High performance of ultrasonic-assisted synthesis of two spherical polymers for enantioselective catalysis

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A B S T R A C T

Chiral polymers have aroused great attention in among chiral supramolecular materials based on their features. Herein, for the first time, the synthesis of chiral polymeric composites (CMNPs/1,4-Zbtb & 1,3-Zbtb) have been reported with entrapment through three strategies: ultrasonic irradiation, solvothermal, and mechanical stirring. According to the obtained results, it is found that ultrasound-assisted synthesis can be considered as an inexpensive and efficient method than the others, from the point of view of energy and time consuming. In this strategy, encapsulation of chiral magnetic nanoparticles (CMNPs) by using tetrazole-based polymers (Zbtbs) happens, in-situ. These chiral sphere-like inorganic–organic polymers can be considered as core and shell composites with catalytic activity due to their acidic (semi unsaturated 2za: open metal sites) and basic (abundant basic nitrogens) centers. In these structures, the unprecedented chirality induction can happen from the core to shell by non-covalent interaction, easily. They could catalyze symmetric oxidation and asymmetric Henry condensation to give chiral β-nitroalkanol. Circular dichroism and chiral gas chromatography were used to characterize the produced enantiomers. These chiral polymeric materials can be considered as unique acid-base bifunctional catalysts with efficient properties such as high stability, enantiomeric excess, enantioselectivity to the main product, and protecting from CMNPs leaching.

1. Introduction

The existence of chirality in living systems and material science is one of the reasons for research in this field [1–6]. Considering the numerous applications of chiral materials such as pharmacy, magnetism, enantioselective separation, and nonlinear optics, exploring their potential for asymmetric catalysis have been also developed. These abilities will surly motivate the researchers to devote much research effort on synthesis [7–11].

Generally, producing chiral materials especially at the supramolecular level can happen by enantiopure reagent [12,13] or spontaneous resolution [14–18]. In contrast to the above-mentioned strategies, using chiral external agent is a fast, low-cost and controllable pathway for chiralization of supramolecular especially metal–organic hybrid materials and chiral coordination polymers (CCPs) [19–24]. Until now, a lot of chiral catalysts have been successfully developed [25–27] such as polymer-supported chiral catalysts but lack of protection of chiral species is one of the disadvantages that it should be regarded. Since there are weaknesses in these catalytic systems such as leaching and decreasing of catalysts activity than their homogeneous counterparts. There is a method to solve problems: encapsulation strategy. This approach can be proposed and designed due to the production of the core–shell polymeric capsules [28]. Usually, this method entails the following requirements: use of a stabilizing agent (e.g. surfactants) [29,30] or functional linker in host synthesis [31], however these factors are not always possible to use. Among other protective agents as host, infinite coordination polymers (ICPs) can be good options due to their dependent properties [32–37]. Mirkin’s groups, Dong, and Wang are pioneers of ICPs development by using different bifunctional ligands [32]. In the following years, the synthesis of homochiral ICPs was investigated by using chiral salen-based ligand, for the first time [38]. Regarding the topic of porosity in ICPs, it should be mentioned that in contrast to porous coordination polymers as well as metal–organic frameworks, infinite coordination polymers lack acceptable porosity. Indeed, crystalline MOFs showed fairly good gas storage, separations, and heterogeneous catalysis owing to their high porosity and large surface areas. While in ICPs are amorphous supramolecular particles that self-assembled in mild conditions. So, due to the lack of growth of regular

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crystalline plates and a well-defined framework, porosity in these compounds is usually not observed or is reported in small amounts [56,57]. However, according to the nature of ICPs structure, their physical and chemical properties, they can be served as a good candidates in gas storage, mixture separations, asymmetric catalysis, and biosensing [37], but from 2006 to now, only one ICP has been reported with catalytic activity [36]. Until now, many chiral polymers have been explored, [39–42] but so far, no chiral ICP has been reported by induction method. In our previous work, we reported two strong tetrazolyl infinite coordination polymers by using 1,3-btmbenzene and 1,4-btmbenzene (Fig. 1) with hollow sphere morphology through three methods including solvothermal, ultrasonic irradiation, and mechanical stirring [43–45]. Herein, for the first time, we prepared two chiro-magnetic nanoparticles/polymer composites (CMNPs/Zbtb) with the assistance of ultrasonic irradiation. Among the above-mentioned synthetic methods for the preparation of chiral magnetic capsules, we chose the sonochemical method as a simple, fast, environmental, and low-cost strategy [46,47]. Ultrasound (US) has an important effect on the synthesis of our spherical polymers because we observed that in the other methods, no core–shell polymer was obtained through the mixing of Zn, btb, and CMNPs. From past investigations to date, ultrasonic irradiation shows a vital role in the synthesis of metal–organic polymers and frameworks [48–50]. In general, several methods have been introduced to perform chemical reactions. Most of these methods are based on heating the precursors to reach the desired product. Although, some of them have problems such as long reaction time, non-satisfactory yields, more solvent, toxic/costly reagents, and high temperatures [51,52]. It seems that ultrasonic as a non-conventional energy source can be a good alternative to conventional energy sources [53]. Moreover, ultrasound waves can provide the conditions for achieving smaller sizes of metal–organic polymers and frameworks in milder conditions without the need for high temperature and long time. So far, ultrasound as a major agent in synthesis has been used in the creation of many materials due to reducing time, producing higher surface area, crystallinity, and pore volume [54,55].

