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Nanostructured MgO-enhanced catalytic ozonation of petrochemical wastewater

Cerámica

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ABSTRACT

Petrochemical wastewater, which is a complex solution of various chemicals, can cause severe environmental problems after being disposed to environment. Here, we investigate the removal of petrochemical contaminants at different periods and pH levels using three distinct processes: conventional ozonation, adsorption and nanostructured MgO-enhanced catalytic ozonation. The latter is found to be an economical and efficient new approach to treat petroleum contaminants, outperforming the other methods. For instance, after 30 min of the treatment using the conventional ozonation, the removal percentages of benzene, toluene, ethylbenzene, and xylene are 35, 58, 81, and 92%, respectively; whereas the corresponding figures are 93.8, 83.5, 94.6, and 99.2% using the catalytic ozonation.

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Tratamiento catalítico de ozono de aguas residuales petroquímicas impulsado por nanoestructuras MgO

RESUMEN

Las aguas residuales petroquímicas, que es una solución compleja de diversos productos químicos, pueden causar graves problemas ambientales después de ser eliminadas en el medio ambiente. Aquí, investigamos la eliminación de contaminantes petroquímicos en diferentes periodos y niveles de pH utilizando tres procesos diferentes: zona convencional, adsorción y ozono catalítico mejorado por nanoestructuras MgO. Este último es visto como un nuevo enfoque económico y eficiente para el tratamiento de los contaminantes del petróleo, superando a otros métodos. Por ejemplo, después de 30 minutos de tratamiento

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utilizando ozono convencional, las tasas de eliminación de benceno, tolueno, etilbenceno y xileno son 35, 58, 81 y 92%, respectivamente; mientras que las cifras correspondientes son 93, 8, 83,5, 94,6 y 99,2% utilizando ozonación catalítica.

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Introduction

Hazardous chemicals such as benzene, toluene, ethylbenzene, 3502 and xylene, which naturally exist in crude oil and gasoline, 36 can easily be transferred into surface and groundwater due to 37 either the leakage from industrial petrochemical equipment 38 or inappropriate waste disposal [1,2]. Considering that the 39 petroleum industry consumes large volumes of water, espe-40 cially for cooling purposes, a large amount of petrochemical 41 wastewater is produced [3,4]. Therefore, the appropriate treat-42 ment of polluted water sources produced in such industries 43 is a challenge. Not to mention that the disposal of untreated 44 petrochemical wastewater can cause grave environmental 45 issues [5], expanding the presence of contaminants such as 46 phenols and toxic hydrocarbons in the environment [6]. 47

In general, wastewater from petroleum refineries contains 48 large quantities of environmentally problematic materials; 49 somewhere around 150-250 mg/L BOD (Biological Oxygen 50 Demand), 150-250 mg/L COD (Chemical Oxygen Demand) and 51 20-200 mg/L phenols, as well as 100-300 mg/L other petroleum 52 products and 5000 mg/L salts [7], disrupting habitats of human 53 and also wildlife communities [8,9]. The disposal of such 54 wastewaters might destroy a considerable number of the 55 biotic elements, leading to the gradual elimination of aquatic 56 plants and animal species. It also simplifies the food chain 57 through reducing the number and diversity of species [10]. 58 Therefore, organic and inorganic contaminants present in 59 petrochemical wastewaters must be carefully treated and 60 removed, before the wastewater can be discharged into the 61 environment [11]. A variety of methods have been sug-62 63 gested, and used, for the removal of petroleum contaminants 64 from water resources, including chemical coagulation and sedimentation, filtering, photocatalytic oxidation, and adsorp-65 tion [12]. Unfortunately, the appropriate implementation of such technologies is often very expensive, complicated and 67 time-consuming, providing various challenges particularly in 68 developing countries [13]. 69

Advanced oxidation processes (AOPs) are among the most 70 effective methods for the removal of organic contaminants. 71 These methods are based on the production of highly oxi-72 dizing free radicals such as hydroxyl radicals, capable of 73 mineralizing various toxic organic compounds [14]. AOPs are 74 operationally easy, low cost, and efficient while being able to 75 stabilize organic compounds through converting them into 76 water and carbon dioxide [15-18] often using ozone [19]. APOs 77 are suitable for treating contaminants that are resistant to 78 biological decomposition [20,21]. 79

Catalytic ozonation processes (COPs) are APOs in which solid catalysts play a role in overall ozonation process, facilitating the decomposition of ozone into free radicals. In fact, the use of catalysts may eliminate the most important limitations of simple ozonation, including the low solubility and stability of ozone in water, and its slow reaction with organic compounds [22,23]. The high efficiency, as well as low operation costs influenced by the short duration of COPs [24,25] also facilitates the implementation of these technologies, particularly, in developing countries.

