



Article

Reversible Redox Property of Co(III) in Amorphous Co-doped SiO₂/γ-Al₂O₃ Layered Composites

Shotaro Tada ¹, Shota Saito ¹, Akito Mori ¹, Hideki Mizuno ¹, Shiori Ando ¹, Toru Asaka ¹, Yusuke Daiko ¹, Sawao Honda ¹, Samuel Bernard ² and Yuji Iwamoto ^{1,*}

- ¹ Department of Life Science and Applied Chemistry, Graduate School of Engineering, Nagoya Institute of Technology, Gokiso-cho, Showa-ku, Nagoya 466-8555, Japan; s.tada.341@stn.nitech.ac.jp (S.T.); shota.saito@cgco.co.jp (S.S.); a_mori_10029@murata.com (A.M.); m.hideki0318@gmail.com (H.M.); shiori786@outlook.jp (S.A.); asaka.toru@nitech.ac.jp (T.A.); daiko.yusuke@nitech.ac.jp (Y.D.); honda@nitech.ac.jp (S.H.)
- ² Centre Européen de la Céramique, University of Limoges, 12 Rue Atlantis, 87068 Limoges, France ; samuel.bernard@unilim.fr
- * Correspondence: iwamoto.yuji@nitech.ac.jp; Tel.: +81-52-735-5276

Received: 29 October 2020; Accepted: 23 November 2020; Published: 25 November 2020

Abstract: This paper reports on a unique reversible reducing and oxidizing (redox) property of Co(III) in Co-doped amorphous SiO₂/ γ -Al₂O₃ composites. The Fenton reaction during the H₂O₂catalyzed sol-gel synthesis utilized in this study lead to the partial formation of Co(III) in addition to Co(II) within the composites. High-resolution transmission electron microscopy (HRTEM) and high-angle annular dark-field scanning transmission electron microscopy (HAADF-STEM) analyses for the composite powder sample with a composition of Al:Si:Co = 85:10:5 showed the amorphous state of the Co-doped SiO₂ that modified γ -Al₂O₃ nanocrystalline surfaces. In situ X-ray absorption fine structure (XAFS) spectroscopic analysis suggested reversible redox reactions of Co species in the composite powder sample during heat-treatment under H₂ at 500 °C followed by subsequent cooling to RT under Ar. Further analyses by in situ IR spectroscopy combined with cyclic temperature programmed reduction/desorption (TPR/TPD) measurements and X-ray photoelectron spectroscopic (XPS) analysis revealed that the alternating Co(III)/(II) redox reactions were associated with OH formation (hydrogenation)-deformation (dehydrogenation) of the amorphous aluminosilicate matrix formed in situ at the SiO₂/ γ -Al₂O₃ hetero interface, and the redox reactions were governed by the H₂ partial pressure at 250–500 °C. As a result, a supported mesoporous γ -Al₂O₃/Co-doped amorphous SiO₂/mesoporous γ -Al₂O₃ three-layered composite membrane exhibited an H₂-triggered chemical valve property: mesopores under H₂ flow (open) and micropores under He flow (closure) at 300-500 °C.

Keywords: Co(III); aluminosilicate; reversible Co(III)/(II) redox property; membrane; H₂-triggered chemical valve property

1. Introduction

Microporous amorphous silica (SiO₂)-based membranes exhibit excellent hydrogen (H₂)selectivity and higher thermal stability compared to organic polymer membranes [1–5]. The SiO₂ membrane component is also attractive for developing novel mixed matrix membranes combined with a polymer membrane component such as polysulfone [6], which shows better membrane performance with improved mechanical properties. Moreover, SiO₂ membranes can be further modified as SiO₂-based organic-inorganic hybrid membranes for other applications such as liquidphase separation [7]. It has been recognized that high H₂-selectivity can be achieved by the molecular sieving property of amorphous SiO₂ with micro-pores of approximately 0.3 nm in size [3–5]. The microporosity can be formed in situ and is related to the highly branched fractal dimensions having a large amount of hydroxyl (Si–O–H) groups [8]. Accordingly, microporous SiO₂ membranes are moisture sensitive [9] and lack hydrothermal stability [10].

Recently, it has been reported that doping amorphous SiO₂ with metal (M) cations (M = Zr [11], Ni [12], Co [13–15]) improved the thermal or hydrothermal stability of sol–gel-derived SiO₂ membranes. Among the dopants, Co was found to be effective at enhancing H₂ permeance at 500 °C [16]. Nanostructure observation of the Co-doped SiO₂-based membranes revealed that CoO and Co₃O₄ nanoparticles of approximately 5–20 nm in size were distributed within the SiO₂ matrix [16]. More recently, SiO₂-based membranes synthesized from Co-modified ethoxy group-functionalized polysiloxane have been reported to show a unique reversible gas molecular sieving property above 300 °C [17]. The reversible gas molecular sieving property was suggested to be governed by alternating volume shrinkage/expansion due to the reversible change in the reducing/oxidizing (redox) state of the Co oxide particles (Co(OH)₂ and CoO/Co₃O₄) finely distributed within the polymer-derived SiO₂ matrix [17]. These results reveal that the divalent(II) or trivalent(III) Co cations that remained under H₂ atmosphere contributed to the enhanced H₂ permeance at 300–500 °C.

On the other hand, reactions between H₂ and various metal cations ($M^{m+} = Ni^{2+}$ [18], Fe³⁺ [19], Ce⁴⁺ [19], Sn⁴⁺ [19], V⁺⁵ [20], Cu⁺¹ [21], Ag⁺ [21]) in SiO₂ glass have been reported: The cations are reduced to lower valence state, whereas protons in the form of hydroxyl are introduced into SiO₂ glasses,

$$2(Si-O-Si)_{n}-O^{-}+2M^{m+}+H_{2} \rightarrow 2(Si-O-Si)_{n}-OH+2M^{(m-1)+}$$
(1)

$$2(Si-O-Si)_n-O^- + Sn^{4+} + H_2 \rightarrow 2(Si-O-Si)_n-OH + Sn^{2+}$$
 (2)

Analogous reactions have been reported for Cu^{2+} in zeolite derivatives [22], and other metal cations ($M^{m+} = Eu^{3+}[23-26]$, $Mn^{3+}[25]$, $Ce^{4+}[27]$) in Al₂O₃–SiO₂ glasses,

$$2(Si-O-Al)_{n}-O^{-}+2M^{m+}+H_{2} \rightarrow 2(Si-O-Al)_{n}-OH+2M^{(m-1)+}$$
(3)

In this study, Co cation-doped SiO₂/ γ -Al₂O₃ composite materials were synthesized; then, the reversible redox properties of Co cations around the hetero interface in the composites were intensively studied. First, a Co cation-doped SiO₂ thin film was deposited on a mesoporous γ -Al₂O₃ layer and an equilibrium chemical composition of the Al, Si, and Co within the mesoporous γ -Al₂O₃ layer was determined. Then, redox properties of the Co cations in the Co-doped SiO₂/ γ -Al₂O₃ composite as powders with a defined equilibrium chemical composition were investigated by cyclic heat treatment at 300–500 °C under alternative flow gas change of H₂ and Ar. Moreover, a unique H₂-triggered chemical valve property of the γ -Al₂O₃/Co-doped SiO₂/ γ -Al₂O₃ three-layered composite membrane is discussed based on the unique reversible redox reactions of the Co cation for the purpose of developing novel hydrogen production, storage, and transportation systems essential for the establishment of a hydrogen-based society.

2. Experimental Procedures

2.1. Sample Synthesis

Tetraethoxysilane (Si(OC₂H₅)₄, purity > 99.0%), aluminum nitrate nonahydrate (Al(NO₃)₃·9H₂O, purity 98.0%), cobalt(II) nitrate hexahydrate (Co(NO₃)₂·6H₂O, purity 98.0%), ethanol (C₂H₅OH, purity > 99.5%), and hydrogen peroxide (H₂O₂, 30% aq.) were purchased from Kishida Chemical Co. Ltd., Osaka, Japan.