To explain the effect of ultrasound waves on the synthesis of metal–organic supramolecular, we should mention that the US can affect in the heterogeneous systems, chemically, and mechanically. Chemical and mechanical effects are related to the presence of high temperature and strong shear gradients, respectively. The localized hot spots with exceedingly high transient temperatures promote chemical reactions and the formation of nano-sized structures as results of acoustic cavitation within collapsing bubbles. In other words, cavitation (bubbles formation, growth and collapsing) in ultrasonic bath, can be considered as the main factor for these chemical and physical effects, and acoustic cavitation is also responsible for promoting organic reactions [53]. During this mechanism (ultrasonic irradiation) the migration of nanoparticles and self-assembly of metal–organic polymer are accelerated. It should also be noted that the consequences of the cavitation are many times more common in heterogeneous systems than homogeneous ones [56,57]. However, the number of published articles about preparation of polymers by ultrasonic irradiation is not low but they are limited in the synthesis of composites. PTA-MIL-53(Fe) [58], Fe3O4@TMU-17-NH2 [59], and Lipase@ZIF-8 by Rathod [60] are some examples of ultrasound irradiated composites. Recently, we also reported a chiral magnetic MOF as a basic framework that was synthesized via the sonochemical method [61].

So, we aimed to use the present stable tetrazole-based polymers (1,3-Zbtb & 1,4-Zbtb as the shell) as nitrogen-rich conjugated materials for heating the precursors to reach the desired product. Although, some of them have problems such as long reaction time, non-satisfactory yields, more solvent, toxic/costly reagents, and high temperatures [51,52]. It seems that ultrasonic as a non-conventional energy source can be a good alternative to conventional energy sources [53]. Moreover, ultrasound waves can provide the conditions for achieving smaller sizes of metal–organic polymers and frameworks in milder conditions without the need for high temperature and long time. So far, ultrasound as a major agent in synthesis has been used in the creation of many materials due to reducing time, producing higher surface area, crystallinity, and pore volume [54,55].

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So, we aimed to use the present stable tetrazole-based polymers (1,3-Zbtb & 1,4-Zbtb as the shell) as nitrogen-rich conjugated materials for
the synthesis of stable chiral catalysts and asymmetric catalysis owing to their features such as polarity, basicity, and large cavity. They have stable catalytic centers in their structures and, in fact, they are not an inert material. To chiralize active surfaces of these polymers, we have also chosen CMNPs as the core, [62,63] that are prepared by using l-(-)-tartaric acid (chiral carboxylic acid) stabilized nano-magnetite with high chirality induction[64]. As above mentioned, three synthetic methods were employed to synthesize these chiral magnetic materials by encapsulation of CMNPs. The results showed that the best method is using ultrasonic irradiation. However, in several reports, the effect of ultrasonic in encapsulation of various molecules has been shown and proved [48,58,65,66]. In fact, carrying out reactions in ultrasonic-bath not only leads to the dispersion of magnetic NPs and especially magnetic NPs, but also causes accelerates binding of metal to organic linkers around NPs. Therefore, we observed that in the absence of ultrasonic irradiation, functionalized-Fe3O4 nanoparticles definitely cannot well disperse well and the majority of them are located on the outer surface of hollow spherical polymers. The produced catalysts by sonochemical synthesis have smaller dimension and higher active surface than two other synthetic methods. So, these features can induce high catalytic performance to chiral ICPs and show the effect of ultrasonic irradiation on the structure, respectively. Fig. 2 shows the schematic diagram of the preparation process for CMNPs/1,4-Zbtb as the main catalyst. Since, it showed better results than CMNPs/1,3-Zbtb, preserving its structural integrity and catalytic activity (synthesis of CMNPs/1,3-Zbtb similar to CMNPs/1,4-Zbtb). Apart from the presence of unsaturated Zn atoms at the shell that can act as active acidic centers in the oxidation, nitroaldehyde reaction (as a type of C=C bond-forming) reaction can be proceeded by Lewis-acid or base [67–71]. To the best of our knowledge, this is the first example of chiral ICP in asymmetric catalysis with high enantiomeric excess [72,73] by chiral agent protection.

