Postprint of: Xing X., Zhao X., Li Z., Du L., Wang C., Feng D., Geng D., Bogdanowicz R., Yang D., Oxygen vacancy-enriched V₂O₅·nH₂O nanofibers ink for universal substrates-tolerant and multi means-integratable NH₃ sensing, CHEMICAL ENGINEERING JOURNAL, Vol. 478 (2023), 147233, DOI: 10.1016/j.cej.2023.147233

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Journal Pre-proofs

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Xiaxia Xing, Xinhua Zhao, Zhenxu Li, Lingling Du, Chen Wang, Dongliang Feng, Dongsheng Geng, Robert Bogdanowicz, Dachi Yang

PII:	\$1385-8947(23)05964-8
DOI:	https://doi.org/10.1016/j.cej.2023.147233
Reference:	CEJ 147233
To appear in:	Chemical Engineering Journal
Received Date:	16 July 2023
Revised Date:	22 September 2023
Accepted Date:	7 November 2023



Please cite this article as: X. Xing, X. Zhao, Z. Li, L. Du, C. Wang, D. Feng, D. Geng, R. Bogdanowicz, D. Yang, Oxygen vacancy-enriched V_2O_5 ·nH₂O nanofibers ink for universal substrates-tolerant and multi means-integratable NH₃ sensing, *Chemical Engineering Journal* (2023), doi: https://doi.org/10.1016/j.cej.2023.147233

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Oxygen Vacancy-enriched V₂O₅•nH₂O Nanofibers Ink for Universal Substrates-tolerant and Multi Means-integratable NH₃ Sensing

3 Xiaxia Xing^a, Xinhua Zhao^a, Zhenxu Li^a, Lingling Du^a, Chen Wang^a, Dongliang Feng^a,

4 Dongsheng Geng^b, Robert Bogdanowicz^c, Dachi Yang^{a,*}

5

^a Tianjin Key Laboratory of Optoelectronic Sensor and Sensing Network Technology,
 Engineering Research Center of Thin Film Optoelectronics Technology, Ministry of
 Education and Department of Electronics, College of Electronic Information and Optical
 Engineering, Nankai University, Tianjin 300350, P. R. China

- 10 E-mail: yangdachi@nankai.edu.cn
- ^b School of Materials Science and Engineering, University of Science and Technology Beijing, Beijing, 100083, P. R. China
- 13 ^c Department of Metrology and Optoelectronics, Faculty of Electronics,
- 14 Telecommunications and Informatics, Gdansk University of Technology, 11/12G.
- 15 Narutowicza St., 80-233 Gdansk, Poland

16

17 Abstract : Universal substrates-tolerant and multi means-integratable ammonia (NH₃) 18 sensing is highly desired in future Internet of Things in environmental monitoring, food security 19 and early diagnosis of human diseases, however, is still less than satisfactory. Here, an oxygen 20 vacancy-governed NH₃ sensing has been developed with V₂O₅·nH₂O nanofibers (NFs) ink, via combined thermal decomposition of ammonium metavanadate and dilution. As-obtained NH₃ 21 sensing ink takes on red colloids, in which the V2O5 nH2O NFs around 14 nm in diameter are 22 23 interconnected. Beneficially, the fabric fiber decorated with V2O5 nH2O NFs ink displays 24 excellent selectivity and ppb-concentration detection limit. Remarkably, V₂O₅ nH₂O NFs ink 25 is integrated over "hard" and "flexible" substrates such as glass, wood, paper, leaf and fabric with excellent tolerance by multi-integratable means such as writing, dipping and sewing. 26 Theoretically, such NH₃ sensing is interpreted that the bonding between V₂O₅ NFs and H₂O 27 28 modulates oxygen vacancy and thus adsorption sites, and the incorporation between crystal water and free one contributes to stable ink. Practically, A sensing device built with 29 V₂O₅·3.1H₂O NFs ink has been simulated to communicate with a smartphone with reliable NH₃ 30 31 sensing.

Keywords: Oxygen vacancy; V₂O₅·nH₂O nanofibers sensing ink; Universal substrates tolerant; Multi means-integratable; Ammonia sensing

35 **1. Introduction**

36 Ammonia (NH₃), as a promising energy carrier [1, 2], may damage human organs if the longterm exposure to NH₃ is larger than 25 ppm due to its corrosive and toxic nature [3, 4]. Instead, 37 38 NH₃ may serve as a tracer of food spoilage [5] and an exhaled biomarker of impaired kidney 39 [6] and liver function. As such, NH₃ sensing is potentially utilized in intelligent environmental 40 monitoring, food security and early diagnosis of human diseases, which is simultaneously 41 required with excellent selectivity and stability, and ppb-level detection limit. Generally, a 42 universal substrates-tolerant and multi means-integratable NH₃ sensing may contribute to 43 intelligent monitoring in the upcoming Internet of Things, although great progress has been 44 made, it needs further exploring.

