Review
Synthesis of Biologically Relevant 1,2,3- and 1,3,4-Triazoles: From Classical Pathway to Green Chemistry

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Abstract: Green Chemistry has become in the last two decades an increasing part of research interest. Nonconventional «green» sources for chemical reactions include micro-wave, mechanical mixing, visible light and ultrasound. 1,2,3-triazoles have important applications in pharmaceutical chemistry while their 1,2,4 counterparts are developed to a lesser extent. In the review presented here we will focus on synthesis of 1,2,3 and 1,2,4-triazole systems by means of classical and « green chemistry » conditions involving ultrasound chemistry and mechanochemistry. The focus will be on compounds/scaffolds that possess biological/pharmacophoric properties. Finally, we will also present the formal cycloreversion of 1,2,3-triazole compounds under mechanical forces and its potential use in biological systems.

Keywords: 1,2,3-triazoles; 1,2,4-triazoles; ultrasound; medicinal chemistry; green chemistry; mechanochemistry; biological properties

1. Introduction

One of the main goals in the area of organic synthesis oriented towards biologically active compounds is the research and development of efficient environmentally safe methods. In fact, since the 2000s many regulations for the chemical and pharmaceutical industries have appeared, especially in terms of efficiency, waste management and energy input. All these issues are now addressed and termed «Green Chemistry», a multifaceted field dealing with what we call the twelve principles of P.T. Anastas and J.C. Warner [1]. Most important of them are: atom economy, preventing the use of solvents volatile and/or toxic, minimize chemical waste and minimize energy [2]. Organic reactions and processes are classically conducted in solutions (mostly organic) under reflux or thermal energy to be balanced at the end of the transformation. We focus on the Green Chemistry synthetic aspects, and focus on chemical reactions by using alternative energy sources that appeared and developed since the last two decades; namely, the processes: photochemistry through light excitation, microwave, sonochemistry irradiation, and mechanochemistry [3].

In this article, in order to give an emblematic example of the evolution of synthesis strategies towards ever greener processes, in particular in the pharmaceutical field, we will focus not only on recent classical synthesis of 1,2,3 and 1,2,4-triazoles, but also on sonochemistry and mechanochemical synthesis of these systems in relation to their biological activities. First, we will initially focus on these two alternative energy sources, i.e., sonochemistry/ultrasonic irradiation, and mechanochemistry. Mechanical effects caused by sound irradiation—called sonochemistry—can be applied to liquids. It can induce formation and growth of acoustic cavitations resulting in implosive bubble collapse [4–6]. This leads to intense compressional heating and extremely high pressures in
the resultant so called hot spots (5000 K, 1000 atmospheres) while heating and cooling rates are exceeding $10^{10}$ K s$^{-1}$ (Figure 1) [7,8]. The sonic spectrum ranges from high power to low power ultrasound (20 KHz to 10 MHz). The range from 20 KHz to 1 MHz is used in sonochemistry. As indicated before, ultrasound irradiation can induce formation and growth of acoustic cavitations resulting in implosive bubble collapse that due to their physical properties can substantially improve chemical reactions (catalytic or not) in terms of speed (in some reactions a million fold reactivity increase was observed), selectivity and yield. Ultrasound reactions are not adequate for reactions between solids or solid-gas systems [9]. Ultrasound in organic synthesis has been studied considerably in the past two decades. Especially, various named organic transformations effected through ultrasound irradiation were developed. Among the most important [10] we can point to the coupling reactions, i.e., Heck, Suzuki, Sonogashira, Ullmann, ultrasound-assisted phase transfer catalysis, some named reactions like Reformatsky, Michael, Baylis-Hillmann, but also oxidation/reduction reactions, halogenations. Finally, ultrasound synthesis of ionic liquids and heterocyclic, especially nitrogen contained compounds [11], has gained much success and development. Mechanical energy can also induce chemical transformations [10]. According to IUPAC, a mechanochemical reaction is a “Chemical reaction that is induced by the direct absorption of mechanical energy” [12,13]. Wilhelm Ostwald (Nobel Prize in 1909), was the first who mentioned the term “Mechanochemistry” and defined it as a “branch of chemistry which is concerned with chemical and physico-chemical changes of substances of all states of aggregation due to the influence of mechanical energy”. It is important to mention the pioneering work of Boldyrev et al., on the mechanisms and kinetics in comminuting devices [14–16], serving as a basis for many mechanochemical works. How the absorption of mechanical energy induces chemical transformations in terms of mechanistic understanding is still under investigation and not fully elucidated. Various models were proposed based on solid chemistry knowledge, like “hot spot” and “magma-plasma model” [17–19]. Other also well-known models (spherical, kinetic and impulse . . . ) were equally proposed [20,21]. Many efforts were developed recently towards a mechanistic level understanding of mechanochemical processes [22]. One of the major trends in progress is to research possible links between the mechanical effect and the action of the forces generated at the molecular level [23–27]. In parallel with these recent advances, the topic is still subject to research from the experimental and theoretical points of view [28]. In terms of the experimental view, the traditional grinding by using a mortar and a pestle has been replaced by more sophisticated ball-milling or mechano-milling techniques that are generally conducted in vibration mills or planetary mills at frequencies of 5–60 Hz. The reactions are generally carried out in vessels or jars of different kinds of materials (stainless steel, tungsten carbide, zirconia, agate, etc.). In recent years a deviation from the pure solid status of reactants, named the liquid assisted grinding, gained considerable interest because it offers opportunities to mechanochemistry to reach viable results in comparison to solution synthesis [29,30]. These studies are greatly facilitated by a possible continuous monitoring of mechanochemical reactions [31,32]. One drawback for mechanochemistry is the fact that up to now difficulties exist in practically controlling the air and moisture sensitive reagents. However, Kubota et al. have shown recently [33] that mechanochemistry allows carrying out the syntheses of organometallics sensitive to humidity in air.

