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REVIEW

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Well-defined tricobalt tetraoxide's critical morphology effect on the structure-reactivity relationship

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This review focuses on exploring the intricate relationship between the catalyst particle size and shape on a nanoscale level and how it affects the performance of reactions. Drawing from decades of research, valuable insights have been gained. Intentionally shaping catalyst particles makes exposing a more significant percentage of reactive facets possible, enabling the control of overactive sites. In this study, the effectiveness of Co_3O_4 nanoparticles (NPs) with nanometric size as a catalyst is examined, with a particular emphasis on the coordination patterns between oxygen and cobalt atoms on the surface of these NPs. Investigating the correlation between the structure and reactivity of the exposed NPs reveals that the form of Co_3O_4 with nanometric size can be modified to tune its catalytic capabilities finely. Morphologydependent nanocatalysis is often attributed to the advantageous exposure of reactive crystal facets accumulating numerous active sites. However, experimental evidences highlight the importance of considering the reorganization of NPs throughout their actions and the potential synergistic effects between nearby reactive and less-active aspects. Despite the significant role played by the atomic structure of Co₃O₄ NPs with nanometric size, limited attention has been given to this aspect due to challenges in high-resolution characterizations. To bridge this gap, this review strongly advocates for a comprehensive understanding of the relationship between the structure and reactivity through real-time observation of individual NPs during the operation. Proposed techniques enable the assessment of dimensions, configuration, and interfacial arrangement, along with the monitoring of structural alterations caused by fluctuating temperature and gaseous conditions. Integrating this live data with spectroscopic methods commonly employed in studying inactive catalysts holds the potential for an enhanced understanding of the fundamental active sites and the dynamic behavior exhibited in catalytic settings.

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1. Introduction

A catalyst's principal role is to facilitate the acceleration of a given process without compromising the reaction's integrity. Pt-group metals such as Ir, Ru, Pt, and Pd are acknowledged for their exceptional catalytic activity.^{1,2} However, their high cost and scarcity on Earth have motivated scholars to investigate cost-effective alternatives with an abundance on the planet.^{3,4} Transition metals including cobalt (Co), Fe, and Ni have surfaced as viable substitutes for catalysts based on platinum group metals.⁵⁻⁸ First-row transition metals such as Co exhibit diverse properties, acting as electron inks or sources, existing in various oxidation states, and participating in electron exchange.9-12 With its three unoccupied d orbitals, cobalt bonds with surface-bound chemical species, enhancing catalytic activity, especially structural flaws such as vacancies near the crystal lattice surface.13 Cobalt-based catalysts including tricobalt tetraoxide (Co₃O₄) have gained significant attention in Europe for their extensive use in energy and environmental industries. The catalytic properties of Co, attributed to its partly filled d orbital $(3d^7)$, allow for facile composite creation by combining it with other elements or supports. Co and Co-based nanostructures have been investigated to enhance the surface area of catalysts, therefore exposing a more significant number of active sites and allowing the selective exposure of the most active catalytic centers. Co's ability to transition between the Co²⁺ and Co³⁺ oxidation states based on redox conditions makes it an ideal reagent complex builder.14

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Cobalt's dual oxidation state takes advantage of surplus electrons during a reaction, demonstrating its adaptability. The spinel crystal structure of Co₃O₄ contributes to its multifunctional semiconductor properties, and Co3O4 nanoparticles (NPs) exhibit direct optical band gaps, making them suitable for visible light photocatalysis.15,16 Co3O4's varied spin states, such as high, low, and intermediate spin, make it intriguing from a fundamental and spintronic perspective.¹⁶ Cobalt's versatility extends to its environmental impact, as demonstrated by Co₃O₄'s ability to oxidize various compounds, including carbon monoxide (CO), volatile organic compounds, sulfur dioxide (SO₂), and hydrocarbons. Co₃O₄ is employed in processes such as three-way catalytic conversion, phenol oxidation, diesel soot oxidation, and clean energy production, such as hydrogen through steam reforming methanol and ethanol.¹⁷⁻²⁰ Additionally, Co₃O₄ serves as a commercial catalyst in the oxidation, hydrogenation, and hydrogenolysis of esters.21

This review examines recent advancements in the shape engineering of Co_3O_4 with nanometric size. It focuses on their catalytic performance, which is influenced by the coordination patterns of oxygen and cobalt atoms on their surface. The analysis encompasses progress in this field, exploring the structure–reactivity relationship concerning exposed NPs. The review concludes with a summary and a perspective on future developments, aiming to inform readers about the potential prospects involving Co_3O_4 -based catalysts.

2. Preparation strategy of the morphological Co_3O_4

Much effort has been dedicated to preparing Co₃O₄ with wellcontrolled shapes, sizes, and crystal structures. Co₃O₄ has been engineered into zero-dimensional (0D) NPs,22 onedimensional (1D) structures such as nanorods (NRs),²³ nanowires,24,25 two-dimensional (2D) nanodiscs, or nanosheets,26-29 three-dimensional (3D) nanocubes (NCs),30,31 and even hierarchical nanoflowers or more complex structures.32-34 Some of the more well-liked methods to form these nanostructures are coprecipitation, ultrasonic spray pyrolysis, thermal decomposition, microwave-assisted, hydrothermal, and solvothermal methods are examples of physical and chemical processes that have been used for the preparation of Co₃O₄ with nanometric size.35-40 NPs enclose outstanding features, such as a simple and economical synthesis method, high surface area, good stability, and uncomplicated recovery. These properties put together more approval than other synthesis strategies of the prepared catalysts. Researchers have tried to prepare Co₃O₄ with nanometric size by different shapes using different methods to obtain a cost-effective, simple procedure, shorter time through an effective manner, and rectify the purity of the synthesized prepared sample. These processes include.

