Comparative Study of the Synthetic Approaches and Biological Activities of the Bioisosteres of 1,3,4-Oxadiazoles and 1,3,4-Thiadiazoles over the Past Decade

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Abstract: The bioisosteres of 1,3,4-oxadiazoles and 1,3,4-thiadiazoles are well-known pharmacophores for many medicinally important drugs. Throughout the past 10 years, 1,3,4-oxa-/thiadiazole nuclei have been very attractive to researchers for drug design, synthesis, and the study of their potential activity towards a variety of diseases, including microbial and viral infections, cancer, diabetes, pain, and inflammation. This work is an up-to-date comparative study that identifies the differences between 1,3,4-thiadiazoles and 1,3,4-oxadiazoles concerning their methods of synthesis from different classes of starting compounds under various reaction conditions, as well as their biological activities and structure–activity relationship.

Keywords: 1,3,4-oxadiazole; 1,3,4-thiadiazole; bioisosteres; synthesis; biological activity; structure–activity relationship

1. Introduction

The nitrogen-containing five-membered heterocycles, particularly 1,3,4-oxadiazoles and 1,3,4-thiadiazoles, have received considerable attention because of their versatile biological properties and applications in drug discovery [1,2]. Based on bioisosterism between sulfur and oxygen atoms [3], 2-amino-1,3,4-thiadiazole could act as a bioisostere of 2-amino-1,3,4-oxadiazole and, therefore, produce biological properties broadly similar to those of the latter [4]. Moreover, these moieties are used as pharmacophores due to their favorable metabolic profile and ability to form hydrogen bonds [5,6].

Different substituted 1,3,4-oxadiazole molecules possess antimicrobial [7,8], antitubercular [9], antimalarial [10], analgesic [11], antiinflammatory [12], antiviral [13], antihelminthic [14], anti-severe acute respiratory syndrome coronavirus 2 (anti-SARS-CoV-2) [15], anticancer [10,16], and antioxidant activities [17]. The antioxidants prevent the onset and propagation of oxidative disorders like autoimmune, cardiovascular, neurovascular diseases [2].

The potent pharmacological activity of 1,3,4-oxadiazoles can be attributed to the presence of the toxophoric –N=C–O– linkage, which may react with the nucleophilic centers of the cell [18,19]. Some of the drugs that contain an oxadiazole nucleus (Figure 1) are Furamizole, a nitrofuran derivative with a strong antibacterial activity [20,21]; Nesapridil, a well-established vasodilating agent [22]; Raltegravir, also known as MK-0518, which was the first FDA-approved HIV-1 integrase (IN) inhibitor launched by Merck & Co to treat HIV infection [23]; Tiodazosin, an α1-adrenergic receptor antagonist used to treat hypertension [24,25]; Fenadiazole, a hypnotic drug [26,27]; Zibotentan, an anticancer...
agent [28,29]; and AZD1979, a melanin-concentrating hormone receptor 1 (MCHr1) antagonist [30].

Compared to 1,3,4-oxadiazole, the sulfur atom of 1,3,4-thiadiazole imparts improved lipid solubility, and the mesoionic nature of 1,3,4-thiadiazole makes this class of compounds show good tissue permeability [6]. Thus, the 1,3,4-thiadiazole fragment appears in many clinically used drugs (Figure 2), such as the diuretic drugs Acetazolamide and Methazolamide; Cefazolin CFZL and Cefazedone CFZD, which are first-generation cephalosporins; Atibeprone, an anti-depressant drug; antidiabetic agents Glybuthiazole and Glybuzole; antiprotozoal drug Megazol [31]; and BMS341, a glucocorticoid receptor modulator [32]. Moreover, 1,3,4-thiadiazole derivatives are reported to possess anticancer [33], antioxidant [34], antimicrobial [35], antidepressant [36], anticonvulsant [37], analgesic, and anti-inflammatory activities [38].

At the molecular level, 1,3,4-oxadiazole 1 is a thermally stable neutral aromatic molecule. It has four possible aromatic systems, 1–4: 1,3,4-oxadiazole 1, which is widely exploited for various applications; 1,3,4-oxadiazolines 2; 1,3,4-oxadiazolium cations 3; and the exocyclic-conjugated mesoionic 1,3,4-oxadiazole 4 (Figure 3) [39].

The linked substituents to 1,3,4-oxadiazole aromatic moieties normally appearing at positions C2, C5, and N3 may be represented by the following structural features 1,5–7 (Figure 4) [39].
The electronic distribution in 1,3,4-oxadiazole has been calculated by various Self-consistent Field Molecular Orbital (SCF-MO) methods [40]. Other structural parameters, the dipole moment, and data on its ultraviolet–visible (λ max calculated to be in the region 193–203 nm), NMR, NQR, and microwave spectra have been derived. Studies on 1,3,4-oxadiazole indicate a maximum positive charge at the 2-position [41]. Molecular diagrams for 1,3,4-oxadiazole, 2-phenyl- and 2,5-diphenyl-1,3,4-oxadiazole, and oligomeric
oxadiazoles have been derived, and conjugation between the rings was found to be similar to that in polyphenyls [42].

Tautomerism in 2-hydroxy 8a, 2-mercapto 8b, and 2-amino-1,3,4-oxadiazole 8c is in equilibrium with the tautomeric oxadiazolines 2a, 2b, and 2c, respectively (Figure 5) [43]. Evidence from UV, IR, 1H-NMR, and 13C-NMR spectra supports structure 2b for 1,3,4-oxadiazoline-5-thione [44].

![Figure 5. Tautomerism of 1,3,4-oxadiazole.](image)

The development of 1,3,4-thiadiazole chemistry is linked to the discovery of phenyl-hydrazines and hydrazine in the late 19th century. The first 1,3,4-thiadiazole was described by Fischer in 1882 and further developed by Bush and coworkers [45], but the true nature of the ring system was demonstrated first in 1956 by Goerdler et al. [46].

Similar to 1,3,4-oxadiazoles, the 1,3,4-thiadiazole derivatives are conveniently divided into three subclasses (Figure 6): a Aromatic systems that include the neutral thiadiazoles 9 [45].
b Mesoionic systems that are defined as a poly-heteroatomic system containing a five-membered heterocyclic ring associated with conjugated p and π electrons and distinct regions of positive and negative charges 10. Mesoionic systems are dense and highly polarizable, with a net neutral electron charge; these characteristics allow mesoionic compounds to cross cellular membranes and interact with biological targets with distinct affinities [31].
c 1,3,4-thiadiazol-2-one(thione) 11a,b [47].

![Figure 6. 1,3,4-thiadiazole ring systems.](image)

Further, the 1,3,4-thiadiazole ring system, with three heteroatoms, does not exhibit tautomerism in its fully conjugated form 9. However, when certain substituents are present, tautomerism is possible. 1,3,4-Thiadiazolin-2-ones 11a and -2-thione 11b exist in the oxo 12a and thione 12b forms, respectively [47]. 2-Amino-1,3,4-thiadiazoles exist in the amino form 13 in solution and in the solid state (Figure 7) [48].

![Figure 7. Tautomeric forms of 2-substituted 1,3,4-thiadiazoles.](image)

Various synthetic routes for the synthesis of different classes of 1,3,4-oxadiazoles and 1,3,4-thiadiazoles from diverse starting materials are illustrated in this collective chart (Figure 8) and are discussed in detail in this review. The biological activities of different 1,3,4-oxadiazole and 1,3,4-thiadiazole classes are also summarized herein.
Synthetic Approaches for the Preparation of Different Classes of 1,3,4-Oxadiazole and 1,3,4-Thiadiazole Derivatives

The synthetic approaches adopted for the preparation of 1,3,4-oxadiazole and 1,3,4-thiadiazole derivatives can be classified according to the starting material as follows:

2.1. Synthesis from Sulfonyl Acetic Acid Hydrazide

Sulphonylmethyl-1,3,4-oxadiazole derivatives are synthesized by cyclocondensation of sulfonyl acetic acid hydrazide with derivatives of carboxylic acids or sulfonyl acetic acids. The most common reagent used for cyclocondensation is POCl₃ (Figure 9). The 1,3,4-oxadiazole can then be interconverted to the 1,3,4-thiadiazole analog using thiourea and tetrahydrofuran (THF) [6].

In 2019, Sekhar et al. [6] reported the cyclocondensation of E-aroylethene (14a–c) or E-arylsulfonylethene (15a–c) sulfonylacetic acid hydrazides with substituted cinnamic acid (16a–c) using the conventional method by reflux in POCl₃ to produce the 5-styryl-1,3,4-oxadiazole derivatives (17a–c, 18a–c) in moderate yields. Interconversion of oxadiazole to thiadiazole was affected by the reaction of (17a–c, 18a–c) with thiourea in refluxing tetrahydrofuran to obtain 5-styryl-1,3,4-thiadiazole (19a–c, 20a–c) in moderate yields. However,
all synthesized compounds were obtained in higher yields and in shorter reaction times by the ultrasound irradiation method when compared to the conventional method (Scheme 1).

![Scheme 1](Image)

**Figure 9.** Proposed mechanism for the synthesis of 1,3,4-oxadiazole derivatives using sulfonyl acetic acid hydrazide based on the reported mechanism of formation of oxadiazole from hydrazide acid and carboxylic acid [49].

![Diagram](Image)

**Scheme 1.** Synthetic pathway for compounds 17a–c–20a–c.

Similarly, a new class of sulfone/sulfonamide-linked bis(1,3,4-oxadiazoles) (24a–c, 25a–c) was prepared by the cyclocondensation of arylsulfonylacetic acid hydrazide (21a–c) or arylaminosulfonylacetic acid hydrazide (22a–c) with sulfonyldiacetic acid 23 in the presence of POCl₃. Then, the bis(1,3,4-oxadiazoles) (24a–c, 25a–c) were converted to bis(1,3,4-thiadiazoles) (26a–c, 27a–c) by heating with thiourea in THF (Scheme 2) [2].

The same methodology was used by Padmaja et al., in the preparation of styrylsulfonylmethyl 1,3,4-oxadiazoles (30a–c) through cyclocondensation of the intermediate arylaminosulfonylacetic acid hydrazides (28a–c) with Z-styrylsulfonylacetic acid (29a–c). The obtained oxadiazoles were then treated with thiourea in THF to undergo interconversion to the corresponding thiadiazoles (31a–c) (Scheme 3) [49].
Similarly, a new class of sulfone/sulfonamide-linked bis(1,3,4-oxadiazoles) (32a-c) with different aryl acid hydrazides and arylsulfonylacetic acid hydrazides in the presence of POCl₃, respectively. The obtained oxadiazoles were then treated with thiourea in THF to undergo interconversion (Scheme 3).