In the two last decades, this green chemistry approach has been developed considerably in areas related to inorganic compounds and metal complexes synthesis and related mechanistic aspects [34–36] while less interest was focused on organic mechanochemistry, even after the pioneering work reported by Toda in the 1980s [37] and Kaupp [38]. This is actually changing since the last decade’s focus was essentially on the green chemistry and green processes approach [39]. In recent times, mechanochemical synthetic approaches for creating carbon-carbon, carbon-heteroatom, metal-ligand coordination bonds etc. became important issues and gained considerable attention in the literature [40,41], and many applications were carried out in the field of organic mechanochemistry [42–52]. Among
the many heterocyclic ring structures, especially nitrogen-contained, which were found and/or designed as important scaffolds for inducing biological effects are the triazoles. Triazole is a five membered ring with three nitrogen and two carbon atoms. Depending on the disposition of the five atoms, triazoles exist in two isomeric forms, namely 1,2,3- and 1,2,4-triazoles. Triazoles have become increasingly popular between medicinal chemists and pharmaceutical companies due essentially to their unique properties such as: rigidity, strong hydrogen-bond properties, stability under in vivo, and interesting pharmacokinetic profiles. Due to their importance, much literature data exist for 1,2,3-triazole systems in comparison to the 1,2,4-triazoles concerning either their syntheses or their biological activities see for instance [53–55]. In that respect the review presented here focuses on three recent parts, namely:

(a) Construction of 1,2,3-triazole systems in biologically relevant compounds by means of classical and “green chemistry” conditions involving ultrasound chemistry and mechanochemistry.

(b) Construction of 1,2,4-triazole systems in biologically relevant compounds by means of classical and “green chemistry” conditions involving ultrasound chemistry and mechanochemistry.

(c) The mechanochemical cyclo-reversion of 1,2,3-triazole compounds and the scientific discussion on the topic that it could be extremely stimulating as mechanochemistry seems to provide a method by which reactive azide or alkyne intermediates could be selectively unmasked.

Figure 1. A summary of various approaches toward synthesis of 1,2,3-triazoles.

2. 1,2,3-Triazole Systems

One of the most important five-membered heterocyclic scaffolds due to its extensive biological activity is the 1,2,3-triazole one. The framework can be readily obtained through the click chemistry via reaction of an aryl/alkyl halide, alkynes and NaN₃. Many synthetic methodologies were developed the past few decades, usually partitioned between metal-free and metal catalysed approaches, thus offering new opportunities for introduction of this valuable moiety to biologically relevant compounds designed and developed by
medicinal chemists (Figure 1). A very recent review treats on those methodologies and on the medicinal attributes of 1,2,3-triazoles [56].

We report here notable current examples (year 2018) of classical synthesis of biologically active compounds bearing this frame, but also recent literature from 2014 concerning synthesis of 1,2,3-triazoles by chemical transformations using alternative energy sources (ultrasonic irradiation).

Alexandre et al. reported in 2018 [57] that compounds based on 4-amino-1,2,3-triazole core as potent inhibitors of indoleamine 2,3-dioxygenase (IDO1) are important targets of immuno-oncology research. The authors screened on a recombinant human IDO1 a library of 350,000 compounds and were able to identify a series bearing the 4-amino-1,2,3-triazole core. Upon chemistry optimisation they obtained compound N-(4-chlorophenyl)-2H-1,2,3-triazol-4-amine with a remarkable potency (IC$_{50}$ of 0.023 µM) substantially more potent than any other IDO1 inhibitor. Synthesis of this compound differs from all other methods. The synthesis involves diazotization of 4-chloroaniline 1 by sodium nitrite followed by reaction with 2-aminacetoni trile hydrochloride 2 in order to afford 2-(2-(4-chlorophenyl) iminohydr azino) acetonitrile 3 which upon heating under reflux in ethanol afforded the desired compound 4 (Scheme 1).

![Scheme 1. Synthesis of tacrine and quinoline derivatives 4.](image)

Wu et al. [58] reported the design and synthesis of tacrine 1,2,3-triazole derivatives as potent cholinesterase inhibitors. Tacrine 5, the first drug approved by the FDA that binds at the catalytic active site (CAS) region a potent non selective inhibitor of both bAChE and hBChE was hybridized through various types of linkers bearing the 1,2,3-triazole pharmocophore frame with activities against AChE and BChE with IC$_{50}$ values of 4.89 and 3.61 µM, respectively. The authors point out that although this compound is less potent than ta-

![Scheme 2. Synthesis of tacrine and quinoline derivatives 9.](image)

Among all compounds synthetized, compound 9 (R = H, linker = piperazine) exhibited a potent inhibition against AChE and BChE with IC$_{50}$ values of 4.89 and 3.61 µM,
respectively. The authors point out that although this compound is less potent than tacrine, it has a unique binding mode at both CAS and also to the peripheral anionic site (PAS), as well as less toxicity. They concluded by considering it as a lead compound that could be the basis for the development of more active dual inhibitors of AChE and BChE (Table 1).

Table 1. Inhibition of AChE (Electrophorus electricus) and horse serum BChE by Tacrine 5 and a derivative 9 (with piperazine as linker and R = H).

| Compounds                | Inhibition (%) at 100 µM | IC<sub>50</sub> (µM) |
|--------------------------|--------------------------|-----------------------|
| 9 (R = H, linker = piperazine) | 78.69                    | 91.80                 |
| Tacrine 5                | 86.22                    | 99.63                 |
|                          | 0.316                    | 0.066                 |

Ashok et al. [59] reported the synthesis of a novel prototype that possessed a chromene and a 1,2,3-triazole pharmacophore frame with activities against <i>M. tuberculosis</i>. The strategy adopted by the authors for their synthesis started from substituted acetophenone 10 which upon Kabbe condensation and reduction of the carbonyl group afforded spirochromene 11. Deprotection of 11 and dehydration provided the corresponding spirochromene 12. 1,2,3-triazole-fused spirochromene derivatives 13 were then obtained through a Huisgen cycloaddition in the presence of pyrrolidine as catalyst (Scheme 3).

![Scheme 3](image_url)  
*Scheme 3. Synthesis of 1,2,3-triazole-fused spirochromenes 13.*

Among the compounds tested against <i>M. tuberculosis</i> H37Rv strain, 5 compounds presented strong MIC activities (between 4 and 9 µM). Their cytotoxicity against RAW 264.7 cells was determined and indicated at least one log difference in comparison to their MIC values. These findings indicated that 1,2,3-triazole-fused spirochromene derivatives can have biological significance for further development (Table 2).
Table 2. Anti-tubercular and toxicity evaluation of compounds 13 against M. tuberculosis H37Rv.

| Compounds 13 | MIC (µg/mL) | MIC (µM) | Cytotoxicity in % Inhibition at 50 µg/mL |
|--------------|-------------|----------|----------------------------------------|
| Me H C₆H₅-  | 1.56 | 4.74 | 30.23 |
| Me H 4-Ome-C₆H₅- | 1.56 | 4.34 | 33.14 |
| Cl H Benzyl | 1.56 | 4.11 | 29.36 |
| Cl Me C₆H₅- | 3.125 | 8.60 | 24.90 |
| Cl Me 4-Cl-C₆H₅- | 3.125 | 7.87 | 24.76 |

MIC: minimum inhibitory concentration (the lowest concentration that inhibited the bacterial growth).

