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Sulfur-Tolerant, Exsolved Fe-Ni Alloy Nanoparticles for CO Oxidation

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Abstract

Metallic nanoparticles exsolved at the surface of perovskite oxides have been recently shown to unlock superior catalytic activity and durability towards various chemical reactions of practical importance. For example, for the CO oxidation reaction, exsolved Ni nanoparticles in oxidised form exhibit site activities approaching those of noble metals. This is of particular interest for the prospect of replacing noble metals with earth-abundant metal/metal oxide catalysts in the automotive exhaust control industry. Here we show that for the CO oxidation reaction, the functionality of exsolved Ni nanoparticles can be further improved when Fe is co-exsolved with Ni, as Fe–Ni alloy nanoparticles, eventually forming mixed oxide nanoparticles. As compared to the Ni nanoparticles, the alloy nanoparticles exhibit higher site activities, greatly improved durability over 170 h of continuous testing and increased tolerance towards sulphur-based atmospheres. Similarly to the single metal nanoparticles, the alloys demonstrate outstanding microstructural stability and high tolerance towards coking. These results open additional directions for tailoring the activity and durability of exsolved materials for the CO oxidation reaction and beyond.

Keywords Exsolution · Perovkites · Metal nanoparticles · Sulphur tolerance · CO oxidation

1 Introduction

The carbon monoxide (CO) oxidation reaction is of key importance in the area of automotive exhaust treatment, i.e. in three-way catalytic converters [1]. Recently, it has been shown that the exsolution of base metal nanoparticles from perovskite oxides may unlock superior catalytic activity towards the CO oxidation reaction [2]. For example, exsolved nickel (Ni) nanoparticles can exhibit nominal turnover frequencies (nTOF, number of CO molecules converted to CO_2 per second, per exposed metal atom site) of the order of hundreds s⁻¹. This is orders of magnitude higher

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than the values typically reported for base metal/metal oxide nanoparticles in literature and approaching values observed for noble metals [3]. This unusually high catalytic activity, along with high resistance to carbon deposition, has been assigned to the confined nature of exsolved nanoparticles which are socketed within the perovskite oxide support [4]. The socketed structure is believed to form as a result of the exsolution process itself [5, 6]. That is, initially the base metal ions, e.g. Ni²⁺, are substituted in the perovskite oxide lattice in oxidising conditions, and upon exposure to reducing conditions, they emerge at the surface forming metallic nanoparticles [7].

So far, the approach used to further enhance the catalytic activity of exsolved Ni particles for CO oxidation, has been through particle restructuring via alloying with cobalt (Co) to trigger the Kirkendall effect [8], in a so-called "chemistry at a point" concept [2]. On the other hand, the effect of alloying on its own has yet to be explored, even though many bimetallic catalyst systems generally show superior performance as compared to their single metal counterparts, probably due to changes in electronic and structural parameters [9–15]. While cobalt oxides have been considered as one of the most promising candidates for use in bimetallic systems for CO oxidation due to its high activity [16], the stability of Co-based catalysts may be compromised among

other factors by the presence of poisons, such as sulphur, and further potential restructuring due to the Kirkendall effect. Iron (Fe) has received significant attention as an alternative alloying metal for Ni-based systems because of its lower cost and toxicity, as compared to Co, and high abundance. Furthermore, Fe performs well in catalysing oxidation reactions in rich and lean oxidation environments due to the high capacity of reactive lattice oxygen species [17, 18].

Here we investigate the catalytic activity of exsolved Fe-Ni bimetallic alloy nanoparticles for the CO oxidation reaction and demonstrate that they exhibit enhanced site activity and durability as compared to the Ni variant and other base-metal systems prepared by conventional routes. Additionally, we investigate their catalytic behaviour under CO-rich conditions of the CO oxidation reaction and observe high tolerance against carbon deposition (coking) whilst maintaining high catalytic activity and structural integrity. Furthermore, Fe-Ni nanoparticles also display improved SO₂ tolerance. To illustrate the above, we employ two perovskite oxide systems, $La_{0.8}Ce_{0.1}Ni_{0.4}Ti_{0.6}O_3$ and $La_{0.5}Sr_{0.4}Fe_{0.1}Ni_{0.1}Ti_{0.6}O_3$ in order to enable exsolution of Ni and Fe-Ni alloy nanoparticles, respectively. It should be noted that, whilst the initial exsolved particles were metallic (Ni metal or Fe-Ni alloy), these evolved into oxidised particles to yield the most active phases throughout the series of experiments discussed herein. These systems were labelled Ni/p and Fe-Ni/p, to denote Ni and Fe-Ni-based oxide particles formed after the oxidation of respective exsolved particles from a perovskite host (p). We prepared dense pellets of these two materials and exsolved particles at the top surface, to serve as model catalyst systems (Fig. 1). This catalyst microstructure allowed us to quantify various nanoparticle characteristics (e.g. size, population etc.) by using computed micrograph analysis, and link particle characteristics to catalytic and kinetic behaviour.