In 2010, Padmavathi et al., synthesized a series of 2-aryliminosulfonylethyl-5-aryl-1,3,4-oxadiazoles (33a–c) and 2-aryliminosulfonylethyl-5-arylsulfonylmethyl-1,3,4-oxadiazoles (34a–c) by the cyclocondensation of arylaminosulfonylacetic acids (32a–c) with different aryl acid hydrazides and arylsulfonylacetic acid hydrazides in the presence of POCl₃, respectively. Interconversion of (33a–c, 34a–c) to thiadiazoles (35a–c, 36a–c) was affected by treatment with thiourea in THF (Scheme 4) [50].
of POCl₃, respectively. In terconversion of (33a-c, 34a-c) to thiadiazoles (35a-c, 36a-c) was affected by treatment with thiourea in THF (Scheme 4)

Scheme 4. Synthetic pathway for compounds 33a–c–36a–c.

For the thiadiazole analogs, a two-step route that involved the synthesis of the bis-hydrazide intermediate followed by cyclization was adopted. The bis-hydrazides (46a-c, 47a-c, 48a-c) were prepared by coupling the carboxylic acid derivatives (37a–c, 38a–c, 39a–c) with the acetohydrazide derivatives (40a–c, 41a–c, 42a–c) in the presence of POCl₃ (Scheme 5).

Scheme 5. Synthetic pathway for compounds 43a–c–45a–c.
of 1-[bis(dimethylamino)methylene]-1H1,2,3-triazolo [4,5-b]pyridinium-3-oxid hexafluorophosphate (HATU) and N,N-dimethylpropyl ethylamine (DIPEA) in DMF. Then, cyclo-condensation of (46a–c, 47a–c, 48a–c) with Lawesson’s reagent (LR), propylphosphonic anhydride (TPr) and TEA produced the corresponding 1,3,4-thiadiazole (49a–c, 50a–c, 51a–c) (Scheme 6) [51].

![Scheme 6](image)

Scheme 6. Synthetic pathway for compounds 49a–c–51a–c.

2.2. Synthesis from 1,2-Diacylhydrazines

1,2-Diacylhydrazines (Figure 10) can undergo dehydrocyclization reaction, forming 1,3,4-oxadiazoles using cyclodehydrating agents such as POCl₃, P₂O₅, polyphosphoric acid (PPA) [52], N,N'-dicyclohexylcarbodiimide (DCC) [53], or ethyl-3-(3-dimethylaminopropyl) carbodiimide (EDC) [54] (Figure 11). They can also be converted to the 1,3,4-thiadiazole ring by dehydrosulfurization reaction using Lawesson’s reagent (LR) [55] or P₂S₅ [56] (Figure 12).

![Figure 10](image)

Figure 10. Types of reactions of 1,2-diacylhydrazines forming the 1,3,4-oxa-/thiadiazole ring.
s-Tetrazine-3,6-dicarbohydrazides (52a–c) can be reacted at high temperature with cyclodehydrating reagents such as POCl₃ or DCC in dry toluene to obtain conjugated s-tetrazine-1,3,4-oxadiazole hybrids. This reaction was unsuccessful under conventional conditions; however, employing microwave irradiation led to the desired 1,3,4-oxadiazoles (53a–c) in good yields. s-Tetrazine-3,6-dicarbohydrazides (52a–c) were also treated with LR in dry toluene under conventional heating and microwave irradiation, with both methods resulting in the formation of s-tetrazine-1,3,4-thiadiazole derivatives (54a–c) in satisfactory yields (Scheme 7) [53].

The key intermediate N-(5-methyl-2-nitrobenzoyl)-2-substituted carbohydrazides (55a–e) were treated separately with POCl₃ and LR to obtain the desired 2-aryl-5-substituted-
1,3,4-oxadiazoles (56a–e) and thiadiazoles (57a–e), respectively. The yields of 1,3,4-thiadiazoles were low compared to those of oxadiazoles (Scheme 8) [55].

![Scheme 8. Synthetic pathway for compounds 56a–e and 57a–e.](image)

Two series of novel diosgenin derivatives bearing 1,3,4-oxadiazole (59a–e) or 1,3,4-thiadiazole (60a–e) moieties were designed and synthesized. The cyclodehydration of the N,N'-disubstituted hydrazine intermediates (58a–e) with POCl₃ furnished the corresponding 1,3,4-oxadiazoles (59a–e) in high yields. On the other hand, heating intermediates (58a–e) with LR in dry toluene under reflux conditions induced dehydrosulfurization into the 1,3,4-thiadazole counterparts (60a–e) in moderate yields (Scheme 9) [58].

![Scheme 9. Synthetic pathway for compounds 59a–e and 60a–e.](image)

Similarly, a series of 4-oxo-4'H-pyrido [1,2-a]pyrimidine derivatives containing the 1,3,4-oxadiazole (62a–c) or 1,3,4-thiadiazole (63a–c) ring were synthesized from the corresponding diacylhydrazines (61a–e) by dehydrocyclization using P₂O₅ in toluene and dehydrosulfurization by LR in THF (Scheme 10) [59].
Scheme 10. Synthetic pathway for compounds 62a–e and 63a–e.

Orthogonally protected 1,3,4-thiadiazole and 1,3,4-oxadiazole-tethered dipeptidomimetics were synthesized from a set of protected diacylhydrazines (64a–h) derived from amino acids in good to high yield with good purities and with no detectable enantiomerization. The 1,3,4-thiadiazolo-dipeptides (66a–h) were formed by refluxing (64a–h) with LR in THF. Furthermore, refluxing (64a–h) in dry CH₂Cl₂ in the presence of EDC and TEA resulted in the formation of 1,3,4-oxadiazolo-peptides (65a–h) by cyclodehydration (Scheme 11) [54].

Scheme 11. Synthetic pathway for compounds 65a–h and 66a–h.

A series of mesogenic bent-shaped 2-(4-nitrophenyl)-5-(4-n-alkoxyphenyl)-1,3,4-oxadiazoles (68a–f) was synthesized by refluxing the analogous diacylhydrazine intermediates (67a–f) with POCl₃. The 1,3,4-thiadiazole analogs (69a–f) were obtained by dehydrodossulfurization of the same intermediates through heating under reflux with P₂S₅ in dry pyridine (Scheme 12) [56].
A solvent-free solid-state method was developed to synthesize a series of substituted 1,3,4-oxadiazoles (71a–k) and 1,3,4-thiadiazoles (72a–d,f–j) derivatives in high yields by cyclodehydration of \(N,N\)-bishydrazide derivatives (70a–k). For cyclization to 1,3,4-oxadiazoles (71a–k), three kinds of cyclizing agents, \(P_2O_5\), PPA, and POCl\(_3\), were used. It was found that the three reagents worked well and provided the products with high yields. On the other hand, \(P_2S_5\) and thiourea were used as cyclizing agents to promote the cyclization to 1,3,4-thiadiazoles (72a–d,f–j). Due to the decomposition of thiourea at high temperatures, the reaction yield using thiourea was lower than that of \(P_2S_5\). Generally, \(P_2O_5\) and \(P_2S_5\) act not only as cyclizing agents but also as dehydrating agents, which further promote cyclodehydration (Scheme 13) [52].

Scheme 12. Synthetic pathway for compounds 68a–f and 69a–f.

![Scheme 12](image)

Scheme 13. Synthetic pathway for compounds 71a–k and 72a–d,f–j.

2.3. Synthesis from Acyl Semicarbazides

Aryl amino-1,3,4-oxadiazoles can be clylized from acyl semicarbazides under the effect of acid through treatment with POCl\(_3\) [60]; likewise, acyl thiosemicarbazides can be dehydrocylized to aryl amino 1,3,4-thiadiazoles under acidic conditions using conc. H\(_2\)SO\(_4\) [61], H\(_3\)PO\(_4\) [60], POCl\(_3\) [62], p-TsCl in TEA [63], or acetic anhydride [64] (Figure 13). 1,3,4-Oxadiazoles can be also synthesized from acyl thiosemicarbazides through oxidative desulfurization reaction using one of the following reagents: I\(_2\)/KI in NaOH [61], HgO in ethanol [62], Hg(OAc)\(_2\) in ethanol [66], HgCl\(_2\) in triethylamine (TEA) [67], or EDC.HCl in DMSO (Figure 14) [63].
Figure 13. Proposed mechanism for the synthesis of 2,5-disubstituted-1,3,4-oxa-/thiadiazoles using acyl semi/thiosemicarbazides based on the reported mechanism for synthesis of 2,5-disubstituted-1,3,4-thiadiazoles from acyl thiosemicarbazide [65].

Figure 14. Types of reactions of thiosemicarbazides forming 1,3,4-oxa-/thiadiazole rings.

A novel series of 5-[N-(p-fluorobenzyl)indolyl]propyl-1,3,4-thiadiazoles (75a–d) was prepared via acid-catalyzed dehydrocyclization of thiosemicarbazides (73a–d) in the presence of H₂SO₄ at 0 °C in 86–89% yields. Treatment of 4-N-phenyl thiosemicarbazide 73d with 5% (I₂/KI) in the presence of 10% NaOH produced the corresponding 2-N-phenylamine-1,3,4-oxadiazole 74d (Scheme 14) [61].

Scheme 14. Synthetic pathway for compounds 74d and 75a–d.

Similarly, cyclization of 4-substituted 1-(2-(2-chlorophenyl)-2-[4-(2,3-dichlorophenyl)piperazin-1-yl]acyl)thiosemicarbazide (76a–e) under acidic conditions yielded 1,3,4-thiadiazoles (78a–e), whereas 1,3,4-oxadiazoles (77a–e) were prepared by treating (76a–e) with iodine and KI in 4 N NaOH (Scheme 15) [68].
Scheme 15. Synthetic pathway for compounds 77a–e and 78a–e.

Oxidative cyclization of 2-isonicotinyl hydrazinecarbothioamide derivatives (79a–d) in 4 N NaOH using a mixture of I2/KI (5%) furnished the corresponding 5-(pyridin-4-yl)-N-substituted-1,3,4-oxadiazol-2-amines (80a–d) via elimination of H2S. On the other hand, 5-(pyridin-4-yl)-N-substituted-1,3,4-thiadiazol-2-amines (81a–d) were obtained by cyclization of (79a–d) upon reaction with cold conc. H2SO4 (Scheme 16) [69].

Scheme 16. Synthetic pathway for compounds 80a–d and 81a–d.

Novel coumarin derivatives possessing 2-arylamino-1,3,4-thiadiazole (84a–e) and 1,3,4-oxadiazole (83a–e) moieties were efficiently synthesized through cyclization of thiosemicarbazides (82a–e) using conventional methods. The intramolecular cyclocondensation of (83a–e) in the presence of concentrated H2SO4 formed the corresponding 1,3,4-thiadiazoles (84a–e) in good yields, whereas the cyclocondensation of (82a–e) in the presence of NaOH/KI/I2 in ethanol formed the corresponding 1,3,4-oxadiazoles (83a–e) in lower yields. Furthermore, the same compounds were also prepared by the microwave irradiation method and grinding method, with higher yields and shorter reaction times (Scheme 17) [70].