Metal oxides, such as magnesium oxide (MgO) nanoparticles, have a high potential to be used in wastewater treatments, due to their possible large surface areas and the low cost of recovery and production. MgO also provides advantages of having antibacterial properties and high adsorption and/or photocatalysis performance for toxic compounds [26–28].

Various studies have been conducted on using MgO in COPs for the removal of toluene [29], textile dyes [30] and humic acid [31]. Moosavi et al. [32] studied the effect of MgOenhanced COP on the removal of Reactive Red198, indicating that the removal efficiency improves with the increase of the pH value, due to the increased rate of ozone decomposition into free radicals at higher pH values. The decomposition of the dye molecules was attributed to an indirect oxidation process brought about by the active radicals. The present research intends to study the feasibility of using MgO nanoparticles in COP for the effective treating of petrochemical wastewater.

Materials and methods

Real wastewater samples were provided by a large petrochemical unit located in southern Iran. All other chemicals used in the research were of high purity and manufactured by the German Company Merck. The wastewater experiments were conducted using a laboratory scale semi-continuous ozonation reactor, shown in Fig. 1.

Before performing water treatment processes, the wastewater sample was tested to determine the values of COD, BOD, TS (Total Solid), and turbidity, as well as the amounts of benzene, toluene, ethylbenzene, and xylene. For this, 1 µL of the wastewater was injected into an Agilent's Gas Chromatography (GC) with the GC column HP-5 30m, and 0.50 µm film thickness using a flame-ionization detector. The incubator temperature was maintained at 40 °C for 10 min before the injection. The extraction was performed at $120^{\circ}C$ and 500 rpm. The initial furnace temperature was kept at 40°C for 2 min, gradually raised to 80°C at the rate of 20°C/min, and kept at this temperature for 1 min. The temperatures of the injector and detector were set at 250 and 300°C, respectively, and nitrogen gas flow rate was maintained constant at 3 mL/min. The amount of each component was calculated through comparing the diagram provided by the GC machine with that obtained by injecting standard samples into the GC

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Fig. 1 – Schematic representation of the reactor used for the petrochemical wasterwater treatment in the presence of MgO nanoparticles.

Table 1 – Characteristics of the petrochemical wastewater collected from an Iranian petrochemical industry.					
Value	Parameter				
7.5	рН				
1355 mg/l	COD				
395 mg/l	BOD5				
3.43	COD/BOD5				
2464 mg/l	TS				
780 NTU	Turbidity				

equipment. Table 1 represents the characteristics of the petro chemical wastewater used in this study.

133 Preparation of MgO nanocatalysts

MgO nanoparticles, used as nanocatalysts in the wastewa-134 ter treatment, were prepared by the calcination approach. 135 For this, Mg (NO₃)₂6H₂O was first dried at 100 °C for 8 h. 136 Then, the dried powder was calcined at 500°C for 2h in 137 an electrical furnace to form MgO nanoparticles. The mor-138 phology of nanoparticles was studied using scanning (SEM), 139 and transmission (TEM) electron Microscopy. A Philips X-140 ray diffractometer (XRD) with Cu-K α radiation (1.54 Å) was 141 employed for the structural study. 142

143 Catalytic ozonation process (COP)

A semi-continuous cylindrical reactor with the internal volume of 1500 mL was assembled for the catalytic ozonation of the wastewater (Fig. 1). Using this reactor, ozone could continuously be bubbled into the wastewater. The off-gas containing the excess ozone could then be directed from the reactor to gas scrubbers containing 20% potassium iodide where the excess ozone could be decomposed. The catalytic ozonation experiments were conducted at room temperature using 50 mL of the wastewater samples mixed with 450 mL double distilled water and 0.3 mg MgO nanoparticles. The pH was adjusted at various values of 3, 5, 7 and 9 employing 0.1 M solutions of HCl or NaOH. For a typical experiment, the solution was transported into the glass reactor shown in Fig. 1, connected to a DONALIO3 ozone generator with an ozone production capacity of 0.2 g/h, and treated for 10, 20, and 30 min. After specific periods, the quantity of remaining benzene, toluene, ethylbenzene, and xylene present in the wastewater samples were measured by injecting 1 μ L of the sample into the GC equipment.

