



2D metal oxide nanoflakes for sensing applications: Review and perspective

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ARTICLE INFO

Keywords:

Two-dimensional
Metal oxide
Nanosheet
Biosensor
Gas sensor
Hierarchical architecture

ABSTRACT

In this review the state of the art and future prospects of 2-dimensional (2D) metal oxide nanoflakes used as active sensing elements for the detection of solutes, gases and radiation are discussed. 2D material geometries are particularly interesting for sensing applications because they provide large specific surface areas and are suitable for crystal facet engineering. In addition, unique material properties of atomically thin nanosheets due to quantum size effects provide engineering possibilities beyond the realm of their bulk counterparts. A variety of possibilities in materials, synthesis routes, (hierarchical) sensor architectures and application areas is sketched. The discussion is focused on high-performing sensors and innovative concepts. The scope is limited to nanoflakes with a thickness of up to 50 nm. Special attention is given to sensing based on material properties that are unique to atomically thin nanosheets.

1. Introduction

Many industries and day-to-day technologies rely on sensors for safety, quality control and analysis. Sensors are required to detect i.e. specific gases, biomolecules or radiation with sufficient sensitivity, selectivity, speed and stability ('4S' requirements). Using 2D sensing elements has several advantages as opposed to using bulk 3D units of the same material. Freestanding 2D nanoflakes have a large specific surface area, which enables extensive interaction with the environment even for small amounts of material. Also, rigid 2D building blocks can facilitate an open packing in 3D assemblies when they are immobilized in a non-stacking fashion. In comparison with 3D architectures derived from 1D or 0D components, 2D-in-3D architectures can provide more efficient electron transport because of fewer grain boundaries and they also tend to have a better mechanical stability [1,2]. Using 2D subunits furthermore allows tuning of the surface reactivity and selectivity via crystal facet engineering [1,3,4]. In addition, unique material properties of atomically thin nanosheets due to quantum size effects, e.g. the increased band gap energy of TiO₂ nanosheets [5] and the visible light absorption of MnO₂ nanosheets [6], provide engineering possibilities beyond the realm of their bulk counterparts. To date, literature reports occasionally describe such unique intrinsic material properties for metal oxide-based sensing, but most often the 2D concept is interpreted only from a geometric viewpoint (increased surface area).

In the past year many reviews have been published on a variety of 2D materials for sensing applications. Zhang et al. [7], Bo et al. [8], Campuzano et al. [9], Zhang et al. [10], Fu et al. [11] and Xu et al. [12]

reviewed 2D graphene-based sensors, mostly employing electrochemical mechanisms and for biological applications. Wang et al. did the same and also included 2D transition metal dichalcogenides, graphite carbon nitride and boron nitride [13]. Bollella et al. furthermore included 2D transition metal oxides [14]. A review focusing on 2D transition metal dichalcogenides for sensors was published by Ping et al. [15]. Yang et al. [16] and Liu et al. [17] published a review focusing on gas sensors, describing various sensor devices and synthetic methods and including 2D graphene-based materials, metal (di)chalcogenides, phosphorene, boron nitride, MXenes and metal oxides. Given the yearly number of publications on 2D metal oxides as sensing elements, indicated in Fig. 1, the present review is devoted particularly to 2D metal oxide sensors. The classification in gas, solute and radiation sensors reflects the available literature and was not used to limit the literature search.

The present review discusses the state of the art and future prospects of 2D metal oxide nanoflakes used as active sensing elements. Most of the publications on 2D metal oxide sensors are proof-of-concept reports demonstrating a variety of possibilities without yet reaching competitive performances. These studies will be summarized shortly to sketch the variety of possibilities in materials, synthesis routes, (hierarchical) sensor architectures and application areas. The discussion is focused on highly sensitive sensors and innovative concepts. Notes on sensor selectivities are included as well, but sensor speed and stability (other '4S' requirements) are left out of scope. Special attention is given to sensing based on material properties that are unique to atomically thin nanosheets. The outline of this review is as follows. In Section 2 a variety

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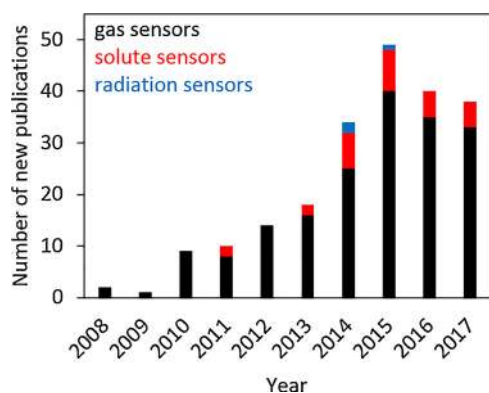


Fig. 1. Indication of the number of new publications per year on sensors with 2D metal oxide nanoflakes of up to 50 nm thick as active sensing elements. The numbers are based on the literature references in Table 1 and 4 and Section 5.

of 2D-based metal oxide architectures and synthesis routes is introduced and described. In Section 3 the use of 2D metal oxide nanoflakes for sensing of solutes is discussed, including sensing mechanisms, used materials and reported sensor performances. The same aspects are discussed for 2D metal oxide nanoflakes for gas sensing in Section 4, with specific attention for rational design approaches. In Section 5 2D metal oxide nanoflakes for sensing of radiation are discussed. In Section 6 a perspective is given on future development and optimization of sensor devices.

2. 2D-based architectures and their synthesis

2.1. Architectures

2D nanoflakes can be employed as active sensing elements both individually and incorporated in 3D architectures. Individual nanoflakes are difficult to handle and are thus mostly employed for sensing in suspended form to detect solutes. Examples of a few-nm thin flexible nanoflake and a stack of rigid nanoflakes of 10–15 nm thickness are

shown in Fig. 2a–b [18,19]. The majority of reports employing 2D nanoflakes as active sensing elements do so by incorporating the nanoflakes in porous 3D architectures. This ensures good exposure of the 2D surfaces to the environment with proper gas diffusion through the assembly. The most common architectures are flower-like structures as shown in Fig. 2c–d [20,21], in which the nanoflakes form a highly porous sphere and are oriented perpendicular to the sphere surface, i.e. emanating from the core. The nanosheets may self-assemble spontaneously or can grow on pre-existing cores. In the tree-like structures of Fig. 2e [22] the nanoflakes are standing on the outer surface of nanorods or tubes (trunks). Nanoflakes vertically standing on a flat substrate form nanowalls as in Fig. 2f [23]. Densely packed nanowalls or randomly oriented nanoflakes can be categorized as nanoforests, shown in Fig. 2g [24]. Films included in this review are not continuous but consist of 2D subunits (crystallites), i.e. layers of nanoflakes that horizontally cover a flat substrate or layers of aggregated nanoflakes without a particular order. The difference between walls, forests and films is somewhat arbitrary. In general it is important that gas molecules can easily diffuse to the active surface, i.e. too densely packed architectures may lose effectiveness.

2D nanoflakes in 3D assemblies generally require a thickness of several nm or more in order to have sufficient mechanical strength to sustain an open architecture under dry conditions. This is usually achieved via bottom-up syntheses and can yield continuous single crystals, but 2D nanoflakes may also consist of 0D [25,26] or 1D [27,28] subunits. Meso- and macropores can be created in the nanoflakes e.g. by a loose connection of 0D subunits [26] or during oxidation or annealing of non-porous precursor nanoflakes [29–31]. Scanning electron microscopy and transmission electron microscopy images of 2D nanoflakes with meso- and macropores are shown in Fig. 3 [2,32]. Porosity facilitates diffusion of target species in the sensing architecture and can increase the surface area.

2.2. Synthesis of 2D nanoflakes

The majority of the 2D nanoflakes is synthesized via hydrothermal or solvothermal procedures. Common advantages of hydrothermal and

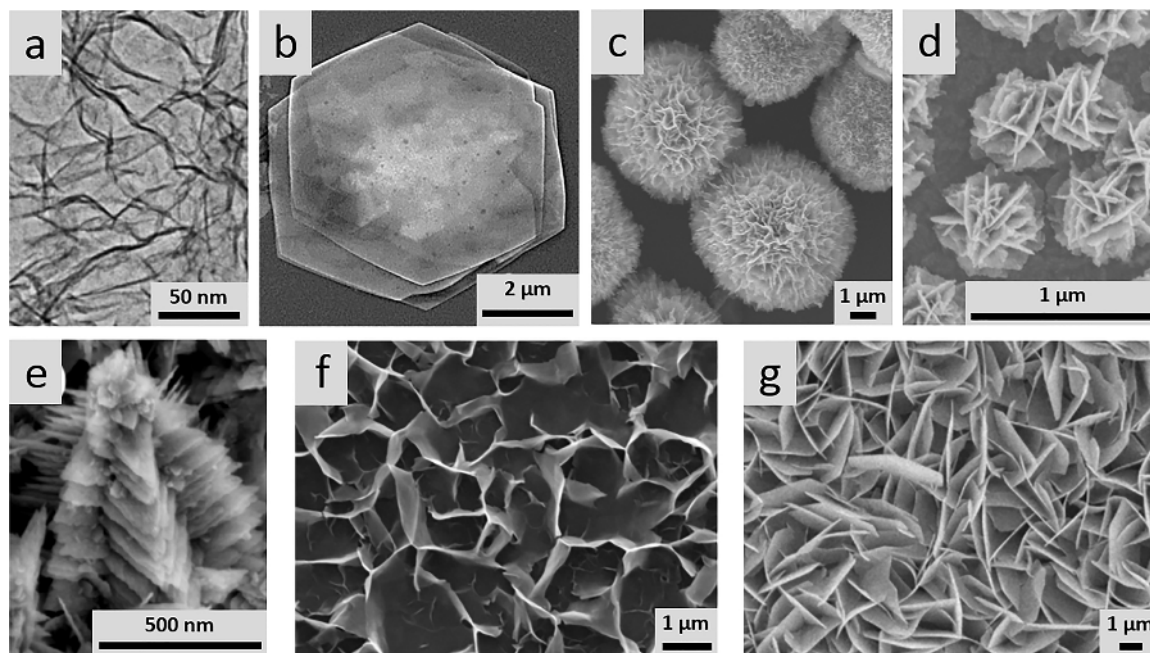


Fig. 2. Transmission electron microscopy image (a) and scanning electron microscopy images (b–g) of individual and assembled nanoflakes. (a) Flexible nanoflake. Reprinted from [18] with permission from Elsevier. (b) Stack of rigid nanoflakes. Reprinted with permission from [19]. Copyright 2014 American Chemical Society. (c,d) Flower-like assemblies. Reprinted from [20,21] with permission from Elsevier and Springer. (e) Tree-like assembly. Reprinted from [22] with permission from Elsevier. (f) Walls. Reprinted from [23] with permission from Elsevier. (g) Forest. Reprinted from [24] with permission from Elsevier.