López-Rojas et al. [60] reported the synthesis of 4-substituted 1,2,3-triazole coumarin-derivatives and evaluated their antimicrobial activity. The strategy adopted by the authors was the synthesis of acetylenic O- 15 or N-propargylated 17 coumarins starting from 4-hydroxy 14 and 4-bromo 16 coumarin respectively. Copper(I) catalyzed Huisgen 1,3-dipolar cycloaddition reaction with synthetized (or commercially available) alkyl or aryl azides and afforded the desired compounds 18 (Scheme 4).

![Scheme 4. Synthesis of 4-substituted 1,2,3-triazole-coumarin derivatives 18.](image)

The authors thus created a focused library of 26 compounds with two isosteric series (hydroxy/ amino) and with different substituents at the triazole moiety. Based on their MIC values against selected microorganisms, 5 out of 26 compounds showed significant antibacterial activity towards Enterococcus faecalis (MIC = 12.5–50 µg/mL) while low cytotoxicity was observed against human erythrocytes (Table 3).

Table 3. Antimicrobial activity of compounds 18 against Enterococcus faecalis (MIC µg/mL) of the synthesized compounds.

| Compounds 18 | Enterococcus faecalis |
|--------------|-----------------------|
| X            | R                     |
| O            | C₆H₅-                 | 50 |
| O            | 2-Ome-C₆H₅-           | 12.5 |
| O            | 4-F-C₆H₅-             | 50 |
| NH           | 3-NO₂-C₆H₅-           | 50 |
| NH           | C₁₁H₂₃                | 50 |

2.1. Ultrasound Assisted Syntheses of 1,2,3-Triazoles

We will refer herein to some relevant publications from 2014 up to now.

In 2014, Mady et al. [61] reported the ultrasound assisted synthesis of diaryl sulfones bearing 1,2,3-triazole moieties as potential antioxidant and antimicrobial agents. Synthesis of disubstituted triazoles is depicted below (Scheme 5). The authors explored three routes: a stepwise approach that allowed a click coupling of two different azides and a second and third one where both alkynes were introduced then allowed to click from the same azide. Ultrasound (US) Barbier type mediated propargylation occurred readily to construct the common intermediate 19. This can undergo CuAAC cycloaddition reaction with different
azides under ultrasound conditions affording a first 1,2,3-triazole containing compound 20. The hydroxy group can be further propargylated and coupled with other azides (or the same), affording final compounds 23.

Scheme 5. Synthesis of mono and bis-1,2,3-triazoles, respectively 20 and 23.

The second route introduces first a second alkyne group via propargylation of the hydroxy group of key sulfone 21 and then CuAAC cycloaddition reaction with the corresponding azide. All syntheses were operated under ultrasound conditions in a very efficient manner. The authors also synthetized bis-triazoles via the one-pot click reaction (third route).

Biological and antioxidant activities of all compounds were also reported. Many of them were found to be most potent antifungal agents with MIC values around 25 µg/mL (Table 4). Moreover, compound 24 (Scheme 6) showed an excellent antioxidant activity (IC₅₀ = 20 µg/mL) using a DPPH free radical scavenging assay.

Table 4. Minimum inhibitory concentration (µg/mL) against A. niger.

| Compounds | R                     | A. niger MIC (µg/mL) |
|-----------|-----------------------|----------------------|
| 20        | C₆H₅                  | 25                   |
| 23        | C₆H₅                  | 25                   |
| 23        | (4-SO₂Ph)C₆H₅PhCO-    | 25                   |
| 23        | (4-F)C₆H₅PhCO-        | 25                   |
| 23        | (4-Br)C₆H₅PhCO-       | 25                   |
Nallapati et al. reported in 2015 [62] synthesis of 1,2,3-triazoles derived from olanzapine. Olanzapine (Zyprexa), a member of the thienobenzodiazepine class, is a confirmed marketed drug used for the treatment of schizophrenia and bipolar disorder. The authors describe modifications of olanzapine and explore their activities. One of the target molecules chosen by the authors being olanzapine decorated with 1,2,3-triazole moieties. In that respect alkyne 26 was first prepared through classical coupling in the presence of NaH of propargyl bromide with the drug olanzapine 25 in THF. The thus prepared alkyne reacted with aryl or alkyl azides at room temperature under ultrasound irradiation and in the presence of disopropylethylamine affording the triazolo derivatives 27 in fairly good yields (Scheme 7).

The authors reported in vitro activities of these compounds against phosphodiesterase 4B protein (PDE4B), a gene family that plays a role in the treatment of schizophrenia. Three of the compounds tested were identified as selective inhibitors of PDE4B (IC50 5 to 6 μM) (Table 5).

Table 5. Inhibition of PDE4B at 10 μM by compounds 27.

| Compounds 27 | Average % Inhibition against PDE4B |
|--------------|----------------------------------|
| R            |                                  |
| -CH3Ph       | 72.82                            |
| -CH2C6H3(Cl-o)(CF3-p) | 75.37                        |
| -C6H4F-p     | 74.83                            |

PDE4B: phosphodiesterase 4B protein.

N. Rezki reported in 2015 [63] synthesis under conventional methods and ultrasound conditions of 1,4 disubstituted 1,2,3-triazoles tethering bioactive benzothiazole nucleus and their antibacterial evaluation. Synthesis (Scheme 8) started from 2-aminobenzothiazole derivatives 28 which were acylated upon reaction with bromoacetyl bromide. Then, azidation in the presence of sodium azide afforded the corresponding azidobenzothiazoles 29. All reactions were performed under classical and ultrasound conditions with better yields in the latter case. Huisgen copper(I) catalysed 1,3-dipolar cycloaddition with appropriate terminal alkynes in the presence of sodium ascorbate in tBuOH/H2O, and was carried out under heat or use of ultrasound at room temperature affording compounds 30. Again, ultrasound conditions revealed to be more favorable.
All compounds were tested against three gram positive and three gram negative bacteria and two fungal strains. Some of them, revealed promising activities in the range of 4–8 μg/mL (Table 6).

Table 6. Antimicrobial activity of compounds 30 expressed as MIC (μg/mL).

| Compounds 30 | Gram-Positive Organisms | Gram-Negative Organisms | Fungi |
|--------------|-------------------------|-------------------------|-------|
| R            | R₁                      | Sp                      | Bs    | Sa    | Pa    | Ec    | Kp    | Af    | Ca    |
| -SO₂Me       | CH(OH)(Ph)              | 8                       | 8     | 4     | 4     | 8     | 8     | 8     | 8     |
| -SO₂Me       | -C₂H₄OH                 | 4                       | 4     | 8     | 4     | 4     | 8     | 4     | 4     |
| -SO₂Me       | -C₃H₆OH                 | 4                       | 4     | 8     | 4     | 4     | 4     | 4     | 4     |

Sp: Streptococcus pneumonia; Bs: Bacillus subtilis; Sa: Staphylococcus aureus; Pa: Pseudomonas aeruginosa; Ec: Escherichia coli; Kp: Klebsiella pneumonia; Af: Aspergillus fumigatus; Ca: Candida albicans.

