A Review of the Recent Development in the Synthesis and Biological Evaluations of Pyrazole Derivatives

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Abstract: Pyrazoles are five-membered heterocyclic compounds that contain nitrogen. They are an important class of compounds for drug development; thus, they have attracted much attention. In the meantime, pyrazole derivatives have been synthesized as target structures and have demonstrated numerous biological activities such as antituberculosis, antimicrobial, antifungal, and anti-inflammatory. This review summarizes the results of published research on pyrazole derivatives synthesis and biological activities. The published research works on pyrazole derivatives synthesis and biological activities between January 2018 and December 2021 were retrieved from the Scopus database and reviewed accordingly.

Keywords: heterocycle; pyrazole; derivatives; synthesis; biological activities; recent development

1. Introduction

Heterocycles are a fundamental and unique class of compounds; they account for over half of all known organic compounds and have a broad range of physical, chemical, and biological properties, covering a broad spectrum of reactivity and stability [1]. Furthermore, their synthetic usefulness as synthetic intermediates, the protective groups, the chiral auxiliaries, the organocatalysts, and the metallic ligands in the asymmetric catalysts in pharmaceutical agents have rendered them multiple units of interest. Among heterocyclic compounds, five-membered rings containing nitrogen atoms constitute a vast and differentiated group with a broad spectrum of biological activity [2–4]. The members of this group, such as pyrazole, imidazole, oxazole, triazole, tetrazole, oxadiazole, thiazole, and isoxazole, are particularly important antibacterial and antifungal agents [3–5]. The pyrazole ring is a five-membered heterocycle containing two adjacent nitrogen atoms. It is a moiety found in many molecules that possess many applications. Additionally, naturally occurring pyrazoles and their synthetic derivatives are well-known to have a broad spectrum of biological properties (Figure 1). In recent years, some of the FDA-approved and commercialized drugs, including patented ones, have been developed from pyrazole derivatives (Figure 2), which implies ample usage of these groups in new-fangled bioactive molecules. This review focuses on a concise overview of the pyrazole pharmacophore synthesis and biological activities reported between 2018 and 2021. Thus, it will serve as a helpful reference guide for researchers interested in the field. This review is loosely categorized into chemical synthesis and biological applications. The first section includes the synthesis of pyrazole derivatives, and the second section describes the biological applications.
2. Synthesis of Pyrazole Derivatives

2.1. Condensation of Hydrazine’s or Similar Nuclei with Carbonyl Functional Group Compounds

2.1.1. Pyrazoles from Vinyl Ketones

The tandem reactions between amine-functionalized enamiones 1 and aryl sulfonyl hydrazine or tosylhydrazone derivatives 2 and 3 in the absence of a metal catalyst have
been reported [6]. The synthesis of the substituted pyrazoles 4 and 5 occurred in water, TBHP, and NaHCO₃, respectively. In addition, when alkyl-based sulfonyl hydrazine such as methyl sulfonyl hydrazine was incorporated, the reaction was not successful. Additionally, when the reaction was carried out in EtOH and DMF, all the analogs were obtained with a good yield. The expected products with a lower yield were formed by introducing double-ethyl functionalized enaminone (R₁ = R₂ = ethyl). (see Scheme 1).

Scheme 1. Synthesis of the substituted pyrazoles using hydrazine [6].

Wan et al. [7] reported a different synthetic route to forming-substituted pyrazole derivatives, including celecoxib (7a), mavacoxib (7b), and deracoxib (7c), respectively. The compounds were synthesized using enaminones and aryl hydrazines in ethanol with acetic acid. The reaction produced regioselective compounds with a high yield. Notably, compared to the other synthetic methods, using fluoroalkylated pyrazoles [8], β-diketones [9], and ynones [10], this method gives an excellent yield, a regioselective product. The synthetic route can be explored to synthesize pyrazole derivatives that are not easy to get from fluoroalkyl β-diketones. (see Scheme 2).

2.1.2. Pyrazoles from 1,3-Diketones

A silver-catalyzed synthesis of 5-aryl-3-trifluoromethyl pyrazoles using N'-benzylidene tolylsulfonylhydrazides 8 with ethyl 4,4,4-trifluoro-3-oxobutanoate 9 as precursors has been reported [11]. The reaction involved consecutive nucleophilic addition, intramolecular cyclization, elimination, and finally, [1,5]-H shift. This led to trifluoromethylated pyrazole derivatives 10 with moderate to excellent yields (see Scheme 3). In optimizing the product, the yield improved by increasing the reaction temperature to 60 °C, but increasing the reaction temperature above 60 °C resulted in a lower yield. The Cu(OTf)₂ transition catalyst afforded 60% yield, while Fe(OTf)₃ was unproductive. THF or dioxane gave a poor yield of the product compared to toluene. Meanwhile, K₂CO₃ was more effective than NaH, KOt-Bu, and NaOt-Bu. Additionally, the use of Me₂phen as a ligand yielded the best performance (>99%), compared to using bpy or phen as a ligand (57% or 92%).
Scheme 2. Synthetic route to the formation of celecoxib, deracoxib, and mavacoxib using hydrazine [7].

Scheme 3. Synthesis of 5-aryl-3-trifluoromethyl pyrazole derivatives in the presence of a silver catalyst [11].

Poletto et al. [12] reported the one-pot synthetic strategy for synthesizing highly regioselective $\alpha$-ketoamide N-arylpurazoles 28, the secondary $\beta$-enamine diketone, and arylhydrazines as precursors. Notably, the intermediate 4-acyl 3,5-dihydroxypyrrolone, produced in situ, went through nucleophilic substitution at C-5 by arylhydrazine. Afterward, heterocyclization occurred at the carbonyl carbon of the acyl group (see Scheme 4).
2.1.3. Pyrazoles from Acetylenic Ketones

The merging of substrates with five electron-rich heteroaromatic nuclei interacts with arylhydrazines with the carbonyl group and triple carbon bond. The cyclo-condensation of cross-conjugated enynones 13 with hydrazines has produced pyrazole derivatives 14 and 15 in good yield [13] (see Scheme 5).

2.2. Dipolar Cycloadditions

2.2.1. Pyrazoles from Diazoester

Fang et al. [14] reported the designed and synthesized polysubstituted 4-trifluoromethylpyrazoles using ketones 16 and trifluoroacetyl diazoester 17. The ketones reacted with the terminal nitrogen atom of the trifluoroacetyl diazoester, followed by cycliza-
tion, forming 4-trifluoromethylpyrazole derivatives 18–27. Meanwhile, the replacement of DBU with NEt₃, NaOT-Bu, CsF, Cs₂CO₃, and Na₂CO₃ resulted in a poor yield (see Scheme 6). Notably, dialkyl ketones did not afford the corresponding 4-(trifluoromethyl)pyrazoles 26 and 27, using DBU and NEt₃, which could be due to the relative lower reactivity of the α-hydrogen of dialkyl ketones.

Scheme 6. Synthesis of polysubstituted-4-trifluoromethylpyrazoles using ketones and trifluoroacetyl diazoester [14].

Chen et al. [15] reported a synthetic approach for synthesizing polysubstituted 4-difluoromethyl 28 and perfluoroalkyl 31 pyrazole derivatives. The authors utilized a Lewis acid and base co-mediated reaction of perfluoroacetyl diazoester with ketones (see Scheme 7). Catalysts such as Cu(OTf)₂, CuCN, Sc(OTf)₃, NiCl₂, FeC₁₃, Fe(OTf)₃, CoCl₂, and ZnI₂ were explored for the optimization of the reaction. Particularly, the Sc(OTf)₃ catalyst displayed the best performance with 97% yield in the presence of DBU as the base. Meanwhile, the base, such as Et₃N, t-BuOK, K₂CO₃, and K₃PO₄, did not give the expected product.
Scheme 7. Synthetic route to 4-difluoromethyl pyrazole derivatives using Sc(OTf)_3 as the catalyst [15].
2.2.2. Pyrazoles from Vinyldiazo Ketones

The synthesis of pyrazole-based triarylmethanes using 2-(1-alkynyl)-2-alken-1-ones and vinyldiazo ketones has been reported [16], as shown in Scheme 8. The first stage of the reaction involved heating vinyldiazo ketones \(32\) in dichloroethane, which produced \(1H\)-pyrazoles \(33\), followed by a reaction with enynones \(34\) to produce pyrazole-based triarylmethanes \(35\).

Scheme 8. Synthesis of pyrazole-based triarylmethanes using a gold catalyst [16].

2.2.3. Pyrazoles from Hydrazones

Zhu and colleagues [17] investigated the oxidative coupling reaction of phenylhydrazone \(36\) and maleimide \(37\) to synthesize pyrazoles derivatives. The reaction was carried out with CuCl as the catalyst, and dimethylformamide (DMF) was used as a solvent (see Schemes 9 and 10). In the reaction, it produced 12% yield in the presence of 20 mol% Cu(OAc)\(_2\) in dimethylsulfoxide (DMSO) at 80 °C for 2 h. Additionally, other Cu(II) salts did not enhance the reaction. The reaction yielded 86% when CuCl as the catalyst in DMSO was utilized, while Cu(I) salts, mainly CuOAc, CuBr, CuI, and CuSCN, led to product reduction. Moreover, catalysts such as Mn(OAc)\(_3\), Ag\(_2\)CO\(_3\), FeCl\(_3\), and Pd(OAc)\(_2\) were inefficient.

An effective protocol for synthesizing pyrazoles derivatives \(46\) and \(49\) via an iodine-catalyzed reaction of aldehyde hydrazones with electron-deficient olefins has been reported [18] (see Scheme 11). The transformation of the reaction produced a 35% yield in the presence of 20 mol% I\(_2\) and 3.0 equiv of TBHP in DMF at 80 °C. The solvents, such as CH\(_3\)CN, displayed a moderate yield. When different oxidants were utilized, BPO displayed a superior performance up to an 81% yield compared to TBHP, K\(_2\)S\(_2\)O\(_8\), DTBP, BTI, H\(_2\)O\(_2\), and m-CPBA. The product was reduced by replacing molecular iodine with other iodides, such as NaI, NIS, or TBAI.
Scheme 9. Synthesis of pyrazole derivatives via copper-catalyzed an oxidative coupling reaction [17].

Scheme 10. Synthesis of the pyrazole derivatives from aldehyde hydrazones via a copper-catalyzed oxidative coupling reaction [17].
Scheme 11. Synthesizing of pyrazoles derivatives via an iodine-catalyzed reaction [18].

A facile one-pot, copper-catalyzed aerobic cyclization has been consecutively used to synthesize the pyrazole derivatives (51 and 54) by Fan and coworkers [19]. In this reaction, β and γ-unsaturated hydrazones were readily available substrates. While O₂ acted as the terminal oxidant and economic Cu(I) salt was used as the catalytic agent, CuOTf showed the best performance compared to other employed catalysts such as CuOAc, CuBr, Cu(acac)₂, CuOTf, and Cu(OTf)₂ (see Schemes 12 and 13).

Scheme 12. Synthesis of the pyrazole derivatives in the presence of terminal oxidant and copper salt [19].
Scheme 13. Synthesis of the pyrazole derivatives from β,γ-unsaturated hydrazones [19].

The intermolecular, thermally activated, and DBU-aided [3 + 2] cycloaddition of pyridin-2-yl-[1,2,4]-triazine dipolarophiles 55 with structurally varied 4-methylbenzenesulfonohydrazides 24 produced 57 as the major isomer with an excellent yield [20] (see Scheme 14).