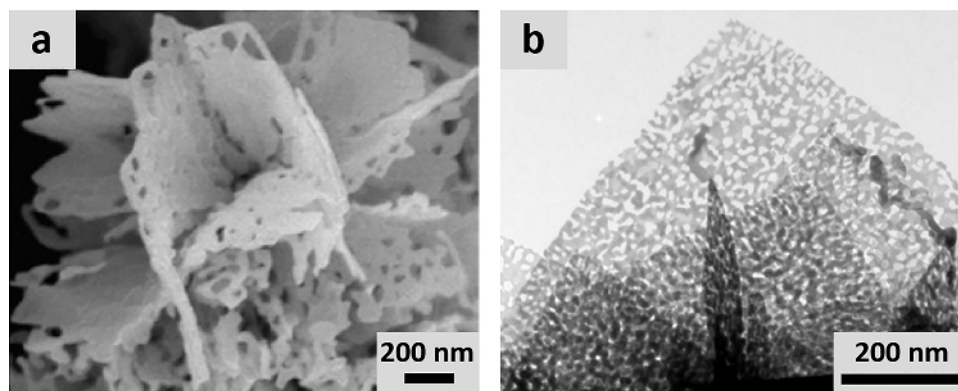


Fig. 3. (a) Scanning electron microscopy image of meso- and macropores in 2D nanoflakes of SnO₂. Reprinted from [32] with permission from Elsevier. (b) Transmission electron microscopy image of mesopores in 2D nanoflakes of SnO₂. Reprinted from [2] with permission from Elsevier.

solvothermal syntheses are simplicity, scalability, low costs, low processing temperatures and compatibility with flexible substrates. Metal oxide precursors are dissolved in water or an organic solvent and the reaction typically takes place at temperatures of 75–200 °C with reaction times of 3–12 h, sometimes extended up to a few days. Common precursors are e.g. metal chlorides, sulfates, nitrates and acetates. For 2D-in-3D hierarchical architectures, the metal oxide may grow in the desired structure spontaneously [33–37] or structure-directing agents such as cetyltrimethylammoniumbromide [38–41], glycerol [39,42], hexamethylenetetramine [23,43–45], polyvinylpyrrolidone [2,40,46,47], urea [25,40,48,49], sodiumdodecylsulfate [50–52] or polyethyleneglycol [52–55] can be added. Also, seed layers may be employed to initiate growth [23,24,43,44,56] and occasionally microwave-assisted procedures are used [57,58]. After the 2D-in-3D architecture is obtained, annealing at 250–500 °C may be required for decomposition of organic residues, dehydroxylation or crystallization. Procedures without annealing are also possible [35,53,59,60]. In some cases the 3D architecture is obtained for sulfide intermediates such as In₂S₃, MoS₂ or SnS₂, which are then converted into the corresponding metal oxide by thermal oxidation [29,61,62]. An uncommon type of hydrothermal fabrication has been reported for ZnSnO₃ structures, involving the hydrothermal dissolution of 3D cubes and subsequent recrystallization into 2D nanoflakes [63].

Alternative synthesis approaches include room-temperature solution procedures, reported for e.g. ZnO [64–66] and MnO₂ [38,67]. Room-temperature sonochemical procedures followed by thermal annealing have been reported for flowers of WO₃ [68] and Fe₂O₃–NiO [69]. The WO₃ flowers were synthesized from Na₂WO₄ in aqueous solution with oxalic acid as capping agent, while the Fe₂O₃–NiO flowers were derived from a coordination polymer precursor with polyethyleneglycol as surfactant. Coordination polymer precursors provide an easy route to multicomponent mixed metal oxides with high composition uniformity [69,70]. Electrodeposition has been reported as a method to prepare Ni flowers that can subsequently be oxidized to NiO [21,71]. ZnO nanoforests and TiO₂ nanowalls have been obtained by oxidation of corresponding metal films [72,73]. 2D metal oxide nanoflakes of only a few nm thickness can be obtained by chemical exfoliation from layered parent compounds, reported for e.g. 1 nm thick RuO₂ [74], 4 nm thick MoO₃ [75–77] and WO₃ [78]. Exfoliation can be assisted by e.g. mild mechanical shaking or sonication and the nanoflakes can be used individually in suspension or be processed into films. More elaborate descriptions of the exfoliation of 2D metal oxides can be found elsewhere [17,79].

The synthesis route determines whether the formed 2D nanoflakes are single-crystalline or polycrystalline and this can profoundly influence the sensor performance. Crystal grain boundaries contain large numbers of surface states that can trap or scatter free charge carriers,

which increases the effective resistivity [80]. Single crystalline nanoflakes tend to have better stability at elevated temperatures due to reduced grain coalescence and resulting alterations of grain boundaries and porosity [81]. Other important synthesis aspects are the obtained crystal orientation and surface morphology, as discussed extensively elsewhere [82]. 2D geometries are particularly suitable for crystal facet engineering.

3. 2D nanoflakes for solute sensing

3.1. Sensing mechanisms

For sensing of solutes, 2D nanoflakes can be immobilized on a support or be used in suspended form. Nanoflakes in suspension can function as individual sensors even if they are only a few atoms thick. Target species in solute sensing are often biomolecules, which can be immobilized onto nanoflakes via e.g. non-specific Van der Waals forces and electrostatic interactions. Sensing based on non-specific interactions is not favorable when analyzing mixtures of biomolecules. In contrast, very high selectivities can be reached for nanoflakes decorated with signaling moieties that interact with the target molecules via e.g. DNA or peptide recognition [38,83,84]. Adsorption-based sensing of solutes often only involves turn-on sensing; once the target molecules are bound by the sensing platform, they do not detach spontaneously anymore. If the nanoflakes are immobilized on a support, regeneration may be possible to enable reuse of the sensor.

Adsorption events can be detected via e.g. fluorescent, plasmonic and/or chemiresistive signals. In fluorescent sensors, light-absorbing metal oxides generally quench the fluorescence of adsorbed signaling species. Upon interaction of the signaling species with the target molecule, e.g. via DNA hybridization, the signaling species (partially) desorbs and the dye label escapes from the quenching influence of the metal oxide nanoflakes, thus retaining its fluorescence [38,83,84]. Alternatively, the fluorescent signaling species can be released via consumption of the metal oxide nanosheets in oxidation reactions with the target species [67,85].

Plasmonic sensors exploit the sensitivity of surface plasmons in metal oxide nanoflakes towards electrical interactions with adsorbed species. The adsorption of charged species causes a change in the free electron density at the surface of the metal oxide, which affects the intensity of plasmonic absorption [75]. Plasmonic sensing particularly benefits from 2D sensing elements, because the existence of one large depolarization factor in 2D geometries reduces the plasmon resonance wavelength as compared to 1D geometries that have two modest depolarization factors [75]. This can shift the resonance wavelength into the visible light and infrared region, which is important for applications as these wavelengths are commonly used in optical systems [75]. A

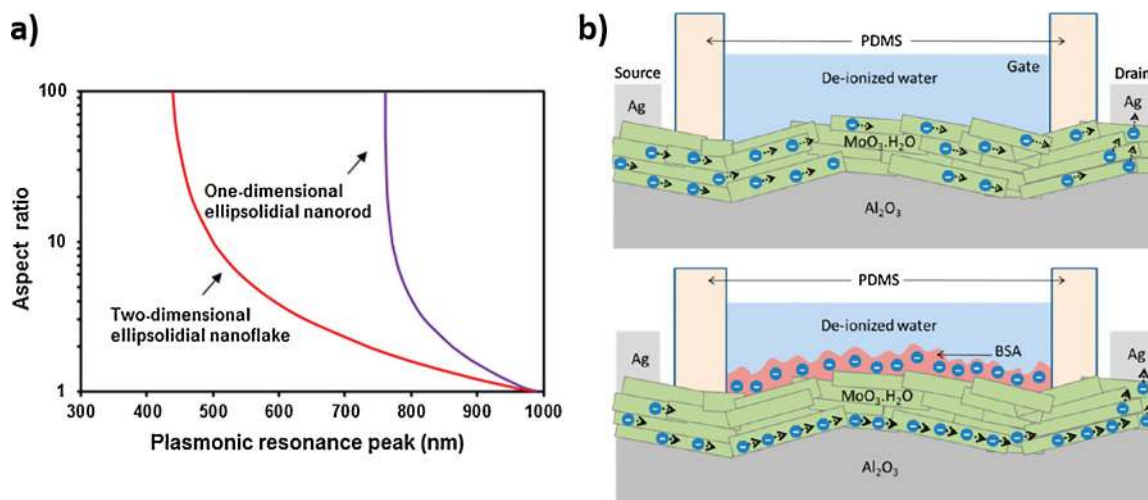


Fig. 4. (a) Theoretical relation between the aspect ratio and plasmonic resonance peak wavelength for 2D and 1D geometries of similar stoichiometry. The relations are based on the Mie-Gans theory, see the supplemental information of [75] for equations and further details. Reprinted with permission from [75]. Copyright 2014 WILEY-VCH Verlag GmbH & Co. (b) Schematic representation of FET sensing of BSA with 2D MoO₃ nanoflakes. Reprinted with permission from [86]. Copyright 2013 American Chemical Society.

schematic representation of the theoretical relation between the aspect ratio and plasmonic resonance peak wavelength for 2D and 1D geometries of similar stoichiometry is shown in Fig. 4a [75].

An example of field-effect transistor (FET) solute sensing employing a film of 2D metal oxide nanoflakes as conduction channel is given in Fig. 4b [86]. The electrical conductivity of the channel varies with the adsorption of target molecules on its surface. In Fig. 4b the adsorption of negatively charged bovine serum albumin (BSA) molecules creates a negative potential on the surface of the n-type semiconductor MoO₃ nanoflake film, which increases the channel resistance.

As an alternative to adsorption-based sensing, solutes may also be detected via electrochemical reactions. Oxidation or reduction of the target species at a nanoflake-covered electrode yields an electric current or potential as response. Electrochemical sensors are especially employed for glucose detection. Drawbacks of commonly used glucose sensors based on enzymes are their poor stability, high cost and weakness to environmental conditions [87], which can be circumvented by non-enzymatic electrochemical sensing.

3.2. Materials

An overview of literature reports on 2D metal oxide nanoflakes that have served as active sensing element for solutes is given in Table 1, indicating the used materials, architectures, syntheses, 2D thicknesses, target species and underlying sensing mechanisms. Individually suspended 2D nanoflakes for sensing generally have a layered structure with a thickness of only a few unit cells. An advantage of layered structures is the possibility of ionic intercalation, which allows very high dopant concentrations (ultradoping) [75]. Ultradoping can increase the free carrier concentration in semiconductors to such an extent that plasmonic resonances shift towards the near-infrared and visible light regions [88]. Metal oxides suitable for plasmonic sensing are e.g. 2D MoO_{3-x} and WO_{3-x}, both of which benefit from having large dielectric constants and are capable of accommodating ultradoping [75,76,89]. For few-nm MoO₃ nanoflakes, irradiation with solar light in the presence of water causes intercalation of the layered structure with H⁺ ions and partially reduces Mo. This results in more oxygen vacancies and causes a transition from semiconducting α -MoO₃ to metallic Mo₄O₁₁. Fig. 5 shows UV/visible light absorption spectra of 2D MoO₃ nanoflakes before and after solar light irradiation, demonstrating the appearance of an absorption band in the near-infrared/visible light region that blue-shifts with increasing irradiation time [75].

