Cobalt-iron Oxide, Alloy and Nitride: Synthesis, Characterization and Application in Catalytic Peroxymonosulfate Activation for Orange II Degradation

Kaixin Zhu 1,2, Changzi Jin 1, Zoltán Klencsár 3, Ayyakannu Sundaram Ganeshraja 1 and Junhu Wang 1,*

1 Mössbauer Effect Data Center, Dalian Institute of Chemical Physics, Chinese Academy of Sciences, Dalian 116023, China; zhukx12@dicp.ac.cn (K.Z.); jinchangzi@dicp.ac.cn (C.J.);
ganeshraja@dicp.ac.cn (A.S.G.)
2 University of Chinese Academy of Sciences, Beijing 100049, China
3 Nuclear Analysis and Radiography Department, Institute for Energy Security and Environmental Safety, Centre for Energy Research, Hungarian Academy of Sciences, Konkoly Thege Miklós út 29-33, 1121 Budapest, Hungary; klencsar.zoltan@energia.mta.hu
* Correspondence: wangjh@dicp.ac.cn; Tel.: +86-411-8437-9159

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Abstract: In meeting the need for environmental remediation in wastewater treatment and the development of popular sulfate-radical-based advanced oxidation processes (SR-AOPs), a series of Co/Fe-based catalysts with confirmed phase structure were prepared through extended soft chemical solution processes followed by atmosphere-dependent calcination. Powder X-ray diffraction (XRD), scanning electron microscopy (SEM), transmission electron microscopy (TEM), X-ray photoelectron spectroscopy (XPS) and 57Fe Mössbauer spectroscopy were employed to characterize the composition, morphology, crystal structure and chemical state of the prepared catalysts. It was shown that calcination in air, nitrogen and ammonia atmospheres generated Co-Fe catalysts with cobalt ferrite (CoFe2O4), Co-Fe alloy and Co-Fe nitride as dominant phases, respectively. The prepared Co/Fe-based catalysts were demonstrated to be highly efficient in activating peroxymonosulfate (PMS) for organic Orange II degradation. The activation efficiency of the different catalysts was found to increase in the order CoFe2O4 < Co-Fe nitride < Co-Fe alloy. Sulfate radical was found to be the primary active intermediate species contributing to the dye degradation for all the participating catalysts. Furthermore, a possible reaction mechanism was proposed for each of the studied catalysts. This study achieves progress in efficient cobalt-iron catalysts using in the field of SR-AOPs, with potential applications in environment remediation.

Keywords: Co-Fe catalysts; advanced oxidation processes; peroxymonosulfate activation; sulfate radical

1. Introduction

Industries such as textiles, printing, plastics, pharmaceuticals, rubber, etc., may bring great threats to the water environment, owing to the production of lots of organic pollutants, most of which are carcinogenic, teratogenic, mutagenic and resistant to biodegradation [1]. Thus, advanced oxidation processes (AOPs) have been developed to solve the wastewater issues, which have been attractive techniques for the realization of powerful degradation ability. As classic active species, hydroxyl radical (·OH) generated during the oxidation processes has been widely investigated because of its remarkable
reactivity towards almost all the organic compounds [2]. However, it may suffer from limitations such as pH value dependence, slow reaction rate, sludge generation and difficulties regarding the safety of storage and transportation of the oxidant—hydrogen peroxide (H$_2$O$_2$). Recently, on account of their superior oxidation ability, good safety, wide range of pH conditions, sulfate-radical-based advanced oxidation processes (SR-AOPs) have emerged as a promising technique and a favorable alternative to H$_2$O$_2$ based organics degradation processes [3].

With respect to other transition-metal-based catalysts, cobalt ions and their composites were reported to be more efficient as peroxymonosulfate (PMS) activators in most organic pollutant degradations [4]. Since the first report about Co$^{2+}$ catalytic decomposition of PMS, homogeneous processes on different Co$^{2+}$ counteranions [5], such as nitrate, sulfate, chloride and acetate, and heterogeneous processes on cobalt oxides and supported composites [6–8], have been explored in succession. Subsequently, metal-free carbocatalysis [9–13]—studied mainly by Wang and Sun’s group—attracted great attention compared to metal heterogeneous catalysts. Even though other transition metals (Mn, Cu, Zn, Ni etc.) [14] were also effective in PMS activation, the cobalt system with no need for activity-enhancing assistance (such as microwave, UV, visible light, thermal activation, etc.) was the most popular choice for investigation. Iron possesses the advantages of low cost, rich supplies and environmental friendliness, so it has been widely applied in water remediation. The addition of Fe into the cobalt compound can not only enhance the reactivity of PMS activation through enriching hydroxyl groups on the surface, but also endow the catalyst with easier separation and recovery properties due to its magnetism, as well as suppressing cobalt leaching owing to the strong Co-Fe interactions [15]. Hu et al. prepared the catalyst by one-step incipient wetness impregnation and put forward that in comparison with Co/SBA-15, CoFe/SBA-15 maintained higher catalytic activity and better stability, and its recovery was also more easily accomplished. Iron-cobalt oxide catalysts coupled with PMS have been reported to be efficient in organics degradation [16]. More recently, octahedral CoFe$_2$O$_4$ was synthesized using a one-pot hydrothermal method and displayed excellent PMS activity [17]. Different series of Co-Fe oxides were prepared by thermal oxidation [15], sol-gel [18], co-precipitation [19,20], hydrothermal [21] and precipitation-oxidation [22] methods, with their application areas including PMS activated pollutant degradation and adsorption processes.

