Isothiazoles in the Design and Synthesis of Biologically Active Substances and Ligands for Metal Complexes

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Abstract The chemistry of isothiazoles is being intensively developed, which is evidenced by the wide range of selective transformations involving the isothiazole heterocycle and the high biological activity of its derivatives that can be used as effective new drugs and plant protection chemicals. Some representatives of isothiazoles have proven to be synergists of bioactive substances, which opens the way to lower the doses of drugs used and is especially important in cancer chemotherapy. In the framework of the present review, the accomplishments in the chemistry of isothiazoles over the past 18 years are examined, whilst current strategies for the synthesis of isothiazole-containing molecules and key directions of studies in this field of heterocyclic chemistry are discussed. Considerable attention is paid to chlorinated isothiazoles and strategies for their use in the synthesis of biologically active substances. In addition, a comprehensive review of existing literature in the field of metal complexes of isothiazoles is given, including the results and prospects for the practical use of isothiazole–metal complexes as catalysts for cross-coupling reactions in aqueous and aqueous–alcoholic media (‘green chemistry’).

1 Introduction

The search for new chemical compounds with useful properties and the development of rational ways to synthesize them are priority tasks in chemical science, with the synthesis of biologically active chemical compounds for medicine and agriculture being of particular importance. In this respect, isothiazole derivatives, which have demonstrated high potential in the design and synthesis of a wide range of biologically active substances, are of great interest. Among natural bioregulators, isothiazole-containing compounds are present in only a few examples: the phytoalexins (brassilexin1 and sinalexin),2 a prostaglandin release inhibitor (pronkodin A),3 and a cytotoxin (aulosirazol).4 However, this does not impede the use of the isothiazole nucleus in the creation of a wide variety of bioactive substances; for example, compounds exhibiting anti-poliovirus activity have been synthesized.5 Promising compounds for the treatment of Parkinson’s disease have also been identified,6 and recently, derivatives suitable for cancer7–12 and diabetes13–15 therapy have been obtained, as well as microbiocides.16,17 The isothiazole heterocycle in microbicidics can protect the molecule from the action of enzymes,18,19 thereby extending the time of the molecule’s action, which in turn makes it possible to administer additional doses of the drug less frequently. A promising class of isothiazole-containing inhibitors of the nuclear bile acid receptor FXR has been found, representatives of which can be used in the treatment of liver diseases.20 Some isothiazoles can be successfully used to create competitive antagonists of insect GABA receptors,21 and among the isothiazoles, effective plant growth regulators22 and fungicides23,24 have been found.
Biographical Sketches

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Fedor Ivanovich Zubkov earned his Master’s degree in chemistry at the Peoples’ Friendship University of Russia in 1997, followed by his Ph.D. in chemistry in 2000. From 2001 until present, he has worked as a docent at the Organic Chemistry Department of the Peoples’ Friendship University of Russia. His scientific interests include cycloaddition reactions, transformations of oxygen- and nitrogen-containing heterocycles, and total synthesis.

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Vladimir Ivanovich Potkin graduated from Gorky State University (now Nizhny Novgorod, Russia) in 1975. He received his Ph.D. in 1982 from the Institute of Physical Organic Chemistry of the Belorussian Academy of Sciences, and his D.Sc. in 1996 from Belarusian State University. In 2000 he was elected as a Corresponding Member of the National Academy of Sciences of Belarus, and in 2009 he earned the title of Professor of Chemistry. Currently, he is the Head of the Organic Chemistry Department at the Institute of Physical Organic Chemistry, National Academy of Sciences of Belarus. His scientific interests include heterocyclic chemistry, the synthesis and study of biologically active compounds, and metal complexes of isothiazoles and isoxazoles and their properties.
In a few reports, isothiazoles have been studied for possible use as ligands for metal complexes. At the same time, despite the fact that metal complexes in general have proven themselves useful in creating a wide range of new technologies and biologically active substances, ways to apply isothiazole complexes in practice have rarely been investigated. Some recent publications, in which the promise of using metal complexes of isothiazole for the catalysis of organic reactions and the development of pesticides has been shown, are the exception.

The above facts demonstrate the high potential of using isothiazole heterocycles in the design and synthesis of compounds for different purposes and indicate broad prospects for the further development of isothiazole chemistry, especially coordination chemistry. Within the framework of the present review, achievements in isothiazole chemistry over the past 18 years will be covered, and references from previous years will be mentioned only to create a holistic picture of the recent research progress in this field.

This review deliberately does not include data on isothiazolinones, 1,1-isothiazoledioxides, and dihydroisothiazoles, and only some examples related to condensed isothiazole-containing systems are considered. This is due to the fact that, in our opinion, such compounds, although being related, differ significantly in their chemical properties from 1,2-thiazole, and a consideration of their chemical transformations as well as their practical use would lead to a significant increase in the volume of this paper and distract from the subject being reviewed. However, special attention is paid to halogen-containing isothiazoles, and in particular, chlorinated isothiazoles, which are convenient synthetic building blocks that allow the ability to obtain a wide variety of substituted isothiazoles capable of acting both as biologically active substances and as ligands for metal complexes.

In the general context of the chemistry of heterocyclic compounds, data on the chemical properties of isothiazoles are presented in review articles, whilst isothiazoles have been directly reviewed and covered in a monograph devoted to biologically active isothiazoles. There are also reports summarizing data on isothiazole-ring-containing compounds in crop protection, the involvement of the isothiazole core in cross-coupling reactions, and [4+2] cycloadditions of isothiazole derivatives.

There are a significant number of methods for isothiazole synthesis, and so in order to present the material consistently as well as to form a holistic view of isothiazole chemistry, it would be rational to first consider the approaches to isothiazole cycle design, highlighting the main synthetic strategies. Next, some methods for the functionalization of isothiazoles will be reviewed, and after that, targeted syntheses of biologically active agents that are promising for practical use will be considered. The available information on metal complexes with isothiazole ligands is considered in the final part of the literature review, and will, in our opinion, provide a comprehensive picture of developments in the field of isothiazole metal complexes, including information that has escaped the attention of other authors for one reason or another.

## 2 Synthesis by Ring-Forming Reactions

Different approaches can be used to classify the methods for producing isothiazoles, and the classification proposed herein by us is based on retrosynthetic analysis. This allows us to build a consistent and logical understanding of the approaches to the production of substituted isothiazoles, and in some cases, to provide the most rational methods for their synthesis. However, the predictive efficiency of this retrosynthetic classification is low in the case of complex reactions with an unobvious mechanism, as the proposed classification does not always reflect the true mechanism of the reaction. Yet, it does allow us to visualize the possible conditions for the preparation of isothiazoles, representing the formation of an isothiazole ring from fragments containing a certain number of atoms. It certainly does not cover all possible methods for the preparation of substituted isothiazoles, but it reflects the key stages of most approaches to their synthesis. Overall, retrosynthetic analysis allows us to distinguish four rational approaches for the formation of the isothiazole ring.

### 2.1 Intramolecular Cyclization

An example of such an approach is the synthesis of substituted isothiazoles 1 by forming an S–N bond in an S–C–C–N fragment via oxidative cyclization of substituted 3-aminopropenethiones 2 under the action of various reagents. The classic version of this approach uses iodine; however, methods using other oxidizing agents, such as hydrogen peroxide, have recently become very popular (Scheme 1). Hydrogen peroxide, in particular, was used in the formation of the isothiazole nucleus at the penultimate stage in the synthesis of a series of allosteric antagonists of the mGluR1 receptor, and which are promising compounds for the treatment of various pain syndromes.

![Scheme 1](image-url)
Along with iodine, bromine can be used for effective oxidative cyclization. Thus, the isothiazoles 4, which are inhibitors of hepatitis C polymerase NS5B as well as precursors of the inhibitors of the important cancer therapy kinases Chk2 and MEK1, have been synthesized by oxidizing N-substituted propanethioamides with molecular bromine in ethyl acetate (Scheme 2).42–44

From a preparative point of view, a promising variation of this approach is the oxidation of isothiazole precursors without the use of a solvent. Thus, it has been shown that the solvent-free oxidative cyclization of 3-aminopropenethiones 5 can be carried out by using chromium trioxide supported on silica gel (Scheme 3).45 It should be noted that the yields of the corresponding 4-cyanoisothiazoles 6 did not depend on whether the reaction was performed at room temperature for 2–3 hours, or for 1–2 minutes under microwave irradiation.

The closure of the C–N–S–C–C fragment with the formation of a C–C bond can serve as another interesting example of intramolecular cyclization. Using this approach, the formation of an isothiazole ring in the synthesis of the VEGFR1 and VEGFR2 inhibitors was carried out.46 The intramolecular condensation of an activated methylene fragment with a cyano group in compounds 7 led to isothiazoles 8, of which further directed modification allowed the target compounds to be obtained (Scheme 4).

As an important example of the formation of the isothiazole nucleus by the closure of a five-membered fragment, a recently proposed method for the synthesis of reactive 3-chloroisothiazole (9) should be mentioned. The 3,3′-disulfanyldipropanoic acid diamide 10 can be cyclized to give 3-chloroisothiazole (9) by the action of phosphorus oxychloride in toluene (Scheme 5).47

2.2 (4+1)-Heterocyclization

In the framework for this approach, we can consider the heterocyclization of synthetic equivalents of synthons, which are formed when one atom is removed from the isothiazole ring, with the synthetic equivalents of the corresponding atom. The most relevant examples of the heterocyclization of compounds corresponding to synthons with one heteroatom removed from a 1,2-thiazole ring are given below.

Recently, an original method for producing substituted furo[2,3-c]isothiazoles 11 from 2-[amino(3-aryl-2-cyanooxiran-2-yl)methylene]malononitriles 12 in an aqueous dioxide medium with sodium thiocyanate as a sulfur donor was described (Scheme 6).48 It is noteworthy that in previous studies the only proposed method for the synthesis of furo[2,3-c]isothiazoles was via intramolecular nucleophilic substitution with the participation of azide and thione fragments.49,50

It was found that an S₂Cl₂ molecule can act as a sulfur donor in the synthesis of isothiazoles, as was shown when preparing dinitrobenzoyl[3]isothiazole 13 from 2-amino-4,6-dinitrotoluene (14) (Scheme 7).51 The key value of the
nature of the base used in the first stage of the reaction was revealed: when N-ethyldisopropylamine or triethylamine were used individually, the yield of the desired isothiazole 13 did not exceed 18%. However, with the sequential introduction of 1,4-diazabicyclo[2.2.2]octane (DABCO), and then triethylamine, a 90% yield of isothiazole 13 was achieved.