Fluorescent sensors require light-absorbing materials such as MoO₃ and MnO₂ [38,67,83,84] to quench the fluorescence of the signaling species. Most 2D transition metal oxides show intensive light absorption due to strong *d-d* transitions [109]. Metal oxides can be beneficial for fluorescent sensing as compared to metal sulfides and selenides because the oxygen atoms yield stronger electrostatic interactions with the signaling species, which causes closer proximity and more efficient quenching [84]. An advantage of atomically thin MnO₂ nanoflakes is that they are also strongly oxidative towards organic compounds and this feature can be employed in combination with their light absorption [67,85].

Electrochemical sensors require metal oxides that are oxidative or reducing towards the target species. Many 2D transition metal oxides have high isoelectric points, which facilitates the immobilization and subsequent charge transfer for a wide range of biomolecules with lower isoelectric points [109,110]. NiO is used for electrochemical glucose sensors because of its low cost, good electrocatalytic activity, biocompatibility and electron transfer capability [25,100]. The absence of poisoning by oxidation products under alkaline conditions enables repeated usage of NiO-based sensors [111]. Other electrochemically active materials are Ni(OH)₂, which is also a precursor for NiO, and CuO and Co₃O₄ [28,90,103,106].

3.3. Performance

3.3.1. Glucose sensors

Most glucose sensors are based on non-enzymatic electrochemical detection with 2D NiO, Ni(OH)₂, CuO and Co₃O₄ nanoflakes, of which the performances are summarized in Table 2. The lowest detection limits (< 1 μ M) and linear response onsets (1 μ M with a window up to several thousand μ M) have been reported for Ni(OH)₂ sensors [101,103,104]. The highest sensitivity has been reported by Cui et al., reaching over 30 mA mM⁻¹ cm⁻² with flower-like structures of NiO on a core of carbon [25]. The addition of urea, ascorbic acid, L-leucine, L-lysine and NaCl did not significantly interfere with the glucose signal. The widest linear response range has been reported by Ngo et al. using NiO flowers on a film of reduced graphene oxide and decorated with Ag particles [100]. The sensitivity was 1.9 mA mM⁻¹ cm⁻² in the linear response region of 50 μ M – 7.5 mM and 116 μ A mM⁻¹ cm⁻² in the linear response region of 10–25 mM. No significant interference was observed from uric acid, ascorbic acid, dopamine, fructose, lactose and sucrose. Solute interference can be suppressed not only by tuning the sensor

Table 1

Overview of literature reports on 2D metal oxide nanoflakes that have served as active sensing element for solutes. Syntheses denoted as ‘hydrothermal’ include at most 50% organic solvent. Syntheses denoted as ‘solution’ include temperatures up to 45 °C.

Reference	2D metal oxide	Architecture	Synthesis	2D thickness [nm]	Target species	Sensing mechanism
[90]	Co ₃ O ₄	aggregate with Ni(OH) ₂	solvothermal	5–15	glucose	electrochemical
[28]	CuO	random film of stacked flakes	solution	20–40	glucose	electrochemical
[38]	MnO ₂	flower decorated with dye-labeled ssDNA	solution	4	ssDNA	fluorescent
[67]	MnO ₂	individual flake decorated with 7-hydroxycoumarin	solution	1	ascorbic acid	fluorescent
[85]	MnO ₂	individual flake decorated with protein-stabilized Au nanoclusters	solution	probably 1	glutathione	fluorescent
[83]	MnO ₂	individual flake decorated with dye-labeled ssDNA or peptide	solution	probably 1	ssDNA (ochratoxin A), cathepsin D	fluorescent
[91]	MnO ₂	individual flake decorated with dye-labeled DNA	solution	1	microRNA	fluorescent
[92]	MnO ₂	individual flake on doped NaYF ₄ nanoparticles	solution	1–3	H ₂ O ₂ , glucose	fluorescent
[93]	MnO ₂	individual flake on doped NaYF ₄ nanoparticles	solution	1–3	glutathione	fluorescent
[94]	MnO ₂	individual flake on doped Sr ₂ MgSi ₂ O ₇ nanoparticles	solution	probably 1–3	glutathione	fluorescent
[95]	MnO ₂	individual flake decorated with C nanoparticles	solution	probably 1–3	glutathione	fluorescent
[96]	MnO ₂	individual flake on phenol formaldehyde resin particles	solution	< 5	glutathione	fluorescent
[97]	MnO ₂	individual flake decorated with Si quantum dots	solution	probably 1–3	glutathione	fluorescent
[18]	MnO ₂	film	solvothermal	5	H ₂ O ₂	electrochemical
[75]	MoO _{3-x}	individual flake	exfoliation	4–6	BSA	plasmonic
[76]	MoO _{3-x}	individual flake	exfoliation	4–26	BSA	plasmonic
[98]	MoO _{3-x}	flower with MoO ₃ surface	solvothermal	20–30	methylene blue	plasmonic
[84]	MoO ₃	individual flake decorated with dye-labeled ssDNA	exfoliation	probably 1–2	ssDNA (prostate-specific antigen)	fluorescent
[86]	MoO ₃	film coated with 1 nm Au	exfoliation	1–3	BSA	FET
[99]	Na ₂ Ti ₃ O ₇	film	hydrothermal	10	Hg ²⁺	electrochemical
[25]	NiO	flower on C core	hydrothermal	30–50	glucose	electrochemical
[100]	NiO	flower on RGO film and decorated with Ag particles	hydrothermal	< 50	glucose	electrochemical
[101]	Ni(OH) ₂	wall [102] on over-oxidized polypyrrole wires	solution	< 50	glucose	electrochemical
[103]	Ni(OH) ₂	flower	solvothermal	5	glucose	electrochemical
[104]	Ni(OH) ₂	hollow flower	hydrothermal	< 50	glucose	electrochemical
[105]	Ni(OH) ₂	flower aggregated with carbon nanotubes	solution (45 °C)	< 50	glucose	electrochemical
[106]	Ni(OH) ₂	forest on Ni foam	hydrothermal	30	glucose	electrochemical
[107]	TiO ₂	flakes and particles decorating RGO	solvothermal	< 5	dopamine	electrochemical
[108]	ZnO	film decorated with antibodies	solution	20	cortisol	electrochemical

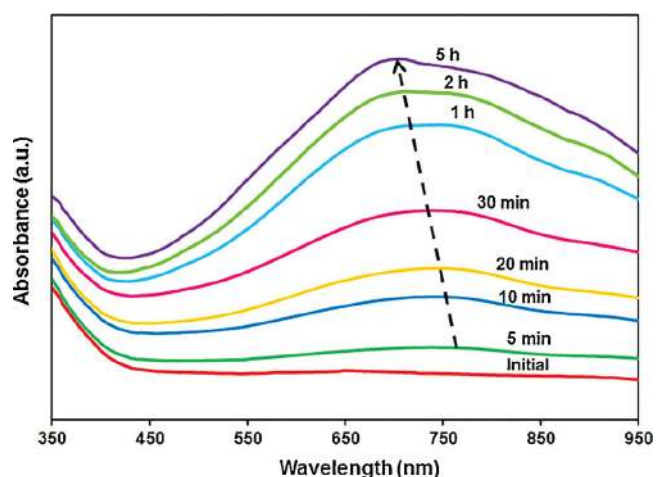


Fig. 5. UV/visible light absorption spectra of 2D MoO₃ nanoflakes before and after solar light irradiation. Reprinted with permission from [75]. Copyright 2014 WILEY-VCH Verlag GmbH & Co.

design, but also by tuning the operation conditions. For example, the operation pH may be chosen such that major interfering species have the same charge as the sensing electrode, i.e. both negative or both positive, and are thus electrostatically repelled [106]. A fluorescent enzymatic glucose sensor employing MnO₂ nanoflakes on doped NaYF₄ upconversion nanoparticles has been reported by Yuan et al. [92]. Glucose was degraded by glucose oxidase, which released H₂O₂ that in turn degraded the MnO₂ nanoflakes and thereby restored the fluorescence of the NaYF₄ nanoparticles.

Table 2

Sensitivity per cm² electrode surface area, lower limit of detection (S/N = 3) and concentration window of linear response for glucose sensors with 2D metal oxide sensing elements.

Reference	2D metal oxide	Sensitivity [μA mM ⁻¹ cm ⁻²]	Lower limit of detection (S/N = 3) [μM]	Linear response window [μM]
[90]	Co ₃ O ₄	1089 / 2145	1.08	5–40 / 40–200
[28]	CuO	26.6 (μA mM ⁻¹)	5	10–7300
[92]	MnO ₂	n.a.	3.7	0–250 / 250–400
[25]	NiO	30190	2	2–1279
[100]	NiO	1869 / 116	2.44	50–7500 / 10000–25000
[101]	Ni(OH) ₂	1049	0.3	1–4863
[103]	Ni(OH) ₂	419	0.08	0.87–10530
[104]	Ni(OH) ₂	223	0.1	0.8749–7781
[105]	Ni(OH) ₂	238.5	0.5	100–1100
[106]	Ni(OH) ₂	2617	2.5	2.5–1050

3.3.2. Glutathione sensors

Glutathione sensors are typically based on composites of fluorescent probes and MnO₂ nanoflakes. The MnO₂ nanoflakes act both as fluorescence quenchers and as oxidizing agents; redox reactions with glutathione consume the nanoflakes and thus restore the fluorescence of the probes. This sensing mechanism only provides turn-on signals, but generally shows good selectivity towards glutathione over other biomolecules and electrolytes. Table 3 lists fluorescent probes, detection limits and concentration windows of linear response reported in literature. All detection limits are well below the typical glutathione concentration in cells (0.5–10 mM [112]). The linear response windows can be tuned by changing the concentration of MnO₂ composites, i.e.

Table 3
Fluorescent probe, lower limit of detection and concentration window of linear response for fluorescent glutathione sensors based on 2D MnO₂ nanoflakes.

Reference	Fluorescent probe	Lower limit of detection [μM]	Linear response window [μM]
[93]	Yb- and Tm-doped core–shell NaYF ₄	0.9	n.d.
[94]	Eu- and Dy-doped Sr ₂ MgSi ₂ O ₇	0.83	0–100
[95]	C nanoparticles	0.022	0.2–600
[85]	BSA-stabilized Au nanoclusters	20	0–500
[96]	phenol formaldehyde resin	0.0076	0–100
[97]	Si quantum dots	0.153	13.3–417

shifting the signal baseline and saturation. Sensing with MnO₂ nanoflakes has been demonstrated in living cells [93,94,97], in human whole blood [95] and in vivo without external excitation [94].

3.3.3. BSA sensors

Bovine serum albumin detection employing individually suspended MoO_{3-x} nanoflakes of 4–6 nm thickness for plasmonic sensing has been reported by Alsaif et al. [75,76]. Initially, response factors of 1.3 and 2.1 were reported for BSA concentrations of 0.15 and 15 mg mL⁻¹, respectively [75]. In following work the response factors were improved to 7 and 24 for BSA concentrations of 0.25 and 10 mg mL⁻¹, respectively [76]. The response factor was found to increase with decreasing lateral flake size and thickness and with decreasing free electron concentration. No data was reported on sensor selectivities. FET-based BSA sensing with drop-cast films of 2D MoO₃ with a 1–1.5 nm Au coating has been reported by Balendhran et al. [86]. The Au coating was added to facilitate charge transfer between film and electrodes and to facilitate the adsorption of BSA. Resistance changes of about 1% and 30% were achieved for concentrations of 1 and 25 mg mL⁻¹, respectively.