2 Experimental

2.1 Catalyst Preparation

The perovskite oxide pellets were prepared by a modified solid state synthesis. High purity precursors including lanthanum oxide (La_2O_3) from Pi-Kem (>99.99%), cerium oxide (CeO₂) from Alfa Aesar (>99.99%), titanium oxide (TiO₂) from Alfa Aesar (>99.6%), nickel(II) nitrate hexahydrate (Ni(NO₃)₂·6H₂O) from Acros (>99%) and iron(III) nitrate nonahydrate (Fe(NO₃)₃·9H₂O) from Alfa Aesar (>99.6%) were used in the appropriate stoichiometric ratios. Oxides were dried at different temperatures (TiO₂—300 °C, La₂O₃—800 °C) and weighed while warm. The reaction mixture was quantitatively transferred to a beaker and mixed with acetone and ~0.05 wt% Hypermer KD1 dispersant. An ultrasonic Hielscher UP200S probe was used to break down agglomerates and homogenize the mixture into a fine, stable dispersion. The acetone was then evaporated at room temperature under continuous stirring and the resulting powder was calcined at 1000 °C for 12 h.

The calcined powder was then pressed into 20 mm diameter pellets (~ 1.5 g of powder per pellet) and fired at 1390 °C for 16 h to form the perovskite phase in dense pellet form (relative density > 95%). The sintered pellets had a diameter of 15–16 mm and were polished on the top side to enable particle exsolution at this side only [2]. Polishing was carried out on a Metaserv 2000 polisher, using MetPrep P1200 polishing paper, followed by cloth polishing with MetPrep 6, 3 and 1 µm diamond paste, respectively. The samples were cleaned in between each steps with acetone in an ultrasonic bath. In order to trigger exsolution, the pellets were then reduced under a continuous flow of 5% H₂/Ar (20 mL min⁻¹), at 900 °C, for 30 h, with heating and cooling rates of 5 °C min⁻¹.

2.2 Catalyst Characterisation

Phase purity and crystal structure of the perovskite oxides were confirmed by X-ray diffraction (XRD) by using a PANalytical Empyrean Diffractometer operated in reflection mode. Catalyst morphology was investigated by scanning electron microscopy (SEM), JEOL JSM-6700F before and after catalytic testing. The principle used to map particles and calculated their size distribution and population is described in our previous work [2]. SEM micrographs were converted to binary images based on pixel contrast. From this, the number of particles as well as the diameter of individual particles can be calculated by using dedicated computer software and code as available e.g. through ImageJ



Fig. 1 Model catalyst samples. Schematics of exsolution of metallic nanoparticles at the top surface of dense pellet perovskite oxide samples. Before and after the catalytic tests the pellet is loaded in the

electron microscope to examine the particle characteristics and link them to the observed catalytic and kinetic behavior

Table 1 Particle characteristics for the Ni/p and Fe-Ni/p catalysts, including: average particle size, particle population and pellet area A and exposed metal particle area per catalyst area A_a

	As-prepared (metallic)		After temperature dependence experiments (oxidised form)	
	Ni/p	Fe–Ni/p	Ni/p	Fe–Ni/p
Catalyst area, A (m ²)	2×10^{-4}	2×10^{-4}	2×10^{-4}	2×10^{-4}
Particle size (nm)	31 ± 3	35 ± 3	36 ± 3	50 ± 3
Particle population (μm^{-2})	123 ± 10	65 ± 5	128 ± 10	63 ± 5
Metal area per catalyst area, A_e $(m^2_{metal} m^{-2}_{pellet})$	0.27 ± 0.01	0.12 ± 0.01	0.39 ± 0.01	0.25 ± 0.01
Unit cell parameter, a (nm)	_	_	0.41752	0.83400
Active sites per unit cell, k	-	-	1	2

software. Based on particle size and population, the exposed area of the particles may be calculated as (assuming the particles have hemispheric geometry). The results of this analysis are listed in Table 1.