Simple ozonation process (SOP) and adsorption process (AP)

In addition to the COP, SOP and AP were also evaluated using petrochemical wastewater samples in order to distinguish the effect of various phenomena which could be involved in the removal, or the reduction, of contaminants present in the wastewater. To perform the SOP, 50 mL of the wastewater was mixed with 450 mL of double distilled water at different pH values of 3, 5, 7 and 9. For this, the pH of wastewater was adjusted with solutions containing 0.1 N of HCl or NaOH. Then, the ozonation process was carried out at reaction durations of 10, 20, and 30 min to evaluate the influence of time on the process.

The AP was performed under the same conditions as mentioned above for SOP, with the difference that no ozone was involved, and instead, around of 0.3 mg MgO nanoparticles was added to the wastewater. Moreover, the solution was aerated to create turbulence in order to achieve an appropriate contact between the solution and MgO nanoparticles.

At the end of both processes, the samples were rested for 5 min, and 5 mL aliquots were collected from the middle level of the reactor, from which $1 \mu \text{L}$ samples were injected into the GC instrument for determining the quantities of contaminants remaining in the solution.

Results and Discussion

Characterization of MgO nanocatalysts

The SEM surface morphology of MgO nanoparticles is shown in Fig. 2a, from which the presence of spherical shaped and agglomerated particles is evident. TEM image of the material (Fig. 2b) clearly indicates that individual MgO nanoparticles have an average dimension of around 30 nm. X-ray diffraction pattern of MgO nanoparticles, presented in Fig. 2c, indicate the reflections at $2\theta = 37^{\circ}$, 43° , 62° , 74° , and 78° corresponding to (111), (200), (220), (311), and (222) characteristic XRD peaks of MgO with a cubic structure (JCPDS 01-075-1525), respectively. Scherrer equation can be employed to evaluate the size of crystals in ceramic materials [33–35]. The mean crystallite size of MgO nanoparticles could be calculated using this approach, based on the most intense (002) XRD peak observed in Fig. 2c, and found to be around 21 nm. This is in agreement with the nanostructured nature of the material.

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Fig. 2 - (a) SEM, (b) TEM, and (c) XRD analysis of MgO nanoparticles synthesized using the calcination method.

It should be mentioned that there are several techniques 203 to prepare ceramic nanoparticles including hydrothermal 204 methods [36,37], sol-gel [38,39], spark discharge [40], elec-205 trochemical discharge machining [41], laser ablation [43,43], 206 ball milling [44], aqueous wet chemical method [45], com-207 bustion synthesis [46-49] and molten salt-based techniques 208 [50-53]. Comparing to these techniques, the simple calcination 209 method used in this study for the preparation of MgO nanopar-210 ticles (reaction (1)) may benefit from relative simplicity, low 211 cost and minimal requirement for complex equipment. 212 $Mg(NO_3)_2 = MgO + 2NO_2(g) + 1/2O_2(g) \Delta G^{\circ}_{500 \circ C} = -80 \text{ kJ} (1)$ 213



Fig. 3 - The particle size distribution histogram of MgO nanoparticles.

It should be mentioned that NO₂ produced as the byproduct of the reaction (1) can be used in the manufacturing of nitric acid by treating with water and air [54–56], improving the economic performance of the process. It is worth noticing that the demand for nitric acid is considerably increasing due to its wide range of applications, including in the processing of fertiliser, mining explosives and gold extraction [57]. The surface area of the MgO nanoparticles produced in our study was measured by N2 adsorption/desorption measurements (TriStar II 3020 Micromeritics) and found to be 221 m^2/g , with BJH Adsorption cumulative pores volume of 0.85 cm3/g, and BJH adsorption average pore diameter of 27.4 nm.