A recently proposed method for the synthesis of 3,4-dichloroisothiazole-5-carboxylic acid 2-cyanoanilide, a class of chlorinated isothiazoles but is also a precursor of ed/annulated isothiazoles esters/esters, was utilized.22,25 This approach involves the interaction of two chemical compounds that act as suppliers of diatomic and triatomic fragments to form an isothiazole ring.

Recently, 3,5-aryl-substituted isothiazoles 21 have been synthesized based on unsaturated N-sulfonfyl ketimines 18, via the cascade addition of an $\text{S}_2\text{Cl}_2$ radical anion and subsequent reductive desylation (Scheme 9).53 The lowest yield (14%) was observed for the thienyl derivative ($R_1 = 2$-thienyl, $R_2 = 4$-MeC$_6$H$_4$).

A new protocol for the synthesis of 3,5-disubstituted/annulated isothiazoles 19 utilizing $\beta$-keto dithioesters/ethyldithioamides 20 and NH$_4$OAc under metal-free conditions was reported in 2016 (Scheme 10).54 The strategic $(4+1)$ annulation initiated by NH$_4$OAc is carbon-economic and relies on a sequential imine formation/cyclization/aerial oxidation cascade forming consecutive C–N and S–N bonds in one pot.

Substituted isothiazoles 21 were able to be isolated as minor products when $\beta$-keto dithioesters 22 were reacted with ammonia, which acts as a donor of the nitrogen atom to construct a 1,2-thiazole heterocycle. Additionally, it was found that copper acetate promotes formation of the isothiazole heterocycle (Scheme 11).55

**2.3 (3+2)-Heterocyclization**

This approach involves the interaction of two chemical compounds that act as suppliers of diatomic and triatomic fragments to form an isothiazole ring.

Recently, a series of 4-arylisothiazoles 24, as inhibitors of the detoxification of the isothiazole-containing brassilex fungicide in L. maculans fungus, was synthesized using this approach. Thus, the corresponding $\alpha,\beta$-unsaturated aldehydes 23 were introduced into the reaction with ammonium thiocyanate in dimethylformamide to give the 4arylisothiazoles 24 (Scheme 12).56 The ammonium thiocyanate in the reaction acts as a donor of the N–S fragment.

According to a similar scheme, isothiazoles 25 were obtained from (Z)-3,4-diyral-3-chloroacrylaldehydes 26. Compounds 25 demonstrate inhibitory activity against COX-1 and COX-2 cyclooxygenases, 5-LOX lipoxygenase,57 and potentially against mitogen-activated p38α protein kinase. They are also considered as promising candidates for in vitro and in vivo testing for anti-neuroinflammatory activity and neuroprotective properties (Scheme 13).58
The 1,3-dipolar cycloaddition reaction, also known as the Huisgen cycloaddition, represents another method that can be used for isothiazole cycle formation and should be mentioned within the framework of the regarded approach. The reaction of functionalized acetylenes with nitrile sulfides—reactive dipoles generated from 2-oxo-1,3,4-oxathiazoles in aprotic polar solvents such as chlorobenzene and toluene—can serve as an example. In particular, 3-phenylsulfamid-5-carboxylic acid 27a and 3-phenylsulfamid-4-carboxylic acid 27b were obtained from phenyloxathiazolone 28 by this method (Scheme 14), and the synthesized products showed low in vitro cytotoxicity along with the ability to protect cells from being infected with HIV-1.39

The efficiency of using microwave irradiation in such reactions has been demonstrated. This method allows avoiding long reaction times and, in some cases, reduces the amount of by-products and increases the yields of the target compounds.60 Thus, phenyloxathiazolone 29 was reacted with dimethyl acetylenedicarboxylate (DMAD) in chloroform under microwave irradiation to form adduct 30 in a yield of 56%, while the reaction with propiolic acid ethyl ester (EP) for 10 minutes resulted in a mixture of regioisomeric isothiazoles 31a and 31b with a total yield of 48% (Scheme 15). The authors noted that a low level of regioselectivity was a common factor for reactions of nitrile sulfides, in contrast to reactions of nitrile oxides, which led to isoxazole-5-carboxylates as the main products. This effect is usually attributed to a higher degree of HOMO-dipole control for nitrile sulfide reactions.

Finally, a very interesting and original example of (3+2)-heterocyclization is the recently proposed method of asymmetric synthesis of condensed isothiazoles 32 through a copper-catalyzed direct asymmetric conjugate addition of allyl cyanide (33) to α,β-unsaturated thioamides 34 (Scheme 16).61

It is important to note that the yield of the target isothiazoles 32a–d increased with the introduction of preliminary isolated products of the allyl cyanides addition 35 to α,β-unsaturated thioamides into the reaction (Scheme 17).

2.4 Syntheses by Ring Transformations

Formally, reactions for the synthesis of substituted isothiazoles on the basis of other heterocycles can be considered within the framework of the three previous approaches, since they usually proceed through the stage of breaking the bonds in the initial cyclic molecule and subsequent heterocyclization of the resulting intermediate. However, in our opinion, it is advisable to single out the reactions that result in the transformation of other heterocycles into isothiazoles as a separate approach. This is due, first of all, to the rather complicated mechanism of such reactions, and to a certain non-observability in choosing possible synthetic equivalents when analyzing the structures of the desired isothiazoles.
An important example of this approach is the synthesis of isothiazoles based on N-substituted 3-isothiazolones. 3-Isothiazolones 36a–i can be transformed into functionalized 3-aminoisothiazoles 37a–i by sequential treatment with phosphoryl chloride and ammonia (Scheme 18).52

Among the methods for the synthesis of isothiazoles based on other heterocycles, the recently proposed method for the synthesis of 3-phenyl(heteroaryl)isothiazole–3-carboxylic acid ethyl esters 38a–d via the tandem photoarylation and photoisomerization of 2-isothiazole-5-carboxylic acid ethyl ester 39 is of considerable interest (Scheme 19).63 It should be noted that no rearrangement occurred in the case of a similar 2-chloro-substituted thiazole.

Recently, an original method for the synthesis of 3,4,5-functionalized isothiazoles 40 was developed through rhodium-catalyzed transannulation of 1,2,3-thiazolones 4144 with alkyl, aryl, and hetaryl nitriles 42 through the formation of a rhodium α-thiavinyl carbenoid. The authors suggest that in the reaction with nitriles, the α-thiavinyl carbenoid acts as an equivalent of a 1,3-dipole with reversed polarity (Scheme 20).

Later, this method was successfully used by the same authors to synthesize a series of bi-, tri-, and tetracyclic isothiazoles 43 with good to excellent yields through the intramolecular transannulation of cyanothiadiazoles 44 (Scheme 21).65

Isothiazoles can be obtained on the basis of 1,2,3-dithiazoles. Recently, improved conditions for the rhodium-catalyzed transannulation of 1,2,3-dithiazoles 45 into isothiazole-5-carbonitriles 46 were reported (Scheme 22).66 Shorter reaction times correlated with higher isothiazole yields and the use of THF as solvent in almost all cases.

### 3 Isothiazoles by Ring Functionalization

**Reactions: Nucleophilic Substitution, Cross-Coupling and Side-Chain Functionalization**

Next, we will consider some methods for the synthesis of isothiazoles in which reagents containing the formed isothiazole ring are involved. The chemical properties of isothiazoles are largely defined by the presence of the delocalized π-orbitals system. Isothiazoles have a high degree of aromaticity in accordance with the HOMA (harmonic oscillator model of aromaticity) criterion. In this respect, among the 1,2-azoles, isothiazoles occupy an intermediate position between isoxazole and pyrazole, being much more aromatic than isoxazoles, but less so than pyrazoles. Isothiazole derivatives are also more aromatic than thiazole derivatives. Positions 3 and 5 of the isothiazole ring are favorable for the attack of nucleophilic reagents, while position 4 is
preferential for the attack of electrophilic agents. It should be noted that position 5 of the isothiazole ring is more active in nucleophilic substitution reactions than position 3. In some cases, interactions with nucleophilic and electrophilic reagents may lead to the ring-opening of the isothiazole.33

The achievements in the development of methods for the functionalization of the isothiazole nucleus and advances in the transformation of the side chains of the heterocycle, without involving the heterocycle itself in the reaction, will be considered separately. Significant attention will be paid to the development of the chemistry of halogen-substituted isothiazoles, which are highly reactive synthetic blocks, allowing the ability to obtain polyfunctional isothiazoles with a wide variety of substituents.

Fairly recently, the possibility of direct oxidative alkenylation of the isothiazole heterocycle at the unsubstituted position 4 was investigated, and it was found that the reactions proceeded with low yields. This was demonstrated by the example of the reaction of 3-methyl-5-phenylisothiazole (47) with n-butyl acrylate (48).67 In addition, along with product 49, 52% of the initial isothiazole 47 was recovered from the reaction mixture (Scheme 23).

It should be noted that the introduction of isoxazole, an isothiazole heteroanalogue, to the reaction under the same conditions led to the formation of the target alkenyl isoxazoles, with yields of 50–68%.

Because of this, the Heck reaction using haloisothiazoles remains the preferred method of alkenylation of the isothiazole heterocycle, despite the moderate yields of the corresponding alkenyl isothiazoles and the formation of a noticeable amount of by-product.68 In recent years, considerable attention has been paid to metatation reactions of the isothiazole core for the subsequent preparation of reactive isothiazole blocks for cross-coupling reactions. Advances in this field of isothiazole chemistry have been discussed in detail by Nutaitis.69

A mild approach to synthesize 3-amino-substituted isothiazole sulfoxides 50 through oxidation of the corresponding 3-aminoisothiazoles 51 with arylsulfonyloxaziridines 52 has been reported (Scheme 24).70 Only minor amounts of dioxides 53 were formed under the described conditions. The reactivity of the resulting isothiazole sulfoxides toward sulfur nucleophiles has been studied and resulted in the formation of 5-sulfanyl-substituted 4,5-dihydroisothiazoles.