Scheme 14. Synthesis of pyrazole derivatives via the dipolar cycloaddition of terminal ethynyl pyridines with tosylhydrazides [20].
Zheng et al. [21] used a metal-free protocol to synthesize pyrazolylthienopyrimidines and other N-heteroaryl pyrazole derivatives 57 from α,β-unsaturated N-tosylhydrazones 103 and N-heteroaryl chlorides 58 under mild reaction conditions. The bi(heteroaryl) pyrazole derivatives were obtained in good to excellent yields (see Scheme 15).

![Scheme 15. Synthesis of pyrazole derivatives from α,β-unsaturated N-tosylhydrazones and N-heteroaryl chlorides [21].](image)

2.2.4. Pyrazoles from Diazo Intermediates and Alkynes

Dimirjian and colleagues [22] developed a synthetic approach for synthesizing fused pyrazoles via an intramolecular reaction. Notably, 1,3-dipolar cycloadditions of diazo intermediates with alkynes 60 produced the spirocyclic product of pyrazole derivatives 62 (see Scheme 16).

![Scheme 16. Synthetic route to the formation of spirocyclic pyrazole via 1,3-dipolar cycloadditions of diazo intermediates with alkynes [22].](image)

2.2.5. Pyrazoles from Vinyl Sulfone

Dihydro-pyrrolo-pyrazoles have been synthesized through a cascade reaction involving cinnamyl azides and vinyl sulfones with moderate to good yields [23]. The protecting group, ethylene sulfone, can be removed by heating the product in pyrrolidine (see Scheme 17). The reaction tolerated a range of solvents such as benzene, acetonitrile, methanol, 1,3-dichloroethene, isopropanol, and dioxane. However, dioxane with triethylamine as a base produced dihydro-pyrrolo-pyrazole excellently compared to dioxane and diisopropylethylamine (DIPEA) or diisopropylamine (DIPA).
Scheme 17. Synthesis of dihydro-pyrrolo-pyrazoles from vinyl sulfones [23].

2.2.6. Pyrazoles from Nitro-Olefins

The Rauhut—Currier cyclization reaction was reported for the synthesized trisubstituted pyrazole derivatives [24]. The trisubstituted tetrahydropyrano [2,3-c]pyrazoles were obtained from the domino Rauhut—Currier cyclization reaction. The reaction occurred between alkylidene pyrazolones and nitro-olefins (see Scheme 18).

Scheme 18. Synthesis of the trisubstituted tetrahydropyrano [2,3-c]pyrazoles domino Rauhut—Currier cyclization reaction [24].

2.2.7. Pyrazoles from Alkynes

A visible light-promoted cascade of Glaser coupling/annulation alkynes and hydrazines has been utilized to synthesize polysubstituted pyrazoles 73 and 75 [25] (see Scheme 19). The replacement of CuI with CuCl, CuCl₂, Cu(OTf)₂, and Cu(OAc)₂ in the reaction using maintaining Ru(bpy)₃Cl₂ as the photocatalyst did not improve the reaction product.

Scheme 19. Synthetic route to forming polysubstituted pyrazoles from Glaser coupling/annulation of alkynes with hydrazines [25].
2.2.8. Pyrazoles from Morita–Baylis–Hillman (MBH) Carbonates

Under mild reaction conditions, phosphine-catalyzed domino Morita–Baylis–Hillman (MBH) carbonates with diazenes yielded tetrahydropyrazole-fused heterocycles 78 and 81 with moderate to excellent yields [26] (see Scheme 20). The substitution of tert-butyl (81g) with MBH carbonate was not reactive, probably due to its steric barrier. It is noteworthy that MBH carbonates derived from other aldehydes, namely benzaldehyde, did not function in this reaction.

Scheme 20. Synthesis of tetrahydropyrazole-fused heterocycles from Morita–Baylis–Hillman (MBH) carbonates [26].
Multicomponent Strategies

A one-pot multicomponent reaction for synthesizing bispyranopyrazole derivatives using MMT K10 as a support heterogeneous catalytic system has been reported [27] (see Scheme 21). Notably, the economic and environmentally friendly catalyst was recycled and reused five times in the reaction, and no lack of activity was observed.

\[
\begin{align*}
\text{CHO} & \quad \text{CHO} \\
\text{CH}_2 & \quad \text{CH}_2 \\
\text{R}_1 & \quad \text{R}_2 \\
\text{R}_3 & \quad \text{N} \quad \text{N} \\
\text{H}_2 & \quad \text{H}_2 \\
\text{O} & \quad \text{O} \\
\end{align*}
\]

\[
\begin{align*}
\text{H}_2 & \cdot \text{OEtOH, reflux} \\
\text{MMT-} & \text{[(CH}_3)_2\text{NH-CHPy]} \cdot \text{Cu(I]} \\
\end{align*}
\]

Scheme 21. Synthesis of bispyranopyrazole under a support heterogeneous catalytic system [27].

Alizadeh-Kouzehrash et al. [28] reported new \textit{N}-fused pyrazole derivatives via an efficient one-pot multicomponent reaction. Using a cheap catalyst, namely 4-toluenesulfonic and ethanol, as a green organic solvent (see Scheme 22), the best yields were reached in ethanol as a solvent under the reflux temperature, and in the absence of a catalyst, the yields percentage of reactions were reduced.

\[
\begin{align*}
\text{O} & \quad \text{N} \quad \text{N} \\
\text{R} & \quad \text{R} \\
\text{NH}_2 & \quad \text{NH}_2 \\
\text{H}_2 & \quad \text{H}_2 \\
\end{align*}
\]

\[
\begin{align*}
\text{CHO} & \quad \text{CN} \\
\text{R} & \quad \text{R} \\
\text{HO} & \quad \text{OH}^+ \\
\text{HO} & \quad \text{HO}^+ \\
\text{NH}_2 & \quad \text{NH}_2 \\
\end{align*}
\]

Scheme 22. Synthesis of fused pyrazole derivatives using 4-toluenesulfonic as the catalyst [28].
3. Miscellaneous

The synthetic route to JNJ-18038683 and other fused pyrazole derivatives were reported by Dvorak et al. [29]. Two synthetic routes were employed to construct the fused pyrazole-azepine heterocyclic core (see Schemes 23–25). In the reaction path to pyrazole triflate, a bi-phasic solvent system of toluene/water was optimal, and no triflate hydrolysis was identified. Meanwhile, the displacement of the BOC-protecting group was achieved by treating with trifluoroacetic acid to give the free base of the amine. Subsequently, the free base was converted into citrate salt, and the salt formation was then performed with a free base and citric acid in methanol. The final recrystallization yielded clinical candidate as a nonhygroscopic free-flow powder.

Scheme 23. Synthetic route to the formation of fused pyrazole [29].

Scheme 24. Synthetic route to the formation of pyrazole triflate using N-phenyltriflamide in pyridine [29].
Scheme 25. Synthetic route to the formation of the JNJ-18038683 heterocyclic compound [29].

1H-pyrazole-5-amines were obtained from the microwave reaction of arylhydrazines 108 with 3-aminocrotonitrile 109 in moderate to excellent yields after 10 min of irradiation [30]. In addition, the reaction of phenylhydrazine 110 with α-cyanoketones 110 under similar conditions produced many functionalized 1H-pyrazole-5-amines. Meanwhile, the m-nitrophenyl and p-nitrophenyl-3-oxopropanenitrile substituents did not react with phenylhydrazine, even at a longer heating time. However, the reaction conditions tolerated other functionalized aromatic groups such as trifluoromethyl and methyl sulfone (see Scheme 26).

Scheme 26. Synthesis of pyrazole derivatives via the microwave reaction of arylhydrazines with 3-aminocrotonitrile [30].
4. Biological Activity of Pyrazole Derivatives

4.1. Anti-Inflammatory

Bhale et al. [31] reported synthesizing 1,3,4,5-tetrasubstituted pyrazole derivatives and their in vitro anti-inflammatory effect (see Figure 3). Compound 117a showed excellent inhibition (93.80%) compared to the standard diclofenac sodium (90.21%) at a 1 mM concentration. El-Karim et al. [32] reported compounds 118a–118f (edema inhibition% = 98.16%, 96.73%, 88.81%, 81.5%, 76.17%, and 76.68%, respectively) as potent candidates producing rapid onset and a long duration of anti-inflammatory activity, as well as a good safety GIT profile. Meanwhile, the analgesic evaluation revealed that 118b–118e produced potent and long-acting analgesia accompanied by a significant inhibition of the inflammatory cytokine TNF-α level compared to the standard drugs. The inhibition in the protein denaturation of bovine albumin with IC_{50} of 34.1 µg/mL using diclofenac sodium as the standard drug (IC_{50} = 31.4 µg/mL). Out of the 15 novel compounds synthesized by Akhtar et al. [33], 123a–123d demonstrated a significant in vitro anti-inflammatory activity, with IC_{50} values of 71.11, 81.77, 76.58, and 73.35 µg/mL, respectively, compared with the standard diclofenac. The benzylidene substituent attached remarkably influenced the anti-inflammatory potency. Abdellatif et al. [34] synthesized a new series of pyrazole derivatives. The inhibition efficacy of the target compounds to ovine COX-1 and human recombinant COX-2 was analyzed using an immune enzyme assay (EIA) kit. Most of the tested compounds showed high COX-2 inhibitory activity with IC_{50} values ranging from 0.02–0.04 µM. Meanwhile, 119a and 119b had the most suitable COX-2 selectivity index (SI = 462.91 and 334.25, respectively), superior to celecoxib (SI = 313.12) and indomethacin (SI = 1.37). Compounds 119a and 119b (SO_{2}NH_{2} as the selective COX-2 pharmacophore) also showed the highest anti-inflammatory activity (ED_{50} = 136 and 126 µmol/kg, respectively). In addition, they have the lowest ulcerogenic liability (Ulcer Index = 1.25 and 1.00, respectively), reflecting their expected safe GI profiles. Shi and coworkers [35] discovered 120 as the most potent anti-inflammatory agent (IC_{50} = 3.17 µM), with low toxicity and strong inhibitory NO release (inhibitory rates (IR) = 90.4% at 10 µM). This compound also showed potent inhibition of iNOS with an IC_{50} value of 1.12 µM. The treatment of compound 120 on acute inflammatory models in AA rats displayed a remarkable inhibitory effect on hind paw swelling and body weight loss, comparable to the effect identified in the aspirin-treated group. Of all the compounds investigated by Sivaramakarthikeyan et al. [36], the para-nitrophenyl moiety linked to a pyrazole conjugate 121 (93.53 ± 1.37%) displayed the highest anti-inflammatory activity in the anti-inflammatory assay using the protein denaturation method. This is superior to the standard, diclofenac sodium (90.13 ± 1.45%). Nayak et al. [37] revealed that compound 122a showed remarkable sodium and celecoxib, which showed IC_{50} values of 55.65 and 44.81 µg/mL, respectively. The potent compounds were further evaluated for their in vitro COX-2 inhibitory activities using an enzyme immunoassay. Compound 123d demonstrated able selectivity toward COX-2 with a selectivity index (SI) of 80.03 compared with the standard celecoxib, with an SI of 95.84. Dimmito et al. [38] reported that compound 124a displayed a good analgesic effect after subcutaneous and intracerebroventricular management in vivo. Additionally, 124a showed an excellent anti-inflammatory effect after subcutaneous administration, indicating prospective activity at the periphery. Harras and colleagues [39] synthesized a series of pyrazole derivatives and evaluated their in vitro COX-1/COX-2 inhibition and in vivo anti-inflammatory activity using the carrageenan rat paw edema model. It was noted that the targeted compounds exhibited more potent inhibitory activity against COX-2 than COX-1. Meanwhile, all compounds’ selectivity indexes (SI) were analyzed and compared to celecoxib (SI = 8.17). Compounds 125a and 125b displayed an outstanding COX-2 selectivity index of 8.22 and 9.31, respectively. Meanwhile, the histopathological investigation of the rats’ stomach, liver, and kidneys revealed that 125a and 125b triggered minimal degenerative changes, suggesting these derivatives’ safety. Sivaramakarthikeyan et al. [40] reported the anti-inflammatory activity of the pyrazole derivatives. The derivative, lacking substitution on the aryl entity 126, exhibited the highest anti-inflammatory profile. Ab-
dellatif et al. [41] synthesized a series of substituted pyrazole derivatives. The targeted compounds were screened for their COX-1/COX-2 inhibitory activity. Additionally, the carrageenan-induced rat paw edema model and histopathological study were demonstrated to examine their anti-inflammatory effectiveness and gastric safety. Compound 127 was the most potent anti-inflammatory agent (ED$_{50}$ = 65.6 µmol/kg) compared to the reference drug, celecoxib (ED$_{50}$ = 78.8 µmol/kg). In addition, the potent compound possessed minimum ulcerogenic (Ulcer Index = 7.25) Figure 3.