Fig. 3 shows the particle size distribution histogram of MgO nanoparticles calculated from TEM analysis, which was built up by counting more than 100 particles. From this, an average diameter of 22.6 nm can be obtained. We can compare this value with those presented in the literature. Wong et al. [58] prepared MgO nanoparticles employing an ultra-sonication incorporated Pechini method [59] using magnesium acetate tetrahydrate and citric acid as the raw materials. The value of average size of MgO nanoparticles produced under different conditions were measured to be around 21 -122 nm. In another work, Hajengia et al. [60] produced MgO nanoparticles with an average particle size of 16 nm by the microwave heating of a mixture comprising Mg(CH₃COO)₂·4H₂O and benzylamine to form Mg(OH)₂ which was separated, washed and calcined at 550 °C for 5 h. The particle size distribution histogram of the sample demonstrated that the most of the particles were in the range of 12–22 nm. Furthermore, Kumar et al. [61] produced MgO nanostructures with particle size of 20-25 nm by a combustion route using $Mg(NO_3)_2$.6H₂O as the magnesium source and vetiver as the fuel. Overall, the characteristics of MgO nanoparticles produced in this study by a simple calcination technique are compatible with those produced by alternative, generally more complicated/expensive techniques.

The effect of variation time of wastewater treatment

The effects of time variation on the efficiency of COP, SOP and AP to remove chemicals from the petrochemical wastewater are exhibited in Fig. 4a-c. As can be depicted from Fig. 4a, the benzene removal efficiency in the SOP improved from 9% to 33% when the reaction time increased from 10 to 30 min. At this time slot, the improvement in the removal efficiency of toluene, ethylbenzene, and xylene could be recorded to be 18

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Fig. 4 - Influence of the variation of time and pH on the performance of SOP, AP and COP for the removal of contaminates from the petrochemical wastewater. The variation of reaction durations at the pH value of 7.5 (Table 1) for (a) SOP, (b) AP, and (c) COP. The variation of the pH solution at the contact time of 30 min for (d) SOP, (e) AP, and (f) COP.

to 58%, 25 to 81%, and 28 to 92%, respectively. Likewise, in the 257 AP, the increase in reaction time from 10 to 30 min improved 258 the benzene removal efficiency from 5 to 18%. For toluene, 259 ethylbenzene, and xylene, the improvement in the removal 260 efficiency was measured to be 15 to 50%, 24 to 81%, and 27 261 to 88%, respectively (Fig. 4b). As shown in Fig. 4c, the removal 262 efficiency in the COP for benzene rose from 47 to 93% when the 263 reaction time increased from 10 to 30 min. The improvement 264 recorded for the removal efficiencies of toluene, ethylbenzene, 265 and xylene was from 42 to 83%, 47 to 94%, and 50 to more than 266 99%, respectively. The COP procedure presents the highest effi-267 cient of removal for all contaminants, denoting the positive 268 influence of the catalytic process promoted by MgO nanopar-269 ticles. 270

The effect of pH on the wastewater treatment efficiency 271

Since pH is an important factor in ozonation process, its 272 effect on the efficiency of SOP, AP and CPO for the removal of various contaminants from the petrochemical wastewater was evaluated, and the results can be observed in Figs. 4d-f. As 274 shown in Fig. 4d, the removal efficiency increases in the SOP by 275 increasing the pH value. For p-Xylene, the removal efficiency increased from about 28% at pH 3 to 94.5% at pH 9. As shown in 277 Fig. 4e, in the AP, the adsorption efficiencies of MgO nanopar-278 ticles for all the chemicals increase when the pH increases to 279 about the neutral value, but considerably decline by further 280 increasing the pH value to 9. As presented in Fig. 4f, in the 281 COP, the efficiencies for the removal of chemicals increase by increasing the pH values. For instance, the removal efficiency 283 of ethylene, benzene and xylene increased from 45-50% at pH 284 3 to more than 90% at pH 9. As presented in Fig. 4f, by increas-285 ing the pH level of the solutions from 3-9, the efficiency of COP process to remove various contaminants from the petrochemical wastewater increases. It is noteworthy that more than 90 percent of all the petroleum compounds present in the wastewater could be decomposed at pH = 9 after 30 min of 290 treatment. It should be noticed that the positive relation of pH

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with SOP efficiency to degraded organic compounds was also
previously observed [23]. Such a relation can also be indicated
for the case of COP as evident in Fig. 4f.