2. Experimental section

2.1. Materials and instrumentation

Materials: l-(-)-Tartaric acid (2R,3R)(-)-tartaric acid, >99%, Merck), iron(III) chloride hexahydrate (FeCl3·6H2O, 98% Fluka), diethylene glycol (99%, Merck), 1,4-phenylenediacetonitrile and 1,3-phenylenediacetonitrile were purchased from Sigma–Aldrich (Milwaukee, WI, USA). Sodium azide and zinc bromide were commercially obtainable from Merck Company (Darmstadt, Germany) and other reagents and solvents were purchased and used without further purification from commercial suppliers (Sigma Aldrich, Alfa Aesar, TCI, and others).

IR spectra were recorded using a Nicolet 100 Fourier Transform IR spectrometer in the range of 500–4000 cm⁻¹ using the KBr disk technique. The inductively coupled plasma (ICP) analyses were achieved on a Varian ICP-OES VISTA-PRO CCD instrument. Melting points were measured on an Electrothermal 9100 apparatus. UV–vis absorption spectra were obtained by UV–vis Perkin Elmer Lambda 25 spectrophotometer. CD spectra were recorded on a Jasco J-715 spectropolarimeter (Japan). Gas chromatography (GC) runs were performed on an Echrow GC A90 gas chromatograph. The samples were also characterized by using a field emission scanning electron microscope (FE-SEM) TESCAN MIRA with gold coating. TEM (transmission electron microscopy) analysis was performed on a JEOL 2100 electron microscope at an operating voltage of 200 kV. Powder X-ray diffraction (PXRD) measurements were performed using a Phillips Xpert diffractometer with monochromated Cu-Kα (λ = 1.54056 Å) radiation. The contact angle was measured by using an OCA 15 plus apparatus. Magnetic measurements were performed with a Quantum Design vibrating sample magnetometer (VSM) at room temperature in an applied magnetic field sweep. Ultrasonic generation was carried out in an ultrasonic bath SONICA-2200 EP (frequency of 40 kHz). The sonicator used in this study was a SONICA-2200 EP with adjustable power output (maximum 305 W at 40/60 kHz). A horn-type tube Pyrex reactor was custom-made and fitted to the sonicator bath.

3. Preparation of CMNPs/1,3-Zbtb & CMNPs/1,4-Zbtb composites via solvo thermal strategy.

The CMNPs (as reported in S1) (0.04 mmol) were well dispersed (10 mL DMF) by ultrasonic irradiation. Then the solution of 1,4-btmbenzene ligand (1 mmol) or (1,3-btmbenzene, S2) in DMF (15 mL) and methanolic solution (5 mL) of Zn(NO3)2·6H2O (296 mg, 1 mmol) was added to dispersed nanoparticles. This mixture was transferred to a 30 mL Teflon-lined stainless-steel vessel, sealed, and heated at 120 °C for 12 h.

3.1. Preparation of CMNPs/1,3-Zbtb & CMNPs/1,4-Zbtb composites via mechanical stirring method

The CMNPs (0.04 mmol) were fully dispersed in DMF (10 mL) by ultrasonic irradiation. Then the solution of 1,4-btmbenzene ligand (1 mmol) or (1,3-btmbenzene, S2) in DMF (15 mL) was added to it. Then, during stirring by a mechanical stirrer, a methanolic solution (5 mL) of Zn(NO3)2·6H2O (296 mg, 1 mmol) was added dropwise to them. After stirring for one hour, the produced product was separated and washed with methanol and DMF.

3.2. Preparation of CMNPs/1,3-Zbtb & CMNPs/1,4-Zbtb composites by using of ultrasonic irradiation

The CMNPs (0.04 mmol) was dispersed well in DMF (10 mL) by use of ultrasonic irradiations. Then the solution of 1,4-btmbenzene ligand (1 mmol) in DMF (15 mL) was added to dispersed nanoparticles and then both of them were sonicated for specific times (30–45 min) at room temperature. After homogenization, a methanolic solution (5 mL) of Zn(NO3)2·6H2O (296 mg, 1 mmol) was added dropwise to the mixture of CMNPs and ligand in the ultrasonic bath. Subsequently, the light-brown micro sphere-like polymers as shell with chiral magnetic nanoparticles within their core were generated. Finally, they were purified by magnetic separation and washed with DMF through filtration-redispersion cycles.IR data (ν/cm⁻¹):427 (s), 657 (w), 777 (w), 1106 (w), 1430 (m), 1658 (s), 1732 (s), 2349 (w), 2830 (w), 2935 (w), 3428 (m). Preparation of CMNPs/1,3-Zbtb was similar to CMNPs/1,4-Zbtb without any change in ratios. IR data (ν/cm⁻¹):446 (w), 690 (w), 765 (m), 1108 (m), 1434 (m), 1653 (s), 1730 (s), 2351 (w), 2810 (m), 2920 (m) and 3430 (m).