**Figure 3.** Structures of pyrazole derivatives with anti-inflammatory activity.
A new series of thiazolidindione 128 and thiazolidinone 129 containing a pyrazole core has been synthesized as hybrid structures [42]. The synthesized compounds were further evaluated for COX-1/COX-2 in vitro anti-inflammatory activity and ulcerogenic liability (Figure 3). The most COX-2-selective derivatives 128a and 128b and 129a and 129b showed the highest anti-inflammatory activities and the lowest ulcerogenic. Among the potent compounds, the thiazolidindione with a methoxy substituent 129b displayed excellent activity against COX-2 (IC$_{50}$ = 0.88 µM) with the highest COX-2 selectivity index (SI = 9.26). While compound 128c with a methoxy substituent displayed the highest potent inhibitory against COX-2 (IC$_{50}$ = 0.62 µM) with the highest COX-2 selectivity index (SI = 8.85). The highest anti-inflammatory (AI) activities were observed in 129a and 128b (after 1 h, AI = 82.34 and 81.15%; after 3 h, AI = 79.00 and 97.68%; and after 5 h, AI = 80.15 and 97.68%, respectively). Additionally, 128a was slightly more potent (ED$_{50}$ = 79.12 µmol/kg) than celecoxib (ED$_{50}$ = 82.2 µmol/kg), while 128b showed a superior ED$_{50}$ value of 5.63 µmol/kg with a more than 14-fold effectiveness of celecoxib. Some pyrazolopyrimidine hybrids were prepared using Schiff base by Abdelall and coworkers [43]. All the synthesized compounds were evaluated in vivo against carrageenan-induced rat paw edema as anti-inflammatory agents. Regarding the anti-inflammatory activity compounds, 130 and 131 showed excellent activity compared to celecoxib. Thangarasu et al. [44] reported the anti-inflammatory effect of pyrazole moieties, and compound 132b was found to have dominated activity potentials with an IC$_{50}$ value of 3.5 nM in the COX-2 inhibition studies.

Murahari et al. [45] designed and synthesized novel pyrazole-based derivatives using the ligand-based approach. Among the synthesized compounds, 133 showed excellent in vivo anti-inflammatory activity with 0.8575 mmol/kg as ED$_{50}$. The design and synthesis of novel thiophene–pyrazole hybrids have been investigated [46]. The thienopyrimidine analogs 134a–134b and the thiophene derivative 135 are promising nontoxic, gastrointestinal-safe anti-inflammatory candidates with good oral bioavailability and physicochemical properties. A series of 1,2,3-triazole-linked 3-(1,3-diphenyl-1H-pyrazol-4-yl)acrylates was synthesized following a multi-step reaction [47] (Figure 4). Three of the evaluated compounds demonstrated significant anti-inflammatory activity, with IC$_{50}$ values of 60.56, 57.24, and 69.15 µg/mL for compounds 136a–13c, respectively, comparable to the standard diclofenac sodium with an IC$_{50}$ value of 54.65 µg/mL.

Taher and colleagues [48] reported the synthesis and pharmacologic evaluation of novel pyrazole and pyrazoline derivatives. The study presents the effect of lengthening the carbon chain in different pyrazole derivatives bearing various amine moieties. Their results showed that lengthening of the aliphatic chain in 137a–137c (26.19%, 30.95%, and 28.57%, respectively) led to higher activity. Meanwhile, the cyclization of chalcones into pyrazolines were more potently anti-inflammatory in compounds 138, 139a and 139b (21.43%, 26.19%, and 28.57%. Compounds 138 and 140 exhibited the highest analgesic activity among all the examined compounds (75.9% and 84.5%, respectively). Mustafa et al. [49] presented a novel series of celecoxib derivatives. The in vivo anti-inflammatory activity of the synthesized compounds was evaluated using celecoxib as a reference standard by the paw oedema model on albino Wistars. Most of the compounds showed higher in vivo anti-inflammatory activity compared to celecoxib. Different substituents on the triazole moiety played a crucial role in the percentage inhibition of anti-inflammatory effects at 1h. Derivatives with chlorine atoms 141a–141d and the nitro derivative 141e showed good anti-inflammatory potency (Figure 4). A series of novel benzophenones conjugated with an oxadiazole sulfur bridge pyrazole has been designed, synthesized, and characterized [50]. It was afterward evaluated for anti-inflammatory and analgesic effects. Among the series, compound 142 (65.38% edema inhibition) with an electron-withdrawing group (fluoro) at the para position of the benzoyl ring of benzophenone was characterized by great activity compared to the standard drug. The analgesics activity data also revealed that compound 142 was the highest potent compound among the compounds evaluated for an analgesic effect on the acetic acid-induced writhing response and thermal pain (see Figure 4).
Figure 4. Structures of the pyrazole derivative with anti-inflammatory activity.
A novel series of pyrazole hybrids, such as pyrazole-thiohydantoin and pyrazole-methylsulfonyl, was synthesized by Abdellatif et al. [51]. The hybrids were evaluated in vivo for their anti-inflammatory activity (Figure 5). Compounds 143a–143d were found to have the most active anti-inflammation. The unsubstituted 143b and 143d showed comparable ED50 (78.90 and 88.28 µmol/kg) with celecoxib (ED50 = 78.53 µmol/kg), while the methoxy-substituted compounds 143a, 143c, and 143e (ED50 = 62.61, 55.83, and 58.49 µmol/kg, respectively) showed superior activity to celecoxib. Thirteen pyrazole derivatives were synthesized and evaluated for their anti-inflammatory activity (in vitro and in vivo) and ulcerogenic liability [52]. Nine compounds 144–146 exhibited a moderate to high edema inhibition percentage (78.9–96%) than celecoxib (82.8%). Additionally, they were found to have potent COX-2 inhibitory activity, with the IC50 values ranging from 0.034 to 0.052 µM. Compound 145a was the benign pyrazole with respect to the ulcerogenic effect (UI = 0.7) on the stomach, which may be ascribed to its high COX-2 enzyme selectivity (SI = 353.8), while compounds 144a, 145b, and 146a–c exhibited ulcer index values (UI = 0.8–2) comparable to celecoxib. Sulphonyl derivatives 147 and 148 have been reported to be selective for the COX-2 isozyme with COX-2 selectivity indexes of 9.78, 8.57, 10.78, and 10.47, respectively, compared to celecoxib (SI = 8.68) [53]. Meanwhile 147 and 148 were observed as excellent anti-inflammatory derivatives with ED50 = 51.51, 46.98, 53.65, and 54.45 µmol/kg better than celecoxib (ED50 = 76.09 µmol/kg). Gedawy et al. [54] reported novel pyrazole sulfonamide derivatives as dual COX-2/5-LOX inhibitors. The benzothiophen-2-yl pyrazole carboxylic acid derivative 149 showed the most potent analgesic and anti-inflammatory activity superior to celecoxib and indomethacin. It showed potent COX-1, COX-2, and 5-LOX inhibitory activities, with IC50 of 5.40, 0.01, and 1.78 µM, respectively, showing a selectivity index of 344.56 superior to the reference standards (see Figure 5).

A new pyrazole sulfonate series has been reported [55]. Among the series, 4-iodophenyl 5-methyl-3-(p-tolyl)-1H-pyrazole-1-sulfonate 150a and phenyl 5-methyl-3-(4-(trifluoromethyl) phenyl)-1H-pyrazole-1-sulfonate 150b displayed superior anti-inflammatory activity (% inhibition of auricular edemas = 27.0 and 35.9, respectively); while the in vivo analgesic activity of 150c and 150d was more effective with an inhibition of 50.7% and 48.5% separately, and compounds 150a, 150c, and 150d were identified as selective COX-2 inhibitors (SI = 455, 10,497, and >189, respectively). In addition, the acute oral toxicity in vivo analysis showed lethal doses of 50 (LD50) of 150a and 150d to mice to be more than 2000 mg/kg. A novel series of 1,5-diaryl pyrazole-3-carboxamides was synthesized and evaluated against COX-1, COX-2, and sEH enzymes as dual COX-2/sEH inhibitors [56]. The anti-inflammatory activities of compounds 151a–c were superior (edema inhibition percentages of 62%, 71%, and 65%) to the reference drug celecoxib (22%). Compounds 151a–c, the most potent dual COX-2/sEH inhibitors in vitro, displayed the highest analgesic activity, with a % inhibition of 62.68, 71.64, and 67.16, respectively, and potencies 4.66, 5.33, and 5, respectively. Furthermore, compounds 151b and 151c substantially decreased the serum concentration of TNF-α with a % inhibition of 77% and 75%, respectively, when compared to celecoxib (64%). Compounds 152 and 153 have been reported as promising anti-inflammatory agents [57]. The compounds inhibited the lipoygenase enzyme with IC50 values of 2.17 ± 0.12 and 2.53 ± 0.06 µM, respectively, compared to the standard quercetin (IC50 value = 3.35 ± 0.01 µM) (see Figure 5).
4.2. Anticancer

Compound 5-(5-Bromo-1-methyl-1H-indol-3-yl)-1-(4-cyano-phenyl)-3-methylsulfanyl-1H-pyrazole-4-carbonitrile 117b showed significant cytotoxicity against MCF 7 (GI$_{50}$ = 15.6 µM) with low cytotoxicity against a normal Vero cell line [31] (see Figure 4).

Sivaramakarthikeyan et al. [36] reported that the evaluation of the anticancer potency of the synthesized pyrazole-benzimidazole hybrids revealed that the hybrids bearing a para-fluorophenyl unit tethered at the pyrazole nucleus (121b) showed the highest activity against both the pancreatic cancer cells (SW1990 and AsPC1) with IC$_{50}$ of 30.9 ± 0.77
and 32.8 ± 3.44 µM compared to the reference compound, gemcitabine 35.09 ± 1.78 and 39.27 ± 4.44 µM. Akhtar et al. [33] revealed that 123b was active against A549, SiHa, COLO205, and HepG2 cancer cell lines, with IC₅₀ values of 4.94, 4.54, 4.86, and 2.09 µM. The potent compound 123b was also nontoxic against normal cells (cell line HaCaT), with an IC₅₀ value greater than 50 µM (see Figure 6).