3.3.4. Molecular recognition sensors

DNA- or peptide-based molecular recognition of various biomolecules in suspension has been achieved with fluorescent sensors. He et al. have reported the use of MnO₂ nanoflakes of approximately 4 nm thickness that were assembled in flower architectures and decorated with adsorbed dye-labeled ssDNA signaling species [38]. A linear relation between the fluorescence and target molecule concentration was obtained over the range of 0–5 nM with a detection limit of 300 pM. The sensor was demonstrated to be of general applicability by simply changing the signaling probe for molecular recognition. Yuan et al. used single-layer MnO₂ nanoflakes decorated with dye-labeled ssDNA and peptides for the detection of ochratoxin A and cathepsin D, respectively [83]. The response to ochratoxin A was linear in the range of 0.02–2.0 ng mL⁻¹ and was > 1000 times larger than the response to its analogues aflatoxin B1 and B2. For cathepsin D detection a linear response was obtained for concentrations of 1–100 ng mL⁻¹. Yang et al. used single-layer MnO₂ nanoflakes decorated with dye-labeled DNA for the detection of microRNA inside living cells [91]. The MnO₂ nanoflakes not only participated in the fluorescence signaling with a linear response between 0 and 100 pM and a detection limit of 1 pM, but also facilitated cell internalization and protected the DNA label against enzymatic degradation. Dhenadhayalan et al. have reported the ultrasensitive detection of ssDNA with MoO₃ nanoflakes for the application of prostate-specific antigen sensing in live cells [84]. A detection limit of 13 pM was reached.

3.3.5. Other sensors

In vivo detection of ascorbic acid was demonstrated by Zhai et al. using single-layer MnO₂ nanoflakes decorated with fluorescent 7-hydroxycoumarin [67]. The oxidation of ascorbic acid consumed the

MnO₂ nanoflakes and thus restored the probe fluorescence. Though the oxidation is not selective towards ascorbic acid, the contribution of ascorbic acid was determined by reference measurements in which all ascorbic acid was removed by ascorbic acid oxidase. A linear response was obtained in the range of 0.5–40 μM with a detection limit of 0.09 μM and a sensitivity of 26 μM^{-1} . Hu et al. used MnO₂ nanosheets for electrochemical oxidative sensing of H₂O₂ with a sensitivity of 3261 mA M⁻¹ cm⁻² and linear responses between 25 nM – 2 μM and 10–454 μM [18]. The detection limit was 5 nM, which enabled detection of H₂O₂ released by living cells. The previously discussed fluorescent glucose sensor reported by Yuan et al. also detected H₂O₂, yielding a detection limit of 0.9 μM and linear responses between 0 and 150 μM and 180–350 μM [92].

Vabbina et al. have reported the electrochemical detection of cortisol with antibody-decorated ZnO nanoflakes, exploiting the favorable catalytic activity and charge transfer capability of the ZnO (0001) crystal plane [108]. This study is one of few that explicitly includes crystal facet engineering in sensor design. Yuan et al. have reported the electrochemical detection of Hg²⁺ cations via ion exchange with layered Na₂Ti₃O₇ [99]. No interference was observed from Cd²⁺, Mn²⁺, Ni²⁺, Pb²⁺ and Cu²⁺ cations and the detection limit was as low as 0.005 ppb (~ 25 pM). Tan et al. have reported flowers of MoO_{3-x} with an oxidized surface of MoO₃ that were used for the detection of methylene blue via surface-enhanced Raman scattering (SERS) [98]. The plasmonic MoO_{3-x} core electromagnetically increased the Raman scattering, while the non-plasmonic MoO₃ shell prevented the photocatalytic degradation of the analyte. A detection limit of 0.1 μM was reached, which is high for semiconductor SERS substrates, and the enhancement factor was comparable to values observed for noble metals, i.e. 1.42×10^5 . How et al. have reported reduced graphene oxide (RGO) flakes that were decorated with (001)-faceted TiO₂ nanoflakes for the electrochemical detection of dopamine [107]. This study is another example of sensor design based on crystal facet engineering; TiO₂ (001) facets were created because of their higher surface energy and reactivity than TiO₂ (101) facets [113,114]. A linear response was obtained between 2 and 60 μM with a detection limit of 6 μM [107].

4. 2D nanoflakes for gas sensing

4.1. Sensing mechanism

Most metal oxide gas sensors are based on a chemiresistive sensing mechanism. In-depth descriptions of this sensing mechanism can be found elsewhere [115–117], but in short the electrical resistance of a metal oxide sensing unit is related to the ionosorption of oxidative species from the atmosphere on its surface. In oxygen-containing environments O₂ adsorbs on the metal oxide as O₂⁻, O⁻ or O²⁻ and depletes electrons from the metal oxide surface region. This increases the electrical resistance in n-type semiconductor metal oxides and decreases the resistance in p-type semiconductors (hole accumulation). If the target species of a sensor is reducing such as CO, H₂ or various hydrocarbons, the gas molecules are oxidized by the anionic oxygen species at the metal oxide surface and the resulting electrons are injected back into the metal oxide lattice. This again changes the charge carrier density and thus the electrical resistivity of the sensing unit. Operation temperatures of 100–500 °C are generally required to achieve sufficient ionosorption and activation of O₂. If the target species of a sensor is itself oxidative, such as NO₂, no intermediate adsorbents are required and ionosorption of NO₂ directly affects the metal oxide resistivity. Sensing of oxidative species favors lower operation temperatures to suppress competitive O₂ ionosorption. As alternative to this oxygen ionosorption model, chemiresistive sensing can also be described by an oxygen vacancy model [117,118]. This model focuses on changes in the oxygen stoichiometry as a result of reactions between oxygen vacancies and gas molecules. Chemiresistive sensors are not intrinsically selective towards specific target molecules; the electronic

properties of the metal oxide sensing element can be affected by any oxidizing or reducing species and even by merely physisorbed molecules [119].

4.2. Materials

In general, n-type semiconductors provide higher gas responses than p-type semiconductors because of the difference in localization of conductive and resistive regions [115,117]. The ionosorption of oxidative species withdraws electrons from the metal oxide surface and thus creates a resistive electron-depleted shell around the semiconducting core for n-type semiconductors. An applied current will travel primarily through the less resistive core of the particle, but needs to cross a resistive barrier when travelling from one particle to another. This barrier height is sensitive to the number of ionosorbed oxidative species on the metal oxide surface. The sensor sensitivity increases dramatically with the 2D thickness decreasing to below twice the thickness of the electron depletion layer, i.e. with the electrical resistivity of the metal oxide becoming fully surface-dependent. The depletion layer thickness varies with the charge carrier density, band bending and relative permittivity of the metal oxide [120]. For p-type semiconductors the ionosorption of oxidative species creates a conductive hole-accumulated shell around the semiconducting core. Now the applied current will travel primarily along the conductive shell of the particle and is still affected by the amount of ionosorbed surface species, but to a lesser extent than for n-type materials. Also the 2D thickness and the number of interparticle contacts have less influence on the sensitivity of p-type materials. For identical morphologies the response of p-type materials only equals the square root of that of n-type materials [121]. Advantages of p-type materials tend to be their stronger affinity for oxygen, higher catalytic activity and reduced humidity dependence [115,122].

Electronic processes vary significantly with minor changes in chemical composition, stoichiometry and crystal quality. It is generally believed that oxygen vacancies facilitate the formation of oxygen species on the crystal surface and/or offer active sites for gas adsorption [117]. Metal cation defects are important especially in p-type semiconductors, but studies on their role are rare [117,123]. For 2D NiO sensors the sensitivity towards NO₂ has been reported to increase with increasing nickel vacancy concentrations [123]. Although the majority of 2D metal oxide sensors contains only one metal species, multicomponent structures can be obtained via internal doping, external decoration or composite formation. Secondary components can improve the sensor performance e.g. by changing the charge carrier concentration (electronic sensitization) [124], creating *p-n* junctions [125–127] and catalyzing chemical interactions [20] or increasing the adsorbed gas concentration via the spillover effect [128] (chemical sensitization). More information on the physics of multicomponent structures can be found in a review on nanoscale metal oxide-based heterojunctions for gas sensing by Miller et al. [129].

Despite increasing insight, the rational design of gas sensing materials is still hindered by the lack of detailed knowledge on the relation between intrinsic material properties, processing details and their performance. This issue was recently discussed by Zhang et al. in a review on metal oxide semiconductors for gas sensing, sketching the tremendous number of possibilities in material engineering and highlighting efforts on high-throughput material screening for sensor applications [117]. Rational design is especially difficult with respect to the selectivity of chemiresistive gas sensors, which hinders widespread application for the detection of low-concentration and low-reactive targets such as indoor air pollutants (e.g. benzene, xylene, toluene) and breath biomarkers (e.g. H₂S, CO, acetone, NH₃ and NO) [130]. Various examples of suitable catalyst-target gas combinations are given in a review by Woo et al. [130].

An overview of literature reports on 2D metal oxide nanoflakes that have served as active gas sensing element is given in Table 4, listing the

used metal oxides, architectures, syntheses, nanoflake thicknesses and molecules towards which the sensing performance was tested. The differentiation between various metal oxides is schematically shown in Fig. 6; the most widely used metal oxides are ZnO, SnO₂, NiO, WO₃ and In₂O₃. All of these materials have a wide band gap, which is beneficial for reducing background signals due to thermally activated conduction. Other advantages include the high dopability of In₂O₃ [20], the low cost and high chemical and thermal stability of SnO₂ [131] and the large exciton binding energy of ZnO [132,133]. Less commonly used metal oxides include Co₃O₄, MoO₃, CuO, Fe₂O₃, Bi₂WO₆, MnO₂, V₂O₅, Cu₂O, RuO₂, Sn₃O₄, TiO₂, V₄O₉, ZnFe₂O₄, ZnSnO₃ and layered double hydroxides of Mg-Al and Ni-Fe-Al.

4.3. Performance

Among the various types of 2D metal oxide gas sensors, the ones with predominant responses towards NO₂, acetone, ethanol and formaldehyde have been reported most. For each of these gases an overview of reported maximum sensor responses at concentrations up to 200 ppm is given in Fig. 7, together with an indication of which materials are used most (based on the number of publications). Table 5 lists the details of 2D metal oxide sensors with the highest responses for a variety of other target gases. Because of the intrinsically unselective nature of the chemiresistive sensing mechanism (see Section 4.1), many approaches to increase the sensor response towards either NO₂ or acetone, ethanol and formaldehyde will have a qualitatively similar effect on the sensor response towards other oxidizing or reducing target gases, respectively. In this review, the response of gas sensors is defined as the resistance ratio $R_{\text{air}}/R_{\text{gas}}$ or $R_{\text{gas}}/R_{\text{air}}$, depending on the metal oxide (p-type or n-type) and target species (oxidizing or reducing). Literature values based on different definitions were converted to facilitate comparison. Occasionally, reported data is based on a reference atmosphere of N₂ instead of air.

