Chem. Chem. Technol., 2019, Vol. 13, No. 2, pp. 163–169

HYPERACTIVE MAGNETICALLY SEPARABLE NANO-SIZED MgFe₂O₄ CATALYST FOR THE SYNTHESIS OF SEVERAL FIVE-AND SIX- MEMBERED HETEROCYCLES

Shobha Bansal¹, Yogendra Kumar¹, Dipak Kumar Das¹, Prabal Pratap Singh^{1, *}

https://doi.org/10.23939/chcht13.02.163

Abstract. MgFe₂O₄ nanoparticle ferrites were synthesized by combustion technique using pure ferric nitrate and magnesium carbonate. The magnetically separable MgFe₂O₄ MNP's were found to be hyper active catalyst for the synthesis of a wide range of biologically active five and six-membered heterocyclic moieties at refluxing conditions. Reaction times are lowest in comparison to all reported in literature with excellent yields. Strong electron pull of Fe^{3+} is responsible for its hyper activity, which has been substantiated by substitution of Fe³⁺ by other trivalent metal ions. Mg^{2+} has a unique role because replacement of Mg^{2+} has poor catalytic activity. The developed protocol has been efficiently utilized for the synthesis of a series of substituted mono/bis pyrimidines, pyrimidin-2-ol, pyrimidin-2-thiol, pyrazoles and isoxazoles by condensing monochalcones/1,4-bischalcones with various bis-nucleophiles in the presence of catalytic amount of heterogenous magnetic MgFe₂O₄ nanoparticles. The structure of these synthesized compounds was determined by FTIR, ¹H, ¹³C and mass spectra. The catalyst can be removed easily from reaction mixture by using a simple external magnet. Nanoparticles of ferrite were recovered and reused with no appreciable change in the activity even after the five runs. Nanoparticles are characterized by XRD, TEM and IR spectroscopy.

Keywords: magnetic $MgFe_2O_4$ nanoparticles, mono chalcones, 1,4-bischalcones, pyrimidines, pyrazolines, isoxazolines, external magnetic separation, heterogeneous and reusable catalyst.

1. Introduction

The heterogeneous catalyst has received a remarkable amount of interest from scientific and industrial perspectives because of its enormous impact on

17km stone, NH-2 Delhi-Mathura Highway,

Chaumuhan, Mathura UP 281406, India

world's economy. The use of functionalized magnetic nanoparticle (MNP's) as heterogeneous catalyst has attracted most of the researchers for its easy separation from reaction mixture with the aid of external magnetic field [1-8].

Magnetic nano catalyst [9] is rapidly developing area. Researchers have reported striking novel catalytic properties including greatly enhanced reactivities and selectivities for nanocatalyst compared to bulk counter parts. Recently functionalized magnetic nano particle have been used as efficient catalytic system in many chemical transformation including synthesis of α -amino nitriles [10], 1-1 diacetates form aldehyde [11], diazepiene derivatives [12], indazalo[2,1-b] phthalazine triones, pyrazoles [1,2-b] phthalazine-diones [13], 3,4 dihydro pyrimidines-2(1H)ones [14], 1,4 dihydropyrimidines [15], and pyrrole synthesis [16]. Also a series of organic reaction such as Michael addition [17], Suzuki/Heck cross-coupling [18], asymmetric aldol reaction [19], Suzuki coupling [20], acetalization reaction [21], Ritter reaction [22], cyanosilylation of carbonyl compounds [23], Henry reaction [24], enantioselective direct addition of terminal alkynes to imines [25] have been done using functionalized nanostructure.

In the context of our research of considerable interest is the synthesis of various heterocyclic organic compounds such as substituted pyrimidines, pyrimidin-2ol, pyrimidin-2-thiol, pyrazoles, and isoxazoles for their biological activities [26-33] including antiproliferative, anticancer, antimalerial, antimicrobial, antihypertensive, herbicidal, ACE inhibitors, antitubercular, and others. We are in the process of development of effective and environment friendly basic nanocatalyst in organic synthesis. Although there are so many reports for the synthesis of heterocylic compounds in the absence of catalyst [34-40], all of them have reported higher reaction time which is a matter of concern for industrial applicability of the procedure.