Similarly, benzimidazole derivatives featuring 1,3,4-oxadiazol-2-amine (86a–c) and 1,3,4-thiadiazol-2-amine (87a–c) heterocyclic moieties were prepared from acyl thiosemicarbazide intermediates (85a–c) via intramolecular cyclization with I2/KI/4N NaOH and conc. H2SO4, respectively (Scheme 18) [71].
Series of substituted 1,4-dihydro-4-oxoquinoline bearing 2-arylamino-1,3,4-oxadiazole (89a,b) or 1,3,4-thiadiazole (90a,b) were synthesized by El-Essawy et al. [72]. The 1,3,4-oxadiazole derivatives (89a,b) were obtained through oxidative cyclization of the acyl thiosemicarbazide intermediates (88a,b) using I$_2$/KI in ethanolic NaOH. Intramolecular dehydration of (88a,b) with cold-concentrated H$_2$SO$_4$ furnished the 1,3,4-thiadiazole counterparts (90a,b) (Scheme 19).
Scheme 19. Synthetic pathway for compounds 89a,b, and 90a,b.

In a different study, the tetrahydropyrimidinone derivative bearing a 2-amino-1,3,4-thiadiazole motif at position 5 93 was synthesized by treatment of the key intermediate tetrahydro pyrimidine-(5)-3-carbothioamide 91 with conc. H₂SO₄. Furthermore, the oxadiazole derivative 92 was prepared by heterocyclization of 91 when treated with I₂/KI in a basic medium (10% NaOH) (Scheme 20) [73].

Scheme 20. Synthetic pathway for compounds 92 and 93.

A series of N-substituted 2-amino-1,3,4-thiadiazoles (96a–c) and 1,3,4-oxadiazoles (95a–c) carrying an (R) 5-(1-(4-(5-chloro-3-fluoropyridin-2-yloxy)phenoxy)ethyl) moiety were synthesized from acyl thiosemicarbazide intermediates (94a–c) by cyclodehydration using conc. H₂SO₄ and cyclodesulfurization with alkaline iodine, respectively (Scheme 21) [74].

Scheme 21. Synthetic pathway for compounds 95a–c and 96a–c.
In a similar approach, Zoumpoulakis et al., synthesized a series of N,N-dimethylsulphonamide-based 1,3,4-thiadiazoles (99a–f) and 1,3,4-oxadiazoles (98a–f) utilizing the acylthiosemicarbazide intermediates (97a–f) through dehydrative cyclization in concentrated H₂SO₄ and desulfurative cyclization with I₂/NaOH, respectively (Scheme 22) [75].

![Scheme 22. Synthetic pathway for compounds 98a–f and 99a–f.](image)

Thermal intramolecular cyclodehydration of the bis-acyl thiosemicarbazide intermediates (100a–c) was performed by the action of H₂SO₄ at 0 °C to furnish the corresponding 2-aminoalkyl/aryl-1,3,4-thiadiazoles (102a–c). Alternatively, compounds (100a–c) were oxidatively cyclized to the corresponding 2-alkyl/arylamino-1,3,4-oxadiazoles (101a–c) in the presence of I₂/KI in refluxing (4%) NaOH (Scheme 23) [76].

![Scheme 23. Synthetic pathway for compounds 101a–c and 102a–c.](image)

A series of 2-aminotetrahydrobenzothiophene linked to arylamino 1,3,4-oxadiazole (104a–e) and 1,3,4-thiadiazole (105a–e) moieties were synthesized from the precursor acylthiosemicarbazides (103a–e) by cyclization using I₂/KI in NaOH and orthophosphoric acid, respectively (Scheme 24) [77].

![Scheme 24. Synthetic pathway for compounds 104a–e and 105a–e.](image)
In 2020, Abu-Hashem synthesized a variety of heterocyclic compounds through acyl thiosemicarbazide intermediates (106a–c). Desulfurization of (106a–c) through refluxing with HgO in ethanol afforded 1,3,4-oxidiazoles (107a–c) in moderate yields. The treatment of (106a–c) with POCl₃ led to the formation of 1,3,4-thiadiazoles (108a–c) in higher yields than those of oxadiazoles (Scheme 25) [62].

![Scheme 25. Synthetic pathway for compounds 107a–c and 108a–c.](image)

The synthesis of some 1,3,4-thia/oxadiazole derivatives containing arylsulphonylphenyl and 4-trifluoromethylphenyl or 2,4-difluorophenyl moieties was achieved through cyclization of hydrazinecarbothioamides (109a–f). Treatment of (109a–f) with POCl₃ resulted in dehydrative cyclization obtaining the 2-arylamino-1,3,4-thiadiazoles (111a–f) in moderate yields, while on refluxing of the same intermediates with HgO, desulfurative cyclization took place, resulting in the corresponding 2-arylamino-1,3,4-oxadiazoles (110a–f) being obtained in fair yields (Scheme 26) [78,79].

![Scheme 26. Synthetic pathway for compounds 110a–f and 111a–f.](image)

Treatment of N-(furan-2-ylmethylidene)-4,6-dimethyl-1-H-prazolo [3,4,b]pyridine-1-yl acetyl thiosemicarbazide 112 with ice-cooled concentrated H₂SO₄ resulted in the formation of the respective 1,3,4-thiadiazole derivative 114. On the other hand, heating intermediate 112 with HgO in methanol resulted in desulfurative cyclization, which afforded the 1,3,4-oxadiazole derivative 113 (Scheme 27) [80].
Burbuliene et al., developed concise and efficient procedures for the synthesis of 1,3,4-oxadiazoles and 1,3,4-thiadiazoles bearing 4-pyrimidinylthio- and 2-pyrimidinylthiоieties by intramolecular cyclization of substituted acyl thiosemicarbazide precursors (115a–d). The cyclodesulfurization to form 1,3,4-oxadiazoles (116a–d) was conducted by treating (115a–d) with Hg(OAc)_2 in ethanol. On the other hand, cyclodehydration of (115a–d) in acidic media using concentrated H_2SO_4 at ~0 °C followed by neutralization with NH_4OH provided the thiadiazoles (117a–d) in lower yields than those of the oxadiazole analogs (Scheme 28) [66].

Series of 1,3,4-thiadiazoles (120a–c) and 1,3,4-oxadiazoles (119a–c) containing isomeric pyridyl and cyclohexyl moieties were synthesized by intramolecular cyclization of 1,4-disubstituted acyl thiosemicarbazides (118a–c). The latter underwent cyclodehydration in the acid medium of conc. H_2SO_4 to form 1,3,4-thiadiazoles (120a–c). The 1,3,4-oxadiazoles (119a–c) were obtained from the corresponding thiosemicarbazides in the presence of Hg(OAc)_2 as an oxidizing agent (Scheme 29) [81].
Novel heterocyclic derivatives of 2-arylamino-5-(pyridin-2-ylmethyl)-1,3,4-oxadiazoles (122a–g) and thiadiazoles (123a–g) were efficiently synthesized through (pyridin-2-ylacetyl)hydrazinecarbothioamide derivatives (121a–g). The method for obtaining 1,3,4-thiadiazole derivatives (123a–g) required cyclization under acidic conditions (conc. H₂SO₄), while oxadiazoles (122a–g) were prepared by a desulfurization/cyclization reaction of (121a–g) using HgCl₂ with optional catalytic amounts of TEA (Scheme 30) [67].

Scheme 29. Synthetic pathway for compounds 119a–c and 120a–c.

Polymer-bound 2-amino-1,3,4-oxadiazoles (125a–d) and 1,3,4-thiadiazole 126 core skeleton resin were synthesized by solid-phase cyclization of acyl thiosemicarbazides (124a–e) with EDC·HCl and p-TsCl, respectively, followed by a functionalization reaction of the resulting core skeleton with various electrophiles such as alkyl halides and acid chlorides to generate N-alkylamino and N-acylamino-1,3,4-oxadiazole (127a–x and 128a–x) and 1,3,4-thiadiazole (129a–j and 130a–j) resin, respectively. Finally, the desired 2-amino and 2-amido-1,3,4-oxadiazole (131a–x and 132a–x) and 1,3,4-thiadiazole (133a–j and 134a–j) were then generated in high purities by cleavage of the resin under trifluoroacetic acid (TFA) in dichloromethane (DCM) (Scheme 31) [63].

In 2013, a regioselective, reagent-based method for the cyclization reaction of acyl thiosemicarbazide (135a–m) into 2-amino-1,3,4-oxadiazole (136a–m) and 2-amino-1,3,4-thiadiazole (137a–m) core skeletons was described. The study showed that the use of EDC·HCl in DMSO as a desulfurizing agent led to 2-amino-1,3,4-oxadiazoles (136a–m) in quantitative yields (regioselectivity ratio = 100:0). However, the use of p-TsCl as a dehydrating agent and triethylamine in N-methyl-2-pyrrolidone as a polar solvent gave 2-amino-1,3,4-thiadiazoles (137a–m) as a major product with regioselectivity of 96:4 and a high yield (Scheme 32) [82].
In 2013, a regioselective, reagent-based method for the cyclization reaction of acyl thiosemicarbazide (135a-m) into 2-amino-1,3,4-oxadiazole (136a-m) and 2-amino-1,3,4-thiadiazole (137a-m) core skeletons was described. The study showed that the use of EDCl·HCl in DMSO as a desulfurizing agent led to 2-amino-1,3,4-oxadiazoles (136a-m) in quantitative yield (regioselectivity ratio = 100:0). However, the use of p-TsCl as a dehydrating agent and triethylamine in N-methyl-2-pyrrolidone as a polar solvent gave 2-amino-1,3,4-thiadiazoles (137a-m) as a major product with regioselectivity of 96:4 and a high yield (Scheme 32) [82].

Scheme 31. Synthetic pathway for compounds 125a–d–134a–j.

= Merrifield resin

R² = ethenyl, isobutyl, cyclohexyl, phenyl, 4-methylphenyl, 4-fluorophenyl, 4-methoxyphenyl, 4-nitrophenyl

R³ = methyl, tertbutyl, penty1, neopentyl, phenyl, 4-fluorophenyl, 4-chlorophenyl, 4-trifluoromethylphenyl, 4-methoxyphenyl, 2-furyl, 3-quino1inyl.

Scheme 31. Synthetic pathway for compounds 125a–d–134a–j.
Yavus et al. synthesized some endoethenotetrahydrothebaine derivatives incorporating 1,3,4-oxadiazole (140a,b) or 1,3,4-thiadiazole moiety (141a,b) by cyclodehydration of the intermediate acylsemicarbazides (138a,b) and acylthiosemicarbazides (139a,b) using the cyclizing agents POCl$_3$ and H$_3$PO$_4$, respectively (Scheme 33) [60].

Long-chain 5-alkenyl/hydroxyalkenyl-2-phenylamine-1,3,4-oxadiazoles and 1,3,4-thiadiazoles were synthesized from acyl semicarbazides and thiosemicarbazides, respectively. Refluxing semicarbazide (142a–d) with POCl$_3$ yielded 2,5-disubstituted 1,3,4-oxadiazoles (143a–d) in good yields. The dehydrative cyclization of thiosemicarbazides (144a–d) by Ac$_2$O produced 2,5-disubstituted 1,3,4-thiadiazoles (145a–d) in excellent yields (Scheme 34) [64].
Scheme 34. Synthetic pathway for compounds 143a–d, 145a–d.