![Figure 6. Pyrazole hybrids with anticancer activity.](image)

Sivaramakarthikeyan et al. [40] showed that 126 exhibited significant activity against both the pancreatic cell lines—namely, AsPC1 and SW1990—with IC₅₀ values of 30.3 ± 0.45, 32.4 ± 0.65 µM and noncancerous cell—namely, MRC5 with an IC₅₀ value of 55.5 ± 3.50 µM (Figure 3). The synthesis and anticancer evaluation of a series of 1,2,3-triazole linked 3-(1,3-diphenyl-1H-pyrazol-4-yl)acrylates have been carried out [47]. Compound 136b showed the most promising anticancer effects among the synthesized compounds, with IC₅₀ values of 1.962, 3.597, 1.764, and 4.496 µM against the A549, HCT-116, MCF-7, and HT-29 cell lines, respectively. The sulphamoyl derivatives 148a and 148b exhibited the most potent activity, with IC₅₀ of 5.34 and 6.48 µM against A549 and IC₅₀ of 4.71 and 5.33 µM against MCF-7. Additionally, the compounds showed potent activity, with IC₅₀ values of 4.39 and 5.12 µM against HCT-116 and IC₅₀ of 3.66 and 4.37 µM against PC-3 [53]. A further analysis disclosed that compounds 148a and 148b arrested the cell cycle activity on the PC-3 cell line with greater selectivity. Meanwhile, the antiproliferation potency of compounds 148a and 148b on PC-3 cells is due to cell cycle arrest and apoptosis-inducing activity categorized by the Bax/Bcl-2 ratio increase (see Figure 7).

Compounds 154–156 were effective VEGFR-2 kinase inhibitors with IC₅₀ of 913.51, 225.17, and 828.23 nM, respectively, compared to sorafenib (IC₅₀ = 186.54 nM) [58]. Further, the cellular mechanistic studies of 156 revealed its promptness towards pre-G1 apoptosis and cell growth termination at the G2/M phase.

A new series of pyrazolo[1,5-a]pyrimidine derivatives has been designed and evaluated for their cytotoxic activities on a human breast adenocarcinoma cell line (MCF-7) and colon cancer cell line (HTC-116) [59]. The results revealed that 157 was the most potent among the tested compounds against HTC-116, with an IC₅₀ value of 1.51 µM, while 158 (IC₅₀ = 7.68 µM) displayed an excellent cytotoxic effect superior to reference doxorubicin against MCF-7. The tetrahydrothiochromeno [4,3-c]pyrazole derivatives were synthesized and evaluated for anticancer activity using MTT [60]. Most of these compounds showed potential anticancer activity and low cytotoxicity on the normal cells in vitro. Compounds 159a and 159b displayed excellent anticancer activity, with IC₅₀ values of 15.43 µM and 20.54 µM towards MGC-803, respectively. Additionally, the potent compounds 159a and 159b induced G2/M cell cycle arrest and apoptosis in MGC-803 cells. New pyrazole Schiff bases containing azo groups 160 have been reported as promising anticancer agents [61]. A new series of novel pyrazole-containing imide derivatives were synthesized and evaluated for their anticancer activities against the A-549, Bel7402, and HCT-8 cell lines [62]. Among the evaluated compounds, 161a–161d exhibited potent inhibitory activity against the A-549 cell line, with IC₅₀ values at 4.91, 3.22, 27.43, and 18.14 µM, respectively, superior to
5-fluorouracil (IC$_{50}$ = 59.27 µM). Additionally, 161a–161c exhibited substantial inhibitory activity towards the HCT-8 and Bel7402 cell lines (see Figure 7).

Figure 7. Structures of pyrazole derivatives with anticancer activity.
The cytotoxic activity of spirocycloadducts and N-arylpurazole hybrids against the HeLa cancer cell line was evaluated using an MTT assay [63]. Spiro[indenooxinoaline-pyrrolizidine]-N-arylpurazole conjugate 162 bearing a p-chlorophenyl substituent exhibited the highest antiproliferative activity against cancer cell line HeLa (IC$_{50} = 1.93$ µM). The IC$_{50}$ is comparable to camptothecin’s standard drug (IC$_{50} = 1.66$ µM) (see Figure 7). A novel series of 1,3-triazole-pyrazole hybrids were designed and synthesized using the Cu-catalyst [64]. The synthesized compounds were evaluated for anticancer activity using three cancer cell line panels. Compound 163 was the most potent cytotoxic candidate for HepG-2, HCT-116, and MCF-7, with IC$_{50}$ = 12.22, 14.16, and 14.64 µM, respectively, comparable to the standard drug doxorubicin (IC$_{50} = 11.21$, 12.46, and 13.45 µM). Ragab et al. [65] revealed compounds 164 and 165 as promising leads for colon cancer treatment. Compounds 164 and 165 were active against the KM12 cell line, with an IC$_{50}$ value of 1.73 and 1.21 µM and high selectivity index (SI) (18.82 and 35.49, respectively). Compared to the standard drug 5-FU with an IC$_{50}$ value of 12.26 µM and SI value of 1.93. The potent compound displayed selective cytotoxic activity against KM12 cells in the annexin V-FITC staining assay.

Mohamady et al. [66] designed and synthesized diarylpyrazole derivatives. The compounds were screened against the MCF7 and HepG2 cell lines. Among the evaluated compounds, 166, which contained a thiophene ring, was observed to have the highest antiproliferative activity against HepG2 cells, with an IC$_{50}$ of 0.083 µM. The compound caused cell cycle arrest at the G2, and the 7.7-fold increase in caspase-3 confirmed its apoptotic effect on HepG2 cells. Additionally, 166 caused a noticeable decrease in Hsp90 proteins (Akt, c-Met, c-Raf, and EGFR) and a 1.57-fold upsurge in Hsp70.

Pyrazolo-thiazole-substituted pyridine conjugates were synthesized and evaluated for cytotoxicity activity [67]. Compound 167—namely, 4-amino-7-(2-(1,5-dimethyl-1H-pyrazol-3-yl)-4-methylthiazol-5-yl)-2-oxo-5-(thiophen-2-yl)-1,2-dihydro-1,8-naphthyridine-3-carbonitrile—has the highest cytotoxicity activity towards PC-3 (IC$_{50} = 17.50$ µM), NCI-H460 (IC$_{50} = 15.42$ µM) and Hela (IC$_{50} = 14.62$ µM), comparable to the anticancer potential of standard drug etoposide (IC$_{50} = 17.15$, 14.28, and 13.34 µM, respectively).

Wang et al. [68] investigated a new series of pyrazole-naphthalene derivatives. The synthesized compounds were evaluated for their anticancer activity against breast cancer cell lines (MCF-7). Compound 168 (IC$_{50} = 2.78$ ± 0.24 µM), with substituted ethoxy at position 4 of the phenyl ring, exhibited the highest activity, and the activity was 5-fold more active than the reference drug cisplatin (IC$_{50} = 15.24$ ± 1.27 µM). Additionally, 168 showed inhibited tubulin polymerization with an IC$_{50}$ value of 4.6 µM. Mohamed and coworkers [69] recommended cyanoacrylamide compound 169 as a new promising chemotherapeutic agent. The compound exhibited high cytotoxic activity toward colorectal carcinoma. Compounds 170 and 171 displayed excellent antiproliferative activities towards HDAC2, with IC$_{50}$ values of 0.25 and 0.24 nM, respectively, and CDK2 with IC$_{50}$ values of 0.30 and 0.56 nM, respectively. The potent compounds 170 and 171 pointedly inhibited the movement of the A375 and H460 cells, arrested the cell cycle in the G2/M phase, and promoted apoptosis in A375, HCT116, H460, and Hela cells, related to proliferating the intracellular reactive oxygen species (ROS) levels. Notably, 170 has good pharmacokinetic properties, with an intraperitoneal bioavailability of 63.6% in ICR mice. Additionally, the compound was effective in vivo for antitumor activity in the HCT116 xenograft model. Thus, the authors proposed compound 171 as a favorable agent for treating malignant tumors. Burgart and colleagues [70] found 4-aminopyrazole derivatives 172a and 172b to be cytotoxic against HeLa cells and human dermal fibroblasts cancer cells. A novel series of pyrazole-arylacemamide hybrids has been synthesized and evaluated for cytotoxicity activity. Compounds 173a and 173b exhibited a potent cytotoxic effect on the MCF-7 cancer cell line, with IC$_{50}$ values of 0.604 µM and 0.665 µM compared to the standard drug cisplatin (0.636 ± 0.458 µM) (see Figure 7).

Answer et al. [71] synthesized some novel pyrazole hybrids using 5-amino-3-(4-(dimethylamino) phenyl)-1-phenyl-1H-pyrazole-4-carbonitrile as a precursor. Including
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different nucleophilic and electrophilic compounds, among the synthesized compounds, the anticancer activities of the synthesized compounds 174, 175, 176, and 177 (6.50 ± 0.5, 3.74 ± 0.3, 3.18 ± 0.2, and 8.67 ± 0.9 µM, respectively) exhibited strong cytotoxic activity against MCF-7 and HCT-116 (7.80 ± 0.6, 4.93 ± 0.3, 4.63 ± 0.4, and 10.02 ± 1.0 µM, respectively) (see Figure 7). Hassan et al. [72] synthesized a novel series of pyrazolopyrimidines and screened the compounds against a panel of 60 human cancer cell lines. Compounds, 5-amino-1H-pyrazole-4-carbonitrile derivative 177, pyrazolo[5,1-b]quinazoline-11-carbonitrile derivative 178, and 1-amino-2,4-dihydro-5H-benzo[4,5]imidazo[1,2-c]pyrazolo[4,3-e]pyrimidin-5-one 179 exhibited anticancer activity against some cancer cell lines (Figure 7). Compounds 180a–180c have been reported to displayed anticancer [73] compounds 180a–180c bearing an electron-donating group, such as methoxy substituent at the para position; the 3,4 dimethoxy and 3,4,5 trimethoxy derivatives demonstrated noticeable cytotoxic activity with IC_{50} values of 0.604 µM, 1.057 µM, and 0.665 µM respectively against the MCF-7 cancer cell lines.

Thiazolyl pyrazole carbaldehyde hybrids have been synthesized and screened for their in vitro anticancer activity by Mamidala and colleagues [74]. Compound 181 exhibited the highest antiproliferative activity against the HeLa, MCF-7, and A549 cancer cell lines, with IC_{50} values of 9.05 ± 0.04, 7.12 ± 0.04, and 6.34 ± 0.06 µM, respectively. Raghu et al. [75] designed and synthesized a new series of 1,3,5-triazine-based pyrazole hybrids with anticancer activity targeting the epidermal growth factor (EGFR) tyrosine kinase. Compounds 182a–182c exhibited potent anticancer activity against the MCF-7 (human breast), HepG2 (human liver), HCT116 (human colorectal), PC-3 (human prostate), LoVo (human colon), and LoVo/DX (doxorubicin-resistant) cancer cell lines. According to the EGFR tyrosine kinase test, 182a–182c demonstrated excellent activity, with an IC_{50} value of 395.1, 286.9, and 229.4 nM. Compared to the reference doxorubicin (63.8 nM) and erlotinib (103.8 nM), compound 182c, with a trifluoromethyl group at the para position on the phenyl rings, exhibited the most potent anticancer activity. Thus, the anticancer activity of the tested compounds was affected by the physicochemical properties of the substituent on the pyrazole-bound phenyl nucleus. Compounds 183a–183c have been reported as promising antiproliferative agents [76]. Suryanarayana et al. [77] synthesized a novel series of dinitrophenylpyrazole-bearing triazole and further investigated their anticancer activity using three tumor cell lines—namely, MCF-7, HeLa, and HeLa Caco-2. Among the synthesized compounds 184a–184c with the methoxy group on the phenyl ring at the ortho, meta, or para position exhibited excellent inhibitory activity against the HeLa (IC_{50} = 4.0 µM, 5.0 µM, and 6.0 µM) compared to the standard drug combretastatin-A4 (IC_{50} = 9.0 µM). Compound 185c exhibited excellent inhibitory activity against the MCF-7 cell line, with an IC_{50} value of 8.0 µM. Compound 185 has been reported as a promising anticancer agent that reduced the level of CDK2, stopped MCF-7 cells in the G0/G1 phase, caused ROS growth, damaged the MMP, and accelerated the apoptosis of MCF-7 cells [78] (see Figure 8). Among the new library of pyrazole derivatives investigated for antiproliferative activity by Signorello and coworkers, compounds 186a and 186b exhibited antiproliferative effectiveness [79]. Compound 186a displayed moderate inhibition (25–30% at 10 µM) toward melanoma (SK-MEL-5, UACC-62) and renal cancer cell lines (UO-31 cancer cell lines). Compound 186b displayed a broad spectrum of action (25–30%) towards leukemia (CCFR-CEM and RPMI-8226), non-small cell lung (NCI-H522), CNS (SF-295 and SNB-75), ovarian (OVCAR-4) and the breast (BT-549 and MDA-MB-468) cancer cell line. Additionally, compound 186b demonstrated 50–60% inhibition towards melanoma (SK-MEL-5 and UACC-62), renal (CAK-1 and UO-31), and prostate (PC-3) cancer cell lines. The water-soluble, BBB4-loaded NPs (BBB4-G4K NPs, 187), achieved from BBB4 in a non-bioactive, polyester-based, lysine-containing fourth-generation cationic dendrimer (G4K) has been reported to have an excellent antibacterial profile and highly selective toward the Staphylococcus genus [80]. The synthesis and biological screening of 5- pyrazolyl urea as potential antiangiogenic compounds were investigated by Morretta et al. [81]. Among the targeted compounds, compound 188a displayed 100% inhibition on leukemia cancer cell...
lines, while 188b and 188c impeded non-small cell lung cancer cell lines. Compound 188d, similar to the STIRUR-41 pharmacophore, exhibited 40% inhibition in the non-small cell lung cancer cell lines. Additionally, compounds 188e and 188f, containing a trifluoromethyl substituent on the urea moiety, displayed excellent inhibitory activity. Meanwhile, the mechanism of action detailed that 188e may likely exert its antiproliferative activity by targeting different signaling pathways, including ERK/MAPK and phosphatases, or the crosstalk between these two associated intracellular mechanisms. Compound 188e can regulate ERK1/2 phosphorylation and PPIg action.

