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Copper-Catalyzed [3 + 2] Cycloaddition/Oxidation Reactions between Nitro-olefins and Organic Azides: Highly Regioselective Synthesis of $NO₂$ -Substituted 1,2,3-Triazoles

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S Supporting Information

[AB](#page-2-0)STRACT: [A new coppe](#page-2-0)r-catalyzed $[3 + 2]$ cycloaddition/ oxidation reaction of nitro-olefins with organic azides has been developed to afford $1,4(-NO₂),5$ -trisubstituted $1,2,3$ -triazoles. This reaction sequence has a broad substrate scope and affords NO2-substituted 1,2,3-triazoles with high regioselectivities and in good to excellent yields. The involved oxidative process

overcomes the elimination of $HNO₂$ for general cycloaddition of nitro-olefins with organic azides, which shows a high atom economy and potential applications.

1,2,3-Triazoles are an important class of heterocyclic compounds with wide applications in medicinal, 1 material, 2 and synthetic organic chemistry.³ For example, some 1,2,3-triazoles show a broad spectrum of biological acti[vi](#page-2-0)ties suc[h](#page-2-0) as antifungal, antiviral, antialler[gi](#page-2-0)c, anti-HIV, and antimicrobial activities.⁴ Because of their reliability, the diverse syntheses of 1,2,3-triazoles have gained much attention. The most popular methods f[or](#page-3-0) these kinds of compounds are based on copper $(I)^{5}$ and ruthenium(II)-mediated° regioselective azide−alkyne cycloaddition reactions, which give 1,4- or 1,5-disubstituted [1](#page-3-0),2,3 triazoles, respectively. [Re](#page-3-0)cently, some fully substituted 1,2,3 triazoles were also synthesized by different approaches, such as 1,4,5-trisubstituted 1,2,3-triazoles, which were obtained by metal-catalyzed cycloaddition reactions of organic azides with internal alkynes⁷ and by organocatalytic cycloaddition reactions or multicomponent reactions of organic azides with activated enones, keto e[st](#page-3-0)ers, nitriles, or aldehydes.⁸ At the same time, other 1,2,3-triazoles were also synthesized by post-functionalization of $NH-1,2,3$ -triazoles or 1,4-disubstitu[te](#page-3-0)d 1,2,3-triazoles.⁹

Nitro-olefins also serve as an important partner with azides to form the 1,2,3-triazoles (Scheme 1).¹⁰ Cyclization reactions [o](#page-3-0)f nitro-olefins with NaN_3 can give the $NH-1,2,3$ -triazoles (Scheme

1a).^{10a−c} TMSN₃ as a nitrogen source with tetra-*n*-butylammonium fluoride is an alternative method for the formation of NH-1,2,[3-tr](#page-3-0)i[a](#page-3-0)zoles (Scheme 1b).^{9d} In general, organic azides react with nitro-olefins to give the N-substituted 1,2,3-triazoles with low efficiency (Scheme 1c).1[0e](#page-3-0) Interestingly, Wang et al. reported that nitro-olefins could also react with organic azides in the presence of $Ce(OTf)$ ₃ to [giv](#page-3-0)e 1,5-disubstituted 1,2,3-triazoles (Scheme 1d).^{10f} For these four cases, the formation of 1,2,3triazoles involved an elimination of HNO₂ process. Nitrogengroup-substit[uted](#page-3-0) 1,2,3-triazoles are very important for their applications; most of them were synthesized by the click reaction of electron-deficient ynamides with organic azides; 11 even so, the diverse nitrogen-group-substituted 1,2,3-triazoles are still scarce. The $NO₂$ group [is](#page-3-0) a good nitrogen source, which is widely used for synthesis of N-containing compounds.¹² We propose that, if the dehydrogenation process is involved instead of the elimination of $HNO₂$ between the cycli[zat](#page-3-0)ion of nitro-olefins with organic azides, it can give the $NO₂$ -substituted 1,2,3triazoles. The challenge for this design is to inhibit the elimination of HNO₂ effectively. Herein we present a coppercatalyzed highly selective synthesis of $NO₂$ -substituted 1,2,3triazoles from nitro-olefins and organic azides, which involved the loss of two hydrogens instead of $HNO₂$ for nitro-olefins (Scheme 1e).