3.3. Typical reaction procedure of Henry reaction

In a typical reaction, chiral catalyst (CMNPs/1,3-Zbtb & CMNPs/1,4-Zbtb) (0.05 g) was added to a solution of MeOH (5 mL) in 5 mL screw-capped vial. Afterwards, benzaldehyde (2 mmol) and nitromethane (5 mmol) were added into above solution. The reaction mixture was allowed to stir at 70 °C. After that, upon filtration by an external magnet, the catalyst was separated and washed with MeOH. The solution was collected and analyzed by chiral GC.

3.4. Recycling of catalyst

After the first step of the catalytic cycle, the heterogeneous catalyst was separated by external magnet, rinsed several times by MeOH and dried for 5 h in oven (80 °C). Then, the recovered catalyst was reused in the subsequent recycling experiments under identical reaction conditions.

3.5. Typical reaction procedure of benzyl alcohol oxidation to benzoaldehyde

In a typical procedure, BzOH (2 mmol) and deionized water (5 mL),
and were added to a 25 mL two-neck flask containing magnetic catalyst (0.025 g). This system was equipped with a condenser. The reaction mixture was vigorously stirred at 85 °C for 5 h and H₂O₂ (5 mmol, 30 wt % in water) was added dropwise in three steps. Then, the solid catalyst was separated from the reaction mixture by an external magnet. The product was extracted by ethyl acetate (5 mL), washed with saturated sodium thiosulfate solution (2 × 3 mL) and analyzed by a gas chromatography system.

4. Results and discussion

4.1. Characterization of chiral magnetic composites (CMNPs/Zbtbs)

In this research work, the catalytic results showed that the efficiency of the optimized chiral catalyst through ultrasonic irradiation is very sensible. In the following, the further observations are used to support this claim. The generation of our novel chiral polymeric composites (CMNPs/Zbtbs) is done in two steps: in first step, tartaric acid as chiral inducer was stabilized on Fe₃O₄ nanoparticle. In next step, Fe₃O₄-tartaric acid nanoparticles were dispersed in solution by the help of ultrasonic irradiations, and eventually they were encapsulated within Zbtb colloidal particles through in situ method, in a few minutes (Fig. 2). We previously characterized our sphere-like polymer structures with different methods. On the other hand, our purpose in this report was the introduction of the best chiral Lewis base to the promotion of asymmetric nitroaldol reaction. As we discussed and showed in the following, CMNPs/1,4-Zbtb worked better than compared to CMNPs/1,3-Zbtb. So, our optimized catalyst was CMNPs/1,4-Zbtb and specific characterization about it was carried out. It should also be noted that due to the high structural similarities of the two ICPs, almost identical results in early characterization are not unexpected, although full characterization about Fe₃O₄/1,3-Zbtb was carried out in our previous work [43–45]. Powder x-ray diffraction (PXRD) pattern was used first as direct evidence to confirm the formation of CMNPs/1,4-Zbtb. For this purpose, powders of 1,4-Zbtb, CMNPs/1,4-Zbtb, and CMNPs nanoparticles are deposited in the hollow of a silicon zero-background plate 0.2 mm deep (PW1812, Philips company). The diffractometer of this apparatus equipped with an X-ray tube (CuKα, λ = 1.5418 Å), and the generator was set at 40 kV and 40 mA (2θ range: from ~10 to 80°). The comparison of CMNPs/1,4-Zbtb with hallow 1,4-Zbtb revealed that polymer preserved its amorphism and several reflections appeared at 2θ = 30.3°, 35.4°, 43.2°, 53.5°, 57.2° and 62.9° corresponding to Fe₃O₄ in XRD-pattern matched well with the presence of Fe₃O₄ (Fig. 3a) [74]. PXRD pattern of CMNPs showed that intensity of Fe₃O₄ peaks decreases due to functionalization of Fe₃O₄ by tartaric acid. Also, in comparison of XRD-patterns of the as-prepared composites by all three methods, no noticeable differences were observed (Fig. S9).