2.1. Coprecipitation

The coprecipitation method is a simple, efficient, and economical method for the mass production of ultrafine nanopowders. Homogeneity, purity, and reactivity of the prepared oxide are the other advantages of this method. This method was used to prepare Co_3O_4 with nanometric size.³⁵ First, $Co(NO_3)_2$ · $6H_2O$ was dissolved in deionized water. Secondly, ammonium oxalate was added to the solution with continuous stirring. The precipitate was then washed with deionized water and dried at room temperature. Finally, it was calcined at 400–500 °C for 3 h. The average size of the obtained NPs was from 40 to 350 nm, and the Co_3O_4 NPs have an average diameter of 100 nm.

2.2. Utrasonic spray pyrolysis

Ultrasonic spray pyrolysis is an efficient, controlled, and versatile synthesis method. It is frequently used to prepare transition metal oxides, particularly Co_3O_4 ,³⁶ through high purity and narrow size distribution.

Three different precursor solutions were prepared by dissolving cobalt acetate, cobalt chloride, or cobalt nitrate in distilled water with a concentration of cobalt salt as 0.5 mol L^{-1} .³⁶ The starting solution was aerosolized using an ultrasonic nebulizer (Omron, model NB-150U) with a frequency of 1.75 MHz. The spray pyrolysis temperature was kept at 750 $^\circ$ C. The obtained powders were collected at the reactor exit. The prepared Co₃O₄ samples from cobalt acetate, cobalt chloride, and cobalt nitrate are denoted as A-Co₃O₄, C-Co₃O₄, and N- Co_3O_4 . According to the X-ray diffraction (XRD) data of A- Co_3O_4 , C-Co₃O₄, and N-Co₃O₄ samples, all the prepared samples adopted a spinel-type cubic structure. The characteristic diffraction peaks are sharp, and no impurities or a second phase were detected, affirming that high-purity Co₃O₄ was obtained. Scanning electron microscopy (SEM) was used to examine the shapes of the A-Co₃O₄, C-Co₃O₄, and N-Co₃O₄ samples. For the A-Co₃O₄ powders, the dimple and wrinkle surface can be observed. C-Co₃O₄ sample has a porous spherical morphology, and microspheres are developed from various closely packed primary particles; moreover, abundant voids are left among adjacent particles. The N-Co3O4 sample has a durian-like shape with a 0.5-3 µm size distribution, suggesting a hollow inner structure.

2.3. Thermal decomposition

The thermal decomposition of metal oxides performed in high boiling point organic solvents and the existence of surfactants are highly relevant. This process is mainly recognized for preparing excellent-quality NPs with small sizes, high crystallinity, and narrow particle size distributions, although the resulting NPs are very stable in organic solvents.⁴¹⁻⁴⁵ Nonetheless, this approach has some associated drawbacks, *e.g.*, it requires preparation at high reaction temperatures, an inert atmosphere, and long processing times, resulting in increased energy and time consumption.

For example, cobalt oxalate was used as a precursor for synthesizing Co_3O_4NRs by thermal decomposition.³⁷ 0.6 g of cobalt oxalate and 5 mL oleylamine (as a surfactant) were placed in a 50 mL two-neck distillation flask and heated up to 140 °C for 1 h. The resulting solution was added to 5 g of triphenylphosphine (as a surfactant) at 240 °C. The black solution was maintained under stirring at 240 °C for 45 min and then cooled to

room temperature. The final sample was washed with ethanol several times to remove the excessive surfactant. Transmission electron microscopy (TEM) was used to verify the size and shape of the prepared samples. The TEM images of Co_3O_4NRs demonstrated that the materials had rod-like shapes. The length of NRs was 400–550 nm, and their diameters were about 20 nm.

2.4. Microwave-assisted methods

Microwave-assisted chemistry is becoming essential in every area of synthetic chemistry since it can boost some competitive advantages over other preparation methods. It could reduce the processing times and enhance the crystallization level of the particles. These advantages of microwave-hydrothermal methods over conventional hydrothermal methods arise from the direct interaction of the microwaves with the ions or molecules in the solution and with the solid phases dispersed in the liquid medium. In effect, it is essential to underline that the efficiency in the conversion capacity of microwave energy into thermal energy is governed by the physics variables: loss tangent, relaxation time, and penetration depth.⁴⁶ Non-aqueous solvents (glycerol, ethylene glycol (EG), propylene glycol) have been frequently used^{47,48} to avoid or minimize the agglomeration process between the particles.

This method produces high yields, simple to operate, and efficient in terms of being environmentally friendly and energyconsuming. Also, it has been extensively applied to prepare inorganic nanostructured materials⁴⁹⁻⁵⁶ with applications, e.g., electrodes,57 humidity sensors,58 or catalytic devices.52 The method's versatility for synthesizing NPs has been especially reported.59 The microwave-assisted hydrothermal route has been developed to prepare Co₃O₄ with NRs' shape.⁴⁰ The method involved two steps: first, NRs of cobalt hydroxide carbonate were prepared by mixing 50 mL of 0.6 M $Co(NO_3)_2 \cdot 6H_2O$ and 2.4 g of $CO(NH_2)_2$ under 500 W microwave irradiated for 3 min. Subsequently, the cobalt hydroxide carbonate NRs were calcined under air at 400 °C for 3 h to fabricate Co₃O₄NRs. After the thermal decomposition of cobalt hydroxide carbonate precursor under 400 °C for three hours, a single phase of well-crystallized Co₃O₄ with the cubic structure was obtained, and no peaks of the other phase were detected, indicating that the sample was of high purity. The as-prepared sample was bamboo-like NRs with a diameter varying from 30 to 60 nm and a length of 100 to 1000 nm.

2.5. Hydrothermal and solvothermal methods

The hydrothermal method is one of the best-used processes for preparing nanomaterials. It is essentially a solution reactionbased approach. To control the shape of the prepared materials, either low-pressure or high-pressure conditions can be used depending on the vapor pressure of the main composition in the reaction. It has numerous advantages over the other conventional methods such as energy saving, simplicity, costeffectiveness, acceleration interaction between solid and species, better nucleation control, higher dispersion, pollutionfree (as the reaction is done in a closed system), higher rate of the reaction, and lower temperature of operation in the presence of a suitable solvent. Also, it provides highly crystalline particles with better control over their size and shape.