Previously, our group synthesized TiO$_2$ and metal-doped TiO$_2$ through a soft chemical solution process, which exhibited good photocatalytic activity in Methylene Blue (MB) degradation [23–25]. Afterwards, zinc ferrite was obtained through the same method and showed good photo-Fenton-like activity in Orange II degradation by PMS activation under visible light irradiation [26]. With the aim of avoiding the energy-consuming usage of visible light, we set out to fabricate a highly active cobalt-iron catalyst through such a soft chemical solution process. In the frame of this study, for the first time, a series of cobalt-iron catalysts including CoFe$_2$O$_4$ spinel, Co-Fe alloy and Co-Fe nitride, were prepared through soft chemical solution method followed by calcination under different atmospheres (air, N$_2$ and NH$_3$). The catalytic performances of the corresponding catalysts were investigated in PMS catalytic activation for Orange II degradation, which displayed high efficiency in organics treatments. Most importantly, Co-Fe nitride was found to have good activity and excellent stability in catalytic activation of PMS for dye degradation for the first time.

2. Results and Discussion

2.1. Basic Characterization of Cobalt-Iron Catalysts

The crystalline structures of the synthesized cobalt-iron catalysts were determined by XRD technique. Based on the XRD patterns (Figure 1), all the diffraction peaks of CFA-500 (referring to cobalt-iron catalyst calcined under air at 500 °C) and CFA-600 (referring to cobalt-iron catalyst calcined under air at 600 °C) can be attributed to CoFe$_2$O$_4$ spinel ferrite (JCPDS 00-022-1086). For the CFN samples (referring to cobalt-iron catalysts calcined under N$_2$), distinct peaks at 2θ values of 44.8° (110) and 65.3° (200) can be indexed to Co-Fe alloy (Wairauite, JCPDS 00-044-1433). When calcined under
ammonia, the main diffraction peaks at 37.5, 40.9, 42.9, 56.7, 67.7 and 75.9° were assigned to Co-Fe nitride (having a crystal structure analogous to that of Fe$_2$N, JCPDS 01-072-2126) for CFNH-600 (referring to cobalt-iron catalyst calcined under ammonia at 600 °C). But CFNH-500 (referring to cobalt-iron catalyst calcined under ammonia at 500 °C) calcined at lower temperature still contained alloy beside the main nitride phase (having similar crystal structure to that of FeN$_{0.09}$, JCPDS 01-075-2140) with low N content. Higher temperature promoted the formation of nitride phase. In addition, it was found that with the increase of the calcination temperature, the intensity of the diffraction peaks increased, indicating better crystallinity and larger particle size. According to the Scherrer equation, the grain size is 13.0, 41.7, 35.8 and 47.7 nm for CFA-500, CFA-600, CFN-500 (referring to cobalt-iron catalyst calcined under N$_2$ at 500 °C), and CFN-600 (referring to cobalt-iron catalyst calcined under N$_2$ at 600 °C), respectively. The nitride grain size is about 18.2 nm in the CFNH-500 sample while the Co-Fe alloy has grain size of about 30 nm. The main nitride phase of CFNH-600 gives grain size of 24 nm calculated from its (011) crystal plane. It can be deduced that the crystal structure of the prepared cobalt-iron catalysts mainly depends on the calcination atmosphere. Phase-pure samples of cobalt-iron oxide, alloy and nitride were obtained by calcination at 600 °C, which were chosen as catalysts for performance evaluation as discussed below.

![XRD patterns of cobalt-iron catalysts prepared under different calcination atmospheres and temperatures](image)

**Figure 1.** XRD patterns of cobalt-iron catalysts prepared under different calcination atmospheres and temperatures (a) CFA-500; (b) CFA-600; (c) CFN-500; (d) CFN-600; (e) CFNH-500 and (f) CFNH-600.

The morphologies presented by SEM for the prepared Co/Fe-based catalysts are shown in Figure 2. Obviously, the air-calcined sample of CFA-500 exhibited sphere-like nanoparticles with the size of 10–20 nm (Figure 2a), which was consistent with the XRD results. Increasing the calcination temperature led to particle aggregation (Figure 2b). When the calcination was conducted in a nitrogen atmosphere, the resultant CFN samples exhibited bright, highly dispersed sphere-like nanoparticles representing the Co-Fe alloys, along with some petal-like layers on the surface (Figure 2c,d). According to the element analysis, the distinctive morphology on the surface of CFNH-500 and CFN-600 could be attributed to the formation of cobalt-iron oxide resulting from the slight surface oxidation during the passivation process, which is consistent with the Mössbauer results below. With the increase of calcination temperature, the particles tended to grow larger. Relatively large nanoparticles with irregular shape can be found in CFNH-500 (Figure 2e), while an increase of the calcination temperature up to 600 °C resulted in a clearly different morphology. Namely, in the latter case, severe sintering took place accompanied by formation of little porous structure (Figure 2f). It is worth mentioning that N was successfully introduced into the samples during the calcination process, with its concentration...
increasing from 23% to 34% with the increase of temperature according to the energy spectrum analysis results. Correspondingly, the relative contents of Co and Fe increased but those of C and O decreased with increasing temperature. The SEM images demonstrated that different kinds of morphologies were developed under different calcination atmospheres.

**Figure 2.** SEM images of cobalt-iron catalysts prepared under different calcination atmospheres and temperatures (a) CFA-500; (b) CFA-600; (c) CFN-500; (d) CFN-600; (e) CFNH-500 and (f) CFNH-600.