Figure 8. Structures of pyrazole derivatives acting as anticancer agents.
4.3. Antibacterial

Nayak et al. [37] reported 122a–122g as promising antibacterial agents (Figure 3). Ebenezer et al. [82] reported a designed and synthesized library of novel pyrazole–imidazo [1,2-α]pyridine scaffolds through a one-pot three-component tandem reaction. All selected compounds (zone of inhibition 9 mm) showed excellent bactericidal activity. In most cases, except for MRSA, the activity of the compounds was better than that of the standard ciprofloxacin. Compounds 189a–189e had excellent activity against S. aureus, E. coli, S. typhimurium, K. pneumoniae, and P. aeruginosa, a with minimum bactericidal concentrations (MBC) <0.1 μg/mL. Pyrazoloquinoline derivatives have been synthesized and their antibacterial activity evaluated using the Agar diffusion method [83]. The ketonic compounds 190a and 190b showed activity percentages of 112% and 95% (Streptococcus pneumoniae) and 86% and 83%, respectively (Bacillus subtilis), compared to the standard control. The halogenated ketonic derivative 190b showed improved activity compared to gentamicin (109%), and no activity was observed in Pseudomonas aeruginosa. Hansa and coworkers [84] reported 4-(anilinomethyl)-3-[4-(trifluoromethyl)phenyl]-1H-pyrazol-1-ylbenzoic acid derivatives as potent anti-Gram-positive bacterial agents. Many of the evaluated compounds are potent growth inhibitors of Gram-positive bacteria and showed low toxicity of human cultured cells. Among the compounds, 191a and 191b exhibited excellent inhibition against Staphylococcus aureus. A novel series of multi-substituted benzo-indole pyrazole derivatives with antibacterial activity targeting DNA gyrase has been investigated [85]. Compound 192 exhibited excellent antibacterial activity against four drug-resistant E. coli bacteria strains (E564e, E68d, E48e, and E109, respectively) with IC50 values of 7.0 and 17.0, 13.5, and 1.0 μM, respectively. The substitution of fluorine or chlorine at R2 enhanced the bacteriostatic effect. The derivative bearing a Cl atom at R1 and R2 exhibited a superior antibacterial effect. Additionally, compound 192 displayed potent inhibition against DNA gyrase, with IC50 values of 0.10 μM in the in vitro enzyme inhibitory assay. A library of twenty-three novel pyrazole–phenylthiazole hybrids was synthesized and screened for antimicrobial activity against five bacterial species and two fungi [86]. Compound 193 displayed a promising antibacterial effect against the Gram-positive methicillin-resistant Staphylococcus aureus (MRSA) strain with a MIC value of 4 μg/mL. Compound 193 was nontoxic to mammalian cells—namely, human embryonic kidney cells and human red blood cells. All the synthesized compounds except 194 (moderate growth inhibition of 40.8%) showed poor antifungal activity. Analogs of pyrazole–thiazolidinone and pyrazole–thiosemicarbazone were designed using a molecular hybridization approach and further synthesized through a Vilsmeier–Haack approach [87]. The compounds were tested for antimicrobial activity against two Gram-positive bacteria, such as Staphylococcus aureus and methicillin-resistant Staphylococcus aureus. Additionally, four Gram-negative bacteria such as Escherichia coli, Salmonella typhimurium, Klebsiella pneumonia, and Pseudomonas aeruginosa were used in the biological assay. Derivatives 195 and 196 appeared as the most active antimicrobial compounds, with an MBC value of <0.2 μM against MRSA and S. aureus. The presence of 2,4-dichloro group on 195 enhanced its antibacterial activity. A new pyrazole containing isonicotinoyl derivatives from substituted chalcones and isoniazid by using sulfamic acid and their pharmacological activity evaluation have been investigated [88]. All examined compounds showed inferior activity against E. coli. The MIC values divulged that compounds 197a, 197c, and 197d (MIC values = 14, 17, and 14 μM) exhibited good antimicrobial activity against Staphylococcus aureus, while compounds 197a and 197b (MIC values = 14 and 29 μM) displayed superior antimicrobial activity against Pseudomonas aeruginosa. Compounds 197b and 197d (MIC values = 117 and 114 μM) exhibited noticeable activity against Salmonella typhi. Notably, the electro-donating group at the R position improved the antimicrobial activities more than the insertion of the electro-withdrawing group. Additionally, electron-donating substituents at position three enhanced the antimicrobial activity compared to position four. The one-pot reaction of bis-hydrazonoyl bromide with active methylene reagents furnished new bis-thiazolyl-pyrazole derivatives [89]. The insertion of acetyl (COCH3) and methyl (CH3) groups improved the
activity of compound 198 against Gram-positive strains with MIC values 2, 8, and 8 µM towards S. aureus, B. subtilis, and E. faecalis, respectively. In comparison, 198 showed moderate antifungal and no substantial activity against Gram-negative strains with MIC > 32 µM. The most potent compound against the Gram-positive bacterial strains was 200 with MIC values of 0.12, 1, and 0.5 µM for S. aureus, B. subtilis, and E. faecalis, respectively, compared to the standard drug vancomycin (1 to 2 µM). The lipophilic aryl substituent at position five and cyano at position four in the pyrazole ring improved the antibacterial activity of 200. Compounds 198, 199b, and 200 exhibited more potent inhibitory activity of DHFR with IC₅₀ values (6.34 ± 0.26, 7.49 ± 0.28, and 3.81 ± 0.16 µM), respectively, compared with Trimethoprim (8.34 ± 0.11 µM). However, bis-1-(thiazol-2-yl)-5-(amino)-1H-pyrazole-4-carbonitrile derivative 199a was revealed to be the least inhibited toward DHFR in comparison to Trimethoprim and other tested derivatives, with an IC₅₀ value 19.38 ± 0.68 µM, and that may be related to the presence of the carboxamide group in position four at the pyrazole ring rather than acetyl or carbonitrile as pyrazole derivatives 198, 199b, and 200. Desai and colleagues [90] synthesized analogs of pyrazole, pyrazoline-clubbed pyridine compounds, and examined their antibacterial and antifungal activities. Among the test compounds, 201a (3-OH) and 201b (4-F) exhibited good activity (MIC = 50 µg/mL) against S. aureus and E. coli, respectively. Compound 201c (2,4-dichloro) showed superior activity (MIC = 12.5 µg/mL) against P. aeruginosa and very good activity (MIC = 25 µg/mL) against S. pyogenes compared to the standard drug ampicillin (100 µg/mL) and chloramphenicol (50 µg/mL). The derivatives bearing electron-donating groups (2-OH, 3-OH, 4-CH₃, and 4-OH-3-OCH₃) showed significant antifungal and antibacterial activity, while the derivatives bearing electron-withdrawing groups (4-F and 2,4-dichloro) showed an augmentation in the antibacterial potency. Compound 202 has been reported as a promising antibacterial agent (B. cereus, S. aureus, P. aeruginosa, and E. coli) with the closest inhibition zones (17–20 mm) [91] (see Figure 9).
Figure 9. Structures of promising pyrazole derivatives with antibacterial activity.
4.4. Antifungal

Compounds 201d (4-CH₃) and 201e (4-OH-3-OCH₃) showed significant antifungal activity towards diverse fungal strains [90]. Compound 201d (4-CH₃) displayed significant activity (MIC = 12.5 µg/mL) towards A. niger, and compound 201e (4-OH-3-OCH₃) showed excellent activity (MIC = 12.5 µg/mL) against C. albicans and A. clavatus (Figure 9). Othman et al. [92] reported novel heterocyclic hybrids of pyrazole and their antimicrobial activity. Compounds bearing a benzenesulphonamide group fused with 3-methyl-5-oxo-1H-pyrazol-4(5H)-ylidene) hydrazine 203 and 6-amino-7-cyano-3-methyl-5H-pyrazolo [4,3-c]pyridazine 204 showed significant and broad-spectrum antimicrobial activity. Compound 204 displayed excellent antifungal activity toward A. fumigates with a MIC value of 0.98 µg/mL. While 203 and 205 are equipotent with the reference, amphotericin B against A. fumigates with a MIC value of 1.95 µg/mL exhibited 2-fold decrease in the effectiveness compared to the standard, ciprofloxacin against S. pneumonia (MIC = 1.95 and 0.98 µg/mL, respectively). Compound 203 was equipotent in the antifungal activity with the reference (MIC = 3.9 µg/mL) against C. albicans.

