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# **ARTICLE**

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# Designing a nickel (II) thiourea-formaldehyde polymer/nanocarbon bifunctional molecular catalyst with superior ORR, OER activities and its application to Zn-air battery†

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As the efficient electrodes in energy conversion and storage devices, we focused on the development of a polymer-type non-precious metal-coordinated eco-friendly catalyst with a high performance, which is of importance in flexible wearable energy storage electronic devices with adaptable shapes. In the present study, we describe the design and synthesis of a nickel (Ni)-coordinated poly(thiourea-formaldehyde) polymer, then a nanocarbon (Vulcan, porous nanocarbon CNovel™, 2differernt multi-walled carbon nanotubes, or single-walled carbon nanotubes) as the conducting support was combined to prepare seven different molecular catalysts. The chemical structure of the Ni-coordinated polymer was characterized by elemental analysis, 13C NMR, FT-IR and XPS. Furthermore, EXAFS studies and theoretical calculations revealed that this polymer is consists of two Ni-O (2.17 Å) and two Ni-S (2.17 Å) coordination bonds. This Ni-coordination polymer was stable in an oxidative and reductive environment in alkaline medium for long term due to its strong Ni-coordination. The catalysts were found to act as an efficient catalyst for an oxygen reduction reaction (ORR) and oxygen evolution reaction (OER). Especially, among the catalysts, the CNovel<sup>TM</sup>-based catalyst showed the highest oxygen electrode performance for the ORR:  $E_{1/2}$ : 0.81V vs. RHE, and OER: 1.57 V vs. RHE at 10 mA/cm<sup>2</sup>. The catalyst acted as an efficient and durable cathode for a rechargeable Zn-air battery (charge-discharge overpotential gap of 0.45 V). Such an outstanding air-cathode performance is explained by the cooperative mechanism between water on the axial site of the Ni in the polymer and superoxide ion on the porous carbon. The present study is of importance for the development of advanced energy materials in batteries and molecular catalysts.

### 1. Introduction

As fossil fuel energy is soon going to run out, the world is looking for alternative energy sources and devices. The metal-air battery and water electrolyzer are some of such successful candidates for future energy requirement. 1-8 Recently, transition metal chalcogenides that exhibited a high electrocatalytic activity with low overpotentials have been reported as successful cost-effective cathode materials for batteries.9-17 Interestingly, the electrochemical transformation of transition metal chalcogenide cathodes showed a high activity and durability compared to their pristine transition metal oxide analogs9-12; namely, such modified metal chalcogenide performances have competed with those of the state-of-the-art catalysts, such as Pt, IrO<sub>2</sub>, and RuO<sub>2</sub>. 11-13 However, the performance and durability of such catalysts are still not appreciable compared to those of state-of-the-art catalysts. 12-15 Recent reports showed that the surface oxidation of the transition metal sulfides is responsible for the lack of durability.<sup>13-25</sup> In order to reduce the surface oxidation during long-term battery operations, several methods including the use of a stable spinel metal sulfide structure, incorporation of iron in the catalysts, etc., have been reported.<sup>26-28</sup> It will be hard to design a catalyst that is highly structurally stable during charging and discharging. Hence, use of the polymer-based catalyst in a fuel cell is one of the efforts to solve this issue.

Recently, we reported that the iron-nickel-sulfides/carbon nanotube hybrid catalyst exhibited a high ORR ( $E_{1/2}=0.82$  V), OER (potential @ 10 mA cm<sup>-2</sup> = 1.55 V), and Zn-air battery and water electrolyzer performances<sup>26</sup>. In this study, to further improve the stability of the metal sulfide active sites, we focused on a polymer-type catalyst, which is used for the development of a flexible eco-friendly catalyst. The presented catalyst is a Ni-coordinated flexible polymer-type molecular catalyst. The Ni is an earth-abundant inexpensive non-precious metal, and it is well-known that the Ni provides efficient catalyst sites for both ORR and OER<sup>26</sup>. Along this line, we newly designed and synthesized a nickel-coordinated thiourea formaldehyde-based polymer, denoted as (Ni-TUF)n, by a polycondensation reaction, then fabricated catalysts by combining it with a carbon support. As the carbon support, we selected carbon black (Vulkan), porous carbon (CNovel<sup>TM</sup> denoted as CN) and single-

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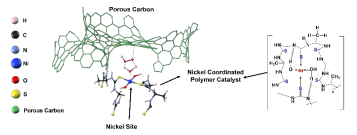
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walled carbon nanotubes (SWCNTs: ZEONNANO SG101: the so-called "Super-growth SWCNTs and multi-walled carbon nanotubes (Nikkiso-MWCNTs<sup>29-31</sup> and Flotube-9000). CN is a carbon black having a high surface area (~1800 m<sup>2</sup>/g), pore volume of ~3 mL/g with a particle size of 3.5~150 nm and tap density of 0.02~0.2 g/mL in its inter-connected mesopores with a 3-D structure. Notably, CN has a unique nanostructure; namely, it has a continuous porous structure (see SEM in the ESI, Fig. S1). Fig. 1 shows a possible structure of a catalyst composed of the nickel coordinated thiourea polymer with a porous carbon (left) and a possible chemical structure of the Nicoordinated polymer. Such a structure is demonstrated based on <sup>13</sup>C NMR, FT-IR, XPS and extended X-ray absorption fine structure (EXAFS) studies. The synthesized (Ni-TUF)n polymer-based catalysts shows a superior oxidation reduction reaction (ORR), and oxidation evolution reaction (OER). Especially, the catalyst composed of the polymer and CN exhibits better performances. Thus, it was used as a cathode of a Zn-air battery, and the fabricated battery showed a superior performance. We discuss the cooperative mechanism of H<sub>2</sub>O on the nickel and superoxide ion on the carbon that are responsible for the outstanding performance.