The solvothermal process is similar in its technology to the hydrothermal one, as it is carried out in autoclaves at high temperatures and pressure, through just one difference: instead of water, the synthesis is carried out in organic solvents. Co_3O_4 nanostructures with different morphologies (NCs, nanowires, nanobundles, nanoplates (NLs), and nanoflowers) have been prepared,^{38,52} and the experimental details of the preparation of Co_3O_4 nanostructures with different shapes are summarized in Table 1.

Tuning the morphology of Co₃O₄ for the catalytic reaction

Cobalt oxide is used mainly as a catalyst. Co₃O₄ was used as a model in oxidizing CO.60-65 An initial investigation revealed that Co₃O₄ might facilitate the oxidation of CO at temperatures as low as -54 °C. The activity was significantly decreased, however, when the reaction gas included trace amounts of moisture (3-10 ppm), which obscured the active Co³⁺ sites.^{66,67} Cobalt oxide's activity and durability in the CO oxidation process were increased by changing its form from spherical NPs to NRs, demonstrating a solid morphology-dependent impact.64 NR-shaped cobalt hydroxycarbonate was generated by precipitating cobalt acetate with sodium carbonate in EG. As seen in Fig. 1a-c, further calcination at 450 °C in air converted this precursor into rod-shaped Co₃O₄ NR measuring 200-300 nm in length and 10-20 nm in diameter. The CO oxidation method using spherical NPs yielded an initial CO conversion of 30% at -77 °C. However, as the time-on-stream increased, this conversion decreased to around 10% (Fig. 1h). More active and stable than Co₃O₄ NP catalysts, NR catalysts demonstrated 100% CO conversion in the first 6 h and maintained an 80% CO conversion for \sim 12 h after the reaction.

In contrast to the spherical NPs, Co_3O_4 NR demonstrated an approximately one-order-of-magnitude increase in the rate of CO oxidation. At -77 °C, the Co_3O_4 NR reaction rate was 3.91 ×

Table 1 Experimental parameters of the preparation of different shapes of Co_3O_4 nanostructures

Shape	Cobalt salt (mM)	Temperature (°C)	Reaction time (h)	Structure-directing agents	
Nanocubes (NCs)	2	180	12	15 mL of ammonia (6%)	
Nanowires	2	150	5	30 mL ethanol (99.9%) and 3 mmol of urea	
Nanobundles	2	120	12	2 mmol urea	
Nanoplates (NLs)	2	150	15	3 mL NaOH solution (3.25 mM) with 2 mL ammonia (6%)	
Nanoflowers	2	180	12	30 mL ethanol and 15 mL ammonia (6%)	



Fig. 1 Transmission electron microscope (TEM) and high-resolution transmission electron microscope (HRTEM) images of cobalt-based nanostructures: (a, and b) cobalt hydroxide carbonate; (c-f) Co_3O_4 nanorods (NRs), (c) low-magnification bright-field view, and (d-f) high-resolution views at {110}, {1-10}, and {100}; (g) NR morphological illustration. Catalytic performance of Co_3O_4 : (h) CO conversion efficiency over time for Co_3O_4 nanoparticles (NPs) and NRs in a continuous-flow reactor at -77 °C; (i) reaction rate (r_{CO}) vs. CO or O_2 concentrations for Co_3O_4 NRs; (j) Arrhenius plots (based on ref. 64, Copyright 2009, Nature Publishing Group).

 $10^{-6} \text{ mol}_{CO} \text{ g}^{-1} \text{ s}^{-1}$. Conversely, the value of the NPs was just $4.66 \times 10^{-7} \text{ mol}_{CO} \text{ g}^{-1} \text{ s}^{-1}$. The high-resolution transmission electron microscope (HRTEM) analysis revealed that the Co₃O₄ NPs were enclosed by a configuration consisting of eight {111}

and six {001} planes. Conversely, the Co_3O_4 NR preferred to reveal the {110} planes, constituting an estimated 40% of their overall surface area (Fig. 1g). It was found that Co^{3+} species functioned as active sites for CO oxidation on the {110} plane.



Fig. 2 Scanning electron microscopy (SEM) and high-resolution transmission electron microscope (HRTEM) analysis with structural models of Co_3O_4 nanostructures: (a and b) Co_3O_4NS ; (c and d) Co_3O_4NB ; (e and f) Co_3O_4NC . (g) Methane conversion efficiency vs. temperature for Co_3O_4 at a GHSV of 40 000 h⁻¹ (based on ref. 30, Copyright 2008, American Chemical Society).

The performance of Co₃O₄ nanobelts (NBs) and NCs in CO oxidation has been investigated.⁶³ The reaction rate of Co₃O₄ NC, which mostly exposed the ~{001} facets, was 0.62 µmol g⁻¹ s⁻¹, as opposed to the 0.85 µmol g⁻¹ s⁻¹ seen on NBs terminated by the {110} plane. The specific conversion rate indicates

that at 56 °C, Co_3O_4 NB exhibits 1.37 times the activity of CO_3O_4 NCs, demonstrating that the Co_3O_4 NB are significantly more active than Co_3O_4 NC. As shown by these studies, the activation of the surface layer lattice oxygen on the {110} planes is more pronounced in the presence of Co^{3+} species compared to the



Fig. 3 Morphological and catalytic characteristics of Co_3O_4 nanostructures: scanning electron microscopy (SEM) and high-resolution transmission electron microscope (HRTEM) images of (a–d) Co_3O_4NR {110}, (e–h) Co_3O_4NC {100}, and (i–l) Octahedra {111}; (m) phenol oxidation reaction rates using peroxydisulfate and Co_3O_4 at pH 11; (n) comparison of rate constants and Brunauer–Emmett–Teller (BET)-normalized rate constants for different Co_3O_4 facets (adapted from ref. 16, © 2020 Elsevier Ltd).