TEM images of the obtained samples are presented in Figure 3. The particle size characteristic of CFA-500 was ~10 nm (Figure 3a), and about 40 nm for CFA-600 (Figure 3b) showing that the higher calcination temperature promoted the formation of larger particles, in agreement with corresponding SEM and XRD results. Nevertheless, metal particles dispersed on carbon support were obtained by N$_2$ calcination (Figure 3c,d). Carbon in CFN samples is generated from the carbonization of organic species present in reactants during calcination under nitrogen. One can discern that many dark-colored nanoparticles (metal species) were dispersed in the light-colored matrix, which was ascribed to carbon species formed during the carbonization process. The nanoparticles grew larger, even to 50 nm, when the calcination temperature was increased from 500 to 600 °C. As for the CFNH-500, non-uniform nanoparticles were observed, interestingly, with the appearance of a core-shell structure (Figure 3e). However, irregular morphology was present on the Co-Fe nitride (Figure 3f). It is clear that higher calcination temperature leads to severe particle aggregation and larger particle sizes.

**Figure 3.** TEM images of cobalt-iron catalysts prepared under different calcination atmospheres and temperatures (a) CFA-500; (b) CFA-600; (c) CFN-500; (d) CFN-600; (e) CFNH-500 and (f) CFNH-600.
with the increase of calcination temperature, the BET specific surface area and pore volume decrease. (A) site (sextet), octahedral (B) site (sextet with broad peaks reflecting binomial distribution of cobalt atoms at the tetrahedral site) and the smallest particles with superparamagnetic nature (quadrupole doublet). In case of samples CFN-500 and CFN-600, the spectra (Figure 5c,d) indicate the presence of the bcc alloy phase with an Fe:Co ratio that is close to 2:1 in both cases (The fit of the corresponding component was carried out as described in the previous work [27,28]). We can observe three components corresponding to Fe$^{3+}$ in tetrahedral (A) site (sextet), octahedral (B) site (sextet with broad peaks reflecting binomial distribution of cobalt atoms at the tetrahedral site) and the smallest particles with superparamagnetic nature (quadrupole doublet). There is a more complete level of inversion (i.e., a lower concentration of Co at tetrahedral sites) present in the sample CFA-600, than in CFA-500, which is presumably connected to the higher calcination temperature applied in the former case. A less intense Fe$^{3+}$ doublet component, accounting for less than 1% of the spectral area, may also be present in the spectra (Table S2). Its contribution is smaller in the case of CFA-600, suggesting that higher calcination temperature and/or higher particle size tends to diminish its presence.

In case of samples CFN-500 and CFN-600, the spectra (Figure 5c,d) indicate the presence of the bcc Fe$_x$Co$_{1-x}$ alloy phase with an Fe:Co ratio that is close to 2:1 in both cases (The fit of the corresponding component was carried out as described in the previous work [29]). A heavily broadened spectral component can be seen in CFN-500 as being associated with the smallest Co-Fe alloy particles that undergo magnetic relaxation on a time scale which is comparable to the Larmor precession time of the $^{57}$Fe nucleus. This component is missing from the spectrum of CFN-600 (Figure 5d), in agreement with the larger characteristic particle size of the latter sample. A paramagnetic Fe$^{3+}$ doublet component can

Figure 4 displays the $N_2$ adsorption-desorption isotherms and corresponding pore size distribution of the prepared Co/Fe-based catalysts. Corresponding textural parameters are displayed in Table S1. Except CFNH-600, other samples give isotherms with hysteresis loops at relatively high pressures of 0.4–1.0, which indicates the existence of mesopores generated due to the accumulation of nanoparticles. This is confirmed by the calculated pore size distribution that exhibits mesopores (mainly in the range of 2–40 nm), and some macropores (>50 nm) as well. The absence of accumulation pores in CFNH-600 can be attributed to its bulk particles, which were generated via aggregation during calcination. The large particle size of CFNH-600 also leads to its lower BET surface area (13.0 m$^2$ g$^{-1}$) and pore volume (0.019 cm$^3$ g$^{-1}$) relative to other samples. Regardless of the calcination atmosphere, with the increase of calcination temperature, the BET specific surface area and pore volume decrease markedly for all the samples, which is another consequence of the particle aggregation, in accordance with SEM and TEM results. The large surface area of the $N_2$-calcined samples, especially for CFN-500, is mainly due to the existence of carbon in samples.

2.2. $^{57}$Fe Mössbauer Spectroscopy Characterization

Mössbauer spectroscopy is an excellent tool for examining the local environment of Fe in cobalt-iron catalysts. Room temperature $^{57}$Fe Mössbauer spectra of the six samples are shown in Figure 5a–f, together with their fit. In the case of CFA-500 and CFA-600, the spectra (Figure 5a,b) mainly reflect the presence of Fe$^{3+}$ in the CoFe$_2$O$_4$ spinel phase with a partial degree of inversion. CoFe$_2$O$_4$ is known to prefer a partially inverse spinel structure where majority of the A (tetrahedral) site is taken by Fe$^{3+}$ [27,28]. We can observe three components corresponding to Fe$^{3+}$ in tetrahedral (A) site (sextet), octahedral (B) site (sextet with broad peaks reflecting binomial distribution of cobalt atoms at the tetrahedral site) and the smallest particles with superparamagnetic nature (quadrupole doublet). This component is missing from the spectrum of CFN-600 (Figure 5d), in agreement with the larger characteristic particle size of the latter sample. A paramagnetic Fe$^{3+}$ doublet component can
also be detected in the spectrum of these samples, with a considerably smaller relative spectral area for CFN-600 (~2%) than for CFN-500 (~7%) (Table S3), suggesting that this amount can be reduced by increasing the calcination temperature.