About 1,3-Zbtb and CMNPs/1,3-Zbtb, similar to 1,4-Zbtb and CMNPs/1,4-Zbtb, powder x-ray diffraction analysis was also performed to prove the existence of CMNPs in composite form with polymeric
capsules (Figs. S9 and 3a) [43]. Despite all the knowledge about poor porosity in ICPs, we examined the uptake of N\textsubscript{2} gas for our desired composition. As expected, we obtained a very low N\textsubscript{2} BET surface area (close to zero) which proved the non-porosity of our structure. The 1,4-Zbtb modification was inspected by Fourier transform infrared spectroscopy (FTIR) (Fig. 3b). Comparing FTIR spectra both of hollow 1,4-Zbtb capsule and CMNPs/1,4-Zbtb revealed the presence of the characteristic C=O stretching band of tart at 1732 cm\textsuperscript{-1} corresponding to chemisorption of tartaric acid on the surface of Fe\textsubscript{3}O\textsubscript{4}[75] in CMNPs/1,4-Zbtb. Carbon–oxygen stretching of tartaric acid carboxylate groups

Fig. 4. SEM images of the synthesized CMNPs/1,4-Zbtb by (a) ultrasonic irradiation, (b) solvothermal method, and (c) mechanical stirring.
Optimization of Henry reaction conditions.

| Entry | Catalyst           | Solvent | Temperature(°C) | Time (h) | Conversion (%) | Selectivity (%) | Ee(%) |
|-------|--------------------|---------|----------------|----------|----------------|----------------|-------|
| 1     | CMNPs/1,4-Zbtb     | MeOH    | 70             | 2        | 15             | 80             | 23(S) |
| 2     | CMNPs/1,4-Zbtb     | MeOH    | 70             | 5        | 26             | 78             | 51(S) |
| 3     | CMNPs/1,4-Zbtb     | MeOH    | 70             | 10       | 58             | 82             | 78(R) |
| 4     | CMNPs/1,4-Zbtb     | MeOH    | 70             | 12       | 85             | 81             | 90(R) |
| 5     | CMNPs/1,4-Zbtb     | MeOH    | 40             | 12       | 20             | 72             | 46(S) |
| 6     | CMNPs/1,4-Zbtb     | Water   | 70             | 12       | 19             | 42             | 39(S) |
| 7     | CMNPs/1,4-Zbtb     | EtOH    | 70             | 12       | 9              | 49             | 41(R) |
| 8     | 1,4-Zbtb           | MeOH    | 70             | 12       | 80             | 78             | –     |
| 9     | CMNPs/1,3-Zbtb     | MeOH    | 70             | 10       | 51             | 73             | 70(R) |
| 10    | CMNPs/1,3-Zbtb     | MeOH    | 70             | 12       | 75             | 77             | 85(R) |
| 11    | CMNPs/1,3-Zbtb     | Water   | 70             | 12       | 15             | 45             | 40(R) |
| 12    | CMNPs/1,3-Zbtb     | EtOH    | 70             | 12       | 7              | 35             | 28(S) |
| 13    | 1,3-Zbtb           | MeOH    | 70             | 12       | 69             | 75             | –     |
| 14    | CMNPs/1,4-Zbtb     | MeOH    | 70             | 12       | 35             | 69             | 61    |
| 15    | CMNPs/1,4-Zbtb     | MeOH    | 70             | 12       | 63             | 78             | 79    |

* unless otherwise specified, all reactions were conducted with 5 mmol of nitromethane, 2 mmol of benzaldehyde, 5 mL of solvent, and 0.05 g of catalyst in the ultrasonic bath.

† Related to nitroalcohol product.

‡ Enantiomeric excess of major isomer was determined by GC analysis using a chiral column.

§ 0.03 g.

‖ prepared by solvothermal method.

¶ prepared by mechanical stirring.

appeared at 1550 cm⁻¹ and 1422 cm⁻¹ indicate the asymmetric νas(CO₂) and symmetric νs(CO₂), respectively [76]. Fe-O vibration represents broad bands at 668, 630, 442 cm⁻¹. Besides, the presence of two distinct, typically narrow peaks at 1652 and 1432 cm⁻¹ associated with deprotonation of tetrazolate groups and absence of broadband from 2200 to 2600 cm⁻¹ confirmed the formation of polymer and coordination of N-H of tetrazole groups as described previously. Moreover, our composite exhibits aromatic signals associated with the benzene ring. Some factors are responsible for the broad characteristic band at 3441 cm⁻¹ such as: OH, stretching vibration owing to the Fe-OH groups, the carboxylic groups of tartaric acid, and adsorbed water [77].

Electrochemical studies were investigated under optimum conditions (solvent, temperature, and catalyst amount) (Table 1). Before performing the catalytic tests, the stability of CMNPs/1,4-Zbtb in various solvents such as MeOH, EtOH and H₂O was monitored by PXRD (Fig. S2). As shown, these new composites are stable under the essayed conditions for at least 24 h (Fig. S2).