4.3.1. NO₂ sensors

For NO₂ detection the highest responses have been reached with In₂O₃ as 2D sensing materials. Hu et al. have reported In₂O₃ flower architectures that were decorated with PdO particles and reached responses of up to 4080 for 50 ppm NO₂ at the optimal operation temperature of 110 °C [20]. The response towards NO₂ was approximately 2500–4000 times larger than that towards CH₄, H₂, CO and ethanol and gas concentrations down to 500 ppb could be detected. This excellent performance is explained by the catalyzing effect of Pd, which activates the dissociation of both environmental O₂ and targeted NO₂ on the sensor surface. This causes electron depletion in In₂O₃ and thereby increases its electrical resistance. The undoped In₂O₃ sensor reached a maximum NO₂ response of 1310 for 50 ppm, which is still very high as compared to the other metal oxides (see Fig. 7). Another high-performance gas sensor for NO₂ detection has been reported by Liu et al. based on In₂O₃ flower architectures decorated with reduced graphene oxide sheets to create local *p-n* junctions [125]. The highest response was 1337 for 1 ppm NO₂ at 74 °C and the NO₂ response was 20–1000⁺ larger than the response towards NH₃, H₂S, O₃, SO₂, Cl₂, ethanol and acetone. NO₂ detection was achieved down to a concentration of 10 ppb and room temperature responses of up to 1098 were reached for 1 ppm. Xu et al. have reported pure In₂O₃ flowers with porous petals that reached a room temperature response of 1210 for 1 ppm NO₂ with 30% relative humidity and could detect concentrations down to 50 ppb [156]. NO₂ sensors based on WO₃ flower architectures have been reported by Wang et al., reaching responses of 150–250 for 0.8 ppm NO₂ at 90–120 °C while being able to detect concentrations down to 2–40 ppb [216,217].

The reason for the observed superiority of In₂O₃ as compared to other metal oxides for NO₂ detection is unclear. Hu et al. have proposed the high electrical conductance of In₂O₃ as reason for its popularity in gas sensor development [20]. However, the charge carrier

Table 4 Overview of literature reports on 2D metal oxide nanoflakes that have served as active gas sensing element. All sensors are based on the chemiresistive sensing mechanism unless denoted otherwise with a footnote. Syntheses denoted as ‘hydrothermal’ include at most 50% organic solvent. Syntheses denoted as ‘solution’ include temperatures up to 45 °C. The molecules towards which the sensors showed the highest responses are indicated with ‘X’ or listed in underlined text. Other molecules towards which the response was tested are indicated with ‘+’ or listed in plain text.

Reference	2D metal oxide	Architecture	Synthesis	2D thickness [nm]	methanol	ethanol	propanol	butanol	formaldehyde	acetone	benzene	toluene	xylene
[134]	AlOOH	film decorated with Co ₃ O ₄ particles	exfoliation	< 5?	+								
[135]	Bi ₂ WO ₆	flower	hydrothermal	10–20	+	X					+		
[136]	Bi ₂ WO ₆	flower	hydrothermal	< 50							+		
[137]	CdO-ZnO	flower	hydrothermal	20–40	+	X	+			+	+		
[43]	Co ₃ O ₄	wall on carbon foam	hydrothermal	20	+	X				+	+		
[138]	Co ₃ O ₄	flower	hydrothermal	20–25	+	X				+	+		X
[139]	Co ₃ O ₄	flower with hollow core and decorated with Pd particles	solvothermal	13	+	+				+	+	X	X
[140]	Co ₃ O ₄	flower	solvothermal	50		X				X			
[141]	Co ₃ O ₄	film	hydrothermal	20		X				X		+	
[142]	Co ₃ O ₄	film	solvothermal	30	+					X			
[143]	Co ₃ O ₄	film	hydrothermal	20						X			
[144]	Co ₃ O ₄	film	hydrothermal	< 30									
[127]	CoO-SnO ₂	flower	hydrothermal	5–10	+	X	+			+	+		
[145]	SnO ₂	flower	solvothermal	50	+	X				X			+
[146]	CuO	flower	solvothermal	50		X				X			
[147]	CuO	film	hydrothermal	20–25		X				X			
[148]	CuO	film	solution	< 50									
[41]	Cu ₂ O	flower	hydrothermal	5–10									
[149]	Fe ₂ O ₃	rod-shaped flower with hollow core	solution	3	+	X				X			
[150]	Fe ₂ O ₃	flower	solution (40 °C)	40		X				X			
[69]	Fe ₂ O ₃ -NiO	flower	sonochemical	20–30	X	X	X			X	X		
[151]	Fe ₂ O ₃ -ZnO	film	solvo-thermal + atomic layer deposition	30–50		+	+			+			
[20]	In ₂ O ₃	flower decorated with PdO	hydrothermal	10–30		+							
[125]	In ₂ O ₃	flower decorated with RGO sheets	hydrothermal	20–40		+				+			
[152]	In ₂ O ₃	flower doped with Ce	hydrothermal	15–20	X	X	X			X	+	+	+
[51]	In ₂ O ₃	flower decorated/doped with MoO ₃ /Mo	hydrothermal	25	+	X							
[153]	In ₂ O ₃	flower	hydrothermal	20–40	+	X				+	+		
[154]	In ₂ O ₃	flower	hydrothermal	20–40	+	+				+	+	+	
[29]	In ₂ O ₃	flower	hydrothermal + oxidation	10–15	X					+	+	+	
[155]	In ₂ O ₃	flower	solvothermal	20		+							
[156]	In ₂ O ₃	flower	hydrothermal	15–30		+							
[157]	In ₂ O ₃	flower	solvothermal	10	+	X				+			
[27]	In ₂ O ₃	flower	hydrothermal	10–20		+							
[158]	In ₂ O ₃	forest	solvothermal + flame spraying	4		+							
[26]	In ₂ O ₃	random film of stacked flakes	solvothermal	20–40	+	+						+	
[159]	Mg-Al LDH	flower	hydrothermal	20–30									

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Table 4 (continued)

Reference	2D metal oxide	Architecture	Synthesis	2D thickness [nm]	methanol	ethanol	propanol	butanol	formaldehyde	acetone	benzene	toluene	xylene
[160]	MnO ₂	forest decorated with Pd	sputtering	30–40									
[161]	MnO ₂	film	solution	few nm	+	+				+		+	
[47]	MoO ₃	flower	hydrothermal	10–20	X	X							
[162]	MoO ₃	flower	hydrothermal	15–18	X	X							
[61]	MoO ₃	flower	hydrothermal + oxidation	5–10	+	X				+			
[77]	MoO _{3,x}	film coated with 1 nm Au/Pd	exfoliation	4									
[163]	MoO ₃	film	exfoliation	1–2	+	X				+	+		
[164]	MoO ₃	film	exfoliation	1–2									
[165]	Ni-Fe-Al LDH	flower	hydrothermal	20–30									
[23]	NiO	wall on RGO film	hydrothermal	20–30		X							
[40]	NiO	flower	hydrothermal, solvothermal	13–32									
[34]	NiO	flower	solvothermal	30	+	+			X	+	+	+	
[39]	NiO	flower	hydrothermal	15–25	X	X							
[33]	NiO	wall	hydrothermal	14		+				+			
[48]	NiO	flower on RGO sheet	hydrothermal	10–15									
[36]	NiO	flower	solvothermal	10	+	X				+	+	+	
[126]	NiO	tree on ZnO trunk	hydrothermal	30–40	+	+				X			
[132]	NiO	wall with ZnO nanorods	hydrothermal	10–30									
[166]	NiO	flower	hydrothermal	20		X							
[167]	NiO	flower	solvothermal	10–20		X							
[50]	NiO	flower	hydrothermal	10–20		X							
[46]	NiO	flower	hydrothermal	15		X							
[168]	NiO	flower	hydrothermal	15		X							
[21]	NiO	flower	electrodeposition	20	X	X				X			
[58]	NiO	flower decorated with Fe ₂ O ₃	solvothermal	< 5	X	X						+	
[169]	NiO	flower doped with Sn	hydrothermal	< 20	+	+				+	+		X
[45]	NiO	flower with hollow core and W doping	solvothermal	50	+	+				X	+		
[71]	NiO	flower	electrodeposition	20–25		X						X	
[170]	NiO	flower decorated with Fe ₂ O ₃ rods	solvothermal	12	+	+						X	
[171]	NiO	flower	hydrothermal	30	+	X				+			
[172]	NiO	flower	solution	< 5	+	X				+			
[173]	NiO	flower doped with Cr	solvothermal	4–8	+	+						+	X
[174]	NiO	flower with hollow core and decorated with Pd/PdO	solution	< 20		+							
[175]	NiO	film/forest	hydrothermal	30		X							
[176]	NiO	film doped with Al	solvothermal	1–2									
[177]	NiO	film	electrochemical	1–15					X		+	+	
[178]	NiO	film	hydrothermal	20		+				+			
[123]	NiO	film	hydrothermal	< 50									
[179]	NiO-ZnO	flower	hydrothermal	5–10		X							
[180]	NiO-WO ₃	film	hydrothermal	< 40									
[74]	RuO ₂	film	exfoliation	1									
[2]	SnO ₂	flower	hydrothermal	20–30		X							

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Table 4 (continued)

Reference	2D metal oxide	Architecture	Synthesis	2D thickness [nm]	methanol	ethanol	propanol	butanol	formaldehyde	acetone	benzene	toluene	xylene
[181]	SnO ₂	flower	hydrothermal	5–15	+	X			+	+	+	+	
[182]	SnO ₂	flower decorated with PdO	solvothetmal	10					+	+	+	X	
[183]	SnO ₂	flower decorated with Fe ₂ O ₃ particles	hydrothermal	20–40	+	+			X				
[184]	SnO ₂	flower	hydrothermal	20–30		X							
[22]	SnO ₂	tree	hydrothermal	10–20	+	+			X	+	+	+	
[185]	SnO ₂	flower	hydrothermal	10–20		X							
[57]	SnO ₂	flower decorated with Pd particles	hydrothermal	10–20									
[54]	SnO ₂	flower	hydrothermal	30		X							
[186]	SnO ₂	flower doped with Cu	hydrothermal	18	+	+			X				
[187]	SnO ₂	flower	hydrothermal	35–75		+			X				
[188]	SnO ₂	flower	solution	20		X							
[189]	SnO ₂	flower	hydrothermal	20	+	+			X				
[32]	SnO ₂	flower	hydrothermal	20		+			+				
[190]	SnO ₂	flower	hydrothermal	25–62		X							
[191]	SnO ₂	flower	hydrothermal	10–15		X							
[192]	SnO ₂	flower	hydrothermal	10–20	+	X			+	+			
[193]	SnO ₂	flower	hydrothermal, solvothetmal	9		+			+	+			
[194]	SnO ₂	flower doped with Zn	hydrothermal	20		+			+	+	+		+
[195]	SnO ₂	flower decorated with rods	hydrothermal	35		X			X			+	
[55]	SnO ₂	flower	hydrothermal	< 50		X							
[196]	SnO ₂	flower doped with Zn	hydrothermal	30	+	X			+			+	
[197]	SnO ₂	flower	hydrothermal	6–45	+	+		X	+	+	+		
[3]	SnO ₂	flower doped with Sn ²⁺	hydrothermal	20–30		+			+	+			
[198]	SnO ₂	flower	hydrothermal	50		X			+	+	+	+	
[199]	SnO ₂	flower on ZnSnO ₃	hydrothermal	25	+	X							
[200]	SnO ₂	core	hydrothermal	15	+	X			+	+		+	
[62]	SnO ₂	flower	hydrothermal + oxidation	10		X			X				
[131]	SnO ₂	flower doped with Zn	hydrothermal	20–40	+	+			+	+	+		+
[201]	SnO ₂	flower decorated with Au particles	hydrothermal	10–20	+	+	+	X	+	+			
[202]	SnO ₂	film decorated with Au particles	hydrothermal + oxidation	50		X			+	+			
[203]	SnO ₂	forest	hydrothermal	< 20		X							
[204]	SnO ₂	forest	hydrothermal	10		X							
[205]	SnO ₂	film	hydrothermal	15		X							
[206]	SnO ₂	film	hydrothermal	10		X			+	+		+	+
[59]	Sn ₃ O ₄	flower	hydrothermal	8–15	+	X			+	+			
[73]	TiO ₂	wall decorated with SnO ₂ particles	oxidation	20–30		X			X				
[207]	V ₂ O ₅	flower	solvothetmal	10–30	+	+			+	+			
[208]	V ₂ O ₅	film	hydrothermal	2–3					+	+			
[35]	V ₂ O ₅	flower	solvothetmal	1–2 (?)	X ^a								
[37]	WO ₃	flower	hydrothermal	25–40	+	+						X	
[56]	WO ₃	flower	hydrothermal	20–40	+	+		+					
[209]	WO ₃	flower	hydrothermal	10–30		X							