**Fig. 1.** Pictorial representation of a catalyst composed of the nickel coordinated thiourea polymer with a porous carbon (CNovel<sup>™</sup> or CN), (left) and a possible chemical structure of the Ni-coordinated thiourea polymer (right).

## 2. Experimental section

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**2.1.** Materials and Methods. Vulkan (XC-72R) and CNovel<sup>TM</sup> were obtained from Cabot Chemical Co., Ltd, and Toyotanso Co.,LTD, respectively. Flotube 9000 (CNano) was purchased from the Hohsen Corp. The Nikkiso-MWCNTs and SWCNTs (ZEONNANO SG101) were provided from Nikkiso Co., Ltd, and Zeon Corp., respectively. Thiourea, nickel (II) nitrate hexahydrate and formaldehyde were purchased from the FUJIFILM Wako Pure Chemical Corporation. Potassium hydroxide, dimethyl acetamide and Nafion® were obtained from Sigma Aldrich. The PTFE membrane with a 3-μm pore diameter was purchased from Advantec Co., Ltd.

The catalysts were characterized by elemental analysis, nuclear magnetic resonance (NMR), and X-ray diffraction (XRD, Rigaku Smartlab, 9kW AMK model). The structural morphologies of the catalysts were investigated by field-emission scanning electron microscopy (FE-SEM, Joel, JSM-7900F, accelerating voltage = 5 kV). For the SEM characterization, the samples were coated by osmium sputtering. Transmission electron microscopy (TEM) measurements were carried out using a JOEL (JEM-ARM200CF model) microscope (accelerating voltage: 80 kV for carbon samples and 200 KV for non-carbon samples). The catalysts were dispersed in isopropanol by

sonication using a bath-type sonicator (Branson 5510) and fewedrops of the suspension were drop-cast onto a carbon-coated copper field, and then dried for the TEM measurements. The X-ray photoelectron spectroscopy (XPS) of the catalysts was performed in a 10-9 mbar vacuum using an AXIS-ULTRA (Shimadzu, Co., Japan).

The XAFS (in fluorescence-detection mode) measurements were carried out using the Kyushu University beam line, BL 06, in the Saga Light Source (Tosu, Japan), in which a homemade *in situ* electrochemical XAFS cell was used. The EXAFS fitting was conducted using the equations described below.

$$\chi(k) = \frac{X_{j} S_{0}^{2} N_{j} f_{j}(k) e^{-2Rj/\lambda(k)} e^{-2k \cdot 2 \cdot \sigma \cdot 2 \cdot j} \quad X \sin[2kRj + \delta j(k)]}{kR_{j}^{2}}$$

$$k = 2\pi\lambda = \frac{\sqrt{2me(E - E0)}}{h^2}$$
 .....(2)

where k = photoelectron wavenumber, E = X-ray energy,  $E_0$  = energy of the absorption edge,  $S_0^2$  = absorption reduction factor,  $N_j$  = Coordination number,  $R_j$  = interatomic distances,  $\sigma_j$  = mean-square disorder in the distance for the j<sup>th</sup> shell (here j = K-edge is followed),  $E_j$  = photoelectron (back-) scattering amplitude,  $E_j$ (k) = back-scattering phase for the j<sup>th</sup> atomic shell,  $E_j$ (k) = photoelectron inelastic mean free path (ranges 1-100Å),  $E_j$ (k) = dimensionless, and  $E_j$  = unit of 1/k,  $E_j$ (k) ≈ -2a<sub>0</sub>k (a<sub>0</sub> is the Bohr radius).

$$R_{(EXAFS)} \approx R_j + (\Delta u^2)/2R_j$$
 .....(3

where  $\Delta u$  is the harmonic approximation.

- **2.2.** Synthesis of Nickel/Thiourea Formaldehyde Polymer (NiTUF)n. A 0.2 mol of thiourea was mixed with 0.2 mol of formaldehyde (5mL) to which acetic acid was dropped until the pH of the mixture reached pH 3. A 1 mol sample of Ni(NO $_3$ ) $_2$  was added to the mixture, stirred well, then refluxed for 6 h at  $90^{\circ}$ C. The obtained viscous product was collected and washed well with 1 M NaOH, de-ionized water and methanol, then dried at room temperature for 1 day to obtain a light green compound, (Ni-TUF)n. Also, we synthesized (TUF)n not containing Ni.
- **2.3. Synthesis of (Ni-TUF)n/Carbon Composites.** A typical synthetic procedure is as follows. The prepared (Ni-TUF)n was sonicated in 40 mL of dimethyl acetamide (DMAc) for 2 h to which a 50 mg of a carbon support material listed in Table 1 was added, then sonicated for 30 min. The obtained material was then filtered to remove the solvent, washed with methanol, followed by vacuum drying at room temperature overnight. Table 1 lists the obtained materials/catalysts, which were used for further structure characterization and electrochemical performance measurements including the ORR, OER and Zn-air battery performances in alkaline media.