The reaction conditions were tested by using a model reaction between nitrostyrene $(1a)$ and benzyl azide $(2a)$, including different metal salts, solvents, and additives, and the results are shown in Table 1. The $Cu(I)$ catalysts were first tested including CuI and CuCl because the organic azides can coordinate with copper comple[xe](#page-1-0)s, especially the $Cu(I)$ complex. Conversions were up to 60% using dimethylformamide (DMF) as solvent under air atmosphere; however, the yields for the proposed oxidative cycloaddition product 3a were less than 10%, and the

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Table 1. Optimization of the Reaction Conditions^{a}

Ph	BnN₂ 2a 1a	Ph conditions O ₂ N $N = N$ 3a		Ph
entry	catalyst	solvent	concn (%)	yield of 3a/4a $(\%)^b$
1	CuI	DMF	67	7/23
$\overline{2}$	CuCl	DMF	64	5/22
3	CuO	DMF	66	35/19
$\overline{4}$	CuCl ₂	DMF	78	23/41
5	CuBr ₂	DMF	80	28/43
6	Cu(OAc)	DMF	73	27/36
7	Cu(NO ₃) ₂	DMF	81	56/17
8	CuSO ₄	DMF	86	77/4
9	Cu(OTf),	DMF	90	83/trace
10	Cu(OTf),	toluene	23	trace/20
11	Cu(OTf) ₂	MeOH	67	16/34
12	Cu(OTf),	DMSO	76	68/trace
13 ^c	Cu(OTf),	DMF	95	85 / < 5
14	$Cu(OTf)$ ₂	DMF/AcOH (10%)	95	90/trace
15	Cu(OTf),	DMF/AcOH (25%)	100	96/0
16	FeCl ₃	DMF	$<$ 5	$-/-$
17	PdCl ₂	DMF	38	trace/22
18	Pd(OAc),	DMF	37	trace/30
19	PdCl ₂ (PPh ₃)	DMF	68	trace/27

^aReactions conditions: nitrostyrene $(1a)$ (1 mmol) , benzyl azide $(2a)$ (1.2 mmol), and catalyst (0.05 mmol), air (1 atm), solvents (5 mL for each, 0.2 M), 110 °C, 14 h. $\rm ^{b}Yields$ of products purified by column chromatography on silica gel. ${}^{c}O_{2}$ (1 atm) was used.

major product was $4a$, which formed via elimination of $HNO₂$. We then tested the $Cu(II)$ catalyst; it is gratifying that the expected 3a was obtained with 35% separated yield by using CuO as catalyst (entry 3), and in this case, 4a was the minor product with 19% yields. Other Cu(II) salts were tested for improving the conversion and selectivity, including CuCl₂, CuBr₂, Cu(NO₃)₂, $Cu(OAc)₂$, $CuSO₄$, and $Cu(OTf)₂$ (entries 4–9). By using $CuCl₂$, $CuBr₂$, and $Cu(OAc)₂$, the conversions were increased to a certain extent; however, the yields of 3a were not modified, while the yields of 4a were increased. When $Cu(NO₃)₂$ and CuSO4 were used, the yields of 3a were increased obviously, and the formation of 4a was inhibited greatly when $CuSO₄$ was used. When more acidic $Cu(OTf)_2$ was chosen as catalyst, the conversion increased to 90% and the yields of 3a increased to 83%. Other solvents were also tested for the optimization conditions, but they were not better than DMF. Because there is an oxidative process for the formation of 3a, the reaction was tested at the oxygen atmosphere (entry 13); however, no improvement was observed in the conversion and yield. By contrast with different $Cu(II)$ salts, we proposed that the acidities of the catalysts may be crucial for this transformation, so AcOH was introduced for this reaction system. The yield reached up to 90% if 10% AcOH was introduced, and when 25% AcOH was added, it gave 100% conversion and the yield was up to 96% (entry 15). Other metal salts were also tested for this reaction, including $FeCl₃$ and $Pd(II)$ salts, and they did not give satisfying results.

With the optimized reaction conditions in hand, we studied the scope and limitations of this transformation with a wide range of nitro-olefins and organic azides (Scheme 2). Both electronScheme 2. Substrate Scope for the Cycloaddition Reaction of Nitro-olefins 1 and Azides 2^a

^aReactions conditions: 1 (1 equiv), 2 (1.2−1.5 equiv), and $Cu(OTf)_2$ (5 mol %), air (1 atm), DMF/AcOH (4:1, v/v, 0.2 M), 14−20 h. Yields of isolated products are given.

donating functional groups, such as methoxy (3d, 3h, and 3r), methyl substituents (3e, 3j, 3l, and 3q), and electronwithdrawing groups, such as halogens moieties (3b, 3c, 3f, 3g, 3i, 3m, 3o, and 3p), on the aromatic rings of the nitro-olefins were compatible with this transformation. The structure of 3b was identified by X-ray single-crystal analysis. Furthermore, acidsensitive furanyl-substituted nitro-olefins were also suitable substrates to react with azides (3s and 3t), and yields (78 and 75%) were somewhat lower than those of other nitro-olefins; we ascribe this to the decomposition of the furanyl groups. In addition, different organic azides, including aliphatic and aromatic azides, were also suitable substrates, and the corresponding products were obtained in good to excellent yields. The structure of 3p was also identified by X-ray singlecrystal analysis (see Supporting Information). However, the reaction for nitro-olefins with the aromatic azides provided the corresponding prod[ucts in slightly dimin](#page-2-0)ished yields in comparison with benzyl azides; furthermore, excess azides (1.5 equiv) were needed for full conversions of nitro-olefins (see Supporting Information).