(continued on next page)

Table 4 (continued)

Reference	2D metal oxide	Architecture	Synthesis	2D thickness [nm]	methanol	ethanol	propanol	butanol	formaldehyde	acetone	benzene	toluene	xylene
[210]	WO ₃	flower doped with Co	solvothetmal	20–80		+	+			X			
[211]	WO ₃	flower	hydrothetmal	20–30		X			+				
[212]	WO ₃	flower	hydrothetmal	20		X							
[213]	WO ₃	flower	hydrothetmal	40–80	+	X			+		+		
[214]	WO ₃	flower	hydrothetmal	15–30									
[215]	WO ₃	flower with hollow core	hydrothetmal	50–70		+			+				
[216]	WO ₃	flower	solution (40 °C)	20–30		+			+				
[217]	WO ₃	flower	hydrothetmal	30–60		+			+				
[218]	WO ₃	flower	solvothetmal	10–20		X							
[219]	WO ₃	forest	hydrothetmal	25		X							
[68]	WO ₃	flower	sonochemical	10–20									
[220]	WO ₃	flower	hydrothetmal	< 50	+	+			+		+		
[221]	WO ₃	forest	solvothetmal	6									
[222]	WO ₃	forest	electrochemical	10–50									
[223]	WO ₃	film	hydrothetmal	10		+							
[78]	WO ₃	film	exfoliation	< 10	+	+	+	X					
[224]	WO ₃	film	solution	10–30									
[225]	WO ₃	film	hydrothetmal	10									
[226]	WO ₃	film decorated with Pd particles	solution (30 °C)	2–3									
[42]	ZnFe ₂ O ₄	flower with hollow core	solvothetmal	20	+	+			+	X	+	+	
[60]	ZnO	flower	hydrothetmal	43		X		+					
[49]	ZnO	flower	hydrothetmal	< 50		X			+		+	+	
[30]	ZnO	flower	solution	30–40		+			X		+	+	
[227]	ZnO	flower	hydrothetmal	20–30		X				X			
[52]	ZnO	flower	hydrothetmal	5–10		X			+		+		
[228]	ZnO	flower	hydrothetmal	30	+	+			+	X		+	
[229]	ZnO	flower decorated with rods	hydrothetmal	30–50	+	+			+		+		
[230]	ZnO	flower	hydrothetmal	20–50		X				X	+		
[64]	ZnO	flower	solution	50	+	X	+			+			
[31]	ZnO	flower	hydrothetmal	40–50	+	X							
[231]	ZnO	flower	hydrothetmal	< 50	X	X							
[232]	ZnO	flower	hydrothetmal	10–40		X					+		
[53]	ZnO	flower	hydrothetmal	20–40		X							
[233]	ZnO	flower	solvothetmal (microemulsion-based)	20–30		+			+				
[234]	ZnO	film	hydrothetmal	18		+							
[235]	ZnO	flower and sheet/aggregate	hydrothetmal	50		X			+		+	+	
[65]	ZnO	flower decorated with In ₂ O ₃ particles	solution	30–50					X				
[66]	ZnO	flower decorated with Pd _{0.5} Pd _{0.5} O ₄ particles	solution	30–40	X								
[236]	ZnO	flower decorated with Ag particles and mixed with RGO sheets	hydrothetmal	6									
[236]	ZnO	flower	hydrothetmal	6									
[237]	ZnO	flower	hydrothetmal	5–10?						X			

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Table 4 (continued)

Reference	2D metal oxide	Architecture	Synthesis	2D thickness [nm]	methanol	ethanol	propanol	butanol	formaldehyde	acetone	benzene	toluene	xylene
[238]	ZnO	flower	hydrothermal	10–20		X							
[124]	ZnO	flower doped with Fe	hydrothermal	30					X		+		
[239]	ZnO	flower decorated with Fe ₂ O ₃ rods	hydrothermal	30–40		+			+			+	
[240]	ZnO	flower decorated with Ag particles	hydrothermal	10		X			X				
[241]	ZnO	flower decorated with Au particles	hydrothermal	< 50	X								
[242]	ZnO	flower/aggregate	hydrothermal	20–40	+	X			X			+	
[243]	ZnO	wall	hydrothermal	10–20		X				X			
[244]	ZnO	wall	hydrothermal	10–20		X			+	X			
[245]	ZnO	wall/forest	hydrothermal	20		X		+					
[246]	ZnO	wall	hydrothermal	10–20		X							
[72]	ZnO	forest	evaporation + oxidation	20–30									
[247]	ZnO	forest	solvothermal	10–30	X				X				
[248]	ZnO	forest doped with Sn	hydrothermal	10–20					X				
[249]	ZnO	forest	hydrothermal	50									
[250]	ZnO	forest	hydrothermal	50–100			+				+		
[24]	ZnO	forest	solvothermal	30		X						+	
[251]	ZnO	rod-shaped flower with hollow core	solution	5–16	+	X	+						+
[252]	ZnO	tree	hydrothermal	30–40		X			+	X		+	
[4]	ZnO	film	hydrothermal	tens		X				+		+	
[253]	ZnO	film	solvothermal + oxidation	15–30			+		X				
[133]	ZnO	film decorated with Au particles	hydrothermal	30–60	+			X		+			
[254]	ZnO	film decorated with Pd particles	hydrothermal	15	+					X		+	
[63]	ZnSnO ₃	forest	hydrothermal dissolution-recrystallization	15–20		X			X	X			

Reference	NO ₂	NO	NO _x	NH ₃	CO	H ₂	CH ₄	CO ₂	O ₂	H ₂ S	Other gases
[134]			X	+	+						Other gases
[135]						+					CH ₂ Cl ₂ , C ₆ H _{1,4}
[136]		X									
[137]				+	+						
[43]	+			+	+						ethylene glycol, benzyl alcohol
[138]				+							CHCl ₃ , diethyl ether
[139]	+			+		+					
[140]											
[141]				+	+						
[142]											
[143]					+						
[144]					X		X				
[127]					+						
[145]											
[146]											
[147]											gasoline

(continued on next page)

Table 4 (continued)

Reference	NO ₂	NO	NO _x	NH ₃	CO	H ₂	CH ₄	CO ₂	O ₂	H ₂ S	Other gases
[148]	+			+	+	+			+	X	N ₂
[41]	X			+	+						DMF
[149]				+							
[150]											
[69]											
[151]		+					+				
[20]	X			+	+	+	+			X	Cl ₂ , SO ₂ , O ₃
[125]	X										
[152]				+	+	+					
[51]				+	+	+					
[153]				+	+	+					
[154]				+	+	+	+				LPG, C ₆ H ₁₂
[29]											
[155]	X			+	+					+	SO ₂ , Cl ₂ , O ₃
[156]	X			+	+					+	SO ₂ , Cl ₂ , O ₃
[157]										+	THF, CH ₂ Cl ₂
[27]	X			+	+	+	+			+	O ₃
[158]			X								C ₃ H ₆ O
[26]			X	+	+	+					
[159]				+	+	X				+	THF, ethyl acetate, 1,2-dichloroethane, DMSO, cyclohexane, acetonitrile
[160]				+	+	X					
[161]											
[47]											
[162]											
[61]				+		X ^b					glycol
[77]											
[163]											HCOOH, CHCl ₃ , CH ₃ CHO, C ₄ H ₈ O
[164]						X					
[165]			X								
[23]	X			+		+				+	
[40]											
[34]											
[39]											
[33]				+	+	+				X	(C ₂ H ₅) ₃ N, C ₆ H ₅ Cl,
[48]	X			+						+	ethanoic acid, DMF
[36]											
[126]				+		+				+	
[132]	X			+							
[166]											
[167]											
[50]											
[46]											
[168]											
[21]											
[58]				+							methanamide, dimethylacetamide, DMF
[169]					+		+				
[45]				+	+						
[71]				+							C ₆ H ₅ Cl
[170]											CHCl ₃
[171]											C ₆ H ₁₂ , 1,2-dichloroethane
[172]						+	+				

(continued on next page)

Table 4 (continued)

Reference	NO ₂	NO	NO _x	NH ₃	CO	H ₂	CH ₄	CO ₂	O ₂	H ₂ S	Other gases
[173]											
[174]	X	+		+	+	+					SO ₂
[175]				+	+		+			+	
[176]	X			+	+						H ₂ O
[177]				+		+					
[178]	X				+						
[123]	X										
[179]											
[180]	X	+			+	+					SO ₂
[74]	X										
[2]											
[181]				+							methanal, diethyl ether
[182]				+							
[183]				+	+	+					
[184]							+				
[22]	+			+							
[185]											
[57]					X						
[54]											
[186]				+							acetic acid, DMF
[187]				+							acetic acid, DMF
[188]											
[189]				+							DMF
[32]											acetic acid
[190]											
[191]											
[192]				+							acetic acid, DMF
[193]				+							acetic acid, DMF
[194]				+	+	+		+			glycol, H ₂ O
[195]				+							
[55]											
[196]				+	+						CHCl ₃
[197]											CHCl ₃
[3]											
[198]	X										
[199]					+						CHCl ₃
[200]											butanone
[62]											ethyl ether, CHCl ₃
[131]						+					glycol, H ₂ O
[201]											
[202]											
[203]				+							
[204]											
[205]						+		+			C ₂ H ₄ , C ₃ H ₆ O
[206]										+	butanone
[59]											
[73]						+					
[207]				+							<u>1</u> -butylamine, pyridine
[208]				X							
[35]											H ₂ O ₂
[37]											
[56]										+	SO ₂
[209]		X		+	+						acetylene, ethane
[210]	+										gasoline
[211]					+			+			

(continued on next page)

Table 4 (continued)