Fig. 4 Morphological analysis of 3D Co_3O_4 : (A) scanning electron microscopy (SEM) images of (a and b) Co_3O_4NC , (c and d) Co_3O_4 nanoplates (NLs), (e and f) Co_3O_4NN , and (g and h) Co_3O_4NF . (B) Transmission electron microscope (TEM) images with electron diffraction patterns of (a–c) Co_3O_4NC , (d–f) Co_3O_4NLS , (g–i) Co_3O_4NN , and (j–l) Co_3O_4NF . (C) Comparative analysis of Co_3O_4 catalysts in oxone activation for 5-sulfosalicylic acid degradation of (a) Co_3O_4NC , (b) Co_3O_4NLS , (c) Co_3O_4NN , and (d) Co_3O_4NF (adapted from ref. 71, © 2020 Elsevier BV).

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{001} planes. Furthermore, it was shown that Co_3O_4 nanowires (NWs) enclosed in {111} planes and measuring around 3 nm in diameter had a notably increased rate of CO oxidation at 248 °C, amounting to 161.75 μ mol CO g⁻¹ s⁻¹.⁶¹ The enhanced

performance resulted from the increased surface area and the profusion of Co^{3+} cations on the surfaces.

A catalytic study for CO oxidation⁶² indicates that Co_3O_4 NR exposed to {111} planes exhibited enhanced activity at an activation energy of 40 kJ mol⁻¹, whereas Co_3O_4 NLs exposed to the



Fig. 5 Structural and compositional analysis of Co_3O_4 and Co_3S_4 nanostructures: (A) FE-scanning electron microscopy (FE-SEM) image of Co_3O_4NO ; (B) $Co_3S_4@Co_3O_4NO$; (C) Co_3S_4 nanoneedles. (D) Transmission electron microscope (TEM) and high-resolution transmission electron microscope (HRTEM) images of $Co_3S_4@Co_3O_4$. (E) Selected area electron diffraction pattern of $Co_3S_4@Co_3O_4$. (F) EDX elemental mapping of $Co_3S_4@Co_3O_4$. (G) Schematic of heterojunction-assisted $Co_3S_4@Co_3O_4$ for oxygen reduction reaction (ORR) and CO_2 reduction reaction (CRR) (adapted from ref. 72, © 2017 WILEY-VCH Verlag GmbH & Co. KGaA, Weinheim).



Fig. 6 Co_3O_4 catalysts synthesized *via* various methods and their photocatalytic performance: (a) SEM images of coatings prepared by PLD, (b) particle size distribution histogram for PLD coatings, (c) time-dependent photocatalytic degradation of MB using Co_3O_4 NPs assembled coating *via* PLD and cobalt nitrate and (d–g) SEM images respectively of coatings prepared by (d) electroless, (e) electron beam, (f) sol–gel depositions, and powder form; (h) comparative photocatalytic efficiency of powder Co_3O_4 and coatings by different methods (adapted from ref. 73 and 74 © 2012 Elsevier BV).

same planes had superior activity at a reduced activation energy of 21 kJ mol⁻¹. Significant morphology-dependent effects on CO oxidation have been observed, contradicting prior hypotheses to some degree (maybe due to the porous structures amid cracks and interspaces in the Co₃O₄ nanostructures). The formation of Co₃O₄ NS, Co₃O₄ NB, and Co₃O₄ NC by hydrothermal synthesis of a cobalt hydroxide precursor followed by direct thermal breakdown was investigated in kinetic experiments for methane (CH₄) combustion (Fig. 2a-f).³⁰ The specific rates (r_{CH4}) for Co₃O₄ NC (343 °C), Co₃O₄ NB (319 °C), and Co₃O₄ NS (313 °C) were 1.25, 2.28, and 2.72 μ mol g⁻¹, respectively, as shown in Fig. 2g. Additionally, the T_{50} , representing the temperature at which half of the methane conversion occurred, exhibited a decreasing trend in the same sequence. The structural study indicated that the most prevalent planes on Co_3O_4 NS, Co_3O_4 NB, and Co_3O_4 NC were {112}, {110}, and {001}, respectively. Beyond these crystal planes, the methane combustion process persisted in the following order: $\{112\} > \{110\}$ \rangle $\{001\}$. It can be deduced that manipulating the structure of nanostructured cobalt oxides leads to a substantial display of catalytically active sites. This is supported by the enhanced CH₄ combustion activity observed in Co₃O₄ as a nanosheet, which exposes the more reactive {112} planes. The catalytic activity of Co₃O₄ supported on stainless steel wire mesh, produced by the ammonia evaporation process, was investigated with the preferred oxidation (PROX) of CO.⁶⁸ The 500 nm-diameter mesoporous Co_3O_4 nanowires' diameter is 3.4 nm, and they have a Brunauer-Emmett-Teller (BET) surface area of 71 m² g⁻¹. This structured catalytic system is very stable over the whole temperature range of 100–175 °C due to its low-pressure drop and high heat exchange rate; furthermore, its exceptional catalytic activity is twice that of the highest-performing Co_3O_4 catalyst previously documented.

Although PROX was believed to have an active Co^{3^+} site, its mechanism may have been distinct from the low-temperature oxidation of the CO reaction. Researchers⁶⁹ stated that the turnover frequency of Co₃O₄ NC, composed of six 100-facet facets, was 3.5 to 4 times more than that of Co₃O₄ NS, Co₃O₄ NB, and Co₃O₄ NP. Besides reducing Co²⁺ in hydrogen-rich environments, spectroscopic investigations revealed that Co₃O₄ NC's bulk and surface Co³⁺ sites were only modestly stabilized. For selective CO oxidation, the optimum pair Co³⁺/Co²⁺ was used. By using a sequence of Co₃O₄ catalysts, including exposed {111}, {110}, and {100} planes, it was verified that Co³⁺ functioned as the active site. As shown from the linear relationship between the number of Co³⁺ surface areas and the quantity of CO₂ produced,⁷⁰ the 100 facets positively impacted the PROX. Analyses of different Co₃O₄ attributes indicate that the phase