In 2019, the same group developed an advanced approach for the synthesis of chiral 3-amino-substituted isothiazole sulfoxides by utilizing (+)- and (–)-[8,8-dichlorocamphoryl]sulfonyloxaziridine under microwave irradiation.71

Relatively little work in the literature is devoted to the transformations of chlorine-substituted isothiazoles; however, these compounds enter into similar reactions to their bromine and iodine analogues, but often under more severe conditions and react at slower rates.

Thus, 3,5-dichloro-4-cyanoisothiazole (54) reacts regio- and stereospecifically with phenylboronic acid by substitution of the chlorine at position 5 and gives 5-phenyl-3-chloro-4-cyanoisothiazole (55) in a high yield. In the same way, substitution of the chlorine at position 5 proceeds with the use of potassium phenyltrifluoroborate (Scheme 25).3,5-Dibromo-4-cyanoisothiazole reacts similarly, but the reaction proceeds at a faster rate.72

In some cases when using chlorine derivatives of isothiazoles, it is possible to synthesize the target compounds with yields greater than those obtained when using bromo derivatives.

Thus, it was found that 3-bromo(chloro)isothiazole-4-carbonitriles 56a and 56b allow direct arylation at position 5 of the heterocycle in the presence of silver fluoride to give the corresponding 5-phenylisothiazoles 57a and 57b (Scheme 26).73 Conditions for the preparative synthesis of disisothiazole derivatives 58a and 58b were also developed. Chlorine derivatives of isothiazole were formed in high yields, both in the case of arylation and in the synthesis of disisothiazole derivatives.

Recently it has been shown that with the introduction of pyrroldine into the nucleophilic substitution reaction of bromine in 3-bromo-4,5-dicyanoisothiazole (59), the main product was 2-[dl(pyrroloidin-1-yl)methylene]malononitrile (60), while the expected product, 4,5-dicyano-3-(pyrrolidin-1-yl)isothiazole (61), was either not formed at all or...
was only formed in small amounts (Scheme 27). By varying the temperature and duration of the reaction, the maximum yield of isothiazole 61 reached 11%.

At the same time, with the introduction of 4,5-dicarbonitrile-3-chloroisothiazole (62) into a similar reaction, a 37% yield of 4,5-dicarbonitrile-3-(pyrrolidin-1-yl)isothiazole (61) was achieved by varying the temperature and reaction time (Scheme 28).

It is noteworthy that Kalogirou and Koutentis found that the most likely precursor of dinitrile 60 is 2-(pyrrolidin-1-yl)ethene-1,1,2-tricarbonitrile (64), which was formed from the substituted isothiazoles 59 and 62 when interacting with pyrrolidine via nucleophilic attack on the sulfur. The resulting 2-(pyrrolidin-1-yl)ethene-1,1,2-tricarbonitrile 65 then gives trinitrile 64 on eliminating the sulfur atom. The subsequent interaction of intermediate 64 with pyrrolidine leads to the formation of dinitrile 60 (Scheme 29).

The chemistry of chlorine-substituted isothiazoles was developed in the Laboratory for Organoelement Compounds of the Institute of Physical Organic Chemistry at the National Academy of Sciences of Belarus. In the course of the research conducted at the laboratory, it was shown that isothiazoles containing chlorine atoms at positions 4 and 5 of the heterocycle and an electron-withdrawing group at position 3, successfully enter into a nucleophilic substitution reaction. When this occurs, the chloride atom at position 5 of the heterocycle is replaced, while position 4 remains unaffected. Research on the transformation pathways of perchloroisothiazoles was initiated with 4,5-dichloro-3-trichloromethylisothiazole, on the basis of which many previously undescribed substituted 4-chloroisothiazoles were synthesized.

Thus, the reactions of 4,5-dichloro-3-trichloromethylisothiazole (66) with N,S-nucleophiles were studied, and it was found that the interactions with heterocyclic amines in dimethylformamide resulted in 5-amino-substituted 3-trichloromethylisothiazoles 67a–d, while reactions with alkyl(aryl)thiolates in ethyl alcohol led to the formation of the corresponding 5-alkyl(aryl)sulfanyl-3-trichloromethyl-4-chloroisothiazoles 68a–c (Scheme 30).
Starting from perchloroisothiazole 66, 5-ethoxy-4-chloro-3-trichloromethylisothiazole (69) and 5-benz oxy-4-chloro-3-trichloromethylisothiazole (70) were synthesized by reactions with sodium ethylate and benzylate in tetrahydrofuran, and were used in the development of new synthetic methods toward 5-hydroxy-4-chloroisothiazoles (Scheme 31).77 Thus, treatment of the 5-ethoxy derivative 69 with hot sulfuric acid allowed 5-hydroxy-4-chloroisothiazolecarboxylic acid (71) to be obtained in a good yield, which was then esterified under acidic conditions to form the methyl ester 72. 5-Benzoylisothiazole 70 was transformed into 5-hydroxy-4-chloro-3-trichloromethylisothiazole (73) in preparative yields by the action of hydrogen in the presence of palladium on carbon (A) or by treatment with trifluoroacetic acid (B).

The reactions of 4,5-dichloroisothiazole-3-carboxylic acid (74), synthesized from 4,5-dichloro-3-trichloromethylisothiazole (66),32 with S,N-nucleophiles have been studied. Reactions with thiols were carried out in diethyl ether in the presence of pyridine, which resulted in 5-alkyl(phenyl)sulfanyl-4-chloroisothiazole-3-carboxylic acids 75a–c in yields of 52–65% (Scheme 32).76

Scheme 31 A new synthetic approach toward 5-hydroxy-4-chloroisothiazoles

Subsequently, 4,5-dichloroisothiazole-3-carboxylic acid tert-butyl ester 76, obtained from acid chloride 77, was used to study the possibility of carrying out reactions with N-nucleophiles (Scheme 33). The possibility of promoting the nucleophilic substitution of the chlorine atom at the 5-position of the heterocycle with amines in the presence of fluorine ions was demonstrated.78 The yield of the corresponding 5-aminoisothiazoles 78 varied depending on the solvent selected, on the presence of fluoride ions in the reaction mixture, on the temperature, and on the nature of the amine added to the reaction.

The reactivity of amines in the nucleophilic substitution of the chlorine atom at position 5 decreases in the series shown in Figure 1.

Scheme 32 Nucleophilic substitution in 4,5-dichloroisothiazole-3-carboxylic acid

Scheme 33 Nucleophilic substitution in tert-butyl ester of 4,5-dichloroisothiazole-3-carboxylic acid

Other derivatives such as 4,5-dichloroisothiazole-3-carboxylic acids 79,79 metalloccenic80 and macrocyclic examples81 as well as various amides 80 were also obtained starting from acid chloride 77 (Scheme 34).

Scheme 34 Synthesis of amides and esters of 4,5-dichloroisothiazole-3-carboxylic acid

Aryl ketones 81a–c were synthesized through the Friedel–Crafts acylation reaction using acid chloride 77 (Scheme 35).82 (4,5-Dichloroisothiazol-3-yl)ferrocenylketone was also prepared in a similar manner.83

Scheme 35 Preparation of 3-ketodervatives of 4,5-dichloroisothiazole

Figure 1
Reactions with O-, N- and S-nucleophiles were investigated using ketone 81b. The reactions of this ketone with O-nucleophiles – sodium alkoxides – in a medium of the corresponding alcohol at 60 °C resulted in alkoxy derivatives 82a and 82b, while reactions with S-nucleophiles – sodium thiolates, generated in situ in anhydrous methanol, led to the corresponding sulfides 83a and 83b. The reaction with piperidine, as an example of an N-nucleophile, was carried out in ethanol with a double excess of piperidine, and the yield of the corresponding 5-piperidinyl derivative 84 was 85% (Scheme 35).

Some of the synthesized isothiazole-containing amides, esters, and ketones were able to exhibit potentiating activity in compositions with the insecticides imidacloprid and/or α-cypermethrin, with larvae of the Colorado potato beetle.84–87 Some derivatives with fungicidal activity were also found.88 In the case of α-cypermethrin, when used in a composition with an insecticide, the maximum synergistic effect was achieved by using a 4,5-dichloroisothiazole derivative with 5–10% of the active component of the insecticidal composition.89

Based on 4,5-dichloroisothiazole-3-carboxylic acid amide (85) and azide 86, both obtained from 4,5-dichloroisothiazole-3-carboxylic acid chloride (77), 3-amino-4,5-dichloroisothiazole (87) was synthesized by the Hoffmann and Curtius reactions.90 It was possible to achieve a 92% amine yield via the preliminary in situ preparation of ethyl carbamate 88, which was then subjected to alkaline hydrolysis (Scheme 36).

4,5-Dichloroisothiazole-3-carboxylic acid nitrile (89) was synthesized by the action of phosphorus pentoxide on 4,5-dichloroisothiazole-3-carboxylic acid amide (85), both without solvent and in tetrachloroethylene. In both cases, the target nitrile was obtained in a quantitative yield.91 Subsequently, reactions of the synthesized nitrile with O-, N- and S-nucleophiles were studied (Scheme 37).

The reaction with O-nucleophilic sodium methoxide led to the formation of a mixture of 5-methoxy-4-chloro-3-cyanoisothiazole (90a) and the product of its alcoholysis 90b, whilst the reaction with piperidine in methanol led to the formation of 5-piperidinyl-substituted isothiazole 91. Reactions with thiolates using benzylmercaptan and thiophenol as thiols proceeded in methanol with satisfactory yields, with the formation of derivatives 92a and 92b, respectively. When n-butylmercaptan was used, a mixture of the thiobutyl derivative 93a and the product of its alcoholysis 93b was formed. In isopropyl alcohol, the yield of isothiazole 93a was 68%.

