Metal-Doped Aerogels Based on Resorcinol Derivatives

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Declaration:
Hereby I declare that this doctoral thesis, my original investigation and achievement, submitted for the doctoral degree at Tallinn University of Technology has not been submitted for any academic degree.

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Resortsinooli derivaatidel põhinevad metalli sisaldavad aerogeelid

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LIST OF PUBLICATIONS

This thesis is based on the following original publications:


OTHER PUBLICATIONS IN RELATED FIELD

AUTHOR’S CONTRIBUTION TO THE PUBLICATIONS

Publication I
The author designed the experiments and prepared all the materials. She interpreted the results of experiments and wrote the manuscript.

Publication II
The author designed and prepared all the materials. She interpreted the results of the materials characterisation experiments and wrote the major part of the publication. The author participated in the designing of the experiments for testing the materials as catalysts and also took part in the interpretation of the results.

Publication III
The author prepared all the materials, interpreted most of the results of the materials characterisation experiments and wrote that part of the publication. The author also participated in discussing the results of the electrocatalytical tests.
INTRODUCTION

Carbon materials due to the different allotropes of carbon have very versatile properties and can be used for a variety of applications. Some examples of these diverse materials include activated carbon, carbon black, graphite, carbon foam, carbon nanotubes, carbon fiber, etc. These materials can be used as adsorbents, electrode materials, insulators, catalyst supports, etc.

The demand for carbon materials to fit specific applications calls for the need to fine-tune the properties of these materials. A promising class of such materials is carbon aerogels (CAs), which are known for the ability to adjust their porous structure. Carbon aerogels have high surface area, open porous structure with pores ranging from micro- to macropores and electrical conductivity. Doping these materials with metals enables one to further modify their structure, improve electrical conductivity or add the desired catalytic activity. The main applications for metal-doped carbon aerogels are as electrode materials and catalysts.

Different methods can be employed to introduce metals into the carbon aerogel structure. The ion-exchange method used in this work enables to evenly distribute metal species to the surface of the porous material. This method involves the use of an ion-exchange moiety containing monomer for aerogel preparation. Each repeat unit of the material will therefore be able to chemically bind metal species.

Different phenolic compounds together with formaldehyde are mostly used for carbon aerogel preparation. In this work 5-methylresorcinol (MR) and 2,6-dihydroxy-4-methylbenzoic acid (dHMBA), which has the ion-exchange moiety, are chosen for aerogel preparation. These compounds are suitable because 5-methylresorcinol is a by-product of the local oil shale processing industry and 2,6-dihydroxy-4-methylbenzoic acid can be easily synthesized from 5-methylresorcinol [1,2].

The goal of this work is to develop a method for doping metals into aerogels prepared from 5-methylresorcinol and 2,6-dihydroxy-4-methylbenzoic acid, to study the properties of the obtained materials and to test them for desired applications. The prepared materials are doped with various metals such as Co, Cu, Ni, La, Ce, Nd, Pr. The structure and morphology of metal-doped carbon aerogels are characterised by methods such as nitrogen adsorption analysis, scanning electron microscopy (SEM), transmission electron microscopy (TEM), X-ray diffraction (XRD) analysis and different elemental analysis techniques. The relationships between the conditions of preparation and properties of the obtained materials are discussed.

The metal-doped aerogels are tested for use as heterogeneous catalysts. At first, lanthanide-doped organic and carbon aerogels are tested as catalysts for the Michael addition reaction. This is relevant because the preparation and application of lanthanide-doped aerogels as catalyst materials has been poorly studied.
Secondly, the metal-doped carbon aerogels prepared in this work, are tested as electrocatalysts for the oxygen reduction reaction (ORR). This reaction occurs on the cathode in proton-exchange membrane (PEM) fuel cells, which are devices that use hydrogen as fuel to produce electricity. The need for less expensive and more durable catalysts is one of the main obstacles for a wider commercial use of these devices. In recent years the development of non-precious metal catalysts (NPMCs) has attracted a lot of attention due to them being good substitutes for expensive Pt-based catalysts which are currently used in fuel cell cathodes [3,4,5,6]. NPMCs can be prepared from different carbon nanomaterials. In this work cobalt- and nitrogen-doped carbon aerogels are developed for use as ORR catalysts. The nitrogen-doping, is achieved by using melamine (Mel) as one of the precursors for aerogel preparation. The physicochemical properties of carbon aerogels are characterised and the relationships between their properties and catalytic activity are discussed.
AIMS OF THE STUDY

The main goal of this work is to show that metal-doped organic and carbon aerogels can be successfully prepared from oil shale processing by-product 5-methylresorcinol and its derivative 2,6-dihydroxy-4-methylbenzoic acid and that these aerogels have potential for use as heterogeneous catalysts. More specific aims of individual studies are the following:

• developing a method for preparation of metal-doped organic and carbon aerogels from 5-methylresorcinol and 2,6-dihydroxy-4-methylbenzoic acid by using the ion-exchange method for metal doping;

• preparing aerogels doped with various metals and determining the relationships between the conditions for preparation and properties of materials;

• showing the potential of lanthanide-doped aerogels for use as heterogeneous catalysts;

• developing metal-doped carbon aerogels for use as catalysts in the oxygen reduction reaction and also characterising the materials;

• finding the relationships between the properties and catalytic activity of carbon aerogels in oxygen reduction reaction.
ABBREVIATIONS

CA  carbon aerogel
MR  5-methylresorcinol
dHMBA  2,6-dihydroxy-4-methylbenzoic acid
SEM  scanning electron microscopy
TEM  transmission electron microscopy
XRD  X-ray diffraction
ORR  oxygen reduction reaction
PEM  proton-exchange membrane
NPMC  non-precious metal catalyst
Mel  melamine
OA  organic aerogel
R  resorcinol
FA  formaldehyde
R-FA  resorcinol - formaldehyde
SOLV  solvent
Cat  catalyst
Mel-FA  melamine - formaldehyde
R-Mel  resorcinol - melamine
DHBA  2,4-dihydroxybenzoic acid
HRTEM  high-resolution transmission electron microscopy
XPS  X-ray photoelectron spectroscopy
DHBAF  2,4-dihydroxybenzoic acid - formaldehyde
MR-FA  5-methylresorcinol - formaldehyde
Ln  lanthanide
PEMFC  proton-exchange membrane fuel cell
M/N/C  metal-containing nitrogen-doped carbon
AAS  atom absorption spectroscopy
SBET  Brunauer-Emmett-Teller surface area
Vmic  microporous volume
TGA  thermogravimetric analysis
AM  aromatic monomer
NaOAc  sodium acetate
Co-NCA  cobalt- and nitrogen-doped carbon aerogel
Co-CA  cobalt-doped carbon aerogel
NCA  nitrogen containing aerogel
Smic  microporous surface area
VTOT  total pore volume
PSD  pore size distribution
BSE  backscattered electron
ICP OES  inductively-coupled plasma optical emission spectrometry
CNT  carbon nanotube
RDE  rotating disk electrode
1. LITERATURE OVERVIEW

Aerogels are solid materials which are prepared by replacing the liquid component of gels with gas. They were first prepared by Kistler [7], who in 1931 published an article in which he dried silica aerogels by using a new supercritical drying method. The method involved increasing pressure and temperature over the critical parameters of the solvent, so that during drying the crossing of the boundary from liquid to gas, which causes capillary stress to emerge, could be avoided.

Aerogels are known for their low density, open porous structure and high surface area. The properties of aerogels depend on the nature of the precursor gel, which can be prepared from different compounds. Besides silica aerogels, different metal oxides and organic compounds can be used for aerogel preparation. Depending on precursors and the preparation method used, aerogels can have various favourable properties such as low thermal and sound conductivity, good electrical conductivity, high hydrophobicity and low refractive index. Some of the applications for aerogels include thermal insulators, adsorbents, electrode materials, drug delivery systems, platforms for chemical sensors, etc [8].

Carbon aerogels (CAs) are mainly composed of carbon as the name suggests. These aerogels are usually prepared by pyrolysing organic aerogels (OAs) which themselves are obtained by drying organic gels. The properties of CAs are largely dependent on the structure and morphology of the parent organic gel. Therefore the study of the properties of carbon aerogels starts with elucidating the relationship between the gel preparation conditions and properties of the gels obtained. The drying method and pyrolysis conditions used also affect the properties of resulting CAs.

The most studied procedure for preparation of organic aerogels was introduced in 1989 by Pekala [9], who used resorcinol (R) and formaldehyde (FA) for gel preparation, followed by the supercritical drying of the gel. Subsequently, the preparation of carbon aerogels by pyrolysing resorcinol-formaldehyde (R-FA) organic aerogels was described [10]. Over the years several other precursors have been used for carbon aerogel preparation, like melamine, phenol, cresol, polymeric isocyanate, polyvinyl chloride, phloroglucinol, ovalbumin/glucose, cellulose, tannins aso., but none of them have been nearly as popular and thoroughly studied as R-FA aerogels [8,11]. The effect of gel preparation conditions, different drying methods and pyrolysis conditions on the properties of the obtained aerogels will be reviewed based on the literature.

1.1. Preparation of organic and carbon aerogels

Since resorcinol and formaldehyde are the most studied precursors for organic aerogel preparation and 5-methylresorcinol and 2,6-dihydroxy-4-methylbenzoic acid are derivatives of resorcinol and form gels in a similar way
[1], the preparation and properties of carbon aerogels will be explained on the basis of resorcinol-formaldehyde aerogels which have been thoroughly studied in the literature. The structure and properties of carbon aerogels are affected by all the stages of the material preparation, so the overview of carbon aerogel preparation starts with explaining the gel formation.

1.1.1. Gel preparation

The preparation of carbon aerogels starts with obtaining an organic gel through sol-gel polymerisation. For R-FA aerogels the monomers resorcinol (R) and formaldehyde (FA) are mixed together into the solvent (SOLV) along with a catalyst (Cat) (acidic or basic). The formation of the gel is shown in Figure 1. Resorcinol is able to bind formaldehyde in positions 2-, 4- and 6- in its aromatic ring. When sodium carbonate is used as catalyst, the polymerisation reaction between R and FA consists of two main reactions: the addition reaction where hydroxymethyl derivatives are formed and the condensation reaction of the hydroxymethyl derivatives to form methylene (-CH2-) and methylene ether (-CH2OCH2-) bridged compounds (Fig. 1) [12]. The polymerisation reaction results in the formation of clusters, which aggregate into colloidal particles and then form a three-dimensional interconnected structure (gel). The pores are filled with the solvent [12].

Figure 1. Diagram of the formation of the R-FA gel through sol-gel polymerisation (the figure is adapted and modified from [1]).
The molar ratios of substances can be varied during preparation, which greatly affects the properties of the obtained gel. The molar ratios of R to FA, SOLV to R and R to Cat can all be changed. According to the literature data, the molar ratio of R to FA is usually kept at 0.5 [12] and tuning of the aerogel structure and properties is carried out by changing SOLV to R and R to Cat ratios.

Increasing the amount of solvent (increasing the SOLV to R molar ratio) results in the dilution of the mixture and decreases the density of the obtained gel. Diluting the sol too much will hinder the gel formation though [13]. Decreasing the SOLV to R molar ratio leads to a more dense gel. The higher density of the gel can lead to a decrease in porosity and surface area of the subsequently obtained aerogel [12].

The amount of catalyst (R to Cat molar ratio) has the biggest effect on the properties of R-FA aerogels. Aerogels prepared using high catalyst amounts (the molar ratio of R/Cat is about 50) result in small particles (about 3-5 nm in diameter) that are joined together by large necks, which are referred to as “polymeric” R-FA aerogels. This produces higher density gels that exhibit substantial shrinkage even during supercritical drying and have high surface areas [8,10,12,14]. Lower catalyst amounts (high R to Cat molar ratios (R/Cat = 200)) result in larger particles (ca 11-14 nm in diameter) that are connected by narrow necks and are referred to as “colloidal“ R-FA aerogels [8,10,12,14]. These gels exhibit little shrinkage during supercritical drying, have lower surface areas and weaker mechanical properties [8].

1.1.2. Drying of gels

To obtain an aerogel, the solvent in the pores of the gel needs to be removed in a way that does not involve crossing the boundary between liquid and gaseous phase, which results in the formation of capillary stresses inside the pores that lead to the destruction of the gel structure. Drying in the atmospheric conditions (the liquid is allowed to evaporate and that causes considerable shrinkage) would lead to the formation of a xerogel, which has higher density and lower surface area. Different drying methods can be used to avoid the collapse of the structure during drying. Among these methods the most popular is supercritical drying.

Supercritical drying involves bringing the solvent in the pores of the gel to the supercritical state. The phase diagram in Figure 2 shows that the idea of the drying is to remove the solvent by at first bringing it from the liquid state (A) to the supercritical state (B) by raising the pressure and temperature above the critical parameters (P_c and T_c). After that the pressure is lowered and the fluid is brought to the gaseous state (C). This avoids crossing the liquid-gas boundary and formation of capillary forces.

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For supercritical drying the use of CO₂ is preferred over the other solvents due to the relatively low critical parameters and safety [16]. The solvent inside the pores of the gel needs to be miscible with CO₂ for it to be extracted out of the pores. For example, if the gel has been prepared with water as the solvent, which is not miscible with CO₂, then it needs to be replaced before drying with another solvent like acetone. This process is diffusion limited and thus time consuming. Later the acetone in the pores of the gel is replaced by liquid CO₂ and then supercritical drying follows.

1.1.3. Pyrolysis and carbon aerogels

Carbon aerogels are prepared by pyrolysing organic aerogels. Pyrolysis (also called carbonization) is a process during which the material is heated in inert (N₂, Ar or He are most commonly used) atmosphere. This transforms the organic aerogel into a material which is mostly composed of carbon. The properties and structure of the carbon aerogel are affected by the pyrolysis temperature (temperatures from 600 - 2100 °C can be used) [12]. The initial structure of the organic aerogel is largely preserved during pyrolysis, but shrinkage and mass loss (about 20-50 wt%) occur [8,12].

Al-Muhtaseb and Ritter [12] have described the effect of pyrolysis temperature on the properties of the resulting CAs in a review article. Carbon aerogels have a greater density than R-FA organic aerogels. The density of CAs is decreased with increasing pyrolysis temperature, eventually reaching a limiting value at 800 °C [17].

During pyrolysis the burnout of the organic material leads to the formation of new small pores in the solid network of the aerogel. Due to these new micro- and mesopores the surface area of the aerogel increases. However, at higher pyrolysis temperatures (above 600 °C) the surface areas of R-FA aerogels tend to reduce. [10,12,17]. This is because of the shrinkage of the meso- and macropores that were present in the organic aerogel.

Pyrolysis changes the pore sizes of R-FA aerogels, but does not change the trend of pore size distribution [17]. It is possible to control the amount of micropores and mesopores independently, which is one of the advantages of
CAs as a porous carbon material [18]. Micropores can be related to the intra-particle structure, which form during pyrolysis inside the solid aerogel network due to the burn-off of the organic material, whereas mesopores and macropores are related to the inter-particle structure of the aerogel that is formed during gelation [18]. When R-FA aerogels were pyrolysed at 900 °C, micropores were formed due to the formation of new pores in the aerogel structure, but the mesopore volume of the carbon aerogel was smaller than that of the original R-FA OA (probably due to shrinkage of the initial pores in OA) [19].