**Table 1.** List of Synthesized Catalysts.

Amount	Carbon Support (each	Abbreviation of catalyst
of (Ni-	50mg)	
TUF)n		

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10mg	Carbon Black (Vulcan XC72R)	(Ni-TUF)n/Vulkan w/w:1/5 )
10 mg	MWCNTs (Flotube 9000)	(Ni- TUF)n/MWCNT(Flotube) w/w:1/5)
10 mg	MWCNTs (Nikkiso)	(Ni- TUF)n/MWCNT(Nikkiso) w/w:1/5)
10 mg	SWCNTs	(Ni-TUF)n/Zeon-CNT- w/w:110
5 mg	C-Novel™ (MH-00)	(Ni-TUF)n/CN-w/w:1/10
10 mg	C-Novel™ (MH-00)	(Ni-TUF)n/CN- w/w:1/5
30 mg	C-Novel™ (MH-00)	(Ni-TUF)n/CN- w/w:3/5

2.4. Electrochemical measurements. The ORR and OER were evaluated by linear sweep voltammetry (LSV) in an  $O_2$  saturated Ar atmosphere using an ALS CH instrument (model 704 EA). A typical three-electrode system with working, reference and counter electrodes were immersed in 0.1 and 1 M potassium hydroxide (KOH) aqueous electrolytes. Mercury/mercurous oxide (Hg/HgO) and a platinum coil purchased from BAS, Inc., were used as the reference electrode and the counter electrode, respectively. Both the OER and ORR measurements were carried out at the scan rate of 5mVs-1. To compare the OER and ORR activities of the catalysts, commercial Pt/C (40%) and RuO2 catalysts were used. The catalyst ink was prepared using 5 mg of the catalyst dispersed in a 200 µL solution mixture consisting of isopropanol, DI water, and Nafion (5%) in the ratio of 16:3:1, and dispersed using a bath-type sonicator (Branson 5540) for 30 min to obtain a homogeneous ink. A 1.5  $\mu L$  aliquot was uniformly dispersed on a glassy carbon disk with an area of 0.07 cm<sup>2</sup>, then dried. The potentials were converted to the reversible hydrogen electrode (RHE) scale using the relationship RHE = E (Hg/HgO) + 0.91 V in 0.1 M KOH and RHE = E (Hg/HgO) + 0.98 V in the 1 M KOH electrolyte. The 0.91 and 0.98 V were obtained from the calibration of the Hg/HgO electrode under complete H<sub>2</sub> saturation conditions in 0.1 and 1 M KOH, respectively.

**2.5. Zn-air Battery Assembly.** The Zn-air battery was assembled using an air battery set-up (Rigaku Corporation). We used gas a diffusion layer (GDL) for the electrode preparation. A zinc sheet and Whatman filter paper (51-55-μm pore size) in 6M KOH was used as the anode and separator. The other set of the catalyst dispersion was then poured into the same membrane filter to make another layer to obtain a two-sided catalyst-coated cathode electrode, which was used as the Zn-air battery assembly (working area = 28.27 cm²). The mass of the zinc metal was calculated by weighing the Zn metal before (for each battery assembly, we used 400 mg of Zn metal) and

after the battery performance measurements. The specific capacity was calculated by the following eg. specific capacity with a specific capacity with the following eg. specific capacity with the specific capacity with the specific capacity was calculated by the following eg. specific capacity with the specific capacity was calculated by the following eg. specific capaci

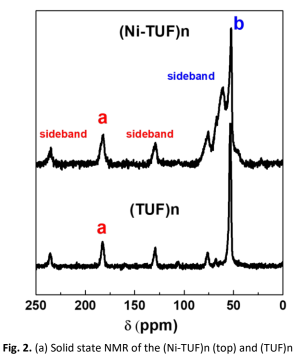
### 3. Results and discussion

**3.1. Synthesis and Structural Characterization**. The synthesis of a nickel coordinated thiourea-formaldehyde polymer (Ni-TUF)n was carried out using the method involving the synthesis of the thiourea-formaldehyde polymer (TUF)n without Ni. $^{31-38}$  The functional group unit in our polymer synthesized in the presence of Ni ions is expected to be -CH<sub>2</sub>-NH-C(=S)-NH-CH<sub>2</sub>- like the reported polymer synthesized in the absence of Ni. We carried out an elemental analysis of (Ni-TUF)n, and the result was: C = 27.64 %, H = 5.19 %, N = 25.62%, S = 29.7%, which indicated the elemental composition of (Ni-TUF)n as  $C_{15}H_{22}N_{12}S_6NiO_2$  (molecular weights,1937.86) because the calculated composition percentage for  $C_{15}H_{22}N_{12}S_6NiO_2$  is: C = 27.56.3 %, H = 3.40%, N = 25.72%, S = 29.44 %. For (TUF)n without Ni: C = 25.18 %, H = 5.07 %, N = 29.15%, S = 33.7%. For Calcd. of  $C_2H_5N_2S_1O_{0.5}$ , C = 24.72 %, H = 5.20 %, N = 28.84 %, S = 33.00 %.

We then measured the solid state C13 NMR spectra of (Ni-TUF)n and (TUF)n, and the result is shown in Fig. 2, in which we observed two kinds of carbons in the (TUF)n and three kinds of carbons in the (Ni-TUF)n. For the (TUF)n, the characterized peak as 'a' in Fig. 2 is due to the C=S splitting (182.34 ppm) and the 'b' carbon is due to the CH<sub>2</sub> splitting (53 ppm<sup>33,34</sup> For the (Ni-TUF)n with a nickel coordination, the chemical shift value observed at 183.1 ppm ('a' carbon) is due the C=S group, while the chemical shift at 52 ppm denotes the type 'b' carbon of the methyl group. 33 There are two bands in the 'b' carbon in the (Ni-TUF)n at 76 and 66 ppm which are broadened, while such bands are not evident in the (TUF)n). This would be due to the electronic environmental changes in the neighboring nitrogen atoms. The chemical shifts at 61 and 53 ppm of (Ni-TUF)n are due to the methyl groups with and without branching, respectively. The other chemical shifts at 129.6 and 236 ppm of (TUF)n are characterized as the spinning sidebands of the C=S carbon, and the chemical shifts at 76 and 69 ppm are the sidebands of the methyl group (CH<sub>2</sub>) carbon. Since the sidebands of the C=S carbon in the (Ni-TUF)n appeared at almost the same chemical shifts as the (TUF)n, the coordination center of the polymer was not clear based on the analysis of the solid state C13 NMR spectral data. The nature of the carbon in (Ni-TUF)n and (TUF)n was further investigated using FT-IR spectroscopy, and the results are shown in Fig. 3a and Table 2. The (Ni-TUF)n exhibits a strong band at 3318 cm<sup>-1</sup>, which is attributed to the stretching vibrations of the OH as depicted in Fig. 3b. 30-33 The (TUF)n shows a broad band in the range of 3270-2865 cm<sup>-1</sup>, which is due to the existence of adsorbed H<sub>2</sub>O molecules.