2.3 Catalytic Testing

A continuous-flow, single-chamber reactor was used for the catalytic tests of the pellet samples. All experiments were conducted at atmospheric pressure. A schematic diagram of the reactor assembly is shown in Fig. 2a. Sample temperature was measured by using a K-type thermocouple, placed in proximity (0.5 cm away) to the catalyst surface. The flow of the feeding gases was controlled by electronic mass flow controllers. The gases used were high purity 20% CO/He, 20% O₂/He, and CP grade He (N5, 99.999% minimum purity) provided by BOC Ltd. The total gas flow rate was kept constant at 1×10^{-4} mol $s^{-1} m^{-2} (150 cm^3 min^{-1})$ measured at the outlet using a Varian digital flow meter (1000 series). The volumetric flow rates are given at normal temperature and pressure (NTP).

The CO₂ mole fraction in the outlet steam was measured by an XTREAM-CO₂ analyser (minimum detectable CO₂ mole fraction was 1 ppm, which corresponds to a minimum measurable rate of 1×10^{-10} mol s⁻¹ with a flow rate of

 1×10^{-4} mol s⁻¹). The production rate of CO₂, r_{CO_2} , expressed in $mol(CO_2)s^{-1}$, is calculated as shown in Eq. 1:

$$r_{\rm CO_2} = y_{\rm CO_2} \cdot \dot{n} \tag{1}$$

where y_{CO_2} is the measured CO₂ mole fraction at the gas outlet, n is the molar flow. The production rate of CO₂ was measured under "gradientless" conditions where the reactor was operated under differential conversion of CO (30% conversion of CO or less).

The effect of temperature on the r_{CO_2} (henceforth referred to as a temperature dependence experiment), was studied by heating the catalysts under inlet gas mixture of 0.5% CO and 0.25%, 1.33% or 3% O₂, from 80 °C up to 460 °C, with 20 °C temperature steps and with a heating rate of 2 °C min⁻¹. At each temperature step, the temperature was held constant until steady-state was achieved, i.e., the r_{CO_2} did not vary by more than $\pm 5\%$ over 60 min. The effect of CO mole fraction on the r_{CO_2} (henceforth referred to as a CO mole fraction dependence experiment) was studied by holding the O_2 inlet mole fraction constant at 0.25, 1.33 and 3%, while the CO inlet mole fraction was varied between 0.5 and 18.9% at 292 °C. The stability of the pellet samples was also investigated as a function of time. The r_{CO_2} was recorded over 170 h using an inlet feed gas mixture of 0.5% CO and 0.25% O₂ at 450 °C. It is worth noting that the same pellet

Fig. 2 Catalytic reactors. Schematics of a the continuous-flow single-chamber reactor and b of the fix packed-bed reactor. The reactor tube is made from quartz (inner diameter: 7 mm) while equal amounts of quartz wool were placed in either side of the powder sample to immobilize the catalyst bed



catalyst was used for the temperature dependence, CO mole fraction dependence and long-term stability studies.

The effect of sulphur poisoning on the r_{CO_2} was studied on powder samples (10 mg). The study was performed in a fix packed-bed reactor (Fig. 2b), by introducing 50 ppm SO_2 a the inlet gas mixture of 0.6% CO and 1% O_2 , at time intervals of 15, 30 and 60 min, at 400 °C for Ni/p and at 480 °C for Fe–Ni/p. The higher temperature was chosen for the Fe-Ni/p catalyst in order to demonstrate the exceptional stability where deactivation due to particle agglomeration is reported in literature for base metal catalysts but still the sulphur poisoning effect is pronounced. For comparison, a commercial 66 ± 5 wt% Ni/Al₂O₃-SO₂ catalyst (Alfa Aesar) was tested under the same inlet gas mixture at 400 °C. Volumetric dilution within the catalyst bed was made by mixing each catalyst powder with Al₂O₃ powder (10 wt% of each catalyst is diluted with Al₂O₃ to get a total weight of 100 mg). It was found that this amount of catalyst sample mixed with Al₂O₃ resulted in a reasonable pressure drop within the bed. The total gas flow rate was kept constant at 3×10^{-4} mol s⁻¹ (450 cm³ min⁻¹, GHSV = 26,000 h⁻¹). The CO_2 mole fraction in the outlet steam was measured by an XTREAM-CO₂ analyser (minimum detectable CO₂ mole fraction was 1 ppm, which corresponds to a minimum measurable rate of 3×10^{-10} mol s⁻¹ with a flow rate of 3×10^{-4} mol s⁻¹). The production rate of CO₂ was measured under "gradientless" conditions where the reactor was operated under differential conversion of 30% CO, hence

the different operating temperatures for the three different powder samples.