We herein report a hyperactive heterogenous magnetic nanocatalyst for the synthesis of many biologically active five and six-membered heterocycles such as pyrimidines, pyrimidin-2-ol, pyrimidin-2-thiol,

¹ Department of Chemistry, GLA University,

^{*} prabal.singh@gla.ac.in

[©] Bansal S., Kumar Y., Das D., Singh P., 2019

pyrazoles, and isoxazoles having lowest reaction time with high yield. The catalyst is recyclable 5-6 times without any change of reaction time and yield.

2. Experimental

2.1. General Considerations

All the solvents and reagents were used as supplied from commercial sources. FTIR spectra were recorded in KBr on a Schimadzu FTIR 8401 spectrometer and Perkin Elmer version 10.03.06 for the liquid samples. ¹H and ¹³C spectra were recorded on a Bruker DRX 300 spectrometer operating at 300 MHz for ¹H NMR and 75 MHz for ¹³C NMR as solutions in CDCl₃ and DMSOd₆. The ESI mass spectra were measured on waters UPLC-TDQ spectrometer. TLC was performed on silica coated glass plates; spots were developed in I₂ chamber or visualized in UV chamber. The morphology of the catalyst was studied by high resolution electron microscopy HRTEM-300 KV Technai G2 30S TWIN with gold coating equipped with energy dispersive X-ray spectroscopy.

2.2. Preparation of Catalyst

Nanoparticles of $MgFe_2O_4$ were synthesized by combustion technique. 2.0 mmol of $Fe(NO_3)_3 \cdot 9H_2O$ and 1.0 mmol of $MgCO_3$ dissolved in HNO_3 were taken together. After uniform mixing, 1.0 mmol of monoethanolamine was added to the reaction mixture followed by 1.0 mmol of each of sucrose and excess of nitric acid, respectively. The resulting liquid reaction mixture was then put on a hot plate at 353 K till it completely dried to a black residue. This black residue was then kept in muffle furnace at 1073 K for 4–6 h to obtain nanoparticle of MgFe₂O₄.

2.2.1. Typical procedure for the synthesis of 1,4-bis pyrimidines, pyrimidin-2-ol, pyrimidin-2-thiol, pyrazoles and isoxazoles (3d-f, 5d-e)

The starting material monochalcone/1,4-bischalcone **1** (Tables 4 and 5) was prepared as per the reported literature procedure [41] by Claisen-Schmidt condensation between acetophenone and aromatic aldehydes in ethanolic solution of sodium hydroxide.

To a well stirred solution of 3,3'-(1,4-phenylene)bis(1-phenylprop-2-en-1-one)**1**(1 mmol), differentbifunctional nucleophiles**2/4**(2 mmol) in 10 ml absoluteethanol, catalyst (10 mmol %) was added and the reactionmixture was refluxed in oil bath for the appropriate periodof time. The progress of the reaction was monitored byTLC. On completion, the reaction mixture was allowed tocool at room temperature, the catalyst was separated by using external magnet and the solvent was evaporated up to the dryness under reduced pressure to obtain a crude solid product. The solid obtained was stirred in water, filtered and recrystallized from EtOAc:hexane solution.

2.2.2. Typical procedure for the synthesis of pyrimidines, pyrimidin-2-ol, pyrimidin-2-thiol, pyrazoles and isoxazoles (3a-c, 5a-c)

To a well stirred solution of benzylidineacetophenone 1 (1 mmol), different bifunctional nucleophile 2/4 (1 mmol) in 10 ml absolute ethanol, catalyst (10 mmol %) was added and the reaction mixture was refluxed in oil bath for appropriate period of time. The progress of the reaction was monitored by TLC. The reaction mixture was allowed to cool at room temperature, catalyst was separated by external magnet and solvent was evaporated up to the dryness under reduced pressure to obtain a crude solid product. The solid was stirred in water, filtered and recrystallized from EtOAc:hexane solutions.