In this study, for the sample syntheses of Co⁻-doped SiO₂/ γ -Al₂O₃ composites and Co⁻-doped aluminosilicate, Co⁻-doped SiO₂ precursor solutions with Co/Si = 1/8, 1/4, and 1/2 were prepared according to a previously reported procedure [16]. Here, as an illustration, we describe the preparation of the Co-doped SiO₂ sol with Co/Si = 1/8: A 10 mL round-bottom flask was charged with Co(NO₃)₂·6H₂O (0.343 g) and C₂H₅OH (4.2 mL). To this solution, 1.92 mL (8 mmol) of Si(OC₂H₅)₄ and

 H_2O_2 (2.3 mL, 30% aq) were added and maintained at RT with stirring for 1 h, then heated at 60 °C for 2 h.

Prior to the composite sample syntheses, the Co/Si ratio of the precursor-derived Co-doped SiO₂ was examined as follows: The Co-doped SiO₂ sols with different Co/Si ratios were heated with stirring to 80 °C and kept at this temperature overnight to give dried Co-doped SiO₂ gel powder samples. The dried gel powder samples were placed on a quartz boat in a quartz tube furnace (Model ARF-50KC, Asahi Rika Seisakusho Co. Ltd., Chiba, Japan), and heat-treated in air at 600 °C for 3 h with a heating/cooling rate of 20 °C·min⁻¹ to afford Co-doped SiO₂ powder samples. Then, elemental analyses were performed for Si and Co (ICP spectrometry, Model ICPS-7510, Shimazu Co., Kyoto, Japan), and carbon (C, non-dispersive infrared method, Model CS844, LECO Co., St Joseph, MI, USA). The residual carbon in the powder samples were negligible (<0.1%), and as shown in Table S1, we confirmed that the Co/Si ratios in the range of 1/8 to 1/2 were well controlled through the present sol–gel route using H₂O₂.

2.1.1. Co-Doped SiO₂/γ-Al₂O₃ Layered Composite Sample

Boehmite (γ -AlOOH) coating solution was prepared using the previously reported procedure [28] and spin-coated on a SiO₂ substrate (10 mm × 10 mm × 3 mm, Meijo science Co. Inc., Nagoya, Japan) at 3000 rpm using a spin coater (Model, MS-A100, Mikasa Co. Ltd., Tokyo, Japan). The γ -AlOOH-coated SiO₂ substrate was placed on an Al₂O₃ plate and loaded in an electric furnace (Model FUW220PA, Advantec Toyo Kaisha Ltd., Tokyo, Japan) then heat-treated in air at 600 °C for 3 h with a heating/cooling rate of 100 °C·h⁻¹ to afford the formation of a thin layer of γ -Al₂O₃ layer on the SiO₂ substrate by using the same manner as described above. The spin-coated sample substrate was placed on a quartz boat and loaded in a quartz tube furnace (Model ARF-50KC, Asahi Rika Seisakusho Co. Ltd., Chiba, Japan) then heat-treated in air at 600 °C for 20 h with a heating/cooling rate of 20 °C·min⁻¹. The sample was labelled CoSiOAlcoat.

2.1.2. Co-Doped SiO₂/γ-Al₂O₃ Composite Powder Sample

 γ -Al₂O₃ powder was prepared from γ -AlOOH sol using a previously reported procedure [28]. γ -Al₂O₃ powder was mixed with Co-doped SiO₂ sol with Co/Si = 1/2 at the Al:Si:Co ratio of 85:10:5. The resulting mixture was heated with stirring to 80 °C and kept at this temperature overnight to give a dried Co-doped SiO₂ gel/ γ -Al₂O₃ powder sample. The dried powders were placed on a quartz boat and set in a quartz tube furnace (Model ARF-50KC, Asahi Rika Seisakusho Co. Ltd., Chiba, Japan) then heat-treated in air at 600 °C for 20 h with a heating/cooling rate of 20 °C·min⁻¹. The sample was labelled CoSiOAlpow.

2.1.3. Homogeneous Co-Doped Aluminosilicate Powder Sample

To study the redox behavior of the Co cation in an amorphous aluminosilicate matrix, Co-doped aluminosilicate sol was prepared according to an atomic ratio of Al (in Al(NO₃)₃·9H₂O):Si (in Si(OC₂H₅)₄):Co (in CoNO₃·6H₂O) = 8:8:1. A 50 mL round-bottom flask equipped with a magnetic stirrer was charged with Co(NO₃)₂·6H₂O (0.343 g), Al(NO₃)₃·9H₂O (3.262 g), and C₂H₅OH (20 mL). To this solution, 1.92 mL (8 mmol) of Si(OC₂H₅)₄ and H₂O₂ (2.3 mL, 30% aq) were added and maintained at RT with stirring for 1 h, then heated at 60 °C for 2 h to give a Co-doped aluminosilicate sol. Then, the resulting sol was dried at 80 °C overnight to give the dried gel powder sample. The samples were placed on a quartz boat in a quartz tube furnace (Model ARF-50KC, Asahi Rika Seisakusho Co. Ltd., Chiba, Japan) and heat-treated in air at 600 °C for 3 h with a heating/cooling rate of 20 °C·min⁻¹. This sample was labelled CoSiOAlpow2.

As reference data, the textural properties were characterized for the composite powder samples. The surface morphology and the pore size distribution (PSD) of the CoSiOAlpow and CoSiOAlpow2 samples are shown in Figures S1 and S2, respectively. The CoSiOAlpow sample kept the morphology of the γ -Al₂O₃ polycrystallites (Figure S1a), while the CoSiOAlpow2 sample exhibited a featureless

Materials 2020, 13, 5345

glassy surface (Figure S1b). The N₂ adsorption and desorption isotherms of the CoSiOAlpow sample exhibited Type-IV isotherms according to the IUPAC classifications [29] and formed a hysteresis loop indicating the presence of mesoporosity, while those of CoSiOAlpow2 sample presented Type-II isotherms [29] without a distinct hysteresis loop (Figure S2a). The PSD curve characterized by the BJH method [30] for each powder sample was consistent with the N₂ adsorption-desorption behavior: The CoSiOAlpow sample exhibited a PSD curve with a dominant peak at 8 nm, while the CoSiOAlpow2 sample showed a broad and weak peak ranging from 20 to 100 nm (Figure S2b). The resulting Brunauer–Emmett–Teller (BET) surface areas of the CoSiOAlpow and CoSiOAlpow2 samples were measured to be 249 m²g⁻¹ and 94 m²g⁻¹, respectively.

2.1.4. γ-Al₂O₃/Co-Doped SiO₂/γ-Al₂O₃ Layered Composite Membrane Sample

A mesoporous γ -Al₂O₃ layer was formed on an outer surface of a macroporous α -Al₂O₃ support (tubular-type, 6 mm outer diameter, 2 mm thickness, and 60 mm length, porosity 40%, Noritake Co., Ltd., Aichi, Japan) according to a published procedure [28]. The mesoporous γ -Al₂O₃-modified tubular support was dipped into Co-doped SiO₂ sol with Co/Si = 1/4 at 10 mm·s⁻¹, kept in the sol for 30 s, then pulled out at 1 mm·s⁻¹ before heat treatment at 600 °C for 20 h under the same manner as that for the CoSiOAlpow sample synthesis. This dip-coating-heat treatment sequence was repeated once to form a Co-doped SiO₂ thin film on the supported mesoporous γ -Al₂O₃ layer. Then, the Co-doped SiO₂ thin film was coated with γ -AlOOH sol (diluted as 0.5 mol·L⁻¹) and heat-treated in air at 600 °C for 3 h by using the same manner as described above to give γ -Al₂O₃/Co-doped SiO₂/ γ -Al₂O₃ three-layered composite membrane. The sample was labelled CoSiOAlmemb.