Upstream to both reactors, a fixed-bed reactor with alumina powder heated at 90 °C was used to capture possible carbonyl species.

3 Results and Discussion

3.1 Catalyst Microstructure

Figure 3 shows the SEM micrographs of Ni/p and Fe–Ni/p catalysts, in as-prepared state, as well as after temperature dependence catalytic experiments. The insets show the corresponding particle size histogram as measured by image analysis from the SEM micrographs, according to a method described in the Sect. 2 and in our previous work [2]. This method allowed us to calculate additional particle characteristics, such as particle size, population and exposed metal area per catalyst area, as summarised in Table 1. From Fig. 3 and Table 1 it is apparent that both Ni/p and Fe-Ni/p catalysts display similar average particle size, with uniformly distributed particles, although the population of the Fe-Ni particles is roughly half that of the Ni particles. The difference in particle population is, due, in part, to the Fe-Ni/p sample containing roughly half the amount of exsolvable ions as compared to Ni/p, and, in part due to the presence of Sr which is known

Fig. 3 Catalyst microstructure. SEM micrographs of the catalyst surface and corresponding particle size histograms for Ni/p, **a** as-prepared (metallic state) and **b** after temperature dependence study (oxidised state), and for Fe–Ni/p, **c** asprepared (metallic state) and **d** after temperature dependence study (oxidised state)



to limit exsolution (compare the initial stoichiometries, $La_{0.8}Ce_{0.1}Ni_{0.4}Ti_{0.6}O_3$ vs. $La_{0.5}Sr_{0.4}Fe_{0.1}Ni_{0.1}Ti_{0.6}O_3$) [4, 7].

3.2 Catalytic Activity

Figure 4a shows the CO_2 production rate for the CO oxidation reaction, as a function of temperature, at three different inlet gas mixtures, for the Ni/p and Fe-Ni/p catalysts. These rates were measure on cooling down, that is, after particles have undergone oxidation (see Fig. 3; Table 1 for particle microstructure and characteristics). The minimum temperature for a measurable r_{CO_2} was around 260 °C for both samples. Below 240 °C, the r_{CO_2} was below the minimum measurable limit $(5 \times 10^{-6} \text{ mmol} \text{ s}^{-1})$. For all the inlet gas mixtures, and for both catalysts, the activity increased upon increasing temperature. Under stoichiometric inlet feed conditions (0.5% CO, 0.25% O₂) and below 380 °C, the CO oxidation rates are similar for Ni/p and Fe-Ni/p catalysts, while above 400 °C, the Fe–Ni/p catalyst showed slightly higher CO₂ production rates. Interestingly, under oxidising conditions (0.5% CO, 1.33% O₂ or 3% O₂) the Fe-Ni/p catalyst exhibited up to twofold higher catalytic rates at temperatures above 400 °C as compared to the Ni/p catalyst.

However, it should be noted that, as revealed by the microstructural analysis, the Fe–Ni/p sample exhibits roughly half the particle population as compared to Ni/p (see Fig. 3; Table 1). In order to better compare the site activities of Ni/p and Fe–Ni/p catalysts with each other and to the literature, we calculate their respective nTOFs for the CO oxidation reaction. For this, we assume that the active sites are associated with the metal oxide particles since, in

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the absence of exsolved nanoparticles, the samples displayed virtually no activity. In addition, lacking any further mechanistic evidence, we assume that each surface metal atom provides one active site. Furthermore, in order to calculate the nTOFs we combined the kinetic data in Fig. 4 with the particle analysis data obtained from SEM image analysis and summarised in Table 1. Thus, the number of reacted CO molecules per second, $N_{\rm CO}$, and the number of nominal active sites, $N_{\rm as}$, are given by Eqs. (2) and (3), respectively:

$$N_{\rm CO} = r_{\rm CO_2} \cdot N_A \tag{2}$$

$$N_{\rm as} = \frac{\left(A_e \cdot A\right)}{a^2 \cdot k} \tag{3}$$

where r_{CO_2} is the rate of CO₂ production, N_A is Avogadro's number, A is the catalyst area with exsolved nanoparticles in oxidised form (top of the pellet, see Fig. 1), A_e is the exposed metal area per catalyst area (also in oxidised form), *a* is the crystal lattice unit cell parameter of the particles and *k* is the average number of metal sites per unit cell face (see Table 1). The crystalline facets of the particles were considered to be in a (100) termination, thus, for the Ni, "NiO"-type rock-salt structure *k*=1 and for the Fe–Ni spinel oxide, AB₂O₄-type structure *k*=2. The nTOF (in s⁻¹) can then be calculated by:

$$nTOF = N_A \cdot r_{CO_2} \cdot a^2 / (A_e \cdot A \cdot k)$$
(4)