3. Results and Discussion

From the FTIR spectra of MgFe₂O₄ it is depicted that the sample show two prominent absorption bands v_1 and v_2 in the range of 560–420 cm⁻¹ (Fig. 1). Band v_1 is caused by stretching of tetrahedral cation and oxygen bonding, while v_2 is due to the vibrations of oxygen in a perpendicular direction to the axis joining the tetrahedron ion and oxygen.

The high-resolution transmission electron microscopy (HRTEM) of NP'S is shown in Fig. 2. The particle size of the magnesium ferrite NP's sample is typically in the range of 100–200 nm. The crystals are irregular in shape and are attached to each other along the grain boundaries. The material was verified by XRD data, which matched very well with standard data (JCPDS file no.737960). The peaks are indexed as (220), (311), (222), (400), (331), and (440), Fig. 3. Size of crystallite was found to be 100 nm from analysis of XRD profile by Delay Shapes per 0.91





Fig. 1. FTIR spectrum of magnesium ferrite MNPs



Fig. 2. X-ray diffraction pattern and HRTEM image of MgFe₂O₄ MNP's

In order to check the catalytic activity of the magnesium ferrite catalyst benzylidine acetophenone **1** and guanidine nitrate **2** were taken as starting materials (Scheme 1). The reaction was first performed in the presence of catalytic amount of Fe_2O_3 (20 mmol %) in the refluxing ethanol to afford the formation of 4,6 diphenyl-pyrimidine-2-ylamine **3c** (Scheme 1, Table 1, Entry 1). However, the synthesis was achieved in poor yields. Then the same reaction was repeated with magnesium ferrite MNP's, and the synthesized product was obtained in the excellent yield and in considerably less time (Table 1, Entry 13). The catalytic study was also carried out in the

presence of various other catalysts (Table 1, Entries 2-12) but we got unsatisfactory results with all of them.



Fig. 3. Powder XRD pattern for the MgFe₂O₄ nanoparticle

Only MgFe₂O₄ (Table 1, Entry 13) was found to have the best results in comparison with other metal ion oxides catalysts. During the model study the presence of various solvents like MeOH, MeCN, CHCl₃, PhMe, and DMF was also investigated using 10 mmol % of MgFe₂O₄ MNPs (Table 2). It was observed that ethanol stands out to be the solvent of choice with its fast conversion, high yields and easy removal (Table 2, Entry 2). This is due to efficient solubility of reactants and products and poor coordination with surface metal ions.

We next examined the effect of catalyst loading on to the model reaction (Table 3, Entries 1-5). It was observed that minimum 10 mmol % of $MgFe_2O_4$ MNPs afforded the product with best result (Table 3, Entry 3).



Scheme 1. Optimized (model) reaction condition for synthesis of 4,6-diphenylpyrimidine-2-amine (3c)

Screening of catalyst for the synthesis of 3c

Table 1

Entry	Catalyst [*]	Time, min	Yield, % ^{**}
1	Fe ₃ O ₄	120	20
2	SrFe ₂ O ₄	120	10
3	ZnO	120	<10
4	$ZnO@Fe_2O_4$	120	NP
5	$MgSO_4$	120	NP
6	SrCr ₂ O ₄	120	Traces
7	Bi(NO ₃) ₃ .5H ₂ O	120	NP
8	CoFe ₂ O ₄	120	NP
9	CaFe ₂ O ₄	120	70
10	CuFe ₂ O ₄	120	NP
11	$ZnCr_2O_4$	120	20
13	MgFe ₂ O ₄	25	92

Notes: *reaction conditions: chalcone (1 mmol), guanidine nitrate (1 mmol) and catalyst (10 mmol %) refluxed in ethanol (10 ml) solvent for the mentioned time; ** isolated yields.