The possibility of further transformation of the nitrite group was illustrated by synthesizing disubstituted 1,2,4-oxadiazoles 94 after obtaining the 4,5-dichloroisothiazole-3-carboxylic acid amidoxime 95, and its subsequent acylation and cyclization (Scheme 38).92

The yields of acylamidoximes 96 were 85–95% for all derivatives, except for the case when the acylation was carried out with trichlorovinylacetic acid, for which the yield of the corresponding acylamidoxime was 71%. Method B provided slightly higher yields of the target acylamidoximes compared with method A. The acylamidoximes were further cyclized in the presence of acetic acid at reflux in yields of

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**Scheme 36** Pathways to the synthesis of 3-amino-4,5-dichloroisothiazole

**Scheme 37** Synthesis and modifications of 4,5-dichloroisothiazole-3-carboxylic acid nitrile
The possibility of using alkyl (4,5-dichloroisothiazol-3-yl)ketones for the synthesis of functionally substituted isothiazoles through replacement of the chlorine atom at position 5 of the heterocycle and the transformation of alkylcarbonyl fragment was evaluated with (4,5-dichloroisothiazol-3-yl) ketones for the synthesis of functionally substituted isothiazoles through replacement of the chlorine at position 5 with significant tar formation, leading to bromomethyl(4,5-dichloroisothiazol-3-yl) methyl ketone 100 in yields of 32% and 46%, respectively. Substitution of two hydrogen atoms in the methyl group for bromine atoms was not observed. The reaction in carbon tetrachloride was complete in 2 hours, whilst the bromination in acetic acid was complete in 1 hour. It was noted that replacement of the solvent with chloroform or dichloromethane did not lead to positive results; in both cases, the bromination did not proceed. The reaction of bromoketone 100 with thiourea in an ethanol solution at 60 °C proceeded smoothly over 3 hours and resulted in 2-amino-4-(4,5-dichloroisothiazol-3-yl)thiazole (101) in a yield of 82%.

First, reactions with amines were investigated. It had been previously shown that the reaction of (4,5-dichloroisothiazol-3-yl) phenyl ketone with amines proceeded with substitution of the chlorine atom at position 5 of the heterocycle by the amine residue. In aprotic bipolar solvents (N-methylpyrrolidone and DMSO) at 130–140 °C, the process was complete in 6 hours.94 Compared to (4,5-dichloroisothiazol-3-yl)(4-methylphenyl) ketone, the reactions of ketone 97b with amines (morpholine and piperidine) were more active. In a DMSO solution at 70 °C, nucleophilic substitution of the chlorine atom at position 5 of the heterocycle was complete within 5 hours, with good yields being obtained. The ketones 97b and 98a were further reduced to secondary alcohols 99a and 99b by the action of sodium borohydride in isopropanol (Scheme 40).

In addition, the reactions of (4,5-dichloroisothiazol-3-yl) methyl ketone (97b) with elemental bromine in carbon tetrachloride and anhydrous acetic acid were studied (Scheme 41). Bromination in these two solvents proceeded with significant tar formation, leading to bromomethyl(4,5-dichloroisothiazol-3-yl) ketone 100 in yields of 32% and 46%, respectively. Substitution of two hydrogen atoms in the methyl group for bromine atoms was not observed. The reaction in carbon tetrachloride was complete in 2 hours, whilst the bromination in acetic acid was complete in 1 hour. It was noted that replacement of the solvent with chloroform or dichloromethane did not lead to positive results; in both cases, the bromination did not proceed. The reaction of bromoketone 100 with thiourea in an ethanol solution at 60 °C proceeded smoothly over 3 hours and resulted in 2-amino-4-(4,5-dichloroisothiazol-3-yl)thiazole (101) in a yield of 82%.

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In addition, the reactions of (4,5-dichloroisothiazol-3-yl) methyl ketone (97b) with elemental bromine in carbon tetrachloride and anhydrous acetic acid were studied (Scheme 41). Bromination in these two solvents proceeded with significant tar formation, leading to bromomethyl(4,5-dichloroisothiazol-3-yl) ketone 100 in yields of 32% and 46%, respectively. Substitution of two hydrogen atoms in the methyl group for bromine atoms was not observed. The reaction in carbon tetrachloride was complete in 2 hours, whilst the bromination in acetic acid was complete in 1 hour. It was noted that replacement of the solvent with chloroform or dichloromethane did not lead to positive results; in both cases, the bromination did not proceed. The reaction of bromoketone 100 with thiourea in an ethanol solution at 60 °C proceeded smoothly over 3 hours and resulted in 2-amino-4-(4,5-dichloroisothiazol-3-yl)thiazole (101) in a yield of 82%.
The reaction of chalcone 105 with thiourea allowed 3,4-dihydro- and 5,6-dihydropyrimidine-2(1H)thiones 108a and 108b to be obtained in an overall yield of 84%, and the reaction with guanidine hydrochloride afforded 2-amino-pyrimidine derivative 109 (Scheme 43).

Reaction of aldehyde 106 with 1,1′-diacetylferrocene (110) allowed 3-(4,5-dichloroisothiazol-3-yl)[5]ferrocenophane-1,5-dione 111 to be obtained (Scheme 44).

The conversion of various isothiazoles 112 into pyrazoles on treatment with hydrazine was investigated. The influence of various C-3, C-4 and C-5 isothiazole substituents and some limitations of this ring transformation were examined in detail. It was found, that when the isothiazole C-3 substituent was a good nucleofuge, 3-aminopyrazoles 113 were obtained. However, when the 3-substituent was not a leaving group it was retained in the pyrazole product 114 (Scheme 45).

4 Selected Synthesis of Biologically Active Isothiazole Derivatives

As already mentioned, compounds containing an isothiazole fragment have a broad spectrum of biological activity, including pesticidal, anticancer, antimicrobial, and antiviral. The mechanism of the biological action of these compounds has only been studied for some of their representatives.

For example, compound 115 (Figure 2) is a vascular endothelium growth factor receptor inhibitor (VEGFR-2) of tyrosine kinase, and it has been proposed as an anticancer therapy drug that prevents the growth and metastasis of tumors. Although being an inhibitor of a protein significant for the treatment of cancer diseases, a reinforcing effect was not observed during clinical trials in combination with known antitumor drugs. Isothiazole-modified nucleoside 116 (Figure 2) is effective against the first type of herpes virus (HSV-1), and the interaction mechanism involves its binding to thymidine kinase HSV-1.

Recently, it has been shown that the isothiazole nuclei of a number of derivatives can interact with target biomolecules, not only through van der Waals interactions and hydrogen bonds, but also by covalent binding at the unsubstituted position 4 of the heterocycle as a result of bioactivation. This is an important factor that must be considered when designing and synthesizing new biologically active compounds containing an isothiazole heterocycle.
The available literature data on the synthesis of biologically active isothiazoles includes not only information about the relationships between the structures of compounds and their biological activities, but also important information on the approaches used in multistep targeted syntheses of isothiazole-containing molecules. Considering the target functionalization of isothiazoles, it seems appropriate to pay attention not only to individual methods of isothiazole heterocycle formation and ways of modification, but also to synthetic strategies and approaches used in multistage syntheses of target isothiazoles. As examples of such strategies and approaches, the synthetic schemes toward some biologically active isothiazoles appear particularly useful and informative. At the same time, attention will be paid to the stages that include the formation or involvement of the isothiazole heterocycle in chemical transformations.

We can identify three rather conventional strategies for the targeted production of biologically active isothiazoles:

1. Introduction of the isothiazole fragment at the final stage.
2. Convergent synthesis of functionalized isothiazoles.
3. Multistage synthesis based on the isothiazole nucleus.

As an example of the first strategy, we can cite the final stages of the synthesis of nucleoside 116 (Scheme 46). 3',5'-Di-O-acetyl-2-deoxy-5-iodouridine 117 was introduced into the Sonogashira reaction with 3,3-diethoxyprop-1-yne (118) and subsequent hydrolysis of the obtained alkyne derivative 119 gave aldehyde 120. This in turn was reacted with sodium thiosulfate and the resulting thiosulfate derivative 121 was treated with liquid ammonia without purification to give nucleoside 116.

![Scheme 46 Synthesis of a nucleoside bearing an isothiazole ring](image)

In this example, the formation of the isothiazole ring is a factor limiting the yield of the target nucleoside 116. However, low yields of the target products can be justified during initial research and bio-testing, and if necessary (for example, when the desired biological activity is detected), further optimization of the procedure can be carried out.

Another example of this strategy is the formation of substituted isothiazoles via the transformation of other functionalized heterocycles such as isoxazoles. Thus, in the synthesis of cyclooxygenase inhibitor 1 (COX-1) analogues, isothiazole derivative 122 was obtained starting from isoxazole inhibitor 123 through reductive opening of the isoxazole ring with hydrogen in the presence of Raney nickel, followed by cyclization of the resulting enamine ketone 124 by the action of phosphorus pentasulfide in the presence of chloranil (Scheme 47). 103

![Scheme 47 Synthesis of cyclooxygenase inhibitor 1 (COX-1) analogues with the fragment of isothiazole](image)

This method has several disadvantages; however, the yield of the corresponding enamine ketones (38–90%) strongly depends on the substituents present on the isoxazole ring, and their cyclization into isothiazoles usually proceeds with rather low yields (21–51%). 104 Regardless, this pathway can be useful in the synthesis of promising biologically active compound libraries for biological screening, subject to the availability of the corresponding substitute isoxazoles.

The strategy associated with the convergent synthesis of isothiazoles, including demonstrating the potential of isothiazole halogen derivatives in preparative organic synthesis, was used in the final stage of the preparation of a selective inhibitor of the metabotropic glutamate receptor 5 (mGluR5) 125 (Scheme 48). 105,106 3,4-Dibromoisothiazole amide 126 was introduced into a Suzuki cross-coupling reaction with boric acid pinacol ester 127. The product 128 thus obtained was then subjected to a Stille reaction with the pyridine organotin derivative 129 to deliver the target product 125.

Perhaps the most common strategy is sequential multistep synthesis based on the isothiazole core with reactive substituents. This method is used in the synthesis of a number of substituted 2-(isothiazol-3-yl)-1,3,4-oxadiazoles 130 possessing significant antimitogenic activity, achieved through the depolymerization of microtubules (Scheme 49). 107

Hydrazide 131, obtained from 4-aminoisothiazole-3-carboxylic acid amide (132), was further reacted with aryl isothiocyanates. The substituted acylureas thus synthesized were transformed by the action of N,N'-dicyclohexylcarbodiimide (DCC) into 2,4-oxadiazoles 133, which were subjected to reductive amino group alkylation.
An alternative method for the synthesis of the desired isothiazol-3-yl-oxadiazoles consisted of the initial reductive alkylation of compound 132, transformation of the amide groups of the resulting compounds 134 into hydrazide groups, subsequent formation of semicarbazides, and then formation of the 1,3,4-oxadiazole fragments.