The (Ni-TUF)n exhibits absorption peaks in the range of 2903-3005 cm<sup>-1</sup>, which are attributed to the -CH- stretching vibrations.<sup>32-33</sup> We observed a band at 3143 cm<sup>-1</sup> for the (Ni-TUF)n, which would be due to the hydrogen bonding of the -NH- group and the hydrogen of the hydroxyl group coordinated to the nickel atom as presented in Fig. 3b.<sup>33,34</sup> This nitrogen in the -NH-group involved in the hydrogen bonding is responsible for the sideband broadening of the NMR spectrum of (Ni-TUF)n.<sup>34</sup> In Table 2, we list the assignment of the FT-IR vibration peaks of -NH-, N-C(=S)-N, -CH<sub>2</sub>-, and -N-CH<sub>2</sub>-N- for (Ni-

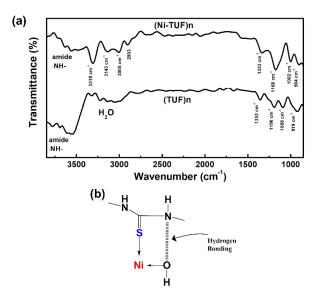
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(bottom).

TUF)n and (TUF)n analyzed based on the reported data in the literature.  $^{\rm 32\text{-}36}$ 

An X-ray photoelectron spectroscopy (XPS) study was carried out for (Ni-TUF)n and (TUF)n to obtain the binding energies of the S2p, N1s and C1s. As shown in Fig. 4a, the S2p binding energy bands of (Ni-TUF)n and (TUF)n appear at 161.0 and 160.0 eV, respectively, which corresponds to the S $^{2-}$  band. We observed a 1.0-eV positive shift in the (Ni-TUF)n, which would be derived from the complex formation between the sulfur and Ni. $^{13,19}$ 



**Fig. 3.** (a) FT-IR spectra of (Ni-TUF)n and (TUF)n and (b) suggested hydrogen bonding and Ni-S coordination.

•			· · · · · · · · · · · · · · · · · · ·		
	Wavenumber /cm <sup>-1</sup> for (Ni- TUF)n)	Wavenumber/c m <sup>-1</sup> for (TUF)n	Wavenumb er /cm <sup>-1</sup> for each attributed group taken in the literatures <sup>24-27</sup>	Attributed group	
	3143	Not found			
	1333	1358	1340	-NH-	
	1171	1196	1140	N-C(=S)-N	
	1088	1002	1086	-CH <sub>2</sub> -	
	945	902	919	-N-CH <sub>2</sub> -N-	
	812	828	Not found		
	710	717	709	-N- C(=S)-	

Table 2. Assignment of the FT-IR of (Ni-TUF)n and (TUF)n Article Online

In the N1s region, the (Ni-TUF)n and (TUF)n exhibit binding energies at 398.3 and 397.8 eV, respectively (Fig. 4b).<sup>38</sup> The 0.4 eV positive binding energy shifts in the (Ni-TUF)n compared to the (TUF)n would be explained by the coordination of the S and Ni through a possible hydrogen bonding between the hydrogen of the hydroxyl group coordinated to the nickel and nitrogen in the polymer chain as depicted in Fig. 5.

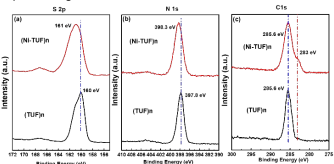
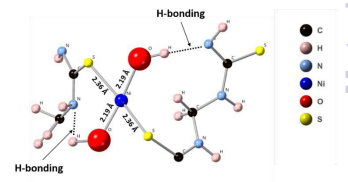


Fig. 4. (a), (b) and (c) are, respectively, the S2p, N1s and C1s XPS spectra of (Ni-TUF) n (red) and (TUF)n (black).



**Fig. 5.** A possible coordination structure and hydrogen bonding in Ni-(TUF)n.

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The (Ni-TUF)n and (TUF)n show binding energies at 285.6 eV (Fig. 4c) in the C1s region, which is due to the characteristics of the sp<sup>2</sup> carbon bonded to nitrogen. Unlike (TUF)n, the (Ni-TUF)n shows a shoulder peak at ~283 eV, which is due to the C-C bond, confirms the presence of the branched methyl group in the polymer chain. The FT-IR peak that appeared at 3143 cm<sup>-2</sup> supports this explanation.<sup>38-39</sup> We observed binding energies at 530.61 and 530.53 eV for (Ni-TUF)n and (TUF)n, respectively, suggesting that both compounds contain the oxygen moiety as water or a hydroxyl group (see the ESI, Fig. S2). The morphologies using SEM and TEM are shown in the ESI, Fig. S3, in which the formation of agglomerated structures is obvious. In Fig. S3 (c), the black-colored part would be Ni-sites and the other area would be CN.

