exhibited effective anti-diabetic properties. Hit compounds were additionally confirmed using *in vitro* enzymatic studies by analysis of its 11- β -hydroxysteroid dehydrogenase-1 (11- β -HSD1) inhibitory property to explain the mechanism of action of these compounds. The outcomes indicated that the 11- β -HSD1 inhibitory property of the molecules was stable and efficient. Molecular docking studies revealed that compound **75** showed a stable binding pattern (-9.758) with human 11- β -HSD1 (Scheme 76).¹⁰⁶

2.5. Agrochemical applications

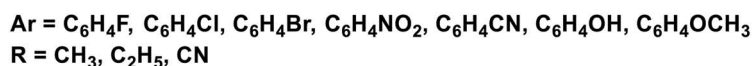
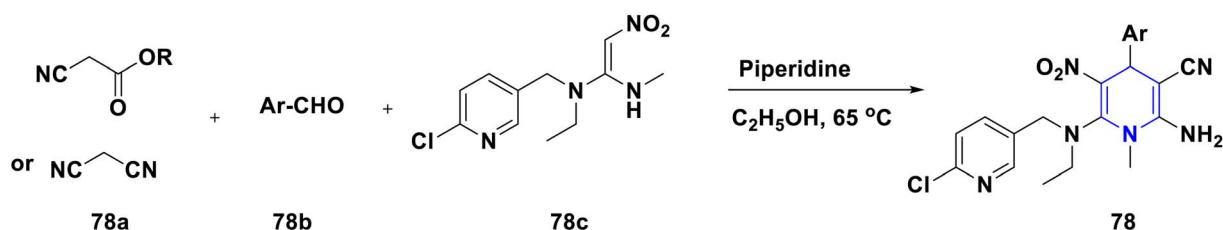
2.5.1. Insecticidal activity. Sun *et al.* (2012) synthesized 1,4-DHP and *cis*-nitenpyram molecular hybrids **76**, and examined the impact of different functional group variations on the insecticidal activity. The functional group variations include the nitro group at the fifth position, CN or ester groups at the third

position, various substituted aryl systems such as F, Cl, Br, and OCH_3 at the fourth position and *cis*-nitenpyram **76a** at the sixth position. The synthesis uses *cis*-nitenpyram **76a**, where the secondary amine and nitro group are sealed in a *cis*-configuration, and aldehydes **76b** and malononitrile **76c** serve as reactants in the piperidine-catalyzed, one-pot synthesis. The insecticidal activity of *cis*-configured 1,4-DHPs against the most frequent pest, *Aphis medicaginis*, was enhanced (Scheme 77).¹⁰⁷ In a related study, Rao and Parthiban (2014) successfully synthesized the *cis*-nitenpyram neonicotinoid insecticide analogue **77** by substituting 1-(6-chloropyridin-3-yl)-*N*-methylmethanamine **77b** on **77a** with good yield. The NO_2 group at the neonicotinoid's fifth position is crucial for enhancing the insecticidal efficacy, according to the literature. In this work, NO_2 groups were linked to 1,4-DHPs at the third and fifth locations in the hopes that this would increase their insecticidal activity.⁶³ The effect of temperature on the product is another intriguing discovery of this study. The ArCH_2NMe and NO_2 groups are in steric hindrance at room temperature, and tend to rotate slowly around the C2–N bond. When the temperature was increased to 80°C , the spectrum became clear with attaining a particular orientation without steric hindrance (Scheme 78). The synthesis and insecticidal activities of 1,4-DHPs and *cis*-nitenpyram molecular hybrids have been carried out by Liu *et al.* (2012). The MW reactor-assisted, piperidine-catalyzed one pot MCR synthesis was achieved by using methyl cyanoacetate **78a**, benzaldehyde, **78b** and nitenpyram, **78c**. The compound **78** has exhibited insecticidal activity at 500 mg L^{-1} against *Aphis*

Scheme 77 *cis*-Nitenpyram analogues containing 1,4-DHPs.