2.4. Synthesis from Semi-/Thiosemicarbazides

Semi-/thiosemicarbazides are reacted with aldehydes in the presence of Na acetate in ethanol, forming the corresponding semi-/thiosemicarbazones, which can be used for the synthesis of 1,3,4-oxa-/thiadiazoles by oxidative cyclization using I$_2$ /K$_2$CO$_3$ [83], Br$_2$ /CH$_3$COOH [84], or nitroalkanes in PPA [85] (Figure 15).

![Proposed mechanism of electroxidative cyclization of semi-/thiosemicarbazones into 1,3,4-oxa-/thiadiazoles based on the reported mechanism in literature [86].](image)

The reaction of trifluoromethyl benzaldehyde 146 with semi- or thiosemicarbazide hydrochloride 147 and 148 formed the corresponding semi- and thiosemicarbazone 149 and 150. Cyclization using I$_2$ /dioxane produced the 1,3,4-oxadiazol-2-amine 151 and 1,3,4-thiadiazol-2-amino 152, respectively, which were further reacted with aroyl chlorides to give the desired 1,3,4-oxadiazol-2-yl/1,3,4-thiadiazol-2-yl benzamides (153a–c) and 154b. Oxadiazoles were produced in higher yields than thiadiazoles (Scheme 35) [87].

![Scheme 35. Synthetic pathway for compounds 153a–c and 154b.](image)
In 2015, 2-amino-substituted 1,3,4-oxadiazoles (159a–c) and 1,3,4-thiadiazoles (160a–c) were smoothly produced via condensation of semicarbazide hydrochloride/thiosemicarbazide 147 and 156 and the corresponding aldehydes (155a–c), followed by I$_2$-mediated oxidative C–O/C–S bond formation using a mixture of I$_2$ and K$_2$CO$_3$ in 1,4-dioxane. The 1,3,4-oxadiazoles were produced in higher yields than the 1,3,4-thiadiazoles (Scheme 36) [83].

$$\text{R}_1\text{CHO} + 147 \rightarrow 157a-c, X=O \quad 158a-c, X=S$$

R$_1$= a) alkyl, b) alkenyl, c) aryl

Scheme 36. Synthetic pathway for compounds 159a–c and 160a–c.

In 2017, Arafa and Abdelmagied reported that a rapid ultrasound accelerated the conversion of bis-semicarbazone derivatives (162a–f) into diverse novel substituted bis-2-amino-1,3,4-oxadiazoles (164a–f) in quantitative yields using molecular iodine and potassium carbonate in dioxane as a solvent. Products (164a–f) could also be obtained in one-pot synthesis without the isolation of intermediates (162a–f). Replacing semicarbazide hydrochloride 147 with thiosemicarbazide 156 produced the desired bis-2-amino-1,3,4-thiadiazoles (165a–f) in excellent yields (Scheme 37) [88].

$$\text{OHC}_2\text{CHO} + 156 \rightarrow 162a-f, X=O \quad 163a-f, X=S$$

Scheme 37. Synthetic pathway for compounds 164a–f and 165a–f.

The reaction of pyrazole-4-carboxaldehyde derivatives bearing 2-hydroxyphenyl or 7-hydroxycoumarin-8-yl moiety (166a–k) with (semi)thiosemicarbazide hydrochloride 147 or 148 in the presence of sodium acetate in ethyl alcohol under reflux conditions afforded the corresponding semicarbazones (167a–k) and thiosemicarbazones (168a–k). Their oxidative cyclization using bromine as an oxidant in acetic acid at room temperature yielded the corresponding 1,3,4-oxadiazoles (169a–k) and 1,3,4-thiadiazoles (170a–k) (Scheme 38) [84,89].
Scheme 38. Synthetic pathway for compounds 169a–k and 170a–k.

In 2020, Askenov et al., discovered an unusual reaction (Figure 16) for the synthesis of 2-amino-1,3,4-oxa-/thiadiazoles via the electrophilic activation of nitroalkanes in the presence of polyphosphoric acid followed by a subsequent nucleophilic attack with semi/thiosemicarbazides. It was found that nitroalkanes 172 in PPA converted into phosphorylated nitronates showing strong electrophilic properties. Semicarbazides 171 and thiosemicarbazides 174 were employed as nucleophilic components in this cyclization reaction to access 2-amino-1,3,4-oxadiazoles (173a–d) and 2-amino-1,3,4-thiadiazoles (175a–g), respectively (Scheme 39) [85].

Figure 16. Mechanistic rationale for the formation of compounds 173a–d and 175a–g [85].

The 2-(β-D-glucopyranosyl)-5-(substituted-amino)-1,3,4-oxa-/thiadiazoles were formed by the oxidative ring closure of (semithio)semicarbazones. Semicarbazone 176 was reacted with Pb(OAc)₄ in glacial AcOH to furnish 1,3,4-oxadiazole 177. On the other hand, acylation of thiosemicarbazone 178 in dry pyridine resulted in the formation of 1,3,4-thiadiazolines (179a–d) as inseparable mixtures of two diastereoisomers in moderate yields. Furthermore, semicarbazone 180 was treated by acid chlorides in CH₂Cl₂ in the presence of sodium acetate in ethyl alcohol under reflux conditions to give intermediates (181a–d), which were then reacted with phenyliodine (III) diacetate (PIDA) in CH₂Cl₂ to obtain 1,3,4-oxadiazoles (182a–d) in good yields. Under these
conditions, no reaction took place with thiosemicarbazone 178. However, 1,3,4-thiadiazoles (183a–d) were directly derived in moderate yields from thiadiazolines (179a–d) by oxidation (Scheme 40) [90].

\[
\begin{align*}
R & = \text{phenyl, 2-pyrazolyl} \\
R_1 & = \text{H, methyl, ethyl} \\
\end{align*}
\]

\[
171 + 172 \xrightarrow{86\% \text{PPA, 120°C, 2.5h}} 173a-d \text{ (yield=62-75%)}
\]

- a) R=phenyl, R_1=ethyl 
- b) R=2-pyrazolyl, R_1=H 
- c) R=2-pyrazolyl, R_1=methyl 
- d) R=2-pyrazolyl, R_1=ethyl

\[
\begin{align*}
R_1 & = \text{H, methyl, ethyl, ethylcarboxylate} \\
R_2 & = \text{H, methyl, phenyl} \\
R_3 & = \text{H, benzyl} \\
\end{align*}
\]

\[
174 + 172 \xrightarrow{86\% \text{PPA, 120°C, 2.5h}} 175a-g \text{ (yield=38-88%)}
\]

- a) R_1=R_2=R_3=H 
- b) R_1=methyl, R_2=R_3=H 
- c) R_1=ethyl, R_2=R_3=H 
- d) R_1=ethylcarboxylate, R_2=R_3=H 
- e) R_1=H, R_2=phenyl, R_3=H 
- f) R_1=ethyl, R_2=phenyl, R_3=H 
- g) R_1=ethyl, R_2=methyl, R_3=benzyl

Scheme 39. Synthetic pathway for compounds 173a–d and 175a–g.

\[
\begin{align*}
R_1 & = \text{H, methyl, ethyl, ethylcarboxylate} \\
R_2 & = \text{H, methyl, phenyl} \\
R_3 & = \text{H, benzyl} \\
\end{align*}
\]

\[
\begin{align*}
\text{Scheme 40. Synthetic pathway for compounds 177, 182a–d, and 183a–d.}
\end{align*}
\]

\[
\begin{align*}
\text{Ar= a) methyl, b) phenyl, c) 4-tertbutylphenyl, d) 2-naphthyl}
\end{align*}
\]
2.5. Synthesis from Acid/Thionyl Hydrazide

To date, various methods have been reported in the literature for generating 1,3,4-oxa-/thiadiazoles from acid hydrazides (Figure 17). 1,3,4-Oxadiazoles can be synthesized from acid hydrazides through treatment with tetramethylthiuram disulfide (TMTD) in DMF [91], cyanogen bromide (CNBr) in aq. Dioxane [91], coupling agent (CDI)/THF/POCl₃ [92], CS₂/KOH in ethanol [93], or by reaction with triethyl orthoesters in glacial acetic acid [53]. On the other hand, 1,3,4-thiadiazoles can be synthesized through treatment of acid hydrazides with TMTD/DMF [91], isothiocyanates/TEA [91], CDI/DMF/P₂S₅ [92], by treatment with conc. H₂SO₄ through the formation of acyl thiosemicarbazide intermediate [93], or by reaction with triethyl orthoesters/LR in glacial acetic acid [53]. Furthermore, TiCl₄-mediated facile synthesis [94] and the palladium-catalyzed one-pot method [95] are used for the synthesis of 1,3,4-oxa-/thiadiazoles from acid/thionyl hydrazide.

![Figure 17. Different methods used for the synthesis of 1,3,4-oxa-/thiadiazoles from acid hydrazides.](image_url)

A series of 1,3,4-oxadiazole and 1,3,4-thiadiazole bearing an (R) 5-(1-(4-(5-chloro-3-fluoropyridin-2-yl oxy)phenoxy)ethyl) unit were synthesized from the corresponding acid hydrazide through the formation of the intermediate formylhydrazide. Prolonged reflux of hydrazide 184 in ethyl orthoformate generated formylhydrazide 185 with subsequent cyclization to oxadiazole 186. Alternatively, sulfurative cyclization of formylhydrazide 185 to thiadiazole 187 was affected by heating with P₂S₅ in xylene under reflux conditions. Oxadiazole was formed in a higher yield than thiadiazole (Scheme 41) [74].

A series of 4,5-dihydro-1,5-diaryl-1H-pyrazole substituted at position 3 with functionalized 1,3,4-oxadiazole or 1,3,4-thiadiazole ring was synthesized from carboxyhydrazide precursors (188a,b) using variable reagents and conditions. The 2-hydroxy-1,3,4-oxadiazoles (189a,b) were prepared by reaction of carboxyhydrazide with (188a,b) in tetrahydrofuran using triethylamine as a base. The 1,3,4-oxadiazole-2-thiols (190a,b) were obtained by reaction of (188a,b) with carbon disulfide employing KOH as a base catalyst. The 1,3,4-oxadiazole-2-azines (191a,b) resulted from the reaction of cyanogen bromide and NaHCO₃ with precursors (188a,b). The 5-methylthio-1,3,4-thiadiazoles (192a,b) were synthesized by reaction of hydrazide (188a,b) with carbon disulfide and KOH followed by methylation using methyl iodide. These compounds were further cyclized in refluxing toluene using p-toluene sulfonic acid (Scheme 42) [96].
Aggarwal et al., synthesized nalidixic acid-based 1,3,4-(oxa)thiadiazoles through the reaction of nalidixic acid hydrazide 193 with ice-cooled carbon disulfide in the presence of potassium hydroxide to produce potassium dithiocarbazinate intermediate 194. Dithiocarbazinate intermediate 194 was then cyclized in acidic or in basic medium forming 1,3,4-thiadazole 196 or 1,3,4-oxadiazole 195, respectively. Thiadiazole was formed in a higher yield than its oxadiazole analog (Scheme 43) [93].
Scheme 43. Synthetic pathway for compounds 195 and 196.