3.2. Theoretical simulations. First principle simulations were performed using the periodic boundary condition, density functional theory with generalized gradient approximation, Pardew, Burke, Erzhofen (PBE) functional and numerical atomic orbitals wavefunction (NAO) as implemented in the QuantumATK simulation package.<sup>40</sup> Geometry optimization was carried out until the forces were minimized below 0.05 eV/Å<sup>2</sup>. In our simulations, we performed optimization of the linear polymer unit cell with relaxation of the atom positions and unit cell parameters. This optimization was followed by optimization of the 2 X 2 X 4 super cell which includes four polymer chains with four-unit cells each. This relaxation was

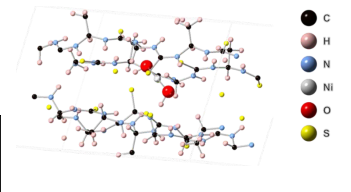


Fig. 6. Optimized supercell with four polymer chains containing Ni(OH)2 in Ni-(TUF)n.

performed to account for inter-chain interactions. Finally, Ni(OH)<sub>2</sub> was placed in the center of the simulations model. The geometry optimization, depicted in Fig. 6, resulted in a four-coordinated Ni center which leaves two free coordination sites in the sixcoordinated Ni structure for the catalytically active sites. The four ligands are two hydroxyl groups and two sulfurs from the polymer chain. Strong hydrogen bonds were formed between the OH group and -NH- groups from the polymer chain.

3.3. Extended X-ray Absorption Fine Structure (EXAFS) on Interder to reveal the bonding nature and geometry of the (NPTUF)/P, We can fed out the extended X-ray absorption fine structure (EXAFS) measurements, and the results were analyzed using the nickel as an observer atom, oxygen and sulfur as the scatter atoms (Fig. 6), and using the Ni-O and Ni-S paths. The variables of  $E_0$ ,  $S_0^2$ ,  $N_i$ ,  $R_i$ , and  $\sigma_i$ listed in Table S1 and the model equation described in Experimental (Materials and Methods section). For all the EXAFS data, the interatomic distances were obtained from solving the multiple path functions (Ni-O and Ni-S) and fitted using the first shell model in the radial distribution function results. The EXAFS analysis of the (Ni-TUF)n and (Ni-TUF)n/CN-w/w:1/5 suggested the 4 coordination number with the Ni-O and Ni-S of which 2 are from Ni-O and 2 are from Ni-S for both catalysts. Hence, both (Ni-TUF)n and (Ni-TUF)n/CN:w/w-1/5 exhibited the +2 oxidation state of nickel and planar geometry by the John-Teller distortion. Therefore, the axial ligands remain uncoordinated due to the elongation of the higher energy dx2-y2 orbital in the Ni+2 (d8-coordination geometry). These results agree with the theoretical simulation described above.

The EXAFS results of (Ni-TUF)n using the first shell fitting (Fig.7) showed that the Ni-O and Ni-S of (Ni-TUF)n exhibited the bond lengths of 2.18 and 2.35 Å, respectively, and for those of (Ni-TUF)/CN:w/w-1/5, the bond lengths of 2.17 and 2.38 Å, respectively. Previous reports<sup>41-43</sup> have suggested that the both the Ni-O and Ni-S bonds possess a similar bonding nature, but the Ni-O is slightly high by polar For the (Ni-TUF)n and (Ni-TUF)n/CN-w/w:1/5 in the ex situ mode, we observed a significant difference in the bond lengths of the Ni-O (2.19Å) and Ni-S (2.36Å) of about ±0.17Å (Table S2). This would be due to the hydrogen bonding of the hydrogen atom in the (Ni-TUF)n which is bonded to the highly electronegative oxygen with the nitrogen in the polymer chain (Fig. 3b), and this is further evident from the FT-IR vibrational frequency at 3143 cm<sup>-1</sup> that corresponds to the hydrogen bonding (Fig. 3b) as well as the theoretical studies

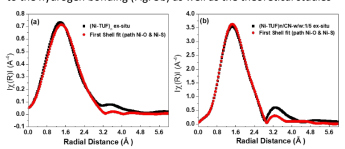


Fig. 7. Ex situ EXAFS (black color) of (a) (Ni-TUF)n and (b) (Ni-TUF)n/CN:w/w:1/5, and their first shell fit model (red color).

described above. Hence, the geometry in the (Ni-TUF)n and (Ni-TUF)n/CN-w/w:1/5 are not completely planar, instead, there are suggested to be slightly deviated from the planar structure. This was also suggested by the XRD of (Ni-TUF)n (see the ESI, Fig. S4), in which no crystalline peak was evident (very weak).

3.4. Electrochemical Studies. We measured the ORR and OER performances of the synthesized catalysts together with Pt/C or Ir/C for comparison in 0.1 M KOH aqueous solutions using 1600 rpm rotating disk electrodes to obtain the electrocatalytic activity. As shown in Fig. 8a, the ORR activity showed a strong dependence of

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the used carbon supports; namely, the half-wave potential  $(E_{1/2})$  of (Ni-TUF)n/CN-w/w:1/5 is 0.82 V vs. NHE, which is an excellent performance among the state-of-the-art non-Pt catalysts<sup>11,18</sup>, and superior compared to those of (Ni-TUF)n/MWCNT(Nikkiso)-w/w:1/5  $(E_{1/2}=0.65 \text{ V})$ , and (Ni-TUF)n/Vulcan--w/w:1/5  $(E_{1/2}=0.73 \text{ V})$  as listed in Table 3. As shown in Fig. 8b, the OER shows a slightly different behavior; that is, when comparing the performance of the voltage at 10 mA cm<sup>-2</sup>, (Ni-TUF)n/CN-w/w:1/5 provided 1.57 V, which is comparable but slightly lower than those of (Ni-TUF)n/Vulkan-w/w:1/5 (1.51V) and (Ni-TUF)n/MWCNT(Nikkiso)-w/w:1/5 (1.52 V) (Table 3).