45 Actually, an NH₃ sensing material with modulated sensing performance plays a crucial 46 role in the compatible integration over universal substrates by available means. As the NH_3 sensing materials, semiconducting metal oxides (SMOs) have been widely investigated [7-9], 47 48 however, their challenging issues may limit their future applications. Firstly, oxygen vacancy 49 may contribute to gas sensing of SMOs materials. Theoretically, the reaction between reducing 50 gas such as NH₃ and ionized oxygen species would be boosted due to the enhanced adsorption of O₂ on oxygen vacancy [10, 11]. Accordingly, the means that can generate more oxygen 51 vacancies such as H₂ plasma treatment [12], doping [13] and annealing [10, 11] have been 52 53 utilized to improve the sensing performance, however, the strategies needs further developing. 54 Secondly, the nano/micro-structured NH₃ sensing SMOs are usually endowed with powder 55 form, and their suspension in an aqueous solution may agglomerate and peel off the utilized 56 substrate [14, 15]. Even being temporarily integrated, further mechanical manipulation may 57 also cause similar peeling off [16]. Thirdly, the tolerance of the sensing materials to universal 58 substrates by facially integrating means is still less than satisfactory. Conductive polymers (CPs) as NH₃ sensing materials have been integrated over "hard" substrates such as glass [17] 59 and ceramic [18] and "flexible" substrates such as polyethylene terephthalate (PET) [19] and 60 61 paper [20]. Nevertheless, the substrates are still limited and their available integratable means require either complicated procedures or proficient technicians [6, 19]. Ideally, a NH₃ sensing 62 63 material is tolerant to various substrates by multi-integratable means and its sensing 64 performance can be improved by an ingenious strategy, however, little has been reported so far.

65 V_2O_5 as a transition metal oxide presents unique electrical and sensing performance [21], in which vanadium ions (V^{5+}) with an oxidation state generate the active sites for adsorbing 66 gaseous molecules and catalyze reactions [22]. Compared with crystalline V_2O_5 , V_2O_5 $\cdot nH_2O_5$ 67 has been investigated with a low crystallization, which is subjected to less mechanical stress 68 69 and thus offers more active sites than their crystalline counterparts during reaction [23]. Notably, 70 the presence of crystal water has been reported to boost the electrochemical reaction kinetics 71 [24]. Being inspired, an oxygen vacancy-enriched V₂O₅ nH₂O nanofibers (NFs) ink with a sol 72 form in this study has been developed for universal substrates-tolerant and multi meansintegratable NH₃ sensing at room temperature. As characterized, the V₂O₅·nH₂O NFs of ~14 73 74 nm in diameter are interconnected to form red and highly dispersed ink with a zeta potential of 75 ~ 38.8 mV. Beneficially, the response of diluted V₂O₅·3.1H₂O NFs fabric to 10 ppm NH₃ have been improved (S = 17.8%) compared with that of pristine one (S = 8.6%). Furthermore, the 76 77 diluted V₂O₅·3.1H₂O NFs fabric fiber shows 100 ppb detection limit of NH₃ and excellent 78 selectivity. Remarkably, the V_2O_5 nH₂O NFs ink has been integrated on various substrates such 79 as ceramics, glass, wood, paper, fabric and leaf, by which multi-integratable means of writing, 80 dipping and sewing have been applied. Such sensing ink would contribute to the diversification 81 of NH₃ sensors in future intelligent sensing.

83 2. Experimental section

84 **2.1 Synthesis of V₂O₅·2.3H₂O NFs ink** [25]

Firstly, 1 g ammonium metavanadate (NH₄VO₃) was ground with deionized (D.I.) water, and then the fluid was mixed with 10 mL of 1M HCl under continuous stirring. Secondly, when the suspension turns red, D.I. water was added to make the total volume of 20 mL, the supernatant

88 was removed after precipitation. Thirdly, the red precipitate was dispersed into 80-90 °C hot

- 89 water to a total volume of 20 mL, the supernatant was removed after stirring and precipitating.
- 90 Finally, the dark red $V_2O_5 \cdot 2.3H_2O$ NFs dispersions were filled with 80-90 °C hot water to a
- 91 total volume of 40 mL for the subsequent utilization.

92 2.2 Synthesis of sensing fabrics and fabrics fiber integrating V₂O₅·nH₂O NFs ink

- 93 Synthesis of V₂O₅·nH₂O NFs fabrics is briefly described as follows. Initially, 0.5 mL, 2 mL
- and 5 mL of the above synthesized $V_2O_5 \cdot 2.3H_2O$ NFs ink were ultrasonically dispersed in 10
- 95 mL D.I. water, respectively. Correspondingly, they are denoted as ink-0.5, ink-2 and ink-5 in
- 96 Fig. 3a, respectively. Secondly, the rectangular polyester fabric $(2 \text{ cm} \times 0.5 \text{ cm})$ and fabric fiber
- 97 (Diameter: ~ 207 μ m, Length: ~ 2 cm) was immersed in the above synthesized V₂O₅·nH₂O NFs 98 dispersion for 1 min. Finally, the V₂O₅·nH₂O NFs fabrics and fabric fiber were dried at room
- temperature. In the same way, the sensing ink was integrated over the PET and paper in Fig. 5e.
- 100 It should be noted that 2 mL of the pristine $V_2O_5 \cdot 2.3H_2O$ NFs ultrasonically dispersed in 10
- mL D.I. water was defined as diluted $V_2O_5 \cdot 3.1H_2O$ NFs ink, which was taken as an example
- 102 for deep investigation.

103 2.3 Synthesis of V₂O₅·2.3H₂O NFs aerogel and powder, and V₂O₅ NFs powder

104 The V₂O₅·2.3H₂O NFs ink was firstly frozen at -18 °C and then lyophilized at -51 °C in a 105 freeze-drier (FD-1A-50, Henan Brothers Instrument and Equipment Co., Ltd., China) to obtain 106 V₂O₅·2.3H₂O NFs aerogel. The V₂O₅·2.3H₂O NFs aerogel was grounded using an agate mortar 107 to obtain V₂O₅·2.3H₂O NFs powder, its resistivity was tested under various pressures (2-30 108 MPa) in Fig. S1, in which the resistivity mean is ~ 7823 Ω cm. The V₂O₅·2.3H₂O NFs powder 109 was annealed in air at 600 °C for 2 h to remove the crystal water, then the V₂O₅ NFs powder 100 was collected.