Similarly, Knyazyan et al., synthesized 3H[1,3,4] (oxa)thiadiazole-2-thiones (199a,b and 200a,b) under base- and acid-catalyzed heterocyclization of 2-(3-alkyl-4-methyl-2-thioxo-2,3-dihydro-thiazole-5-carbonyl)-hydrazinecarbodithioic acid potassium salts (198a,b) (Scheme 44) [97].

Scheme 44. Synthetic pathway for compounds 199a,b and 200a,b.

A series of benzosuberone analogs embedded with 1,3,4-oxadiazole or 1,3,4-thiadiazole moieties were synthesized in high yields. The reaction of the hydrazides (201a,b) with phenyl isothiocyanate in ethanol gave the corresponding acylthiosemicarbazide (203a,b), which on treatment with conc. sulphuric acid yielded the required 1,3,4-thiadiazoles (204a,b), while oxadiazole analogs (202a,b) were obtained from (201a,b) directly by refluxing with carbon disulfide in the presence of alcoholic potassium hydroxide (Scheme 45) [98].
In 2020, (+)-sclareolide-based homodrimane sesquiterpenoids bearing 1,3,4-oxa-/thiadiazole-2-thiols (206,207) were synthesized from acetohydrazide 205 through treatment with TMTD in DMF. Products were obtained selectively or in a mixture where the yield ratio between oxadiazole 206 and thiadiazole 207 was dependent on the amount of TMTD used in the reaction (Scheme 46) [91].

To obtain 2-aminosubstituted 1,3,4-thiadiazole and 1,3,4-oxadiazole units, homodrimane hydrazide 205 was reacted with isothiocyanate derivatives, without any isolations of the intermediate compounds, in the presence of Et$_3$N in H$_2$O, giving substituted 2-amino-1,3,4-thiadiazoles (209a–c) in moderate yields. Additionally, acetohydrazide 205 was treated with CNBr in aqueous dioxane to form unsubstituted 1,3,4-oxadiazol-2-amine 208 in an 80% yield (Scheme 47) [91].
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Scheme 47. Synthetic pathway for compounds 208 and 209a-c.

In 2019, Zhang et al. developed an efficient synthesis of 2,5-disubstituted 1,3,4-oxadiazoles (213a–o) and 1,3,4-thiadiazoles (214a–d) from the reaction of acid hydrazides (210a–i) or thionyl hydrazide 211 with N,N-dimethyl amide derivatives (212a–e) utilizing TiCl4 as a single dehydration reagent for both coupling and cyclodehydration events. This protocol is a facile and economical method that provides the target oxadiazoles/thiadiazoles in good yields and with a wide substrate scope (Scheme 48) [94].

Scheme 48. Synthetic pathway for compounds 213a–o and 214a–d.

In 2011, Padmavathi et al., prepared 5-aryl-2-(chloromethyl)-1,3,4-oxadiazoles (216a–c) by the reaction of chloroacetic acid 215 with benzoic hydrazides (210a–c) in the presence of POCl3. However, they obtained the 5-aryl-2-(chloromethyl)-1,3,4-thiadiazoles (217a–c) by the interconversion of compounds (216a–c) with thiourea in THF (Scheme 49) [99].

Scheme 49. Synthetic pathway for compounds 216a–c and 217a–c.
Xiang et al., reported an efficient one-pot Pd-catalyzed carbonylative coupling reaction for the synthesis of 1,3,4-oxadiazoles (221a–o, 222a–j) from the reaction of benzohydrazides (219a–p) and iodobenzene (218a–j) with tert-butyl isocyanide 220 via isocyanide insertion/cyclization sequential reaction. The optimal reaction conditions were PdCl₂ and DPPP as the catalyst systems, with NaOAc as the base and DMF as the solvent, which furnished the products in moderate yields. Obtaining 1,3,4-thiadiazoles (224a–e) required replacement of benzohydrazides (219a–o) with benzothiohydrazides (223a–e), as well as changing the base from NaOAc to Na₂CO₃ (Scheme 50) [95].

\[
\begin{align*}
218a & \quad + \quad 219a-o & \quad + \quad t-Bu-N=NC \quad \overset{\text{PdCl}_2, \text{DPPP, NaOAc}}{\longrightarrow} & \quad 220 \\
221a-o & \quad (\text{Ar= a) methyl, b) cyclohexyl, c) 2-methylphenyl, d) 4-methylphenyl, e) 4-dimethylaminophenyl, f) 4-chlorophenyl,} \\
& \quad g) 4-fluorooctacetylophenyl, h) 4-methoxyphenyl, i) 3,4,5-trimethoxyphenyl, j) 1,1'-biphenyl, k) naphthyl,} \\
& \quad l) 3-furyl, m) 3-thienyl, n) benzo[d][1,3]-dioxol-5-yl, o) N-methylindol-3-yl)
\end{align*}
\]

\[
\begin{align*}
218a-j & \quad + \quad 219p & \quad + \quad t-Bu-N=NC \quad \overset{\text{PdCl}_2, \text{DPPP, NaOAc}}{\longrightarrow} & \quad 222a-j \\
222a-j & \quad (\text{R= a) H, b) 3-methyl, c) 4-methyl, d) 3,5-dimethyl, e) tertbutyl, f) 4-chloro, g) 4-fluoro, h) 3,5-difluoro,} \\
& \quad i) trifluoromethyl, j) 3-methoxy)
\end{align*}
\]

\[
\begin{align*}
218a & \quad + \quad 223a-e & \quad + \quad t-Bu-N=NC \quad \overset{\text{PdCl}_2, \text{DPPP, Na}_2\text{CO}_3}{\longrightarrow} & \quad 224a-e \\
224a-e & \quad (\text{Ar= a) phenyl, b) 4-methylphenyl, c) 3,5-dimethylphenyl, d) 3-methoxyphenyl, e) naphthyl}
\end{align*}
\]

Scheme 50. Synthetic pathway for compounds 221a–o, 222a–j and 224a–e.

A series of benzimidazole derivatives bearing a 1,3,4-oxadiazole/thiadiazole core were synthesized by the precursor carboxyhydrazone 225 with substituted benzoic acid using the coupling agent CDI and POCl₃ and P₂S₅ as cyclizing agents to yield the oxadiazole (226a–j) and thiadiazole (227a,c,f,i) cores, respectively (Scheme 51) [92].

\[
\begin{align*}
\text{CDI, THF, POCl}_3, \quad \text{reflux 5h} & \quad \rightarrow \quad 225 \\
\text{CDI, DMF, P}_2\text{S}_5, \quad \text{reflux 5h} & \quad \rightarrow \quad 227a,c,f,i \\
\end{align*}
\]

\[
\begin{align*}
226a-j & \quad (\text{R= a) H, b) 4-methyl, c) 4-butyl, d) 4-cyano, e) 3-fluoro,} \\
& \quad f) 4-fluoro, g) 4-bromo, h) 3-methoxy, i) 4-nitro, j) 4-phenyl}
\end{align*}
\]

Scheme 51. Synthetic pathway for compounds 226a–j and 227a,c,f,i.
In 2020, s-tetrazine-1,3,4-oxadiazole hybrids (229a–d) were obtained by reaction of s-tetrazine-3,6-dicarbohydrazide 228 and triethylorthoesters in glacial acetic acid via conventional heating. S-tetrazine-1,3,4-thiadiazoles (230a–d) were synthesized by the same method with the addition of LR. The products were also obtained by microwave irradiation in similar yields, but in a very short reaction time (Scheme 52) [53].

![Scheme 52. Synthetic pathway for compounds 229a–d and 230a–d.](image)

### 3. Biological Activities

Over the past few decades, many 1,3,4-oxadiazoles and 1,3,4-thiadiazoles have been intensively studied for their broad biomedical applications, such as antimicrobial, antiviral, anticancer, antidiabetic, antioxidant, and analgesic activities (Figure 18).

![Figure 18. Some biological activities for 1,3,4-oxa-/thiadiazoles.](image)

#### 3.1. Antimicrobial Activity

In 2020, thirteen novel sclareolide-based homodrimane sesquiterpenoids bearing 1,3,4-oxadiazole or 1,3,4-thiadiazole moieties were synthesized and demonstrated promising antifungal and antibacterial activities toward fungal species Aspergillus niger, Fusarium solani, Penicillium chrysogenum, Penicillium frequentans, and Alternaria alternata and Gram-positive Bacillus species and Gram-negative Pseudomonas aeruginosa bacteria strains. On the basis of the SAR correlations, it was noticed that compounds having a 1,3,4-thiadiazole moiety were more active compared with those having a 1,3,4-oxadiazole moiety.
Among the thiadiazole derivatives, compound 231 (Figure 19), featuring the 2-mercapto group, was the most potent, with promising broad-spectrum antifungal (MIC 0.032 μg/mL) and antibacterial (MIC 0.094 μg/mL) activities [91].

![Structure of compound 231.](image)

In 2012, nalidixic acid-based 1,3,4-(oxa)thiadiazol-2-thione, their Mannich bases, S-alkylated/arylated sulfides, and sulfones were synthesized and screened against Gram-positive bacteria (S. aureus and B. subtilis) and Gram-negative bacteria (E. coli, P. aeruginosa, and K. pneumonia) using Streptomycin as the reference standard drug. According to evaluation results, the Mannich base 232 (Figure 20) emerged as a potent antibacterial agent against all test microorganisms except E. coli, with an MIC range of 6.25–12.5 μg/mL [93].

![Structure of compound 232.](image)

A series of 1,3,4-oxadiazoles/thiadiazoles bearing pyrazole scaffolds were screened for their in vitro antimicrobial activity against S. aureus, E. coli, P. aeruginosa, A. niger, and A. flavus bacteria, and C. albicans fungal species using Ciprofloxacin and Fluconazole as reference drugs. The tested compounds showed moderate to excellent antibacterial and antifungal activities. The chloro-substituted oxadiazole 169d was equipotent to reference drugs Ciprofloxacin and Fluconazole MIC 12.5–25 μg/mL, while the chloro-substituted 1,3,4-oxadiazole 169j showed greater activity against Gram-negative E. coli and C. albicans fungi (MIC 12.5 μg/mL) than Ciprofloxacin and Fluconazole (MIC 25 and 50 μg/mL, respectively). On the other hand, the chloro-substituted thiadiazoles (170d,j) showed a higher activity against Gram-positive S. aureus (MIC 12.5 μg/mL) compared with standard Ciprofloxacin (MIC 25 μg/mL). Moreover, 170j was more potent against Gram-negative E. coli (MIC 12.5 μg/mL) than Ciprofloxacin (MIC 25 μg/mL).