4.2. Catalytic activity for asymmetric Henry reaction

As mentioned above, catalysts with rich basicity can be the efficient catalysts for Henry reaction. So, the activity of CMNPs/1,4-Zbtb and CMNPs/1,3-Zbtb composites was investigated as basic catalysts for this kind of aldol condensation through large numbers of tetrazole nitrogen atoms on the surface of spherical composites. The catalytic experiments were investigated under optimum conditions (solvent, temperature, and catalyst amount) (Table 1). Before performing the catalytic tests, the stability of CMNPs/1,4-Zbtb in various solvents such as MeOH, EtOH and H₂O was monitored by PXRD (Fig. S2). As shown, these new composites are stable under the essayed conditions for at least 24 h (Fig. S2). To investigate effect of solvent, we examined this transformation in several solvents. EtOH and water gave only 13% and 19% of total yield, respectively. Whereas, a significant acceleration in the rate of reaction and considerable yield were achieved in MeOH, so it was used as an optimized solvent. The influence of temperature was also investigated on the conversion of reaction as main parameter. At first,
temperature 40 °C was examined in the model substrate catalysis that the conversion 20% was achieved (entry 5). Increasing the temperature from 40 to 70 °C could affect on conversion, from 20 to 85% for 12 h. The comparison of composites demonstrated that CMNPs/1,4-Zbtb and CMNPs/1,3-Zbtb catalysts have ability for transformation of aldehyde to chiral alcohol with 85%, 75% conversion and 90%, 85% enantiomeric excess to R enantiomer, respectively (entry 4 and 10). Also, we investigated the hollow achiral form of our catalysts that they had potential to transform benzaldehyde to β-nitroalcohol with 80% (1,4-Zbtb) and 69% (1,3-Zbtb) conversion due to strong interaction between carbonyl groups of aldehyde and catalytic active sites of catalysts surface.

This difference may be attributed to the situation and orientation of the tetrazole groups in polymeric structures (Fig. 1). It seems that the superiority of 1,4-Zbtb to 1,3-Zbtb is due to its structure, physical-chemical properties and more accessible catalytic active centers on its surface. Therefore, it can be concluded that although these composites are almost the same with similar components, but CMNPs/1,4-Zbtb showed higher catalytic activity compared to CMNPs/1,3-Zbtb. After the confirmation of the high performance of 1,4-Zbtb in catalysis, we investigated and compared the activity of the synthesized CICPs that were prepared by three different methods. They have same PXRD patterns but different morphologies that affect on their catalytic activity. The catalytic results show: ultrasonic irradiation > mechanical stirrer > solvothermal (Table1). This point demonstrates that in ultrasonic method, there are uniform polymeric spherical particles with smooth morphology. In this case, catalytic active sites can be accessed by substrates. Mechanical stirrer and solvothermal processes can be considered as bulk syntheses. According to SEM images, the obtained compounds by these methods are not more regular and do not have high surface area than ultrasonic-assisted synthesis. In fact, using autoclave and synthesis in high pressure and temperature are important agents to produce large capsules with fewer active sites on their surface (35% conversion, 69% ee). Or mechanical stirrer caused the agglomeration of spheres and reducing their active catalytic centers (63% conversion, 78% ee), too. So, it has been concluded that produced CICPs by ultrasonic can show higher activity in basic nitroaldol reaction because a) and they have small size, so rich accessible nitrogen groups as active sites and b) spheres agglomeration is likely much less due to nanoparticles dispersion (owing to presence of ultrasound waves).

About enantiomeric excesses, certainly, in the obtained catalysts by mechanical and solvothermal methods, a high percentage of CMNPs were not encapsulated and probably placed on the outer surface of polymeric particles. To demonstrate the high ability of our chiral catalyst, we compared the activity of CMNPs/1,4-Zbtb with blank sample without any catalyst, 1,4-Zbtb, CMNPs and Fe₃O₄ for reaction between benzaldehyde and nitromethane, too (Fig. 7a). Given that basic groups are considered as the main driving force for this reaction so, there is no significant difference in conversions of achiral and chiral spheres as catalyst. Also, due to the influence of acidic catalyst in promotion of henry reaction, low conversions were also observed for the Fe₃O₄ NPs and CMNPs, although carboxylic groups of tartaric acid led to semi higher result. We know that the nature of substrates has important influence on yields and selectivity. Thus, to show this fact, we compared the reaction of aromatic and aliphatic aldehydes (branched and unbranched) with nitromethane to prepare the respective β-nitroalcohols (yields ranging from 54 to 89% and selectivity up to ca. 90%). Acetaldehyde as an aliphatic aldehyde demonstrated 89% conversion, 63% enantiomeric excess and 75% selectivity. In addition, in comparison to acetaldehyde and isobutyraldehyde, it can be understood that short chain without steric hindrance leads to higher yield and selectivity. Using the same reaction conditions, the obtained nitroalcohol product from the isobutyraldehyde catalysis was isolated in 54% conversion and 70% ee.