This strategy is a direct method for synthesis of $NO₂$ [substituted 1,2,3-triazo](#page-2-0)les via introduction of an oxidative dehydrogenative process instead of the elimination of $HNO₂$, which shows higher atom economy. More importantly, the corresponding $NO₂$ -substituted 1,2,3-triazoles are difficult to obtain by nitration of $C4(5)$ −H 1,2,3-triazoles.¹

To get insight into the mechanism of the reaction, some control experiments were done (Scheme 3). [The](#page-3-0) Cu/TEMPO (2,2,6,6-tetramethylpiperidine-N-oxyl) system was widely used for aerobic oxi[d](#page-2-0)ation reactions, 14 and TEMPO was first

Scheme 3. Control Experiments for the Cyclization of Nitroolefins with $BnN₃$

introduced for this reaction; however, it was found that formation of 3a was totally inhibited. In this case, 4a via elimination of $HNO₂$ also formed with 58% yield. When AcOH was used, 3a still cannot be detected, but the elimination of $HNO₂$ may be inhibited to a certain extent (only 36% yield for 4a). We proposed that the formation of 3a should involve a radical process: TEMPO served as a radical inhibitor,¹⁵ while formation of 4a was not related to the oxidation process. Furthermore, the reaction was run under Ar atmospher[e u](#page-3-0)sing 2 equiv of CAN (ceric ammonium nitrate) as oxidant or a stoichiometric amount (2.5 equiv) of $Cu(OTf)_{2}$, and 3a was normally formed with 90 and 95% separated yields, respectively. So the air just served as an oxidant to promote the regeneration of the Cu(II) catalyst. We also checked the alkyl-substituted nitro-olefin 1b under standard conditions. The reaction gave 100% conversion; however, the corresponding product was 1,5 disubstituted 1,2,3-triazole 4b.

According to these control experiments, a plausible mechanism for this Cu(II)-catalyzed oxidative $[3 + 2]$ cycloaddition reaction is outlined in Scheme 4. The first step of the reaction is

Scheme 4. Plausible Mechanim for Cu(II)-Catalyzed $\lceil 3 + 2 \rceil$ Cycloaddition/Oxidation Reaction

the regioselective 1,3-dipolar cycloaddition of nitro-olefin 1 with azide 2 to form the triazoline intermediate I (attempts to synthesize intermediate I were unsuccessful); this process can be promoted by transition metal complexes. $^{10d-\rm f}\rm{The}$ a radical **II** is formed with the help of the $Cu(II)$ catalyst; this radical can be stabilized by an Ar group, which facilit[ates t](#page-3-0)he loss of another electron to give the cation intermediate III, which loses a proton to form $NO₂$ -substituted 1,2,3-triazole 3. Formation of the same kind of radical II for the alkyl-substituted nitro-olefin is not as favorable in energy as that of Ar-substituted nitro-olefins, so it tends to form 1,5-disubstituted 1,2,3-triazoles via elimination of $HNO₂$ under the same conditions. The Cu(II) can be regenerated from Cu(I) species with oxygen under acidic condition. The $Cu(I)$ species react with oxygen to generate $[Cu(II)-O-O$ •] or $[Cu(II)-O-OH]$ ¹⁶ then dissociates to form H_2O_2 and regenerates the Cu(II) catalyst.

In conclusion, we report a copp[er-](#page-3-0)catalyzed $[3 + 2]$ cycloaddition/oxidation reaction of organic azides with nitroolefins. The reactions give $1,4(-NO₂),5$ -trisubstituted 1,2,3triazoles with high selectivity in good to excellent yields, and it proceeds from simple building blocks and tolerates a broad range of functional groups as well as readily available reagents. The potential viability and generality of this reaction for the construction of various 4-amino 1,2,3-triazoles may be anticipated (reduction of the $NO₂$ group); derivatives such as 4-amino 1,2,3-triazoles have already been found to increase potency as antibacterial agents.¹⁷ The detailed reaction mechanism and synthetic applications of this reaction to synthesize other diverse 1,2,3-t[ria](#page-3-0)zoles and other $NO₂$ substituted heterocycles together with biological activities are under investigation.

■ ASSOCIATED CONTENT

S Supporting Information

General experimental procedure, characterization data of the compounds, and CIF data for 3b and 3p. This material is available free of charge via the Internet at http://pubs.acs.org.

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Notes

The authors declare no competing financial interest.

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