N. Rezki and M.R. Aouad reported in 2017 [64] synthesis of hybrid compounds bearing fluorinated 1,2,4-triazole, 1H-1,2,3-triazole and also a benzothiazole functionality. Construction of the 1,2,4-triazole substituted frame started from reaction of 2-fluorobenzoyl chloride 31 with hydrazine hydrate.

Subsequent treatment was administered with diverse alkyl/aryl isothiocyanates, which upon basic reaction conditions underwent an oxidative ring closure affording the thione derivatives 32. The latter reacted with propargyl bromide in the presence of triethylamine under ultrasound conditions, furnishing the thiopropargylated 1,2,4-triazole precursors 33 required for the click reaction. On the other hand, acylation of the appropriate 2-aminobenzothiazoles 34 followed by the azidolysis reaction allowed obtention of the azidoacetamide derivative 35. The Huisgen cycloaddition reaction was then performed between the two coupling reagents in the presence of CuSO₄ and Na-ascorbate as catalysts in DMSO-H₂O. The ultrasound conditions were less time consuming and much more efficient with almost quantitative yields (Scheme 9).

Almost all compounds showed activities with MIC values in the range 6.45–33.2 μmol/L against S. pneumoniae. In addition, compound 37 (Figure 2) showed the strongest antifungal activities among all compounds with MIC values of 6.45 μmol/L against A. fumigatus and C. albicans.

2.2. Mechanochemical Syntheses of 1,2,3-Triazoles

Praveen et al. reported in 2017 [65] the synthesis of new hybrid pharmacophores under ball milling conditions through two well established named reactions, namely a Baylis-Hillman [66] and a Huisgen’s click chemistry [67]. The authors aimed to prepare potential medicinal targets bearing a 3-substituted-3-hydroxy-2-oxindole frame present in many natural products and medicinal agents [68–70] and a 1,2,3-triazole scaffold. The authors successfully combined a Baylis-Hillman and a click reaction by using DABCO as a base and copper oxide nanoparticles as catalysts. By milling together a mixture of N-propargyl isatin, N-methylmaleimide, benzyl azide, DABCO in the presence of CuONP catalyst (5%) they were able to find optimal conditions of achieving the synthesis of the target compound 38a in 96% of yield (Scheme 10).
Scheme 9. Synthesis of fluorinated-1,2,4-triazole 36.

Figure 2. 2-(4-(((3-(2-fluorophenyl)-4-phenyl-4,5-dihydro-1H-1,2,4-triazol-5-yl)thio)methyl)-1H-1,2,3-triazol-1-yl)-N-(5-(methylsulfonyl)-2,3-dihydro-1H-inden-2-yl)acetamide 37.

Scheme 10. Mechanochemical synthesis of oxindole-triazole hybrid pharmacophores 38.
They then applied the conditions found for creating a small focused library (Table 7) as their methodology and accommodated a large variety of substituted starting compounds. In addition, the authors proved the recyclability of the catalyst and its total recovery. Biological studies along with molecular docking demonstrated the rational efficiency of the compounds as antibacterial and antifungal. The best activities were found for compound 38m (Figure 3), which was most active against S. aureus (with a MIC value of 16 µg mL⁻¹), for compounds 38a,d,i,l active against E. coli and for compounds 38e,h,k,m,p active against C. albicans.

Table 7. Compounds 38: Structures and mechanochemical yield.

| Compounds 38 | R   | R₁  | R₂    | Yield (%) |
|--------------|-----|-----|-------|-----------|
| 38a          | Me  | H   | benzyl| 96        |
| 38b          | Me  | H   | p-methylbenzyl | 98        |
| 38c          | Me  | H   | p-methoxybenzyl | 99        |
| 38d          | Me  | H   | hexyl | 80        |
| 38e          | Me  | H   | phenyl| 70        |
| 38f          | Me  | H   | ethoxycarbonylmethyl | 90        |
| 38g          | Me  | H   | p-nitrobenzyl | 93        |
| 38h          | Me  | 5-Cl| benzyl | 99        |
| 38i          | Me  | 5-Cl| p-methylbenzyl | 90        |
| 38j          | Me  | 5-Cl| p-nitrobenzyl | 90        |
| 38k          | Me  | 5-Cl| p-methoxybenzyl | 98        |
| 38l          | Me  | 5-Cl| phenyl | 71        |
| 38m          | phenyl | H   | benzyl | 90        |
| 38n          | benzyl | H   | benzyl | 93        |
| 38o          | Me  | 5-Br| benzyl | 96        |
| 38p          | Me  | 5-Me| benzyl | 88        |

Figure 3. Most active compound 38m of the series.

Sahu et al. published in 2019 [71] the synthesis of quinine-triazole systems with the aim to find new compounds via molecular hybridization [72] that can be addressed to two antiprotozoal targets that are malaria and leishmaniosis [72,73]. The synthetic route (Scheme 11) started with activation through mesylation of the secondary alcohol of quinine generating the compound 39 and subsequent substitution with the azide group via a solution-based methodology [74]. The generated azido dehydroxyquinine 40 was then allowed to react via a copper catalyzed cycloaddition reaction with a variety of alkynes. These reactions were carried out under mechanochemical conditions in ball mill at 300 rpm, affording the triazolyl compounds 41 in 45% to 91% yields (Table 8). Screening results showed that from the 19 synthetized compounds, 5 showed significant antimalarial and antileishmanial activities (Table 8) and four of them did not reveal any in vivo (rodent animal model) toxic manifestation at doses as high as 1000 mg/Kg.
Finally, S. Sampath et al. reported last year [75] the synthesis of 1,2,3-triazole tethered 3-hydroxy-2-oxindoles under ball milling conditions (Scheme 12) as corrosion inhibitors and antimicrobials. 3-Functionalized oxindoles can be obtained from the valuable heterocyclic scaffolds isatins. Among the different 3-substituted oxindoles, the 3-hydroxy-3-substituted-2-oxindoles are present in many natural products [76]. In addition, they are considered as valuable key intermediates in organic synthesis [77–79], leading to compounds with pronounced pharmaceutical properties [80–82]. The authors synthesized a set of new derivatives of this family by combining an aldol condensation and a click reaction using ball milling conditions. A mixture of N-propargyl isatin, acetophenone, benzyl azide in the presence of DABCO and copper oxide nanoparticles CuONPs (2.5 mol%) was reacted in a ZrO\textsubscript{2} jar material at a speed of 400 rpm, affording the desired products 42 in 87\% to 92\% yields, except for azide possessing the strong electron withdrawing NO\textsubscript{2} group (80\% yield). Among the compounds synthesized, derivative 42b displayed a remarkable corrosion inhibition potency (for corrosion inhibition in acidic media see references [83,84]), while compound 42a showed appreciable antifungal (C. albicans) and antibacterial (S. aureus) effects (Table 9). The authors consider that the biological results are quite encouraging for triggering a detailed structure-activity study and the comprehension of their activity.
Finally, S. Sampath et al. have found enormous applications in medicinal and agricultural sciences. A great number of drugs are extensively used in clinics. Among them, we can point to the antifungal fluconazole 43, antitumoral letrozole 44, and the antiviral ribavirin 45, (Figure 4) [85], while several triazole based compounds play an important role in agriculture ensuring harvest and crops [86]. Their extensive medicinal, agrochemical potential, resulted in an overwhelming effort to develop synthetic methods that include three categories of synthetic objectives: (a) cyclizations to form the triazole ring, (b) transformations of heterocyclic compounds to construct the triazole ring, and (c) substitutions on the 1,2,4-triazole ring.