For carbon aerogels [20] and xerogels [21] to be electrically conductive a pyrolysis temperature above 750 °C is recommended [12]. The formation of graphitic structures in R-FA carbon aerogels can be observed at around 1050 °C [12,22].

1.2. Aerogels prepared from 5-methylresorcinol, 2,6-dihydroxy-4-methylbenzoic acid and melamine

Since instead of resorcinol its derivatives MR and dHMBA and also melamine are used in this work for aerogel preparation, an overview of the previously published results significant for this thesis is given.

1.2.1. 5-methylresorcinol for aerogel preparation

The preparation of organic and carbon aerogels from 5-methylresorcinol and their properties have previously been studied in our laboratory. MR is a phenolic compound which is obtained from the local oil shale industry where it is available as a by-product of oil shale processing. MR is produced by extraction from oil shale retort liquor [23].

MR-FA aerogels were synthesized using a doubly catalysed polycondensation (after gelation the gel was immersed in an acetic acid solution), with Na2CO3 as catalyst and supercritically dried with CO2. Pyrolysis was carried out at temperatures 700 and 1000 °C [24]. 5-methylresorcinol is a trifunctional monomer which is able to react with FA in the 2nd, 4th and 6th positions of the aromatic ring. The formation of the gel was proven to be faster for the MR-FA gel than for R-FA gels and at lower temperature. This was due to an additional directing group (-CH3) in the 5th position of the aromatic ring. The gel was formed at 25 °C within 100 minutes and was ready for handling after 24 hours [25].

The obtained CAs had properties comparable to those of R-FA aerogels. CAs with a density of 0.13 g/cm3 and specific surface area of over 500 m2/g can be obtained [24]. Aerogels were prepared at different MR to catalyst molar ratios, MR to water molar ratios and pyrolysis temperatures. The highest surface areas for CAs were obtained at a pyrolysis temperature of 900 °C and at an MR to Cat molar ratio of 90 [24].

The morphology of MR-FA carbon aerogels was evaluated by SEM analysis and proved to be similar to that of R-FA CAs as demonstrated in Figure 3. Both
are composed of interconnected particles. The sizes of particles depend on the preparation conditions [1,12].

Overall, MR behaved in a similar way to R during aerogel preparation with similar properties being achieved for aerogels.

![SEM pictures of carbon aerogels](image_url)

**Figure 3.** SEM pictures of carbon aerogels: a) a “conventional” R-FA carbon aerogel. Reproduced from [26], with permission from Elsevier; b) a typical MR-FA carbon aerogel. Reproduced from [24], with permission from Elsevier.

### 1.2.2. 2,6-dihydroxy-4-methylbenzoic acid for aerogel preparation

dHMBA is obtained by synthesizing it from MR [2]. dHMBA is a bifunctional monomer which is capable of forming bonds with FA in two positions of the aromatic ring. It can also act as an acidic catalyst to promote polymerisation [27].

dHMBA and FA without adding a trifunctional monomer like MR will not form a gel. Figure 4 shows that varying the amount of MR and dHMBA changes the aerogels morphology. At higher MR amounts (85/15), the structure of OAs is composed of uniform spherical particles (diameters 24-25 nm), which is similar to that of MR-FA aerogels [1,27]. Increasing the percentage of dHMBA leads to the formation of strings of particles and formation of large macropores (pore diameter > 50 nm) as is the case with aerogels 50/50 and 25/75 [1,27].

The specific surface area of OAs was shown to increase when dHMBA was added. The highest specific surface area and the largest total pore volume, 600 m²/g and 800 mm³/g, respectively, were achieved at a MR/dHMBA ratio of 75/25. Compared to MR-FA OAs, these values are higher [1,27].
1.2.3. Melamine for aerogel preparation

Its cheap price and high nitrogen content makes melamine a suitable substance for aerogel preparation. Melamine-formaldehyde (Mel-FA) organic aerogels with low densities, high surface areas, continuous porosity and optical clarity were first prepared by Pekala [10,28]. The Mel-FA OAs can also be pyrolyzed in an inert atmosphere to obtain carbon aerogels.

Figure 5 shows a schematic diagram of the formation of the cross-linked polymer network by the reaction of melamine and formaldehyde. Melamine is a hexafunctional monomer which is capable of reacting at each of the amine hydrogens [28]. During polymerisation diamino methylene (-NHCH2NH-) and diamino methylene ether (-NHCH2OCH2NH-) bridges form [28]. The structure and properties of the aerogel are dependent on the polymerisation conditions. Factors such as pH, reactant ratio, and temperature influence the cross-linking chemistry and growth processes taking place prior to gelation [28].
Melamine-formaldehyde aerogels have been prepared by adding other monomers like phenol [29], cresol [30] and resorcinol [31,32,33,34,35] to the sol. The aim is usually to obtain nitrogen containing materials whose microstructure can be fine-tuned.

Carbon aerogels with various molar ratios of resorcinol to melamine have been prepared [33]. Changing the molar ratio of R to Mel allowed control of the nitrogen content in the materials. However, adding melamine reduced the surface areas and microporosity of materials [33]. The addition of melamine also lead to significant changes in the material morphology as seen from SEM pictures (Figure 6). Figure 6a shows the morphology of a typical R-FA CA [12,33], with a particle size ranging from 20-30 nm. A material to which melamine has also been added (Fig. 6b) shows the presence of “interconnected sphere-like nanoparticles“. It was also noted that particle size tended to increase with increasing melamine content [33]. In the case of 1:0.8 R-Mel CA, the decomposition of the spherical structure was observed (Fig. 6c) [33].

1.3. Metal-doped organic and carbon aerogels

The preparation of metal-doped aerogels has earlier been reviewed once by Moreno-Castilla and Maldonado-Hódar [18]. The purpose of adding metals to the aerogels is to modify their structure, conductivity and catalytic activity. These aerogels can be used as catalysts and electrode materials.

The methods used for doping can be divided into three categories [18]. First method involves the addition of the metal precursor to the initial mixture before gelation. Secondly, the ion-exchange method can be used. In this approach the
gel, which has ion-exchange moieties, is immersed in the solution of metal ions. Lastly, the metal precursor can be deposited on the organic or carbon aerogels by using various deposition methods. Each doping method affects the structure and properties of the resulting materials differently.

An overview of the methods used for metal-doping will be given in the next sub-chapters, with emphasis on the ion-exchange method. The properties of OAs and CAs prepared by using this technique will be discussed in detail. A short description of the other doping methods will be given.

1.3.1. Addition of the metal-precursor to aerogels before gelation

When the metal precursor is dissolved in the sol before gelation, then after gelation the dopant will be trapped within the gel structure. It has been suggested that the metal ions can be chelated by the functional groups of the polymer matrix [8,18,36]. Several different metal ions have been doped into aerogels this way, including Cr-, Mo-, W-, Fe-, Co-, Ni-, Pd-, Pt-, Cu- and Ag-ions [37,38,39,40].

The metal precursors (salts) added to aerogels can also act as catalysts for the polymerisation reaction. An early work by Bekyarova and Kaneko [36] showed that the concentration of the metal dopant and the pH of the reaction mixture can have a profound effect on the structure, porosity and physical properties of aerogels. It was suggested that both the nature of the metal dopant and the change in pH due to adding metal salts affect the sol-gel chemistry. It was found that the Ce- and Zr-doped CAs had developed a larger micropore volume and surface area due to the metals added.

The preparation of metal-doped aerogels by adding metal precursor before gelation has been extensively studied by Moreno-Castilla, Maldonado-Hódar and co-workers. Pt-, Pd- and Ag-doped carbon aerogels were prepared by adding the metal salts to the sol instead of a “typical“ Na2CO3 catalyst and it was found that this changed the porous structure of the materials [39]. Furthermore, while studying Fe-, Co-, Ni- and Cu-doped organic aerogels, it was found that depending on the metal salt used, the surface areas, pore volumes and sizes either increased, did not change or decreased compared to the undoped aerogel [40]. This means that each metal affects the polymerisation reaction in a different way.

The fact that metal salts used for doping also influence the polymerisation reaction and therefore the structure of the materials has been confirmed in many subsequent papers [41,42,43]. Adding the metal salt before gelation changes the porous structure of the aerogel and can make its porosity more difficult to control.

Generally a good dispersion of metal particles has been observed in carbon aerogels prepared by this doping method. It has been suggested that it is due to dopant elements being anchored to the carbon structure and thus preventing their migration and growth [18,36]. On the other hand, it has also been claimed that some metal particles can be encapsulated by the carbon matrix and this can
reduce the accessibility of gases and reactants to the metal when they are required for use as catalysts [18,43].

1.3.2. The ion-exchange method for doping with metals

This approach has been pioneered by Baumann and co-workers [44]. The method involves using 2,4-dihydroxybenzoic acid (DHBA) instead of resorcinol. The strategy is to use a resorcinol derivative that contains an ion-exchange moiety which for DHBA is the carboxylic group. This means that each repeat unit of the organic gel is able to bind metal ions and thus ensures a uniform distribution and better control over the content of metal ions. Since the metal precursor is introduced after gelation, it will not interfere with the sol-gel process and the resulting structure of the aerogel.

In a typical experiment, the aerogels would be prepared adding DHBA to the solvent (mostly water) and then K$_2$CO$_3$ is added to form a potassium salt of DHBA. Formaldehyde is then added to crosslink the DHBA molecules together and finally the catalyst is added, which is usually K$_2$CO$_3$. The sol is allowed to gel and cure. After that the K$^+$ doped hydrogels are immersed in an aqueous solution of a desired metal ion. This process is carried out for three days and the metal ion solution is changed every 24 h. Finally the solvent change and drying are carried out.

The copper content in the organic aerogel, which was prepared in the above mentioned procedure, was measured to be 4.4 wt% and in the carbon aerogel 9.2 wt%. For comparison an R-FA aerogel was immersed in the copper solution to determine the role that the ion-exchange moiety plays in the metal ion incorporation [44]. This approach proved ineffective for metal-doping, because most of the copper ions had been washed out during the acetone exchange.

The morphology of Cu-doped DHBA aerogels can be seen in TEM pictures (Fig. 7). The organic aerogel (Fig. 7a) consists of a network of spherical particles with diameters ranging from 15 to 30 nm. No copper particles can be seen, which indicates that the copper ions are distributed throughout the material and have not aggregated [44].

Figure 7. High-resolution transmission electron microscopy (HRTEM) pictures of Cu-doped aerogels: a) organic aerogel; b) carbon aerogel with 1% of Cu after carbonization at 1050 ºC. Figure is reprinted with permission from [44]. Copyright 2005 American Chemical Society.
After carbonization at 1050 °C the general morphology of the material is retained (Fig 7b), although the particles forming the aerogel network appear to have slightly shrunk (particle diameter 10-15 nm). In Figure 7b, copper nanoparticles with a diameter from 10 to 50 nm can be seen. This happens due to the reduction of Cu$^{2+}$ ions during pyrolysis. The size of copper nanoparticles depends on the copper content in aerogels, with larger particles forming with higher copper loading. As seen from Fig. 7b the distribution of copper nanoparticles is uniform.

The Cu-doped OAs and CAs exhibited a high surface area (ca 700 m$^2$/g) and pore diameters from 2 to 20 nm. The metal-doped materials retained the overall porous structure of the metal-free carbon aerogels [44].

The evolution of the morphology of metal-doped carbon aerogels during carbonization was more thoroughly studied for Co- and Ni-doped aerogels [45]. OAs were pyrolysed at temperatures from 300 to 1050 °C and characterised by X-ray diffraction (XRD) and transmission electron microscopy (TEM). The metal contents in the samples were from 9 to 10 wt%. The formation and size of metal nanoparticles were dependent on the carbonization temperature. At 450 °C small Co nanoparticles (ca 2 nm) started to form. With increasing carbonization temperature the size of Co nanoparticles also increased (up to 60 nm at 1050 °C). Based on X-ray photoelectron spectroscopy (XPS) and XRD analysis it was concluded that the Co$^{2+}$ ions were not directly reduced to metal particles at lower pyrolysis temperatures. The XRD analysis results show that at pyrolysis temperatures below 400 °C the peaks corresponding to CoO are present and at 450 °C the peaks corresponding to Co start to emerge as well (Fig. 8a). At 1050 °C the peaks for CoO are not visible, indicating its full reduction (Fig. 8c) [45].

For Ni-doped CAs, small nanoparticles (ca 2-4 nm) started to form at 400 °C and at 1050 °C nanoparticles in the size range of 200-400 nm was observed from TEM pictures [45]. The formation of nickel oxide was not observed by XRD analysis (Fig. 8b,c).

![Figure 8. XRD patterns: a) Co-doped aerogels pyrolysed at temperatures 300-600 °C; b) Ni-doped aerogels pyrolysed at temperatures 300-600 °C; c) Co- and Ni-doped aerogels pyrolysed at 1050 °C. Figure is reprinted with permission from [45]. Copyright 2007 American Chemical Society.](image-url)
The formation of highly ordered carbon structures in Co- and Ni-doped aerogels pyrolysed at 1050 °C was observed [45]. Figure 9a,b shows that carbon structures had formed around Co nanoparticles. Graphitic nanoribbons with width ranging from a few nanometers to 20 nm were observed throughout the CA structure (Fig. 9b). At lower temperatures (600 °C) the formation of graphitic structures in metal-doped aerogels did not occur. Despite the formation of graphitic structures, the surface areas, pore volumes and average pore diameters of the metal free sample, Co-and Ni-doped CA remained similar at 1050 °C [45]. The tendency for some metal-doped aerogels to catalyse the formation of carbon structures at relatively low temperatures has been used before to grow carbon nanotubes [46].

Figure 9. a,b) HRTEM pictures of Co-doped CAs pyrolysed at 1050 °C showing metal nanoparticles and graphitic structures. Figure is reprinted with permission from [45]. Copyright 2007 American Chemical Society.

The differences in the porosity and surface area for metal-doped organic aerogels and their non-metal containing counterparts is quite small when the ion-exchange doping method is used. This was noted for Fe-, Ni- and Cu-doped aerogels [47]. The changes that take place when metal precursors were added before gelation were far more significant [40].

2,4-dihydroxybenzoic acid-formaldehyde (DHBAF) organic aerogels have been prepared by also adding resorcinol to the materials [48]. The results showed that for DHBA and R containing organic aerogels (DHBARF) the surface area, pore volume, pore sizes and ion exchange capacity changed in a systematic fashion when the DHBARF ratio was varied. For DHBARF aerogels prepared with an equal amount of both R and DHBA in the mixture (the molar ratio of R to DHBA was 1:1), the mesopore volume and pore diameters were lower than in R-FA aerogels, but higher than in DHBAF aerogels. Compared to both RF and DHBAF aerogels the surface areas of DHBARF aerogels were the highest. It was also found that in all cases (R-FA, DHBAF and DHBARF aerogels) higher solids content in organic aerogels decreased pore volumes and surface areas [48].