Table 2

Entry	Solvent [*]	Time, min	Yield, % ^{**}	
1	EtOH	25	72	
2	MeOH	25	92	
3	MeCN	180	<20	
4	CHCl ₃	180	<20	
5	PhMe	180	<10	
6	DMF	35	60	

The model study in various solvents using MgFe₂O₄ MNP's for the synthesis of 3c

Notes: ^{*}reaction conditions: chalcone (1 mmol), guanidine nitrate (1 mmol) and catalyst (10 mmol %) refluxed in ethanol (10 ml) solvent for the mentioned time; ^{**} isolated yields.

Table 3

Entry	Catalyst, mmol %	Time, min	Yield, % ^{**}		
1	20	30	92		
2	15	32	91		
3	10	25	<92		
4	5	50	<80		
5	3	>60	45		

The effect of catalyst loading on the model reaction

Notes: *reaction conditions: benzalidene acetophenone 1 (1 mmol), 2 (1 mmol), MgFe₂O₄, ethanol reflux; ** isolated yields.

Table 4

Synthesis of substituted mono and bis pyrimidines, pyrimidin-2-ol and pyrimidin-2-thiol (3a-f)



To check the scope and generality of this methodology, several diverse chalcones were treated with various bifunctional nucleophiles to give substituted sixmembered mono- and bis-pyrimidines, pyrimidin-2-ol and pyrimidin-2-thiol heterocycles in good to excellent yields in lesser reaction time. The results of the experiments are listed in Table 4. It was observed that present methodology could be applied to various substrates successfully.

After successful synthesis of six-membered heterocylces we next targeted to synthesize five membered heterocycles like substituted pyrazoles and isoxazoles. Interestingly, using the same protocol we were able to synthesize the desired products in very good yield with lesser reaction time (Table 5).

In order to check the advantages of developed strategy we compared our protocol with the various published reports and it has been found that the catalyst reduced the reaction time drastically from 2-12 h (in the reported processes) to 30-120 min for all reactions with good yields in the present process. The following few examples illustrate the advantages of the process over using the different catalyst Table 6.

In the mechanistic study of chalcone reaction with guanidine nitrate in the presence of MgFe₂O₄MNP's, it is believed that Mg has a well established spinel. In the crystal structure there are two sites of accommodation of metal ions: one is octahedral site and another one tetrahedral site. Tetrahedral site is occupied by Mg^{2+} and octahedral site is occupied by Fe³⁺ ions. All the metal ions are bridged through oxide anions. Thus Mg^{2+} is bridged with Fe³⁺ through oxide ion (O²⁻). Mg^{2+} and Fe³⁺ ions both are Lewis acids but the Fe³⁺ ion is a stronger Lewis acid than Mg²⁺ion. As a result, electron cloud of oxygen anion moves towards Fe³⁺ ion depriving partially Mg^{2+} . Thus Mg^{2+} become more electron deficient leading to its hyperactivity. Role of Fe³⁺ ion is well understood through its substitution by Cr^{3+} and Mn^{2+} , where the catalytic activity of spinel is found to be very poor. The role of Mg^{2+} is very dominant, which is understood by substitution of Mg^{2+} by Ca^{2+} and Sr^{2+} producing no catalytic activity. Therefore, the combination of Mg and Fe is unique for its hyper catalytic activity.





Synthesis of substituted mono and bis pyrazoles and isoxazoles (5a-f)

Compound No	Our protocol		Reported methods		Reaction conditions	Ref.
	Time, min	Yield, %	Time, min	Yield, %		
3c	25	92	60	45	H ₂ O ₂ /KOH/EtOH/Reflux	[42]
3f	55	85	720	69	EtONa/EtOH/Reflux	[43]
3b	45	67	360	90	Amberlyst 15/EtOH/Reflux	[44]
5a	120	85	200	90	PW12/PA/EtOH/318 K	[45]
5a	120	85	300	92	H3PW12O40/EtOH/318 K	[46]
5a	120	85	100	90	Fe ₂ O ₃ @SiO ₂ /PW ₁₂ NP/EtOH/Reflux	[47]
5a	120	85	25	82	PhNHNH ₂ /HCOOH/EtOH/353 K	[48]
5b	120	70	1440	57	NH ₂ NH ₂ /MeOH/Reflux	[49]
5b	120	70	540	70	NH ₂ NH ₂ /EtOH/Reflux	[50]