2.2. Characterizations

2.2.1. Characterization Techniques

X-ray diffraction (XRD) measurements were performed on powder samples (Model X'pert Pro 1, Philips Ltd., Amsterdam, The Nederlands).

X-ray photoelectron spectroscopic (XPS) analysis was performed on the CoSiOAlpow2 sample with an Al K α X-ray source operated at 14 kV and 14 mA (Model PHI-5000, Ulvac-phi, Kanagawa, Japan). An alignment on the C 1s peak (at 284.8 eV) was performed before survey scans. To investigate the reversible redox property of the Co cation in the aluminosilicate, the CoSiOAlpow2 sample was heat-treated at 350 °C for 1 h with a heating/cooling rate of 5 °C·min⁻¹ under H₂ flow. Then, under Ar flow, the sample was heat-treated at 350 °C for 10 h with a heating/cooling rate of 5 °C·min⁻¹. The heat treatments were cyclically performed under flowing of H₂–Ar–H₂ or H₂–Ar–H₂–Ar using a catalyst analyzer (BELCAT-A, MicrotracBEL Corp., Osaka, Japan). Then, Co 2p spectrum was recorded for the heat-treated sample.

To investigate the distribution of Co cations within the CoSiOAlcoat sample, the depth-profiling XPS analysis was carried out by combining Ar ion gun etching cycles at 3 kV. The etching rate was calibrated as 3.73 nm min⁻¹ by measuring that of the SiO₂ substrate.

Surface morphology of CoSiOAlpow and CoSiOAlpow2 samples and the cross-section of the CoSiOAlmemb sample were observed by a scanning electron microscope (SEM, model JSM-6010LA, JEOL Ltd., Tokyo, Japan) operated at a voltage of 15 kV.

The BET surface area of the CoSiOAlpow and CoSiOAlpow2 samples was evaluated by measuring N₂ adsorption and desorption isotherms at –196 °C under relative pressure ranging from 0 to 0.99 (Model Belsorp Max, BEL Japan Inc., Osaka, Japan).

Nanostructure of the CoSiOAlpow sample was characterized by transmission electron microscopy (TEM) using a high-angle annular dark-field-scanning transmission electron microscope (HAADF-STEM, Model JEM-ARM200F, JEOL Ltd., Tokyo, Japan, operated at an accelerating voltage of 200 kV). The distribution of the constituent elements of Si, Al, Co, and O was measured and analyzed by the energy dispersive X-ray spectrometer (EDS, Model JED-2300, JEOL Ltd., Tokyo, Japan) mounted on a JEM-ARM200F.

Local structure of γ -Al₂O₃ powder sample was studied by ²⁷Al solid-state magic angle spinning nuclear magnetic resonance (MAS NMR) spectroscopic analysis with a 600 MHz NMR spectrometer (Model JNM-ECA600II, JEOL Ltd., Tokyo, Japan) operating at a static magnetic field of 14.01 T (155.4 MHz). Single gas permeances of the CoSiOAlmemb sample were evaluated for He and H₂ by the constant-volume manometric method [31]. The evaluations were performed at 100–500 °C according to the procedure as reported elsewhere [32,33]. The pressure on the gas feed side in this study was maintained at 120 kPa. On the gas permeate side, the gas line and the buffer tank with the total volume of 3.11×10^{-4} ·m³ were completely evacuated by using an oil rotary vacuum pump. After the evacuation was terminated, the pressure increase rate at the inside of the buffer tank was measured three times. The average of the three measurements was used for calculation of the permeance of gas-*i* (*Q*_{*i*}).

2.2.2. In Situ Characterizations

The hydrogenation/dehydrogenation reactions were monitored in situ for the CoSiOAlpow sample by diffuse reflectance infrared Fourier transform spectroscopy (DRIFTS, Model Spectrum 100, Perkin Elmer, Waltham, MA, USA). The Co-doped SiO₂/ γ -Al₂O₃ composite powder sample was loaded in a diffuse reflection cell (Model STJ900C Diffuse IR Heat Cham, S.T. JAPAN Inc., Tokyo, Japan), and IR spectrum was recorded after each treatment:

- heat-treatment to remove adsorbed water at 500 °C for 8 h under a flow of Ar (4 mL·min⁻¹).
- subsequent heat treatment at 500 °C for 0.5 h under a flow of H₂ (4 mL·min⁻¹).
- final heat treatment at 500 °C for 8 h under a flow of Ar (4 mL·min⁻¹).

Temperature-programmed reduction (TPR) and desorption (TPD) experiments were performed on the CoSiOAlpow sample using a catalyst analyzer (BELCAT-A, MicrotracBEL Corp., Osaka, Japan) fixed with a quadrupole mass spectrometer (BEL Mass, MicrotracBEL Corp., Osaka, Japan). The TPR profile was measured during heating from 100 °C to 500 °C (5 °C·min⁻¹) under a flow of 10% H₂/Ar (50 mL·min⁻¹). Then, the sample powder was maintained under pure-H₂ flow (50 mL·min⁻¹) for 1 h and cooled down to room temperature. The subsequent TPD measurement was carried out by heating up to 500 °C (5 °C·min⁻¹) under He flow (50 mL·min⁻¹). This cyclic TPR/TPD measurement was repeated 3 times.

In situ X-ray adsorption fine structure (XAFS) spectroscopic analyses were conducted at the BL5S1 of Aichi Synchrotron Radiation Center (AichiSR, Aichi Science & Technology Foundation, Seto, Aichi, Japan). The experimental set up is shown in Figure 1. The absorption spectrum can be obtained based on Equation (4), where μ and t are absorption coefficient and sample thickness, respectively.

$$\mu t = -\ln(I_1/I_0) \tag{4}$$

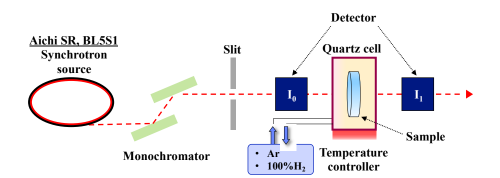


Figure 1. Experimental set up for in situ XAFS spectroscopic analysis.

The CoSiOAlpow sample (3.1 mg) was diluted with hexagonal boron nitride powder (100 mg, provided by the AichiSR, Seto, Aichi, Japan) using an agate mortar and pestle, then it was uniaxially pressed into a disk with a diameter of 7 mm. The disk sample was loaded in a quartz cell equipped with an electric heater and gas lines (Model 2000-1431, Makuharikagaku Garasu Seisakusho Inc., Chiba, Chiba, Japan) and maintained for 30 min under Ar flow. Then, under H₂ flow, the disk sample was heat-treated at 500 °C for 30 min with a heating rate of 8 °C·min⁻¹. Subsequently, under Ar flow, the furnace was cooled down to RT with a cooling rate of 8 °C·min⁻¹ then maintained at RT for additional 1 h. During this treatment, the spectrum was recorded stepwise at each measurement as follows: (1) at RT under initial Ar flow, (2) after H₂ treatment at 500 °C for 30 min, (3) at RT after being cooled down from 500 °C under Ar flow, and (4) at RT after being maintained under Ar flow for 1 h. In this study, the measurements were also performed on Co₃O₄ (4N, Kojundo Chemical Laboratory Co., Ltd., Saitama, Japan), CoO, and Co-foil (stored at AichiSR) at RT as reference samples. Then, to investigate the reversible redox property of the CoSiOAlpow sample, X-ray absorption near edge structure (XANES) spectroscopic analyses were performed using software (ATHENA, ver. 0.9.24/Demeter [34]).