Fig. 7 Microscopic analysis of 5% Pd-doped Co_3O_4NS : (a) scanning electron microscopy (SEM), (b) transmission electron microscope (TEM), and (c) high-resolution transmission electron microscope (HRTEM) images highlighting of PdO {002} and 0.466 nm of Co_3O_4 {111} (reproduced with permission from ref. 75, Copyright 2011, WILEY-VCH Verlag Gmbh& Co). Detailed TEM and scanning transmission electron microscopy (STEM) analysis of Co_3O_4NR catalysts: (d) TEM, (e) HRTEM, and (f) STEM image of Pt atoms singularly dispersed on Co_3O_4NR (reproduced with Permission from ref. 77, Copyright; American Chemical Society).

and surface characteristics, including shape, surface area, and facets, significantly affect the catalytic activity. As seen in Fig. 3a–l,¹⁶ the synthesis of Co_3O_4 , including a variety of Co_3O_4 NR {110}, Co_3O_4 NC {100}, and nano-octahedron {111} (NO) facets has been completed. The catalytic reactivity of Co_3O_4 NR, Co_3O_4 NC, and Co_3O_4 NO was the highest for phenol oxidation by the persulfate (PS) process. Fig. 3m and n demonstrated that the Co_3O_4 NR exhibited the lowest adsorption energy estimated by the density functional theory (DFT). This confirms that PS is more easily activated *via* a non-radical pathway on the Co_3O_4 {110} plane.¹⁶

To degrade 5-sulfosalicylic acid, four distinct 3D Co_3O_4 catalysts were fabricated, each with a unique morphology (Fig. 4): Co_3O_4 NC {111}, Co_3O_4 NLs {110}, Co_3O_4 NNs (nanoneedles, {110}), and Co_3O_4 NFs (nanoflowers, {112}).⁷¹ Primarily, Co_3O_4 NF ({112} facets) is the most beneficial 3D Co_3O_4 catalyst for the oxidation activation to degrade 5-sulfosalicylic acid⁷¹ due to its plentiful Co^{2+} and more reactive surface, in addition to its most excellent surface area (121.1 m² g⁻¹). The core-shell contrast ratio of the as-prepared $Co_3S_4@Co_3O_4$ core-shell octahedron catalyst *via* hydrothermal and post-surface lattice anion exchange is comparatively less than that of the other core-shell structures.⁷² This is because the concentrations of Co_3S_4 and Co_3O_4 are close. The hexagonal shape of the selected

area electron diffraction pattern, as seen in Fig. 5E, corresponds to both the {111} facet exposure and the close-packed hexagonal pattern observed in the inset of Fig. 5D in HRTEM. The lattice spacing of the {220} pattern is 0.33 nm. As seen in Fig. 5G, electrochemical CO₂ reduction reaction (CRR) and oxygen reduction reaction (ORR) were investigated using a core-shell configuration of Co₃O₄ NO coated with a Co₃S₄ surface. A distinctive electronic configuration is bestowed by the heterojunction separating the p-type Co₃O₄ core and the n-type Co₃S₄ shell, enabling both catalytic processes.

To solve the recovery issue and make a reusable, eco-friendly "green" catalyst, the optimum catalyst is Co_3O_4 with nanometric size attached to a particular substrate with solid adhesion. Chemical (sol–gel), physical (pulsed laser deposition, or PLD), and electrochemical (electroless) methods have been used to create coatings that are reconstructed with Co_3O_4 with nanometric size. Fig. 6a and b shows that the Co_3O_4 with nanometric size. Fig. 6a and b shows that the Co_3O_4 NPs generated using the PLD approach without post-annealing treatment have a mixed amorphous–nanocrystalline phase, a tiny average size of 18 nm, a narrow size distribution of $\sigma = 3$ nm, a perfectly spherical form, and allow a degree of accumulation.^{73,74}

In a methylene blue (MB) solution, the activity of a homogeneous catalyst generating Co^{2+} ions was compared to that of a thin coating catalyst constructed with heterogeneous Co_3O_4



Fig. 8 Transmission electron microscope (TEM) images of Co_3O_4NR (a), Co_3O_4NH (d), and Co_3O_4NC (g). High-resolution transmission electron microscope (HRTEM) images of Au/Co₃O₄NR (b, and c), Au/Co₃O₄NH (e, and f), and Au/Co₃O₄NC (h, and i). Source: reprinted with permission from ref. 78, ©2011 Elsevier BV.

with nanometric size. Complete mineralization of MB dye was achieved in 240 min, indicating a far greater degradation rate than the 40% removed by Co^{2+} ions (Fig. 6c). In the same study, researchers⁷³ found that coatings made of assembled Co_3O_4 with nanometric size had a slightly lower catalytic activity but still demonstrated good recycling capability. Fig. 6d–g shows that PLD-deposited Co_3O_4 coatings have the superior photodegradation rate of MB dye when compared to Co_3O_4 coatings made using other processes (*i.e.*, electro-beam deposition, solgel, and electroless) that have almost equal particle-like morphology (Fig. 6h).

4. Co₃O₄-supported metal nanoparticles (NPs)

 Co_3O_4 has been considered an active support for heterogeneous catalysis for a very long time and is distinguished by its solid metal support interactions. As stated, Co_3O_4 with nanometric

size has an evident morphological influence on CH_4 combustion in the following sequence: Co_3O_4 NS, Co_3O_4 NB, and Co_3O_4 NC.³⁰ Despite applying the same quantity of Pd NPs to these materials, the Pd/Co₃O₄ NS catalyst continued to produce the most methane combustion. The PdO {111} and Co_3O_4 NS formed a geometrically advantageous match, particularly on the {112} facet (Fig. 7a–c), which enhanced the solid metal support interactions and subsequently facilitated the activation of C–H bonds.⁷⁵ The number of missing neighbors of a Co_3O_4 unit cell on a plane {112} is five for the NS shape. PdO must be sited in the 5-fold center of the surface of Co_3O_4 NS as a thin discrete film through a matching geometry and strong coordination rather than a top or bridge site.⁷⁶