This phenomenon is due to the fact that substitution in carbonyl compounds that leads to steric crowding, increasing the activation barrier [78] and reducing the selectivity (60%) [79]. About model substrate, GC analysis with chiral SGE-CYDEX-B capillary column indicated that no further conversion had existed after 12 h and even for 36 h. The obtained results demonstrated the necessity of catalysts for achieving high conversion and enantiomeric excess. As known, the important parameter in the asymmetric catalysis is enantiomeric excess that is used to determine purity of the major chiral isomer with range of 0% to 100%. It is calculated based on following equation:

\[ \text{enantiomeric excess} = \text{ee} = \frac{A - B}{A + B} \times 100 \]

A and B are the enantiomers mass that are determined by the chiral GC data. On the other hand, for investigation of the enantiomeric nature of produced β-nitroalcohol, we utilized CD spectrum in the wavelength range 370–800 nm (Fig. 5) that showed the nature of the obtained enantiomer that was similar to the obtained chiral GC [96] data in this work and CD of R-Limonene.

Interestingly, we observed the change in the kind of enantiomeric excess, from S- to R-enantiomer. Until 5 h, S-enantiomer was the major isomer that was determined with GC but the next 5 h, R-enantiomer was the main isomer. As known, during asymmetric henry reaction, the racemic mixture of S and R enantiomers of nitroalcohol is produced. In this case, the formation rate and energy of these enantiomers are equal.
Surely, before or after racemate creation as intermediate, the growth of one enantiomer is seen further, R or S, due to the stability of intermediate and preferred transition state. So far, many examples have been reported with chiral core–shell structures that chiral shell as host entrapped inorganic material as guest. In this case, chirality can be induced from shell to core by various interactions in confined space [80,81]. In this respect, we believe that similar interaction has occurred in the opposite direction (chirality information transfer from core to shell by the same interactions). One of the works has been done in this area was about chirality activity of unprotected Au25 and Au55. Gold cluster was intrinsically chiral due to the stability in energy state. In the second step, in comparison of bare gold cluster with protected manner, it was found that protection by methyliol monolayer can enhance the degree of chirality of the metallic core. It is suggested that interaction between chiral core and achiral shell is responsible for chirality induction from metal center to ligand [82,83]. About mechanism, recently we published a work about the basic effect of nitrogen groups for promoting henry reaction [61]. Similarly, in this work it should be said that the most prevalent mechanism for Henry reaction is based on the deprotonation of nitroalkane under basic condition. In second step, C–C bond is formed upon nucleophilic addition and subsequently β-nitroalkanol is produced by means of protonation [79]. In our report for the evaluation of the mechanism and stereoselectivity, we must suppose several hypotheses, first N moieties of tetrazole would deprotonate nitromethane, so in the next step, bidentate nature of nitronate anions as hydrogen-bond acceptor led to the formation of chiral tetrazolium nitroate ion pair. Eventually, nucleophilic addition has occurred in high stereo-selectivity manner. It can be concluded that responsibility for the second sequence, is N–H⋯O as non-covalent interactions. Moreover, we suggest that other interactions like C-H⋯π has an auxiliary role in stabilizing the transition state (TS) and increasing the reaction rate that the way π system of the phenyl scaffold can activate aldehyde through C-H⋯π interaction and subsequently affect the transition state (Fig. 6). To illustrate protection power and to determine heterogeneity of our composite, the leaching test of active species into the catalytic reaction medium was carried out. After 8 h and removal of catalyst from reaction mixture, the acquired results demonstrated that catalyst was necessary to promote the reaction (Fig. 7b).

4.3. Catalytic activity for oxidation of alcohols

In the field of oxidation of alcohols to achieve carbonyl compounds, the best way is using non-precious and environmentally friendly precursors. Except for oxidant-free photocatalytic instances, it is well known that the oxidation processes can be occurred in the presence of the various oxidants in most cases normally [84]. Among them, hydrogen peroxide can be used as an ideal green oxidant due to its low cost, high content of active oxygen, non-toxic and corrosive. On the other hand, Zinc as an easily available metal is relatively inexpensive and non-toxic than precious metals (mostly Pd, Ir, Ru, and Au) which makes it as a catalyst for practical applications particularly on an industrial scale [85]. In addition to, the role of metal in oxidation [86], we can suggest that tetrazole groups of btmbenzene can act as a base. In this state, tetrazole moieties as weak bases can promote oxidation of alcohols through generating alkoxy [87]. The benzaldehyde is produced by optimized 1,4-Zbtb (prepared in the ultrasonic bath) through oxidizing benzylalcohol in the presence of H2O2. After evaluation of H2O2 amount (Fig. S6), the optimal combination is achieved by applying four equivalents of H2O2 (in three portions). This reaction (benzylalcohol to benzaldehyde) was also performed in H2O as the green solvent with low percent yield (35%). Successive adding of H2O2 improved the yield but increasing the temperature decreased it. The mechanism of the conversion of alcohols to aldehyde in the presence of H2O2 as oxidant and

Fig. 7. a) Optimization of catalyst in Henry reaction of benzaldehyde and nitromethane. b) Leaching test confirming no contribution from homogenous catalysis of active species leaching into the reaction solution.