3. Results and Discussion

3.1. Distribution of Co Cations within the Co-Doped SiO₂/ γ -Al₂O₃ Layered Composite Sample

To investigate the chemical composition at the hetero interface between the Co-doped SiO₂ and γ -Al₂O₃, a Co-doped SiO₂ thin layer was formed on a mesoporous γ -Al₂O₃ layer supported on a SiO₂ substrate. This sample labelled CoSiOAlcoat was deposited by spin-coating of the Co-doped SiO₂ sol with Co/Si = 1/8 followed by heat treatment in air at 600 °C for 20 h as detailed in the experimental section. Figure 2 presents the results of depth-profiling XPS analysis for the constituent elements of Si, Co, and Al.

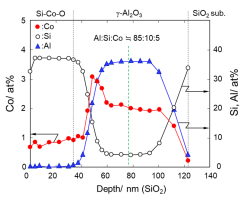


Figure 2. Depth-profiling XPS analysis for Co, Si, and Al within the CoSiOAlcoat sample.

The thickness of the Co-doped SiO₂ top layer was approximately 37 nm, estimated based on the Ar sputtering rate determined for the SiO₂ substrate. Then, almost all of the mesopore channels of the γ -Al₂O₃ layer with a thickness of about 85 nm were infiltrated by the Co-doped SiO₂. During the spincoating process, Co-doped SiO₂ sol solution easily penetrated into the mesoporous γ -Al₂O₃ layer. However, the depth profile of Co revealed that the Co/Si ratio of the top layer was approximately 1/37, much lower than the nominal composition of Co-doped SiO₂ coating sol (1/8). On the other hand, around the Co-doped SiO₂/ γ -Al₂O₃ hetero interface (around 50 nm in depth), the Co content increased to approximately 3 at.%, then, within the mesoporous γ -Al₂O₃ layer at 60–100 nm in depth, the Co content was shown to be constant at a value of ca. 2 at.% and the resulting Al:Si:Co ratio was 85:10:5.

The high Co concentration at the interface of the γ -Al₂O₃ layer was consistent with the previous report on the synthesis of Co-doped SiO₂ membrane on a porous Al₂O₃ support by Diniz Da Costa et al. [14]. The observed high concentration of Co around the hetero interface revealed preferential diffusion of Co cations from the SiO₂ matrix to the γ -Al₂O₃ surface. As shown in Figure S3, ²⁷Al solid-

state MAS-NMR spectroscopic analysis reveals that the γ -Al₂O₃ prepared in this study was composed of AlO₄ and AlO₆ units with the AlO₆/AlO₄ unit ratio of about 2.4, which is consistent with reported data [35]. Moreover, our previous study on mesoporous γ -Al₂O₃ revealed that a considerable amount of the Al–OH group existed at the γ -Al₂O₃ surface, which led to the highly hydrophilic property of γ -Al₂O₃ [28]. Accordingly, the driving force for the Co diffusion is due to the charge compensation for the AlO₄ site within the amorphous aluminosilicate formed in situ at the hetero interface, m[(SiO₄)(Al-O₄)]Co^{m+} (m = 2 or 3). Based on these results, the Al:Si:Co ratio of 85:10:5 was selected as an equilibrium chemical composition to investigate the redox property of Co cations at the SiO₂/ γ -Al₂O₃ hetero interface.

3.2. Properties of Co-Doped SiO₂/y-Al₂O₃ Composites

To investigate the later redox properties of Co cations at the SiO₂/ γ -Al₂O₃ hetero interface, we prepared powders labelled CoSiOAlpow with the equilibrium chemical composition (Al:Si:Co = 85:10:5). We first investigated the structural and microstructural changes of the material after H₂ treatment (the reductive condition) at 500 °C. As shown in Figure 3, the X-ray diffraction pattern of the CoSiOAlpow sample is composed of XRD peaks of γ -Al₂O₃, indicating that the Co-doped SiO₂ layer is X-ray amorphous. Then, after the H₂ treatment at 500 °C, it retains the X-ray amorphous state of the Co-doped SiO₂ phase.

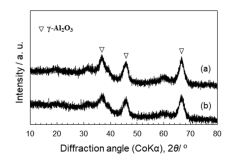


Figure 3. XRD patterns of the CoSiOAlpow sample. (**a**) As synthesized and (**b**) after H₂ treatment at 500 °C for 1 h.

TEM observation of the 500 °C H₂-treated CoSiOAlpow sample (Figure 4) also resulted in the detection of γ -Al₂O₃ as a single crystalline phase by the selected area electron diffraction ring pattern analysis (inset in Figure 4a), and the composite powder sample was composed of polycrystalline γ -Al₂O₃ of several nanometers in size (Figure 4b). Then, STEM-EDS mapping for the constituent elements of Al, Si, Co, and O within the area of the annular dark-field (ADF)-STEM image (Figure 5a) revealed homogeneous distribution of Co and Si as well as Al and O (Figure 5b–e). Accordingly, the composite sample powders were characterized as γ -Al₂O₃ polycrystalline homogeneously modified with Co-doped amorphous SiO₂.

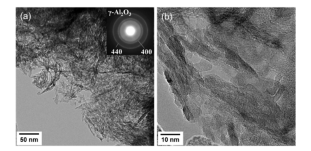


Figure 4. (a) TEM image of the CoSiOAlpow sample and the corresponding selected area electron diffraction pattern obtained, and (b) HRTEM image showing γ -Al₂O₃ polycrystalline of several nanometers in size.

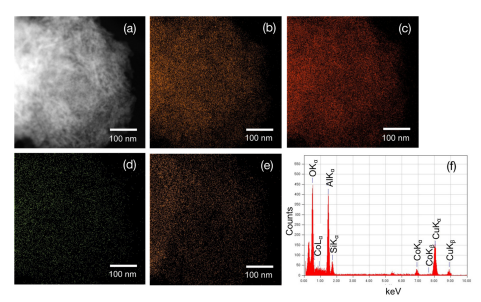


Figure 5. STEM-EDS mapping analysis for the CoSiOAlpow sample. (a) Annular dark field (ADF)-STEM image and the results of EDS mapping of (b) Al, (c) O, (d) Co, and (e) Si. (f) Typical EDS spectra recorded for the mapping analyses.

3.3. Redox Behavior of Co Species

Redox properties of the Co cation in the CoSiOAlpow sample were assessed by in situ using XAFS spectroscopic analyses. The CoSiOAlpow sample was heat-treated at 500 °C under H₂ flow followed by cooling to RT under Ar flow (Figure 6a). The spectra recorded at the measurement points shown in Figure 6a are shown in Figure 6b.

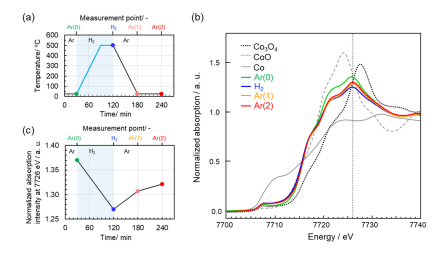


Figure 6. Results of in situ XAFS spectroscopic analysis for the CoSiOAlpow sample. (a) Heat treatment conditions and spectrum measurement points in this study, (b) XANES spectra of the CoSiOAlpow and reference samples, and (c) change in normalized absorption peak intensity at 7726 eV shown in (b).

The CoSiOAlpow sample at RT under the initial Ar flow (Ar(0) in Figure 6a) exhibited a spectrum with a peak at 7726 eV, and this spectrum was different from those of references, Co₃O₄, CoO, and Co-foil recorded at RT. The unique behavior is that the peak intensity at 7726 eV changes reversibly (Figure 6c): After the 500 °C H₂-treatment (H₂ in Figure 6a), the peak intensity decreases, i.e., approaches the intensity level of the Co-foil at 7726 eV, then increases in a stepwise manner under

Ar flow after cooling down to RT (Ar(1) in Figure 6a) followed by being maintained at RT for 1 h (Ar(2) in Figure 6a).