A comparative study of the developed protocol with the earlier reported methods



Scheme 2. A mechanistic pathway for the formation of 4,6diphenylpyrimidine-2-amine in presence of MgFe₂O₄ MNPs

The separation of MgFe₂O₄ nano catalyst from reaction mixture could be very easily achieved by applying external magnetic field owing to the supermagnetic nature of Fe₂O₃ nanoparticles at room temperature. The reusability is one of the important criteria of heterogeneous magnetic catalyst. The recyclability of magnesium ferrite NP's was investigated for the model reaction of Scheme 1. After completion of reaction, the separated catalyst was reused after washing with acetone and drying at 343 K for 5 h. The reusability of the catalyst was examined by repetitive use of catalyst. It was found that catalyst showed no appreciable change in activity even after five cycles (Fig. 4).



Fig. 4. Recyclability of catalyst for the formation of various heterocycles

4. Conclusions

We have synthesized magnesium ferrite MNP's through facile and simple combustion method that act as highly efficient and reusable heterogeneous catalyst. This MNP are highly active to catalyse hetero condensation of mono- as well as bis-chalcones and various bifunctional nucleophiles to give five- and six-membered heterocylces. The advantages of this catalytic system are milder reaction conditions, shorter reaction times, high yield, easy catalyst preparation, simple and clean work up, and reusability.

Acknowledgements

All the authors humbly thank GLA University, Mathura India for providing the financial assistance and infrastructure for all the research work done.

References

[1] Gardimalla H., Mandal D., Stevens P. et al.: Chem. Commun., 2005, 4432. https://doi.org/10.1039/b504128g [2] Abu-Rezig R., Alper H., Wang D., Post M.: J. Am. Chem. Soc., 2006, 128, 5279. https://doi.org/10.1021/ja060140u [3] Phan N., Gill C., Nguyen J. et al.: Angew. Chem. Int. Ed., 2006, 45, 2209. https://doi.org/10.1002/anie.200503445 [4] Baruwati B., Guin S.: Org. Lett., 2007, 9, 5377. https://doi.org/10.1021/ol702064x [5] Shokouhimehr M., Piao Y., Kim J. et al.: Angew. Chem. Int. Ed., 2007, 46, 7039. https://doi.org/10.1002/anie.200702386 [6] Chouhan G., Wang D., Alper H.: Chem. Commun., 2007, 4809. https://doi.org/10.1039/b711298j [7] Liu J., Peng X., Sun W. et al.: Org. Lett., 2008, 10, 3933. https://doi.org/10.1021/ol801478y [8] Zheng X., Luo S., Zhang L., Cheng J.: Green Chem., 2009, 11, 455. https://doi.org/10.1039/b823123k [9] Lu A-H., Salabas E., Schuth F.: Angew. Chem. Int. Ed., 2007, 46, 1222. https://doi.org/10.1002/anie.200602866 [10] Kassaee M., Masrouri H., Movahedi F.: Appl. Catal. A, 2011, 395, 28. https://doi.org/10.1016/j.apcata.2011.01.018 [11] Esmaeilpour M., Sardarian A., Javidi J.: Appl. Catal. A, 2012, 445-446, 359. https://doi.org/10.1016/j.apcata.2012.09.010