The corresponding nTOF values are plotted against temperature on the right y-axis in Fig. 4a for both catalysts. Consistent with a previous report [2], the nTOF values for both

Fig. 4 Catalytic activity. CO_2 production rates for CO oxidation, for Ni/p and Fe–Ni/p as a function of **a** temperature and **b** CO mole fraction y_{CO} , at 292 °C. Inlet feed gas mixtures were 0.5% CO, with 0.25%, 1.33% and 3.0% O₂ and total gas flow rate of 150 mL min⁻¹. The right y-axis shows corresponding nTOF values calculated based on the characteristics of the particles on oxide form (Table 1)



exsolved catalysts are of the order of several hundred of s⁻¹. Interestingly though, under the same conditions, the nTOF values for Fe–Ni/p are roughly three times higher as compare to Ni/p. This difference largely originates in the fact that the particle population of the Fe–Ni/p sample is roughly half that of the Ni/p sample (see Table 1). These results indicate that the activity of Ni-based exsolved nanoparticles for the CO oxidation reaction can be further enhanced by alloying with other earth-abundant metals such as Fe. The Fe–Ni/p catalysts exhibited nTOF values of up to 1000 s⁻¹ at 460 °C (0.5% CO, 1.33% O₂). This is a few orders of magnitude higher than values reported in literature for base metal and metal oxide nanoparticles [3].

The microstructure of the samples after the temperature dependence experiment is shown in Fig. 3. The nanoparticles expanded and displayed more faceted shapes as a result of undergoing oxidation. For Ni/p, the average particle size increased from ~31 to ~35 nm. The size increase (35 nm/31 nm = 1.13) is similar to what is expected from the lattice expansion of Ni⁰ to NiO following oxidation (4.175 Å/3.523 Å = 1.18). For Fe–Ni/p, the average particle size increased by a larger factor, from ~ 35 to ~ 50 nm (50 nm/35 nm = 1.43). This increase is consistent with a lattice expansion from Fe-rich, Fe-Ni alloys in a body centred cubic structure (e.g. ~ 2.88 Å) to mixed oxides of Fe–Ni (>8.34 Å), which would give a ratio of 1.45 [19]. It is also worth noting that the nanoparticle populations in Ni/p and Fe-Ni/p remain essentially unchanged following the temperature-dependence catalytic tests, indicating that the particles did not agglomerate as reported for other systems prepared by conventional routes [20].

Figure 4b shows the CO₂ production rate (left y-axis) and the corresponding nTOF values (right y-axis) as a function of CO mole fraction (y_{CO}) , for both catalysts, at 292 °C, under three different inlet gas mixtures. Upon increasing the CO mole fraction, both catalysts displayed monotonically increasing catalytic rates. For Ni/p, the corresponding nTOF values varied between 75 and 125 s^{-1} when the mole fraction of O₂ (y_{O_2}) varies from 0.25 to 3% (at $y_{CO} = 18.8\%$). Under the same conditions the oxidized Fe-Ni/p catalyst displayed similar reaction rates but higher nTOF values $(230-370 \text{ s}^{-1})$ as compared to the Ni catalyst. It is worth mentioning that no change in particle characteristics and no carbon formation were detected by SEM (not shown here) after this experiment for either Ni/p and Fe-Ni/p. This is in contrast to base metal/metal oxide nanoparticles prepared by conventional methods which generally agglomerate and/or coke under such experimental conditions [4, 21, 22].