As discussed above, the metallic Co was not detected by our intensive HRTEM-STEM analyses. Thus, the spectra shown in Figure 6b suggested that the Co species in the CoSiOAlpow sample were Co(III) and Co(II) with the Co(III)/Co(II) ratio lower than that in Co_3O_4 (Co(III)/Co(II) = 2/1). Moreover, the peak intensity change shown in Figure 6c suggests the reversible redox property of the Co cation species in the CoSiOAlpow sample under the present cyclic condition of Ar (RT)-H₂ (500 °C)-Ar (RT) treatment.

To investigate the nature of the Co-doped SiO₂/ γ -Al₂O₃ hetero interface during the redox reactions of Co species, in situ DRIFT spectroscopic analysis was performed on the CoSiOAlpow sample under a cyclic heat-treatment condition under Ar flow, followed by 10% H₂/Ar flow and subsequent Ar flow at 500 °C. The results are shown in Figure S4. Due to the heterogeneous structure with the Co-doped SiO₂/ γ -Al₂O₃ hetero interface, the CoSiOAlpow sample exhibited broad spectra of OH groups in the range of 3800–3200 cm⁻¹: The initial DRIFT spectrum under Ar flow at 500 °C exhibited a broad peak around 3736 cm⁻¹ attributed to free Si–OH groups [36,37] together with a broader one from 3640–3300 cm⁻¹ assigned to hydrogen-bonded X–OH (X = Si or Al) group [23,38] (Figure S4a). After heat treatment under 10% H₂/Ar flow for 5 h at 500 °C, the intensity of the broader peak at 3640–3300 cm⁻¹ increased (Figure S4b), then decreased to be close to the initial intensity level after the subsequent Ar treatment at 500 °C (Figure S4c). Accordingly, it is suggested that the reduction-oxidation of Co cation species in the CoSiOAlpow sample is associated with OH formation-deformation, i.e., hydrogenation-dehydrogenation.

To identity the hydrogenation-dehydrogenation reactions, cyclic TPR/TPD analysis was performed on the CoSiOAlpow sample. In this attempt, the first TPR/TPD was conducted to initialize the oxidation state of the composite powder sample, then the TPR/TPD profiles were recorded for the 2nd and 3rd cycles. As shown in Figure S5, both the 2nd and 3rd TPR profiles showed a weak broad peak for H₂ up-take at 250–300 °C, and a larger one at 400–500 °C, then the H₂-uptake continued to some extent under the isothermal H₂-treatment at 500 °C. Both the 2nd and 3rd TPD profiles showed appropriate desorption peaks, a very weak peak at 250–350 °C, and another larger one during the isothermal Ar-treatment at 500 °C. The simultaneous MS analysis resulted in the detection of a trace amount of H₂ as a single desorption component at the two temperature regions appropriate to the desorption peaks.

These results suggest the following:

- The reduction-oxidation of Co cation species in the CoSiOAlpow sample is associated with the formation-deformation of OH groups.
- The redox reactions begin to proceed at relatively low temperatures around 250–350 °C.
- The redox reactions are reversible and governed by the H₂ partial pressure at $T \ge 250$ °C.

For further investigation on the redox reactions of the Co cation species that proceeded around 250–350 °C, we prepared another powder sample labelled CoSiOAlpow2 with homogeneous structure having almost same Co content but a higher Co/Al of 1/8 compared with that of CoSiOAlpow (1/17).

The redox behaviors of the Co cation species were investigated by cyclic heat treatment up to 350 °C. The CoSiOAlpow2 sample kept its X-ray amorphous state after the cyclic heat treatment at 350 °C under flow of H₂–Ar–H₂ or H₂–Ar–H₂–Ar (Figure 7A). As shown in Figure 7B, the XPS spectrum of as-synthesized CoSiOAlpow2 sample (Green line) exhibited a broad peak centered at 782 eV attributed to Co²⁺ together with another broad peak around 788 eV corresponding to a Co²⁺ satellite peak (labeled * in the graph) [39]. In addition, on the right side of the Co²⁺ peak, there was a peak shoulder at 779.9 eV assigned to Co³⁺ [39]. The Co³⁺ peak intensity at 779.9 eV apparently decreased after the second H₂ treatment of the H₂–Ar–H₂ run (blue line), while that due to metallic Co (777.8 eV) [39] was not observed. Then, after the subsequent Ar treatment (H₂–Ar–H₂–Ar run), the Co³⁺ peak intensity increased to some extent (orange line).

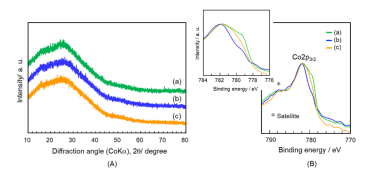


Figure 7. (**A**) XRD patterns and (**B**) XPS spectra of the CoSiOAlpow2 sample. (**A**) As-synthesized, after (**B**) heat treatment at 350 °C for 1 h under H₂ flow and the subsequent heat treatment at 350 °C for 10 h under Ar flow.

The oxidation state of the Co cation in the starting $Co(NO_3)_2$ ·6H₂O was divalent (II). Moreover, metallic Co was not formed under the present heat treatment conditions, and thus formation of Co(III) via the disproportionation reaction ($3Co^{2+} \rightarrow 2Co^{3+} + Co$) was excluded. It should be also noted that the oxidation state of the Co cation in the polymer-derived Co-doped amorphous SiO₂ was completely divalent [40].

It is well known that the Fenton reaction is the Fe(III/II)-catalyzed oxidation of organic compounds (RH) via generation of highly reactive radical species (\bullet OH) in the presence of H₂O₂ [41,42]:

$$Fe^{2+}(aq) + H_2O_2 \rightarrow Fe^{3+}(aq) + OH^- + \bullet OH$$
(5)

$$RH + \bullet OH \rightarrow \bullet R + H_2O$$
 (6)

Recently, Ling reported [43] that Co(II) analogously generates •OH in the presence of H₂O₂:

$$\operatorname{Co}^{2+}(\operatorname{aq}) + \operatorname{H}_2\operatorname{O}_2 \to \operatorname{Co}^{3+}(\operatorname{aq}) + \operatorname{OH}^- + \bullet\operatorname{OH}$$
(7)

In this study, H₂O₂ was used to prepare the precursor solutions for the Co-doped SiO₂ system, which resulted in the partial formation of Co(III) cations within the sample powders via the Fenton reaction. The results obtained in the present study reveal that reversible Co(III)/Co(II) redox reactions are associated with OH formation-deformation, which is governed by the H₂ partial pressure at 250–500 °C:

$$2(Si-O-Al)_{n}-O^{-} + \xrightarrow[-H_{2} \text{ in } Ar]{} 2Co^{3+} \cdot 2(Si-O-Al)_{n}-OH + 2Co^{2+}$$
(8)

The H₂ up-take/desorption of the CoSiOAlpow sample was mainly detected at higher temperatures around 500 °C (Figure S2). This could be explained by the kinetic factor necessary for H₂ diffusion through the amorphous SiO₂ network between the top surface and the SiO₂/ γ -Al₂O₃ hetero interface where Co cations mainly existed.

3.4. Gas Permeation Properties of γ -Al₂O₃/Co-doped SiO₂/ γ -Al₂O₃ Layered Composite Membrane

The cross-sectional SEM image of the layered composite membrane sample labelled CoSiOAlmemb is shown in Figure 8. In this study, the initial Co content of the SiO₂ layer was reduced from Co/Si = 1/2 to 1/4, then the Co-doped SiO₂ layer (Co/Si = 1/4) was placed between the two mesoporous γ -Al₂O₃ layers to investigate the relationship between the Co(III)/(II) redox reactions at the SiO₂/ γ -Al₂O₃ hetero interface and gas permeation properties. Initially, He permeance was measured at 100–500 °C. As shown in Figure 9a, initial He permeance (Q_{He} -1st) at 100 °C was 3.1 × 10⁻⁶ mol·m⁻²·s⁻¹·Pa⁻¹, then it slightly increased with the permeation temperature. On other hand, after H₂-treatment at 500 °C, the H₂ permeance (Q_{H2}) at 500 °C was slightly higher than that of Q_{He} -1st, and the Q_{H2} showed an opposite temperature dependency, with decreasing temperature, the Q_{H2} increased. Finally, after the subsequent He-treatment at 500 °C for 8 h, the He permeance (Q_{He} -2nd)

at 500 °C was close to that of Q_{He} -1st, then showed a positive temperature dependency similar to that observed for Q_{He} -1st.