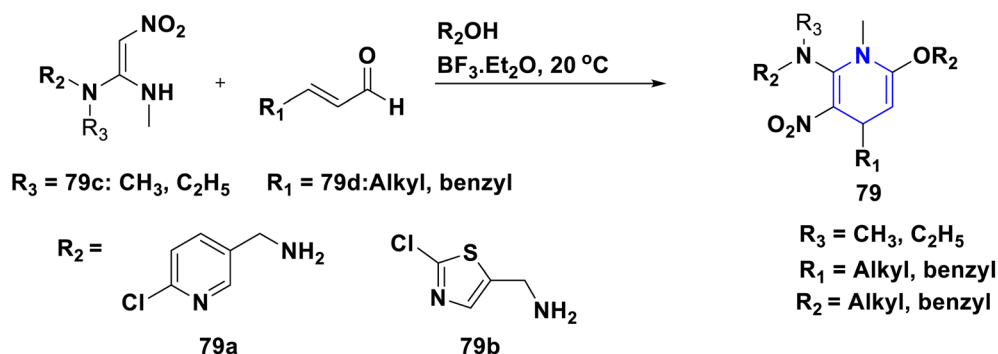
Scheme 78 Synthesis of neonicotinoid insecticide analogue.

Scheme 79 Synthesis of 1,4-DHPs and *cis*-nitenpyram hybrid analogues.

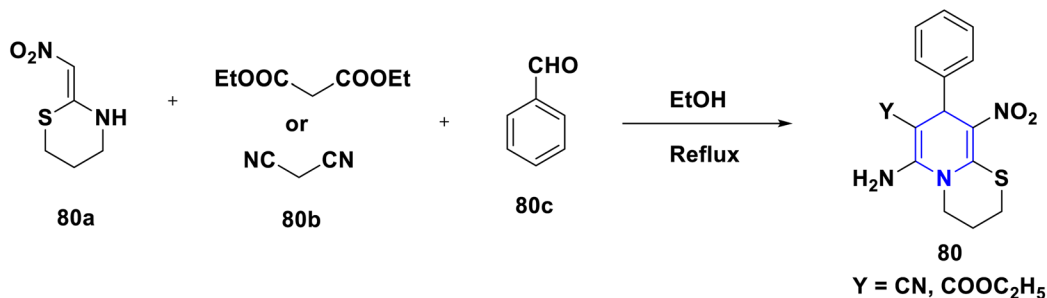
craccivora (Scheme 79).¹⁰⁸ In addition, Lu *et al.* (2012) explored the molecular hybrids of 1,4-DHPs and nitenpyram, and the molecular hybrids of 1,4-DHPs and clothianidin. Among these two, clothianidin nitromethylene neonicotinoids showed good insecticidal properties against cowpea aphid (*Aphis craccivora*) and armyworm (*Pseudaletia separata* 'Walker') (Scheme 80).¹⁰⁹

A novel series of nithiazine-containing 1,4-DHP analogues **80** was designed and synthesized by Tian *et al.* (2014). The MCR reaction involves nithiazine **80a**, malononitrile or ethyl cyanoacetate **80b**, and benzaldehyde **80c** as reactants. The synthesized compounds have been screened for insecticidal activity, and showed moderate activity against *Aphis craccivora*. When compared with the ester group, the CN at the fifth position of 1,4-DHP had greater insecticidal activity owing to the stronger electron-withdrawing nature of the substituents (Scheme 81).¹¹⁰ The works of Bakhite *et al.* (2014) designed,

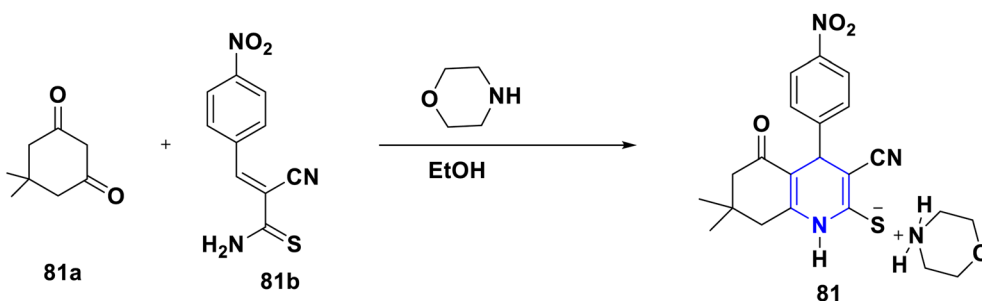
synthesized and identified the insecticidal activities of novel piperidinium salts of novel 1,4-DHPs **81**. In their bioassay studies, the compounds showed moderate to strong aphidicidal activities. Interestingly, one of the compounds **81** showed 4-fold higher insecticidal activity than acetamiprid because of the nitro group present at the fourth position aryl of the 1,4-DHP (Scheme 82).¹¹¹ Yang *et al.* (2018) created a new set of 1,4-DHPs **82A–C**, and tested their insecticidal effectiveness against *M. persicae*, *B. brassicae*, *T. cinnabarinus*, *B. cinerea*, *M. oryzae*, *S. sclerotiorum*, and *F. oxysporum*. The LC_{50} values were 0.011, 0.0015, and 0.0007 μM . The most active compound, **82B**, showed greater insecticidal efficacy against *B. brassicae*, *M. persicae*, and *T. cinnabarinus*, respectively. At 50 mg L^{-1} , compound **82A** inhibited *F. oxysporum* and *S. sclerotiorum* by 45% and 65.83%, respectively. They determined through SAR experiments that functional group differences in 1,4-DHP have