In the sample that was calcined at 500 °C under NH₃ atmosphere, one can detect both Co-Fe alloy (magnetic sextet component) and Co-Fe nitride (paramagnetic doublet components) (Figure 5e, Table S4). In contrast, only one paramagnetic iron doublet is observed (Figure 5f) when the calcination was carried out at 600 °C: the Mössbauer parameters of the doublet are similar to those of Fe₂N [30], which is consistent with the XRD results.

2.3. Surface Elements Analysis

XPS measurements were carried out on the cobalt-iron samples to investigate the surface characteristics. Full XPS survey in Figure 6a shows the compositional information of the sample CFA-600, indicating the presence of Co, Fe, O, as well as C. The high-resolution XPS spectra of Co 2p, Fe 2p and O 1s are shown in Figure 6b–d, respectively. The Co 2p₃/₂ peaks at 779.5 eV and 781.5 eV were ascribed to Co²⁺ at octahedral (B) and tetrahedral (A) sites of the spinel structure, respectively, with a satellite peak of Co 2p₃/₂ at ~785.5 eV [31]. There were also two peak signals for Fe 2p₃/₂ at 710.0 and 712.7 eV, suggesting the occupation of octahedral and tetrahedral sites, respectively. Additionally, one satellite peak appeared at 719.1 eV, which was further proof for the existence of Fe³⁺. The O 1s peaks at 529.5 and 531.1 eV were assigned to lattice oxygen and surface adsorbed oxygen, respectively. The results are consistent with the Mössbauer fit results that spinel structured cobalt ferrite with some degree of inversion was observed. Similar spectra were obtained on the sample CFA-500 (Figure S1), revealing that the main phase of samples obtained under air was cobalt ferrite.
The peaks at 284.5, 285.5, 286.5 and 288.3 eV were assigned to C=C (or C–C), C–O, C=O or O–C–O, respectively [32]. However, the Co 2p and Fe 2p peaks could be distinguished evidently on CFN-600 (Figure 7), maybe because of the relatively higher contents of Co (2.86%) and Fe (3.87%) on the surface. The Co 2p$_{3/2}$ peaks at 778.2 and 781.1 eV were ascribed to Co$^0$ and Co$^{2+}$, respectively, with a satellite peak at 786.6 eV. As for iron, beside Fe$^{3+}$, a small amount of Fe$^0$ can be observed through the peak of 708.1 eV [17]. Thus, we can discern that the metals on the surface were oxidized by the ambient air to some extent. Furthermore, only three peaks at about 284.6, 285.4 and 286.2 eV were observed showing a dramatic decrease in carbon-bonded oxygen groups along with the absence of carboxyl group. The results indicate that the carbon species obtained at different carbonization temperatures display different characteristics. In order to illustrate the variation between the CFN-500 and CFN-600, thermal analysis under inert atmosphere with a heating rate of 10 °C min$^{-1}$ was conducted, as shown in Figure S3a. The TGA plots revealed dramatic weight loss from 200 to 600 °C, which can be assigned to the removal or destruction of oxygenated functional groups. Two main endothermic peaks can be observed at about 300 and 530 °C, accompanied by the release of CO and CO$_2$ as determined with TG-MS technique (Figure S3b), suggesting that carbonization occurred and two different carbon-containing complexes were decomposed during these processes. The appearance of CO indicated that self-reduction occurred during the calcination, resulting in the formation of Co-Fe alloy. Importantly, the second endothermic peak appeared at about 533 °C, clearly revealing that the reducing degree under 600 °C was higher than that of 500 °C and that different carbon-bonded groups were found in CFN-500 and CFN-600.

Figure 6. XPS spectra of CFA-600 in different regions (a) survey; (b) Co 2p; (c) Fe 2p and (d) O 1s.
Chemical states of surface elements in cobalt-iron nanomaterials CFNH-500 and CFNH-600 obtained under ammonia atmosphere are presented in Figure S4 and Figure 7, respectively. From the survey in Figure 8a, clear peaks of Co, Fe, C, N and O were observed, further certifying the successful introduction of element N through NH$_3$ calcination. High resolution Co 2p$_{3/2}$ peaks present at 780.4 and 782.7 eV can be ascribed to Co$^{2+}$ and Co$^{3+}$, respectively, with a satellite at 786.6 eV (Figure 8b).