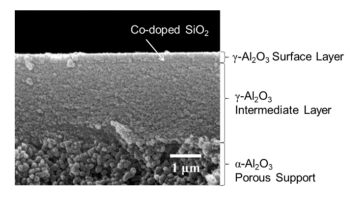


Figure 8. Cross-sectional SEM image of the CoSiOAlmemb sample.

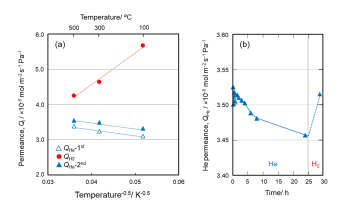


Figure 9. (a) Single gas permeation behaviors through the CoSiOAlmemb sample, and (b) time dependence of Q_{He} after H₂ treatment at 300 °C for 4 h.

The temperature dependency of the Q_{He} -1st and Q_{He} -2nd reveals that He dominantly permeates through micropore channels with a pore size close to the kinetic diameter of He (0.26 nm) [44], and both Q_{He} -1st and Q_{He} -2nd are governed by activated diffusion, theoretically expressed as,

$$Q_i = Q_0 \exp\left(-\frac{Ea}{RT}\right) \tag{9}$$

where Q_i is the permeance of permeate gas-i, Q_0 is the pre-exponential factor (mol·m⁻²·s⁻¹·Pa⁻¹), *Ea* is the activation energy (J·mol⁻¹), *R* is the gas constant (8.314 J·mol⁻¹·K⁻¹), and *T* is the temperature (K) [45,46].

On the other hand, as shown in Figure 9a, the Q_{H2} is proportional to the inverse square root of permeation temperature ($T^{-0.5}$), and is thus suggested to be governed by the Knudsen's diffusion mechanism. The theoretical gas permeance based on the Knudsen diffusion through a membrane having pore radius (d_p), porosity (ε), tortuosity (τ), and thickness (L) is formulated using the molecular weight of the permeating gas-i (M_i) and the permeation temperature (T) as follows,

$$Q_{i} = \frac{\varepsilon_{i} d_{p,i} \rho_{i}}{\tau_{i} L_{i}} \sqrt{\frac{8}{\pi M_{i} R T}}$$
(10)

where ρ is the probability of diffusion through the pore channel, ρ_i , ε_i , dp_i , τ_i , and L_i are membrane structural factors due to possible dependence on the size of permeate gas-i [47].

Knudsen diffusion occurs when the mean free path of the permeate gas molecule is relatively longer than the pore diameter, so the permeate gas molecules collide frequently with the pore wall. Generally, Knudsen diffusion is dominant for mesopores (2–50 nm) [28,47]. Since the kinetic diameter of He (0.26 nm) is smaller than that of H₂ (0.289 nm) [44], prior to the activated diffusion through micropores, He molecules should exhibit Knudsen diffusion characteristics by permeating through the mesopore channels where H₂ molecules dominantly permeate. These results suggest that the size of the gas permeable channel pores alternatively change from micropores under He flow to mesopores under H₂ flow, then back to micropores under He flow.

Time dependence of the Q_{He} was monitored after maintaining the CoSiOAlmemb sample under H₂ flow for 4 h. In this attempt, H₂ treatment and subsequent Q_{He} measurements were performed at the lower temperature of 300 °C to highlight the time dependence. As shown in Figure 9b, the initial Q_{He} of $3.53 \times 10^{-6} \text{ mol}\cdot\text{m}^{-2}\cdot\text{s}^{-1}\cdot\text{Pa}^{-1}$ slightly decreased with time and reached $3.45 \times 10^{-6} \text{ mol}\cdot\text{m}^{-2}\cdot\text{s}^{-1}\cdot\text{Pa}^{-1}$ after 24 h. Then, after the second H₂ treatment for 4 h, the Q_{He} regained close to its initial level. Accordingly, the size of pore channels under the H₂ flow became larger, and then reduced to be close to the initial size. This reversible change in pore size could be explained by the reversible redox of Co(III)/(II) associated with the OH group formation/deformation (Equation (8)), i.e., bond cleavageregeneration of the Si–O–Al amorphous network governed by the H₂ partial pressure at 300–500 °C.

4. Conclusions

In this study, Co-doped SiO₂/ γ -Al₂O₃ composite samples were synthesized by conventional solgel methods using H₂O₂. The redox behaviors of the Co cations in the composite samples were studied and the results can be summarized as follows:

- XPS depth profile analyses for the Co-doped SiO₂/mesoporous γ-Al₂O₃ layered composite sample revealed preferential Co diffusion from the SiO₂ top-layer matrix to the γ-Al₂O₃ surface, and the equilibrium chemical composition of the Al, Si, and Co within the mesoporous γ-Al₂O₃ layer was determined as Al:Si:Co = 85:10:5.
- XRD, HRTEM, and HAADF-STEM analyses revealed that the Co-doped SiO₂/γ-Al₂O₃ composite powder sample with the equilibrium composition of Al:Si:Co = 85:10:5 kept its amorphous state without crystallization of Co oxides after H₂ treatment at 500 °C.
- In situ XANES spectroscopic analyses during the 500 °C heat treatment under H₂ flow and subsequent cooling to RT under Ar flow exhibited reversible redox properties of Co cations in the Co-doped SiO₂/γ-Al₂O₃ composite powder sample.
- 4. XPS and DRIFT spectroscopic analyses and cyclic TPR/TPD measurements concluded the reversible Co(III)/Co(II) redox reactions were associated with OH formation (hydrogenation)deformation (dehydrogenation) within amorphous aluminosilicate, which was governed by the H₂ partial pressure at 250–500 °C.
- 5. The Co(III) cations in the present composite samples formed via the Fenton reaction in the presence of H₂O₂ during the sample syntheses.
- 6. Gas permeation measurements for the γ-Al₂O₃/Co-doped SiO₂/γ-Al₂O₃ layered composite membrane under the cyclic He–H₂–He flow suggested that H₂ triggered a chemical valve property: micropores under He flow (closure) and mesopores under H₂ flow (open), which could be explained by the reversible redox reactions of Co(III)/(II) associated with bond cleavage (hydrogenation)-regeneration (dehydrogenation) of the Si–O–Al amorphous network formed in situ at the hetero interface of the Co-doped SiO₂/γ-Al₂O₃.

The observed H₂-triggered chemical valve response needs a long time to function, especially for closure that takes approximately 8 h at 500 °C under Ar flow. The relations between the relative amount of Co(III) in the Si–O–Al amorphous network, recyclability of the redox reactions, and the valve response remain as future subjects.

Supplementary Materials: The following are available online at www.mdpi.com/1996-1944/13/23/5345/s1, Figure S1: SEM images of (a) CoSiOAlpow and (b) CoSiOAlpow2 samples., Figure S2: (a) N₂ adsorptiondesorption isotherms at –196 °C of CoSiOAlpow and CoSiOAlpow2 samples, and (b) the pore size distribution curves characterized by BJH plot., Figure S3: ²⁷Al solid-state MAS-NMR spectrum of γ -Al₂O₃ powder sample showing two peaks at 66 and 1 ppm assigned to AlO₄ and AlO₆ units, respectively., Figure S4: *In-situ* DRIFT

Materials 2020, 13, 5345

spectroscopic analysis for Co-doped silica/ γ -Al₂O₃ composite powder sample (CoSiOAlpow sample) with nominal composition of Al:Si:Co = 85:10:5. (a) as-synthesized, (b) after heat-treatment at 500 °C under 10%H₂-Ar flow for 8 h, and (c) after subsequent heat-treatment at 500 °C under Ar flow for 8 h., Figure S5: Results of cyclic TPR/TPD analysis for Co-doped silica/ γ -Al₂O₃ composite powder sample (CoSiOAlpow sample) with nominal composition of Al:Si:Co = 85:10:5. (a) 2nd and 3rd TPR profiles, (b) 2nd and 3rd TPD profiles and (c) H₂ ion current detected during the 3rd TPD profile measurement, Table S1: Chemical composition of 600 °C heat-treated Si–Co–O powder samples.