111 2.4 Characterization

The samples were characterized by field emission scanning electron microscopy (FE-SEM, 112 JSM-7800) with energy dispersive X-ray spectroscopy (EDS, Oxford), transmission electron 113 114 microscopy (TEM, JEM-2200FS), X-ray diffraction (XRD, Rigaku Smart Lab 3kW) using Cu 115 Ka radiation, Raman spectra (SR-500I-A, a wavelength of 532 nm as the excitation), 116 Ultraviolet-visible diffuse reflectance spectra (Shimadzu UV-3600), Mott-Schottky test (electrochemistry workstation VersaSTAT 4, AMETEK Princeton), thermogravimetric 117 118 analysis (TGA) (Netzsch STA449F5 instrument, temperature range 30-600 °C, heating rate 119 10 °C/min, in nitrogen atmosphere), Automatic powder resistivity tester (ST2742B), Zeta 120 potential analyzer (Malvern Zetasizer Nano ZS ZEN3600, UK), electron paramagnetic resonance (EPR) spectroscopy (Bruker EMXPLUS) and X-ray photoelectron spectroscopy 121 (XPS, Thermo Scientific ESCALAB 250Xi). The XPS spectra on binding energies of various 122 123 elements have been calibrated with C 1s at 284.8 eV.

124 **2.5 Gas sensing measurement**

The gas sensing was tested at room temperature (RT, ~ 25 °C) in air atmosphere. In detail, the 125 two ends of fabric were connected to the Data Acquisition System (KEITHLEY 2701) by two 126 127 gold clamps, which were placed in a homemade test chamber of 18 L with two air fans and a 128 vaporizer. Notably, the NH₃ sensing is *in-situ* detection directly without other electrodes. 129 Additionally, the gaseous and dry NH₃ with high-purity was adopted. The calculation of NH₃ 130 concentration is conducted by the gas distribution formula (equation 1), in which C (ppm) and 131 φ represent the target gas concentration and volume fraction, respectively, and V₁ (mL) and V₂ (mL) are denoted as the volume of target gas and testing chamber ($V_2 = 18$ L). The sensing 132 response is expressed by $S = (R_g/R_a-1) * 100\%$, of which R_a and R_g are the resistances in the air 133 134 and target gas, respectively. The response/recovery time is defined as the time taken by the 135 sensor to reach 90% of the final steady-state resistance after injecting or switching off the target 136 gas.

(1)

$$137 \qquad V_1 = \frac{V_2 \times C}{\varphi} \times 10^{-6}$$

138

139 **3. Results and Discussion**

140 **3.1. Synthesis and characterization**

141 In Fig. 1a, the pristine V_2O_5 nH₂O NFs ink was diluted and integrated over fabrics, and the synthetic details of pristine V₂O₅·nH₂O NFs ink were provided in above experimental section. 142 143 Meanwhile, the three-dimensional (3D) crystal structure of V₂O₅ nH₂O NFs was simulated by Visualization for Electronic and STructural Analysis (VESTA) [26]. Also, the X-ray 144 diffractions (XRD) of pristine and diluted V₂O₅·nH₂O NFs were conducted (Fig. S2a) with the 145 146 diffractive peak of $V_2O_5 \cdot nH_2O$ at ~ 10° [27]. Meanwhile, the crystal water was removed by 147 annealing the pristine V₂O₅·nH₂O NFs, and was then confirmed as the V₂O₅ (PDF#89-0612) in 148 Fig. S2b. Further, Raman spectra of pristine V₂O₅·nH₂O NFs (Fig. S2c) show the V-O Raman 149 scattering peaks with the orthorhombic crystalline [28]. Remarkably, the thermogravimetric analysis (TGA) was carried out to determine the "n" value in V2O5 nH2O NFs. In Fig. 1b, 150 151 weight loss of 23.4% and 18.6% occur at 100 - 600 °C, which is attributed to the loss of crystal 152 water, and the "n" values corresponding to diluted and pristine $V_2O_5 \cdot nH_2O$ NFs are 3.1 and 2.3, respectively. Specifically, the detailed calculation of "n" value in V₂O₅·nH₂O is described as 153 154 follows. M, m_p and m_d represent relative molecular mass, mass of pristine V₂O₅ \cdot nH₂O NFs and 155 mass of diluted one, respectively. Therefore, the "n" values corresponding to the diluted and 156 pristine ones are calculated by the bellow proportional formula of the chemical equation.

 $V_2O_5 \cdot nH_2O \rightarrow V_2O_5$ M: 181.88 + n*18 181.88
mp: 4.198 g 4.198 g*(1-18.6%)

3.933 g

160

m_d:

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163

164 165 In Fig. 1c, $V_2O_5 \cdot 3.1H_2O$ NFs ink is observed dense and overlapped in a lower magnification with scanning electron microscopy (SEM). While in a closer observation under transmission electron microscopy (TEM), the diameter of $V_2O_5 \cdot nH_2O$ NFs in Fig. 1d is measured ~ 14 nm (Fig. S3). Moreover, the high-resolution TEM (HR-TEM) image and selected area electron diffraction (SAED) pattern in Fig. 1f and Fig. S4 show the (102) plane of

3.933 g*(1-23.4%)

166 V_2O_5 (PDF#89-0612). Further, the elemental mappings under TEM (Fig. S5) verified the 167 existence of V and O elements, and the diameter of the fabric fiber integrated with diluted 168 $V_2O_5 \cdot 3.1H_2O$ NFs ink was measured ~ 207 µm in Fig. 1g₁. By comparing with the shape of 169 pristine fabric fiber in Fig. 1h₁, the flake shape in Fig. 1g₂ reveals that the V₂O₅ · 3.1H₂O NFs 170 ink has been integrated over fabrics. Interestingly, the Tyndall effects of pristine and various 171 diluted ink were compared in Fig. S6, the dispersibility of pristine V₂O₅ · 2.3H₂O NFs ink can 172 be improved via dilution.