Results obtained indicate that the duration of ozonation
and contact time enhance the overall performance of SOP, AP
and COP. The highest removal efficiencies achieved can be
summarized as 94.5, 88, and 99.2% for the SOP, the AP, and
COP, respectively, obtained after 30 min of treatments. Therefore, the order of removal efficiency of organic pollutants can
be regulated as AP < SOP < COP.

The adsorption proprieties and catalytic performance of 302 MgO for the degradation of n-alkanes have been evaluated in 303 the literature [62,63], highlighting the electron-acceptor and 304 electron-donor abilities of the MgO surface. Those studies 305 identify that the dye adsorption performance of MgO nanopar-306 ticles is maximized at the pH value of 6, which is in an 307 approximate agreement with our results obtained for AP pre-308 sented in Fig. 4e. 309

For the case of COP, the removal efficiencies are approximately same at the pH values of 7 and 9, providing the maximum pollutants degradation performances. This results is in agreement with those obtained by Moussavi et al. [64].

These authors suggested a mechanism for the degradation of phenol by COP, based on which, in the presence of ozone, the Lewis acid sites on the surfaces of MgO particles (MgO-s) react with ozone, and eventually with phenol, leading to the degradation event. This mechanism may also shed light on the degradation of pollutants in the current study, as explained by the occurrence of reactions (1)-(4).

 $O_3 + MgO-s \rightarrow MgO-s^{O3}$ (1)

 $MgO-s^{O3} \rightarrow MgO-s^{O\bullet} + O_2 (2)$

MgO-s^{O•} + pollutant \rightarrow CO₂ + H₂O + intermediates (3)

MgO+pollutant + $O_3 \rightarrow CO_2 + H_2O$ + intermediates (4)

Reaction (4) can be regarded as the direct oxidation of pollutant with O₃ molecules.

On the other hand, regarding the degradation of organic molecules in the presence of O_3 as an oxidizer ($E^0 = + 2.07 \text{ eV}$), it is known that under alkaline conditions, the decomposition of O_3 molecules occurs with the production of hydroxyl radicals (\bullet OH) [65]. These radicals have an even greater oxidative potential (E0 = + 2.8 eV) for decomposition of organic molecules [66,67].

In the presence of a catalyst, the production of \bullet OH can be stimulated. Also, the increase in the pH value of the reaction medium can accelerate the mass transfer of O₃ and its decomposition rate, promoting the formation of highly reactive free radicals, such as \bullet OH, \bullet O₂H and \bullet O₃H, leading to the increase of pollutants degradation efficiency [32,68,69].

The formation of ${}^{\bullet}$ OH and MgO-hydroxyl (MgO-s^{OH}) radicals have been explained to occur based on reactions (5) and (6) [32].

 $MgO-s^{O\bullet} + 2H_2O + O_3 \rightarrow MgO-s^{OH} + 3 \bullet OH + O_2$ (5)

•OH + O₂ + pollutant \rightarrow CO₂ + O₂⁻ + H₂O (6)

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The ●OH radicals could also be produced in COP, providing a
more effective environment for the oxidation of organic pollutants present in the petrochemical wastewater, in comparison
with radicals produced by SOP. The reactions (3), (4) and (6)
can occurs simultaneously, improving the overall degradation
efficiency.

Treatment of the petrochemical wastewater at the pH value of 12

Inspired by the experimental results, the petrochemical wastewater was further treated using the three distinct approaches; SOP, AP and COP, to reduce its contamination level under deviated conditions from those exhibited in Fig. 4, i.e. the enhanced pH value of 12, and extended contact duration of 50 min. The results obtained are presented in Fig. 5. As indicated, in the SOP, the xylene removal percentage is higher compared to that of ethylene benzene, toluene, and benzene. Moreover, the removal percentage of xylene and ethylbenzene in the AP are very close to each other and both higher than those of benzene and toluene. The lowest removal efficiency is related to that of benzene (33%). These observations can be explained by the combination of effects, comprising the lower solubility of benzene and toluene compared to ethylbenzene and xylene, and the higher molecular weights and melting points of xylene and ethylbenzene compared to benzene and toluene [70-72].