Concerning iron, only Fe$^{3+}$ was detected with a satellite peak at 719.1 eV (Figure 8c). However, peaks of 778.5 and 707.9 eV were also found in the case of CFNH-500 (Figure S4), which can be assigned to metallic Co$^0$ and Fe$^0$, consistent with the XRD result that Co-Fe alloy was formed under 500 °C. The appearance of high valence cobalt and iron suggested that the surface of the sample was oxidized when exposed to air, similarly to the case of the CFN samples. In addition to Co and Fe, high resolution XPS peaks of C, N and O were also observed. Peaks centered at 284.6, 285.8, 286.9 and 288.6 eV (Figure 8d) can be attributed to the C=C or C–C, C–O or C–N, C=O and O–C=O, respectively [33,34]. The N1s spectrum (Figure 8e) can be deconvoluted into five peaks centered at 397, 398.6, 400.1, 402.7 and 404.7 eV, representing different kinds of nitrogen species and confirming the presence of N in the sample. The peaks at 397, 400.1, 402.7 and 404.7 eV could be assigned to the pyridinic, pyrrollic, graphitic N and oxidized pyridinic N [35,36]. The peak at 398.6 eV was ascribed to the pyridinic N coordinated with Fe or Co, i.e., N–Fe or N–Co, according to the literatures [36,37]. Less than 10% of graphitic N was found. The majority of pyridinic N resulting from the coordination with metal (Fe or Co) [38], verified the formation of metal nitride. The O 1s peaks at 530.0 and 531.7 eV (Figure 8f) were ascribed to the lattice oxygen coordinated with metal and surface hydroxyl oxygen, respectively [39]. It should be noted that a peak at 536.0 eV, which may originate from organics residuals such as carboxyl groups, was also detected for both CFNH-500 and CFNH-600.
which may have resulted from the existence of carbon. After PMS addition, the removal rate of Orange II on the catalysts calcined under N\textsubscript{2} (CFN-600) was significantly higher than that of CFA-600 and CFNH-600. CFN-600 was able to effectively degrade Orange II through activating PMS to produce oxidizing species, with more than 90% removal within 10 min. For the CFNH-600, whose catalytic activity measurements the Co-Fe alloy provided the highest Orange II removal efficiency and the activity order of the different catalysts is CFN-600 > CFNH-600 > CFA-600. Besides, according to the ICP results (Table S5), Co-Fe alloy gave more metal ions leaching, which could also contribute to the degradation activity through homogeneous catalysis. In comparison to the spinel, the nitride catalyst displayed superior activity in PMS activation, which may be due to the different adsorption surfaces with PMS molecules endowing the nitride with stronger adsorption energy, longer O-O bond lengths and more transferred electrons [40]. On the whole, according to the activity measurements the Co-Fe alloy provided the highest Orange II removal efficiency and the activity order of the different catalysts is CFN-600 > CFNH-600 > CFA-600.

In order to estimate the kinetic parameter of the degradation reactions, first order kinetics was employed for PMS activated pollutant degradation, with the equation

\[ \ln \left( \frac{c_0}{c} \right) = kt \]  

(1)
where $k$ is the apparent first order rate constant of Orange II removal, $c_0$ is the original concentration of Orange II aqueous solution, $c$ is the real time concentration of Orange II during the reaction, and $t$ is the reaction time. The reaction rate constants (Figure 9B) for CFA-600, CFN-600 and CFNH-600 were estimated to be 0.007 ($R^2 = 0.976$), 0.322 ($R^2 = 0.939$) and 0.024 ($R^2 = 0.907$) min$^{-1}$, respectively. Based on the appropriate regression coefficients for the reaction system, the pseudo-first-order kinetic was basically thought to fit well with the experiment. These catalysts yielded results that were similar to those of the previous studies [41,42], where the kinetic rate of organic pollutants was explored in the presence of PMS. The rate constant of CFN-600 was 46 times and 13.4 times higher than that of the CFA-600 and CFNH-600 samples, respectively, suggesting the excellent performance of the Co/Fe-based catalysts obtained under N$_2$ atmosphere. The catalytic efficiency of Co-Fe nitride was 3.4 times better than the corresponding CoFe$_2$O$_4$ in this study, clearly revealing the superiority of nitride. The sequence of the catalytic activity of different types of cobalt-iron samples is Co-Fe alloy > Co-Fe nitride > Co-Fe oxide.

Apart from the excellent catalytic activity, the stability and reusability of a heterogeneous catalyst is also crucial in the catalytic reaction processes. Figure 10 shows the successive test results of the three cobalt-iron catalysts calcined at 600 °C for Orange II degradation. It can be seen that deactivation took place on the spinel and alloy catalysts during the measured time (Figure 10a,b). At the same time, the nitride kept the original activity, indicating its high stability and reusability (Figure 10c). Previous literature made the point that the formation of Fe$_3$C contributed to a better stability in the Fe$^0$/Fe$_3$C@CS catalyst [43]; in our case, nitride may play a similar role. Besides, manganese nitride (Mn$_4$N) was reported to possess good stability compared to the zero valent Co, Fe and Mn in PMS activation for BPA degradation [40]. From the point of view of reusability and stability, Co-Fe nitride exhibited the best performance. According to the XRD patterns (Figure S5) of the used cobalt-iron catalysts, no variation took place in the phase composition compared to the fresh catalysts. However, surface chemical compositions and states of the used catalysts experienced partial oxidation for CFA-600 and CFN-600. In the used CFA-600, a new peak appeared at 783.1 eV representing Co$^{3+}$ in octahedral site appeared (Figure 11a), indicating the oxidation of Co$^{2+}$ to Co$^{3+}$ during the catalytic reaction. This transformation was also reported by Ren et al. who found that Co$^{3+}$ emerged after PMS activated degradation [18]. Besides, no significant variation took place in the valence state of Fe before and after reaction, since only Fe$^{3+}$ was observed (Figure 11b). As for CFN-600, the peak of Co$^0$ disappeared, at the same time, the peak at 783.3 eV representing Co$^{3+}$ turned up compared to the fresh catalyst (Figure 11c). It was demonstrated that metallic Co$^0$ was totally oxidized, and Co$^{2+}$ was partially oxidized into Co$^{3+}$ during the catalytic oxidation reaction. Concerning iron, only Fe$^{3+}$ was
present in the used catalyst (Figure 11d), manifesting the oxidation of metallic Fe\(^0\). It was found that Fe\(^0\) was reactive in catalytic PMS activation, being converted to Fe(II)/Fe(III) after the reaction [43,44]. These results demonstrated that the active species on the catalyst surface were oxidized to some extent, which was one of the proofs for the deactivation. Besides, according to the XPS results in Figure S6, element sulfur was discovered for the used CFN-600 sample, suggesting that organic compounds were adsorbed on the catalysts, which could be another reason for its deactivation. From the adsorption curves in Figure 10b, the adsorbed amounts of Orange II decreased from run1 to run3, suggesting the decreased adsorption ability and the coverage of the catalyst surface. More importantly, after extending the reaction time to 80 min (Figure 10d), the CFN-600 sample presented good reusability with no deactivation in run2 and run3, further verifying that adsorbed organics were the main reason for the deactivation testing in a short time (Figure 10b). Thus, surface active sites may be covered by the adsorbed organic compounds during reaction, resulting in the decrease of activity. Similar results were obtained on the CFNH-600 sample (Figure 12a), where sulfur was detected only after reaction. But the small BET surface area of the nitride sample may inhibit the adsorption of organic species from having appreciable impact on the activity of the catalyst, endowing it with better reusability. No significant changes were found on the variation of total pyridinic N (59 ± 2%), pyrrolic N (25 ± 2%), graphitic N (10 ± 2%) and oxidized nitrogen species (5 ± 1%) in the fresh and used catalyst (Table S6), certifying the superior stability of nitrides.