The comparative analysis illustrated that, among oxadiazoles and thiadiazoles, the sulfur moiety in 1, 3, 4- thiadiazoles acts as a potent inhibitor, while compounds containing moderately electronegative elements such as sulfur and chlorine (170d,j) showed better in vitro activity than compounds with more electronegative elements, such as oxygen (169d,j) (Figure 21) [84,89].
Long-chain 5-alkenyl/hydroxyalkenyl-2-phenylamine-1,3,4-oxadiazoles and thiadiazoles were synthesized and screened as antimicrobial agents against *S. pyogenes*, *S. aureus*, *P. aeruginosa*, *K. pneumoniae*, and *E. coli* bacterial strains and *C. albicans*, *A. fumigatus*, *T. mentagrophytes*, and *P. marneffei* fungal strains. The study results showed no difference in activity between the bioisosteres of oxadiazoles and thiadiazoles. Examination of SAR indicated that the presence of a hydroxyalkyl chain at the fifth position of the oxadiazole ring increased the antibacterial activity in compounds 143c, 143d, 145c, and 145d, as they were equally as potent as the reference drug chloramphenicol (MIC = 6.25 μg/mL). Moreover, the compounds with an internal double bond in the long alkenyl substituent of synthesized oxadiazoles 143b and 145b were found to be potent antifungal agents (Figure 22) [64].

A series of 5-[N-(p-flurobenzyl)]indolyl]propyl-1,3,4-(oxa)thiadiazoles were tested in vitro for their antibacterial and antifungal activities. Only the thiadiazole derivatives showed promising results. The 2-phenylamino derivative 75d was more potent than the reference drug Gentamicin against Gram-negative *E. coli*, with MIC = 0.24 and 3.9 μg/mL, respectively. The 2-ethylamino analog 75b was equipotent to Gentamicin against *E. coli* (MIC = 3.9 μg/mL), and was four times more potent than Gentamicin against *P. aeruginosa* (MIC 3.9 μg/mL and 15.6 μg/mL, respectively). The 2-cyclohexyl amino counterpart 75c showed promising results against *E. coli* (MIC 0.49 μg/mL) compared to Gentamicin and displayed equivalent potency to Amphotericin B against the *S. racemosum* fungus (MIC = 3.9 μg/mL) (Figure 23) [61].

The potential antimicrobial effect of 5-(benzimidazol-1-yl methyl)-2-arylamino-1,3,4-(oxa)thiadiazoles was investigated against microorganisms representing Gram-positive bacteria (*S. aureus* and *S. epidermidis*), Gram-negative bacteria (*E. coli* and *P. aeruginosa*), and fungi (*C. albicans*). The oxadiazole derivative 86b (inhibition zone diameter (IZD) = 26–28 mm), as well as the thiadiazole derivative 87b (IZD = 25–27 mm), carrying bromophenyl amino rest, and the methoxyphenyl amino thiadiazole 87c (IZD = 26–28 mm) showed comparable antibacterial activity to the reference drug Cefotaxime (IZD = 28–35 mm) against the tested strains. Additionally, they elicited superior antifungal activity (IZD = 28–30 mm) compared to Nystatin (IZD = 24 mm) (Figure 24) [71].

**Figure 21.** Structures of compounds 169d,j and 170d,j.

**Figure 22.** Structures of compounds 143a–d and 145a–d.
Screening the antibacterial activity of 5-(arylamino)-1,3,4-(oxa)thiadiazol-2-yl)quinolin-4-one derivatives against three different bacterial species, namely, Gram-negative *P. aeruginosa*, Gram-positive *B. subtilis*, and actinomycete *Streptomyces* sp., revealed variable degrees of inhibitory actions. The highest degrees of inhibition were recorded for the oxadiazole compound **89b** and the thiazole compound **90a**, and they were equipotent. The most susceptible organism was *B. subtilis* (IZD = 20–22 mm), followed by *Streptomyces* sp. (IZD = 12–18 mm), whereas the lowest inhibitory effect was encountered in the case of *Pseudomonas* sp. (IZD = 4 mm) (Figure 25) [72].

Several 1,3,4-oxa-/thiadiazole derivatives were designed and synthesized as analogs for 4-chloro-N-(5-(4-fluorophenyl)-1,3,4-oxadiazol-2-yl)benzamide **KKL-35**, which is one of the most active oxadiazole compounds, showing potent bactericidal activity against various non-related bacteria [100]. The designed compounds were tested for antibacterial activity against a selection of 24 bacterial species, using **KKL-35** as a reference. 6-Chloro-N-(5-(4-(trifluoromethyl)phenyl)-1,3,4-oxadiazol-2-yl)nicotinamide **153c**, and its 3-chlorobenzamide analog **153a** exhibited a wide spectrum of action and strong antimicrobial activity. They were active against almost all of the tested Gram-positive strains except *C. perfringens* and *S. pyogenes*. However, the only Gram-negative strain affected was *B. fragilis* (by **153c**). They also affected the growth of the mycobacterium, *M. fortuitum*, and **153c** had an effect against *M. abscessus*. Compared to **KKL-35**, **153c** displayed lower
MICs for *E. faecalis* (MIC 2–4 mg/L), *E. faecium* (MIC 2 mg/L), and *M. fortuitum* (MIC 8 mg/L). In addition, 153a showed a superior effect against *E. faecalis*, *E. faecium*, and *S. epidermidis* (MICs = 0.0625–1 mg/L). Furthermore, these compounds show synergistic activity with conventional antibiotics and have low toxicities (Figure 26) [87].

![Figure 26. Structures of compounds KKL-35 and 153a,c.](image)

Karabanovich et al., evaluated the activity of a series of substituted benzylsulfanyl-1,3,4-oxadiazoles and 1,3,4-thiadiazoles as antitubercular agents. The majority of these compounds exhibited outstanding in vitro activity against *Mycobacterium tuberculosis*. The activity results revealed that there was no difference in activity between oxadiazoles and thiadiazole analogs. Most of the compounds bearing the 3,5-dinitrobenzylsulfanyl moiety (233 and 234 series) exhibited excellent activity against drug-susceptible and multidrug-resistant M.tb. strains, with no cross-resistance with first- and second-line anti-TB drugs. An SAR study indicated that removal or substitution of one nitro group or changing the positions of both nitro groups resulted in a decrease or loss of activity (Figure 27). The anti-TB effect of the compounds was selectivity with low toxicity, genotoxicity, and mutagenicity [101].

![Figure 27. SAR for compounds 233 and 234.](image)

According to the theory that the incorporation of different pharmacophores into a single molecule can change the activity of the newly obtained compound, a series of cyclic imides containing both 1,3,4-oxadiazole and 1,3,4-thiadiazole heterocycles was prepared and tested for their antimicrobial activity against four strains of bacteria (*S. aureus*,...
S. pyogenes, E. coli, and P. aeruginosa) and C. albicans fungi. The results indicated that compounds 235 (IZD = 14–20.5 mm), 236 (IZD = 12.6–21 mm), and 237 (IZD = 12.9–20.1 mm) were highly active against all types of tested bacteria compared to Ampicillin (IZD = 12–17 mm). Only compounds 235 (IZD = 18.6 mm) and 237 (IZD = 22.2 mm) were also highly active against C. albicans fungi (Fluconazole IZD = 18 mm) (Figure 28) [102].

![Figure 28. Structures of compounds 235–237.](image)

This study supports the same theory, whereby the potential antibacterial activity of a series of 1,3,4-thiadiazole derivatives tagged to oxadiazole and thiadiazole units were evaluated against E. coli, S. typhimurium, L. monocytogenes, K. pneumonia, S. typhi, S. aureus, and B. subtilis species. The assay results revealed that compounds 238 and 239, with a strong electron-withdrawing fluorine atom at the para position, were more potent than Ampicillin on all tested strains (MIC = 62.5–100 μg/mL and 100–250 μg/mL, respectively). It was also noticed that the 1,3,4-thiadiazoles linked with 1,3,4-thiadiazole units 239 were more active than 1,3,4-thiadiazoles linked with 1,3,4-oxadiazole 238 (Figure 29) [103].

![Figure 29. Structures of compounds 238 and 239.](image)

A series of 2,5-disubstituted-1,3,4-thiadiazole tethered 1,3,4-(oxa)thiadiazole derivatives were synthesized as antimicrobial agents. Their activities were evaluated against a panel of standard strains of pathogenic microorganisms, including Gram-positive bacteria (S. pneumonia, B. subtilis, and S. aureus), Gram-negative bacteria (P. aeruginosa, E. coli, and K. pneumonia), and fungi (A. fumigatus, C. albicans, and G. candidum), and against reference drugs Ciprofloxacin and Fluconazole. The antimicrobial activity of the thiadiazoles (102a–c) revealed that the tested compounds showed promising activity against all bacterial and fungal strains at 8–31.25 μg/mL. However, the oxadiazole isosteres (101a–c) showed greater antibacterial activity against Gram-positive bacteria, at MIC = 8–16 μg/mL, but they exhibited lower antifungal activity (MIC = 31.25–62.5 μg/mL). Oxadiazole 240 functionalized with a thiol group at position 2 exhibited excellent antibacterial activities against all bacterial strains at MIC = 4–8 μg/mL and good activity towards fungal strains at MIC =16–31.28 μg/mL (Figure 30) [76].

Antimicrobial activity screening of 2-(bis(1,3,4-oxadiazolyl/1,3,4-thiadiazolyl)methylene)malononitrile derivatives (241a–c, 242a–c) against S. aureus, B. subtilis (Gram-positive bacteria) and E. coli, K. pneumoniae (Gram-negative bacteria) revealed that compounds having a thiadiazole moiety (242a–c) exhibited high activity against both Gram-positive and
Gram-negative bacteria (IZD = 17–35 mm). However, the oxadiazole compounds (241a–c) displayed moderate activity towards Gram-positive bacteria (IZD = 10–20 mm). Similarly, compounds having thiadiazole rings (242a–c, IZD = 18–34 mm) exhibited higher antifungal activity than those with an oxadiazole moiety (241a–c, IZD = 12–22 mm) against C. albicans, A. niger, and F. solani (Figure 31) [99].

Sekhar et al., prepared new classes of methylthio-linked pyrimidinyl 1,3,4-oxadiazoles and 1,3,4-thiadiazoles. Screening the antibacterial activity indicated that the title compounds were more effective against Gram-negative bacteria than Gram-positive ones. The pyrimidinyl bis-thiadiazoles 243 exhibited greater activity than pyrimidinyl bis-oxadiazoles. The presence of an electron-withdrawing group on the aromatic ring increased the activity. The p-chlorophenyl 243a and p-nitrophenyl 243b derivatives (MIC 6.25 μg/well) were effective, particularly against P. aeruginosa, and were equipotent to Chloramphenicol (MIC 6.25 μg/well) (Figure 32) [104].
In 2019, a new class of aroylethenesulfonylmethyl/arylsulfonylethenesulfonylmethyl styryl 1,3,4-oxadiazoles, 1,3,4-thiadiazoles (Figure 33), was synthesized and evaluated for antimicrobial activity. It was observed that compounds having thiadiazole (19c, 20c) moiety exhibited higher antimicrobial activity than those with the oxadiazole moiety (17c, 18c). Additionally, aroylethenesulfonylmethyl styryl azoles (18c, 20c) exhibited greater antimicrobial activity than aroylethenesulfonylmethyl styrylazoles (17c, 19c). Moreover, compounds with an electron-withdrawing chloro substituent on the phenyl ring possessed enhanced activity [6].