A series of vanillin-containing isothiazole derivatives was synthesized by successive transformations of the substituents on the isothiazole nucleus. It has previously been shown that 4,5-dichloroisothiazole-3-carboxylic acid vanillin ester exhibits potentiating activity in a composition with the pyrethroid insecticide cypermethrin, and its azomethine and amine derivatives possess potentiating activity when combined with the neonicotinoid insecticide imidacloprid. An optimized method was used for producing 5-butyl[benzyl, phenyl]thio-substituted 4-chloroisothiazole-3-carboxylic acids 75a–c. The methyl ester of 4,5-dichloroisothiazole-3-carboxylic acid 135 was treated in situ with the corresponding sodium thiocyanates in anhydrous methanol, followed by hydrolysis of the intermediate methyl-5-alkyl[benzyl, phenyl]thio-4-chloroisothiazole-3-carboxylates 136. The synthesis of vanillin esters 137a–c was carried out by the acylation of vanillin with 5-butyl[benzyl, phenyl]thio-4-chloroisothiazole-3-carbonyl chlorides 138a–c, which were obtained by boiling the acids 75a–c with thionyl chloride in the presence of a catalytic amount of DMF.

![Scheme 48](image-url) A selective inhibitor of the metabotropic glutamate receptor 5 (mGluR5) with the isothiazole central nucleus

![Scheme 49](image-url) Synthesis of isothiazol-3-yl-oxadiazoles possessing antimitogenic activity

![Scheme 50](image-url) Synthesis of isothiazolic esters of vanillin and their azomethines
Azomethines 139a–c were obtained by condensation of esters 137a–c with p-toluidine in dry methanol, with yields of 78–92%. Compounds 139a–c were further reduced to the corresponding amines 140a–c with high yields by the action of sodium borohydride and acetic acid in benzene.

Vanillin ether 141, containing a 4,5-dichloroisothiazole moiety, and also azomethine 142 based on it, were prepared in order to study the effect of the nature of the linker between the isothiazole and vanillin fragments on the biological activity of vanillin derivatives of isothiazole (Scheme 51).110 (4,5-Dichloroisothiazol-3-yl) carbinol 143 was reacted with thionyl chloride in carbon tetrachloride to form (4,5-dichloroisothiazol-3-yl) chloromethane 144 in a satisfactory yield. Alkylation of vanillin with chloromethane 144 in dimethylformamide using sodium hydride as the base led to the formation of vanillin ether 141 in a 71% yield. The resulting ether was introduced into a condensation reaction with p-toluidine in methanol in the presence of a catalytic amount of acetic acid to form azomethine 142 (Scheme 51). Isothiazolic derivatives of comenic acid were prepared in a similar manner.111

Another example of such a strategy is the synthesis of the Aurora kinase inhibitor, serine/threonine kinase, which plays a crucial role in cell proliferation (Scheme 52).112 Methyl ether 145 was reacted with intermediate 146 in the presence of sodium hydride, and the carboxyl group in the obtained product 147 was reduced by lithium triethylborohydride in tetrahydrofuran. The resulting alcohol 148 was then mesitylated to form ester 149. Transformation of the ester group into a tertiary amino group by reaction with 2-(ethylamino)-2-methylpropan-1-ol led to the formation of compound 150. After deprotection of the pyrazole nitrogen by the action of hydrochloric acid in dioxane, the desired inhibitor 151 was obtained.

Finally, using this strategy, a collection of 4-chloroisothiazole-containing carbamides113,114 and amides,115 as vascular endothelium growth factor receptor inhibitor (VEGFR-2) 115 analogues (see Figure 2), was synthesized for bio-testing in compositions with known antitumor drugs. Some of the obtained compounds were able to potentiate activity of the known anticancer drugs cisplatin, cytarabine, and etoposide in vitro.113

The approach chosen for the synthesis of isothiazole-containing carbamides included the initial preparation of the corresponding (isothiazol-3-yl)carbonyl azides 86 and 152. To obtain 5-benzylthio-4-chloroisothiazole-3-carbonyl azide (152), two approaches were tested: using acid chloride 138b and 5-benzylthio-4-chloroisothiazole-3-carbox-
Scheme 53 Approaches to the synthesis of 4-chloroisothiazole-containing azides

Acrylic acid hydrazide (153). Treatment of hydrazide 153 with nitrous acid (a mixture of NaNO₂ and HCl) resulted in the target 5-benzylthio-4-chloroisothiazole-3-carboxylic acid azide (152). However, due to the low solubility of hydrazide 153 under the reaction conditions, the process proceeded slowly and the yield of azide 153 did not exceed 65%, whereas the reaction of 5-benzylthio-4-chloroisothiazole-carbonyl chloride (138b) and NaN₃ gave the target azide 152 in a yield of 82% (Scheme 53).

The hydrazide 153 was synthesized in a 95% yield by the action of hydrazine hydrate on ester 136b in methanol. Subsequently, azides 86 and 152 were used to synthesize carbamates, as immediate precursors of ureas (Scheme 54).

The synthesized isothiazolylcarbamates 154a and 154b were further reacted with aliphatic amines, including functionalized examples (n-hexylamine, 6-aminohexanol, 2-aminoethoxyethanol, N',N'-dimethylpropan-1,3-diamine). In addition to the p-fluorophenol carbamate, ethyl carbamate 155 was obtained from azide 152; it had greater solubility but did not react with amines under the utilized conditions, even with prolonged boiling of the reaction mixture. In the case of γ-aminobutyric and ε-aminocaproic acids, which are poorly soluble in chloroform, the reactions with urethanes 154a and 154b were carried out in aqueous ethanol. The obtained carbamides 156 were then transformed into the corresponding salt forms 157 by treatment with sodium carbonate (Scheme 55).

The amide synthesis involved the acylation of selected amino acids with 4,5-dichloroisothiazole-3-carboxylic acid azide 86 in methanol, followed by conversion of the synthesized amides 158 into their salt forms 159 by reaction with sodium bicarbonate (Scheme 56).

Among biologically active isothiazoles, benz[d]isothiazoles occupy an important place. The benz[d]isothiazole moiety has been successfully used in the design and synthesis of neuroleptic ziprasidone as well as in the synthesis of a number of ureas and thioureas, which showed both in vitro antiglycosylation activity and the ability to inhibit urease. Some ureas and thioureas containing benzo[d]isothiazole fragments show high inhibitory activity against Mycobacterium tuberculosis GyrB DNA gyrase. In this regard, it is necessary to mention the recently proposed original approach to the synthesis of substituted benz[d]isothiazoles based on the Diels–Alder reaction, which has been successfully demonstrated with the example of 3-phenyl-4,5-bis(carbomethoxy)isothiazole (160) as the starting compound (Scheme 57).

The carboxyl groups of isothiazole 160 were transformed into carbinols by the action of borohydride in an ethanol/tetrahydrofuran medium, forming the diol 161. The carbinal groups of diol 161 were further converted into bromomethyls by reaction with phosphorus tribromide and pyridine in dichloromethane. The reaction of the
The formation of by-products 169 was observed during the formylation stage of the reaction, and so the imines 168 together with impurities were introduced into the oxidation reaction with iodine, and the impurities were removed during purification of the target products 166.

An alternative strategy was proposed for the synthesis of brassilexin (166a) through the formylation of indolin-2-one (170), followed by heterocyclization of the resulting 2-bromo-1H-indole-3-carbaldehyde (171) in a reaction with urea and sodium thiocyanate under microwave irradiation (Scheme 59). This strategy has also been successfully applied in the synthesis of isothiazole-containing steroids.

Isothiazoles condensed with pyrazole rings were found to be promising biologically active compounds with fungicidal properties. For their synthesis, a strategy has been reported consisting of the oxidation of 1-aryl-5-aminoo-3-methylpyrazole-4-carboxylic acid thioamides 172 with hydrogen peroxide in pyridine, leading to good yields of the desired 3-aminopyrazolo[3,4-c]isothiazoles 173 (Scheme 60). In this case, nitrile 174 was formed in trace amounts. At the same time, the use of iodine as an oxidizing agent in a tetrahydrofuran medium resulted in the formation of either only product 174 or a mixture of products with a predominance of the 4-cyano derivative 174 in a ratio of 95:5. In the case of a 2,5-dichlorophenyl derivative, the corresponding nitrile was the main oxidation product. Some of these synthesized 5-aminopyrazolo[3,4-c]isothiazoles have been shown to be promising as safe antidermatophytic drug candidates.
The interaction of 5-aminopyrazoles 175 with Appel's salt (176) represents another approach to the synthesis of condensed heterocyclic systems (Scheme 61), and pyrazole derivatives of 1,2,3-dithiazole 177 and substituted pyrazolo[3,4-c]isothiazole-3-carbonitriles 178 have been observed,124 rather than pyrazolo[3,4-d]thiazole-3-carbonitriles as previously believed.125

\[
\begin{align*}
\text{Scheme 61} & \quad \text{Investigation of the reaction between 5-aminopyrazoles and Appel's salt}
\end{align*}
\]

The yields of compounds 177 and 178 depend on the temperature, reaction time, acidity of the reaction mixture, and substituents on the pyrazole ring. When substituted 1H-pyrazoles were introduced into the reaction, the main products were pyrazole derivatives of 1,2,3-dithiazole, regardless of the reaction conditions, and pyrazolo[3,4-d]isothiazole-3-carbonitriles did not form. This is particularly interesting because from the corresponding pyrazole derivatives of 1,2,3-dithiazole, it is possible to synthesize thiazole-3-carbonitriles did not form. This is particularly interesting because from the corresponding pyrazole derivatives of 1,2,3-dithiazole, it is possible to synthesize thiazole-3-carbonitriles regardless of the reaction conditions, and pyrazolo[3,4-d]isothiazole-3-carbonitriles as previously believed.125

5 Isothiazoles in the Synthesis of Transition-Metal Complexes and in Metal-Complex Catalysis

The properties manifested by metal complexes are largely determined by the ligand environment of the metal, which ensures the creation of a specific distribution of the electron density on the metal atom and the formation of a spatial framework around it. Information about the metal complexes of the isothiazole series is sparse and fragmentary, and techniques for their practical application have been studied only as isolated examples.