To explore the variation in the chemical state of Fe, \(^{57}\)Fe Mössbauer spectra of the used catalysts were also measured, as shown in Figure 13. For CFA-600, there was no appreciable change detected concerning the Fe\(^{3+}\) species in the spinel, but the relative area fraction of the paramagnetic Fe\(^{3+}\) doublet component increased up to ~2.5% (Table S2). Similarly, the paramagnetic Fe\(^{3+}\) doublet in CFN-600 increased from ~2% to 9% during reaction (Table S3), with a corresponding decrease in metallic Fe\(^0\) in the Co-Fe alloy phase, which also refers to the oxidation of catalysts, in agreement with the XPS results. No variation occurred in the state of Fe in CFNH-600, providing further confirmation in the stability of this catalyst. Hence, the good stability and reusability made the nitride more promising in the environmental remediation.

![Figure 10](image_url)

**Figure 10.** Reusability in Orange II degradation on different samples (a) CFA-600; (b) CFN-600 (14 min); (c) CFNH-600 and (d) CFN-600 (80 min). (Conditions: Orange II, 50 mg L\(^{-1}\); Catalysts, 0.1 g L\(^{-1}\); Oxone, 0.5 g L\(^{-1}\)).
Figure 11. XPS spectra of the used CFA-600 (a) Co 2p$_{3/2}$; (b) Fe 2p and CFN-600 (c) Co 2p$_{3/2}$; (d) Fe 2p.

Figure 12. High resolution XPS spectra of the used CFNH-600 in different regions (a) S 2p before and after reaction; (b) Co 2p; (c) Fe 2p; (d) C 1s; (e) N 1s and (f) O 1s.
Figure 13. $^{57}$Fe Mössbauer spectra of the used CFA-600 (a); CFN-600 (b) and CFNH-600 (c).

2.5. Reaction Mechanism Study

Previous investigations have pointed out that in the cobalt-iron homogeneous and heterogeneous catalytic activation of the peroxymonosulfate reaction, sulfate radical (SO$_4^{2-}$) and hydroxyl radical ($\cdot$OH) can always be identified as the main reactive species [45]. To ascertain the main intermediate active species in the three different systems, radical scavenger tests were taken in this study. It is widely accepted that alcohols with an $\alpha$-hydrogen such as ethanol readily react with both SO$_4^{2-}$ and $\cdot$OH (1.6–7.7 × 10$^7$ M$^{-1}$ s$^{-1}$ and 1.2–2.8 × 10$^9$ M$^{-1}$ s$^{-1}$, respectively), while those without $\alpha$-H, such as tert-butyl alcohol (TBA), react more easily with $\cdot$OH (3.8–7.6 × 10$^8$ M$^{-1}$ s$^{-1}$), but in a much more slower reaction rate with SO$_4^{2-}$ (4.0–9.1 × 10$^5$ M$^{-1}$ s$^{-1}$) [46,47]. Thus, TBA as a particular scavenger for $\cdot$OH and ethanol as a universal scavenger for both SO$_4^{2-}$ and $\cdot$OH were used to differentiate the contribution of SO$_4^{2-}$ from $\cdot$OH. In the organic degradation process, where radicals played a decisive role when the quenching agent was added into the solution, the degradation would be either...
role when the quenching agent was added into the solution, the degradation would be either prevented or significantly reduced. As shown in Figure 14, degradation was inhibited for all three samples with the addition of ethanol. More importantly, there was hardly any effect observed on the CFN-600 sample when using the same amounts of TBA as a scavenger, suggesting that sulfate radical was the main reactive species. As for CFA-600 and CFNH-600, the addition of TBA caused some inhibiting effects in the Orange II degradation efficiency, but ethanol made a more significant contribution. Therefore, we could conclude that sulfate radical played the key role in the CFA-600/PMS and CFNH-600/PMS systems, with a certain amount of hydroxyl radical; only sulfate radical played a role as reactive species in the CFN-600/PMS system.