The metal content was experimentally measured for OAs containing 40 to 100 molar percent of DHBA (explanation: 40 mol% of DHBA means there is 60 mol% of R). For both Ni-and Cu-doped OAs the ion-exchange capacity
stayed at its maximum value when at least 50 mol% of DHBA was in the material [48].

1.3.3. Deposition of the metal precursor onto the organic or carbon aerogel

When the metal precursor is deposited onto the already prepared organic or carbon aerogel, then the metal will not affect the formation of the aerogel structure. Different methods for deposition can be used, among them are incipient wetness, adsorption, sublimation and supercritical deposition [18,43,49,50,51,52].

After deposition of the metal precursor, heating of the doped aerogel in an inert atmosphere is usually necessary to reduce the metal precursor to metal particles. Since temperatures high enough for carbonization are not needed when ready-made CAs are doped with metal precursors, then materials with smaller metal nanoparticles can be obtained when lower temperatures (around 300 °C) are used for reduction of metal precursors [51]. The growth and aggregation of metal nanoparticles mainly occur at higher temperatures. For instance, it has been observed that Pt particles of size of a few nanometers were obtained when reduced at 500 °C, but at 1000 °C, the sizes and size distribution of metal nanoparticles had grown to 2-10 nm [52].

In several publications it has been noted that deposition of metal species onto carbon aerogels reduces their porosity [41,51,53]. In an early work Miller and Dunn showed that after doping a CA with a surface area of 700 m²/g with ruthenium precursor, the surface area had decreased to 280 m²/g [51]. Furthermore, the comparison of pore size distributions showed that all the pores with diameters under 2.5 nm had disappeared. The authors suggested that this could be due to the blocking of smaller pores by metal nanoparticles [51].

1.4. Metal-doped organic and carbon aerogels for various applications

As organic and carbon aerogels are materials with highly versatile properties, they show great promise in very different areas of technology. The main application areas for metal-doped aerogels are as catalysts or in various electrical applications. A short description of these application areas will be presented in the next chapter. Also, since in this thesis the lanthanide-doped aerogels have been characterised and tested as supported catalysts and Co-doped aerogels have been studied for use as electrocatalysts for ORR, then an introduction to these areas will also be given.

1.4.1. Metal-doped aerogels as catalysts

The use of carbon aerogels as catalysts has been reviewed before [18]. Their tunable surface area, porosity and pore size distributions make them promising materials for use as catalyst supports. Doping the materials with suitable metals
gives them desired catalytic activity. A few of the reactions where metal-doped CAs have been tested as catalysts are the isomerization reaction of 1-butene [38], synthesis of methyl-tert-butyl ether from methyl alcohol and tert-butyl alcohol [54], toluene combustion reaction [43] and photooxidation or ozonation of micropollutants [55,56]. In all these cases the metal-doped aerogels showed good catalytic activity and were proposed as new promising catalyst materials.

However, the tendency for carbon aerogels to burn-off at elevated temperatures in oxidizing conditions can be a drawback. This has been studied for Pt-doped CAs pyrolysed at 500 and 1000 °C [43]. Under air flow the burn-off of the materials started at temperatures between 300 to 400 °C, with the onset being at a bit higher temperature for the CAs pyrolysed at 1000 °C.

Organic and carbon aerogels have been compared to silica aerogels for use as catalyst supports. The catalytic activity of Pd-doped organic and carbon aerogels prepared from DHBA were compared to the Pd-doped silica aerogel in the Mizoroki-Heck coupling reaction [57]. It was found that the Pd-doped organic and carbon aerogel were more active catalysts than the Pd-doped silica aerogel. The authors suggested that since the use of DHBA enables each repeat unit of the organic polymer to bind metal ions, the uniform distribution of the metal dopant maximizes the contact of the catalyst with the reactants and can explain the difference in catalytic activity between doped organic/carbon and silica aerogels [57].

Different metal-doping methods affect the structure and properties of the resulting catalyst materials differently and thus the materials catalytic activity can be different. This has been studied before by preparing Pt-doped materials in two different ways. The Pt-doped carbon material was prepared by either impregnating of OA in an aqueous solution of the Pt-precursor or by adding the precursor to the initial sol during preparation [43]. The catalytic activity of the materials towards oxidation of toluene was studied and the results showed that both materials proved to be active catalysts and there was only a slight difference in the starting temperature of the combustion. The authors suggested it could have been due to different Pt-particle sizes.

However, when the ion-exchange method is used for metal-doping, the need for a temperature high enough for carbonization, may be a drawback. The catalytic activity of Pt-doped CAs for the oxygen reduction reaction was studied and it was found that at high carbonization temperatures (1050 °C) the platinum nanoparticles became blocked and thus the Pt-doped CA lost its catalytic activity [58]. At lower pyrolysis temperatures some activity of the Pt-doped CA remained. For comparison, when the Pt-doped material was prepared by immersing the CA with the Pt precursor, a chemical reduction instead of high temperatures could be used, so the Pt nanoparticles had small uniform sizes (2-5 nm) and the Pt particles were accessible [58].

Previously in our laboratory the preparation of the Pd-doped 5-methylresorcinol-formaldehyde (MR-FA) carbon aerogel has been studied. The material was prepared by impregnating the MR-FA carbon aerogel with Pd(C4HF6O)2 in supercritical CO2 and using H2 for reducing the metal [25].
Pd-doped CA showed total conversion in the reduction of cyclooctene into cyclooctane and the material could be used several times.

1.4.2. Lanthanide-doped organic and carbon aerogels

Lanthanides are widely used in organic synthesis because lanthanide-mediated reactions display high chemo- and stereoselectivities [59,60]. Use in the Michael addition reaction is one example.

Lanthanides(III) supported on polymers or ion-exchange resins are developed and studied to take advantage of the catalytic activity of lanthanides on the one hand and the ability of the solid phase to simplify products isolation on the other. The potential use of lanthanide(III) catalysts supported on ion-exchange resins has been demonstrated in routine organic reactions [61]. The preparation of resin-supported catalysts is straightforward, and the resultant catalysts are stable and active under common conditions (acidic conditions and room temperature) in organic synthesis [61].

The support or entrapment of lanthanides or lanthanide species in aerogels or xerogels has been mostly performed with the goal of preparing luminescent materials [62]. The doping of aerogels with lanthanide species for use as supported catalysts has been poorly studied [63].

The europium-doped organic gels, and organic and carbon aerogels have previously been characterised and used as catalysts for the Michael addition reaction [63]. The gels were prepared from DHBA and the ion-exchange method was used for doping with Eu(OTf)_3. The TEM pictures showed that the gel, OA and CA all had nanoparticles in their structure. The presence of various mixed europium oxides, sulphides and fluorides in CAs were confirmed by XRD analysis, as shown in Figure 10 [63]. The gel and OA were amorphous. The materials were tested as catalysts for the Michael addition reaction and the most active were the organic gels with over 90% yield after five cycles. The carbon aerogels did not catalyse the reaction.

![Figure 10. XRD pattern of Eu-doped materials. Reproduced from [63].](image-url)
1.4.3. Metal-doped carbon aerogels for electrical applications

Carbon aerogels are appropriate materials for electrochemical storage of energy in supercapacitors. Adding metal species to the carbon structure can increase the capacitance values due to pseudocapacitance effects [8,64]. Carbon aerogels doped with the oxides of Ru, Co, Cu, Fe, Mn and Zn have been studied for use in supercapacitors [51,53,65].

For metal-doped carbon aerogels another promising application is as electrocatalysts in proton-exchange membrane fuel cells (PEMFC). The reaction that takes place on the cathode is the oxygen reduction reaction (ORR). Currently platinum on porous carbon materials is used in almost all commercially available PEM fuel cells. Due to the high cost of Pt, the development of non-precious metal catalysts for PEM fuel cells is one of the most important goals [3,4,5,6,66].

A promising class of catalysts is transition metal-containing nitrogen-doped carbon materials (M/N/C catalysts) that can be prepared by pyrolysis of the carbon support in the presence of the nitrogen precursor and a transition metal (usually Co or Fe) salt [3,4,5]. Since the early studies of ORR on this type of pyrolysed materials by Yeager’s group in 1989 [67] the electrocatalytic activity of M/N/C catalysts has been significantly improved and it currently approaches that of commercial Pt/C catalysts in acidic solution [5]. In some cases, their activity can even surpass that of Pt-based catalysts [68,69,70].

Various different carbon materials such as carbon nanotubes, carbon black, graphene and other porous carbon structures can be used for electrode preparation. Carbon aerogels show great promise for use as electrode materials in low-temperature fuel cells because of their high surface area, good electrical conductivity, chemical stability and it is possible to control the porous structure of the material. CAs for use as electrocatalysts in ORR have previously been prepared from various phenolic compounds like resorcinol and its derivatives [58,71,72,73,74,75], melamine [76,77,78], ployacrylonitrile [79], cellulose [80] and sustainable carbohydrate based derivatives like glucose, D-glucosamine aso [81,82].

The use of Pt-doped carbon aerogels for PEM fuel cell applications has been studied previously [58,71,72,73,80,83,84,85]. However, as nowadays most of the research is directed towards development of non-precious metal catalysts, there have also been attempts to prepare non-precious metal-containing carbon aerogels that would have high activity for ORR. This can be done by choosing the right precursors for aerogel preparation or through further modification of aerogels [70].

There are examples where carbon aerogels have previously been doped with nitrogen [81,82] or dual-doped with nitrogen and sulfur [86]. The metal free carbon aerogels showed increased activity towards ORR compared to non-doped materials, but this activity was still rather low [81,82,86]. Carbon aerogels or xerogels can be doped with nitrogen by either using different monomers which contain nitrogen in their structure, like melamine or
polyacrylonitrile \[70,76,77,79,87,88\] for aerogel preparation or by further modifying aerogels for instance by pyrolysis in a NH$_3$ atmosphere \[74,75\].

Doping CAs or xerogels with transition metals has been shown to increase their activity for ORR \[74,75,76,77,78,79,89\]. The transition metal was introduced either by adding the precursor to the sol before gelation \[74,75,76,77\] or was added later to the already existing aerogel \[89\].

For example, nitrogen-and cobalt-doped organic xerogels have been prepared from resorcinol and formaldehyde and tested as catalysts for ORR \[74,75\]. Cobalt nitrate was added to the materials before gelation. The content of cobalt in the carbonized xerogels was very low however (not higher than about 0.5 wt%). The materials were carbonized under NH$_3$ atmosphere at 800 °C. These materials showed similar activity to commercial Pt/C in alkaline solution \[75\], but somewhat lower activity in acidic media \[74\]. In both cases the catalysts showed good stability. The physical characterisation of the catalysts that were tested in alkaline solution showed that the amorphous carbon materials had metallic cobalt nanoparticles and the prevalent nitrogen species were pyridinic and pyrrolic, which have previously been observed in other active ORR catalysts \[75,90,91\].

Rather similar activities for ORR in acidic solution were observed for Co- or Fe-doped xerogels prepared from melamine instead of resorcinol \[76,77\]. Melamine is a good choice because of its affordable price and the high nitrogen content in the molecule.

It has also been shown that varying the melamine/resorcinol ratio in the synthesis of aerogels enables one to adjust the microstructure of the materials \[70,78\].
2. EXPERIMENTAL

2.1. Reagents

5-methylresorcinol with a reported a purity > 99% was provided by AS VKG, Estonia and 2,6-dihydroxy-4-methylbenzoic acid, with a purity > 99% was provided by AS Carboshale OÜ, Estonia. Sodium acetate (CH₃COONa*3H₂O), with a purity of 99.7 %, was obtained from Fisher Chemical, USA. The water used was purified employing a Milli-Q water system. Acetonitrile, HPLC purity, was obtained from Rathburn Chemical Ltd., Germany. Formaldehyde (37% w/w solution in water) was purchased from Sigma-Aldrich. CO₂ (99.8%) was obtained from AS Eesti AGA. Melamine, with a purity > 99%, was purchased from Aldrich Chemistry, USA. Anhydrous sodium carbonate with a purity of 99.8% was purchased from Riedel-de Haen, Germany.

Lanthanide (La, Ce, Pr, Nd) carbonates were obtained from AS Silmet (Molycorp), Estonia and were turned into nitrates by adding the carbonates to a stoichiometric amount of nitric acid and evaporating the excess solution.

Cobalt(II)nitrate hexahydrate with a purity > 98%, nickel(II) nitrate hexahydrate with a purity > 97% and copper(II)chloride with a purity > 97% were purchased from Sigma-Aldrich, USA.

2.2. Instrumentation and characterisation

The supercritical extraction system with a double clamp autoclave, 100 ml in volume, was constructed by NWA analytische Meßgeräte GmbH, Germany.

The MTF 12/38/400 pyrolysis oven with a maximum temperature of 1200°C was from Carbolite, England.

The bulk densities of carbon aerogels were calculated by measuring the dimensions and mass of each monolithic sample. Elemental analyses were performed by Spectra AA 220F flame (Agilent, USA) atomic absorption spectrometer (AAS), inductively coupled plasma optical emission spectrometry OPTIMA 7300 DV ICP OES (Perkin Elmer, USA) or using the Vario Micro Cube (Elementar, Germany).

Lanthanide contents were measured using an OPTIMA 7300DV ICP OES (Perkin Elmer). Nitrogen adsorption analyses were performed using a Sorptometer KELVIN 1042 built by Costech International. Helium was used as a carrier gas, nitrogen as an adsorptive gas. The specific surface area (S_BET) was calculated according to the Brunauer-Emmett-Teller theory. The specific micropore volume (V_mic) was determined via the t-plot and the pore size distributions were determined using the Barrett-Johner-Halendar method from the desorption branch. The XRD pattern was recorded with a Rigaku Ultima IV diffractometer (Japan) using a line detector DtexUltra. Cu-Kα radiation was used with a Ni filter for removing Kβ radiation. The recorded diffractograms were analysed by Rigaku PDXL software. Energy-filtered transmission electron
microscopy (TEM) was carried out with the Leo 912 Omega microscope (Zeiss, Germany) at 120 kV, using a LaB$_6$ gun. Raman spectroscopy was carried out using a Raman spectrometer (Horiba Jobin Yvon LabRam HR800, Japan). SEM images were taken using ZEISS ULTRA 55 (Germany) microscope.

Thermogravimetric analysis (TGA) was conducted using a Labsys Evo TG-DTA 1600 °C (Setaram, France) and heating the samples in an Ar atmosphere at 10 °C/min to 800 °C.

2.3. Aerogel preparation

2.3.1. Metal doped aerogels prepared from 5-methylresorcinol and 2,6-dihydroxy-4-methylbenzoic acid

The gels were prepared by adding dHMBA and MR (the molar ratio of MR to dHMBA was either 1:1, 3:1 or 9:1) to the solvent (70 wt% acetonitrile and 30 wt% distilled water). The molar ratio of the solvent to aromatic monomers (MR and dHMBA) was 35:1. The moles of aromatic monomers (AMs) were calculated adding together the moles of MR and dHMBA and the moles for the solvent were calculated adding together the moles of acetonitrile and water. To the solution of aromatic monomers and the solvent a stoichiometric amount of sodium acetate (NaOAc) to dHMBA was added (the molar ratio of NaOAc to dHMBA was 1:1) to obtain the sodium salt of dHMBA. After that formaldehyde (FA) was added (the molar ratio of AM to FA was 1:2) and finally another amount of NaOAc was added as a catalyst (the molar ratio of AM to NaOAc was 50:1).