Reference	NO ₂	NO	NO _x	NH ₃	CO	H ₂	CH ₄	CO ₂	O ₂	H ₂ S	Other gases
[212]											
[213]											
[214]	X				X						
[215]	X			+	+	+				+	Cl ₂
[216]	X			+	+					+	Cl ₂
[217]	X			+	+						
[218]											
[219]											
[68]	X									X	
[220]						X					
[221]						X					
[222]						+					
[223]	X			+	+						
[78]											
[224]	X										
[225]	X										
[226]						X ^c	+				ethene
[42]					+						
[60]											
[49]											
[30]				+							
[227]											
[52]				+	+	+					
[228]				+	+	+					
[229]				+	+	+	+				
[230]											
[64]											
[31]	+			+	+						
[231]											
[232]					+	+					
[53]											
[233]	X	+			+	+					
[234]					+	+					
[235]				+	+	+	+				
[65]											
[66]											
[236]	+				+	+		+	+		
[236]	+				+	X		+	+		
[237]											
[238]											
[124]				+	+						
[239]											
[240]											
[241]											
[242]											
[243]	X			+		+				X	
[244]	+	+		+		+					
[245]	X										
[246]											
[72]						X					

(continued on next page)

Table 4 (continued)

Reference	NO ₂	NO	NO _x	NH ₃	CO	H ₂	CH ₄	CO ₂	O ₂	H ₂ S	Other gases
[247]		+									C ₆ H ₅ Cl
[248]											H ₂ O
[249]											H ₂ O
[250]	+			+	+	+					SO ₂ , Cl ₂
[24]	+			+	X	+					hexamethylene acetonitrile, C ₆ H ₅ Cl
[251]				+							
[252]											
[4]											
[253]				X							CHCl ₃
[133]											C ₆ H ₅ Cl, acetonitrile
[254]											
[63]											

^a Based on electrochemical sensing.

^b Based on plasmonic sensing.

^c Based on either chemiresistive or gasochromic sensing.

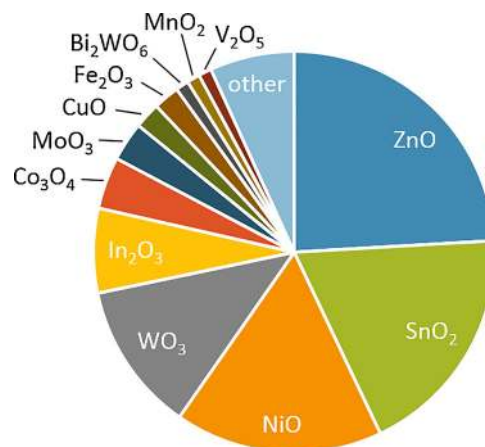


Fig. 6. Differentiation between metal oxides used for the 2D subunits in gas sensing elements (based on Table 4). For composites each individual metal oxide is counted.

concentration and mobility in a semiconductor metal oxide depend massively on crystal defects and doping levels, which suggests that the intrinsic conductance of the metal oxide matrix is of inferior importance. Alternatively, the surface of In₂O₃ may be favorable for interaction with NO₂. The particular response of In₂O₃ extends to other oxidizing gases such as O₃ [255], for which it is known that surface hydroxyl groups promote O₃ dissociation [256].

4.3.2. Acetone sensors

For acetone detection the highest responses have been reached with TiO₂, ZnO and NiO as 2D sensing materials. Maziarz et al. have reported SnO₂-decorated TiO₂ walls with an extraordinary response of 348 to 8 ppm acetone at 308 °C [73]. Though the SnO₂ surface was believed to be the primary sensing surface, TiO₂ donates electrons to increase oxygen ionosorption on SnO₂ and offers rapid electron transport [73,257,258]. Non-decorated TiO₂ walls yielded a response of about 2 towards an acetone concentration of 2 ppm. Xie et al. have reported ZnO flower architectures that reached a response of 362 to 100 ppm acetone at 300 °C and could detect concentrations down to 1 ppm [227]. The response towards acetone was about 12 times larger than the response towards ethanol. Lu et al. have reported the use of 2D NiO nanoflakes to decorate 1D nanorod-assembled flowers of ZnO [126]. The local *p-n* junctions between NiO (p-type) and ZnO (n-type) increase the electron depletion and thus the electrical resistivity in ZnO. A response of 205 for 100 ppm acetone was reached at 240 °C and concentrations were detected down to 10 ppm. The response towards acetone was 2 times larger than that towards ethanol and 4–15 times larger than that towards ethanoic acid, methanol, DMF and NH₃. Wang et al. have reported W-doped NiO flowers with a similar response of 198 for 100 ppm acetone at 250 °C, though the response towards acetone was only about 1.2 times larger than that towards ethanol [45]. Doping with W decreases the hole concentration and thus the conductivity in NiO. Also, an increasing specific surface area was observed as compared to undoped NiO flowers.

4.3.3. Ethanol sensors

For ethanol detection many different materials and sensor designs have been reported. The highest responses were reached with ZnO sensors and with a composite of CoO, SnO and SnO₂. Responses to 100 ppm ethanol of about 340 and 285 have been reported for ZnO flower architectures, respectively, at 370 °C by Xie et al. [227] and at 400 °C by Zhang et al. [232]. In both cases the gas diffusion inside the 3D assembly was facilitated by mesopores in the 2D ZnO nanoflakes and successful detection was demonstrated down to 0.1–1 ppm. Guo has reported a ZnO flower architecture with similarly high performance at low ethanol concentrations, i.e. responses of respectively 17 and 112

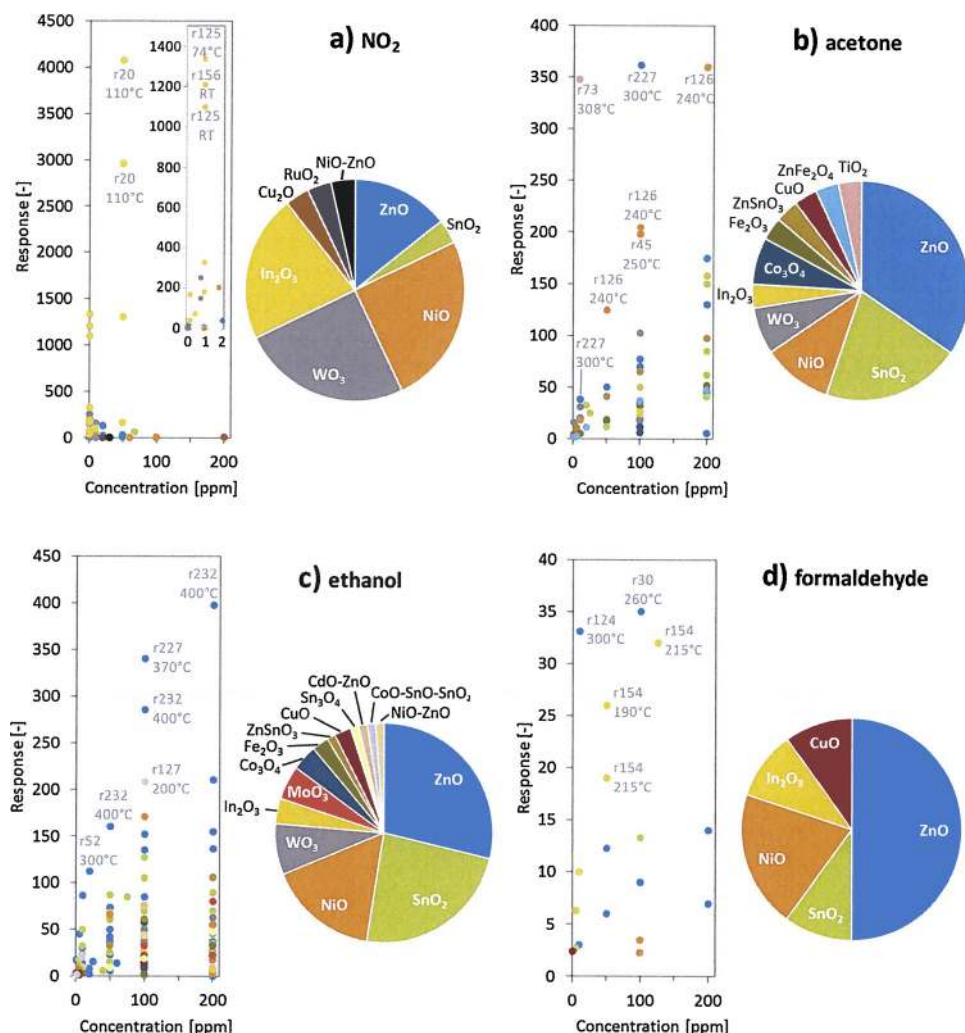


Fig. 7. Overview of reported maximum responses and used materials (based on the number of publications) for 2D metal oxide gas sensors for (a) NO_2 detection with the inset zooming in on low concentrations, (b) acetone detection, (c) ethanol detection and (d) formaldehyde detection. All sensors from Table 4 with selective behavior towards NO_2 , acetone, ethanol and/or formaldehyde are included except [39,69,135,147,185,199].

for 1 and 20 ppm at 300 °C, but for higher concentrations its response leveled off at about 130 [52]. Wang et al. have reported a flower architecture of a composite of CoO, SnO and SnO_2 [127]. The many *p-n* junctions between CoO (*p*-type) and SnO_2 (*n*-type) yield depletion of charge carriers and thus an increasing resistance in the sensing body. Also, CoO was believed to catalyze the dissociation of ethanol molecules on the sensor surface. A Co:Sn ratio of 1 yielded a response of 208 to 100 ppm ethanol, which is lower than the responses of ZnO flowers mentioned above, but an important advantage of the Co/Sn oxide sensor is its significantly lower sensing temperature of 200 °C. Room-temperature ethanol sensing has been reported by Alenezi et al. using ZnO nanodisks that were activated with UV irradiation [4]. Though UV irradiation decreases the overall amount of oxygen adsorbed on the sensor surface, the presence of excitons promotes oxygen adsorption in the more active atomic form (O^-) rather than the less active molecular form (O_2^-). A response of about 1.3 was reached for 200 ppm ethanol under 1.6 mW cm^{-2} UV irradiation.

4.3.4. Formaldehyde sensors

Formaldehyde detection has been reported with the highest responses for ZnO and In_2O_3 sensors. Cao et al. reached a response of 35 for 100 ppm formaldehyde at 260 °C with a ZnO flower architecture [30]. The response towards formaldehyde was 2.3 times larger than the response towards ethanol, 4 times larger than the response towards acetone and > 10 times larger than the response towards NH_3 , toluene

and benzene. ZnO flowers with Fe doping have been reported by Guo, yielding a response of 33 for a formaldehyde concentration of only 10 ppm at 300 °C [124]. Substituting Zn^{2+} in the ZnO lattice with Fe^{3+} causes the release of more electrons in the conduction band. This facilitates the reduction of O_2 at the sensor surface and thereby increases the number of sites where formaldehyde can be oxidized. Also, Fe doping is believed to enhance gas sensing via increasing the oxygen vacancy concentration. Zhang et al. have reported In_2O_3 flowers with mesoporous petals that detected 50 ppm formaldehyde at 190 °C with a response of 26 [154]. At 215 °C the responses were 19 for 50 ppm and 32 for 125 ppm.