Scheme 80 Synthesis and insecticidal activities.



Scheme 81 Synthesis of insecticides.

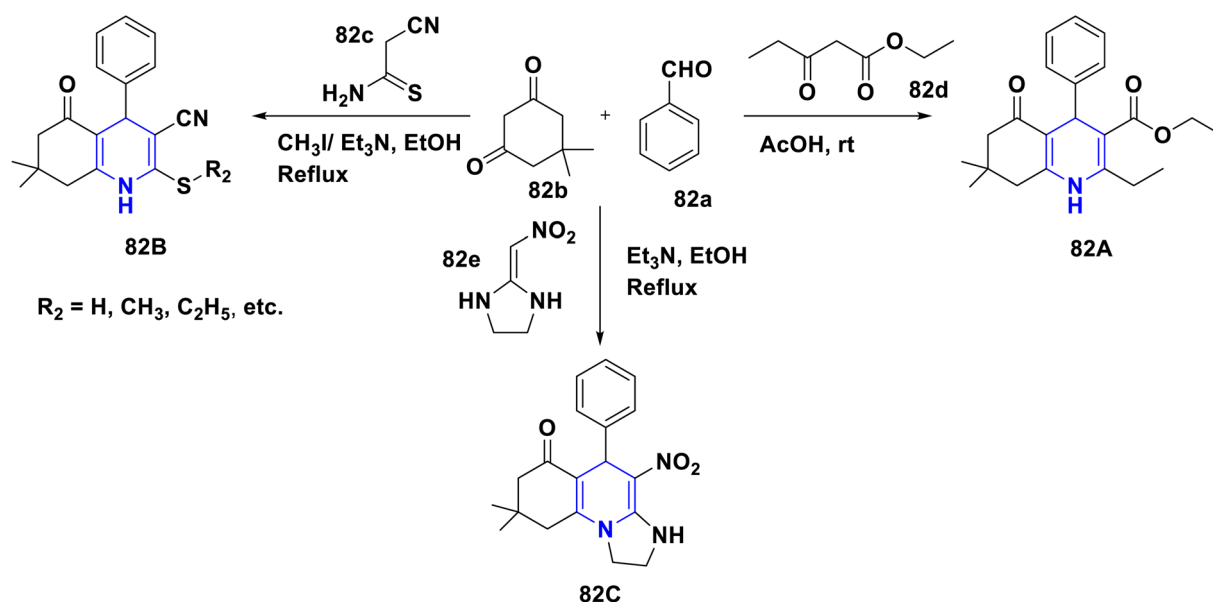


Scheme 82 Synthesis of insecticides.

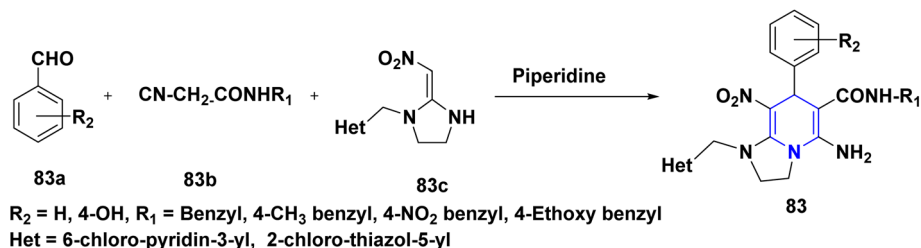
a significant influence on insecticidal activity (Scheme 83).¹¹² He *et al.* (2013) developed the synthesis of neonicotinoid and 1,4-DHP hybrids **83**, as well as their insecticidal and antibacterial activities. At 500 mg L⁻¹, compound **83** has insecticidal activity against *N. lugens*, and at 200 mg L⁻¹, they have antibacterial activity against *Pseudomonas solanacearum* (Scheme 84).¹¹³

The enhanced π - π stacking-based neonicotinoid 1,4-DHPs **84** have been designed and synthesized by Zhang *et al.* (2010).