Glass molds containing the solutions were sealed and the mixture was allowed to gel and cure at 40 °C for seven days. The obtained Na$^+$-doped gels were orange in colour and transparent. The gels were then soaked for three days in 0.1 M aqueous solutions of CuCl$_2$, Co(NO$_3$)$_2$ * 6H$_2$O, Ni(NO$_3$)$_2$ * 6H$_2$O or lanthanide nitrates, respectively. Each time the solution was renewed after every 24 hours. After that the gels were washed by soaking them in water for 24 hours, followed by acetone exchange, which lasted four to five days. Acetone was changed every 24 hours. The gels were dried with supercritical CO$_2$. A more detailed description of the drying method is published elsewhere [27]. Carbonization was performed in a N$_2$ atmosphere by raising the temperature to 700 °C (2.5°C/min), from then on to 900 °C (5 °C/min) and was kept at 900 °C for one hour.

2.3.2. Preparation of metal-and nitrogen-doped aerogels

The amounts of substances (in grams) to be used for gel preparation were calculated by the following equation (1):

\[
\frac{\text{substances}}{\text{water}} = \frac{\text{monomers+FA+Na}_2\text{CO}_3}{\text{water added+water from FA solution}} = 0.25 \tag{1}
\]
The ratio of substances to water was always kept at 0.25. During preparation the amounts of monomers (Mel, MR and dH MBA) were varied from 10 to 75 mol%:

\[
\text{mol\%}_1(\text{Mel}) + \text{mol\%}_2(\text{MR}) + \text{mol\%}_3(\text{dH MBA}) = 100\%(\text{monomers}) \quad (2)
\]

The amount of sodium carbonate used was calculated adding the amount needed for converting dH MBA into the sodium salt of dH MBA (the molar ratio of dH MBA to Na\textsubscript{2}CO\textsubscript{3} was 2) and the catalytic amount (the molar ratio of monomers to Na\textsubscript{2}CO\textsubscript{3} was 100). The molar ratio of FA to melamine was 3.7 and that of FA to resorcinol derivatives (MR and dH MBA) was 2. At first two separate solutions were prepared in two vials. In one vial a mixture of Mel, the FA solution and half of the water used was prepared. The mixture was kept at 70 °C for about 25 min to allow melamine to dissolve. In the second vial a mixture of MR, dH MBA, sodium carbonate and half of the water was prepared. Both homogeneous solutions were poured into one vial, mixed and poured into glass test tubes and allowed to gel and cure for four days at 85 °C.

To obtain Co\textsuperscript{2+}-doped gels, the hydrogels were soaked in 0.1 M solution of cobalt nitrate for three days (the solution was changed every 24 hours), after which the sodium ions bound to the ion-exchange moiety (the carboxylic group) were exchanged for cobalt ions.

In order to use the supercritical CO\textsubscript{2} drying for aerogel preparation, the exchange of the solvent for acetone was carried out changing the solution every 24 hours, starting with a 25:75 mixture of acetone and water and proceeding with 50:50 and 75:25 mixtures and finally with 100% acetone for two days. A more detailed description of the method used for supercritical drying has been published elsewhere [27].

Carbon aerogels were prepared by pyrolysis in a N\textsubscript{2} atmosphere by heating at 10 °C/min to 800 °C and keeping at this temperature for one hour. The Co-containing N-doped aerogels prepared using different amounts of monomers are denoted as Co-NCA1, Co-NCA2, etc. (Table 4). For comparison, the nitrogen-free Co-containing carbon aerogel (Co-CA) and the cobalt-free material (NCA) were also prepared. In the latter case, the ion-exchange procedure was not performed.
3. RESULTS

3.1. Co-, Cu- and Ni-doped aerogels

In a previously developed method for the preparation of aerogels from MR and dHMBA, it was suggested that dHMBA in addition to being a monomer also acted as an acidic catalyst [27]. This was not the case with the materials prepared in this work, because NaOAc was also added to the sol, first to exchange the hydrogen proton for sodium ion in the carboxylic moiety and secondly an extra amount of NaOAc was added to act as a catalyst.

In the method developed by Baumann et al [44], potassium carbonate was used to neutralize the solution and as a catalyst, but in our case a stronger gel was obtained using sodium acetate instead. The use of NaOAc in our case resulted in the sol being slightly acidic (pH 5.5) due to the formation of acetic acid and therefore incomplete neutralization of the solution.

The gels containing Na⁺ ions were immersed in the solution of copper ions, after which the Na content in the obtained organic aerogels was measured and was found to be about 0.005 %. For comparison, in the same aerogel that had not been immersed in the solution of copper ions, the Na content was 6.40 %. It confirms that the exchange of sodium ions for copper ions takes place.

The effect that dHMBA has on the metal content in aerogels was studied. Gels at MR to dHMBA molar ratios of 1:1, 3:1 and 9:1 were prepared and dried. The results are shown in Figure 11. It can be seen that by increasing the dHMBA content in aerogels the metal content also increases, which is due to the presence of more ion-exchange moieties. Therefore, changing the MR to dHMBA molar ratio is an easy way to control the metal content in aerogels. The metal content in CAs was mostly higher than in the organic aerogels due to the burn-off of the organic material. CAs prepared at a 1:1 molar ratio of MR to dHMBA had about twice as high metal content as OAs.

In a previous study the ratio of the two phenolic compounds used (R and DHBA) was varied, to evaluate the ion-exchange capacity. Though it was also observed that at lower relative mole fractions of the ion-exchange monomer the ion-exchange capacity decreased, a change in metal content to the same extent as ours was not shown [48].
Table 1 presents the densities, specific surface areas and pore volumes of metal-doped carbon aerogels. The densities of organic and carbon aerogels were respectively in the range of 0.45-0.62 g/cm$^3$ and 0.83-1.26 g/cm$^3$. These values were higher than those of metal-doped carbon aerogels prepared from DHBA [92]. A possible explanation is that in our case we used higher solids content to prepare aerogels, which is known to cause higher densities [12].

Table 1 presents the specific surface areas and pore volumes of carbon aerogels doped with different metals. The surface areas and pore volumes of CAs doped with transition metals (Co, Cu, Ni) were larger than those of the corresponding blank sample (containing Na). Similar results have been obtained before for aerogels prepared from DHBA (the blank sample contained potassium) [92].

Table 1. Properties of Co-, Cu- and Ni-doped aerogels

<table>
<thead>
<tr>
<th>Element</th>
<th>$\delta$(OA) (g/cm$^3$)</th>
<th>$\delta$(CA) (g/cm$^3$)</th>
<th>$S_{\text{BET}}$ (m$^2$/g)</th>
<th>$S_{\text{mic}}$ (m$^2$/g)</th>
<th>$%S_{\text{mic}}$ (%)</th>
<th>$V_{\text{TOT}}$ (mm$^3$/g)</th>
<th>$V_{\text{mic}}$ (mm$^3$/g)</th>
<th>$%V_{\text{mic}}$ (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Na</td>
<td>0.58</td>
<td>0.83</td>
<td>286</td>
<td>133</td>
<td>47</td>
<td>429</td>
<td>47</td>
<td>11</td>
</tr>
<tr>
<td>Cu</td>
<td>0.45</td>
<td>1.01</td>
<td>464</td>
<td>291</td>
<td>63</td>
<td>607</td>
<td>103</td>
<td>17</td>
</tr>
<tr>
<td>Co</td>
<td>0.62</td>
<td>1.03</td>
<td>519</td>
<td>353</td>
<td>68</td>
<td>610</td>
<td>124</td>
<td>21</td>
</tr>
<tr>
<td>Ni</td>
<td>0.59</td>
<td>1.26</td>
<td>379</td>
<td>273</td>
<td>72</td>
<td>446</td>
<td>96</td>
<td>22</td>
</tr>
</tbody>
</table>

Remark: All the samples were prepared at MR to dHMBA molar ratios of 1:1. All the properties measured correspond to those of carbon aerogels, unless otherwise indicated. $\rho$ is the calculated density, $S_{\text{BET}}$ is the Brunauer-Emmett-Teller surface area, $S_{\text{mic}}$ is the microporous surface area, $%S_{\text{mic}}$ is the percent of microporous surface area, $V_{\text{TOT}}$ is the total pore volume, $V_{\text{mic}}$ is the volume of micropores, $%V_{\text{mic}}$ is the percent of microporous volume.
Transition metal-containing aerogels showed higher microporous surface area and microporous pore volume values compared to the Na-containing sample. The percent of microporous area and volume was almost twice as high for aerogels that contained transition metals. The increase in the micropore volume of transition metal-containing CAs could be attributed to the metals contributing to the formation of new microporosity during carbonization [36], or causing a more significant shrinkage of mesopores which due to that become micropores during carbon aerogel preparation [18,47]. The increase in the pore volume of transition metal-containing aerogels was due not only to the increase in microporosity, but also that the non-microporous volume of Cu and Co containing carbon aerogels was larger.

Pore size distributions (PSD) in Figure 12 show that for the Na-containing aerogel the peak in pore size distribution occurs around the pore diameters of 10-15 nm, whereas in the case of all the other doped aerogels the peak maximum is around 5-10 nm. Therefore, in this case, the addition of transition metals resulted in the reduction of pore diameters of carbon aerogels. This result is different from the finding of a previous study, which established that in the case of carbon aerogels made from DHBA, the incorporation of transition metals did not have a significant effect on pore size distributions [92].

![Figure 12. Pore size distributions of metal-doped carbon aerogels, molar ratio of MR to dHMBA was 1:1. The legend shows the metal the carbon aerogel contains.](image)

The X-ray diffraction analysis results for carbon aerogels are shown in Figure 13. The diffractograms indicate the presence of peaks corresponding to metals (oxidation state 0) for all CAs. This implies that the metal ions were reduced during carbonization. For the copper-doped carbon aerogel there is also a peak corresponding to the copper(I)oxide present on the diffractogram. This could be caused by either incomplete reduction of copper during pyrolysis or oxidation afterwards.
TEM pictures of aerogels were taken to study the morphology of the materials. Figure 14a shows an organic aerogel, which before drying was soaked in the solution of cobalt nitrate. No metal nanoparticles were observed in this case, which is in agreement with previous studies where it was also concluded that metal ions were uniformly distributed throughout the organic aerogel [44]. When the same aerogel in presented in Figure 14a was pyrolysed (Figure 14b) the nanoparticles emerged. These were uniformly distributed in the carbon aerogel matrix and their size was in the range of 10-60 nm. Figure 14c and 14d show nickel- and copper-doped carbon aerogels prepared under the same conditions as the previously mentioned Co-doped carbon aerogel. The size of metal nanoparticles in the samples ranged of 20 to 150 nm for nickel-containing carbon aerogels and 3-100 nm for copper-containing carbon aerogels, showing that the metal nanoparticle diameters for different metals varied. In copper-containing carbon aerogels the nanoparticles had also clustered together. The pictures taken of metal-doped CAs from different locations showed that the metal nanoparticles were evenly distributed throughout the material.

Figure 14e shows a carbon aerogel prepared using a smaller amount of dHMBA (the molar ratio of MR to dHMBA was 9:1). It can be seen from the figure that the metal nanoparticles are smaller in size (about 2 to 40 nm) and are
not clustered together. This is due to the smaller amount of metal in the carbon aerogel. It has been observed before that higher metal content leads to larger metal nanoparticles in CAs [44].

Figure 14. a-d) TEM pictures of MR/dHMBA 1:1 metal-doped aerogels. a) Co-doped organic aerogel; b) Co-doped carbon aerogel; c) Ni-doped carbon aerogel; d) Cu-doped carbon aerogel. e) copper-doped CA with a MR/dHMBA molar ratio of 9:1.

In the SEM picture (Fig 15a) of the Co-doped CA prepared at an MR to dHMBA molar ratio of 1:1, the cobalt nanoparticles are also distinguishable in the aerogel carbon network. A picture taken from the same location using backscattered electron (BSE) imaging shows the areas composed of heavier elements light up, confirming that these are cobalt nanoparticles. The morphology of the carbon aerogel matrix in Fig 15a and c indicates that the structure of the aerogel is similar to that of R-FA aerogels which are described as “polymeric”, meaning that they are composed of small particles that are conjoined by large necks [12].

Figure 15. SEM pictures of Co-doped CAs prepared at a MR/dHMBA molar ratio of 1:1. Picture b was taken using backscattered electron imaging.
3.2. Lanthanide-doped aerogels

The same method that was used to prepare Cu-, Co-, and Ni-doped aerogels from MR and dHMBA was employed to make lanthanide-doped (Ln-doped) organic and carbon aerogels. The Ln-doped aerogels were prepared at MR to dHMBA molar ratio of 1:1.

The amount of lanthanides in aerogels was determined by inductively coupled plasma optical emission spectrometry (ICP OES) and the results are given in Table 2. The amount of lanthanides in OAs was about 11 wt% and in CAs 21 wt%, which was higher than in the corresponding Co-, Cu- and Ni-doped aerogels. This can be explained by the higher molecular mass of lanthanide compounds.

Table 2. Properties of La-, Ce-, Pr- and Nd-doped organic and carbon aerogels.

<table>
<thead>
<tr>
<th></th>
<th>OA</th>
<th>CA</th>
<th>OA</th>
<th>CA</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>$\rho$ (g/cm$^3$)</td>
<td>$S_{BET}$ (m$^2$/g)</td>
<td>$V_{TOT}$ (mm$^3$/g)</td>
<td>$V_{mic}$ (mm$^3$/g)</td>
</tr>
<tr>
<td>La</td>
<td>0.54</td>
<td>429</td>
<td>889</td>
<td>2.0</td>
</tr>
<tr>
<td>Ce</td>
<td>0.52</td>
<td>393</td>
<td>734</td>
<td>2.5</td>
</tr>
<tr>
<td>Pr</td>
<td>0.53</td>
<td>371</td>
<td>721</td>
<td>3.1</td>
</tr>
<tr>
<td>Nd</td>
<td>0.53</td>
<td>378</td>
<td>659</td>
<td>2.0</td>
</tr>
</tbody>
</table>

Remark: $\rho$ is the calculated density, $S_{BET}$ is the Brunauer-Emmett-Teller surface area, $V_{TOT}$ is the total pore volume, $V_{mic}$ is the volume of micropores, $V_{mic}$% is the percent of microporous volume and metal% is the weight percent of metal in aerogels.

Nitrogen adsorption analysis results of organic and carbon aerogels (Table 2) show that the materials were mostly mesoporous. The total pore volume and specific surface area of organic aerogels were higher than those of the corresponding carbon aerogels. The decrease in porosity during pyrolysis can be explained by the shrinkage of pore diameters. Figure 16 shows the pore size distribution for La-doped organic and carbon aerogels. It can be seen that the overall shape of the PSD curve remains the same but it is shifted towards smaller diameters for La-doped CAs.