173 **3.2.** The stable V₂O₅·nH₂O NFs ink for dilution-modulated NH₃ sensing

174 To get insight into the role of water in $V_2O_5 \cdot nH_2O$ NFs ink, the free water was initially removed 175 by freezing and drying $V_2O_5 \cdot 2.3H_2O$ NFs ink, and then the lyophilized ones were annealed to remove crystal water and obtain V₂O₅ for subsequent comparison. In Fig. S7, the color of 176 $V_2O_5 \cdot 2.3H_2O$ NFs powder changed from its pristine dark red to orange after annealing. The 177 178 V₂O₅ and V₂O₅·2.3H₂O NFs powder were ultrasonically dispersed into D.I. water and pure 179 ethanol, respectively. Correspondingly, various dispersions were dripped over interdigital electrodes in Fig. 2d for comparing their NH₃ sensing performance, and the real-time resistance 180 181 curves were shown in Fig. 2a.

In Fig. 2b-c, the V₂O₅·2.3H₂O NFs exhibit a higher response to 10 ppm NH₃ than that of 182 V₂O₅ NFs in both water and pure ethanol solvent, revealing the crystal water-boosted NH₃ 183 184 sensing. Actually, water solvent may contribute to lower baseline resistance in both V₂O₅ and 185 V₂O₅·nH₂O NFs (Fig. 2c). Further, free water is required in preparing V₂O₅·nH₂O NFs ink in 186 Fig. 2e. Otherwise, uneven and unstable dispersion can be obtained. Meanwhile, the 187 $V_2O_5 \cdot 2.3H_2O$ NFs ink and ethanol dispersion were dipped over fabric in Fig. 2d₁. In Fig. S8, the resistance value of V₂O₅·2.3H₂O NFs ink fabric is ~ 0.49 MΩ, however, the one with 188 189 ethanol dispersion is larger than 20 M Ω , which reveals that the uniform and stable V₂O₅·2.3H₂O 190 NFs ink contribute to integrating conductive fabric. Notably, if one deliberately removed the 191 crystal water in V₂O₅·2.3H₂O NFs or replaced the dispersion medium from water to pure 192 ethanol, the dispersed phase is obviously separated from dispersion medium (Fig. 2e), rather 193 than obtaining stable ink. As such, the incorporation of crystal water bonded by V₂O₅ with free 194 water in the dispersion medium plays a pivotal role in the formation of sensing ink.

195 The diluted $V_2O_5 \cdot 3.1H_2O$ and pristine $V_2O_5 \cdot 2.3H_2O$ NFs inks were observed with the 196 variation of Tyndall effect in Fig. 2f and g, in which the light path penetrates after diluting with 197 high dispersibility of colloid [29]. Meanwhile, the simulated 3D crystalline structures with 198 various oxygen vacancies of diluted and pristine V_2O_5 nH₂O NFs are shown in Fig. 2f₁ and g₁, 199 respectively. With the pristine $V_2O_5 \cdot 2.3H_2O$ NFs ink for comparisons, the diluted $V_2O_5 \cdot 3.1H_2O$ NFs ink was integrated over the fabric (2 cm×0.5 cm) and the fabric fiber (Diameter: ~ 207 μ m, 200 201 Length: ~ 2 cm), respectively. In the photographs of Fig. 2f and g, the color of diluted 202 $V_2O_5 \cdot 3.1H_2O$ NFs fabrics was seen lighter than that of pristine $V_2O_5 \cdot 2.3H_2O$ ones. To further 203 gain insight into the role of dilution, the NH₃ sensing performance of the above integrated 204 various pristine V₂O₅·2.3H₂O and diluted V₂O₅·3.1H₂O NFs fabric were investigated with 205 comparison. In Fig. 2h, the recovery speed of pristine $V_2O_5 \cdot 2.3H_2O$ NFs fabric is improved by 206 both diluting and adopting the fabric fiber. Meanwhile, the responses toward 5 ppm and 25 ppm 207 NH₃ were summarized in Fig. 2i, and show that the sensing responses of diluted $V_2O_5 \cdot 3.1H_2O$ 208 NFs ink onto both fabric and fabric fiber are higher than those of pristine ones. Moreover, the 209 real-time responses to 1-50 ppm NH₃ were evaluated in Fig. 2j, which further reveals the 210 dilution improved NH₃ sensing performance.

211 **3.3. Evaluation of the NH₃ sensing performance**

212 The content of $V_2O_5 \cdot nH_2O$ NFs in the sensing ink governs the NH₃ sensing. In Fig. 3a, the responses of pristine and various diluted V₂O₅ · nH₂O NFs fabric to 10 ppm NH₃ were evaluated, 213 214 and the $V_2O_5 \cdot nH_2O-2$ mL NFs fabric manifested the highest response (S = 17.8%) compared 215 with pristine ones (S = 8.6%), and was thus chosen for subsequent evaluation and renamed by 216 $V_2O_5 \cdot 3.1H_2O$ NFs fabric. The response and recovery time were evaluated to ~ 75 s and 36 s 217 toward 1 ppm NH₃ in Fig. S9, respectively. Remarkably, the flexibility of V₂O₅·nH₂O NFs 218 fabric was investigated by testing their responses to 1 ppm and 10 ppm NH₃ upon the bending 219 angle at 0° , 45° , 90° and 360° , respectively. Excitedly, little difference was observed in Fig. 3b, 220 indicating excellent flexibility.