The potential activity of 2-(arylaminosulfonyl methyl)-1,3,4-oxa(thia)diazole derivatives (compounds 33a–c to 36a–c) as antimicrobial agents was screened in vitro against a panel of Gram-positive bacteria—S. aureus, B. subtilis—and Gram-negative bacteria—K. pneumoniae and P. vulgaris and fungi F. solani, C. lunata, and A. niger. The results showed that compounds with the thiadiazole moiety (35a–c, 36a–c) exhibited higher antimicrobial activity than those with oxadiazole units (33a–c, 34a–c). In particular, compounds (7a–c) carrying the thiadiazole moiety displayed potent activity against the tested Gram-positive bacteria and fungi (IZD > 26 mm) and good activity against Gram-negative bacteria (IZD > 20 mm). Thiadiazole derivatives featuring the 5-arylsulfonylmethane moiety (36a–c) displayed higher activity than their respective 5-aryl counterparts (35a–c) against all of the tested microorganisms. Additionally, the antimicrobial effect was enhanced upon grafting a Cl-substituent to the phenyl ring (Figure 34) [50].

A series of 5-(styryl/pyrrolyl/pyrazolylsulfonylmethyl)-2-(arylaminosulfonylmethyl) 1,3,4-oxa(thia)diazoles (Figure 35) were prepared and tested for antimicrobial activity against S. aureus, B. subtilis (Gram-positive bacteria), P. aeruginosa and K. pneumoniae (Gram-negative bacteria). The data showed that the presence of a chloro substituent on phenyl rings of thiadiazoles (31c, 245c, 247c) enhanced the activity against the tested bacteria. The corresponding oxadiazole analogs (30a–c, 244a–c, 246a–c) proved to be less potent antimicrobial agents. It was also observed that the styryl derivatives (30a–c, 31a–c) displayed higher potency than their pyrrolyl/pyrazolyl counterparts (244a–c–247a–c) [49]. An extract from the antimicrobial structure–activity relationship of substituted 1,3,4-oxa-/thiadiazoles is illustrated in (Figure 36a,b.)
Figure 35. Structures of compounds 30a–c, 31a–c and 244a–c–247a–c.

Ar=Different antimicrobial pharmacophores: long alkyl chain substituted phenyl pyrazolyl indolyl benzimidazolyl quinolinyl naphthyridinone sesquiterpenoids

1,3,4-thiadiazoles are more active antimicrobial agents than 1,3,4-oxadiazoles

(a)

Ar= Alkyl substituted styryl substituted phenyl

substitution of phenyl or styryl with electron withdrawing group like chlorine, increases antimicrobial activity

1,3,4-thiadiazoles are more active antimicrobial agents than 1,3,4-oxadiazoles

(b)

Figure 36. (a) Collective antimicrobial SAR for thioxo-, amino-, and amido-substituted 1,3,4-oxa-/thiadiazoles. (b) Collective antimicrobial SAR for arylsulfide-, sulfone-, and sulfonamide-substituted 1,3,4-oxa-/thiadiazoles.
3.2. Antiviral Activity

A series of 4-oxo-4H-pyrido[1,2-a]pyrimidine derivatives containing 1,3,4-oxa(thia)diazole rings as a part of the metal chelation motif were screened for their in vitro anti-HIV-1 activity by determining their ability to inhibit the replication of HIV-1 in Hela cell cultures. All of the tested compounds were safe with no cytotoxicity at a concentration of 100 μM. Compounds containing thiadiazole rings were more potent than the corresponding oxa(diazole analogs. The highest degrees of inhibition against HIV-1 (NL4-3) were reported for the thiadiazoles 63b and 63c: 51% and 48%, respectively. The docking study revealed that the anti-HIV activity of these compounds might involve a metal chelating mechanism (Figure 37) [59].

![Figure 37. Structure of compounds 63b,c.](image)

The antiviral activity of a series of thioether/sulfone compounds containing 1,2,3-thiadiazole and 1,3,4-oxa(thia)diazole rings was tested against tobacco mosaic virus TMV using Ningnanmycin as a reference drug. The data results revealed that the compounds containing 1,2,3-thiadiazole incorporating 1,3,4-oxadiazole ring 248a, 248b, 248c, and 250a showed equipotent activities to Ningnanmycin with curative effects ranging from 46.8% to 54.1%. However, a slight lowering of activity was noticed with the 1,3,4-thiadiazole isosteres (249, 251). Oxidation of thioethers (248, 249) to their corresponding sulfones (250, 251) slightly reduced the values of antiviral activities (Figure 38) [105].

![Figure 38. Structures of compounds 248a-d–251a-d.](image)

3.3. Anticancer Activity

A series of N-(4-substitutedphenyl)-5-(pyridin-4-yl)-1,3,4-oxa(thia)diazol-2-amines were synthesized and evaluated with respect to their in vitro cytotoxicity against six human cancer cell lines, including cells derived from human gastric cancer (NUGC), human colon cancer (DLD1), human liver cancer (HA22T and HEPG2), nasopharyngeal carcinoma (HONE1), human breast cancer (MCF), and normal fibroblast cells (WI38) using CHS 828, a pyridyl cyanoguanidine, as a standard antitumor drug. The overall results revealed superior activity of 1,3,4-thiadiazoles compared to that of their 1,3,4-oxadiazoles bioisosteres, where only one oxadiazole compound 80b showed selective moderate activity against NUGC and DLD1 cell lines. Among 1,3,4-thiadiazoles, only the compounds bearing 4-chlorophenyl (81b) and 4-bromophenyl 81c pharmacophores were found to be active, which proves that the electronegativity of substituents plays an essential role in the activity. Compound (81c) showed nearly equipotent activity against the NUGC cell line compared to the standard CHS 828, with IC₅₀ = 0.028 μM, while 81c also showed 4-fold higher activity...
against HA22T and HEPG2 cell lines compared to its analog 81b (IC₅₀ = 0.180, 0.264 μM and 0.753, 0.970 μM, respectively) (Figure 39) [69].

Figure 39. Structures of compounds 80b and 81b,c.

Rajak et al., prepared two series of hydroxamic acid-based 1,3,4-oxa-/thiadiazoles as histone deacetylase inhibitors and examined their antiproliferative activities in vitro using histone deacetylase inhibitory assay and MTT assay. They were also tested for antitumor activity against Ehrlich ascites carcinoma cells in Swiss albino mice. The results showed that among the oxadiazole series, compound 252 was the most potent, displaying the maximum HDAC inhibitory activity with an IC₅₀ = 0.017 μM against HDAC-1 and an IC₅₀ = 0.28 μM in HCT-116 cell proliferation assay, as well as %TWI = 85.7 and %TCI = 77.7 against Ehrlich ascites carcinoma cells in Swiss albino mice. In the thiadiazole series, compound 253 was the most potent, displaying the maximum HDAC inhibitory activity, with an IC₅₀ = 0.018 μM against HDAC-1 and an IC₅₀ = 0.31 μM in HCT-116 cell proliferation assay, and %TWI = 75.0; %TCI = 76.8 against Ehrlich ascites carcinoma cells. An overview of the results shows that thiadiazoles were found to be more active than the corresponding oxadiazole (Figure 40) [106].

Figure 40. Structures of compounds 252 and 253.

The in vitro antiproliferative activity of a series of benzosuberones embedded with 1,3,4-oxa(thia)diazole moieties was examined against four human cancer cell lines (cervical HeLa, breast MDA–mB-231, pancreatic PANC1, and alveolar A549), where Paclitaxel, Nocodazole, Colchicine, Combretostatin, and Doxorubicin were used as standard drugs. The reported data showed that compounds 204a, 254, and 255 showed potent activity, with GI₅₀ values ranging from 0.079 μM to 0.957 μM against the four human cancer cell lines. However, compound 254 was nearly equipotent to Colchicine against the HeLa cell line with GI₅₀ = 0.079 μM. Based on the overall results, it was concluded that benzosuberones attached to oxadiazoles were more active than the corresponding thiadiazoles (Figure 41) [98].

The cytotoxic activities of a series of 2-(5-methyl-2-nitrophenyl)- 5-(substituted)-1,3,4-oxa(thia)diazoles were tested using Podophyllotoxin as a standard drug. The unsymmetrical 1,3,4-oxadiazole compounds with thiophene or chloropyridine substitution at the 5th position (56c, 56e) and the 1,3,4-thiadiazole compound with furan substitution at the 5th position 57b showed significant activity (IC₅₀ = 10–19.45 μg/mL) compared to the standard podophyllotoxin (IC₅₀ of 3.64 μg/mL), in contrast to the inactive symmetrical oxadiazole 56a (Figure 42) [55].
Ar= a) 2-nitro-5-methylphenyl, c) 2-thienyl, e) 6-chloropyrid-3-yl

Figure 42. Structures of compounds 56a,c,e, and 57b.

In the same year, two series of novel diosgenin derivatives bearing 1,3,4-oxadiazole or 1,3,4-thiadiazole moieties were evaluated for their cytotoxicity in four human cancer cell lines (HepG2, A549, MCF-7, and HCT-116) and normal human gastric epithelial cells (GES-1) using the MTT assay in vitro. The data revealed that the 1,3,4-thiadiazole series were more active than the 1,3,4-oxadiazole series against HepG2 and A549 cells. Among the 1,3,4-thiadiazoles, compound 60d (Figure 43) with a 3-pyridyl group at the C5 position was the most potent. It was 6.7-fold more potent than the parent drug diosgenin (DG) against the A549 cell line (IC_{50} = 3.93 \mu M and 26.41 \mu M, respectively), as well as being 2.9-fold more potent than DG against the HepG2 cell line (MIC = 11.73 \mu M and 33.87 \mu M, respectively). Moreover, compound 60d displayed low toxicity against non-cancer cells GES-1 cells (IC_{50} = 420.4 \mu M), showing high selectivity [58].

Figure 43. Structure of compound 60d.

On the basis that combining different heterocyclic nuclei may show synergistic effects, Kamal and Sobhy decided to merge the two bioisosteres, 1,3,4-oxadiazole and 1,3,4-thiadiazole, in one system to enhance the biological activity of the synthesized compounds.
and evaluated their cytotoxicity against a human colon carcinoma cell line (HCT-116) using Doxorubicin as a reference drug. Among all the tested compounds, the most active derivatives were 257d and 258c (IC₅₀ = 0.73 and 0.86 µg/mL, respectively), while the other compounds showed moderate to poor activity compared to Doxorubicin IC₅₀ = 0.42 µg/mL. The assay results and an examination of the structure–activity relationship revealed that, as a substituent at position 2 of 1,3,4-thiadiazole, the acetyl group (COCH₃) resulted in higher activity than the anilide group (CONHPh) (258a–c), which was better than the phenyl moiety (Ph) 256. Additionally, for substituents at position 4 of 1,3,4-thiadiazole, 4-ClC₆H₄ > 4-NO₂C₆H₄ > 4-OCH₃C₆H₄ > 4-CH₃C₆H₄ > 4-BrC₆H₄ > C₆H₅ (Figure 44) [107].