Organic aerogels have almost no microporous pore volume, while the $V_{mic}$% of carbon aerogels can be up to 18%. A certain portion of the microporosity was probably formed during pyrolysis by the burn-off of organic material, but some amount was likely generated additionally due to the shrinkage of mesopores [12].
Figure 16. Pore size distribution of La-doped organic and carbon aerogels (MR/dHMBA molar ratio was 1:1).

The morphology of the prepared materials was determined from the TEM and SEM pictures of OAs and CAs, which were taken from different locations and are shown in Figure 17. Both the TEM and SEM pictures show that lanthanide nanoparticles are not distinguishable in the pictures. In previous studies, nanoparticles in metal-doped organic aerogels were also not observed [44,63]. During pyrolysis metal-ions in aerogels tend to reduce and form metal nanoparticles [18]. In this work it was not observed, showing that lanthanides require stronger reducing conditions to form nanoparticles. Except for the lack of metal nanoparticles, the morphology of La-doped CAs (Fig. 17e) is similar to that of the Co-doped CAs (Fig. 15a).

Figure 17. a,b) TEM pictures of a La-doped organic aerogels; c,d) TEM pictures of La-doped carbon aerogels; e) SEM picture of a La-doped carbon aerogel.
The amorphous nature of Ln-doped aerogels was also confirmed by XRD analysis results as shown in Figure 18a. Both La-doped OAs and CAs had no peaks in the diffractogram, which would have referred to the presence of crystal structures in aerogels. For metal-doped carbon aerogels it is common that various crystalline structures like oxides, metals (oxidation state 0) and graphitic structures are formed during pyrolysis [44,45,93], but in this case the material remained amorphous.

Raman spectroscopy on CAs was performed to evaluate the nature of carbon [94]. In Figure 18b the peak at 1590 cm\(^{-1}\) (G-band) can be attributed to the 2E\(_{2g}\) mode, which is present in both ordered and amorphous carbon materials. The peak at 1330 cm\(^{-1}\) (D-band) can be attributed to the A\(_{1g}\) mode, which is associated with defects in the G layers of the graphenic carbon.

![Figure 18](image.png)

**Figure 18.** a) XRD analysis results for La-doped organic and carbon aerogels; b) Raman spectra for La-doped CA.

### 3.3. Lanthanide-doped aerogels as heterogeneous catalysts

The Ln-doped OAs and CAs were tested for use as heterogeneous catalysts in the Michael addition reaction of ethyl-2-oxocyclopentanecarboxylate 1 with 3-buten-2-one 2 (Scheme 1).

![Scheme 1](image.png)

**Scheme 1.** Michael addition reaction of ethyl 2-oxocyclopentanecarboxylate (1) with 3-buten-2-one (2) in the presence of Ln-aerogels as catalysts. 0.006 mmol of the catalyst (3 mol% Ln with respect to the limiting reagent) was carried into the reaction vessel. 0.4 ml of CH\(_2\)Cl\(_2\), 0.6 mmol (excess of 3 times) of 3-buten-2-one and 0.2 mmol of ethyl 2-oxocyclopentanecarboxylate (the limiting reagent) were added and the mixture was stirred at room temperature for 24 hours. Conversion of the product was determined by gas chromatography. For OAs and CAs catalyst loadings of 7.5 mg and 4.0 mg were used, respectively [95].
The results in Table 3 demonstrate that Ln-doped OAs were active catalysts in the Michael addition reaction. The conversions within 24 hours ranged from 82 to 97%.

La-, Pr- and Nd-doped carbon aerogels also exhibited some activity in the Michael addition reaction, however their activity was lower than that of OAs. In the case of Ce-doped CAs the formation of several by-products was observed. The decrease of catalytic activity of these CAs can be explained by the structural and chemical changes taking place during pyrolysis. After pyrolysis all lanthanide species might not be accessible to the reactants anymore and the lanthanide ions previously bound to the ion-exchange center may have formed new chemical bonds.

In a previous study, a mixture of Eu oxides, sulphides and fluorides had formed during pyrolysis (confirmed by XRD analysis) and the CA did not catalyse the reaction, but the Eu-doped OAs, which did not show crystalline structures in the XRD diffractogram (Figure 18a), was catalytically active [63]. In our case the presence of oxides or other crystalline structures was not detected in the XRD diffractogram of CAs. The TEM and SEM pictures did not show the presence of metal nanoparticles either. The absence of metal oxide nanoparticles could be the reason for our Ln-doped CAs being catalytically active, since it is known that rare-earth metal oxides are mostly inert as catalysts in the Michael addition reaction [63].

Table 3. Conversions for Michael addition reaction of ethyl 2-oxocyclopentanecarboxylate with 3-buten-2-one in the presence of Ln-doped aerogels as catalysts.

<table>
<thead>
<tr>
<th>Element</th>
<th>Conversion (%)</th>
<th>OA</th>
<th>CA</th>
</tr>
</thead>
<tbody>
<tr>
<td>La</td>
<td>84</td>
<td>49</td>
<td></td>
</tr>
<tr>
<td>Ce</td>
<td>82</td>
<td>-</td>
<td></td>
</tr>
<tr>
<td>Nd</td>
<td>97</td>
<td>92</td>
<td></td>
</tr>
<tr>
<td>Pr</td>
<td>91</td>
<td>32</td>
<td></td>
</tr>
</tbody>
</table>

3.4. **Co-and N-doped carbon aerogels as electrocatalysts for ORR**

To obtain active electrocatalysts for ORR, carbon aerogels were doped with nitrogen and cobalt or iron [70]. Besides MR, dHMBA and FA, melamine was also used as a monomer for gel preparation to introduce nitrogen into the materials. Melamine was chosen because of its affordable price and the presence of pyridinic nitrogen atoms in its molecule, which are expected to be active catalytic sites for ORR [90,91].

The first experiments in which the electrocatalytic activity of Co- or Fe- and N-doped CAs was measured, showed that the Co-doped CAs proved to be more active than the Fe-doped ones [70]. Materials pyrolysed at different
temperatures were also tested for electrocatalytic activity, the ones pyrolysed at 800 °C being the most active.

Subsequently, the composition of CAs was varied during the preparation process, using different amounts of melamine, MR and dHMBA, to study the effect this variation has on the properties of CAs and their activity towards ORR. Table 4 presents the molar percentages of monomers used during the preparation of different CAs and the experimentally measured elemental compositions of the samples. Co-containing N-doped CAs prepared using different amounts of monomers are denoted as Co-NCA1, Co-NCA2, etc. (Table 4). For comparison, nitrogen-free Co-containing CA (Co-CA) and a cobalt-free material (NCA) were also prepared. In the latter case, the ion-exchange procedure was not performed.

Table 4. Properties of Co- and N-doped carbon aerogels.

<table>
<thead>
<tr>
<th>Sample</th>
<th>Monomers (mol%)</th>
<th>Elemental composition (wt%)</th>
<th>(S_{\text{BET}}) (m²/g)</th>
<th>(V_{\text{TOT}}) (mm³/g)</th>
<th>(V_{\text{MIC}}) (mm³/g)</th>
<th>MA</th>
</tr>
</thead>
<tbody>
<tr>
<td>Co-NCA1</td>
<td>30 60 10</td>
<td>2.0 1.6 87.0 1.3</td>
<td>370 330</td>
<td>85 2.8</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Co-NCA2</td>
<td>30 45 25</td>
<td>4.3 2.5 82.2 1.0</td>
<td>314 230</td>
<td>70 3.2</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Co-NCA3</td>
<td>40 40 20</td>
<td>4.1 2.9 80.2 1.4</td>
<td>461 399</td>
<td>95 3.4</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Co-NCA4</td>
<td>60 15 25</td>
<td>5.3 2.6 81.0 0.7</td>
<td>316 228</td>
<td>54 14.4</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Co-NCA5</td>
<td>75 0 25</td>
<td>5.6 8.4 68.9 1.2</td>
<td>118 93</td>
<td>1.5 12.5</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Co-CA</td>
<td>0 75 25</td>
<td>2.6 0.03 n.d. n.d</td>
<td>n.d. n.d</td>
<td>n.d. n.d</td>
<td>2.3 0.23</td>
<td></td>
</tr>
<tr>
<td>NCA</td>
<td>60 15 25</td>
<td>0 14.3 65.2 1.3</td>
<td>n.d. n.d</td>
<td>n.d. n.d</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Remark: \(S_{\text{BET}}\) is the Brunauer-Emmett-Teller surface area, \(V_{\text{TOT}}\) is the total pore volume, \(V_{\text{MIC}}\) is the volume of micropores, MA is the mass activity (A/g) at -0.2 V.

The molar percentage (mol%) of dHMBA was varied from 10 to 25 mol%. With the increase in the amount of dHMBA in the composition of the aerogel, the Co content also increased (Table 4). This shows that changing the dHMBA amount in the aerogel enables to control the metal content, which is due to dHMBA having the ionexchange moiety (the carboxylic group) that binds Co²⁺ during immersion in the cobalt-nitrate solution. This result is in accordance with previous results of analysing Co-, Cu- and Ni-doped MR/dHMBA aerogels [93].

It was also possible to vary the nitrogen content in the aerogels by changing the melamine content. The nitrogen content in cobalt-containing samples was changed from 1.6 to 8.4 wt% by increasing the amount of melamine (Table 4). The increase in melamine content also decreased the carbon content of aerogels from 87 to 69 wt%.

\(\text{N}_2\)-adsorption analysis was used to determine the specific surface areas and pore volumes of CAs (Table 4). Samples Co-NCA1 to Co-NCA4 had high surface areas (\(S_{\text{BET}} = 314\text{–}461\text{ m}^2/\text{g}\)) and total pore volumes (\(V_{\text{TOT}}\)) of 228 to 399 mm³/g. Figure 19a shows the pore size distribution of sample Co-NCA2, indicating that in this material pores with a diameter (\(d_p\)) of 2 and 4 nm predominate.

Co-NCA5 had significantly lower \(S_{\text{BET}}\) and \(V_{\text{TOT}}\) values compared to samples Co-NCA1 to Co-NCA4. During preparation, Co-NCA5 had the highest
melamine content and no MR was added during preparation. The $S_{\text{BET}}$ and $V_{\text{TOT}}$ values of the organic aerogel precursor of Co-NCA5 were also measured, the respective figures being 868 m$^2$/g and 1490 mm$^3$/g. This indicates that a drastic reduction in specific surface area and pore volume occurred during pyrolysis.

![Figure 19. a) Pore size distribution analysis result for Co-NCA2; b) TGA results for materials Co-NCA 1 – Co-NCA 5. Samples analysed were pre-pyrolysed (organic aerogels).](image)

Thermogravimetric analysis (TGA) results illustrated in Figure 19b show the mass-loss curves for organic aerogels (precursors of Co-NCA1–Co-NCA5) during heating in an Ar atmosphere using the same heating program that was used during pyrolysis for the Co-NCA materials. The mass loss of those OAs was between 56 and 80%. Samples that were prepared using higher melamine contents (Co-NCA3 to Co-NCA5) had similar mass losses (60–65 wt%). The mass-loss curves for Co-NCA1 and Co-NCA2 were smooth in shape, without abrupt changes. However, for Co-NCA4 and Co-NCA5, there was a steep change in mass at 340–370 °C. From literature it can be found that melamine resins also experienced significant mass loss, owing to the decomposition and evaporation of the products generated in the above temperature range [96]. The decomposition of melamine and, therefore, the structure of the material could explain the smaller surface area and porosity of CAs that were prepared using higher melamine contents.

The surface morphology of CAs was evaluated using scanning electron microscopy (Figure 20). The pictures were taken from different locations on the material. The morphology of the CA that was prepared using the highest MR content (Co-NCA1) was similar to that of resorcinol/formaldehyde CAs [12], with a grainy structure composed of linked spheres (Figure 20a and 20b). The Co-NCA2 aerogel was prepared using more dHMB and less MR than for Co-NCA1. In the structure of Co-NCA2 (Figure 20d), separate particles forming the aerogel network were not visible and the morphology was different from that of resorcinol/formaldehyde CAs. This was also the case with samples prepared using higher melamine contents (Co-NCA4 and Co-NCA5), whose surfaces appear to be composed of particles that have merged together (Fig.
This demonstrates that the addition of resorcinol derivatives is necessary to keep the bead-like structure that is associated with resorcinol/formaldehyde CAs. The structure of melamine-rich aerogels will be affected by the sudden mass loss and decomposition of melamine during pyrolysis [96].

Figure 20c shows a SEM image of Co-NCA1 taken using backscattered electron (BSE) imaging, where brighter spots may be assigned to Co nanoparticles. These were evenly distributed in the sample and had sizes ranging from 5 to 40 nm (Figures 20a–c). The cobalt nanoparticles in Co-NCA3 were also evenly distributed (Figure 20g), but their sizes were larger (10–100 nm) than in Co-NCA1 (Figures 20e and 20f). This was caused by the increased amount of metal in the aerogel [44,93]. Samples Co-NCA4 and Co-NCA5 (Fig. 20h–l) had smaller nanoparticles (10–40 nm) than Co-NCA3, and their SEM images indicated the formation of long cylindrical structures with a diameter of
about 20 nm, which appeared to be carbon nanotubes (CNTs) or nanoribbons (Figures 20h,j,k). The BSE image (Figure 20j,l) shows brighter spots (Co nanoparticles) at the end of the long structures.

Cobalt nanoparticles are known to be used as catalysts for synthesis of CNTs [97]. The growth of nanoribbons has been observed before in cobalt-doped carbon aerogels [45]. From nitrogen-containing precursors, N-doped CNTs have been synthesised previously [98,99], so it is possible that the long carbon nanostructures in Co-NCA4 and Co-NCA5 are also nitrogen-doped CNTs [100].

Figure 21 shows X-ray diffraction (XRD) peaks for crystalline Co in all samples, which is an expected result under these pyrolysis conditions [45,93]. A small peak attributed to Co$_3$O$_4$ indicates the oxidation of the surface of Co nanoparticles. The formation of graphitic structures is visible in samples Co-NCA1 and Co-NCA2. In Co-NCA5, a peak at the same location cannot be identified. This indicates that the increasing melamine content inhibits the formation of graphitic structures in this case. In the diffraction patterns shown in Figure 21, there are a few other small peaks visible at various degrees. The identification of these peaks is problematic, due to the limited amount of data. It is likely that various cobalt–carbon, cobalt–nitrogen or nitrogen–carbon structures have also formed in CAs.

![Figure 21. XRD analysis results for the Co-NCA materials.](image-url)
3.5. Oxygen reduction reaction activity of Co-NCA catalysts

The electrocatalytic activity of the materials for ORR was studied using the rotating disk electrode (RDE) method in 0.1 M KOH solution [100]. RDE voltammetry curves of O₂ reduction for materials Co-NCA1 - Co-NCA5 are shown in Figure 22a. Co-NCA 4 and Co-NCA 5 were the most active. These materials had the highest cobalt content. The increased activity towards ORR has previously been observed for Co₃O₄-containing N-doped graphene [101]. The activity increased with higher Co content. Both the Co-and Co₃O₄ nanoparticles, which were observed in the XRD diffracitogram for Co-NCAs (Fig. 21), could attribute to the ORR activity, according to the literature [101,102,103,104].