**Figure 4.** Antifungal fluconazole 43, antitumoral letrozole 44, the antiviral ribavirin 45.

In this review, we discuss only some examples of the recent cyclization reactions with amidrazones and hydrazides. In addition, it is noteworthy to point out that there are no reported methods to synthetize 1,2,4-triazole frames under green chemistry conditions. We report here our first results concerning the mechanochemical organic synthesis of a valuable annulated 1,2,4-triazole scaffold.

### Table 9. Compounds 42 Structure, Yield and Activity.

| Compounds 42 | R     | R₁    | R₂     | Yield (%) | Biological Effect         |
|--------------|-------|-------|--------|-----------|---------------------------|
| a            |       | H     |        | 90        | Anti-fungal, Anti-bacterial|
| b            |       | 5-Me  |        | 92        | Main activity (f.i.): Corrosion inhibition |

**Scheme 12.** Synthesis of 1,2,3-Triazole Tethered 3-Hydroxy-2-oxindoles.
3.1. From Amidrazones

Amidrazones are the conjugated products of imines and hydrazines; their cyclisation with carbonyl compounds is one of the most important pathways to access 1,2,4-triazole derivatives.

In 2015, Nakka et al. [87] reported an environmentally benign protocol for the synthesis of 3,4,5-trisubstituted 1,2,4-triazoles 48. The authors performed the coupling/cyclisation reaction by heating in polyethylene glycol and in the presence of ceric ammonium nitrate (catalyst, 5%), N-arylamidrazones 46 and aldehydes 47. The authors demonstrated that this protocol could generate good yields of 3,4,5-tri-substituted 1,2,4-triazoles bearing different functionalities, while in addition, the effective recyclability of the medium could make the process industrially interesting (Scheme 13).

\[
\begin{array}{c}
\text{Ph} - \text{N} \quad \text{NH}_2 \\
+ \quad R_1 - \text{CHO} \\
\text{PEG, 80 °C} \\
\text{Ph} - \text{N} \quad \text{N} \quad \text{R}_2 \\
\end{array}
\]

Scheme 13. Synthesis of 3,4,5-trisubstituted 1,2,4-triazoles 48.

Szöcs et al. [87] have already shown that 1,2,4-triazole frames judiciously attached at the 5-position of a 3-C-glucopyranosyl scaffold give access to very efficient inhibitors of glycogen phosphorylase. In that respect they become extremely important hits as potential antidiabetic agents (type 2 diabetes). Szöcs et al. reported in 2015 [88] 38 new developments concerning the synthetic approaches for this class of compounds and in particular the oxidative ring closures of N\(^1\)-alkylidene carboxamidrazones. When glycosyl cyanides 49 and amidrazones 50 were treated under Raney reductive conditions in the presence of NaHP\(_2\)O\(_2\), they afforded the two tautomeric forms 51 of O-peracylated N\(^1\)-(β-D-glucopyranosyl-methylidene)-arenecarboxamidrazones. Bromination of 52 by N-bromosuccinimide (NBS) led to halogenated 47 type derivatives. The latter can then undergo in pyridine or by NH\(_3\)OAc in AcOH, a ring closing reaction to the desired 3-C-glycosyl-5-substituted-1,2,4-triazoles 53 (Scheme 14).

\[
\begin{array}{c}
\text{Gly-CN} + \text{NH}_2 \quad \text{NH}_2 \\
\text{Ni, Raney, NaH}_{2}\text{PO}_2 \\
\text{AcOH, H}_2\text{O, Py, 40°C} \\
\text{tautomer a} \\
\text{Gly} - \text{N} \quad \text{N} \quad \text{R} \\
\text{NH}_2 \\
\text{tautomer b} \\
\end{array}
\]

Scheme 14. Synthesis of 3-C-glycosyl-5-substituted-1,2,4-triazoles 52.

The authors took advantage of their methodology to create 3,5-diaryl-1,2,4-triazoles 55 starting from N\(^1\)-arylidene-arenecarboxamidrazones 54. Reaction of the latter with NBS/NH\(_3\)OAc in AcOH (whatever the order) afforded triazoles 55, thus demonstrating the general applicability of the method (Scheme 15).
Scheme 14. Synthesis of N\textsuperscript{1}-arylidene-arenecarboxamidrazones and their transformation into 3,5-diaryl-1,2,4-triazoles \textsuperscript{55}.

Among the different compounds synthetized and tested, it is noteworthy to point out that compound \textsuperscript{56} (Figure 5) with an inhibition constant \(K_i\) of 0.41 \(\mu\)M against rabbit muscle glycogen phosphorylase could be considered as the starting point for the development of more potent compounds for pharmacological treatment, not only of diabetes but also wherever the regulation of glycogen metabolism plays a significant role (cerebral and cardiac ischémias, and tumor growth).

\textbf{Figure 5.} C-glucopyranosyl 1,2,4-triazoles \textsuperscript{56}.