As shown in Fig. 5, in the AP, the highest and lowest (18%) removal efficiency could be achieved for p-Xylene and benzene, respectively. Moreover, the removal percentage of o-Xylene is lower than those of ethylbenzene and paraxylene.

As indicated by Lu et al. [72], at the initial benzene and toluene concentrations of 60 and 200 mg/L, respectively, toluene adsorption per unit of the adsorbent mass was greater than that of benzene. Moreover, Aivalioti et al. [70] showed that the adsorption capacity of diatomic adsorbents was higher for toluene than benzene. Yakout and Daifullah [73] found that a higher amount of toluene could be adsorbed per unit mass of the activated carbon than benzene. Furthermore, Su et al. [74] showed that the capacity of carbon nanotubes modified by sodium hypochlorite for the adsorption of benzene and toluene are 212 and 225 mg/g of the adsorbent, respectively, at their initial concentration of 200 mg/L, the contact time of 240 min, and adsorbent concentration of 600 mg/L. Other studies show that activated carbon adsorbent could reduce petroleum contaminants from water by up to 80% [75], although it cannot completely remove organic matter from water. Garoma et al. [76] found that the removal efficiencies of petroleum compounds by SOP could be 27%. Furthermore, they noticed that SOP was more efficient than the AP using MgO nanoparticles in removing contaminants, except for ethylbenzene, for which SOP and the AP exhibited almost the same efficiency.

As mentioned earlier, it should be noticed that molecular ozone has a high oxidation-reduction potential of 2.07 V vs. SHE, which is greater than those of most oxidants such as chlorine (1.48 V), but lower than that of hydroxyl radicals (2.8 V) [77]. As shown in Fig. 5, the COP is much more efficient than the SOP and the AP. The reason for this might lie in the fact that the simultaneous use of ozone and adsorbent accelerates the transformation of ozone from its molecular state into hydroxyl radicals. This is in agreement with the work of Dehouli et al. [78], who found that the utilization of ozonation together with the activated carbon could enhance the wastewater treatment performance, and this was attributed to the greater quantities of hydroxyl radicals present in the COP. 386

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Fig. 5 – The efficiency of SOP, AP and COP for the removal of various chemicals from the petrochemical wastewater, after 50 min of the treatment at the pH level of 12.

Table 2 – A summary on the performance of various techniques used for the removal of contaminants from the petrochemical wastewater: Concentrations of poullants in the raw wastewater (CR), and those after the ozonation (CO), the adsorption (C-AP), and the catalytic ozonation (C-COP). The removal efficiency in the ozonation process (E-SOP), adsorption (E-AP), and the catalytic ozonation (C-COP).

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Pollutant	CR	CO	C-AP	C-COP	E-SOP	E-AP	E-COP
Benzene	510.7	341.2	417.1	31.2	33%	18.3%	93.8%
Toluene	725	298.9	357	119.5	58.7%	50.7%	83.5%
Ethylbenzene	436.9	81.3	82	23.4	81.3%	81.2%	94.6%
o-Xylene	497.2	40.3	259.3	2.4	91.8%	47.8%	99%
p-Xylene	341.4	18.6	40.4	6.2	94.5%	88%	99.2%

The performance of COP for the removal of Reactive Red 198 409 in the presence MgO nanocrystals was found to be related to 410 the decomposition of dye molecules due to indirect oxidation 411 by the active radicals [32]. Valdez et al. [79,80] examined the 412 effects of COP together with volcanic rocks in the benzothia-413 zole removal, and related the high efficiency of the process to 414 the intense inclination of ozone to react with Lewis acids of the 415 metal oxides in the volcanic rocks. Moreover, results indicated 416 that the COP performed better than the SOP at contact time of 417 30 min. Since both direct and indirect oxidations are involved 418 in contaminant removal in the COP, the process exhibits a 419 higher performance at shorter contact periods compared to 420 the common SOP. It should also be noticed the ozone decom-421 position can be enhanced on the adsorbent surfaces results 422 in higher concentrations of the free radicals in the system, 423 promoting the decomposition of organic pollutants [80]. 424

It has been found that ozone decomposition can be promoted on the surfaces of crystals of natural zeolite, and particularly on volcanic rocks [79–81]. It was because such materials are capable of adsorbing ozone, followed by its conversion into free radicals.