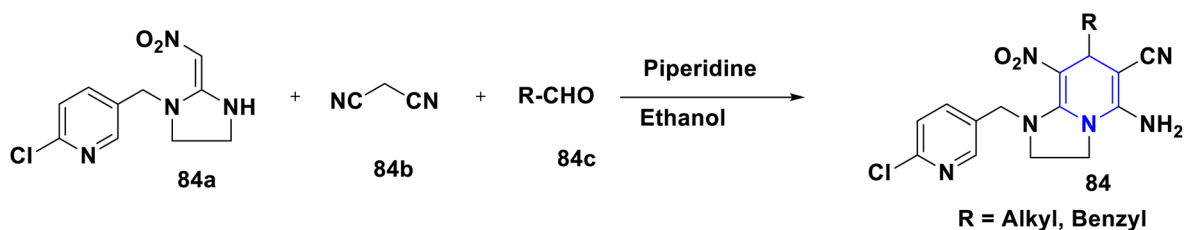
From the biological assays, these compounds were found to possess significant insecticidal properties, especially against pea aphid, *Aphis craccivora* (Scheme 85).¹¹⁴ In their 2015 study, Tian *et al.* (2015) described the design and multicomponent reactions of nitenpyram **85a**, triethoxymethane **85b**, and 4,4,4-trifluoro-3-oxobutaneanilides **85c** under solvent-free and catalyst-free conditions that produced a series of novel neonicotinoids with a dihydro-pyridine ring and a trifluoromethyl



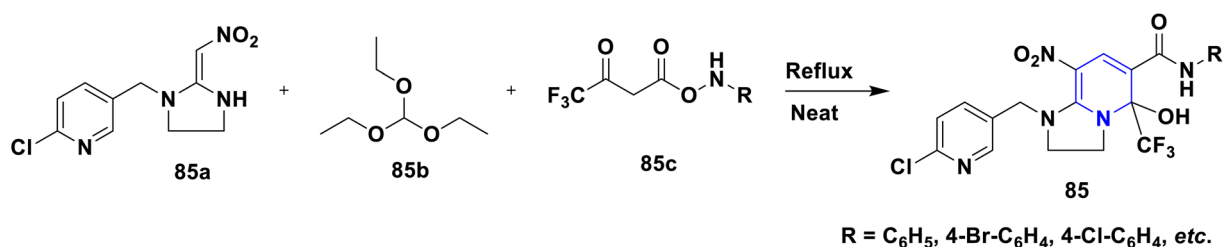
Scheme 83 Synthesis of insecticides.



Scheme 84 Synthesis of insecticidal neonicotinoid and 1,4-DHPs hybrids.



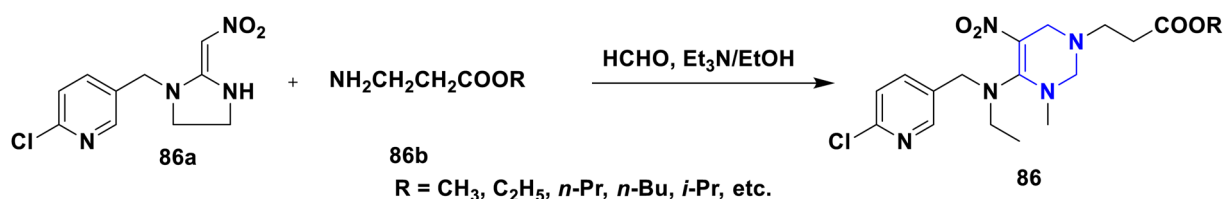
Scheme 85 Synthesis of insecticides.



Scheme 86 Synthesis of neonicotinoids insecticides with trifluoromethyl group.

group 85 (Scheme 86). As an electron-withdrawing group, the trifluoromethyl group at the second position of 1,4-DHPs was crucial in increasing the reaction activity of the molecule. According to bioassays, the majority of the synthetic compounds had mild insecticidal effects on *Aphis craccivora*.¹¹⁵ According to Sun *et al.* (2011) nitenpyram **86a** was modified by adding various straight chain amino acid alkyl esters **86b** to create several unique nitenpyram analogues **86**. By varying the size of the ester group and the length of the flexible ester arm, the analogues displayed diverse insecticide activity in a more manageable and sensible manner (Scheme 87). The test analogues all showed effective insecticidal action at 100 mg L⁻¹. However, analogues **4c** and **4d** had the best *in vitro* activity and 100% mortality at 20 mg L⁻¹.¹¹⁶