In 2018, Aly et al. \textsuperscript{89} reported a general method for the synthesis of 1,3,5-trisubstituted 1,2,4-triazoles \textsuperscript{59, 60} from reaction of amidrazones \textsuperscript{57} with diethyl azodicarboxylate \textsuperscript{58}. The authors performed the coupling/cyclisation reaction between \(N\)-arylamidrazones \textsuperscript{57} and diethyl azodicarboxylate \textsuperscript{58}. The reaction was conducted under reflux in EtOH and catalyzed by a few drops of triethylamine (Scheme 16), thus allowing to get an easy access to the highly diverse triazoles \textsuperscript{59, 60}. The reaction is based on oxidation of ethanol to acetaldehyde via the Mitsunobu reagent; upon reaction with amidrazones the substituted 3-methyltriazoles \textsuperscript{59} could be obtained, while a \([2 + 3]\) cycloaddition reaction between two oxidized forms of amidrazones produced triazoles \textsuperscript{60}.

\textbf{Scheme 16.} Reaction of the Mitsunobu reagent on amidrazones \textsuperscript{57}.

3.2. \textit{From Hydrazides}

Several hydrazides are commercially available and the non-commercial ones are successfully prepared by the reaction of hydrazine with the corresponding ester precursor. Up to now, a lot of works have been done concerning the cyclization of hydrazides or their derivatives. Very recent work in relation to biological activities is presented here.

In 2018, Singh et al. \textsuperscript{90} reported the design and synthesis of new bioactive 1,2,4-triazoles as potential antituberculosis and antimicrobial agents. In that respect, they synthetized a series of functionalized 1,2,4-triazole derivatives through the synthetic scheme presented below (Scheme 17).
Isonicotinic acid hydrazide 61 was transformed to the potassium dithiocarbazinate 62 by reaction with carbon disulfide under basic conditions. Then, treatment with hydrazine hydrate under thermal conditions in water, afforded the 4-amino-1,2,4-triazole-3-thiol 63. The latter reacted on its 4-amino group to form various Schiff base compounds. Some of the compounds were found to have very potent antitubercular activities, even better than isoniazid and also against clinical isolates (Table 10).

| Compounds 64 | Percent Reduction in RLU |
| R | M. tuberculosis H37Rv | Clinical Isolates, S, H R and E resistant M. tuberculosis |
|---|---|---|
| 4-F-C₆H₄- | 80.50 | 50.72 |
| 4-CH₃-C₆H₄- | 82.01 | 52.79 |
| CH=CH-C₆H₅ | 83.49 | 52.47 |

Synthetized compounds were also tested in vitro against representative bacterial and fungi strains, one compound has very potent activity against B. subtilis, while another one is very potent against A. niger and C. albicans fungi (Table 11).

| Compounds 64 | Minimum Inhibitory Concentration (MIC, µg/mL) |
| R | Gram-Positive Bacteria | Fungi |
|---|---|---|
| | S. aureus | B. subtilis | A. niger | C. albicans |
| 4-F-C₆H₄- | 12.5 | 10.2 | 125 | 106.3 |
| -CH=CH-C₆H₅ | 75 | 81.3 | 11.7 | 10.9 |

Sonawane et al. reported in 2017 [91] the synthesis of 1,2,4-triazole-3-thione derivatives as antimycobacterial agents. The two routes employed by the authors are outlined in Scheme 18. The acid chloride 66, prepared by reacting aromatic carboxylic acid 61 with thionyl chloride reacted with thiosemicarbazide, which without isolation and upon thermal heating under aqueous basic conditions, led to the desired compounds 67 (route A). Triazolothiones 69 could not be synthetized by this procedure were obtained by reaction of hydrazide 68 with carbon disulfide, followed by heating in the presence of a 25% ammonia solution (route B).

Two of the compounds synthetized showed high antitubercular activity against the dormant H37Ra strain in vitro and ex vivo; they also showed extremely low cytotoxicity and high solubility indicating the potential of developing these compounds further as novel therapeutics against tuberculosis infection (Table 12).
route A

\[
\text{thiophyl chloride} \quad \overset{\text{dry CH}_2\text{Cl}_2, \text{DMF}, \text{reflux, 4h}}{\longrightarrow} \quad \text{65}
\]

route B

\[
\text{hydrazine hydrate} \quad \overset{\text{reflux, 5h}}{\longrightarrow} \quad \text{68}
\]

Scheme 18. Synthesis of 1,2,4-triazole-3-thione 67 and 68.

Table 12. In vitro anti-tubercular and anti-bacterial (gram-positive and gram-negative bacteria) activities of thione derivatives 67, 69 (IC\text{50} and MIC values—µg/mL).

| Compound | Anti-Tubercular Activity against MTB (µg/mL) | Cytotoxic Activity of Triazole Thiones against Human Cancer Cell Lines (µg/mL) | Aqueous Solubility (µM) |
|----------|---------------------------------------------|---------------------------------------------|------------------------|
|          | Dormant Stage | Active Stage | THP-1 | A549 | PANC-1 |               |
|          | MIC | MIC | GI\text{50} | GI\text{50} | GI\text{50} | GI\text{50} |     |
| 67        | 0.64 | 9.05 | >100 | >100 | >100 | >100 | >1280 |
| 69        | 0.46 | >30 | >100 | >100 | >100 | >100 | >3200 |

MTB H37Ra, Mycobacterium tuberculosis H37Ra; Growth Inhibition (GI): GI\text{50} (concentration which resulted in 90% decrease in cell viability). Expressed in µg/mL. THP-1: acute monocytic leukemia; A549: lung adenocarcinoma; PANC-1: pancreas carcinoma.

Liu et al. [92] reported in 2017 a family of 7-hydroxy-4-phenylchromen-2-linked 1,2,4-triazoles with potent antitumoral activities. The synthetic procedure adopted made use of the coumarin synthetized derivatives 70 that were functionalized by reaction with ethylbromooacetate followed by transformation of the ester group to a hydrazide functionality. The latter, when condensed with dimethylacetal followed by a strong thermal reaction with an amine in the presence of glacial acetic acid afforded the triazole derivatives 73 in good overall yields (Scheme 19).

Scheme 19. Synthesis of 7-hydroxy-4-phenylchromen-2-linked 1,2,4-triazoles 73.

The new 1,2,3-triazole derivatives showed improved antiproliferative activities. Particularly, compound 74 exhibited potent activity with important IC\text{50} values against AGS (2.63 ± 0.17), MGC-803 (3.05 ± 0.29) and HCT-116 cell lines (11.57 ± 0.53 µM) (Figure 6). The authors also demonstrated that these compounds had strong activity against the HeLa cell line, with an IC\text{50} value of 13.62 ± 0.86 µM. All activities were better than those of the non-substituted 7-hydroxy-4-phenyl-2H-chromen-2-one 70 (with R = H) and that of the positive control drug 5-fluorouracil. Moreover, the authors showed that the biolog-
ical activities of the 1,2,4-triazole derivatives were significantly higher than that of the 1,2,3-triazole ones.