A series of novel pyrazole-thiazole carboxamides were designed, synthesized, and investigated for their antifungal activity [93]. The outcomes showed that compounds 206, 207, and 208 have promising in vitro activities against Rhizoctonia cerealis, with EC₅₀ values from 1.1 to 4.9 mg/L, superior to thifluzamide (EC₅₀ = 23.1 mg/L). The antifungal activity of 207 (EC₅₀ value = 1.1 mg/L was ~21-fold more active than thifluzamide and ~ 2-fold more active than compound 206 (EC₅₀ = 2.0 mg/L). Meanwhile, 208 exhibited excellent antifungal activity towards S. sclerotiorum, with an EC₅₀ value of 0.8 mg/L, which was ~6-fold higher than thifluzamide (EC₅₀ = 4.9 mg/L). The conjugates bearing an aniline moiety with a single substituent at the ortho- or meta-position exhibited promising antifungal activity. Additionally, the in vivo antifungal assay showed that 206 (90% at 10 mg/L) exhibited higher antifungal activity than thifluzamide against R. solani (90% at 10 mg/L). The synthetic route to pyrazole-4-formylhydrazine derivatives bearing a diphenyl ether fragment was reported by Wang et al. [94]. The synthesized compounds were evaluated for their antifungal activity by targeting a succinate dehydrogenase. Among the tested compounds, 209a against Rhizoctonia solani, 209b against Fusarium graminearum, and 209c against Botrytis cinerea, exhibited superior antifungal activity. The compounds displayed EC₅₀ values of 0.14, 0.27, and 0.52 µg/mL higher than carbendazim against R. solani (0.34 µg/mL) and F. graminearum (0.57 µg/mL), along with penthiopyrad against B. cinerea (0.83 µg/mL). Compound 209a was ~2- and 15-fold higher than the marketed fungicides carbendazim (0.34 µg/mL) and boscalid (2.21 µg/mL) towards R. solani. The results from the determination of the inhibitory effects of 209a against the SDH collected from the mycelia of R. solani showed IC₅₀ values of 3.99 µM (1.58 µg/mL). The in vivo anti-R. solani control effectiveness of the most potent compound, 209a (73.25% at 200 µg/mL), was significantly superior to carbendazim under the same conditions (59.81%). Compound 210 has been reported as an excellent antifungal agent with equivalent activity to the marketed fungicide drug thifluzamide, and its EC₅₀ value was 0.022 mg/L against R. solani [95] (see Figure 10).
Figure 10. Pyrazole derivative with antifungal activity.
A synthetic route to substituted 3-(trifluoromethyl)-4,5-dihydro-1H-furo [2,3-c] pyrazole conjugates using the [3 + 2] Michael/Alkylation approach was developed by Tan et al. [96]. The antifungal activity of the synthesized compounds was further examined, and 211a exhibited excellent antifungal activity against A. solani with IC₅₀ values of 5.44 µM. Compounds 211b and 212 also displayed good antifungal effects, with corresponding IC₅₀ values of 9.00 and 31.45 µM, respectively. The antifungal effects of the most active compounds 211a and 211b and 212 were relatively superior to the standard compound cycloheximide (IC₅₀ = 71.00 µM). The conjugates bearing the electron-deficient group on the aromatic ring displayed higher inhibitory effects. At the same time, the compound bearing an electron-donating substituent on the aromatic ring showed reduced inhibitory effects. Compounds 213, 214a and 214b, and 215 have been unveiled to exhibit good antifungal activity toward Fusarium Oxysporum f. sp. and Albedinis (FOA) fungus with IC₅₀ values ranging from 25.6 to 33.2 µg/mL [97]. Compound 216 has been reported to have superior antifungal inhibitory activity against C. albicans (MIC ≤ 146 µg/mL) compared to 217, which displayed a MIC = 183 µg/mL. Meanwhile, 218a and 218b exhibited significant antifungal effectiveness concentrations against C. albicans compared to the reference compound cycloheximide with the MIC values of 168 µg/mL and 165 µg/mL, respectively [98]. Piperazine-pyrazole-4-carboxylic acids have shown good antifungal inhibitory effects. Compounds 219a–219e showed equipotent antifungal activity with the reference miconazole against C. albicans (MIC value = 78.1 µg/mL) [99] (see Figure 10).

Dong and coworkers [100] synthesized a series of novel pyrazole-4-carboxamides hybrids and further evaluated their antifungal activity (see Figure 10). Compound 220 was the most potent compound against A. solani in vitro, with an EC₅₀ value of 3.06 µg/mL. It displayed 100% (10 µg/mL) inhibitory activity against A. solani in vivo compared to the standard drug boscalid. Makhanya et al. [101] revealed compounds 221a and 221b as promising antifungal agents. The antifungal assay showed that 221a and 221b exhibited significant inhibitory activity against Saccharomyces cerevisiae (zone inhibition (ZI) = 23 and 20 mm, respectively), with a MIC value of 0.18 µM. Comparable with the standard drug amphotericin, B. Bayazeed et al. [102] detected four chromen-3-yl-pyrazole derivatives: 222a, 222b, 223a, and 223b to have superior antifungal activity. Compared to the standard drug ketoconazole against Aspergillus fumigatus with the corresponding % ZI values of 164%, 147.1%, 158.8%, and 147.1%, respectively. In addition, compounds 222b (% ZI value = 150%) and 223b (% ZI value = 150%) have 1.5-fold superior activity against C. albicans compared to the reference ketoconazole (% ZI value = 150%). Notably, the insertion of two ester groups in compound 222a improved its antifungal activity. Meanwhile, incorporating electron-donating groups (OCH₃ and CH₃) at the para-position of the aromatic ring in 222b, 223a, and 223b enhanced their antifungal activity. Wang et al. [103] reported a novel series of pyrazole-4-acetyhydrazide derivatives targeting fungal SDH, further evaluating their antifungal properties towards R. solani, F. graminearum, and B. cinerea. Among the evaluated compounds, the antifungal activity of 224a against R. solani, 224b against F. graminearum, and 224c against B. cinerea had EC₅₀ values of 0.27, 1.94, and 1.93 µg/mL, respectively. These values were superior to the standard reference boscalid against R. solani (0.94 µg/mL) and fluopyram against F. graminearum (9.37 µg/mL) and B. cinerea (1.94 µg/mL). Additionally, the compounds with hydroxyl group substituents at the R₁ position displayed higher anti-R. solani activity than the corresponding conjugates bearing an ethoxy group substituent. The in vivo studies detailed that compound 224a was effective toward R. solani (79.83% at 200 µg/mL), comparable to validamycin (86.56%) and thifluzamide (83.49%). Compound 224a was predicted as an SDH inhibitor (see Figure 11).
A series of new pyrazole-4-carboxamide conjugates were designed and synthesized by Wu et al. [104]. The synthesized compounds were evaluated for their antifungal activity using four phytopathogenic fungi (G. zeae, F. oxysporum, C. mandshurica, and P. infestans). The EC$_{50}$ values were 1.8 µg/mL for 225a against G. zeae, 1.5 and 3.6 µg/mL for 225b against F. oxysporium and C. mandshurica, respectively, and 6.8 µg/mL for 225c against P. infestans. Meanwhile, the SDH enzymatic effectiveness unveiled corresponding IC$_{50}$ values of 6.9, 12.5, 135.3, and 223.9 µg/mL, for 225c, 222d, 225e, and penthiopyrad, respectively. Incorporating substituents (CH$_3$, F, or Cl) into the 2-phenyl and methyl into 2-pyridinyl positions enhanced the antifungal activity. While the introduction substituent into the 3-phenyl and 3-pyridinyl positions decreased the antifungal properties. Xia et al. [105] reported novel pyrazole carboxylate derivatives bearing thiazole as potent fungicides. The antifungal studies revealed compound 226 displayed superior activities against Botrytis cinerea and Sclerotinia sclerotiorum, with EC$_{50}$ values of 0.40 and 3.54 mg/L, respectively. Compound 227 displayed superb antifungal activity against Valsa mali, with an EC$_{50}$ value of 0.32 mg/L. The in vivo fungicide control studies against B. cinerea and V. mali revealed that compounds 226 and 228 at 25 mg/L, respectively, were influential on cherry tomatoes and apple branches. Compound 227 displayed an inhibitory activity toward SDH, with an IC$_{50}$ value of 82.26 µM. Nevertheless, compounds 226 and 228 lack inhibitory activity toward SDH in the in vivo studies (see Figure 11).
4.5. Antidiabetics

Thiazolindindione 128a and 128b and thiazolidinone derivatives 129a and 129b displayed significant inhibitory activities against α- and β-glucosidase (% inhibitory activity = 62.15, 55.30, 65.37, and 59.08 for α-glucosidase and 57.42, 60.07, 58.19, and 66.90 for β-glucosidase, respectively) than the reference compounds: acarbose with % inhibitory activity = 49.50 for α-glucosidase and D-saccharic acid 1,4-lactone monohydrate with % inhibitory activity = 53.42 for β-glucosidase. Compared to pioglitazone and rosiglitazone, the potent compounds showed good PPAR-γ activation and hypoglycemic effectiveness [42]. Kattimani et al. [106] reported the ring alteration of 3-arylsydnone into 1-aryl-1H-pyrazole-3-carbonitriles via a [3 + 2] cycloaddition reaction and were subsequently converted into 5-(1-aryl-1H-pyrazol-3-yl)-1H-tetrazole. The synthesized compounds were screened for in vivo antihyperglycemic activity using albino Wistar rats. Compounds 229a and 229b and 230a–230d pointedly reduced the blood glucose levels and prevented vascular difficulties in streptozotocin-induced diabetic rats (Figure 12). Compounds 231 and 232 were potent inhibitors of the α-amylase enzyme [107]. Compound 231 showed excellent activity against α-amylase, with an IC50 value of 4.08 µg/mL, followed by 232 with an IC50 value of 7.59 µg/mL. The potency of the compounds was superior to acarbose (IC50 value = 8.0 µg/mL). Pogaku et al. [108] designed and synthesized new pyrazole–triazolopyrimidine hybrids as potent α-glucosidase inhibitors using a one-pot multicomponent approach. Among the evaluated compounds, 233a–c bearing an electron-withdrawing group on the phenyl ring displayed significant inhibitory activity against the α-glucosidase enzyme. Compound 233a with the 4-Cl group showed the highest inhibition, with an IC50 value of 12.45 µM, equipotent to the standard drug acarbose (IC50 value = 12.68 µM). In contrast, 233b with a fluoro substitution at the para position displayed an IC50 value of 14.47 µM, followed by 233c bearing a 4-NO2 group (IC50 value 17.27 µM). Karrouchi et al. [109] designed and synthesized a pyrazole-3-carbohydrazide, 234. The in vitro α-glucosidase inhibition study of 234 showed good activity for a concentration of 0.08 mM with a percent inhibition of 79.83%, superior to acarbose (29%). The β-galactosidase evaluation displayed a good inhibitory activity with a percentage of 64.6%, comparable to quercetin (68%) for a concentration of 3.30 mM, while the α-amylase inhibition results revealed an inhibitory activity of 20.51% comparative to the acarbose with a percentage of 36% for a concentration of 3.53 mM. The Rhodanine–pyrazole conjugates were designed and synthesized by Singh et al. [110]. The compounds were further tested for their antidiabetic activity. Among the evaluated compounds, 235a (IC50 = 2.259 × 10−6 mol/L) was the most potent compound, 42-fold superior to acarbose. The unsubstituted hybrid 235b (IC50 = 2.854 × 10−5 mol/L) was 3-fold superior to acarbose. Meanwhile, 235c (IC50 = 6.377 × 10−5 mol/L) and 235d (IC50 = 1.325 × 10−4 mol/L) exhibited strong inhibitory activity against α-amylase comparable to acarbose. Compound 236 has been reported to showed a promising bifunctional antidiabetic effect [111]. A series of new benzo[d][1,2,3]triazol-1-yl-pyrazole-bearing dihydro-[1,2,4] triazolo[4,3-a]pyrimidine groups have been designed and synthesized [112]. All synthesized compounds were screened in vitro for α-glucosidase inhibition, anticancer (A549 and MCF-7 cell lines), and antioxidant studies. Among all the compounds tested for antidiabetic potential, 237a, 237b, and 237c exhibited substantial inhibition activity, with IC50 values of 20.12 ± 0.19 µM, 21.55 ± 0.46 µM, and 24.92 ± 0.98 µM, respectively, compared to the reference compound acarbose (IC50 = 12.68 µM) (see Figure 12).
Figure 12. Structures of pyrazole hybrids with antidiabetic activity.
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Kaur et al. reported a series of novel isatin–pyrazole hybrids and tested their antidiabetic activity [113]. Among the tested compounds, 238 (Figure 12) appeared to be most potent with IC$_{50}$ = 3.26 ± 0.25 µM, which was ~146-fold more potent than acarbose (IC$_{50}$ = 478.07 ± 1.53 µM). Kausar and coworkers [114] synthesized Celebrex derivatives and investigated their antidiabetic effectiveness. Many of the evaluated compounds exhibited good activity. Compound 239 emerged as the most potent inhibitor of the α-glucosidase enzyme (IC$_{50}$ = 92.32 ± 1.530 µM), comparable to the standard drug acarbose (IC$_{50}$ = 875.75 ± 2.08 µM). Compound 240 exhibited excellent antidiabetic activity (IC$_{50}$ = 5.8 µM) compared to the reference acarbose (IC$_{50}$ = 58.8µM) and other evaluated compounds (241, IC$_{50}$ = 65 µM and 242, IC$_{50}$ = 103 µM) under the same conditions [115]. Shen described a series of novel pyrazolo [1,5-a]pyrimidine derivatives as promising and selective DPP-4 inhibitors [116]. Compound 243 was 2-fold superior to alogliptin (IC$_{50}$ = 4 nM) and notably selective over DPP-8 and DPP-9 (>2000-fold). The in vivo IPGTT assays in diabetics showed that 243 significantly lowers blood sugar by 48% at 10 mg/kg. The in vitro antidiabetic potential of compound 244 was evaluated against α-glucosidase and α-amylase enzymes. The results revealed that 244 with IC$_{50}$ = 60.45 ± 1.23 µM showed superior α-glucosidase effectiveness relative to acarbose (IC$_{50}$ = 89.12 ± 2.08 µM) [117]. Nevertheless, compound 244 was inactive against α-amylase (see Figure 12).