4.3.5. Comparison

When considering the performance of various metal oxides with respect to the number of literature reports (see Fig. 3), In_2O_3 has yielded several of the sensors with the highest responses while it represents only 7% of the gas sensor reports included in this review. In terms of sensitivity (note that speed and stability of the sensors are not considered in the present review) In_2O_3 thus appears promising for further development in 2D-based sensing elements, though the precise reason for its success remains unclear (see Section 4.3.1). ZnO has been used for many of the most responding as well as least responding gas sensors, indicating no intrinsic superiority of ZnO as compared to the other metal oxides. As for SnO_2 , it has been used in about 20% of the literature reports on gas sensors included in this review and has been

Table 5
Details of 2D metal oxide sensors with the highest responses for various target gases.

Target species	Concentration [ppm]	Response	Temperature [°C]	2D metal oxide	Reference
methanol	1/5	2.7/8.4	230	NiO (flower)	[21]
methanol	20/100	30/108	255	NiO (flower, porous petals, Fe ₂ O ₃ -decorated)	[58]
butanol	2/100	31/161	300	WO ₃ (film)	[78]
toluene	5	305	250	Co ₃ O ₄ (hollow flower, Pd-decorated)	[139]
xylene	5	361	250	Co ₃ O ₄ (hollow flower, Pd-decorated)	[139]
xylene	5	164	250	Co ₃ O ₄ (hollow flower)	[139]
chlorobenzene	100	6.9	200	ZnO (flower, porous petals)	[238]
acetic acid	50/100/200	15/132/351	340	SnO ₂ (flower, porous petals)	[32]
acetylene	1/200/1/200	10/164/4/22	400/400/285/285	ZnO (aggregated film, porous flakes)	[234]
acetylene	100	12.3	200	ZnO (flower, Ag- and RGO-decorated)	[236]
dimethylamine	1/200	16.8/2087	370	ZnO (flower, porous petals, rod-decorated)	[229]
trimethylamine	20/200	3.9/11.6	260	ZnO (flower, Fe ₂ O ₃ -decorated)	[239]
glycol	5/100	5/90	240	SnO ₂ (flower, Zn-doped)	[194]
CO	10/40	> 25/ > 100	RT	Co ₃ O ₄ (aggregated film, porous flakes)	[144]
CH ₄	10/40	> 25/ > 100	RT	Co ₃ O ₄ (aggregated film, porous flakes)	[144]
H ₂	100	15	200	ZnO (flower)	[236]
H ₂	100	3.3/11.4/20.5	25/100/300	MnO ₂ (forest, Pd-decorated)	[160]
H ₂ S	2.5/50	164/2185	200	ZnO (wall, porous)	[244]
H ₂ O	50%/90% RH	9/345	25	ZnO (wall, porous)	[244]
H ₂ O	60	2	RT	ZnO (forest)	[249]
NH ₃	50/200	4.3/8.4	250	ZnO (film, porous)	[253]
NO _x	0.1/1/10	2/65/213	120	In ₂ O ₃ (forest, porous)	[158]
NO	20	203 ^a	200	WO ₃ (flower, porous rough petals)	[56]

^a For an oxygen-free reference atmosphere.

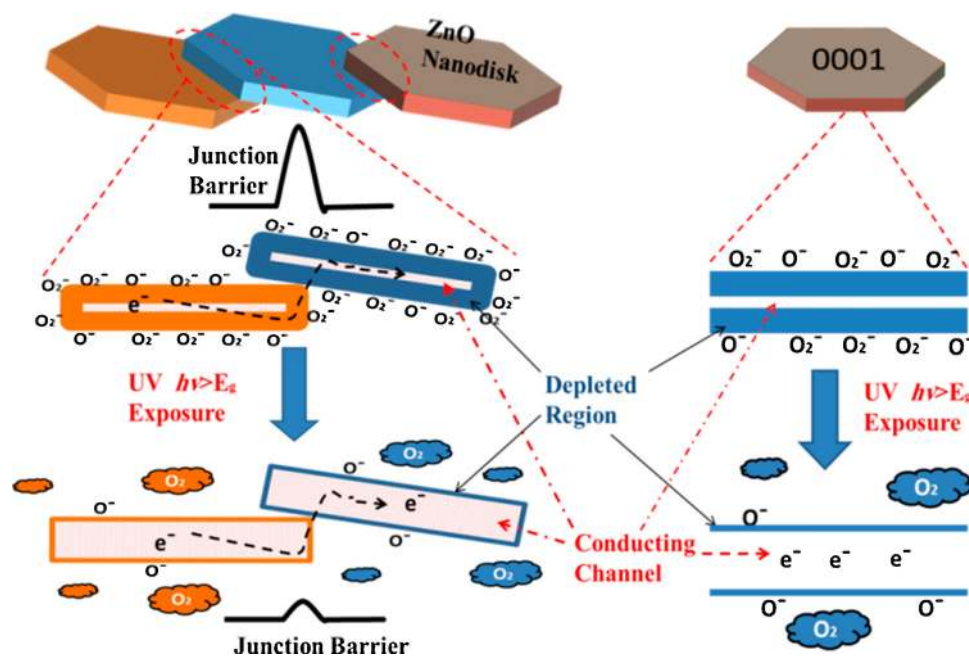


Fig. 8. Schematic representation of the photoelectric sensing mechanism in a stack of ZnO nanodisks and in a single nanodisk. Reprinted with permission from [19]. Copyright 2014 American Chemical Society.

used for the detection of many different target gases, but is almost always outperformed by other metal oxides in terms of response. For selective detection of acetic acid and glycol only SnO₂-based 2D sensors were found.

As for the relation between 2D nanoflake thickness and sensor response, in principle, thinner nanoflakes yield more surface area and thus more active sensing sites per unit volume of metal oxide. However, the used amount of metal oxide is often not of critical importance in sensor design. The active surface area of a sensor can then also be tuned by e.g. changing the dimensions and architecture of the substrate and the packing density of the metal oxide nanoflakes. Since reports on sensor performance generally do not consider the amount of metal

oxide used in a sensor, care must be taken in interpreting correlations between response and nanoflake dimensions or specific surface areas. The various best-performing sensors discussed above have nanoflake thicknesses over the entire size range up to 50 nm (the limit of this review), not indicating a general design rule for sensor optimization.

In addition to the metal oxide loading on a specific device, other fabrication and operation aspects can also significantly influence the sensor performance. The electrical contact depends on the bonding between the sensing material and the device and it may thus be advantageous to grow 2D-based architectures directly onto the sensing device. However, morphological evolution during subsequent processing of the sensor can also be detrimental [259]. As for operation

conditions, sensor responses and selectivities vary with e.g. the gas flow rate [260], the environmental humidity [122] and the operation temperature. Combined signals may be deconvoluted by applying temperature profiles to a single sensor [261] or by using so-called electronic noses, i.e. arrays of different sensors of which the responses are processed with pattern recognition software [262]. Resistive switching, i.e. tuning the metal oxide resistivity by applying a voltage bias, can also be used to increase selectivities and deconvolute combined signals [263].

5. 2D nanoflakes for radiation sensing

Few examples have been reported of 2D nanoflake sensors for detection of radiation. Alenezi et al. have reported on UV sensing with 2D ZnO nanodisks of 10–100 nm thickness, employing a photoelectric sensing mechanism [19]. UV irradiation generates electron-hole pairs in the metal oxide lattice. The holes migrate to the surface and are neutralized by adsorbed oxygen ions, while the electrons increase the conductivity in the metal oxide phase. Sensing was demonstrated with both individual and stacked ZnO nanodisks and the conceptual difference between both cases is schematically indicated in Fig. 8 [19]. Here the junction barriers change because of UV-induced fluctuations in the ionosorbed oxygen population, but the same principle holds for fluctuations of ionosorbed species induced by reactive gases as discussed in Section 4. The tunnel current across the junction barriers between stacked nanodisks is exponentially related to the barrier height and thus highly sensitive to small changes. The sensor response was about 1000 for a single nanodisk and about 1500 for a stack of nanodisks. The single-nanodisk detector showed a photoresponsivity of 3386 A W^{-1} at 3 V. Hoa et al. have prepared a photoresistive sensor with tree-like structures of 1D ZnO nanorods decorated with 10–20 nm thick 2D NiO nanoflakes and thus containing many *p-n* junctions [44]. The response reached over 400% after UV exposure for about 2 min and still increased significantly, yielding a response improvement of 1–2 orders of magnitude as compared to both metal oxides separately. Uthirakumar et al. have increased the UV sensing performance of polymethylmethacrylate films by incorporating 0.2% CuO nanoflakes, resulting in a more than threefold increase of UV absorption capability [264].

6. Perspective

In the field of gas sensors, the concept of using 2D metal oxides is well established. Proof-of-principle sensor devices have been prepared with many different materials via easy and scalable routes and several studies have demonstrated competitive performances. However, further device optimization is needed to enable widespread application in industry and day-to-day life. The geometrical benefits of 2D nanoflakes can be exploited best when they are assembled in 3D hierarchical architectures with moderate packing density to ensure extensive surface exposure. Achieving high selectivities is a difficult challenge with the intrinsically unselective chemiresistive sensing mechanism, but various routes can be explored in addition to the widely investigated approach of chemical sensitization (adding catalysts). For example, illumination of metal oxide sensing elements either above or below their band gap has been proposed to facilitate (selective) desorption of surface species [116]. This increases reversibility and selectivity at relatively low operation temperatures. An alternative approach is to integrate the sensor with a permselective membrane. Kato et al. have reported the use of cross-linked polysiloxane monolayers on top of sensor electrodes to detect NO in solution while preventing interfering species such as nitrite, dopamine, adrenalin and uric acid from reaching the electrode surface [265,266]. Recently, incorporation of zeolites as both filtering and catalytic component has been investigated extensively because it enables discrimination based on molecular size, shape and affinity [117,267]. Membrane layers with somewhat larger pores can induce

discrimination based on molecular weight (exploit Knudsen diffusion) [117]. Finally, exploiting particular surface energy landscapes via crystal facet engineering is a promising road towards higher and fine-tuned sensor selectivities. Though facet engineering has been applied in several cases to promote specific chemical interactions, it is not yet generally adopted as point of consideration.

Solute sensing with 2D metal oxides is still relatively new, but pioneering reports already set the bar in minimizing the detection limit. Sensing in solution is particularly suitable for exploiting mono- or few-layer nanosheets that do not have the rigidity to form and maintain hierarchical assemblies under dry conditions. Keeping these ultrathin crystallites in suspension yields a relatively robust and easy-to-handle system that provides full surface exposure of the nanosheets. Mono- or few-layer crystallites generally provide unsurpassed structural and chemical surface homogeneity, which is favorable for achieving well-defined sensor responses. Also, new sensing strategies may be developed based on quantum confinement effects in ultrathin nanosheets. As for plasmonic sensing, this generally varies with the shape and size of nanoparticles [268] and thus performances may be improved by controlling the size of 2D nanoflakes. A new and still unexplored strategy to modulate the selectivity of 2D sensing elements further is to introduce specific adsorption sites using controlled ion doping strategies. Doping with aliovalent dopants will affect both the nanosheet conductivity and may introduce (redox-active) surface sites for molecular or ionic recognition.

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