Compared to Co-NCA4, the nitrogen content of Co-NCA5 was significantly higher, although their activities were similar. This suggests that only a small fraction of nitrogen is composing the active centres for ORR and the remaining part is either in a passive form or inaccessible to O₂ molecules [100]. According to the literature, only certain types of nitrogen centres are considered to be involved in ORR electrocatalysis [5] and the activity of the catalysts does not directly depend on the total nitrogen content [105].

The surface area and pore volume of Co-NCAs do not seem to have a great effect on the ORR activity of the materials, because Co-NCA5, which has a low surface area and porosity, has a similar activity to Co-NCA4, whose S_{BET} and V_{TOT} values are much higher. However, the nanotubes or nanoribbons observed in Co-NCA4 and Co-NCA5 may contribute to their activity [100]. Both the metal-free and transition metal-containing N-doped CNTs have been shown to be very active catalysts for ORR in alkaline solutions [106,107,108].

The activity of the prepared Co-NCAs for ORR has been compared to that of Co- and N-free materials and also a commercial Pt/C catalyst (20 wt% Pt catalyst supported on Vulcan carbon XC-72(E-TEK)) [100]. Figure 22b shows that the activity of CA which contained nitrogen, but no cobalt (material marked NCA in Fig. 22b), had a much lower ORR activity than the Co-NCA materials. However, the ORR activity of the Co-CA material (contained cobalt, but no nitrogen), was higher than that of NCA, but still lower than for Co-NCA4. This suggests that the synergistic effect of cobalt or cobalt-oxide nanoparticles with nitrogen is what gives the Co-NCAs their high activity for ORR. This has also been proposed previously in literature [101,102,109].

Compared to the commercial Pt/C catalyst the Co-NCA4 was slightly less active (Fig. 22b). However in a short-term stability test the Co-NCA4 proved to be more stable [100].
Figure 22. RDE voltammetry curves of oxygen reduction reaction for a) Co-containing N-doped CAs and b) for various CA-based catalysts, glassy carbon (GC) and Pt/C catalyst in O₂-saturated 0.1 M KOH; n=10 mVs⁻¹, w=1900 rpm [100].
4. CONCLUSIONS

In this work, a method for preparing metal-doped organic and carbon aerogels from 5-methylresorcinol and 2,6-dihydroxy-4-methylbenzoic acid was developed. The specific conclusions drawn are the following:

- The ion-exchange method was used to dope organic and carbon aerogels with Co, Cu, Ni, Ce, La, Nd and Pr.
- It was shown that this method allows one to easily control the metal content in aerogels. In Co-, Cu- and Ni-doped aerogels varying the MR to dHMBA molar ratio during preparation enabled changing the metal content in organic aerogels from around 1 to 8 wt% and in carbon aerogels from around 1 to 15 wt%. Ln-doped aerogels had metal contents of approximately 12 and 22 wt% for OAs and CAs, respectively. All the metal-doped organic and carbon aerogels had high specific surface areas (ca 250-520 m²/g) and pore volumes (ca 300-900 mm³/g). During pyrolysis Co-, Cu- and Ni-ions were reduced and metal nanoparticles formed in carbon aerogels. The size of nanoparticles ranged from 3 to 150 nm depending on the type and amount of the metal used for doping. However, the Ln-doped CAs remained amorphous and metal nanoparticles were not observed in their structure.
- Ce-, La-, Nd- and Pr-doped OAs and CAs were tested as catalysts for the Michael addition reaction. Organic aerogels containing different lanthanides catalysed the Michael addition reaction efficiently, with conversions ranging from 82 to 97 %. For carbon aerogels the conversion was modest, 32 % (Pr-doped) to high, 92 % (Nd-doped).
- To prepare active catalysts for the oxygen reduction reaction, the incorporation of nitrogen into the structure of CAs was necessary. A method for preparing Co- and N-doped carbon aerogels from MR and dHMBA was developed using melamine for nitrogen doping. The results showed that by varying the dHMBA or melamine content, the composition of CAs may be altered. Co-NCAs showed high surface area and had both meso- and microporosity. The increasing amount of melamine in the composition of aerogels decreased their surface area and pore volume. The morphology of the materials also changed drastically with increasing melamine amount. All the CAs had crystalline structures of cobalt and cobalt oxide as well as cobalt nanoparticles present in these materials. In CAs prepared using higher melamine and cobalt concentrations, carbon nanotube structures had also formed.
- The Co- and N-doped carbon aerogels were tested as electrocatalysts for the oxygen reduction reaction. The electrocatalytic activity of the materials for this reaction was rather high and depended on their composition. The materials with the highest Co content showed the highest activity for oxygen reduction reaction, which can be associated with the coexistence of cobalt nanoparticles and N-doped carbon structures in them.
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ABSTRACT

In this thesis a method for preparing metal-doped organic and carbon aerogels was developed, the materials were characterised and tested as heterogeneous catalysts. The aerogels were prepared from 5-methylresorcinol (MR), which is a by-product of the local oil shale processing industry, and 2,6-dihydroxy-4-methylbenzoic acid (dHMBA), which can easily be synthesized from MR. dHMBA has an ion-exchange moiety (carboxylic group) that allowed the ion-exchange method to be used for metal-doping. The materials were doped with metals such as Co, Cu, Ni, La, Ce, Pr and Nd.

The properties of the materials and their dependence on the preparation conditions were discussed. The structure and morphology of metal-doped carbon aerogels were characterised using methods like nitrogen adsorption analysis, scanning electron microscopy, transmission electron microscopy, X-ray diffraction analysis, different elemental analysis techniques, etc.

It was shown that the developed method allowed us to easily control the metal content in aerogels. Varying the MR to dHMBA molar ratio in Co-, Cu- and Ni-doped aerogels during preparation enabled to change the metal content in organic aerogels from around 1 to 8 wt% and in carbon aerogels from around 1 to 15 wt%. Ln-doped aerogels had metal contents of approximately 12 and 22 wt% for OAs and CAs, respectively. All the metal-doped organic and carbon aerogels had high specific surface areas (about 250-520 m²/g) and pore volumes (about 300-900 mm³/g). The materials were micro- and mesoporous. During pyrolysis the Co-, Cu- and Ni-ions present in organic aerogels were reduced and metal nanoparticles formed in carbon aerogels. The size of nanoparticles ranged from 3 to 150 nm, depending on the type and amount of metal used for doping. However, the Ln-doped CAs remained amorphous and metal nanoparticles were not observed in their structure.

Ce-, La-, Nd- and Pr-doped OAs and CAs were tested as catalysts for the Michael addition reaction. The initial results showed that organic aerogels containing different lanthanides catalysed the Michael addition reaction efficiently, with conversions ranging from 82 to 97 %. For carbon aerogels the conversion was modest, 32 % (Pr-doped) to high, 92 % (Nd-doped).

The method developed for making metal-doped carbon aerogels was further modified to prepare active electrocatalysts for the oxygen reduction reaction. The cobalt-doped aerogels were also doped with nitrogen by adding melamine during carbon aerogel preparation. The results showed that by varying the dHMBA or melamine content, the cobalt and nitrogen content in materials could easily be changed. Cobalt- and nitrogen doped carbon aerogels (Co-NCAs) showed high surface area and had both meso- and microporosity, although the increasing amount of melamine in the composition of aerogels decreased their surface area and pore volume. The morphology of the materials also changed drastically with increasing melamine amount. All the CAs had crystalline structures of cobalt and cobalt oxide present in them. Cobalt nanoparticles were present in all Co-NCAs and also carbon nanotube structures.
had formed in those Co-NCAs that were prepared using higher melamine and cobalt concentrations.

The Co- and N-doped carbon aerogels were tested as electrocatalysts for the oxygen reduction reaction. The electrocatalytic activity of the materials for the reaction was rather high and depended on their composition. The materials with the highest Co content showed the highest activity for the oxygen reduction reaction, which can be associated with the coexistence of cobalt nanoparticles and N-doped carbon structures in them.
KOKKUVÕTE

Antud töös töötati välja meetod valmistamaks metalli sisaldavaid orgaanilisi- ja süsinikaerogelee põlevkivitööstuse jääkproduktidest ning uuriti võimalusi nende materjalide kasutamiseks katalüsaatoritena. Aerogeelid valmistati 5-metüülresortsinoolist (MR), mis on fenoolne ühend mida eraldatakse põlevikivi töötlemisel tekkinud ütteveest, ning 2,6-dihüdroksü-4-metüülbensoehappest (dHMBA), mida on võimalik lihtsalt sünteesida 5-metüülresortsinoolist. Kuna dHMBA molekul sisaldab karboksüülrühma, siis oli võimalik kasutada metallide viimiseks materjali ioonvahetus meetodit, mis tagab metalli ioonide ühtlase jaotumise. Selles töös valmistati koobaltit, vaske, niklit ja erinevaid lantanoide (La, Ce, Nd, Pr) sisaldavaid aerogeele.

Valmistatud aerogeelide struktuuri ja omadusi karakteriseeriti kasutades erinevaid meetodeid nagu lämmastik adsorptsioon analüüs, röntgendifraktsioon analüüs, skaneeriv ja transmisioon elektronmikroskoopia, mitmed erinevad elementsisalduse määramise meetodid jne. Tähelepanu pöörati seostele aerogeeli valmistamise tingimuste ja omaduste vahel.

Tulemuste osas on näidatud, et välja töötatud meetodi puhul on võimalik lihtsalt kontrollida metallide sisaldust materjalis. Co-, Cu- ja Ni-sisaldavate aerogeelides oli võimalik MR ja dHMBA moolsuhet valmistamisel muutes varieerida metalli sisaldust orgaanilistes aerogeelides ühest kuni kaheksee protsendini ja süsinikaerogeelides ühest kuni viiteistkümne protsendini. Lantanoide sisaldavad aerogeelides oli metalli sisaldus umbes 12% ning süsinikaerogeelides umbes 22%. Nii orgaanilised kui ka süsinikaerogeelid omasid kõrget eripinda umbes (umbes 250-520 m²/g) ja poorsust (umbes 300-900 mm³/g). Co-, Cu- ja Ni-sisaldavate süsinikaerogeelide puhul oliid tekkinud metalli nanoosakesed pürolüüsi käigus. Nende nanoosakeste suurused, jäädes vahemikku 3-150 nm, sõltusid kasutatud metallist ning selle sisaldusest materjalis. Lantanoide sisaldavad aerogeelide puhul metalli nanoosakeste teksti ei täheldatud.

La-, Ce-, Nd- ja Pr-sisaldavad orgaanilisi- ja süsinikaerogelee testiti katalüsaatoritena Michaeli reaktsioonis. Tulemused näitasid, et lantanoide sisaldavad orgaanilised aerogeelid omasid kõrget katalüüutilist aktiivsust Michaeli reaktsioonis (konversioon oli 82-97 %). Kõik süsinikaerogeelid aga ei osutunud nii aktiivseks (konversioon 32-92%).


Koobaltit ja lämmastikku sisaldavate süsinikaerogeelide katalüütilist aktiivsust testiti hapniku redutseerumis reaktsioonis ning need materjalid omasid kõrget katalüütilist aktiivsust. Parimad olid kõige kõrgemat koobalti sisaldust omavad materjalid.
Publication I

PREPARATION OF METAL-DOPED CARBON AEROGELS FROM OIL SHALE PROCESSING BY-PRODUCTS

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\textbf{Abstract.} Carbon aerogels are nanostructured porous carbon materials, which can be produced from locally available phenolic compounds. A method for the preparation of metal-doped carbon aerogels from oil shale processing by-products 5-methylresorcinol and 2,6-dihydroxy-4-methylbenzoic acid was developed. Aerogels doped with Ni, Co and Cu were characterized by atomic absorption spectroscopy, nitrogen adsorption, transmission electron microscopy and X-ray diffraction. The resulting materials exhibited pore diameters in microporous and mesoporous regions, high surface areas and pore volumes. The metal content in aerogels was found to be dependent on the amount of ion-exchange moiety and metal content as high as 14.5 wt\% was achieved in carbon aerogels, where also metal nanoparticles had formed.

\textbf{Keywords:} aerogel, organic aerogel, carbon aerogel, 5-methylresorcinol, 2,6-dihydroxy-4-methyl benzoic acid, supercritical drying, oil shale, ion-exchange, metal-doped, nanoparticles.

\section{1. Introduction}

Nanoporous materials, due to their unique surface, structural and bulk properties, have attracted a lot of interest and acquired many important applications in several areas. Aerogel, as an example of a nanoporous material, is a low density material with high surface area and porosity. There are various types of aerogels which are prepared from different precursors.

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Carbon aerogels are made by pyrolysing organic aerogels. For example, Pekala was the first to obtain organic aerogels by supercritically drying gels, which were made through sol-gel polymerization by using resorcinol (R) and formaldehyde (FA) as precursors [1, 2].

Previously, the preparation of aerogels from 5-methylresorcinol (MR) and 2,6-dihydroxy-4-methylbenzoic acid (dHMBA), has been studied in our laboratory. MR is a good alternative to resorcinol. Besides MR being a by-product of oil shale industry and therefore a locally available precursor, gels prepared from MR have also shown faster gelation times compared to gels prepared from resorcinol [3–5]. The preparation of aerogels from MR and dHMBA together showed that the structure and properties of aerogels are dependent on the molar ratio of MR to dHMBA. dHMBA differs from MR by being a bifunctional monomer and, in addition to being a monomer, also acts as a catalyst due to its acidic properties [6]. dHMBA can be synthesized from MR.

Doping of carbon aerogels prepared from MR and FA with palladium by using impregnation in supercritical CO₂ has been previously carried out, but to the authors’ knowledge no attempt to dope aerogels prepared from dHMBA has been made before [5].

There is on-going activity to expand potential aerogel applications with the modification of carbon aerogels (CA) through the incorporation of metal species into carbon frameworks. Doping of carbon aerogels with metals modifies their structure, conductivity and catalytic activity [7–12]. There are different methods for preparing metal-doped carbon aerogels, which can roughly be divided into three categories, as was done by Moreno-Castilla and Maldonado-Hódar [13]. First, the metal precursor can be added to the initial mixture before gel formation, secondly, the metal precursor can be deposited onto the already existing organic or carbon aerogel and lastly, the ion-exchange method can be used [13].

The ion-exchange method for preparing metal-doped aerogels was first studied by Baumann et al. [12]. Carbon aerogels were prepared in which instead of resorcinol the researchers used the potassium salt of 2,4-dihydroxybenzoic acid (dHBA) in the sol-gel process, producing K⁺-doped hydrogels. This meant that each repeat unit of the gel contained a binding site for metal ions. The desired metal ion was deposited by exchanging the potassium ion in an ion-exchange process. A common carbon aerogel treatment process for gel drying and carbonization was used to generate metal-doped carbon aerogels.