Due to its low activation barrier of 29.6 kJ mol⁻¹, single Pt atoms attached to Co_3O_4 demonstrate significant catalytic activity in the water gas shift process at 200 °C (turnover frequency = 0.58 mol_{H2} per site_{Pt} per s). The significantly decreased activation energy observed for these individual Pt



Fig. 9 Scanning transmission electron microscopy (STEM) imaging and catalytic performance of Au-doped Co_3O_4 : (a–c) Au particles on Co_3O_4 (001) in Au/ Co_3O_4NC ; (d–f) Au particles on Co_3O_4 (111) in Au/ Co_3O_4 nanoparticles (NPs); (g) catalytic efficiency of Au/ Co_3O_4NPs and Au/ Co_3O_4NC in ethylene glycol (EG) oxidation. Reprinted with Permission from ref. 79, 2021 Royal Society of Chemistry. Performance analysis of Co_3O_4 and Au/ Co_3O_4 in CO oxidation: (h) temperature-dependent catalytic activity for CO oxidation; (i) Arrhenius plots showing rate *vs.* 1/*T* for CO oxidation over Au/ Co_3O_4 ; (j) durability tests at 25 °C and 60 °C with CO conversion rates from 25% to 45%. Reprinted with Permission from ref. 80, 2023 Royal Society of Chemistry.

atoms may indicate that the interaction between the atoms and the 110-faced Co_3O_4 NR substantially customized the chemical environment of the active sites (Fig. 7d–f).⁷⁷

Using straightforward hydrothermal and solvothermal techniques, anion adsorption was employed to deposit gold NPs onto Co₃O₄ materials produced in various forms, including rods, polyhedra, and cubes.78 Au catalysts based on Co3O4 were characterized using TEM and HRTEM. The predicted morphologies of the Co₃O₄ supports are cube-shaped, rodshaped, and polyhedron (NH)-shaped (Fig. 8). Research into the exposed planes of various morphological Co₃O₄ materials has led to the discovery that the morphology of the support plays a crucial role in determining the catalytic activity. Co₃O₄ NR shows {110} planes most of the time on HRTEM, whereas the $\{011\}$ and $\{001\}$ planes are the most prominent on Co₃O₄ NH and Co_3O_4 NC structures, respectively. The {110} plane has the most excellent oxygen vacancies, which are very important for the oxidation of ethylene, in comparison to the $\{011\}$ and $\{001\}$ planes. Consequently, the ethylene conversion rate of 93.7% was achieved by Au/Co_3O_4 NR, demonstrating their exceptional catalytic activity. Ethanol conversion was 85.5% for the Au/ Co₃O₄ NH catalyst. At 0 °C, the ethylene conversion on Au/Co₃O₄ NC was 26.8%, which was the lowest value recorded.

Our prior research⁷⁹ examined the effect of Co_3O_4 crystallization on EG oxidation supports in the form of Co_3O_4 NCs and NLs. As shown in Fig. 9a–c, Au NPs in the Au/Co₃O₄ NCs samples exhibited a quasi-truncated octahedron structure with Au {111} and {100} faces and had an average size of 2.0 nm. As shown by the interplanar distance of 0.29 nm, corresponding to the {220} crystal plane of cubic Co_3O_4 (001) facet. Furthermore,

the inter-planar spacing of 0.46 nm corresponds to the lattice fringes seen in Au/Co₃O₄ NL catalysts and is caused by the Co₃O₄{111} facets of Co₃O₄ NL oxides. The uniform loading of Au particles onto the Co_3O_4 {111} facet resulted in the formation of a quasi-truncated octahedron encircled by Au {111} and {100} facets, as seen in Fig. 9d-f. Under these conditions, the Co₃O₄ NC and Co₃O₄ NL constituents remained dormant during the aerobic oxidation of EG. With the addition of Au NPs, the catalytic activity of EG oxidation processes was substantially enhanced. Therefore, when subjected to glycol oxidation facilitated by intrinsic defects and surface oxygen vacancies, Au/ Co₃O₄ NL {111} exhibited much greater selectivity and catalytic activity than its Au/Co_3O_4 NC {001} counterpart (Fig. 9g). One potential catalyst for the oxidation of EG using Au NPs is Co₃O₄ NL {111}, which facilitates the activation of O_2 via the oxygen vacancies on its surface.

Furthermore, the catalysts Au/Co₃O₄ P were evaluated in the CO oxidation processes.⁸⁰ The catalytic activity was substantially enhanced by adding Au NPs, as shown in Fig. 9h. This resulted in a noteworthy CO conversion of 35% at 20 °C and complete at 80 °C. As depicted in Fig. 9i, the activation energy (E_a) for CO oxidation in Au/Co₃O₄ P is 15.49 kJ mol⁻¹. Therefore, oxygen molecules follow the Langmuir–Hinshelwood mechanism, which catalyzes CO oxidation at low temperatures (20–60 °C) *via* Au/Co₃O₄ P {111}. In particular, rather than traversing the surface lattice oxygen sites, CO should be adsorbed onto oxygen vacancies at the surface and activated by Au NPs. The durability of the Au/Co₃O₄ P catalysts was also evaluated at temperatures of 25 and 60 °C (Fig. 9j). Throughout the twelve-hours CO oxidation process at 25 °C, the Au/Co₃O₄ P catalyst activity decreased from 2.92 to 1.87 mol_{reactedCO} g_{Au}⁻¹ s⁻¹. A minimum



Fig. 10 Schematic representation of CO oxidation on Co₃O₄. Reprinted with Permission from ref. 82, 2018 American Chemical Society.

activity of 5.26–5.39 mol_{reactedCO} $g_{Au}^{-1} s^{-1}$ was recorded for 9 h at 60 °C. This phenomenon might be primarily attributed to the surface oxygen vacancies and inherent defects of Co₃O₄ {111}, which activated O₂. Similarly, the presence of Au⁰, Au^{δ +}, and Au⁺ species on the surface of Au NPs further enhanced the activation of CO.