221 The stability and selectivity are crucial parameters for NH₃ sensing. Remarkably, the 222 sensing evaluation to 5 ppm NH₃ is repeated for 126 days' durations in Fig. 3c with good 223 stability. Furthermore, the responses of various interfering gases and 10 ppm target NH₃ were 224 compared in Fig. 3d, revealing excellent selectivity. Meanwhile, the NH₃ sensing of three 225 $V_2O_5 \cdot 2.3H_2O$ NFs fabrics in Fig. S10 is compared, which shows a slight variation in response 226 to the same concentration NH₃ and takes a good consistency. Additionally, the diluted 227 V₂O₅·3.1H₂O NFs ink was integrated over the fabric fiber in Fig. 3e, and its low detection limit 228 is around 100 ppb NH₃. Towards a low NH₃ concentration (e.g., 100 ppb-1 ppm), the responses 229 show an excellent linear relationship in Fig. 3f. While towards a high one (e.g., 1-50 ppm), 230 excellent repeatability is observed in Fig. 3g. As a result, the V2O5 nH2O NFs fabric simultaneously present ppb-level detection, high selectivity and stability, excellent flexibility 231 232 and low working temperature. Compared with other SMOs NH₃ sensing materials in Table 1, 233 a gel-stated and stable ink of $V_2O_5 \cdot nH_2O$ NFs is prepared, which can be integrated over various "hard" and "flexible" substrates by multi-integratable means. 234

235 In our experiments, both temperature and humidity can influence the NH₃ sensing. In Fig. 236 S11, the temperature-dependent sensing responses were observed to 20 ppm NH₃ at ~ 26-140 237 °C, and the highest one takes place at room temperature (~ 26 °C). Meanwhile, the baseline resistance and the sensing response of V2O5.3.1H2O NFs fabric toward 5 ppm NH3 decrease 238 239 with humidity (Fig. S12), similar to previous SMOs [30] and to other humidity sensors [31]. 240 Such a decrease in the sensing response might be interpreted that H_2O molecules occupy 241 adsorption sites, which weakens the reaction between NH_3 and adsorbed oxygen onto the surface of V₂O₅·3.1H₂O NFs [32], as may be addressed by covering filter membrane [33]. 242

243 **3.4.** The oxygen vacancy governed NH₃ sensing mechanism

244 We experimentally investigated the chemisorbed oxygen to understand the dilution-boosted 245 NH_3 sensing mechanism, and three characterizations on oxygen vacancy (V_0) of diluted 246 V₂O₅·3.1H₂O NFs were performed with pristine V₂O₅·2.3H₂O ones as comparison. Firstly, the 247 O 1s X-ray photoelectron spectroscopy (XPS) in Fig. 4a spectra were deconvoluted into three 248 oxygen species of O_{I} , O_{II} and O_{III} , which are associated with oxygen atoms bound to metals, 249 defect sites with low oxygen coordination and hydroxy species, respectively. Remarkably, the 250 integral-area ratios of OII increase from 20% of pristine V2O5 · 2.3H2O NFs to 52% of diluted 251 $V_2O_5 \cdot 3.1H_2O$ ones, indicating that the diluted $V_2O_5 \cdot 3.1H_2O$ one possesses more oxygen 252 vacancies [34]. Meanwhile, the V 2p spectra in Fig. 4b correspond to the characteristics of V^{5+} , 253 the discrepancy in binding energy (0.3 eV) indicates distinct electronic environments of V ions in the pristine and diluted V2O5 nH2O NFs, which might be interpreted as increased oxygen 254 vacancy in the diluted $V_2O_5 \cdot 3.1H_2O$ ones [10]. Secondly, the presence of oxygen vacancy was 255 256 further studied by electron paramagnetic resonance (EPR) spectroscopic measurements in Fig. 257 4c and symmetrical EPR signals (g = 1.9612) are assigned to the unpaired electrons in the 258 oxygen vacancy sites [10, 35]. The ESR intensity of diluted $V_2O_5 \cdot 3.1H_2O$ NFs is higher than

that of pristine $V_2O_5 \cdot 2.3H_2O$ ones, indicating dilution governed the oxygen vacancy, which result in more chemisorbed oxygen for gas sensing. Thirdly, such result is also evidenced by the narrower optical bandgaps (Eg) of diluted $V_2O_5 \cdot 3.1H_2O$ NFs (1.87 eV) than that of pristine $V_2O_5 \cdot 2.3H_2O$ (2.19 eV) in Fig. 4d.

263 The energy-band variation of V₂O₅ nH₂O NFs was investigated for understanding the NH₃ sensing mechanism. Specifically, the valence band maximum (E_v) of the V₂O₅·3.1H₂O NFs is 264 265 determined to ~ 2.4 eV (Fig. S14). Accordingly, the conduction band minimum (E_c) of $V_2O_5 \cdot 3.1H_2O$ is calculated to ~ 0.53 eV according to Equation (2). Usually, V_2O_5 is reported 266 267 as a n-type semiconductor [36]. However, p-type sensing characteristic with increased 268 resistance was observed in this study (Fig. 2a and h), which is explained as follows. The V2O5·3.1H2O NFs contain abundant oxygen vacancy, which will improve chemisorption of O2 269 270 and H₂O molecule [11, 37], capture more electrons from the conduction band of $V_2O_5 \cdot nH_2O$ 271 NFs and thus bend upward band causing an inversion layer, therefore, the Fermi level (E_F) 272 located below the intrinsic level (E_i) in Fig. 4e [38]. In the surface inversion layer, holes usually 273 serve as the major carriers with p-type feature, which was confirmed by Mott-Schottky with a 274 negative slope in Fig. 4e₁.