The first information regarding such metal complexes, published in two articles in 1971, was devoted to the synthesis and study of the structure of Co(II), Ni(II), Cu(II) and Pt(II) complexes with unsubstituted isothiazole as a ligand.126,127

For cobalt, complexes CoX₂L₂, CoX₂L₄ and [CoL₄](ClO₄)₂ were obtained, where L is an unsubstituted isothiazole and X = Cl, Br, I.126 The characterization of their structures was accomplished by using elemental analysis, diffuse reflectance spectroscopy, magnetic susceptibility, conductivity in solution, and IR spectroscopy. CoCl₂L₂ and CoBr₂L₂ complexes are polymer structures, CoCl₂L₄ and CoBr₂L₄ are octahedral, and CoCl₂L₂ and CoBr₂L₂ are tetrahedral, but in solution all complexes are characterized by a tetrahedral structure. For the Co₄L₂ complex, instability with regard to the action of atmospheric oxygen and decomposition in nitromethane solution with the formation of molecular iodine and the CoI₂L₂ complex were observed. The [CoL₄](ClO₄)₂ complex is assigned an octahedral structure, both in solid form and in nitromethane solution, with the perchlorate anion in the outer sphere of the complex. Based on the electronic and IR spectra, it was suggested that isothiazole is coordinated through a nitrogen atom.126

In the other paper published in 1971,127 the trans-Co(NCS)₂L₂ complex was obtained, to which the octahedral structure was also attributed as a result of magnetic susceptibility data. The hypothesis of the trans-form of the complex is suggested based on the reaction with α,α′-dipyridyl, leading to the formation of a Co(NCS)₂(bipy)₂ complex as a result of ligand exchange. The same paper describes the synthesis of complexes CuCl₂L₂, CuBr₂L₂, Cu(NO₃)₂L₂, [LH⁺]Cl₄CuOCH₂CH₃(NO₃)₂L₂ (unstable), Cu(NO₃)₂L₄, NiCl₂L₄, NiBr₂L₂, Ni(ClO₄)₂L₄, and cis-PtCl₂L₂. The synthesized complexes were characterized using IR spectroscopy, elemental analysis, magnetic susceptibility, and powder X-ray diffraction. For complexes of the MH₂L₂ type [M = Cu(II), Co(II); Hal = Cl, Br] and for NiBr₂L₂, a polymer octahedral structure with bridging halogen atoms and isothiazole in the trans position was proposed, while complexes NiCl₂L₄, Cu(NO₃)₂L₄, Cu(NO₃)₂L₂, and Ni(ClO₄)₂L₄ were characterized as octahedral. For cis-PtCl₂L₂, a cis-planar configuration was proposed with coordination through the nitrogen atom of the cycle. Later, various cobalt and nickel complexes with 3-, 4- and 5-methylisothiazoles were studied in detail.128,129 It was also shown that 3-methylisothiazole was only able to form tetrahedral complexes of the MeX₂L₂ type, which was associated with steric hindrance created by the methyl group at position 3 of the heterocycle. The isothiazole nuclei in all the synthesized complexes are coordinated via a nitrogen atom.

The same researchers obtained Pd(II), Pt(II), Rh(III), and Ir(III) complexes with isothiazole and 3-, 5-methylisothiazoles: PdX₂L₂ (X = Cl, Br; L = isothiazole, 3-methylisothiazole, 5-methylisothiazole), PtCl₂L₂ (L = isothiazole, 3-methylisothiazole, 5-methylisothiazole), RhCl₂L₂ (L = isothiazole, 3-methylisothiazole, 5-methylisothiazole), RhCl₄L₄ (L = 5-methylisothiazole), IrCl₄L₄ (L = 3-methylisothiazole, 5-methylisothiazole), and IrCl₄L₄ (L = isothiazole).130 The unsubstituted isothiazoles in the complexes are coordinated via a sulfur atom, while 3-methylisothiazole is coordinated via a nitrogen atom. 5-Methylisothiazole was coordinated through a nitrogen atom in the Pd(II) and Rh(III) complexes, and in the Pt(II) and Ir(III) complexes through a sulfur atom.

The literature describes only one example of an isothiazole-containing complex with PdCl₂, the structure of which was accurately determined by X-ray crystal structure.
The stability of the complexes varies in the series of 1,2-azoles as follows: pyrazole > isothiazole > isoxazole. It was suggested that the higher stability of the isothiazole complexes in comparison with their isoxazole analogues could be explained by the possibility of additional π-bond formation between the isothiazole and the metal cation due to the presence of an unoccupied d-orbital on the sulfur atom in the heterocycle.

The literature describes bimetallic tetraselenocyanate complexes of cobalt(II), nickel(II), cadmium(II), and mercury(II), in which isothiazole, 4-methylisothiazole, and 4-benzylisothiazole act as ligands.132,133 Assumptions about the structures of these complexes were made based on elemental analysis, UV-VIS and IR spectroscopy, magnetic susceptibility, and the HSAB theory, wherein a single monomer complex $L_2Ni(NCSe)_{n}Hg(NCSe)_{m}$ ($L$= methylisothiazole, with selenocyanate acting as a bridging ligand) was separated, in which nickel was in an octahedral environment, probably due to the axial coordination of the selenocyanate fragments of the upper layers. All other bimetallic complexes of different composition are described as polymeric tetrahedral, where the selenocyanate acts as a bridging ligand, the N-terminus is coordinated on cobalt and nickel, and the S-terminus is coordinated on cadmium and mercury. Coordination of isothiazoles occurs in these cases via a nitrogen atom at the axial positions in the case of Ni(II) and Co(II), or through sulfur in the case of Cd(II) and Hg(II).

Data on isothiazole-containing carbonyl complexes of tungsten $L_xW(CO)_{6-x}$, molybdenum $LMo(CO)_5$, and chromium $L_xCr(CO)_{6-x}$ ($L$ is the isothiazole ligand; $x$ = 1, 2), obtained by ligand exchange from the hexacarbonyl of the corresponding metal and isothiazole, 4-methylisothiazole or 5-methylisothiazole under ultraviolet irradiation, have been published.135,136 At the same time, depending on the irradiation time of the reaction mixture, substitution of one to two carbonyl fragments in the starting metal hexacarbonyl has been observed. Based on IR spectra, it has been established that the synthesized complexes have a $C_4$ symmetry group, and based on mass spectrometric data for the fragmentation of $L_xW(CO)_{6-x}$ and $L_xCr(CO)_{6-x}$ complexes and the starting isothiazole ligands, it has been found that in all cases, the coordination was achieved through the nitrogen atom.136 This was also consistent with the $^1$H NMR data for the synthesized complexes. An attempt was made to synthesize the isothiazole-π-complex based on the triacetonitrile tricarbonyl complex of chromium [(CH$_3$CN)$_3$Cr(CO)$_3$], but an unstable product was separated with an $L_3Cr(CO)_2$ ($L$ = isothiazole) structure, according to IR spectroscopy data.136

Finally, some examples of isothiazole complexes, in which the metal is coordinated not only through ring heteroatoms, but also through heteroatoms of exocyclic groups, are known. Thus, diperchlorate isothiazole complex 179 was obtained by treatment of Δ-bis(ethylenedi-amine)threininatocobalt(III) trifluoromethansulfonate 180 with thionyl chloride in dimethylformamide, followed by ion exchange chromatography. In this case, the isothiazole-3-carboxylic acid acts as a bidentate ligand, chelating cobalt through the heterocyclic nitrogen atom and carboxyl group (Scheme 62).137

The structure of the complex was confirmed by X-ray crystal structure analysis. Threonine serves as the starting material for the formation of the isothiazole ring. In this case, thionyl chloride acts as both a dehydrating agent and a sulfur atom donor for the isothiazole heterocycle. It is assumed that in the first stage of this reaction, cobalt-chelating threonine chloride is formed, with a sufficiently acidic proton on the α-carbon atom, which makes it possible to proceed with subsequent transformations.

In order to obtain the isothiazole complexes, Meyer et al.138 synthesized isothiazoles containing exocyclic chelating fragments. Thus, pyridine-containing azomethines 181a and 181b were prepared from 5-formyl-3-methylisothiazole (182) (Scheme 63).
In addition to this, sulfide 183 was synthesized via the lithiation of 3,5-dimethylisothiazole (184 Scheme 64). Also, 3-bromomethyl-5-methylisothiazole (185) was obtained starting from 3,5-dimethylisothiazole (184) through bromination of the methyl group at position 3. The introduction of isothiazole 185 into the reaction with N,N,N',N'-tetraethylthiolenetriamine led to the formation of amine 186.

Based on the synthesized isothiazoles, binuclear isothiazole-containing silver triflate complexes \([\text{AgOSO}_2\text{CF}_3]_2 \text{ (L = 181a, 181b, 183}) \) and \([\text{Ag}]_2[\text{O}_3\text{SCF}_3]_2 \text{ (L = 186}) \) were obtained. The structures of the complexes were established by X-ray crystal structure analysis in all cases, except for the complex with ligand 181a. In the case of L = 183, 186, ligands were coordinated to the silver atom by the nitrogen atom of the heterocycle and exocyclic heteroatoms (nitrogen or sulfur). In the case of L = 181b, the silver coordination occurs exclusively on the exocyclic heteroatoms (through the nitrogen of the azomethine and pyridine fragments).