[12] Maleki A.: Tetrahedron Lett., 2013, 54, 2055. https://doi.org/10.1016/j.tetlet.2013.01.123 [13] Kiasat A., Davarpanah J.: J. Mol. Catal. A, 2013, 373, 46. https://doi.org/10.1016/j.molcata.2013.03.003 [14] Zamani F., Izadi E.: Catal. Commun., 2013, 42, 104. https://doi.org/10.1016/j.catcom.2013.08.006 [15] Gawande M., Bonifácio V., Varma R. et al.: Green Chem., 2013, 15, 1226. https://doi.org/10.1039/c3gc40375k [16] Mahmoudi H., Jafari A.: Chem. Cat. Chem., 2013, 5, 3743. https://doi.org/10.1002/cctc.201300623 [17] Nemati F., Heravi M., Rad R.: Chin. J. Catal., 2012, 33, 1825. https://doi.org/10.1016/S1872-2067(11)60455-5 [18] Du Q., Zhang W., Ma H. et al.: Tetrahedron, 2012, 68, 3577. https://doi.org/10.1016/j.tet.2012.03.008 [19] Yang H., Li S., Wang X. et al.: J. Mol. Catal. A, 2012, 363-364, 404. https://doi.org/10.1016/j.molcata.2012.07.017 [20] Li W., Żhang B., Li X. et al.: Appl. Catal. A, 2013, 459, 65. https://doi.org/10.1016/j.apcata.2013.04.010 [21] Hu A., Yee G., Lin W.: J. Am. Chem. Soc., 2005,127, 12486. https://doi.org/10.1021/ja0538810 [22] Wang P., Kong A., Wang W. et al.: Catal. Lett., 2010, 135, 159. https://doi.org/10.1007/s10562-010-0271-x [23] Gawande M., Rathi A., Nogueira I. et al.: Green Chem., 2013, 15, 1226. https://doi.org/10.1039/c3gc40375k [24] Atashkar B., Rostami A., Tahmasbi B.: Catal. Sci. Tech., 2013, 3, 2140. https://doi.org/10.1039/c3cy00190c [25] Alizadeh A., Khodaei M., Beygzadeh M. et al.: Bull. Korean Chem. Soc., 2012, 33, 2546. https://doi.org/10.5012/bkcs.2012.33.8.2546 [26] Cheng L., Li H., Sun J. et al.: Bioorg. Med. Chem., 2010, 18, 4606. https://doi.org/10.1016/j.bmc.2010.05.034 [27] Bhatt R.: J. Global Pharma. Tech., 2010, 2, 110. [28] Kappe C., Shishkin O., Uraya G., Verdinoa P.: Tetrahedron, 2000, 56, 1859. https://doi.org/10.1016/S0040-4020(00)00116-2 [29] Amrutkar S., Bhagat U., Pargharmol P. et al.: Int. Pharm. Pharm. Sci.ence 2010, 2, 84. [30] Atwal K., Swanson B., Unger S. et al.: J. Med. Chem. 1991, 34, 2248. https://doi.org/10.1021/jm00106a048 [31] Kudo N., Futura S.: Chem. Pharma. Bull., 1999, 47, 857. https://doi.org/10.1248/cpb.47.857 [32] Bonsei M., Loizzo M., Statti G. et al.: Bioorg. Med. Chem. Lett., 2010, 20, 1990. https://doi.org/10.1016/j.bmcl.2010.01.113 [33] Yar M., Abdullah M., Majeed J.: World Acad. Sci. Eng. Tech., 2006, **55**, 593. [34] Siddiqui S., Azam A.: Med. Chem. Res., 2014, 23, 2976. https://doi.org/10.1007/s00044-013-0877-9 [35] Kanagarajan V., Thanusu J., Gopalakrishna M.: Eur. J. Med. Chem., 2010, 45, 1583. https://doi.org/10.1016/j.ejmech.2009.12.068 [36] Thanh N., Mai N.: Carbohyd. Res., 2009, 344, 2399. https://doi.org/10.1016/j.carres.2009.09.002 [37] Bhat A., Arshad M., Lee E. et al.: Chem. Biodiver., 2013, 10, 2267. https://doi.org/10.1002/cbdv.201300009 [38] Hasan A., Khaleeq M., Raizi U.: J. Org. Chem., 2010, 22, 5581.

[39] Butta R., Donthamsetty S., Adivireddy P., Venkatapuram P.: J.
Heterocycl. Chem., 2017, 54, 524. https://doi.org/10.1002/jhet.2615
[40] Kanagarajan V., How G. A., Siu C. N., Gopalkrishnan M.: J.
Heterocycl. Chem., 2013, 50, 396. https://doi.org/10.1002/jhet.1046.