5. Chemical nature of the oxide particle morphology

Many people think that certain cobalt cations are abundant at the active sites. Co_3O_4 NR, rich in Co^{3^+} cations and having mostly exposed {110} surfaces, is very active in low-temperature CO oxidation.⁶⁴ Moreover, among Co_3O_4 NR, Co_3O_4 NC, and Co_3O_4 NP, Co_3O_4 NS with mostly exposed {111} planes enriched in Co^{2^+} cations are the most active.³⁸ At low temperatures, a Co_3O_4 SiO₂ nanocomposite devoid of ordered planes but abundant in Co^{2^+} proved an exceptionally active catalyst.⁸¹ However, these findings were mainly obtained *via* catalytic research, and direct spectroscopic evidence of the active surface oxidation state was absent.

Contrary to comparable nanostructures, there have been consistent findings on the shape influence of Co₃O₄ with nanometric size in catalyzing oxidation processes (as shown above). The many reaction routes can contribute, including changing the reaction conditions (primarily the gas and temperature). CO may be oxidized by the Langmuir-Hinshelwood method, which requires surface oxygen species, or the Mars-van Krevelen mechanism, which utilizes lattice oxygen species, according to spectroscopic observations⁸² and the spectroscopically examined possible reaction pathways/ elementary steps of CO oxidation on Co₃O₄ are configured in Fig. 10, the former exhibited dominance at over 100 °C due to oxygen vacancy formation and the Co^{3+}/Co^{2+} redox cycle. Conversely, at lower temperatures, the latter demonstrated dominance. One possible reaction mechanism is that CO adsorbs onto Co3+ cations and then absorbs oxygen from the surface lattice coordinated to three Co³⁺ cations. The oxygen vacancy is then filled with oxygen from the gas phase, according to the Mars-van Krevelen mechanism.64

Spectroscopic evidence is lacking, although an interaction between molecularly adsorbed CO and O–O peroxo species has been postulated by analyzing the impact of pretreatment conditions,⁶⁵ although no peroxo O–O species were found using *in situ* Raman spectroscopy.⁸¹ According to *in situ* infrared research, CO adsorbed on Co²⁺ sites interacted with an oxygen atom bound to a nearby Co³⁺ cation, and the gas phase oxygen was used to fill the oxygen vacancy.⁸³ Isotopes are vital in the redox Mars–van Krevelen process and are responsible for CO oxidation.^{84,85}

Theoretical investigations into the CO oxidation pathway on Co_3O_4 have also shown differences.^{86–89} For instance, a Marsvan Krevelen process involving mostly exposed {110} planes in Co_3O_4 has been proposed, as shown in Fig. 11.⁸⁸

Theoretically, the octahedrally coordinated Co^{2+} site in CoO^{90} would be the most active site for the PROX of CO in the



Fig. 11 Three adsorption configurations of CO on $Co_3O_4(110)$: (a) on O^{2f} ; (b) on O^{3f} ; (c) on Co. Bond lengths are in angstroms; bond angle are in degrees. Co, green, O, blue, and C, red. Reprinted with Permission from ref. 88, 2011 Royal Society of Chemistry.

hydrogen-rich stream. According to DFT calculations, the generated carbonates should make the $\{001\}$ facet of Co_3O_4 less reactive by blocking the surface sites on that facet but not on CoO $\{001\}$, as shown in Fig. 12.

Surface and lattice oxygen species interact concurrently in the reaction network, making methane oxidation on Co_3O_4 catalysts more difficult. There were three distinct temperature/ conversion phases in the methane oxidation process, identified by the presence or absence of the adsorbed or lattice oxygen and the catalyst's redox state.⁹¹ At temperatures between 300 and 450 °C, the dominating superficial Langmuir–Hinshelwood structure produces a stoichiometric {100} surface on Co_3O_4 NC





Fig. 12 Potential energy diagrams for (a) the hydrogenation of Co_3O_4 {001} and (b) the oxidation of CO to CO_2 on Co_3O_4 {001} and CoO {001}. For each transition state (hollow boxes), reaction barriers are given in kJ mol⁻¹. Selected intermediates are shown as a side view along [110], using the following color codes: black (C), blue (Co), red (O), and white (H). Reprinted with Permission from ref. 90, 2019 American Chemical Society.

with a regular size of around 40–60 nm and with the preferential exposure {100}, as previously shown for CH₄ combustion on these particles. At temperatures ranging from 450 to 650 °C, where O₂ nearly occupies the oxygen vacancies generated by the emission of CO₂ and H₂O, the imperfect surface area is delineated by the coexistence of the interfacial (Mars–van Krevelen) and suprafacial (Langmuir–Hinshelwood) mechanisms.^{92,93} At temperatures over 650 °C and with a non-stoichiometric surface area, the completion of oxygen vacancies is only partial, resulting in a substantial reduction in catalyst activity and the combustion of CH₄ *via* the Mars–van Krevelen technique.⁹¹