Based on the obtained ORR and OER values, the oxygen electrode potentials of (Ni-TUF)n/CN-w/w:1/5, (Ni-TUF)n/MWCNT(Nikkiso)-w/w:1/5, and (Ni-TUF)n/Vulkan-w/w:1/5 were calculated to be 0.75, 0.87 and 0.79 V, respectively. This result demonstrates that the performance of (Ni-TUF)n/CN-w/w:1/5 is better than those of the other two catalysts. The CN as the support provides the best

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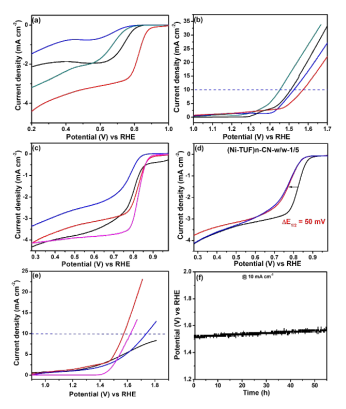


Fig. 8. (a) ORR and (b) OER of (Ni-TUF)n/CB-w/w:1/5 (black line), (Ni-TUF)n/Nikkiso-w/w:1/5 (blue), (Ni-TUF)n/CN-w/w:1/5 (red), and (Ni-TUF)n/Zeon CNT (green). (c) ORR activities of (Ni-TUF)n/CN-w/w:1/10 (black), (Ni-TUF)n/CN-w/w:1/5 (red) and (Ni-TUF)n/CN-w/w:3/5 (blue), together with that of Pt/C (purple), (d) Accelerated ORR durability of Ni-TUF)n/CN-w/w:1/5 (initial (black), after 10000 cycles (red), and after 20000 cycles (blue), (e) OER activities of Ni-TUF)n/CN-w/w:1/10 (black), Ni-TUF)n/CN-w/w:1/5 (red) and Ni-TUF)n/CN-w/w:3/5 (blue), together with that of IrO2 (purple). (f). The OER chronopotentiometric durability tests of Ni-TUF)n/CN-w/w:1/10.

performance among the three catalysts, which might be due to the highly mesoporous carbon structures that provide many efficient catalytic sites.

We then examined the weight ratios of (Ni-TUF)n and CN present in (Ni-TUF)n/CN-w/w:1/5, (Ni-TUF)n/CN-w/w:1/10 and (Ni-TUF)n/CNw/w:3/5. As shown in Fig. 8c, for the ORR, (Ni-TUF)n/CN-w/w:1/5 is better than (Ni-TUF)n/CN-w/w:1/10 and (Ni-TUF)n/CN-w/w:3/5, indicating the optimum ratio between the (Ni-TUF)n and carbon support is 10 mg of Ni-(TUF)n and 50 mg of the CN carbon. Notably, the ORR performance of (Ni-TUF)n/CN-w/w:1/5 is very close to that of the conventional Pt/C. One more issue we like to demonstrate is its high OER performance; namely, as described, when comparing the performance of the voltage at 10 mA cm<sup>-2</sup>, (Ni-TUF)n/CNw/w:1/5 showed better activity than those of (Ni-TUF)n/CNw/w:1/10 and (Ni-TUF)n/CN-w/w:3/5. We emphasize that the OER activity of (Ni-TUF)n/CN-w/w:1/5 is suite superior compared to the conventional IrO<sub>2</sub> (Fig. 8e). Such obtained results are listed in Table 3. Such nanocarbon-species dependence and the molar-ratio effect of the catalysts and carbon as the support on the catalyst on the catalyst performances were observed for a previously published paper using Fe<sup>III</sup>-doped nickel sulfides/nanocarbon hybrid catalyst<sup>23</sup>. In addition to the mesoporous carbon structures that have high porous area, the electronic interaction of the catalytic sites and the nanocarbon as a support would be important for such high performances.

**Table 3.** Oxygen electrode potentials of the synthesized polymer catalysts.

Polymer Catalysts	Oxygen Reduction Reaction $(E_{1/2})$ (ORR) V vs. RHE	Oxygen Evolution Reaction (OER) at 10 mA cm <sup>-2</sup>	Oxygen Electrode potential V vs. RHE
(Ni-TUF)n/CN-w/w:1/5	0.82	1.57	0.75
(NiTUF)n/MWCNT(Nikkiso)- w/w:1/5	0.65	1.52	0.87
(Ni-TUF)n/Vulcan-w/w:1/5	0.72	1.51	0.79
(Ni-TUF)n/CN-w/w:1/10	0.79	none	none
(Ni-TUF)n/CN-w/w:3/5	0.78	1.74	0.96

To further evaluate the ORR performances, the accelerated durability tests were carried out by performing repeated potentio-dynamic cycling for 20,000 cycles in the potential range of 0.2 to 1.0 vs. RHE at the scan rate of 50 mVs<sup>-1</sup> in an O<sub>2</sub>-saturated atmosphere. The result is shown in Fig. 8d, in which even after 10,000, then 20,000 cycles, the  $E_{1/2}$  of (Ni-TUF)n/CN-w/w:1/5 shifted by a negative potential of only ~50 mV, while for 10000~20000 cycles, the ORR

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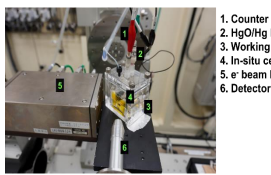
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activity still remained, indicating that the catalyst had a robust durability. This durability would be due to the presence of the S-atom in the thiourea formaldehyde polymer framework that forms  $d\pi$ – $d\pi$ bonding, resulting in the transfer of the 3d-electrons from the Ni atom to the vacant 3d orbital of the S-atom.<sup>27</sup> Such durability on (Ni-TUF)n/CN-w/w:1/5 was higher than those of (Ni-TUF/CN)n-w/w:1/10 and (Ni-TUF/CN)n-w/w:3/5 (see the ESI, Fig. S5a&b), indicating that the complex nanostructures in the catalysts are important factors for their durability. We plotted the Tafel slops of the (Ni-TUF/CN)nw/w:3/5, and IrO<sub>2</sub> for both OER and ORR. For the ORR region, the (Ni-TUF/CN)n-w/w:3/5 and IrO<sub>2</sub> exhibited the Tafel slopes of 72 mV/dec and 98 mV/dec, respectively (see the ESI, Fig. S5c). For the OER region, the (Ni-TUF/CN)n-w/w:3/5 and IrO<sub>2</sub> exhibited the Tafel slopes of 78 mV/dec and 57 mV/dec, respectively (see the ESI, Fig. S5d). Such results indicated the superior bifunctional performances of the (Ni-TUF/CN)n-w/w:3/5.