275
$$E_c = E_v - E_g$$

276 To understand the p-type sensing mechanism, the NH₃ sensing evaluations under various working temperatures were investigated in Fig. S11, the V₂O₅·3.1H₂O NFs show increased 277 sensing resistance to 20 ppm NH₃ at ~ 26-80 °C and decreased ones at ~ 100-140 °C. Such 278 279 phenomenon is explained as follows. At lower temperatures, the strong adsorption of O₂ and 280 H₂O molecules contribute to the formation of an inversion layer on the surface of $V_2O_5 \cdot 3.1H_2O$ 281 NFs, exhibiting p-type semiconductor properties [39]. With the elevating of temperature, an 282 inversion layer would be destroyed without sufficient O₂ and H₂O molecules, n-type sensing 283 behavior would be seen. Further, we conducted additional comparative experiments on NH₃ 284 sensing under insufficient oxygen conditions and air atmosphere in Fig. S15, the significantly 285 decreased response in Fig. S15a indicates that the sufficient surface adsorption of oxygen 286 contributes to NH₃ sensing of V₂O₅·3.1H₂O NFs.

(2)

Accordingly, the NH₃ sensing mechanism of $V_2O_5 \cdot nH_2O$ NFs fabrics is interpreted as follows. In Fig. 4f, when the pristine p-type $V_2O_5 \cdot 2.3H_2O$ ones are exposed to NH₃, the preadsorbed oxygen species (O_2^-) and hydroxy species (-OH) react with NH₃ and release electrons [40, 41], reducing the hole concentration and thus elevating the resistance. Similarly, the diluted $V_2O_5 \cdot 3.1H_2O$ NFs show NH₃ sensing mechanism in Fig. 4f₁. However, the content of their oxygen vacancy is significantly increased thus improved chemisorbed oxygen, and finally present boosted NH₃ sensing.

3.5. V₂O₅·nH₂O NFs ink for universal substrates-tolerant and multi means-integratable NH₃ sensing and the simulation detection of NH₃

The universal-substrates tolerance and multi-means integration of $V_2O_5 \cdot 2.3H_2O$ NFs ink were investigated. The tolerance has been widely examined on hard substrates such as ceramics, stainless steel, glass and wood, and flexible ones such as Chinese "Xuan" paper, leaf, Al foil, plastic wrap and A4 size paper in Fig. 5a. Meanwhile, the adhesive performance of the $V_2O_5 \cdot 2.3H_2O$ NFs ink over the above substrate has been investigated in Fig. S16, one can see that the adhesive properties depend on the substrates and the sensing ink shows a weaker adhesion than that of commercial one on A4 paper (Fig. S17). As for the integratable means, our $V_2O_5 \cdot 2.3H_2O$ NFs ink can be dipped with a paintbrush to draw the school badge and the motto of Nankai University in Fig. 5a and other "dipping-drying" approach in Fig. 5b.

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305 Impressively, the $V_2O_5 \cdot 3.1H_2O$ NFs ink can also serve as a colouring agent with color variation from white of pristine fabric fiber to orange, which can be integrated over the fabric fiber (Fig. 306 5c), and can even be sewed on the clothes with the "NKU" pattern. Particularly, by freezing 307 308 and drying, the V₂O₅·3.1H₂O NFs ink can be transformed into lightweight aerogel, and can 309 even stand on the tip of the reed (Fig. S18). In this case, even being storing 365 days (Fig. 5d) 310 and 608 days (Fig. S19a), the V₂O₅·3.1H₂O NFs ink remains excellent dispersibility and 311 stability, which is verified by characterizing the zeta potential of V_2O_5 ·3.1H₂O NFs ink to ~ 312 38.8 mV after storing 608 days (Fig. S19b).

313 As examples, the PET, Chinese "Xuan" paper and fabric integrated with $V_2O_5 \cdot 3.1H_2O$ 314 NFs ink were examined for their NH₃ sensing performance in Fig. 5e, showing substrates-315 dependent NH₃ sensing, which may be explained that these bare and insulated substrates serve as support and don't participate electron transport. Although previous investigations (Table 1) 316 317 have made great progress, our V₂O₅·nH₂O NFs ink is the one that can be simultaneously utilized for universal substrates-tolerant and multi means-integratable NH₃ sensing. Practically, such 318 319 NH₃ sensing ink enable to be integrated into the feasible substrates such as smocks, mask and 320 food packaging bag for environmental monitoring, exhaled diagnosis of human diseases and inspection of food safety. Herein, we elaborately integrated the V₂O₅·3.1H₂O NFs ink onto the 321 322 polyethylene sample bag (4 cm×6 cm) as an example, to simulate detection of NH₃, which was 323 read by a smartphone (Fig. 5f). Specifically, the microcontroller NodeMCU (ESP8266, 5.8 cm×3.1 cm) with Wireless Fidelity (Wi-Fi) module communicate with the smartphone and 324 325 perform the NH₃ sensing and alarming of the device. In the supplemental video, when 10 ppm 326 NH₃ was injected and the sensing voltage is lower than the alarm threshold (0.5 V), the smartphone read "AlARMING!" (Fig. 5g). Conversely, the NH₃ being released with the one 327 higher than 0.5 V, and "Monitoring" in smartphone is seen. Also, the detailed historical 328 329 information can be read and recorded in Fig. 5h, which is great potential for inspection of food 330 safety.