Over the years, several studies on ion-exchange deposition have been reported [12, 14–20]. In these researches, the effect of different metals, pyrolysis temperature and different applications have been described. Recently, a paper was published in which the ion-exchange capacity of gels was altered by using together dHBA (ion-exchange moiety containing monomer) and resorcinol [21].

The goal of this work is to modify the preparation of aerogels from precursors MR and dHMBA, so that metal-doped carbon aerogels could be
obtained. The doping of materials with metals will be carried out similarly to the ion-exchange method developed by Baumann et al. [12], though in the current case two resorcinol derivates (MR and dHMBA) will be used for aerogel preparation. Since only dHMBA has an ion-exchange moiety (carboxylic group), then the possibility of preparing aerogels with different metal contents, by changing the MR to dHMBA molar ratio, will be discussed. The process for preparation of metal-doped carbon aerogels from MR and dHMBA will be developed and properties of the obtained materials determined.

2. Experimental

2.1. Materials

5-methylresorcinol with a reported purity of > 99% was provided by AS VKG, Estonia and 2,6-dihydroxy-4-methylbenzoic acid, with a purity of >99%, was provided by Carboshale OÜ, Estonia. Sodium acetate (CH$_3$COONa·3H$_2$O), with a purity of 99.7%, was obtained from Fisher Scientific, UK. The water used was purified using a Milli-Q water system. Acetonitrile, HPLC purity, was obtained from Rathburn Chemical Ltd., Germany. Formaldehyde (37 w/w solution in water) was purchased from Sigma-Aldrich. CO$_2$ (99.8%) was obtained from Eesti AGA AS.

The supercritical extraction system with a double clamp autoclave, 100 ml in volume, was constructed by NWA analytische Meßgeräte GmbH, Germany.

The MTF 12/38/400 pyrolysis oven with a maximum temperature of 1200 °C was from Carbolite, England.

2.2. Sample preparation

The gels were prepared by adding dHMBA and MR (MR:dHMBA molar ratio 1:1, 3:1 or 9:1) to the solvent (70 wt% acetonitrile and 30 wt% distilled water). The molar ratio of solvent (SOLV) to aromatic monomers (AM) was 35:1. The moles of aromatic monomers (AM) were calculated by adding up the moles of MR and dHMBA and the moles for the solvent were calculated by adding up the moles of acetonitrile and water. To the solution of aromatic monomers and solvent a stoichiometric amount of sodium acetate (NaOAc) to dHMBA was added (NaOAc:dHMBA molar ratio 1:1) to obtain the sodium salt of dHMBA. After that formaldehyde (FA) was added (AM:FA molar ratio 1:2) and, finally, another amount of NaOAc was added as a catalyst (AM:NaOAc molar ratio 50:1).

Glass molds containing the solutions were sealed and the mixture was allowed to gel and cure at 40 °C for 7 days. The obtained Na⁺-doped gels were orange and transparent. The gels were then soaked for 3 days in 0.1 M aqueous solutions of CuCl$_2$, Co(NO$_3$)$_2$·6H$_2$O and Ni(NO$_3$)$_2$·6H$_2$O, respectively. Each time the solution was renewed after 24 hours. After that the
gels were washed by soaking them in water for 24 hours, followed by acetone exchange which lasted 4 to 5 days. Acetone was changed every 24 hours. The gels were dried with supercritical CO₂. A more detailed description of the drying method has been published elsewhere [6]. Carbonization was performed in N₂ atmosphere by raising the temperature to 700 °C (2.5 °C/min), then to 900 °C (5 °C/min) and was kept at 900 °C for 1 hour.

2.3. Characterization

The bulk densities of carbon aerogels were calculated by measuring the dimensions and mass of each monolithic sample. Elemental analyses were performed by a Spectra AA 220F flame atomic absorption spectrometer (Australia). Nitrogen adsorption analyses were performed using a KELVIN 1042 Sorptometer (Costech International, Italy). Helium was used as a carrier gas, nitrogen as the adsorptive gas. Nitrogen adsorption data was taken at relative pressures from 0 to 1 and at liquid nitrogen temperature. The specific surface area (S_{BET}) was calculated according to the Brunauer-Emmet-Teller theory. The specific micropore volume (V_{mic}) was determined via the t-plot and the pore size distributions were determined using the Barrett-Johnner-Halender (BJH) method from the desorption branch. The X-ray diffraction (XRD) pattern was recorded with a Rigaku Ultima IV diffractometer using the D/teX Ultra line detector (USA). Cu-Kα radiation was used with a Ni filter for removing Kβ radiation. The recorded diffractograms were analysed by Rigaku PDXL software. Energy-filtered transmission electron microscopy (TEM) was carried out with the Leo 912 Omega microscope (Zeiss, Germany) at 120 kV, using a LaB₆ gun.

3. Results and discussion

In the previously developed method for the preparation of aerogels from MR and dHMBA, it was suggested that dHMBA, in addition to being a monomer, also acted as an acidic catalyst [6]. The method developed in this work involved, during gel preparation, the addition of NaOAc, which can also act as a catalyst. Therefore the role of dHMBA as a catalyst is not clear in the present case.

The method developed by Baumann et al. employed potassium carbonate to neutralize the solution and as a catalyst, but in the current work a stronger gel was obtained by using sodium acetate instead. This resulted in a slightly acidic solution (pH 5.5) due to the formation of acetic acid and, hence, incomplete neutralization of the solution.

After immersing the gels containing Na⁺ ions in the copper ions containing solution, the Na content in organic aerogels was measured and found to be about 0.005%. In comparison, the Na content in the aerogel, which had not been immersed in the solution containing copper ions, was 6.40%. It confirms that the exchange of sodium ions for copper ions takes place.
The authors studied which effect dHMBA has on the metal content in aerogels. Gels at MR to dHMBA molar ratios of 1:1, 3:1 and 9:1 were prepared and dried. The results are shown in Figure 1. It can be seen that with increasing dHMBA content in aerogels their metal content also increases. Therefore changing the MR to dHMBA molar ratio is an easy way to control the metal content in aerogels. Previously a study was performed where the ratio between the two phenolic compounds used (R and dHBA) was changed, to evaluate the ion-exchange capacity. Though it was also noted that at lower relative mole fractions of the ion-exchange monomer the ion-exchange capacity decreased, a change in metal content to the same extent as in the current case was not shown [21].

Subsequently, carbon aerogels at an MR:dHMBA molar ratio of 1:1 were prepared (Table). The metal content in carbon aerogels was about twice as high as in the organic aerogel. It can be explained by the decomposition of organic matter during pyrolysis.

The Table also presents data on the density, specific surface area and pore volume of metal-doped carbon aerogels. The density of organic aerogels ranged from 0.45 to 0.62 g/cm³ and that of carbon aerogels from 0.83 to 1.26 g/cm³, which is higher than for metal-doped carbon aerogels prepared from dHBA [16]. This may be explained by that in the present case the authors used higher solids content to prepare aerogels, which is known to cause higher densities [22]. Also, the diameter of gels decreased about 20% during the immersion of gels in metal ion solutions and during solvent exchange.

Nitrogen adsorption/desorption isotherms for metal-doped carbon aerogels are shown in Figure 2. The Table presents the specific surface areas and pore volumes of carbon aerogels doped with different metals. While comparing carbon aerogels that were doped with transition metals (Co, Cu, Ni) to the blank sample (Na containing), it was found that both specific surface areas and pore volumes were larger for the transition metal containing samples. The same effect has been observed before for aerogels prepared from dHBA (the blank sample contained potassium) [12, 16].

Fig. 1. Metal content in organic aerogels.
Table. Properties of metal-doped carbon aerogels

<table>
<thead>
<tr>
<th>Sample</th>
<th>$\rho$(OA), g/cm$^3$</th>
<th>$\rho$(CA), g/cm$^3$</th>
<th>$S_{BET}$, m$^2$/g</th>
<th>$S_{mic}$, m$^2$/g</th>
<th>$S_{total}$, m$^2$/g</th>
<th>%-$S_{mic}$</th>
<th>$V_{mic}$, mm$^3$/g</th>
<th>%-$V_{mic}$</th>
<th>Metal, %</th>
<th>OA</th>
<th>CA</th>
</tr>
</thead>
<tbody>
<tr>
<td>Na</td>
<td>0.58</td>
<td>0.83</td>
<td>286</td>
<td>153</td>
<td>133</td>
<td>46.5</td>
<td>429</td>
<td>47</td>
<td>11.0</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Cu</td>
<td>0.45</td>
<td>1.01</td>
<td>464</td>
<td>173</td>
<td>291</td>
<td>62.7</td>
<td>607</td>
<td>103</td>
<td>17.0</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Co</td>
<td>0.62</td>
<td>1.03</td>
<td>519</td>
<td>166</td>
<td>353</td>
<td>68.0</td>
<td>610</td>
<td>124</td>
<td>20.3</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Ni</td>
<td>0.59</td>
<td>1.26</td>
<td>379</td>
<td>106</td>
<td>273</td>
<td>72.0</td>
<td>446</td>
<td>96</td>
<td>21.5</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

All the samples were prepared at an MR:dHMBA molar ratio of 1:1. OA is organic aerogel and CA is carbon aerogel. All the parameter values obtained apply to carbon aerogels, unless otherwise noted. $\rho$ is the calculated density, $S_{BET}$ is the BET surface area, $S_{mic}$ is the microporous surface area, %-$S_{mic}$ is the percent of microporous surface area, $V_{mic}$ is the total pore volume, $V_{mic}$ is the volume of micropores, %-$V_{mic}$ is the percent of microporous volume, metal % is the weight percent of metal in aerogels.

![Graph showing adsorption/desorption analysis of nitrogen](image)

Fig. 2. Results of adsorption/desorption analysis of nitrogen; (ads) – adsorption isotherm, (des) – desorption isotherm.

Transition metal containing aerogels showed higher values of microporous surface area and microporous pore volume compared to the Na containing sample, the respective percentages being almost twice as high for the aerogels. The increase in micropore volume could be attributed to the transition metals favouring the formation of new microporosity or causing a stronger shrinkage of mesopores, which become micropores during carbon aerogel preparation as a result. The increase in pore volume of transition metal containing aerogels was due not only to the increase in microporosity, but also to the fact that the non-microporous volume of Cu and Co containing carbon aerogels was larger.
From Figure 3 it can be seen that the peak of pore size distribution (PSD) of the Na containing aerogel is at a pore diameter of 10–15 nm, whereas the peak for all the other doped aerogels is at 5–10 nm. This indicates that for carbon aerogels the addition of transition metals resulted in the reduction of pore diameter. This result is different from those obtained in a previous study, which showed that in case of carbon aerogels made from dHBA, incorporation of transition metals had no significant effect on pore size distribution [16].

![Graph showing pore size distribution of metal-doped carbon aerogels](image)

Fig. 3. Pore size distribution of metal-doped carbon aerogels (MR:dHMBa molar ratio 1:1).

The results of X-ray diffraction analysis of carbon aerogels are shown in Figure 4, revealing the presence of peaks corresponding to metals (oxidation state 0). This suggests that metal ions must have been reduced during carbon aerogel preparation. For the copper-doped carbon aerogel there is also a peak corresponding to copper(II)oxide, whose presence could have been caused by either incomplete reduction during pyrolysis or oxidization of copper afterwards. The diffractogram for nickel-doped carbon aerogel has also peaks corresponding to graphite (marked C in the figure). However, no graphitic structures were observed in TEM pictures, possibly due to low resolution.

TEM pictures of aerogels were taken to visualize metal nanoparticles in samples. Figure 5a shows an organic aerogel, which before drying was soaked in the solution of cobalt nitrate. No metal nanoparticles can be observed in this case, which is in agreement with the conclusion of a previous study that metal ions are uniformly distributed throughout the organic aerogel [12]. When the aerogel depicted in Figure 5a is pyrolysed (Fig. 5b), then nanoparticles emerge. These are uniformly distributed in the carbon matrix and their size is in the range of 10–60 nm. Figure 5c and d depict nickel- and copper-doped carbon aerogels prepared in the same conditions as the above-mentioned Co-doped carbon aerogel. The size of metal nanoparticles in these samples is in the range of 20–150 nm for nickel containing...
carbon aerogels and 3–100 nm for copper containing carbon aerogels, showing that the metal nanoparticle diameters of different metals varied. For copper containing carbon aerogels the nanoparticles had also clustered together.

Fig. 4. Results of X-ray diffraction analysis of metal-doped carbon aerogels.

Fig. 5. TEM pictures of metal-doped aerogels. a: Co-doped organic aerogel; b: Co-doped carbon aerogel; c: Ni-doped carbon aerogel; d: Cu-doped carbon aerogel.
4. Conclusions

A method for preparing metal-doped carbon aerogels from oil shale processing by-products 5-methylresorcinol and 2,6-dihydroxy-4-methylbenzoic acid was developed. The ion-exchange process was used to introduce Ni, Co and Cu into carbon aerogels.

The specific surface area of metal-doped carbon aerogels was in the range of 380–520 m²/g and pore volume 446–610 mm³/g. It was shown that the metal content in carbon aerogels can be varied by changing the molar ratio of MR to dHMBA. In carbon aerogels the metal ions had reduced to metals during pyrolysis and formed metal nanoparticles. The size of metal nanoparticles ranged from 3 to 150 nm.

The method developed is suitable for preparing metal-doped carbon aerogels from oil shale processing by-products. The resulting materials have high surface area and porosity and it is possible to control their metal content.

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Publication II

Organic and carbon aerogels containing rare-earth metals: Their properties and application as catalysts

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**ABSTRACT**

Lanthanide-doped organic and carbon aerogels were obtained based on the modified resinolic-formaldehyde aerogel preparation procedure. The prepared materials were tested as catalysts for the Michael reaction. Organic aerogels were prepared by sol-gel polymerization of 2,6-dihydroxy-4-methylbenzoic acid (dHMB) mixed with 5-methylresorcinol (MR) and formaldehyde. The resulting gels were doped with Ln(NO₃)₃ (Ln = Ce, La, Nd, Pr) by ion-exchange and further dried with supercritical CO₂. Pyrolysis of organic aerogels at 900 °C in N₂ atmosphere resulted in metal-doped carbon aerogels. The mostly mesoporous materials contained up to 12 wt% of metal in organic aerogels and 22 wt% in carbon aerogels. The materials were amorphous and crystal structures were not formed during pyrolysis.

Ln-doped aerogels were tested as catalysts for Michael addition reaction. Organic aerogels doped with different lanthanides catalyze the Michael reaction efficiently with conversions ranging between 82 and 97% in 24 h. For Ln-doped carbon aerogels the conversion was modest (32%) to high (92%).

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1. Introduction

Aerogels as nanoporous materials have attracted a lot of interest and found many important applications in several areas due to their unique surface, structural and bulk properties. The most common method for making organic aerogels (OAs) involves the sol–gel polymerization of resorcinol and formaldehyde and subsequent supercritical drying of the gel [1,2]. Upon pyrolysis of OAs, carbon aerogels (CAs) are produced in which the initial morphology of the OAs remains, but surface areas can be higher and also the nonconductive OAs become electrically conductive.