Based on the interpretation of the results, the SAR has been created and is represented visually in Fig. 11. The 1,4-DHP central core is necessary for carrying out the activity. The replacement of hydrogen at the first position –NH– with a methyl group has shown better activity, but extending the alkyl chain beyond the propyl system reduced the activity. It was found that the amine and trifluoromethyl group enhanced the activity more than the thio group at the second position. Numerous functional group changes have been researched at the third and fifth positions. It was found that the nitro and cyano groups have enhanced the activity due to stronger electron-withdrawing activity and smaller volume than the ester and amide. Methyl ester has demonstrated greater activity among the esters than the ethyl ester group. Better activity has



Scheme 87 Synthesis of neonicotinoids insecticides with ester group.

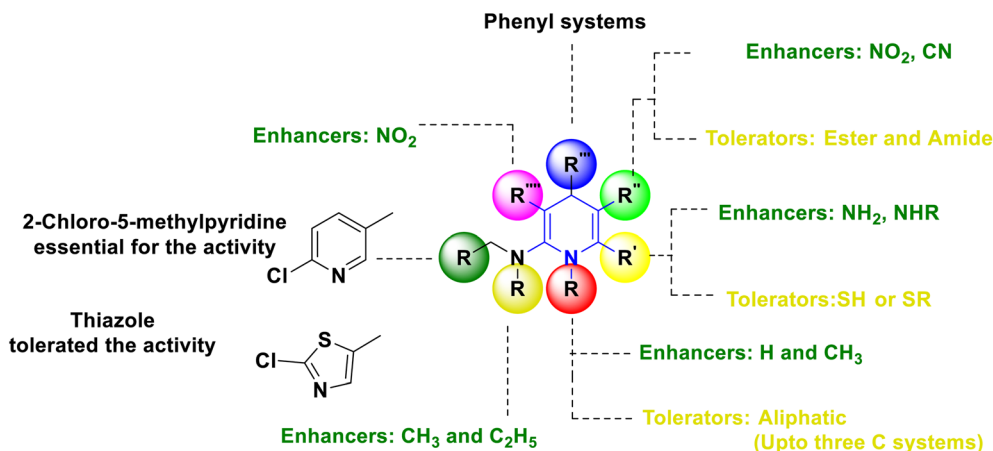


Fig. 11 SAR studies of insecticidal activities.

been seen for phenyl and substituted 4-Cl, 2-F, and electron-donating groups such OCH₃, OH, and CH₃ on phenyl systems at the fourth position. The important element for exerting the insecticidal activity was discovered to be the substitutions at the sixth position. The alkyl amine substituted with 6-chloro-pyridin-3-yl and 5-chloro-pyridin-3-yl, respectively, six-membered and five hetero aromatic systems, have shown enhanced activity at the sixth position. The activity has also been improved by substituting methyl and ethyl groups for hydrogen in the alkyl amine (-NH-) system. At the sixth position, clothianidin nitromethylene neonicotinoids are better than their nitenpyram counterparts in terms of insecticidal activity.

3. Conclusion

The current study concludes that 1,4-DHPs constitute a pharmacologically significant class of compounds. We have systematically described the work done on 1,4-DHP compounds with synthetic methodology developments, biological applications as an agrochemical, and pharmacological activities such as antihypertensive, anticancer, calcium channel blockers, anti-inflammatory, anti-microbial, anti-fungal, anti-oxidant, anticholinesterase, anti-coagulant, neuro-protective, and anti-diabetic activities in this study. However, there are limited publications and no review on the agrochemical applications of 1,4-DHPs. As a result, medicinal chemists are still exploring new synthesis techniques to generate new neonicotinoid-embedded 1,4-DHPs with insecticidal action. This review covers the insecticidal properties of 1,4-DHPs and *cis*-nitenpyram neonicotinoid molecular hybrids. These neonicotinoid 1,4-DHP compounds could be a potential source of chemicals for the pharmaceutical industry. The biological functions and SAR analysis of the compounds were presented in this review to stress their medicinal possibilities for future medicinal chemistry developments. This could be useful and enlightening for academic and industrial researchers working in the fields of 1,4-DHP base medicinal and pharmaceutical chemistry.

Conflicts of interest

The authors declare that there are no conflicts of interest.

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