Figure 6. 7-hydroxy-4-phenyl-2H-chromen-2-one 70 (with R = H) and 7-((4-(4-Chlorophenyl)-4H-1,2,4-triazol-3-yl)methoxy)-4-phenyl-2H-chromen-2-one 74.

3.3. Annulated 1,2,4-Triazole Systems

Among the annulated 1,2,4-triazole systems we will present the triazolophthalazine frame that was developed in our group by conventional and non-conventional means.

Some years ago, De et al. [93] explored the possibility of cinnamic acid derivatives as potential antituberculosis agents. In the course of their studies the authors synthesized 4-alkoxy cinnamoyl derivatives resulting from the coupling of the corresponding acids (or activated ones) with different nucleophiles and among them amines, hydrazines, thiolis.

In the course of their first studies, when reacting under peptide coupling conditions (EDC, HCl, HOBt, and trimethylamine), 1-hydrazinophthalazine hydrochloride 75, and cinnamic acid derivatives for 48 h under reflux in acetonitrile, the authors obtained in good yields the corresponding 3-(4-alkoxystyryl)-1,2,4triazolo[3,4-a]phthalazines 77 (65–90%). This was formed through a coupling-intramolecular cyclization-dehydration sequence (Scheme 20).

Scheme 20. Synthesis of 3-(4-alkoxystyryl)-1,2,4triazolo[3,4-a]phthalazines 77.

All alkoxylated compounds showed good antitubercular activities. More importantly, triazolophthalazine derivative 73 (Figure 7), bearing a 4-isopentenyloxy chain on the phenyl ring, showed excellent antitubercular potency (MIC = 1.4 μM) in addition to a very good cytotoxicity toward HCT116 human cells (IC_{50} = 449 μM; 160 μg/mL and selectivity index SI = 320). It is also noteworthy to point out that this compound does not act on the mycolic acid biosynthesis of mycobacteria and up to now its target is unknown.

Figure 7. (E)-3-[4-(3-Methylbut-2-enyloxy)styryl]-1,2,4triazolo[3,4-a]phthalazine 78.

In order to build a small, focused library of styryltriazolophthalazines, Veau et al. [94] modified the convergent route to a divergent one, by exploring the possibility of the construction of the phenolic precursor 79 that could then lead to various alkoxylated derivatives of type 77 (Scheme 21).
The target precursor 79 was thus obtained by the authors in a two step procedure: 1-hydrazinophthalazine hydrochloride 74 and \( p \)-hydroxy cinnamic acid 80 when reacted in acetonitrile for 1 h under peptidic coupling conditions but under microwave afforded a 69% yield after a simple filtration–recrystallization sequence of the styrylphenolic triazolophthalazine 79. Alkylation by various alkylating agents and under standard conditions led to the desired alkoxylylated derivatives 81 (alkylating agent, \( K_2 CO_3 \), KI, DMF, 60 °C, overnight) (Scheme 22).

![Scheme 21. Convergent (A) and divergent (B) routes for synthesis of compounds 77.](image)

In continuation of this work on the triazolophthalazine frame the authors considered that this could be an interesting pharmacophore to explore. In that respect they also synthetized two compounds bearing either an alkyne group or a bromine at the 2 position of the 1,2,4-triazole frame.

Concerning the alkyne compound, after several attempts, Veau et al. [94] considered the best way to obtain it is a two step sequence. Coupling of trimethylsilyl propiolic acid 82 with 1-hydrazinylphthalazine hydrochloride 74 under mild conditions afforded acyclic precursor 83. Under microwave irradiation in acetonitrile for 1 h, the precursor was transformed quantitatively to the cyclized silylated intermediate 84. Upon standard deprotection conditions (i.e., \( K_2 CO_3 \) in MeOH) compound 84 led to derivative 85 (31% overall yield over three steps) (Scheme 23).

![Scheme 22. Synthesis of alkoxylylated trans-styryltriazolophthalazine derivatives 81.](image)

![Scheme 23. Synthesis of 3-ethynylated derivative 85.](image)
For the first time, Gonnet et al. [95] reported in 2019 the mechanochemical synthesis of 1,2,4-triazoles starting from hydralazine hydrochloride. By using a planetary ball-mill, and in the presence of pyrogenic S13 silica as the grinding auxiliary, total conversion to intermediate hydrazones 86 was achieved in a few minutes (Scheme 24). Iodobenzene diacetate (IBD) was used for optimal conversion of nonphenolic hydrazones to annulated 1,2,4-triazoles 87, while SeO$_2$ was found to be efficient for phenolic compounds (Scheme 25).

Scheme 24. One pot two step mechanochemical synthesis of annulated 1,2,4-triazoles 87.

Scheme 25. One pot two step synthesis of phenolic 1,2,3-triazoles 89.
In addition, for the first time, the one-pot two-step synthesis (Scheme 24) leading to annulated 1,2,4-triazoles was also successfully conducted. Comparison to the conventional syntheses of hydrazone 86 and triazole 87 clearly showed the green metrics overall efficiency of the mechanochemical synthesis (Table 13).

Table 13. Comparison of Green metrics for the synthesis of Hydrazone 86 and Triazole 87, in solution/by mechanochemistry.

| Compound | Time (h) | Yield (%) | E-factor | PMI |
|----------|----------|-----------|----------|-----|
| 86       | 1/0.5    | 100/100   | 14/4     | 15/5|
| 87       | 4/0.75   | 84/97     | 74/12    | 75/13|

Synthesis of the brominated compound 91 was also explored. First a conventional method was performed by Veau et al. [94], by reacting overnight under reflux in ethanol, 1-hydrazinyl-phthalazine hydrochloride 74 with trimethylorthoformate in the presence of some drops of acetic acid in order to obtain unsubstituted triazolophthalazine 90. Then, reaction with bromine in the presence of pure acetic acid under reflux afforded the brominated compound 91 in 53% yield.

More recently, Gonnet et al. [95] reported the two-step bromination reaction successfully conducted by mechanochemistry. Reaction of 1-hydrazinylphthalazine hydrochloride 74 in the presence of trimethylorthoformate and some drops of acetic acid reacted in a planetary ball mill for 1 h affording quantitatively triazolophthalazine 90. Reaction in the same planetary ball-mill (PBM) of the triazolophthalazine with sodium bromide, oxone and some silica afforded after 1 h the brominated compound 91 in quantitative yield (Scheme 26).

Scheme 26. Syntheses of 3-bromo-[1,2,4]triazolo[3,4-a]phthalazine 91 by classical method (above) and mechanochemical condition (below).