Author Contributions: Conceptualization, S.T. and Y.I.; methodology, T.A. and Y.D.; investigation, S.S., A.M., S.A. and H.M.; Formal analysis, T.A. and S.H.; writing—original draft preparation, S.T. and Y.I.; writing—review and editing, S.B.; visualization, S.B. and Y.I.; supervision, S.B. and Y.I.; funding acquisition, Y.I. All authors have read and agreed to the published version of the manuscript.

Funding: This work was partially supported by JSPS KAKENHI (Grant Number JP20K05076). Samuel Bernard and Yuji Iwamoto would like to thank CNRS who financially supported the present research work via the International Research Project (IRP) 'Ceramics materials for societal challenges'.

Acknowledgments: In situ XAFS spectroscopic analyses were conducted at the BL5S1 of Aichi Synchrotron Radiation Center (AichiSR, Aichi Science & Technology Foundation, Aichi, Japan).

Conflicts of Interest: The authors declare no conflict of interest.

References

- Lawal, S.; Kanezashi, M.; Nagasawa, H.; Tsuru, T. Development of an acetylacetonate-modified silicazirconia composite membrane applicable to gas separation. *J. Membr. Sci.* 2020, 599, 117844, doi:10.1016/j.memsci.2020.117844.
- Oyama, S.T.; Aono, H.; Takagaki, A.; Sugawara, T.; Kikuchi, R. Synthesis of Silica Membranes by Chemical Vapor Deposition Using a Dimethyldimethoxysilane Precursor. *Membranes* 2020, 10, 50, doi:10.3390/membranes10030050.
- 3. Prabhu, A.K.; Oyama, S. Highly hydrogen selective ceramic membranes: Application to the transformation of greenhouse gases. *J. Membr. Sci.* **2000**, *176*, 233–248, doi:10.1016/s0376-7388(00)00448-8.
- 4. Da Costa, J.D.; Lu, G.; Rudolph, V.; Lin, Y. Novel molecular sieve silica (MSS) membranes: Characterisation and permeation of single-step and two-step sol–gel membranes. *J. Membr. Sci.* 2002, *198*, 9–21, doi:10.1016/s0376-7388(01)00565-8.
- Iwamoto, Y. Precursors-Derived Ceramic Membranes for High-Temperature Separation of Hydrogen. J. Ceram. Soc. Jpn. 2007, 115, 947–954, doi:10.2109/jcersj2.115.947.
- Miricioiu, M.G.; Iacob, C.; Nechifor, G.; Niculescu, V.-C. High Selective Mixed Membranes Based on Mesoporous MCM-41 and MCM-41-NH₂ Particles in a Polysulfone Matrix. *Front. Chem.* 2019, 7, 332, doi:10.3389/fchem.2019.00332.
- Ren, X.; Tsuru, T. Organosilica-Based Membranes in Gas and Liquid-Phase Separation. *Membranes* 2019, 9, 107, doi:10.3390/membranes9090107.
- 8. Brinker, C.J.; Scherer, G.W. Sol-Gel Science, the Physics and Chemistry of Sol-Gel Processing; Academic Press: Boston, MA, USA, 1990; p. 908. ISBN 0-12-134970-5.
- Tanaka, K.; Sakata, Y. Present and Future Prospects of Hydrogen Production Process Constructed by the Combination of Photocatalytic H₂O Splitting and Membrane Separation Process. *Membranes* 2011, *36*, 113– 121, doi:10.5360/membrane.36.113.
- Miyajima, K.; Eda, T.; Nair, B.N.; Honda, S.; Iwamoto, Y. Hydrothermal stability of hydrogen permselective amorphous silica membrane synthesized by counter diffusion chemical vapor deposition method. *J. Ceram. Soc. Jpn.* 2013, *121*, 992–998, doi:10.2109/jcersj2.121.992.
- Yoshida, K.; Hirano, Y.; Fujii, H.; Tsuru, T.; Asaeda, M. Hydrothermal Stability and Performance of Silica-Zirconia Membranes for Hydrogen Separation in Hydrothermal Conditions. *J. Chem. Eng. Jpn.* 2001, 34, 523–530, doi:10.1252/jcej.34.523.
- 12. Kanezashi, M.; Asaeda, M. Hydrogen permeation characteristics and stability of Ni-doped silica membranes in steam at high temperature. *J. Memb. Sci.* 2006, 271, 86–93, doi:10.1016/j.memsci.2005.07.011.
- Igi, R.; Yoshioka, T.; Ikuhara, Y.H.; Iwamoto, Y.; Tsuru, T. Characterization of Co-Doped Silica for Improved Hydrothermal Stability and Application to Hydrogen Separation Membranes at High Temperatures. J. Am. Ceram. Soc. 2008, 91, 2975–2981, doi:10.1111/j.1551-2916.2008.02563.x.