From Figs. 4 and 5, the enhanced performance of the heterogeneous COP process, comparing to SOP and AP is evident.
The catalyst surface properties, therefore, play an important role in the ozonation process, as also indicated by Zhang et al.
[82]. Based on the results obtained by Mohammadi et al. [29], COP is an efficient and rapid method for removing of toluene

from aqueous solutions. It is known that metal oxides are able to adsorb water molecules, and this causes the formation of surface hydroxyl groups. Such hydroxyl groups exhibit various electric charge patterns at different pH values [83], affecting the decomposition of ozone on the metal oxide's surface. For instance, Qi et al. [84] studied the degradation of 2, 4, 6-trichloroanisole by the ozonation process in the presence of Al₂O₃ particles, and found that the decomposition of ozone on the alumina particles is enhanced when pH level approaches around 8. Furthemore, Garoma et al. [76] concluded that in ozonation experiments, the presence of sufficient amount of iron in groundwater samples would improve the removal of BTEX, MTBE, TBA, and TPHg. These observations are in agreement with our experimental results. Table 2 summarizes the performance of the SOP, AP and COP processes for the removal of the pollutants from the petrochemical wastewater.

It is known that the physical and chemical properties of contaminants significantly influence their removal performance from wastewaters [85].

For instance, in advanced oxidation treatments, xylene can be eliminated more than chemicals such as ethylbenzene, toluene, and benzene. Furthermore, the amount of adsorbed ethylbenzene and xylene is very close, whilst their adsorption values are considerably higher than those of benzene and toluene. These characteristics can be explained based on the properties of these chemicals. Based on this, the relatively high dissolution rate of benzene in water (1780 ppm) in 436

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comparison to other compounds such as toluene (512 ppm),
ethyl benzene (152 ppm), and xylene (175 ppm) [86,87], makes
it more difficult to be removed from wastewasters. The considerably higher molecular weight and boiling points of xylene
and ethylbenzene than toluene and benzene around (around
144, 136, 111, and 80 °C, respectively) can also influence their
removal efficiency [70,88].

470 Remarks and conclusions

Petroleum-derived chemicals can be hazardous and toxic 471 threatening the human health and environment security. 472 473 These compounds may enter the food chain via various routes, one of which is the water resources. Therefore, the removal 474 of petrochemical contaminants from water resources is of 475 significant importance. This can be carried out by various 476 methods, including adsorption and ozonation; considering 477 that each technique has its own advantages and disadvan-478 tages. Catalytic oxidation processes are among promising 479 water treatment methods which have the advantages of both 480 adsorption and oxidation approaches. Here, we investigated 481 the performance of AP, SOP, and COP on the removal of various 482 contaminants (benzene, toluene, ethylbenzene, and xylene) 483 from petrochemical wastewaters, collected from an Iranian 484 petrochemical industry. MgO nanoparticles produced by the 485 calcination of Mg (NO₃)₂6H₂O with a mean particle size of 486 around 23 nm was used as the catalyst. The effects of con-487 tact time and pH were evaluated for each approach. Results 488 showed the considerably higher performance of COP in com-489 parison with those of SOP and AP. The increase in the contact 400 491 time and the pH value of the solutions led to the improvement of removal efficiencies. During 30 min of the COP, the 492 removal efficiencies of benzene, toluene, ethylbenzene, and 493 xylene were recorded to be 93, 83, 94, and 99.2%; which are 494 considerably greater than those obtained in SOP; 33, 58, 81, 495 and 92%, respectively. MgO-enhanced COP can be consid-496 ered a promising and economical approach for the removal 497 of petroleum-based pollutants from aqueous environments. 498

Declaration of interests

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.

The authors declare the following financial interests/personal relationships which may be considered as potential competing interests:

Q3 Uncited reference

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