4.6. Antileishmanial

The incorporation of a heteroaromatic ring coupled with a 1,3,4-oxadiazole moiety improved the antileishmanial activity. Compounds 245, 246, and 247 (Figure 13) proved the dose-dependent killing of the promastigotes with corresponding IC$_{50}$ values of 33.3 ± 1.68, 40.1 ± 1.0, and 19.0 ± 1.47µg/mL, respectively [118]. Additionally, the compounds (245, 246, and 247) displayed IC$_{50}$ values of 44.2 ± 2.72, 66.8 ± 2.05, and 73.1 ± 1.69 µg/mL, respectively, on amastigote infectivity. These compounds depicted a comparable point in dose-dependent parasite killing with the standard drug, pentamidine (IC$_{50}$ = 2.6 ± 0.32 µg/mL). Camargo et al. synthesized a series of novel pyrazole hybrids [119]. The hybrids were investigated in vitro against the promastigote of Leishmania amazonensis. At the same time, the hybrids were examined against the epimastigote of Trypanosoma cruzi (T. cruzi). The S-methyl thiosemicarbazones 248a–248c and 2-amino-1,3,4-thiadiazole pyrazole hybrids 249a–249c displayed significant antileishmanial and antitrypanosomal properties. The substitution of Br, OCH$_3$, or NO$_2$ at the para position of the aryl ring attached at position five of pyrazole favored their activity. The substituent attached to position three of the pyrazole ring also influenced the activity of the evaluated compounds. Silva et al. synthesized and screened a series of 1,5-biaryl 3-arylaminomethyl 4-carboxyethyl pyrazoles and screened against L. amazonensis and T. cruzi [120]. The most active compounds 249, 250, and 251 demonstrated similar profiles against both L. amazonensis and T. cruzi parasites, describing their dual activity. Meanwhile, compound 249 induced morphological and ultrastructural alterations in the promastigote of L. amazonensis (see Figure 13).
4.7. Antimalarial

Compound 252 (Figure 13) has been demonstrated to be an excellent inhibitor of Falcipain-2, with an IC$_{50}$ value of 14 µM [118]. Akolkar et al. [121] have examined the antimalarial potency of 253, 254, and 255. Compounds 253 and 254 (IC$_{50}$ = 0.47 µM) were equipotent regarding efficacy compared to the standard drug quinine (0.83 µM). The inhibition activity of 255 (0.21 µM) was 4-fold superior to the standard drug. Molecular hybrids of the thiophene, pyrazoline, and benzene rings enhanced the antimalarial activity. Strašek and coworkers [122] reported the synthetic route of the tetrahydropyrazolo [1,2-a]pyrazole-1 carboxylates. The reactions yielded mixtures of 7-oxo-2,3,5,6 tetrahydropyrazolo [1,2–a]pyrazole-1-carboxamides 256 and 257. An assessment inhibition of dihydroorotate dehydrogenase of *Plasmodium falciparum* (*Pf*DHODH) was demonstrated. The strongest potency was found in compound 257 with an IC$_{50}$ value of 2.9 ± 0.3 µM. All the evaluated compounds developed selectivity for *Pf*DHODH more than *Hs*DHODH. Gogoi et al. [123] reported dimethoxy pyrazole 1,3,5-triazine derivatives as a novel class of potent antimalarial agents with good toxicity profiles. Within the series compound 258 was observed as a promising antimalarial agent (see Figure 14).
4.8. Antioxidant

Antioxidant activities of pyrazoline derivatives were screened using the 2,2-diphenyl-1-picrylhydrazyl (DPPH) radical scavenging method. All the tested compounds showed antioxidant activity [88]. Compound 197c bearing 4-fluoro and 4-methyl substituents was the most potent antioxidant agent among all the tested compounds at all the concentrations. Compound 234 has been reported as a promising antioxidant agent [109]. Compounds 237d and 237e have been revealed as having effective antioxidant activity (IC_{50} values = 4.25 µM and 5.40 µM, respectively) [112]. The results of an evaluation of synthesized thiazolidine-2,4-dione-pyrazole conjugates as antioxidant agents showed the efficacy of all the examined compounds [107]. Compounds 259a and 259b and 260 (see Figure 14) showed the most potent results, with IC_{50} values of 110.88, 127.18, and 128.55 µg/mL, respectively. The standard drug ascorbic acid showed an IC_{50} value of 81.12 µg/mL. The synthesis of functionalized pyrano[2,3-c]pyrazoles and pyrazolopyran[2,3-d]pyrimidines containing a bioactive chromone moiety has been achieved, along with their antioxidant activity [124]. The in vitro antioxidant activity was determined using DPPH radical scavenging methods. Among the tested hybrids, 261–264 displayed promising activity with all the concentrations in the evaluation with the reference drug. The hybridization of pyranopyrazole with the pyrimidine moiety with substituted NH and OH groups improved the antioxidant properties. Ali et al. [125] reported the synthesis of a novel series of pyrazoline 269a–269e, phenylpyrazoline 270a–270e, isoxazoline 271a–271e, and pyrazoline carbothioamide derivatives 272a–272e using chalcones as a precursor 265a–265e. The hybrid compounds were vetted for in vitro antioxidant activity using DPPH, nitric oxide (NO), and the superoxide radical scavenging (RSA) assay, along with 15-lipoxygenase (15-LOX) inhibition activity. Pyrazoline carbothioamide derivatives 272a and 272e were the most potent anti-LOX compounds, 2.2- and 2.1-fold superior to quercetin, while compounds 269a, 270e, 271b, 271c, 272a, 272c, and 272e exhibited substantial RSA in all the three in vitro assays relative to the ascorbic acid, along with 15-LOX inhibition potency. The presence of electron-donating groups (CH_3 and OCH_3) or halogens (di-Cl) on the benzene ring enhanced the inhibition activity. The potential antioxidant activity of 272a and 272e were comparable in all three assays. Compounds 271b, 271c, and 271e (Figure 15) showed significant in vivo antioxidant potential compared to the standard group at a dose of 100 mg/kg B.W. Meanwhile, there was an increase in CAT activity, the GSH level, and a decrease in lipid peroxidation in the treated rat liver compared to the control treatment. The in vitro antioxidant effectiveness of 4-(arylmethacetyl)-1H-pyrazoles bearing sulfur or 1H-pyrazole groups has been investigated in different assays by Oliveira et al. [126], along with their oxidative stress impacts in biological systems. Compounds 273 and 274 showed significant inhibition in the ABTS assay, revealing that the mechanisms of the antioxidant action of compounds 273 and 274 were connected to their ability to donate electrons. Additionally, compounds 273 and 275 are more potent in the NO scavenging assay, while 274 reduced the lipid peroxidation levels in the brain and the liver after 72 h of treatment remarking on the compound efficacy in oxidative stress. A new series of pyrazole-containing heterocyclic skeletons—namely, pyrimidine, triazole, triazepine, pyrrole, and thiadiazolopyrimidine—along with acylthiourea derivatives, were synthesized from 2-cyano-3-pyrazolylpropenoyl isothiocyanate by Badawy et al. [127]. The antioxidant screening of all the synthesized compounds showed that pyrimidinethione derivatives 276 and 277 were the most potent antioxidant agents. El-Borai et al. [128] achieved a biological evaluation of the cytotoxicity, antimethemolytic and antioxidant activities of some thienopyrazole compounds. The antioxidant activity of the examined compounds was achieved by utilizing the DPPH radical scavenging assay with ascorbic acid as the reference. Compound 278 exhibited excellent radical scavenging activity, with an IC_{50} value of 4.49 µg/mL comparable to an IC_{50} of 4.76 µg/mL. The excellent antioxidant result was obtained due to the existence of the two amino groups on the pyrimidine ring. Additionally, 279 exhibited strong antimethemolytic and antioxidants, justifying that the antioxidant activity may protect red blood cells from hemolysis. The insertion of chlorine atoms, hydroxyl, and
cyanide with a pyrimidine ring in a single moiety enhanced the activity of 279. In addition, 280 was noxious to all the tested cancerous cell lines; however, a lower cytotoxicity activity against the normal fibroblast cell line was observed. Elnagdy and colleagues [129] described a synthetic route to pyrazole analogs by using copper oxide nanoparticles (CuO-NPs) as catalyzed. The compounds were evaluated for their antioxidant activity using the DPPH radical scavenging assay. Most of the compounds tested demonstrated a greater interaction with the DPPH radical relative to the standard compound Trolox (IC$_{50}$ = 11.48 mM). The compounds 281, 282, and 283 showed maximum antioxidant activity in the order of 282 > 281 > 283, with IC$_{50}$ values of 3.06, 3.53, and 5.42 mM, respectively. The ability of 281 and 282 to recover the DPPH radical assay resulted from the prolonged conjugation in compounds, while that of compound 283 was due to a phenolic hydroxyl group at the ortho-position and a fluorne group. The condensation reaction between 1,3-thiazole or aminopyridine derivatives and 1H-pyrazole, 3,5-dimethyl-1H-pyrazole or 1,2,4-triazole was described by Kaddouri and colleagues [130]. The reaction produced novel heterocyclic compounds containing pyrazole, thiazole, and pyridine. Additionally, the DPPH scavenging assay was utilized to investigate their antioxidant activity. Ligand 284 showed the best antioxidant activity, with an IC$_{50}$ value of 4.67 µg/mL, while the IC$_{50}$ value for the reference compound was 2 µg/mL (ascorbic acid). The applicable route for the direct synthesis of (E)-ethyl 2-benzylidene-3-oxobutanoate through the 3 + 2 annulation method, including the investigated in vitro antioxidant vulnerabilities through the DPPH and hydroxyl radical scavenging methods of this compound, have been reported [131]. The assays showed that compound 285 has a strong antioxidant power (see Figure 16).