Metal complexes with Cu(II) were obtained based on 4,5-dichloroisothiazole-3-carboxylic acid, its amide, and benzo triazolamide.\(^{27,139,140}\) Polymer complexes of copper \([\text{LCuCl}_2]_n \), \([\text{CuBr}_2]_n \), \([\text{Cu(L)}(\text{BtaH})\text{H}_2\text{O})_n\text{NO}_3]_n \), and \([\text{L}(\text{CuH}_2\text{O})\text{Cl}0.5\text{H}_2\text{O}]_n \) were synthesized. The complexes were characterized by X-ray crystal structure analysis. In these complexes, isothiazole ligands chelated a copper atom with a nitrogen atom of the heterocycle and with an oxygen atom of an amide or carboxyl group. The complex \([\text{Cu(L)}_2\text{Cl}]_n \) possesses fungicidal properties against the phytopathogenic fungi Botrytis cinerea and Fusarium sp., completely inhibiting the development of the fungi at a concentration of 0.125%, according to biological tests. The Cu(L2)Br2 complex, being added at a 5% amount relative to the neonicotinoid insecticide imidacloprid, increases by almost 2 times the toxicity of the insecticide toward Colorado potato beetle larvae, which increases the effectiveness of the insecticide and reduces its consumption.\(^{139}\) Complexes of 4,5-dichloroisothiazole-3-carboxylic acid ethanolamide with copper bromide and copper chloride were also obtained, and according to XRD data, 4,5-dichloroisothiazol-N-(2-hydroxyethyl)-3-carboxamide was coordinated to copper(II) in a tridentate manner. In the bidentate cyclic-type complex, it is coordinated on the nitrogen atom of the heterocycle and on the exocyclic oxygen atom of the amide group with the formation of the five-membered metalloclcylic CuNCO, and also on the oxygen atom of the substituent hydroxy group, and all these bonds form polymer chains. The complex of 4,5-dichloroisothiazole-N-(2-hydroxyethyl)-3-carboxamide with copper(II) bromide (Figure 4) can enhance the action of the insecticides cypermethrin and imidacloprid, and at the same time, copper bromide and the ligand separately did not show any similar activity with insecticides.\(^{141}\)

In 2011, the Pd(II) complex of composition L2Pd (L = 4,5-dichloroisothiazole-carboxylate) was synthesized from 4,5-dichloroisothiazole-3-carboxylic acid.\(^{26}\) It could not be obtained in crystalline form, and therefore, to describe its structure, quantum chemical modeling using the B3LYP1/MIDI(3d) level of theory was used. According to the quantum chemical calculations, a flat-square structure was attributed to the complex; however, it was not possible to establish exactly whether the isothiazole fragments in the complex were in cis- or trans-form due to the proximity of their calculated formation enthalpies. The complex showed catalytic activity in the Suzuki reaction: on addition of 0.1
mol% of the complex, the reaction of 2,4-difluorophenylboronic acid with 5-bromosalicylic acid, carried out in water exposed to air at 100 °C, was complete in 15 minutes with a quantitative yield of 5-(2,4-difluorophenyl)-2-hydroxybenzoic acid being obtained. The latter is active pharmaceutical substance diflunisal, a nonsteroidal anti-inflammatory drug (NSAID) with analgesic and antipyretic effects.

The possibility of using isothiazole complexes with palladium(II) as a catalyst for cross-coupling reactions is of particular interest because of its relevance both in practical and scientific terms. Cross-coupling reactions catalyzed by palladium complexes are widely used in modern organic synthesis as a reliable method of forming carbon–carbon bonds in order to produce polyfunctional biaryls, arylated olefins, acetylenes, and their heterocyclic analogs. Compounds of this type are key structural elements of a number of modern drugs, and are used to develop special purposes.

Synthesis

As in the case of the palladium complex of acid 74, the complexes of hydrazide 187a and phenylhydrazide 187b exhibit high catalytic activity in the Suzuki reaction, albeit slightly lower than that of the 4,5-dichloroisothiazole-3-carboxylic acid complex with palladium (Table 1). At the same time, the catalytic activity of the hydrazide complex was higher than that of the phenylhydrazide complex.

**Table 1 Characteristics of the Cross-Couplings of 3-Bromobenzoic Acid with 4-Methoxyphenylboronic Acid Catalyzed by Complexes of Palladium Chloride with 4,5-Dichloroisothiazole-3-carboxylic Acid (74)**

| Pd complexa | Time (min) | Yield of 190 (%) |
|-------------|------------|-----------------|
| (74)2·PdP  | 15         | 100             |
| 187a·PdCl2c | 30         | 100             |
| 187b·PdCl2d | 60         | 96              |

a Adapted from ref. 149.
b Complex (74)2·Pd: double salt; 187a·PdCl2: hydrazide; 187a·PdCl2: phenylhydrazide.
c Determined from 1H NMR data.

d Diastereomers.

Detailed studies on the influence of the functional environment of the isothiazole heterocycle on the catalytic activity of the corresponding palladium complexes required the creation of new ligands. For this purpose, several new derivatives were synthesized based on 4,5-dichloroisothiazole-3-carboxylic acid, which contained different substituents at position 3 of the heterocycle as well as additional coordination centers of various nature.

For this purpose, heterocyclic molecules containing 1,2,4-triazole or tetrazole rings together with an isothiazole fragment have been synthesized. To construct a triazole heterocycle and to obtain (isothiazole)triazoles, an
approach including initial synthesis of amidrazone 191 via the reaction of nitrile 89 with hydrazine hydrate was chosen (Scheme 67).

![Scheme 67](image)

Scheme 67 An example of the proposed approach towards (isothiazol-3-yl)triazoles

In order to obtain the target amidrazone, it was found that the reaction should be carried out with a minimal amount of methanol or without solvent at all, because otherwise the hydrazine hydrate does not react with the nitrile 89. Amidrazone 191 was further treated with acetyl chloride in the presence of triethylamine, and the resulting acyl derivative 192 was subjected to cyclization to give the target 1,2,4-triazolylisothiazole 193 by heating in glacial acetic acid followed by neutralization of the reaction mixture with KOH solution (Scheme 67). The yield of 1,2,4-triazolyl(isothiazole) 193 was 85%.

The nitrile group of carbonitriles 89 and 92b was used for the formation of tetrazole heterocycles (Scheme 68). The synthesis of tetrazolyl isothiazolones 194a,b was carried out by the reaction of sodium azide and ammonium chloride with carbonitriles 89 and 92b in methanol, with good yields being obtained.

![Scheme 68](image)

Scheme 68 Synthesis of (isothiazol-3-yl)tetrazines

The derivative 193 and its precursor amidrazone 191 were used for the synthesis of azole complexes. For a qualitative assessment of the mutual influence of 1,2-azole and triazole fragments in mixed ligands, palladium complexes with 1,2,4-triazole were also obtained. For the synthesized complexes, extremely low solubility was noted in organic solvents and in water, possibly due to their polymer structure, and therefore they were characterized by IR spectroscopy and elemental analysis. The model reactions presented in Scheme 66 were carried out in 50% aqueous methanol (at 20 °C and 75 °C) or in water (at 100 °C) in the presence of 0.1 mol% of palladium complexes and potassium carbonate as the base. All reactions were carried out in air in the absence of an inert atmosphere. The results of testing the catalytic activity of the complexes are presented in Table 2.

The resulting complexes, except for (191)2PdCl2, (194b)2PdCl2 and (1,2,4-triazole)2PdCl2, exhibited high catalytic activity at elevated temperatures (i.e., 75 °C or 100 °C). The palladium complexes with the isothiazole ligands 191 and 193 were inert at room temperature, while the (191)2PdCl2 complex remained inactive even when heated, as did the (1,2,4-triazole)2PdCl2 complex. The (194b)2PdCl2 complex exhibited only a slight catalytic activity at 100 °C.

Later, the corresponding 3,5-isoxazolyl(isothiazolyl)-substituted 1,2,4-oxadiazoles 195a–c were synthesized by successive transformations of 5-(4-methylphenyl)isoxazole-3-carboxylic acid amidoxime 95 and 4,5-dichloroisothiazole-3-carboxylic acid amidoxime 196 (Scheme 69).151

Additionally, the 2,5-(5-aryl)isoxazolyl(isothiazolyl)-substituted 1,3,4-oxadiazoles 197a–c were obtained by selective recyclization of (4,5-dichloroisothiazol-3-yl)tetrazole 194a (Scheme 70).

Dichloromethane was the only solvent suitable for the preparation of complexes with representatives of the synthesized ligands 195 and 197; the palladium benzonitrile complex (PhCN)2PdCl2 was used as the source of palladium. The complexes were characterized by their elemental
analyses and IR spectroscopic data. In a model Suzuki reaction between 4-methoxyphenylboronic acid (188) and 3-bromobenzoic acid (189) (Scheme 71), it was shown that palladium complexes with these ligands exhibited high catalytic activity in aqueous and aqueous–alcoholic media (Table 3).

Table 2 Characteristics of the Cross-Couplings of 3-Bromobenzoic Acid with 4-Methoxyphenylboronic Acid Catalyzed by Complexes of Palladium Chloride with Ligands 191, 193 and 194b

| Pd complex                  | Temp (°C) | Time (min) | Yield of 190 (%)b |
|-----------------------------|-----------|------------|-------------------|
| 191·PdCl₂                  | 20        | 20         | traces            |
|                            | 75        | 5          | 97                |
| (191)₂·PdCl₂               | 20        | 20         | traces            |
|                            | 100       | 30         | traces            |
| 193·PdCl₂                  | 20        | 20         | traces            |
|                            | 75        | 5          | 100               |
| (193)₂·PdCl₂               | 20        | 20         | traces            |
|                            | 75        | 5          | 98                |
| (194b)₂·PdCl₂              | 20        | 120        | 0                 |
|                            | 100       | 5          | 5                 |
| 1,2,4-triazole·PdCl₂       | 20        | 20         | 10                |
|                            | 75        | 5          | 100               |
| (1,2,4-triazole)₂·PdCl₂    | 20        | 240        | 0                 |
|                            | 100       | 200        | 0                 |

* Adapted from ref. 150 with permission from Springer Nature.