[41] Singh J., Dulawat M., Jaitawat N. *et al.*: Ind. J. Chem., 2012, 51B, 1623.
[42] Varga L., Nagy T., Kovesdi I. *et al.*: Tetrahedron, 2003, 59, 655. https://doi.org/10.1016/S0040-4020(02)01560-0
[43] Ghoneim A., El-Farargy A.: Org. Chem Curr. Res., 2016, 5, 1. https://doi.org/10.4172/2161-0401.1000159
[44] Jetti S., Verma D., Jain S.: ISRN Org. Chem., 2012. https://doi.org/10.5402/2012/480989
[45] Fazaeli R., Aliyan H., Mallakpour S. *et al.*: Chin. J. Catal., 2011, 32, 582. https://doi.org/10.1016/S1872-2067(10)60203-3
[46] Fazaeli R., Aliyan H., Bordbar M., Mohammadi E.: The Open Catal. J., 2010, 3, 79. https://doi.org/10.2174/1876214X01003010079
[47] Aliyan H., Fazaeli R., Tajsaeed N.: Iranian J. Catal. 2013, 3, 99.

[48] Maleki B., Azarifar D., Moghaddam A. et al.: J. Serb. Chem.

Soc., 2009, 74, 1371. https://doi.org/10.2298/JSC0912371M

[49] Pinto D., Silva A., Ĉavaleiro J., Elguero J.: Eur. J. Org. Chem.,

2003, **4**, 747. https://doi.org/10.1002/ejoc.200390117

[50] Kavitha N., Divekar K., Priyadarshani B., Gajanan S.,

Manjunath M.: Der Pharma Chemica, 2011, 3, 55.

Received: December 06, 2017 / Revised: January 10, 2018 / Accepted: March 30, 2018

ГШЕРАКТИВНИЙ МАГНІТО-СЕПАРАБЕЛЬНИЙ НАНО-КАТАЛІЗАТОР MgFe₂O₄ ДЛЯ СИНТЕЗУ П'ЯТИ- ТА ШЕСТИЧЛЕННИХ ГЕТЕРОЦИКЛІЧНИХ СПОЛУК

Анотація. З використанням чистого нітрату заліза та карбонату магнію методом горіння синтезовано наночастки фериту MgFe₂O₄. Виявлено, що магніто-сепарабельний MgFe₂O₄ є гіперактивним каталізатором для синтезу широкого спектру біологічно активних п'яти та шестичленних гетероциклічних компонентів за умов дефлегмації. Встановлено, що вихід сполук є високим, а час реакції найменшим у порівнянні з літературними даними. Сильний електронний натяг Fe^{3+} відповідає за гіперактивність каталізатора, що було доведено заміщенням Fe^{3+} іншими тривалентними йонами металів. Показано, що заміна Mg^{2+} негативно впливає на каталітичну активність. За допомогою розробленої методики синтезовано ряд заміщених моно/біспіримідинів, піримідин-2ол, піримідин-2-тіол, піразоли та ізоксазоли внаслідок конденсації монохалконів/1,4-бісхалконів з різними біснуклеофілами у присутності гетерогенних магнітних наночастинок $MgFe_2O_4$ як каталізатора. Структуру синтезованих сполук визначено за допомогою спектроскопії $\Phi yp'\epsilon, {}^{1}H, {}^{13}C$ та мас-спектроскопії. Каталізатор можна легко видалити з реакційної суміші за допомогою простого зовнішнього магніту. Показано можливість відновлення та повторного використання наночастинок фериту без помітної зміни в активності навіть після п'яти циклів. Проведено аналіз наночастинок за допомогою рентгено-дифракційного аналізу, трансмісійної електронної та ІЧ-спектроскопії.

Ключові слова: магнітні наночастки MgFe₂O₄, монохалкони, 1,4-бісхалкони, піримідин, піразолін, ізоксазолін, зовнішнє магнітне відділення, гетерогенний відновлювальний каталізатор.