The multicomponent reaction of some heterocyclic compounds with activated acetylenic, alkyl bromides, triphenylphosphine, and hydrazine in water under ultrasonic irradiation yielded pyrazole derivatives in better yields [132]. Additionally, the antioxidant activities of the compounds were examined using DPPH radical scavenging and the ferric-reducing power assay. Compound 286 (Figure 16) exhibited exceptional DPPH radical scavenging activity and greater reducing power compared to the standard reference butylated hydroxytoluene (BHT) and 2-tertbutylhydroquinone (TBHQ). Compounds 287 and 288 have been reported as more potent antioxidant inhibitors than ascorbic acid and butylated hydroxyanisole (BHA) [133]. The corresponding IC$_{50}$ values for 287 and 288 from the DPPH radical assay were 0.245 ± 0.01 and 0.284 ± 0.02 µM, respectively. These compounds have more potent RSA than ascorbic acid (IC$_{50}$ = 0.483 ± 0.01 µM). In the hydroxyl radical scavenging assay, compounds 287 and 288 showed IC$_{50}$ values of 0.905 ± 0.01 µM and 0.892 ± 0.01 µM, respectively. They displayed greater RSA than BHA (IC$_{50}$ = 1.739 ± 0.01 µM).

Patil et al. [134] prepared sulfonic acid functionalized 1,4-diazabicyclo[2.2.2]octane assisted on Merrifield resin, [MerDABCO-SO$_3$H]Cl as a catalyst to synthesized pyrazolopyranopyrimidines in one-pot four-component reactions in an excellent yield. The antioxidant effect of the synthesized compounds was determined using the 1,1-diphenyl-2-DPPH radicals scavenging assay, and ascorbic acid was used as a standard control. Among the evaluated compounds, 289–292 showed excellent antioxidant activity compared to the standard ascorbic acid due to the incorporation of an electron-withdrawing substituent (nitro group) on the phenyl ring, enhancing the resonance impact stabilizing the consistently formed radical. Compounds 293 and 294 have been reported as promising antioxidant agents [135] (see Figure 16).
Figure 15. Structures of promising pyrazole derivatives with antioxidant activity.
Figure 16. Structures of pyrazole hybrids with antioxidant activity.

4.9. Antituberculosis

Jagadale et al. [136] described the pathway to synthesize thiazolyl-pyrazolyl-1,2,3-triazole derivative 295 and bis-pyrazolyl-1,2,3-triazole 296 derivative, along with their
antimycobacterial activity against *M. tuberculosis* (*Mt*b) with H37Ra dormant and active. The antimycobacterial activity revealed that most of the compounds showed moderate to good activity against both strains of *M. tuberculosis*. Compounds 295a–295c and 296a and 296b exhibited good activity against the *Mt*b H37Ra active strain; also, compounds 294d and 295e and 296c–296e displayed good activity against the *Mt*b H37Ra dormant strain. Using Pd/Cu catalyst coupling-cyclization strategy, 3-indolylmethyl substituted (pyrazolo/benzo) triazinone derivatives have been expediently prepared in a one-pot reaction [137]. The synthesized compounds were tested for chorismate mutase (CM) inhibitory properties in vitro using an assay that measured the enzyme’s catalytic activity (MC) in converting chorismate (substrate) to prehenate. The best active compounds, 297 and 298, showed 78% inhibition at 30 µM. Meanwhile the concentration-dependent evaluation resulted in IC<sub>50</sub> values of 0.40 ± 0.05 µM and 0.85 ± 0.10 µM for compounds 297 and 298, respectively. Compound 299 has also been shown to maintain good potency against clinical samples from the four main lines and strains containing isoniazid or rifampin resistance mutations [138]. The mutated strains in MmpL3 were resistant to 299 and under replication conditions, and it exhibited bactericidal activity against *Mt*b. However, compound 299 was not effective in an acute model of tubercular infection. This is likely the result of in vivo exposure remaining above the minimum inhibitory level for a restricted period (see Figure 17).

![Figure 17. Structures of pyrazole hybrids with antitubercular activity.](image)

Hu and coworkers [139] reported novel series of pyrazolo [1,5-alpyridine-3-carboxamide (PPA) conjugates bearing diaryl side chains and their antitubercular activity. Most of the evaluated compounds were highly potent in vitro against *Mt*b strains, including H37Rv (MIC = < 0.002–0.381 µg/mL), rINH (MIC = < 0.002–0.465 µg/mL), and rRMP (MIC = < 0.002–0.004 µg/mL). Notably, compound 300 demonstrated favorable in vitro activity against *Mt*b H37Rv, rRMP, and rINH, with corresponding MIC values ≤ 0.002 µg/mL and a lack of toxicity against Vero cells. Moreover, in vivo studies showed that 300 sub-
stantially lessened the mycobacterial load in a mouse model infected with H37Ra. Other reported antituberculosis pyrazole derivatives and the corresponding references are shown in Table 1.

Table 1. Reported antituberculosis pyrazole derivatives, along with the corresponding references.

| Compound | MIC (µg/mL) | Reference |
|----------|-------------|-----------|
| 302a     | 0.52        | [140]     |
| 302b     | 0.50        | [140]     |
| 302c     | 0.79        | [140]     |
| 302d     | 1.48        | [140]     |
| 302e     | 0.53        | [140]     |

MIC (µg/mL) = 5 against H37RV strain

| Compound | IC50 (µg/mL) | MIC (µM) |
|----------|-------------|----------|
| 302a     | 3.96        |          |
| 302b     | 3.67        |          |

MIC (µM) = 3.96 and 3.67 µM against H37RV
Table 1. Cont.

![Chemical structure 304](image)

MIC(µM) = 6.25, against H37RV

![Chemical structure 305](image)

MIC (µg/mL) = 25 against H37RV

![Chemical structure 306](image)

H37RV MIC(µg/mL)

| Structure  | R1 | R2 | MIC (µg/mL) |
|------------|----|----|-------------|
| 306a       | 4-F-C6H4, 3,4-di-F-C6H4 | 0.8 |
| 306b       | 4-CH3-C6H5 | 4-Cl-C6H5 | 3.12 |
| 306c       | 2-OCH3-C6H5 | 3-CH3-C6H5 | 3.12 |
| 306d       | H | 3,4-di-Cl-C6H4 | 6.25 |

![Chemical structure 307](image)

MIC(µg/mL) H37RV

| Structure  | R1 | R2 | MIC (µg/mL) |
|------------|----|----|-------------|
| 307a       | H | Cl | 0.78 |
| 307b       | 4-Br | H | 1.56 |
| 307c       | 4-Cl | Cl | 1.56 |
| 307d       | 4-Br | Cl | 1.56 |
| 307e       | 4-Cl | Cl | 1.56 |
| 307f       | 4-Cl | Br | 1.56 |
| 307g       | Br | Br | 1.56 |
Table 1. Cont.

|          |          |          |          |
|----------|----------|----------|----------|
| 308a     | R = H    | 86       |          |
| 308b     | R = COPh | 85       |          |
| 309a     | R = H    | 90       |          |
| 309b     | R = COPh | 88       |          |

**Growth Inhibition (%GI) H37R<sub>V</sub>**

![Chemical structures](image)

**MIC(µg/mL) H37R<sub>V</sub>**

|          |          |
|----------|----------|
| 310a     | isobutyl |
| 310b     | tert-butyl |

|          |          |
|----------|----------|
| 1.562    |          |

**MIC(µg/mL) = 1.6, against H37R<sub>V</sub>**

**MIC(µM) = 17, and MBC(µM) = 34 against H37Ra**

![Chemical structures](image)
Table 1. Cont.

![Chemical Structure](image)

| Compound | MIC (µg/mL) | H37Ra | M. bovis BCG |
|----------|-------------|-------|-------------|
| 314      | 2.96        | 0.20  |             |
| 314      | 1.16        | 0.72  |             |
| 314      | 4.65        | 2.3   |             |
| 314      | 2.5         | 2.26  |             |
| 315      | 2.54        | 0.51  |             |
| 315      | 1.72        | 1.33  |             |

4.10. Agrochemical

Series of novel pyrazole–isoindoline-1,3-dione hybrids as favorable 4-hydroxyphenylpyruvate dioxygenase (HPPD) inhibitors were designed by combining 2-benzoylethen-1-ol and isoindoline-1,3-dione into a single moiety [153]. Among the evaluated compounds against *Arabidopsis thaliana* HPPD in vitro, using mesotrione and pyrasulfotole as the positive control, the IC$_{50}$ of 316 (Figure 18) was extended to 90 nM. In addition, 316 was identified as the most promising inhibitor, with a $K_i$ value of 3.92 nM, which was ~10 times superior to pyrasulfotole ($K_i = 44$ nM) and 300 times marginally superior to mesotrione ($K_i = 4.56$ nM). Jiang et al. [154] designed and synthesized novel heptacyclic pyrazolamide conjugates using the scaffold hopping approach. The insecticidal activities of all synthesized compounds were examined against *P. xylostella* in vivo at 500 mg/L. Meanwhile, the marketed insecticide—namely, tebufenozide—was used as the reference drug. Compounds 317 and 318 flaunted excellent insecticidal activities (>90%) against *P. xylostella*. Additionally, compound 317 displayed 100% insecticidal activity at the dose of 200 mg/L. The lower dose and LC$_{50}$ value of 317 (64.13 mg/L) was akin to tebufenozide (LC$_{50} = 33.83$ mg/L). Zhao et al. [155] reported a novel series of fluoro-substituted compounds bearing altered pyrazole and their anti-larvicial effects. The larvicidal activity unveiled fluoro-substituted compounds to have good to excellent activities against *M. separata* and *P. xylostella*. The corresponding LC$_{50}$ values for 320a and 320b against *P. xylostella* were $2.9 \times 10^{-6}$ mg/L and $3.1 \times 10^{-6}$ mg/L, respectively, superior to the LC$_{50}$ of chlorantraniliprole ($4.6 \times 10^{-5}$ mg/L). In addition, fluoro-substituted compounds 320a–320c bearing ether groups at position three of the pyrazole showed better inhibitory effects than compounds with halogen, amide, or ester groups substituents. The
insertion of fluorine atoms on the ethoxy group enhanced the larvicidal activity. Compound 320a exhibited the 50% larvicidal mortality against *M. separata* to 0.1 mg/L. Moreover, 320a displayed 90% larvicidal activity against *P. xylostella* at $10^{-5}$ mg/L, higher than that of chlorantraniliprole. Judge and colleagues [156] revealed substituted 3-hydroxyprozole derivative 321 as a promising herbicidal agent. Pyridylpyrazole-4-carboxamides bearing 1,3,4-oxadiazole rings were designed and synthesized by dehydration of aromatic hydrazine derivatives and formanilides [157]. The synthesized compounds were further evaluated for their insecticidal activities (*Plutella xylostella*). Among the examined compounds, 322 displayed promising activity as follows, 67%, 50%, 34%, 20%, and 17% activity at the concentrations of 100, 50, 10, 5, and 1 µg/mL, respectively (see Figure 18).

Figure 18. Structures of pyrazole a derivative acting as promising agrochemical agents.

5. Conclusions

Pyrazoles are five-membered heterocyclic compounds containing nitrogen. They are an important class of compounds for drug development; they constitute an essential class of hit compounds to develop new pharmacological agents to treat various infections of clinical primacy. With such a diverse range of biological activities, they have attracted much attention from researchers focusing on synthesizing different pyrazole analogs to developing novel and more effective drugs. This literature review documented various synthetic pathways to pyrazole derivatives and the biological potential of some pyrazole derivatives in recent years. Their biological activity properties, such as antibacterial, analgesic, anti-inflammatory, anticancer, antibacterial, antidiabetic, antioxidant, and agrochemical, were detailed in this review. The information presented in this review will assist prospective researchers in further investigating pyrazole derivatives and update scientists with promising biological activities of recently developed derivatives. Additionally, this will allow them to identify other derivatizations that can be explored. However, where the pyrazole unit itself plays a significant role in the compound’s mode of action, including cases where the pyrazole is more of a structural element, still needs to be explored. Additionally, the molecular hybridization of pyrazole with other bioactive compounds will be explored in our future work.

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