Fig. 8. Catalytic cycles for asymmetric Henry reaction of benzaldehyde and nitromethane catalyzed by CMNPs/1,4-Zbtb (5 mol %). Reaction conditions: benzaldehyde (2 mmol), nitromethane (5 mmol) in 5 mL of MeOH at 70 °C.
Zinc(II) as catalyst has been investigated by computational methods in literatures [88]. In the transition state, O-O bond in H2O2 is weakened and coordinated to Zinc at the surface of spheres. These weak O–O bonds are replaced by strong O–H bonds that are also coordinated to Zinc atoms. The generated water molecules as by-product have interactions with aldehyde molecules by hydrogen bonding. Also, they can accelerate separation of aldehyde molecules from metals. Finally, the control experiment was investigated without catalyst that no progress was seen in the catalytic reaction. Given that the importance of asymmetric Henry reaction and its chiral product, chiral 1,3-Zbtb composite was also investigated in this kind of asymmetric catalysis but it showed lower activity than 1,4-Zbtb composite (main catalyst) so, we did not study its catalytic application in oxidation of alcohols. Although, we should remind that 1,3-Zbtb composite showed moderate to poor performance as Lewis-acid in Michael reaction [45].

4.4. Catalyst recyclability

Due to significance of the recycling and life-time of heterogeneous catalysts in practical applications, we investigated recyclability of the optimized CMNPs/1,4-Zbtb catalyst. Investigations showed high stability of catalyst with less chiral species leaching after 5 catalytic consecutive cycles. Not only no significant deactivation was observed, even after the fifth cycle but also high conversion and enantiomeric excess (ee) are observed due to high stability of catalyst along with high protection of CMNPs (Fig. 8). PXRD, SEM, VSM and ICP analysis showed that our sphere-like composite is stable chiral heterogeneous catalyst (Figs. S2, S3, S4, S5). These observations can confirm a) protection of CMNPs by stable polymeric shell and b) the presence of accessible tetrazole groups as catalytic active sites after several cycles. Since, robust shell prevents leaching of CMNPs so, enantioselectivity does not significantly decrease. In general, it can be said that the strength and capability of our polymer are at a very high level compared to other mineral and organic polymers. Also, to compare our catalyst with reported heterogeneous and homogeneous catalysts in asymmetric henry reaction, Table 2 has been presented. Although, relatively good results have been obtained by other catalysts, but the high cost of the used compounds in their synthesis is a fault. For example, Pd NPs and salen-ligands can be mentioned that they limit the use of the synthesized catalysts, in addition long time of catalytic reactions in high or very low temperature that they are main disadvantages of these kinds of catalysts [89,90].

5. Conclusion

In conclusion, we reported the first examples of the CMNPs/1,3- and 1,4-Zbtb core–shell microspheres as promising chiral magnetic spherical polymers via ultrasonic-assisted encapsulation as an effective method. In this synthetic way, chiral magnetic nanoparticles could enter into spherical polymers that due to this process, chirality induction happened from chiral magnetic core to optically inactive polymeric shell through hydrogen-bonding interactions. These compositionally modified polymers by using chiral magnetic nanoparticles showed their capability for enantioselective conversion of aldehydes to chiral alcohols. Catalytic results demonstrated that CMNPs/1,4-Zbtb is better than CMNPs/1,3-Zbtb due to the orientation of the side groups (basic centers) around the benzene ring that has a vital role to convert CMNPs/1,4-Zbtb to optimized catalyst. The other synthetic methods were examined to produce optimized CICP (CMNPs/1,4-Zbtb). The important point is that the synthesis method could directly affect on catalysis performance of polymers. The obtained samples by solvothermal and mechanical stirring did not show high catalytic activity than ultrasonic synthesis. An important property of this hollow catalytic system is high protection of chiral components without any change in chemical structure of the polymer.

With the presence of the nitrogen and unsaturated Zn atoms as basic and acidic active sites at the surface of shell, magnetic and chirality property, our sustainable composite showed ability in nitroalaldol (asymmetric) condensation and oxidative transformation of alcohol to aldehyde with high performance (mild catalytic conditions). Also, other unique advantages of our green catalyst as a low-cost chiral catalyst are high stability, reusing for five successive runs with retention of structure and morphology without obvious loss of activity. So, it is unprecedented among other chiral polymeric catalysts. Our methodology will be able to induce chirality into other hollow structures, especially metal–organic materials. Finally, given its efficiency, we highly expect that this kind of asymmetric catalytic system can have an immediate impact on the asymmetric syntheses and catalysis.

Declaraton of Competing Interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

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Appendix A. Supplementary data

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