![Figure 44. Structures of compounds 256, 257a–f, and 258a–c.](image)

Based on the same idea, a series of novel hybrid molecules containing 1,3,4-oxadiazole and 1,3,4-thiadiazole bearing the Schiff base moiety were designed and evaluated with respect to their in vitro antitumor activities against SMMC7721, MCF-7, and A549 human tumor cell lines by CCK-8 assay. The data revealed that some compounds were more potent than the positive control, 5-Fluorouracil (5-FU), against various cell lines. Among these compounds, compound 259b (4-chloro) was the most potent against SMMC-7721 cells, with IC₅₀ = 2.84 µM. Compounds 259c (4-methoxy) and 259d (4-nitro) displayed highly effective antitumor activities against MCF-7 cells, with IC₅₀ = 4.56 and 4.25 µM, respectively. The unsubstituted phenyl derivative 259a and the 4-nitro-substituted derivative 259d showed significant activity against A549 cells, with IC₅₀ = 4.11 and 4.13 µM, respectively. The pharmacological results suggest that the substituents of the phenyl ring on the 1,3,4-oxadiazole are important for modulating antiproliferative activities against various tumor cell lines (Figure 45) [108].

![Figure 45. Structure of compounds 259a–d.](image)

In 2015, a series of 2,5-Bis[2-substituted-1,3,4-oxa(thia)diazol-5-yl]propylthio]-1,3,4-thiadiazoles were tested for their in vitro antiproliferative activities in four different human cancer cell lines (human breast adenocarcinoma, MCF7, human ductal breast epithelial tumor, T47D, human colorectal adenocarcinoma, Caco-2, and human epithelial carcinoma, He La). All compounds demonstrated relatively high activities against the examined cell lines. There was no difference in activity related to the replacement of oxadiazole with a thiadiazole ring, while changing methyl and phenyl groups in (101a,c) and (102a,c) into ethyl groups 101b and 102b significantly enhanced the cytotoxic activity, with LD₅₀ values...
ranging from 376 ng/µL to 438 ng/µL, which suggests a steric factor mediating either transport or molecular interaction of these compounds with cellular targets. Furthermore, the addition of one more Cl atom into the structure of compound 260a (LD₅₀ = 648–690 ng/µL) gave compound 260b, with double the activity (LD₅₀ = 356–398 ng/µL) (Figure 46) [76].

![Figure 46. Structures of compounds 101a-c, 102a-c and 260a,b.](image)

Ozdemir et al. designed and evaluated several oxadiazole and thiadiazole derivatives with respect to their anticancer effects on A549 human lung adenocarcinoma and C6 rat glioma cell lines and evaluated their inhibitory effects on matrix metalloproteinases (MMPs). Compounds 261, 263, and 264 showed high cytotoxic activity against the C6 cell line, with IC₅₀ ranging from 0.0128 mM to 0.157 mM, compared to Cisplatin, with IC₅₀ = 0.103 mM. Only the benzodioxole-substituted oxadiazoles 263 and 264 had a cytotoxic effect on the A549 cell line, with IC₅₀ values of 0.125 and 0.349 mM, without causing any toxicity towards the NIH/3T3 mouse embryonic fibroblast cell line, which suggests that the (1,3-benzodioxol-5-ylmethyl) amino group enhances the antitumor activity against A549 cells. At the same time, the benzodioxole-substituted thiadiazole analog 262 was inactive against A549 cell lines, which proves that the oxadiazole scaffold is essential for activity against A549 cells. Compounds 263 and 264 were also the most effective MMP-9 inhibitors in this series. Moreover, docking studies pointed out that compounds 263 and 264 had a good affinity for the active site of the MMP-9 enzyme (Figure 47) [109].

![Figure 47. Structures of compounds 261–264.](image)

The structure–activity relationship of the anticancer activity of substituted 1,3,4-oxadiazoles and thiadiazoles is summarized in Figure 48.
Figure 48. Collective anticancer SAR for substituted 1,3,4-oxa-/thiadiazoles.

3.4. Antidiabetic Activity

In 2019, Taha et al., reported a series of 5-(4-hydroxy-2-methoxyphenyl)-1,3,4-oxadiazoles coupled to 5-aryl-1,3,4-thiadiazole motifs through a phenyl ring as potent β-glucuronidase inhibitors. Several analogs were more potent enzyme inhibitors (IC$_{50}$ = 0.96–28.10 µM) than the standard D-saccharic acid 1,4-lactone (IC$_{50}$ = 48.40 µM). Study of the structure–activity relationships indicated that the nature and relative position of substituents on the phenyl ring attached to thiadiazole greatly affected the inhibitory potency. The 2,4,6-trichlorophenyl derivative 265 displayed superior activity (IC$_{50}$ = 0.96 µM) compared to monochloro-substituted analogs, regardless of the position (o, m or p). Moreover, compound 266a with the 3,4-dihydroxyphenyl moiety (IC$_{50}$ = 1.40 µM) was four times more potent than the 2,5-dihydroxyphenyl analog 267a (IC$_{50}$ = 6.20 µM). A docking study revealed that the hydroxyl groups in 6 had higher potential to be engaged in proper hydrogen bonding interaction with the enzyme active site than in 7. In contrast, compound 267b having 2-hydroxy-5-methoxy groups on the phenyl ring was much more potent than the 3-hydroxy-4-methoxy phenyl derivative 266b, with IC$_{50}$ = 12.30 and 28.10 µM, respectively. The steric effect of the methoxy group next to the hydroxy function in 266b decreased the ability of the hydroxyl group to be involved in H-bonding with the enzyme and reduced the activity (Figure 49) [109].
The results revealed that only the compounds with an oxadiazole unit (241a-c) showed high antioxidant activity in both methods. An SAR study on this series of Nrf2 activators revealed that the 1,3,4-oxadiazole showed good potential to penetrate the BBB and reach the CNS as a neuroprotective agent. Thus, compound 226c represents a new chemotype against oxidative stress and provides a new therapeutic strategy for different neurodegenerative diseases, including Alzheimer’s disease. An SAR study on this series of Nrf2 activators revealed that the 1,3,4-oxadiazole derivatives showed higher Nrf2 inductivity than the 1,3,4-thiadiazole analogs as a result of their improved water solubility and lower lipophilicity [92].

In addition to the reported [99] antimicrobial activity of 2-(bis((1,3,4-oxadiazolyl/1,3,4-thiadiazolyl) methylthio)methylene)malononitrile derivatives (241a-c, 242a-c) (Figure 31), the compounds were tested for the antioxidant property by NO and DPPH methods. The results revealed that only the compounds with an oxadiazole unit (241a-c) showed high antioxidant activity in both methods.

Nabhubygari et al. tested the antioxidant activity of a variety of symmetrical 2,4-bis-oxazolyl/thiazolyl/imidazolylacetamido-sulfonylmethyl 1,3,4-oxadiazoles (43a-c, 44a-c, 45a-c) and 1,3,4-thiadiazoles (49a-c, 50a-c, 51a-c) (Figure 51) using 2,2-diphenyl-1-picryl-hydrazyl (DPPH) and nitric oxide (NO) methods. The results showed that oxadiazole derivatives (43a-c, 44a-c, 45a-c) exhibited higher activity than the corresponding thia-
diazoles (49a-c, 50a-c, 51a-c). Among the tested compounds, the bis-(4-methylphenyl oxazolyl)oxadiazole 43b and bis-(4-methylphenyl oxazolyl) thiadiazole 49b were the most potent antioxidants, and they demonstrated higher activity than the standard ascorbic acid in both methods. These results suggest that the presence of an electron-donating methyl substituent on the aromatic ring enhanced the antioxidant activity [51].
Similar results were observed upon evaluating the antioxidant potential of sulfone-/sulfonamide-linked bis(oxadiazoles) and bis(thiadiazoles) using the DPPH and NO methods. The 4-methylphenyl sulfone/sulfonamide-linked bis(oxadiazoles) 24b and 25b presented higher antioxidant activity than Ascorbic acid. The SAR of the tested compounds revealed that compounds with oxadiazole rings (24a–c and 25a–c) showed higher radical scavenging activity than the corresponding thiadiazoles (26a–c and 27a–c). It was also observed that sulfonamide-linked bis(oxadiazoles) and bis(thiadiazoles) (25a–c and 27a–c) showed comparatively higher antioxidant activity than sulfone-linked bis(oxadiazoles) and bis(thiadiazoles) (24a–c and 26a–c) (Figure 52) [2].

NO and DPPH scavenging activities were assayed for a series of 2-(arylaminosulfonyl methyl)-1,3,4-oxa(thia)diazole derivatives (33–36) (Figure 54). The results highlighted that the compounds with an oxadiazole unit (33a–c, 34a–c) exhibited better antioxidant activity than did those with a thiadiazole unit (35a–c, 36a–c), where compounds 33a, 33c, 34a and 34c showed high potency in both methods at a concentration of 100 μM [50].

A summary of the antioxidant structure–activity relationship for substituted 1,3,4-oxa-/thiadiazoles is shown in Figure 53.
3.6. Analgesic and Anti-inflammatory Activities

In 2013, some N-(tetrazol-1H-5-yl)-6,14-endoethenotetrahydrothebaine 7α-substituted 1,3,4-(oxa)thiadiazoles were prepared and screened for analgesic activity in rats, applying hot-plate and tail-flick tests and employing Morphine as a reference drug. At 0.5 and 1 h after administration, the phenylamino-oxadiazole derivative 268 (reaction time in hot plate test = 9.78 s, tail-flick test = 21.45 s) was found to be more potent than Morphine (reaction time in hot plate test = 6.28 s, tail-flick test = 15.90 s). Conversely, the thiadiazole analog 269 was moderately active. The results demonstrated that the oxadiazole group on the 7α position of thebaine increased the analgesic activity (Figure 54) [60].

Figure 54. Structures of compounds 268 and 269.

A series of polyheterocyclic thioethers containing 1,3,4-(oxa)thiadiazoles were examined for anti-inflammatory activity using carrageenan-induced rat paw edema assay and Diclofenac as a reference drug. Compounds 270, 271a, and 271b were more effective than Diclofenac in alleviating carrageenan-induced edema after 2, 3, and 4 h of treatment, with inhibition percentages of 37.5–55%, 30–35%, and 38–48%, respectively. In this study, oxadiazole and thiadiazole isosteres were presented as potent anti-inflammatory agents with no difference in their activities (Figure 55) [110].

Figure 55. Structures of compounds 270 and 271a,b.

4. Conclusions

This study is a review article summarizing most of the published work about the medicinally important pharmacophores 1,3,4-oxadiazoles and 1,3,4-thiadiazoles throughout the past 10 years. The review presents different synthetic approaches to the two bioisosteres using different starting compounds, comparing the differences in the reagents used, the conditions, and the yields of the two classes. This comparative study also identified their pharmacological activities, as well as introducing deduced collective structure–activity relationship charts for antimicrobial, anticancer, and antioxidant activities. In most cases, it was concluded that 1,3,4-thiadiazoles are more active antimicrobial agents than 1,3,4-oxadiazoles, in contrast to antioxidant activity, where 1,3,4-oxadiazoles are more active agents than 1,3,4-thiadiazoles. Concerning anticancer activity, it was noticed that combining the 1,3,4-oxadiazole ring with the 1,3,4-thiadiazole ring enhanced the activity.

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