### 3.5. In situ Electrochemical EXAFS Measurements During **Electrochemical Reactions.**

The experimental in-situ setup using Pt as a counter electrode, Hg/HgO reference electrode, working electrode and electrochemical cell specially constructed for the XAFS studies are displayed in Fig. 9. We analyzed the EXAFS data of (Ni-TUF)n and (Ni-TUF)n-CN-w/w:1/5 during the oxygen reduction reaction region (0.81 ~ 0.91 V vs. RHE), non-faradaic region (0.1V vs. RHE) and oxygen evolution reaction region (1.31 ~ 1.41V vs. RHE). The first shell fitted radial distribution functions of (Ni-TUF)n:w/w-1/5 at the voltages of 0.81, 0.91, 1.01, 1.31 and 1.41V are shown in Fig. 10a, b, c, d and e, respectively. The input parameters ( $E_0$ ,  $S_0^2$ ,  $N_i$ ,  $R_i$ , and  $\sigma_i$ ,) are listed in Table S1 in the ESI. The bond lengths of the Ni-O and Ni-S for both (Ni-TUF)n and (Ni-TUF)n-w/w:1/5 were are in Table 4. Compared to the ex situ bond lengths of (Ni-TUF)n/CN:w/w-1/5, the bond lengths of the Ni-O and Ni-S at 1.01 V vs. RHE are deformed by +0.08 and -0.02 Å, respectively, suggesting that the H<sub>2</sub>O molecules occupy the axial orbital of the nickel atom and there is a cooperative intermediate step between the adsorbed superoxide (OOH-) species on the porous



1. Counter electrode 2. HgO/Hg Electrode 3. Working Electrode 4. In-situ cell 5. e<sup>-</sup> beam line

Fig. 9. Simultaneous electrochemical setup using (Ni-TUF)n/CN:w/w-1/5 electrode at the Kyushu University beamline, BL06, in the Saga Light Source.

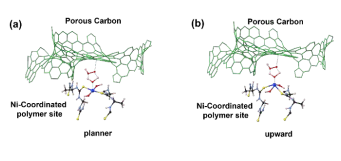
carbon (CN) and H<sub>2</sub>O on the nickel atom (Fig. 11a). In the QRR region (0.71 ~ 0.91V), compared to the Ni-O bond 10f (\$\text{Ni-Trut}\)06/€N1 w/w:1/5,

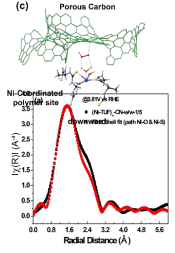
Table 4. Interatomic distances calculated using the EXAFS results taken from Fig.S8a-e.

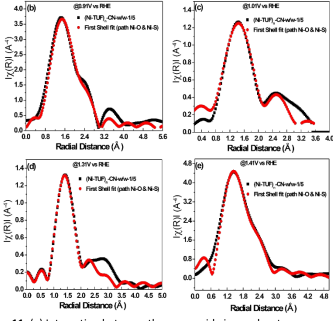
Applied voltages⁴ (V vs. RHE) ↔	Interatomic distances 4 of (Ni-TUF)n 4		Interatomic distances of (Ni-TUF)n-CN-w/w-1/5	
	Ni-O (Å)≪	Ni-S (Å)€	Ni-O (Å)℃	Ni-S (Å)≪
0.81	2.17	2.38	2.20	2.41
0.91	2.24	2.31	2.21	2.41
1.01	2.20€	2.37	2.25	2.36€
Ex situ °	2.18	2.35	2.17	2.38
1.31	2.18	2.40	2.23€	2.32
1.41	2.18	2.40	2.20	2.32

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**Fig. 10.** (a-e) *In situ* electrochemical EXAFS of (Ni-TUF)n/CN:w/w-1/5 electrode at the voltages of 0.81V(a), 0.91V(b), 1.01V(c), 1.31V (d) and 1.41 V (e) vs. RHE, based on the parameters described in the ESI, Table S1. All the black colored curves in the corresponding voltages are the raw XAFS data and all the red colored curves are their first shell fitting curves.



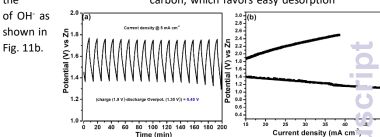




**Fig. 11.** (a) Interaction between the superoxide ion and water involved in the oxygen reduction in the voltage range of 0.81 **~0.91** 

vs. RHE. (b) Cooperative intermediate at the voltage of 1.01 Me sonline RHE. (c) Cooperative mechanism involving of 1.01 Me 1.

the Ni-S bonds elongated to 2.41 Å, which demonstrated that the Ni center is slightly upward from the planar position and the  $H_2O$  on the axial position of the Ni is close to the adsorbed superoxide species on the carbon, which favors easy desorption



Meanwhile, in the OER region, the bond lengths of the Ni-S at the 1.31 and 1.41 V vs. RHE were 2.32 Å, which suggested that the nickel center shifted slightly downward from the planar position during the applied voltage, and the H<sub>2</sub>O on the axial position of the Ni center is not close as in the ORR voltage region, which favors the desorption of the H<sub>2</sub>O and O<sub>2</sub> molecules (Fig. 11c). Furthermore, we analyzed the EXAFS results of the (Ni-TUF)n in the same voltage regions (see the ESI, Table S2 and Fig.s S6). The bond lengths of the (Ni-TUF)n are shown in Table 4. Unlike the carbon supported polymer (Ni-TUF)n-CN-w/w:1/5, (Ni-TUF)n without the carbon support exhibited a similar bond length for both the *ex situ* and 1.01V vs. RHE (nonfaradaic voltage) (see Table 4). This showed that absence of a cooperative mechanism of the H<sub>2</sub>O and OOH- ion. Hence, although

(Ni-TUF)n showed bond length changes at the OER and ORR voltages (Table 4), there is no catalytic performance which is different from the (Ni-TUF)n-CN-w/w.