- 14. Uhlmann, D.; Liu, S.; Ladewig, B.P.; Da Costa, J.C.D. Cobalt-doped silica membranes for gas separation. *J. Membr. Sci.* **2009**, 326, 316–321, doi:10.1016/j.memsci.2008.10.015.
- Yacou, C.; Smart, S.; Da Costa, J.C.D. Long term performance cobalt oxide silica membrane module for high temperature H₂ separation. *Energy Environ. Sci.* 2012, *5*, 5820–5832, doi:10.1039/c2ee03247c.
- Fujisaki, S.; Hataya, K.; Saito, T.; Arai, S.; Iwamoto, Y.; Kuroda, K. Nanostructural characterizations of hydrogen-permselective Si–Co–O membranes by transmission electron microscopy. *J. Mater. Res.* 2009, 24, 372–378, doi:10.1557/jmr.2009.0038.
- 17. Miller, C.R.; Wang, D.K.; Smart, S.; Da Costa, J.C.D. Reversible Redox Effect on Gas Permeation of Cobalt Doped Ethoxy Polysiloxane (ES40) Membranes. *Sci. Rep.* **2013**, *3*, srep01648, doi:10.1038/srep01648.
- Estournès, C.; Lutz, T.; Guille, J. Reduction of nickel in soda—lime—silicate glass by hydrogen. J. Non-Cryst. Solids 1996, 197, 192–196, doi:10.1016/0022-3093(95)00637-0.
- 19. Johnston, W.D.; Chelko, A.J. Reduction of Ions in Glass by Hydrogen. J. Am. Ceram. Soc. **1970**, *53*, 295–301, doi:10.1111/j.1151-2916.1970.tb12111.x.
- Smedskjaer, M.M.; Deubener, J.; Yue, Y. Inward Cationic Diffusion and Formation of Silica-Rich Surface Nanolayer of Glass. *Chem. Mater.* 2009, 21, 1242–1247, doi:10.1021/cm802513r.
- Chahadih, A.; El Hamzaoui, H.; Cristini, O.; Bigot, L.; Bernard, R.; Kinowski, C.; Bouazaoui, M.; Capoen, B. H₂-induced copper and silver nanoparticle precipitation inside sol-gel silica optical fiber preforms. *Nanoscale Res. Lett.* 2012, *7*, 487, doi:10.1186/1556-276X-7-487.
- 22. Bates, S.A.; Verma, A.A.; Paolucci, C.; Parekh, A.A.; Anggara, T.; Yezerets, A.; Schneider, W.F.; Miller, J.T.; Delgass, W.N.; Ribeiro, F.H. Identification of the active Cu site in standard selective catalytic reduction with ammonia on Cu-SSZ-13. *J. Catal.* **2014**, *312*, 87–97, doi:10.1016/j.jcat.2014.01.004.
- Nogami, M. Reduction Mechanism for Eu Ions in Al₂O₃-Containing Glasses by Heat Treatment in H₂ Gas. J. Phys. Chem. B 2015, 119, 1778–1784, doi:10.1021/jp511513n.
- Nogami, M.; Koiwai, A.; Nonaka, T. Control of Oxidation State of Eu Ions in Na₂O-Al₂O₃-SiO₂ Glasses. J. Am. Ceram. Soc. 2016, 99, 1248–1254, doi:10.1111/jace.14111.
- Van Tuyen, H.; Nonaka, T.; Yamanaka, K.-I.; Chau, P.M.; Hai, N.T.Q.; Quang, V.X.; Nogami, M. Fluorescence properties of valence-controlled Eu²⁺ and Mn²⁺ ions in aluminosilicate glasses. *J. Lumin.* 2017, 184, 83–88, doi:10.1016/j.jlumin.2016.12.020.
- Van Tuyen, H.; Nogami, M.; Hung, L.X. Reduction of Sm³⁺ and Eu³⁺ ions-co-doped Al₂O₃–SiO₂ glasses and photoluminescence properties. *Opt. Mater.* 2020, 100, 109639, doi:10.1016/j.optmat.2019.109639.
- Chau, P.T.M.; Hai, N.T.Q.; Quang, V.X.; Van Tuyen, H.; Nogami, M. Control Valence and Luminescence Properties of Cerium Ions in Al₂O₃–SiO₂ Glasses Prepared by Sol–Gel Method. *J. Electron. Mater.* 2019, 48, 6972–6977, doi:10.1007/s11664-019-07508-9.
- Kubo, M.; Kojima, M.; Mano, R.; Daiko, Y.; Honda, S.; Iwamoto, Y. A hydrostable mesoporous γ-Al₂O₃ membrane modified with Si–C–H organic-inorganic hybrid derived from polycarbosilane. *J. Membr. Sci.* 2020, 598, 117799, doi:10.1016/j.memsci.2019.117799.
- 29. Sing, K.S.W. Reporting physisorption data for gas/solid systems with special reference to the determination of surface area and porosity (Recommendations 1984). *Pure Appl. Chem.* **1985**, *57*, 603–619, doi:10.1351/pac198557040603.
- Barrett, E.P.; Joyner, L.G.; Halenda, P.P. The Determination of Pore Volume and Area Distributions in Porous Substances. I. Computations from Nitrogen Isotherms. J. Am. Chem. Soc. 1951, 73, 373–380, doi:10.1021/ja01145a126.
- Barrer, R.M.; Rideal, E.K. Permeation, diffusion and solution of gases in organic polymers. *Trans. Faraday* Soc. 1939, 35, 628–643, doi:10.1039/tf9393500628.
- 32. Iwamoto, Y.; Sato, K.; Kato, T.; Inada, T.; Kubo, Y. A hydrogen-permselective amorphous silica membrane derived from polysilazane. *J. Eur. Ceram. Soc.* **2005**, *25*, 257–264, doi:10.1016/j.jeurceramsoc.2004.08.007.
- Inada, T.; Uno, N.; Kato, T.; Iwamoto, Y. Meso-Porous Alumina Capillary Tube as a Support for High-Temperature Gas Separation Membranes by Novel Pulse Sequential Anodic Oxidation Technique. *J. Mater. Res.* 2005, 20, 114–120, doi:10.1557/jmr.2005.0016.
- Ravel, B.; Newville, M. ATHENA, ARTEMIS, HEPHAESTUS: Data analysis for X-ray absorption spectroscopy using IFEFFIT. J. Synchrotron Radiat. 2005, 12, 537–541, doi.:10.1107/S0909049505012719.
- Barrow, B.N.S.; Scullard, A.; Collis, N. Surface Selective 1H and 27Al MAS NMR Observations of Strontium Oxide Doped γ-Alumina. *Johns. Matthey Technol. Rev.* 2016, 60, 90–97, doi:10.1595/205651316x690943.

- Silverstein, R.M.; Bassler, G.C.; Morrill, T.C. Spectrometric Identification of Organic Compounds; John Wiley & Sons: New York, NY, USA, 1991; pp.135–139. ISBN 4-8079-0506-6.
- Mouline, Z.; Asai, K.; Daiko, Y.; Honda, S.; Bernard, S.; Iwamoto, Y. Amine-functionalized polycarbosilane hybrids for CO₂-selective membranes. *J. Eur. Ceram. Soc.* 2017, 37, 5213–5221, doi:10.1016/j.jeurceramsoc.2017.04.008.
- Chakarova, K.; Drenchev, N.; Mihaylov, M.; Nikolov, P.; Hadjiivanov, K. OH/OD Isotopic Shift Factors of Isolated and H-Bonded Surface Silanol Groups. J. Phys. Chem. C 2013, 117, 5242–5248, doi:10.1021/jp400106s.
- Kengne, B.-A.F.; Alayat, A.M.; Luo, G.; McDonald, A.G.; Brown, J.D.; Smotherman, H.; McIlroy, D.N. Preparation, surface characterization and performance of a Fischer-Tropsch catalyst of cobalt supported on silica nanosprings. *Appl. Surf. Sci.* 2015, 359, 508–514, doi:10.1016/j.apsusc.2015.10.081.
- 40. Tada, S.; Ando, S.; Asaka, T.; Daiko, Y.; Honda, S.; Bernard, S.; Iwamoto, Y. Hydrogen transport property of polymer-derived cobalt cation-doped amorphous silica. *Inorg. Chem. Front.* **2020**, doi:10.1039/d0qi01035a.
- 41. Fenton, H.J.H. LXXIII. Oxidation of tartaric acid in presence of iron. J. Chem. Soc. Trans. 1894, 65, 899–910, doi:10.1039/ct8946500899.
- Puppo, A.; Halliwell, B. Formation of hydroxyl radicals from hydrogen peroxide in the presence of iron. Is haemoglobin a biological Fenton reagent? *Biochem. J.* 1988, 249, 185–190, doi:10.1042/bj2490185.
- Ling, S.K.; Wang, S.; Peng, Y. Oxidative degradation of dyes in water using Co²⁺/H₂O₂ and Co²⁺/peroxymonosulfate. J. Hazard. Mater. 2010, 178, 385–389, doi:10.1016/j.jhazmat.2010.01.091.
- Breck, D.W. Zeolite Molecular Sieves: Structure, Chemistry, and Use; John Wiley & Sons: New York, NY, USA, 1974; pp. 634–636. ISBN 978-0471099857.
- Yoshioka, T.; Tsuru, T.; Asaeda, M. Molecular dynamics study of gas permeation through amorphous silica network and inter-particle pores on microporous silica membranes. *Mol. Phys.* 2004, 102, 191–202, doi:10.1080/00268970310001649383.
- Yoshioka, T.; Kanezashi, M.; Tsuru, T. Micropore size estimation on gas separation membranes: A study in experimental and molecular dynamics. *AIChE J.* 2013, *59*, 2179–2194, doi:10.1002/aic.13966.
- Malek, K.; Coppens, M.-O. Knudsen self- and Fickian diffusion in rough nanoporous media. J. Chem. Phys. 2003, 119, 2801–2811, doi:10.1063/1.1584652.

Publisher's Note: MDPI stays neutral with regard to jurisdictional claims in published maps and institutional affiliations.



© 2020 by the authors. Licensee MDPI, Basel, Switzerland. This article is an open access article distributed under the terms and conditions of the Creative Commons Attribution (CC BY) license (http://creativecommons.org/licenses/by/4.0/).