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332 **4. Conclusion**

To summarize, an oxygen vacancy-enriched V₂O₅·nH₂O NFs ink has been developed by 333 334 combining the thermal decomposition of ammonium metavanadate with subsequent dilution, 335 for universal substrates-tolerant and multi means-integratable NH₃ sensing at room temperature. Experimentally, the $V_2O_5 \cdot nH_2O$ NFs of ~ 14 nm in diameter were observed to be interconnected, 336 337 forming red colloids in an aqueous solution with high dispersibility. Theoretically, the bonding 338 between V_2O_5 NFs and H_2O governs the oxygen vacancy with improved the adsorption sites of 339 NH₃, and the incorporation between crystal water and free water contributes to stable ink. Beneficially, the diluted V₂O₅· 3.1H₂O NFs fabrics show an increased response to 10 ppm NH₃ 340 (S = 17.8%) compared with the pristine ones (S = 8.6%). Also, the V₂O₅·nH₂O NFs ink fabric 341 342 fiber displays excellent selectivity and ppb-level detection limit to NH₃. Remarkably, 343 V₂O₅·nH₂O NFs ink has been integrated over various substrates such as ceramics, glass, wood, paper, fabric and leaf with universal substrates-tolerance. Meanwhile, multiple strategies of 344 345 writing, dipping and sewing have been adopted for integration. As an example of application, 346 the developed oxygen vacancy-enriched V2O5.3.1H2O NFs ink has been integrated into a 347 sensing device and communicates with a smartphone with reliable monitoring and alarming, which is potential in future intelligent sensing of Internet of Things. Future investigations are 348 349 expected to be conducted on theoretical calculations and humidity-dependent NH₃ sensing.

351 Declaration of Competing 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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355 **Data availability**

- 356 Data will be made available on request.
- 357

358 Acknowledgements

359 This work was financially supported by the National Natural Science Foundation of China

- (Grant No. 52072184) and Tianjin Research Innovation Project for Postgraduate Students
 (General Project, Grant No. 2022BKY035).
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Fig. 1. The synthesis and characterization of $V_2O_5 \cdot nH_2O$ NFs. (a) The schematic diagram of diluting and integrating $V_2O_5 \cdot nH_2O$ NFs ink, and the simulated crystal structure of $V_2O_5 \cdot nH_2O$ NFs. (b) The TGA curves of pristine and diluted $V_2O_5 \cdot nH_2O$ NFs. (c) The SEM, (d-e) TEM and (f) HRTEM images of pristine $V_2O_5 \cdot 2.3H_2O$ NFs. (g) The schematic diagram and (g₁-g₂) SEM images of fabric fiber integrated with $V_2O_5 \cdot nH_2O$ NFs ink. (h) The schematic diagram and (h₁) the SEM image of bare fabric fiber.



Fig. 2. The NH₃ sensing comparison between freeze-dried and annealed V₂O₅·2.3H₂O NFs powder being dispersed in D.I. water and pure ethanol, respectively. (a) The real-time resistance and (b) sensing curves to 10 ppm NH₃, and corresponding (c) histogram of baseline resistance and response value. (e) The photograph of above four dispersions and (d) integrated interdigital electrode. (d₁) The freeze-dried V₂O₅·2.3H₂O NFs dispersed in D.I. water and pure ethanol were integrated onto the fabric, respectively. The scale bars in (d₁) are 0.5 cm. The photographs and structures of (f-f₁) diluted and (g-g₁) pristine V₂O₅·nH₂O NFs inks. The "Tyndall effect" of diluted V₂O₅·3.1H₂O NFs ink irradiated by red light ($\lambda = 638$ nm). The comparison on NH₃ sensing performance between diluted and pristine V₂O₅·nH₂O NFs inks integrated on fabric and the fabric fiber, respectively. (h) The real-time resistance curves and (i) the summarized responses. (j) The response curves to various NH₃ concentration. The RH of (a-b, h-j) is at ~ 22%.



529 **Fig. 3.** The systematic NH₃ sensing evaluation of $V_2O_5 \cdot nH_2O$ NFs ink integrated over fabrics: 530 (a) The dilution-dependent sensing response of $V_2O_5 \cdot nH_2O$ NFs ink, (b) the $V_2O_5 \cdot 3.1H_2O$ NFs 531 fabric under various bending angle and their corresponding sensing responses, (c) the long-term stability and (d) selectivity of diluted V₂O₅·3.1H₂O NFs fabric. The NH₃ sensing evaluation of 532 533 V₂O₅·3.1H₂O NFs fabric fiber: (e) The response curve to 0.1-1 ppm NH₃, (f) the relationship 534 between sensing responses and NH₃ concentrations, (g) the response curve to various NH₃ concentrations. The scale bars in (c) and (e) are 0.5 cm. The RH of (a-b, d, e-g) is at ~ 19% and 535 536 RH of (c) is at ~ 19%-57%.



Fig. 4. The various characterizations on oxygen vacancy of pristine $V_2O_5 \cdot 2.3H_2O$ NFs and diluted $V_2O_5 \cdot 3.1H_2O$ NFs. High-resolution XPS spectra are related to (a) O 1s and (b) V 2p, (c) EPR spectra and (d) plots of $(ahv)^2$ vs photon energy (hv). The schematic energy-band variation of $V_2O_5 \cdot nH_2O$ NFs. (e) An inversion layer marked with red rectangle and p-type surface conductivity. (e₁) The Mott-Schottky plot of $V_2O_5 \cdot 3.1H_2O$ NFs. The NH₃ sensing mechanism diagrams of (f) pristine $V_2O_5 \cdot 2.3H_2O$ NFs and (f₁) diluted $V_2O_5 \cdot 3.1H_2O$ NFs.