Figure 14. Radical scavenger tests of (a) CFA-600; (b) CFN-600 and (c) CFNH-600.
In the past years, several investigations have been carried out to recognize the mechanism of Co/Fe-based catalysts in the activation of PMS. It has been reported that when CoFe$_2$O$_4$ was used as a catalyst for PMS activation, cobalt mainly presented as Co$^{2+}$ before reaction, and it underwent partial transformation to Co$^{3+}$ after catalytic reaction, in which the involvement of Co$^{2+}$-Co$^{3+}$-Co$^{2+}$ redox processes were proposed during the reaction [18]. In addition, sulfate radical was found to be generated as the main active species in oxone activation by cobalt-iron oxides [22,48]. Concerning the CFA sample in the present study, the reaction mechanism for Orange II degradation can be proposed as following (Equations (2)~(7)).

\[
\begin{align*}
\equiv & \text{Co}^{2+} + \text{HSO}_5^- \rightarrow \equiv \text{Co}^{3+} + \text{SO}_4^{2-} + \text{OH}^- \quad (2) \\
\equiv & \text{Fe}^{2+} + \text{HSO}_5^- \rightarrow \equiv \text{Fe}^{3+} + \text{SO}_4^{2-} + \text{OH}^- \quad (3) \\
\equiv & \text{Co}^{3+} + \text{HSO}_5^- \rightarrow \equiv \text{Co}^{2+} + \text{SO}_5^- + \text{H}^+ \quad (4) \\
\equiv & \text{Fe}^{3+} + \text{HSO}_5^- \rightarrow \equiv \text{Fe}^{2+} + \text{SO}_5^- + \text{H}^+ \quad (5) \\
\equiv & \text{Fe}^{2+} + \equiv \text{Co}^{3+} \rightarrow \equiv \text{Co}^{2+} + \equiv \text{Fe}^{3+} \quad (6) \\
\text{SO}_4^{2-} + \text{Orange II} \rightarrow & \text{many steps} \rightarrow \text{CO}_2 + \text{H}_2\text{O} + \text{SO}_4^{2-} \quad (7)
\end{align*}
\]

It was reported that the active sites are mainly the cobalt and iron ions in the spinel type CoFe$_2$O$_4$ and the zero-valent cobalt and iron for the octahedral CoFe/CoFe$_2$O$_4$ submicron composite [17]. In the case of the CFN sample, the following reactions (Equations (8)~(11)) may be considered, with the rest of the steps being identical to processes described by Equations (2)~(7) above.

\[
\begin{align*}
\equiv & \text{Co}^0 + 2\text{HSO}_5^- \rightarrow \equiv \text{Co}^{2+} + 2\text{SO}_4^{2-} + 2\text{OH}^- \quad (8) \\
\equiv & \text{Fe}^0 + 2\text{HSO}_5^- \rightarrow \equiv \text{Fe}^{2+} + 3\text{SO}_4^{2-} + 2\text{OH}^- \quad (9) \\
\equiv & \text{Co}^0 + 2\equiv \text{Co}^{3+} \rightarrow 3\equiv \text{Co}^{2+} \quad (10) \\
\equiv & \text{Fe}^0 + 2\equiv \text{Fe}^{3+} \rightarrow 3\equiv \text{Fe}^{2+} \quad (11)
\end{align*}
\]

For the Co-Fe nitride, the redox process of Co$^{2+}$-Co$^{3+}$-Co$^{2+}$ also exist in the system, and one can observe only small changes in the relative amounts of Co$^{2+}$ and Co$^{3+}$ according to the XPS fitting results of the CFNH-600 before and after reaction. The potential mechanism could be speculated to be similar to that mentioned for the CFA sample, except that the introduction of N into the catalyst made its stability appreciable, and at the same time, the activity superior to the oxide.

3. Materials and Methods

3.1. Chemicals

Cobalt acetylacetonate (Co(acac)$_2$), ferric acetylacetonate ((Fe(acac)$_3$), Orange II sodium salt (C$_{16}$H$_{11}$N$_2$NaO$_4$S) and tert-butyl alcohol (TBA, >99.5%) were purchased from Aladdin Chemistry Co. Ltd. (Shanghai, China) 2-ethoxyethanol (C$_4$H$_{10}$O$_2$) and methanol (CH$_3$OH, ≥ 99.5%) were purchased from Tianjin Damao Chemical Reagent Company (Tianjin, China) and H$_2$O$_2$ (30 wt.%) from Tianjin Kermel Chemical Reagent Company (Tianjin, China). PMS was purchased from Alfa Aesar (Tianjin, China). Ethanol (CH$_3$CH$_2$OH, ≥ 99.7%) was obtained from Sinopharm Chemical Reagent Co. Ltd. (Shanghai, China). All the chemicals were analytical grade with no further treatment. Deionized water was used for solution preparation.

3.2. Synthesis of Cobalt-Iron Catalysts

Co/Fe-based catalysts were prepared through an extended soft chemical solution process reported previously [49,50]. Typically, Co(acac)$_2$ and Fe(acac)$_3$ (molar ratio 1:2) were first dissolved
homogeneously together in 50 mL 2-ethoxyethanol within 1 L beaker at 50 °C, followed by slow addition of 30 mL H₂O₂ aqueous solution under magnetic stirring. The suspension was obtained after rotary evaporation and drying. Finally, the catalysts were obtained by calcining the solid precursors at 500 or 600 °C under special atmosphere. The prepared cobalt ferrite samples are denoted as CF-X-T, where X represents the calcination atmosphere of air (A), nitrogen (N) and ammonia (NH) and T represents the final calcination temperature.