All compounds bearing the 1,2,4-triazole frame were evaluated for various biological properties. The 3-aryl substituted 1,2,4-triazole derivatives 87 and 89 do not present valuable activities against *M. tuberculosis* (MIC around 80 μM) [96]. The alkyne derivative 85 present a very good activity (MIC 12.9 μM) while the brominated 91 is much less potent (MIC 40 μM) [94]. The alkyne derivative 85 did not manifest cytotoxicity toward HCT116 human cells while it is equally active against multidrug-resistant *M. tuberculosis* strains. Considering all results starting from compound 73, it seems likely that the triazolophthalazine could be an important scaffold in order to obtain new families of compounds with strong antitubercular activity and an alternative mode of action for compared with standard anti *M. tuberculosis* drugs.

In that respect, the authors consider the possibility of developing focused libraries of triazolophthalazine compounds by using the two important precursors that are the alkyne and the brominated derivatives and developing coupling reactions under conventional and/or mechanochemical means. In addition, further work is necessary to tackle the identification of the protein targeted by this class of potent anti *M. tuberculosis* compounds.
4. Mechnochemical Cycloreversion of 1,2,3-Triazoles

Globally, the cycloaddition process is strongly favored thermodynamically \((\Delta H = -45 \text{ to } -55 \text{ Kcal/mol})\) [97]. The 1,2,3-triazole frame is robust and inert under most thermal chemical treatments but also in aqueous or biological environments. In 2011, Brantley et al. [98] reported the possibility of unclicking the click on specific 1,4-substituted 1,2,3-triazoles by mechanical forces. They first hypothesized that mechanical exogenous forces directed to judiciously chosen scaffolds incorporated in a polymer chain can formally disallow pericyclic reactions. The authors incorporated the triazole ring in polymer chains and one of them was judiciously chosen, when mechanical ultrasound forces were applied (ultrasonication in a Suslick cell at 0 °C) resulted in the cleavage of the triazole ring 92 to its alkyne 93 and azide 94 components (Scheme 27).

![Scheme 27](image)

Scheme 27. Effect of ultrasound on the triazole 92.

They concluded that the ability to selectively deconstruct triazoles might serve to elaborate mechano-responsive materials for potential controlled bioconjugation applications or force responsive fluorescent tags for biological assays.

This stimulated a puzzling publication (that was retracted since by the editor) [99] aroused strong debate. The same authors, based on sonochemical experiments related to extended Bel theory, discussed and concluded on the lowering of the activation energy barrier for cycloreversion [100] through application of an external force to the triazole ring. In the contrary, purely theoretical work, it was shown that the cycloreversion barrier is as high as 70 Kcal/mol [101]. In addition, the mechanochemically induced retro-click of the 1,2,3-triazole ring vs. bond rupture next to it could not be unambiguously concluded when single molecule force spectroscopy experiments were applied [100]. Stauch and Dreuw reported in 2017 [102] a theoretical work where by using the JEDI (Judgment of Energy D1stribution) analysis it was concluded that for 1,4 disubstituted triazoles the unclick reaction is impossible, even when Cu^1 assisted (Scheme 28a). For 1,5-disubstituted triazoles where a parallel alignment of the scissile bond exists, this could be feasible. Nevertheless, the retro click cycloreversion is not selective as it competes with the carbon-nitrogen bond connecting the triazole ring to the linker. During the same year, Krupička et al. [103] also concluded that only in these 1,5-disubstituted 1,2,3-triazole systems are the Gibbs free energy barriers 55 Kcal/mol (unclick reaction) versus 45 Kcal/mol for external C-N bond cleavage. The authors also point out an extremely exciting finding by showing that the calculated Ru-assisted mechnochemical cycloreversion of the 1,5 regioisomer dramatically lowers the activation energy of the rate determining step down to 20 Kcal/mol (the first step), while the decomplexation of the cleaved intermediate readily occurs, leading to the alkyne and azide components (Scheme 28b).
concluded when single molecule force spectroscopy experiments were performed showing that the calculated Ru-assisted reaction is impossible.

There is still a lot to be done, especially in terms of developing new synthetic methodologies under green chemistry approaches based on less energy requirements. It is our feeling that the mechanical approaches will be further developed (mechanochemical synthesis, ultrasound). There is still a lot to be invented and this is a great opportunity for the chemists and medicinal chemists, but also in the pharmaceutical industry.

The same and in a greater extent is also true for the synthesis of the valuable regioisomeric scaffold of 1,2,4-triazole systems. Except for our contribution in the field, there is no other green chemistry (ultrasound or mechanochemistry) developed for this family of compounds. We believe that here also great opportunities exist for all communities of synthetic, physical, theoretical and medicinal chemists, whether they are in the academia or in the industry.

Finally, concerning the mechanochemical (ultrasound) unclicking of the 1,5 disubstituted 1,2,3-triazole, active experimental research work is needed in order to create an extremely selective process that could also confirm the theoretical work. That could pave the way for important biological and other applications.

**Scheme 28.** Reactions of (a) 1,4-dimethyl 1,2,3-triazole leading to mononuclear CuI-catalyzed cycloreversion and (b) 1,5-dimethyl 1,2,3-triazole leading to RuII-catalyzed cycloreversions.

In conclusion, the authors point out that the Ru-assisted mechanochemical unclicking of the 1,5 regioisomer could be an extremely selective process. If this is to be experimentally proved it would open the path for very important potential applications.

5. Conclusions

Among the nitrogen-contained heterocyclic ring structures one of the most important providing long term advancement in the medical field are triazoles. They became in the last decades the heterocycle of choice in all fields of drug discovery receiving much of the attention and offering new opportunities for medicinal chemists.

We must point out that while 1,2,3-triazole systems are very well documented in terms of classical organic synthesis, their synthetic methodologies under green chemistry approaches based on less energy input requirements are beginning to emerge, but are still focused on ultrasound reactions. It is our feeling that the mechanical approaches will be further developed (mechanochemical synthesis, ultrasound). There is still a lot to be invented and this is a great opportunity for the chemists and medicinal chemists, but also in extenso for the pharmaceutical industry.

The same and in a greater extent is also true for the synthesis of the valuable regioisomeric scaffold of 1,2,4-triazole systems. Except for our contribution in the field, there is no other green chemistry (ultrasound or mechanochemistry) developed for this family of compounds. We believe that here also great opportunities exist for all communities of synthetic, physical, theoretical and medicinal chemists, whether they are in the academia or in the industry.

Finally, concerning the mechanochemical (ultrasound) unclicking of the 1,5 disubstituted 1,2,3-triazole, active experimental research work is needed in order to create an extremely selective process that could also confirm the theoretical work. That could pave the way for important biological and other applications.

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