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546 Fig. 5. The universal substrates-tolerant and multi means-integratable NH₃ sensing. (a) The 547 V₂O₅·2.3H₂O NFs ink for drawing the school badge and the motto of Nankai University onto 548 hard and flexible substrates. (b) The pristine $V_2O_5 \cdot 2.3H_2O$ ink was evenly integrated onto 549 various flexible substrates as examples. (c) The diluted V₂O₅·3.1H₂O NFs ink integrated fabric 550 fiber for sewing patterns into clothes. (d) The photographs of V₂O₅·3.1H₂O NFs ink after 551 storing 365 days. (e) The NH₃ sensing of diluted V₂O₅·3.1H₂O NFs ink integrated on flexible 552 substrates with PET, Chinese "Xuan" paper and fabric as examples, and "Con." in the ordinate represents concentration. The scale bars in (a, b, c) are all 1 cm. (f) Simulated detection of NH₃ 553 554 were conducted by intergrating the $V_2O_5 \cdot 3.1H_2O$ NFs ink onto sample bag and communicating 555 with a smartphone. (g) Smartphone reading the real-time sensing parameters and the records of 556 alarming to 10 ppm NH₃ and (h) the historical measurements. The RH of (e) is at $\sim 22\%$.

Table 1. Comparison of various NH₃ sensing materials in both presenting forms and NH₃ sensing.

Materia l type	Materials	Presentin g forms	Substrates	Mechanic al flexibility	Integrating means	W T ^{a)}	Response@Con . ^{c)}	Tres/Tree time (s) @Con. ^{c)}	LOD ^{h)} (ppm)	Refs. ^{k)}
SMOs based NH ₃ sensing material s	Co-Fe ₂ O ₃	powder	ceramic tube	No	spin-coating combined with calcination	27 5 °C	275% ^d)@10 ppm	7.2/5.4@1 0 ppm	0.01 ⁱ⁾	[42]
	MoO3 nanorods	powder	glass	No	spin-coating by mixing with solvent	20 0 °C	36% ^a)@5 ppm	230/267@ 5 ppm	~5 ⁱ⁾	[8]
	WO3@SnO2 Core shell nanosheet	thin film	MEMS	No	dripping- coating by mixing with solvent	20 0 °C	1.5 °)@15 ppm	62/42@15 ppm	5 ⁱ⁾	[43]
	Ni-doped In ₂ O ₃ nanostructure	powder	ceramic tube	No	coating by mixing with solvent	14 0 ℃	2732 ^{e)} @50 ppm	23/10@50 ppm	~1 ⁱ⁾	[9]
	MXene/CuO composite.	solution	ероху	Yes	spraying	RT	24.8 ^{f)} @100 ppm	43/26@10 0 ppm	~1 ⁱ⁾	[41]
	MXene/V2O5/CuW O4	precipitat e	alumina sheet with interdigitated gold electrode	No	coating	RT	53.5 ^{f)} @51 ppm	1.6/4@51 ppm	1 ⁱ⁾ 0.3 ^{j)}	[44]
	V2O5*nH2O NFS	ink	ceramics, stainless steel, glass, wood, paper, leaf, Al foil, plastic wrap, fabric and polyurethan e	Yes	dripping, writing, dipping and sewing	RT b)	~4.2% ^{g)@} 1 ppm	75/36@1 ppm	~ 0.1 ⁱ⁾	This wor k
Carbon based NH3 sensing	PANI/MWCNTs	-	polypropylen e fabric	Yes	spray-coating and chemical polymerizatio n	RT	61.54% ^{g)} @20 ppm	9/30@20 ppm	0.2)	[45]

material s	PEDOT:PSS nanowires	aqueous suspensio n	PET	Yes	spin-coating	RT	~2.2% ^{g)} @6 ppm	96/318@6 ppm	0.1 ^{j)}	[19]
	PSS-PANI/PVDF	-	PVDF membrane	Yes	in-situ polymerizatio n	RT	70% ^{g)} @1 ppm	160/400@ 1 ppm	~ 0.1 ⁱ⁾	[46]
	Pt-NDs/PPy- nanolayer@CNTs	powder	filter paper	Yes	coating by mixing with solvent	RT	~40% ^{g)} @50 ppm	2/~10@2 v/v%	~0.00 5 ⁱ⁾	[20]
	PANI/NiCo2O4	powder	gelatin film	Yes	spin-coating	RT	4.67 ^{f)} @20 ppm	22/62@20 ppm	~ 0.5 ⁱ⁾	[47]
	PANI/MXene	solution	ероху	Yes	dripping	RT	27% ^{g)} @5 ppm	27/5@5 ppm	~ 0.3 ⁱ⁾	[48]
	SnPx/rGO	powder	interdigitated electrodes	No	dripping	RT	117.5% ^{d)} to 40 ppm	126/306@ 10 ppm	0.043 6 ^{j)}	[49]

^{a)} Working temperature, ^{b)} Room temperature, ^{c)} Concentration, ^{d)} Calculated by $(R_a/R_g-1)*100\%$, ^{e)} Calculated by R_a/R_g , ^{f)} Calculated by R_g/R_a , ^{g)} Calculated by $(R_g/R_a-1)*100\%$, ^{h)} Limit of Detection, ⁱ⁾ Experimental measurements, ^{j)} Theoretical calculation, ^{k)} References. 559 560 561

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563 **Declaration of interests**

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It is authors declare that they have no known competing financial interests or personal 565 566 relationships that could have appeared to influence the work reported in this paper.

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568 The authors declare the following financial interests/personal relationships which may be 569 considered as potential competing interests:



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- 582 2. The ink toward universal substrates-tolerant and multi means-integratable NH₃ sensing.
- 583 3. The oxygen vacancy governed NH₃ sensing mechanism is rationally interpreted.
- 584 4. Simulation on detecting NH_3 is conducted with reliable sensing response.