Theoretical computations have led to the notion that the C-H bond in CH₄ would be activated by the doubly coordinated lattice oxygen (O_{2c}) across the {110} surface. Therefore, the {110} surface is expected to exhibit more activity than the {100} surface, devoid of any O_{2c} sites.94 Assuming dissociation of CH₄ on the Co-O pair; researchers95 indicated that the reactivity of methane combustion increased as follows: $\{001\} < \{011\} < \{112\}$. Experimental observation of cubic Co₃O₄ revealed the less active $\{001\}$ facet, while flower-shaped Co_3O_4 exhibited the active $\{111\}$ facet.96 As compared to spherical NPs enclosed in the {001} and $\{111\}$ facets or Co₃O₄ NRs exposed to the $\{110\}$ and $\{001\}$ facets, Co_3O_4 NLs encased in the {112} facet showed higher activity in the CH₄ combustion process.⁹⁷ The surface remodeling during reaction circumstances may contribute to the contradicting findings on the reactive facets. It has been shown by molecular modeling of Co₃O₄ NPs that the form may be maintained; however, when exposed to oxidizing and reducing atmospheres, the relative ratio of {111}/{100}/{110} facets changes dynamically.98 Under conditions rich in hydrogen gas, the faceting {110} plane was preferentially exposed. At the same time, the {111} surface remained untreated due to the development of oxygen surface vacancies and their subsequent diffusion toward the bulk. Nevertheless, the oxygen-rich conditions promoted the {111} termination. Therefore, it was necessary to describe the shape of the active catalysts. Recent breakthroughs in highresolution microscopic and spectroscopic methods have opened the door to studying the functions of shapedsynchronized NPs in terms of their dynamic performance. Nitric oxide (NO) may be reduced with CO by reshaping Co₃O₄ NRs with an exposed {110} surface into non-stoichiometric CoO_{1-x} NR (Fig. 13a and b).⁹⁹ The structure-modified NRs generate nitrogen gas by selectively reducing nitrogen oxides (NO_x) with CO at temperatures ranging from 250 to 520 °C. Environmental transmission electron microscopy (ETEM) and ambient pressure X-ray photoelectron spectroscopy showed that the non-stoichiometric CoO_{1-x} NRs had a rock-salt (RS) structure. The 100% selectivity was brought about by the active phase, which included around 25% oxygen vacancies. Electron transport microscopy measurements in environments rich in hydrogen showed that CO₃ was reduced to CO, indicating the formation of a boundary contact for particles larger than 15 nm but not for smaller ones, showing that smaller NPs undergo rapid reduction.¹⁰⁰ ETEM identified a two-step phase transition during the heating experiment, as shown in Fig. 13c and d. In the low-temperature range of 200 to 280 °C, the wurtzite (WZ) CoO was spontaneously oxidized to spinel (SP) Co₃O₄ owing to the residual oxygen in the TEM. Secondly, under low oxygen partial pressure conditions, SP Co₃O₄ was reduced to RS CoO at temperatures reaching 280 °C.101 These visual results show that the as-prepared oxide NPs changed significantly under response conditions.



Fig. 13 Structural transformation of Co_3O_4NR : (a) high-resolution transmission electron microscope (HRTEM) image; (b) schematic illustration of Co_3O_4 to CoO transformation under reaction conditions; (c) HRTEM image of CoO hexagonal pyramid; (d) illustration of the phase transformation from metastable wurtzite (WZ) CoO to stable rock-salt (RS) CoO *via* the intermediate spinel (SP) Co_3O_4 . Reprinted with Permission from ref. 90 and 101 Copyrights 2013 and 2019, American Chemical Society.

6. Concluding remarks and perspectives

Extensive exploration into the field of nanocatalysis utilizing Co₃O₄ nanometrics has undeniably demonstrated that the size and shape of the catalyst at the nanoscale level profoundly impact its catalytic effectiveness. A growing body of evidence suggests that the configuration of the nanometric Co₃O₄ is always critical in achieving optimal levels of selectivity, stability, and catalytic activity. This technology's advancement has been significant due to the incorporation of morphology-dependent nanocatalysts, an innovative tool for finely adjusting catalytically active sites. Both theoretical and experimental investigations have been extensive into the morphology-dependent nanocatalysis of nanometric Co₃O₄. Specifically, the arrangement of surface Co³⁺/Co²⁺ and O sites,¹⁰²⁻¹⁰⁴ focusing on the oxygen vacancy, has been linked to the catalytic properties of reactive surface facets. However, there are conflicting reports regarding the effectiveness of similar nanostructures in catalyzing different processes or even the same reaction under identical conditions. This suggests that the form-dependency of nanometric Co₃O₄, as documented, is highly susceptible to variations in reaction parameters and established reaction pathways.

The relationship between the catalytic activities of nanometric Co₃O₄ and the selectively exposed facets induced by shape has been demonstrated through experimental evidence. However, it cannot be ruled out that adjacent facets may work together synergistically. Initially designed nanostructures may undergo structure, morphology, and chemistry changes under actual reaction conditions. The catalytic properties observed in the experiments are determined by the dynamic behavior of the catalyst particles in response to temperature and the reactive environment rather than their state when prepared or recently used. In some instances, the activation of species in a multimolecule chemical reaction may occur through diffusion on adjacent facets, resulting in a synergistic effect where the species activated by the adsorbed reactant can adsorb and stimulate a different type of reactant. In situ studies, physical and chemical analyses, and dynamic characterization techniques must be employed in operational environments to fully understand functional nanostructures.

To gain a deeper understanding of the relationships within nanostructured catalysts, further exploration is needed to develop improved experimental and theoretical methods.¹⁰⁵⁻¹⁰⁷ Variations in temperature and reactive gas fluctuations can impact the well-defined form of Co_3O_4 nanometric, leading to changes in its electrical and geometric properties. This, in turn, influences the proportion of active surfaces and the coordination environments of oxygen and cobalt atoms on the surface, ultimately affecting the development of active sites. The lack of published studies on the atomic structure of nanometric Co_3O_4 can be attributed to the limited availability of high-resolution spectroscopic and microscopic characterizations among researchers worldwide. Also, studying active sites' dynamic performance under operational conditions would provide valuable insights into the structure–reactivity relationship. By employing techniques that allow for real-time assessment of size, shape, interfacial structure, and gas-induced structural changes at the active sites of individual nanoparticles, combined with spectroscopic methods, we can significantly enhance our understanding of the inherent active regions and dynamic capabilities of nanostructured catalysts within catalytic environments.

Abbreviation

BET	Brunauer–Emmett–Teller	
CRR	CO ₂ reduction reaction	
DFT	Density functional theory	
EG	Ethylene glycol	
MB	Methylene blue	
NBs	Nanobelts	
NCs	Nanocubes	
NH	Polyhedron	
NLs	Nanoplates	
NPs	Nanoparticles	
NRs	Nanorods	
ORR	Oxygen reduction reaction	
PLD	Pulsed laser deposition	
PS	Persulfate	
PROX	Preferred oxidation	
SEM	Scanning electron microscopy	
TEM	Transmission electron microscopy	
XRD	X-ray diffraction	

Conflicts of interest

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

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