As another promising group of ligands for metal complexes, N-aryl(isothiazol-3-yl)methyleneamines (aromatic azomethines) and amines were synthesized. Conden-

Table 3 Yields of the Suzuki Reaction Product 4′-Methoxybiphenyl-3-carboxylic Acid (190) Depending on the Pd Complex and Reaction Conditionsa,b

| Experiment | Pd complex                  | Temp (°C) | Time (min) | Yield of 190 (%)b |
|------------|-----------------------------|-----------|------------|-------------------|
| 1          | 195c·2PdCl₂                 | 20        | 15         | 30                |
| 2          | 195c·2PdCl₂                 | 75        | 5          | 84                |
| 3          | 195c·2PdCl₂                 | 100       | 1          | 100               |
| 4          | 197a·PdCl₂                  | 20        | 15         | 0                 |
| 5          | 197a·PdCl₂                  | 75        | 15         | 35                |
| 6          | 197a·PdCl₂                  | 100       | 50         | 93                |
| 7          | 197b·2PdCl₂                 | 20        | 20         | 29                |
| 8          | 197b·2PdCl₂                 | 75        | 5          | 79                |
| 9          | 197b·2PdCl₂                 | 100       | 5          | 100               |
| 10         | (PhCN)₂·PdCl₂               | 20        | 15         | 79                |
|            |                             | 4 h        |            | 90                |
| 11         | (PhCN)₂·PdCl₂               | 75        | 15         | 96                |

* Adapted from ref. 151 with permission from Springer Nature.

As another promising group of ligands for metal complexes, N-aryl(isothiazol-3-yl)methyleneamines (aromatic azomethines) and amines were synthesized. Conden-

sation of 4,5-dichloroisothiazol-3-carboxaldehyde (106) with 4-aminobiphenyl and 1-naphthylamine in methanol in the presence of glacial acetic acid resulted in 85% yields of azomethines 198a,b (Scheme 72). In the case of the 4-aminobenzoic acid ethyl ester, this technique was not useful due to only trace amounts of the target azomethine being formed. When carrying out the reaction in boiling benzene in the presence of catalytic amounts of glacial acetic acid, incomplete conversion of the starting aldehyde 106 into azomethine 198c (40%) was observed, therefore the process was carried out with higher-boiling toluene and an increased amount of acetic acid was used. Azomethines 198a–c were further reduced to give amines 199a–c by the action of sodium borohydride and acetic acid in benzene, in 94–95% yields (Scheme 72). Reduction of the isothiazole-containing azomethine 198c proceeded only when the reaction mixture was boiled; at room temperature, there were no signs of the reaction occurring. Amine 199c was further hydrolyzed into isothiazole amino acid 200 (Scheme 73).

As another promising group of ligands for metal complexes, N-aryl(isothiazol-3-yl)methyleneamines (aromatic azomethines) and amines were synthesized. Conden-
Azomethine 198b, amine 199b, and isothiazole-containing amino acid 200 (from the synthesized aromatic azomethines and amines) were used as ligands in order to obtain palladium(II) complexes by reaction with sodium tetrachloropalladate. Complexes of the composition LPdCl₂ were formed by the interaction of equimolar amounts of sodium tetrachloropalladate and the ligands in methanol at 20 °C. Since acid 200 was poorly soluble in both methanol and acetonitrile, which were previously used to produce 1,2-azole palladium complexes, the complex was synthesized in a mixture of methanol and DMF, in which derivative 200 is quite soluble. The obtained palladium complexes 198b·PdCl₂, 199b·PdCl₂, and 200·PdCl₂ were identified based on their elemental analyses as well as IR spectroscopic data, in which the characteristic vibration bands of the C=N and C=C bonds of the isothiazole heterocycle and the corresponding exocyclic fragments were apparent. According to their elemental analyses, they form complexes with palladium chloride with the composition LPdCl₂. The complexes are insoluble in organic solvents and in water, which made it impossible to record their NMR spectra or to grow single crystals for X-ray crystal structure analysis. The assumption of their structure was made by analogy with the previously obtained palladium complexes with oxime ligands of the isoxazole series, as well as by comparing the IR spectra of the ligands and the corresponding complexes. This indicates the participation of the heterocycle and exocyclic fragments (C=N/C–NH) in coordination with palladium. For the complex of the isothiazole amino acid 200 with palladium chloride, calculations of the geometry and IR spectrum were made at the B3LYP/6-31+G*/LANL2TZ(f)ECP(Pd) level of theory. As a result, it was proposed that the isothiazole ligand 200 in the molecules of the complex 200·PdCl₂ is coordinated to palladium in a bidentate cyclic-type manner by the nitrogen atoms of the heterocycle and the exocyclic amino group to form a five-membered metallocycle (Figure 5).

Evaluation of the catalytic activity of the obtained complexes on the model reaction presented in Scheme 66 was performed under conditions similar to those used in previous tests. The results of the testing of the catalytic activity of the complexes are presented in Table 4.

Table 4 Characteristics of the Cross-Couplings of 3-Bromobenzoic Acid with 4-Methoxyphenylboronic Acid Catalyzed by Palladium Chloride Complexes with Ligands 198b, 199b and 200

| Pd complex          | Temp (°C) | Time (min) | Yield of 190 (%) |
|---------------------|-----------|------------|------------------|
| 198b·PdCl₂          | 20        | 30         | 67               |
|                     | 75        | 20         | 96               |
| 199b·PdCl₂          | 20        | 30         | 42               |
|                     | 100       | 30         | 100              |
| 200·PdCl₂ (in situ potassium salt) | 35        | 5          | 100              |
| Na₂PdCl₄            | 100       | 3          | 100              |
| Na₂PdCl₄            | 20        | 240        | 92               |
|                     | 75        | 5          | 99               |

a Adapted from refs. 152 and 153 with permission from Springer Nature.
b Determined from ¹H NMR data.

The reactions were complete in 3–30 minutes with the formation of 4′-methoxy[1,1′-biphenyl]-3-carboxylic acid (190), the target product of the cross-coupling, with yields of 96–100%. In each case, the formation of palladium black did not occur until the end of the reaction. Analysis of the reaction mixtures by TLC at the time of separation of the
palladium black consistently showed the absence of aryl halide. The solution itself remained almost colorless, which indirectly indicates a low content of colloidal (nanoscale) palladium in the reaction mixture and products.

According to the obtained data, the palladium complexes with one ligand \((LPdCl_2)\) were significantly more active than those with two ligands \((L_2PdCl_2)\). The complex with the isothiazole derivative 200 was the most active among all the complexes tested, and its activity was especially noticeable at 20–35 °C. It should also be noted that the formation of a small amount of the arylboronic acid homocoupling product, \(4,4′\)-dimethoxy-1,1′-biphenyl (with yields of 1–3%), was observed in the reactions in all cases. Since the reactions were carried out in the absence of an inert atmosphere, it is possible that the by-product was formed because of oxidation of the starting arylboronic acid by aerial oxygen during the catalysis with palladium, but the contribution of this process is considered negligible. Complex 200 was used for the preparation of a heterogeneous silicon-oxide-supported catalyst, and its catalytic properties were almost as good as the complex itself and it was able to withstand at least 10 catalytic cycles.\(^{154}\)

The observed temperature dependence of the catalytic activity of the described isothiazole complexes can be used to control the selectivity of the cross-coupling reactions in molecules with several reaction centers. Isothiazole palladium complexes are suitable for chemical and chemical-pharmaceutical processes in cases where the synthesis of the target substances is carried out using the Suzuki reaction and high purity requirements are imposed on the final product.

We are confident that the data on isothiazole–metal complexes summarized above, as well as the information on the catalytic and biological activity of their representatives, can stimulate further research in this very promising area. At the same time, we believe that while the synthesis of new metal complexes of the isothiazole series and the study of their properties definitely deserves the attention of modern researchers, so also does the study of the practical utilization of the isothiazole complexes that were previously described in the literature.

The above information on isothiazole complexes, as far as we know, is comprehensive. In an earlier review on the coordination chemistry of thiadiazoles, thiazoles and isothiazoles, isothiazole–metal complexes were given little more than one page, with only five papers on this topic being considered.\(^{155}\)

## 6 Conclusion

The presented and summarized data, published mainly over the past 18 years, demonstrates the intensive development of isothiazole chemistry and highlights the rich potential of isothiazole derivatives both for organic synthesis and practical use.

New data on the design of isothiazole heterocycles and the synthesis of isothiazole derivatives has been considered from the standpoint of retrosynthetic analysis, which allows four rational approaches for the formation of the isothiazole rings to be distinguished: intramolecular cyclization, \((4+1)\)-heterocyclization, \((3+2)\)-heterocyclization and synthesis based on other heterocyclic compounds. Such classification enables a consistent and logical understanding of the available approaches to obtaining substituted isothiazoles, and in some cases, to predict the most rational synthetic methods thereof.

The largest part of the published information is covered in the section dedicated to the target functionalization of substituted isothiazoles. This data demonstrates the rich possibilities of the developed strategies for successive transformations of isothiazole derivatives. Chlorinated isothiazoles seem particularly promising as starting compounds in this regard.

The logical continuation of such functionalizations is a separate section devoted to biologically active isothiazoles. In recent years, a large number of molecules containing the isothiazole fragment have been proposed as bioactive substances capable of selectively conjugating with key enzyme sites. There are various strategies for the formation of such molecules, which have been illustrated in detail. The most common strategy is sequential multistep synthesis based on isothiazole cores with reactive substituents. This method is used for the synthesis of a number of isothiazoles possessing significant antimitogenic activity. Sufficiently high interest is represented by published data on isothiazoles that exhibit synergistic effects in compositions with known anticancer drugs and pesticides. It is hoped that the research in this field will continue to develop rapidly and will unite the efforts of both chemists and specialists in the fields of biochemistry and medicine.

A separate section is devoted to isothiazoles in the synthesis of transition-metal complexes and in metal-complex catalysis. There is a great future for research in this field of isothiazole chemistry, especially in the development of metal-complex catalysts for cross-coupling reactions. The data on the metal complexes of the isothiazole series was previously sparse and fragmentary, and information on their possible practical use had not been summarized. Much remains to be done in this area in terms of improving metal-complex catalysts for effective use in aqueous media (‘green chemistry’) and in finding appropriate methods for...
determining their structures. The biological activity of metal complexes of isothiazoles remains an almost untouched field of research.

We hope that the data presented and analyzed herein will serve as a stimulus for further studies in the field of isothiazole chemistry, and that it will be useful not only to those who work in the field of heterocyclic chemistry and medicinal chemistry, but also to all specialists in modern organic synthesis as an inspiration and a source of ideas for further research.

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