3.6. Zn-air battery performance We then evaluated the Zn-air battery performance using the catalyst, (Ni-TUF)n/CN-w/w:1/5, and the results are shown in Fig. 12. The cathode for the battery assembly was prepared by coating the (Ni-TUF)n/CN-w/w:1/5 on the GDL (Sigracet DGL2288). The primary battery testing of the battery in the initial cycle was carried out by discharging and charge separately as shown (Fig. 12a). As the current density increases from 15 mA cm<sup>-2</sup>, the voltage for charge and discharge potentials are 1.80 and 1.35 V. Further increase in the current density resulted in increase in the charge potentials by ~2.4 V and decrease in discharge potentials by ~1.2 V (Fig. 12b). The open circuit potential of Ni-TUF)n/CN-w/w:1/5 exhibited 1.35 V before evaluating for the charge and discharge tests. Such high charge and discharge activities would be derive. d from the stable Ni-S active sites in the catalyst, Ni-TUF)n/CN-w/w:1/5. However, as the current density increased over 30 mA cm-2, the Ni-S and Ni-O bonding might increase drastically unlike the Ni-O and Ni-S bond length of Ni-TUF)n/CN-w/w:1/5 as shown in Table 4. This is confirmed from the Ni-O and Ni-S bonding trends in the in-situ EXAFS studies (Fig. 9 and Table 4), in which (Ni-TUF)n in Table 4 showed increased bond length for Ni-TUF)n/CN-w/w:1/5 from the potential 1.01 V to 1.41V. The rechargeable battery performance is shown in the Fig. 12b, in which the current density versus voltage

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performances of (Ni-TUF)n/CN-w/w:1/5, in which the battery shows the very low overpotential of 0.45 V, indicating an outstanding Zn-air battery performance when comparing the reported state-of-the-art Zn-air batteries, 13,18 in which in many papers, the overpotentials were 0.62~0.75 V. One of the important factors for the observed very high Zn-air battery using (Ni-TUF)n/CN-w/w:1/5 as the cathode material would be due to the cooperative H<sub>2</sub>O and superoxide ion mechanism of the electrochemical oxygen reduction and evolution reactions. Moreover, this polymer catalyst exhibited 0.45 V chargedischarge potential gap compared to the recently reported nickel iron sulfides (0.66V)<sup>26</sup> where the nickel sulfide is one of the active sites. This decrease in the charge-discharge potentials of Ni-TUF)n/CN-w/w:1/5 due to the polymerized Ni-S sites in this catalyst compared to its crystalline Ni-S analogue site<sup>26</sup> would be due to stabilization of the Ni-S and Ni-O sites in the (Ni-TUF)n polymer.

Fig. 12. (a) Charge-discharge profile of (Ni-TUF)n/CN-w/w:1/5 and (b) Current vs. voltage curve of the (Ni-TUF)n/CN-w/w:1/5.

### 4. Conclusions

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A novel poly(nickel-thiourea-formaldehyde), (Ni-TUF)n, was designed and synthesized in one pot. For comparison, we also synthesized poly(thiourea-formaldehyde), (TUF)n. Their structures were characterized in detail using elemental analysis, solid state NMR, FT-IR and XPS, and determined their chemical structures (see Fig. 1), in which the Ni forms a complex with the sulfur moieties in the polymer. Theoretical calculations and first shell fit of EXAFS showed that nickel is in a four coordination with two Ni-S and two Ni-O. We revealed that there is hydrogen bonding between the coordinated oxygen and nitrogen in the polymer chain. We then prepared seven different catalysts by combining (Ni-TUF)n and the carbon supports including conventional carbon black (Vulkan), unique porous carbon black (CN), and two different MWCNTs (Table 1). The (Ni-TUF)n supported on CN showed the highest ORR-OER bifunctional performances, which would be due to the formation of suitable nano-complex structures for the electrochemical reactions. In addition, the weight ratios of (Ni-TUF)n and CN in the catalysts were important for the higher performances. The Zn-air battery performances using the (Ni-TUF)n/CN-w/w:1/5 as the cathode material exhibited a superior rechargeable charge-discharge performance with the very low overpotential of 0.45 V, and the battery showed a high durability. We have carried out in situ EXAFS during the electrochemical reactions and found that there exist a bond length changes in the Ni-S and Ni-O, which confirms the existence of cooperative mechanism between (Ni-TUF)n and CN carbon. No such behavior was observed in (Ni-TUF)n.

This study provides a guide for the design and one-pot synthesis of a less-expensive nonprecious metal (Ni in this study) bifunctional polymer catalyst and rechargeable Zn-air battery catalyst a with very high performance, which is highly important in a future eco-friendly energy society. We also emphasize that the presented polymercatalyst design is simple and can be applied to many catalyst fields, i.e., replacement of Ni with other metals (or metal alloy) as electrode catalysts and molecular catalysts with different (or similar) objectives in the eco-friendly area is easy.

### Author Contributions

The manuscript was written through contributions of all authors. All authors have given approval to the final version of the manuscript.

### Conflicts of interest

There are no conflicts to declare.

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