3.3. Catalysts Characterization

The X-ray diffraction (XRD) patterns were measured on a PW3040/60 X'Pert PRO diffractometer (PANalytical, Alemlo, The Netherlands) equipped with a copper target (Cu Kα radiation, λ = 1.5406 Å) at an accelerating voltage of 40 kV, a current of 40 mA, and a scanning 2θ range of 10°–80° at room temperature. The average crystallite sizes were calculated by Scherrer equation as below:

\[ \varphi = \frac{K\lambda}{\beta\cos\theta} \]  

(12)

where \( \varphi \) is the average crystallite size, \( K \) is the Scherrer constant (taken to be equal to 0.89); \( \lambda \) is the wavelength of the used X-ray, \( \theta \) is the diffraction angle of the calculated peak, and \( \beta \) is the full width at half maximum height (FWHM) of the mean intensity peak. The morphological feature was observed on a JEM-2000EX electronic microscope (JEOL, Tokyo, Japan) with an accelerating voltage of 120.0 kV. The sample was dispersed in the ethanol solution and then withdrawn to drop on a copper grid for measurement. The field-emission scanning electron microscopy (FESEM, JSM-7800F) (JEOL, Tokyo, Japan) was operated at an accelerating voltage of 3.0 kV, and the sample was put onto a conductive carbon tape to be measured. Textural information was measured by nitrogen adsorption/desorption at 77 K using a Micromeritics ASAP 2460 apparatus (Micromeritics, Norcross, GA, USA). Prior to measurement, all the samples were pretreated in vacuum at 110 °C for 1 h, followed by 300 °C for 6 h. The thermogravimetric and differential scanning calorimetry (TG-DSC) curves in N₂ were investigated on a TGA instrument Q600 (TA, New Castle, PA, USA) with alumina crucible as sample holder, and the heating rate was set at 10 °C min⁻¹ with flow rate of 100 mL min⁻¹. The metal ion leaching amount after reaction was determined by inductively coupled plasma-atomic emission spectrometry (ICP-AES) on an IRIS Intrepid II XSP instrument (rThermoFisher, Waltham, MA, USA). The X-ray photoelectron spectroscopy (XPS) experiment was performed on an ESCALAB250 X-ray photoelectron spectrometer (ThermoFisher, Waltham, MA, USA) with monochromatic Al Kα as the X-ray source, and contaminated carbon was used as the internal standard (C1s = 284.6 eV). \(^{57}\)Fe Mössbauer spectra were measured on a Topologic 500A spectrometer employing a \(^{57}\)Co(Rh) as γ-ray radioactive source moving in a constant acceleration mode at room temperature. The spectra were fitted by using the MossWinn 4.0 program [51]. Room temperature α-iron foil was applied for the calibration of the velocity axis, and \(^{57}\)Fe isomer shift values are quoted relative to that of the standard.

3.4. Activity Evaluation

The prepared cobalt-iron catalysts were applied in catalytic degradation of Orange II by activation of PMS. All the experiments were carried out in a beaker with constant rotation speed. Prior to the degradation experiments, proper amounts of catalyst were mixed with Orange II solution under stirring for 30 min to achieve adsorption-desorption equilibrium, and then PMS was added into the system to initiate the reaction. The typical concentrations of catalyst, PMS and Orange II were 0.1 g L⁻¹, 0.5 g L⁻¹ and 50 mg L⁻¹, respectively. At certain time intervals, 1 mL reaction solution was withdrawn and quenched immediately by adding 1 mL methanol solution. After centrifugation, the supernatant was analyzed by the GBC Cintra apparatus to record variation of the absorption peak intensity (wavelength at 484 nm for Orange II aqueous solution).
4. Conclusions

Three kinds of cobalt-iron catalysts with the dominant phase of CoFe$_2$O$_4$ spinel, Co-Fe alloy and Co-Fe nitride were successfully synthesized through extended soft chemical solution processes followed by atmosphere-dependent calcination. All these obtained catalysts were applied to the Orange II degradation by PMS activation and performed good pollutants removal efficiency. By comparison, Co-Fe alloy exhibited the best activity and Co-Fe nitride possessed very good stability. Sulfate radical was proved to be the main active species in Co-Fe alloy system, while hydroxyl radical is also effective in the Co-Fe nitride and CoFe$_2$O$_4$ participant reactions. Considering the severe leaching problem of the zero valent metals, efforts need to be made to retard the leaching problem and improve the stability. Besides, the detailed properties and mechanisms of the catalysts especially for the nitride-derived catalysts need to be further explored.

Supplementary Materials: The following are available online at www.mdpi.com/2073-4344/7/5/138/S1, Figure S1: XPS spectra of CFA-500 in different regions (a) survey, (b) C 1s, (c) Co 2p, (d) Fe 2p. Figure S2: XPS spectra of CFN-500 in different regions (a) survey, (b) Co 2p, (c) Fe 2p, and (d) C 1s. Figure S3: TG-DSC (a) and the TG-MS (b) curves of the as-prepared cobalt-iron nanomaterials under Ar atmosphere. Figure S4: XPS spectra of CFNH-500 in different regions (a) survey, (b) Co 2p, (c) Fe 2p, (d) C 1s, (e) N 1s and (f) O 1s. Figure S5: XRD patterns of the used CFA-600, CFN-600 and CFNH-600. Figure S6: S2p XPS spectrum of the CFN-600 before and after reaction. Table S1: The BET surface area and pore volume of prepared samples. Table S2: $^{57}$Fe Mössbauer parameters of the samples CFA-500 and CFA-600 as well as CFA-600-after reaction derived on the basis of Mössbauer spectra recorded at room temperature. Table S3: $^{57}$Fe Mössbauer parameters of the samples CFN-500 and CFN-600 as well as CFN-600-after reaction derived on the basis of Mössbauer spectra recorded at room temperature. Table S4: $^{57}$Fe Mössbauer parameters of the samples CFNH-500 and CFNH-600 derived on the basis of Mössbauer spectra recorded at room temperature. Table S5: Metal leaching concentrations detected by ICP. Table S6: XPS fitting results of surface elements for the CFA-600, CFN-600 and CFNH-600 samples before and after reaction.

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