3.3 Catalyst Durability

To investigate the stability of the Fe–Ni system as compared to its Ni variant we measured the catalytic activity of both



Fig. 5 Long term catalytic activity. CO_2 production rates as a function of time at 450 °C over Ni/p and Fe–Ni/p catalysts, for CO oxidation, with inlet gas mixture of 0.6% CO and 1% O_2 and total gas flow rate of 450 mL min⁻¹. The right y-axis shows corresponding nTOF values calculated based on the characteristics of the particles on oxide form (Table 1). The oscillations may be due to temperature and pressure fluctuations in the lab over 24 h periods affecting the volumes of the gasses

systems over 170 h of continuous testing, at 450 °C (Fig. 5). It should be noted that in this test we used the same pellet catalysts used for the temperature dependence test as well as the CO mole fraction dependence test. It can be seen that the catalytic activity of the Ni catalyst degraded over the first 120 h of testing. After this initial loss of activity, the Ni catalyst appeared to stabilise, however an almost twofold drop in the catalytic activity was observed over this period. On the contrary, the catalytic activity of the Fe–Ni/p sample improved over the first 50 h of testing and was maintained over the remaining testing time. It is worth noting that under these testing conditions typical based metal nanostructured catalysts deactivate via agglomeration in a matter of a few hours [23, 24]. However, the Fe-Ni/p exsolved system used in this study outperforms previous base metal systems for oxidation reactions in terms of activity and long-term stability. Our Fe-Ni/p catalyst is able to operate at high temperature for close to 200 h, conditions which are usually not reported in similar studies in literature even though are of practical importance in base metal catalysis [23, 25, 26]. Furthermore, the catalytic activity, in terms of TOFs, even after 170 h of continuous testing is of the order of 600 s⁻¹, which is significantly higher than those generally measured for mixed oxide base metal systems [26, 27].

3.4 Sulphur Tolerance

To investigate the effect of sulphur poisoning, we compared the Ni/p and Fe–Ni/p catalysts against a state-of-the-art Ni/ Al_2O_3/SiO_2 catalyst (henceforth referred as "commercial Ni catalyst") on a weight-by-weight basis (10 mg). Nickelbased catalysts are known to be very sensitive to sulphur



Fig. 6 SO₂ poisoning. nTOF values for CO₂ production during introduction of 50 ppm SO₂ in an inlet gas mixture of 0.6% CO and 1% O₂ at time intervals of 15, 30 and 60 min (grey-marked regions) over **a**

compounds and even a few ppm of sulphur species can cause irreversible poisoning. Although the poisoning mechanism is not well understood, some reports assign the irreversible poisoning effect of sulphur compounds to Ni-based catalysts to the formation of inactive Ni₃S₂ surface species, formation of bulk Ni-S species (at concentrations above 100 ppm) and the S-induced reconstruction of the Ni surface [28, 29]. To study the effect of sulphur poisoning, 50 ppm SO₂ were added sequentially for 15, 30 and 60 min in an inlet gas mixture of 0.6% CO and 1% O₂ and we recorded the nTOFs before, during and after the SO_2 exposure (Fig. 6). We expect that SO_2 at this concentration and temperatures below 500 °C will poison effectively the CO oxidation rates [30, 31]. Prior to poisoning the recorded rates correspond to around 30% CO conversion or less for all tested systems. Upon introducing the SO_2 in the reaction mixture for 15 min, the commercial Ni catalyst showed an instant decrease in nTOF values from 7×10^{-2} to 3×10^{-3} s⁻¹ (~95% decrease) (Fig. 6a). After removing the SO₂ from the feed, the nTOF values did not recover but remained stable at 3×10^{-3} s⁻¹. Under the same conditions, the nTOF values appear to decrease for the Ni/p catalyst from 110 to 18 s⁻¹ (~83% decrease). However, the nTOF values recovered partly, to about 68% (75 s⁻¹) of the initial values, when SO₂ was removed from the feed (Fig. 6b). Interestingly, under the same conditions, for the Fe-Ni/p catalyst, even though the nTOFs decreased by 50% during the 15 min of SO₂ exposure, after removing the SO₂ from the feed, approximately 98% of the initial activity is eventually restored (Fig. 6c). This finding holds great practical importance because it unlocks new areas of application for these systems.

 Ni/Al_2O_3 -SiO₂ at 400 °C, **b** Ni/p at 400 °C and **c** Fe-Ni/p at 480 °C. Total gas flow rate: 450 mL min⁻¹. The nTOF values calculated based on the characteristics of the particles on oxide form (Table 1)

4 Conclusions

In summary, we demonstrated that the catalytic activity of exsolved Ni nanoparticles can be greatly enhanced when Fe is used as an alloy metal. We believe that this superior catalytic activity and long-term stability of the Fe–Ni system is achieved by preserving the initial spatial arrangement and avoiding surface reconstruction in combination with the confined nature of the exsolved nanoparticles. Our findings also demonstrate that alloying and confinement of nanoparticles enhances site activities even under sulphur poisoning environments in base metal systems for CO oxidation reaction at temperature.

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