The addition of metal species to carbon aerogels has been studied before and several methods have been used for the deposition of metal species [3]. The incorporation of metals modifies the structure, conductivity and catalytic activity of the materials obtained [4–7]. One of the methods used for doping is the ion-exchange method in which gels having an ion-exchange moiety are prepared [8].

Lanthanides are widely used as catalysts in organic synthesis, because lanthanide-mediated reactions display high chemo- and stereoselectivities [9]. Due to the high coordination numbers (f-block elements), lanthanides can promote organic reactions catalytically rather than stoichiometrically. Their association with substrates and dissociation of products occurs fast, giving high catalyst efficiency. Lanthanide complexes have great advantages as Lewis acid catalysts for carbon–carbon bond forming reactions [10,11], including the Michael reaction [12,13].

The use of a lanthanide salt as an efficient catalyst for the Michael reaction was first reported in 1993 by Sceti and co-workers [14]. This article broke the previous unwritten rule that the catalyst had to be a complex bearing donor ligands [15], it was shown that these species would be readily generated in situ from the 1,3-dicarboxyl compound. Since then, other lanthanide species of La, Sc and Yb have been studied as catalysts in conjugate additions [16,17].

Due to the various advantages of supported catalysts, it is important to find the right support for lanthanides. Aerogels are good candidates for catalyst carriers, because of their high surface area and continuous porosity. The use of carbon aerogels as catalyst carriers has been reviewed before [3]. Support or entrapment of lanthanides or lanthamide species in aerogels has been rarely studied [6,19]. As far as we know, only one example about the use of lanthanide-aerogels as heterogeneous catalysts in Michael reaction has been described [19].

This paper presents the first known attempt to prepare of La, Ce-, Pr- and Nd-doped organic and carbon aerogels. Their properties were studied and the materials tested as catalysts in a reaction which needs lanthanide ions for catalytic activity. The materials used for making the aerogel are by-products of the local oil-shale processing industry [20,21].
2. Materials and methods

2.1. Materials

5-Methyl Resorcinol (5-MR) with a reported purity of >99% was provided by VSG, AS, Estonia and 2,6-dihydroxy-4-methylenbenzoic acid (dHBMA), with a purity of >99% was donated by Carbosole OY, Finland. Sodium acetate trihydrate, with a purity of 99.7%, was obtained from Fisher Chemical, USA. The water used was purified using a Milli-Q water system. Acetonitrile, HPLC purity, was obtained from Rathburn Chemicals Ltd, Germany. Formaldehyde (37 wt/w solution in water) was purchased from Sigma-Aldrich. CO₂ (99.8%) was obtained from Eesti AGA AS. Lanthanum carbonates were obtained from Molycorp Silmet, Estonia and were turned into nitrates. The supercritical extraction system with a double clamp autoclave, 100 mL in volume, was constructed by NWA Analytische Meßgeräte GmbH, Germany. The pyrolysis oven type MTF 12/38/400 with a maximum temperature of 1200 °C was manufactured by Carbolite, England.

2.2. Sample preparation

The two aromatic monomers (AM) used, 5-MR and dHBMA (molar ratio of 5-MR:dHBMA = 1), were added to the solvent (70 wt.% acetonitrile and 30 wt.% distilled water). The molar ratio of solvent to aromatic monomers (5-MR and dHBMA) was 35. The moles of aromatic monomers were calculated by adding the moles of 5-MR and dHBMA and the moles for the solvent were calculated by addition of the moles of acetonitrile and water. A stoichiometric amount of sodium acetate (NaOAc) was added to dHBMA (molar ratio of NaOAc/dHBMA = 1) to obtain the sodium salt of dHBMA. After that formaldehyde (FA) was added to the solution (AM/FA molar ratio 0.5) and finally another amount of NaOAc as a catalyst was added (AM/NaOAc molar ratio 50).

Glass molds containing the solutions were sealed and the mixture was allowed to gel and cure at 40 °C for 7 days. The obtained Na⁺-doped gels were orange and transparent. The gels were then soaked for 3 days in 0.1 M aqueous solutions of lanthanide nitrates [La(NO₃)₃, Ce(NO₃)₃, Nd(NO₃)₃, Pr(NO₃)₃], respectively. Each time the solutions of lanthanide nitrates were renewed after 24 h. After that the gels were washed by soaking them in water for 24 h, followed by acetone exchange which lasted 4 to 5 days. Acetone was changed every 24 h. The gels were dried with supercritical CO₂. A more detailed description of the drying method is published elsewhere [22]. Carbonization was performed in N₂ atmosphere by raising the temperature to 700 °C (2.5 °C/min), from then on to 900 °C (5 °C/min) and was kept at 900 °C for 1 h.

2.3. Characterization

The bulk densities of organic and carbon aerogels were calculated by measuring the dimensions and mass of three monolithic samples and calculating the average. Standard deviations are added to Table 1. Lanthanide contents were measured using an OPTIMA 7300 DV ICP OES, PerkinElmer. The uncertainty of the results is ±10%. Nitrogen adsorption analyses were performed using a KELVIN 1042 Sorptometer built by Costech International. Helium was used as a carrier gas, nitrogen as the adsorbate. Nitrogen adsorption data was taken at relative pressures from 0 to 1 and at liquid nitrogen temperature. The specific surface area (SBET) was calculated according to the Brunauer-Emmett-Teller (BET) theory. The specific micropore volume (Vmic) was determined via the t-plot and the pore size distributions were determined using the Barrett-Joyner-Halenda (BJH) method from the desorption branch. The uncertainty of the method is ±3%. The X-ray diffraction (XRD) pattern was recorded with a Rigaku Ultima IV diffractometer by using a D-Theta Ultra line detector. Cu-Kα radiation was used with a Ni filter for removing Kα radiation. The recorded diffractograms were analyzed by Rigaku PDF2 software. Energy-filtered transmission electron microscopy (TEM) was carried out with a Leo 912 Omega microscope at 120 kV, using a LaB₆ gun. Fourier transform infrared spectroscopy (FT-IR) was performed using a TENSOR 27 (Bruker) FT-IR spectrometer. Raman spectroscopy was carried out using a Raman spectrometer (Horiba Jobin Yvon LabRam HR800). Thermogravimetric analysis was carried out using TG-DTA Setaram Labsys Evo in Ar atmosphere with the same heating program as was used during pyrolysis.

2.4. Experimental data on Michael reaction

0.006 mmol of the catalyst (3 mol% Ln with respect to the limiting reagent) was carried into the reaction vessel. 0.4 mL of CH₂Cl₂, 0.6 mmol (excess of 3 times) of 3-buten-2-one 2 and 0.2 mmol of ethyl 2-oxocyclopentanecarboxylate 1 (limiting reagent) were added and the mixture was stirred at room temperature for 24 h. Conversion of the product 3 was determined by GC. For OAs a catalyst loading of 7.5 mg and for CA 4.0 mg was used.

---

Table 1

<table>
<thead>
<tr>
<th>p(LOA), g/cm³</th>
<th>p(LOA), g/cm³</th>
<th>SBET (LOA), m²/g</th>
<th>Vmic (LOA), cm³/g</th>
<th>Vtot (LOA), cm³/g</th>
<th>SBET (OA), m²/g</th>
<th>Vmic (OA), cm³/g</th>
<th>Vtot (OA), cm³/g</th>
<th>Vmic (OA), cm³/g</th>
<th>Metal (LOA), %</th>
<th>Metal (OA), %</th>
<th>Metal (CA), %</th>
</tr>
</thead>
<tbody>
<tr>
<td>La</td>
<td>0.54 ± 0.02</td>
<td>1.16 ± 0.08</td>
<td>429</td>
<td>889</td>
<td>0.36 ± 0.1</td>
<td>368</td>
<td>103</td>
<td>131</td>
<td>45.0</td>
<td>11.0 ± 1.1</td>
<td>22 ± 2.2</td>
</tr>
<tr>
<td>Ce</td>
<td>0.52 ± 0.09</td>
<td>1.03 ± 0.1</td>
<td>393</td>
<td>734</td>
<td>2.5</td>
<td>255</td>
<td>533</td>
<td>18.2</td>
<td>11.5 ± 1.2</td>
<td>21 ± 2.1</td>
<td></td>
</tr>
<tr>
<td>Pr</td>
<td>0.53 ± 0.06</td>
<td>1.14 ± 0.06</td>
<td>371</td>
<td>721</td>
<td>2.1</td>
<td>257</td>
<td>301</td>
<td>17.3</td>
<td>11.3 ± 1.1</td>
<td>22 ± 2.2</td>
<td></td>
</tr>
<tr>
<td>Nd</td>
<td>0.53 ± 0.07</td>
<td>1.08 ± 0.08</td>
<td>378</td>
<td>609</td>
<td>2</td>
<td>257</td>
<td>301</td>
<td>17.3</td>
<td>11.3 ± 1.1</td>
<td>22 ± 2.2</td>
<td></td>
</tr>
<tr>
<td>Na</td>
<td>0.58 ± 0.06</td>
<td>0.83 ± 0.1</td>
<td>221</td>
<td>645</td>
<td>0</td>
<td>257</td>
<td>301</td>
<td>17.3</td>
<td>11.3 ± 1.1</td>
<td>22 ± 2.2</td>
<td></td>
</tr>
</tbody>
</table>

Fig. 1. Pore size distributions of OA-s and CA-s.
3. Results and discussion

3.1. Materials characterization

The method for introducing lanthanides into aerogels involves the exchange of sodium ions for lanthanide ions in the ion-exchange moiety. The wt.% of lanthanides in aerogels (measured by ICP-MS) is shown in Table 1. The theoretical (about 11 wt.% if all the ion-exchange moieties were available for binding) and measured wt.% of lanthanides in organic aerogels are similar, enabling one to conclude that the ion-exchange of sodium ions for lanthanides was successful. After pyrolysis the wt.% of lanthanides increased about twice. It can be attributed to the combustion of organic matter.

The results of nitrogen adsorption analysis for organic and carbon aerogels (Table 1) show that the materials are mostly mesoporous. The total pore volumes and specific surface area of organic aerogels were higher than those of carbon aerogels. This indicates that carbonization during pyrolysis decreased the pore volume of aerogels, also their pore size is smaller (Fig. 1). Organic aerogels have a very small microporous pore volume, while the $V_{\text{mic}}$ of carbon aerogels can be up to 18.2%. A certain portion of the microporosity was probably formed additionally during pyrolysis, but some amount was likely generated due to the shrinking of mesopores.

As a reference, a blank sample, which was not immersed in an aqueous solution of lanthanide salt (contained sodium instead), was prepared. The surface areas and pore volumes of lanthanide-doped aerogels are higher for both OA-s and CA-s compared to the blank samples.

![Fig. 2. TEM pictures of aerogels: (a, b) La-doped organic aerogel; (c, d) La-doped carbon aerogel.](image)

![Fig. 3. XRD analysis results of La-doped organic and carbon aerogels.](image)
For carbon aerogels the difference in surface area and pore volume between lanthanide-doped aerogels and the blank sample is vast. During pyrolysis most of the mesoporosity disappeared in the blank sample, whereas for aerogels with lanthanides the decrease was not this drastic.

The morphology of the prepared materials was determined from the TEM pictures of organic and carbon aerogels, which were taken from different locations and are shown in Fig. 2. Images a and b correspond to La-doped organic aerogel and c and d to La-doped carbon aerogels. Judging from the pictures, the materials seem to mostly consist of the amorphous aerogel matrix, without sharply distinguishable nanopr1cles. In previous studies, no distinguishable nanoparticles in organic aerogels were observed [8,21]. During pyrolysis aerogels enriched with metal ions tend to reduce and form metal nanoparticles. In this work it was not observed, which confirms the fact that lanthanides require stronger reducing agents.

This is also confirmed by XRD analysis results illustrated in Fig. 3. Both La-OAs and -CAs are amorphous materials with no peaks referring to crystal structures in aerogels. For metal-doped carbon aerogels it is common that crystal structures are formed during pyrolysis [8,23], but in this case the material remained amorphous.

Raman spectroscopy on CAs was performed to evaluate the nature of carbon [24]. In Fig. 4 the peak at 1590 cm$^{-1}$ (G-band) can be attributed to the 2E2g mode, which is present in both ordered and amorphous carbon materials. The peak at 1330 cm$^{-1}$ (D-band) can be attributed to the A1g mode, which is associated with defects in the G layers of graphitic carbon.

The IR spectra of La-OA and La-CA are shown in Fig. 5. For La-OA there are absorptions present at several wavenumbers. Absorption at wavenumber 1500 cm$^{-1}$ can be attributed to an asymmetric stretching from carboxylic salt. The stretching at 1240 cm$^{-1}$ can be attributed to the C–O bond from phenolic compounds and at 1100 cm$^{-1}$ to the C–O bond from methylene ether bridges. However, for the same aerogel after pyrolysis only a small absorption band at 3420 cm$^{-1}$, which can be attributed to the O–H bond from adsorbed water, can be observed. This shows that the initial organic material underwent transformation during pyrolysis.

To describe the changes in mass that take place during pyrolysis of the organic aerogel, thermogravimetric analysis was conducted in Ar atmosphere for La-OA and the results are shown in Fig. 6. The overall mass loss during pyrolysis was 39%. It can be seen from the first derivative curve that the biggest rate of change takes place before 100 °C when water is being eliminated. After that the changes taking place are smaller.

### 3.2. In-doped aerogels as catalyst for Michael reaction

The Michael addition reaction is one of the most versatile methods for the formation of carbon–carbon bonds, and has been widely used to generate valuable building blocks in organic synthesis.

Herein, we report our preliminary results by testing In-doped aerogels as catalysts in the Michael addition of ethyl 2-oxyoctocyclopentanocarboxylate 1 to 3-buten-2-one 2 (Scheme 1).

The conversions of the Michael donor 1 catalyzed by organic and carbon aerogels are shown in Table 2. In-doped OAs were active catalysts in the Michael reaction with conversions of the starting keto ester ranging from 82 to 97% in 24 h.

La-, Pr- and Nd-doped carbon aerogels also exhibited activity in the Michael reaction. In case of the Ce-doped CAs the formation of several by-products was observed. The activity of CAs was not as high as that of OAs. This is due to the structural and chemical changes that take place during carbonization. After pyrolysis all lanthanide species might not be available for the catalytic reaction anymore and the lanthanide ions previously bound to the ion-exchange center might have formed new bonds.

In a previous study the formation of Eu mixed oxides during pyrolysis, which proved to not catalyze the reaction, was confirmed [19]. However, in our case the presence of oxides could not be confirmed by XRD analysis. This could be the reason for our CAs being catalytically active, since it is known that metal oxides are mostly inert as catalysts in Michael addition [19].

The results obtained with OA-s and CA-s as catalysts in the Michael addition are preliminary. Clearly, the experimental procedure used for determining the conversions needs further development, particularly the method for obtaining a small homogeneous amount of catalyst.
4. Conclusions

A method for preparing Ln-doped (Ln = La, Ce, Nd, Pr) organic and carbon aerogels with catalytic activity in the Michael reaction was developed.

Using 5-methylresorcinol, 2,6-dihydroxy-4-methylbenzoic acid and formaldehyde to produce gels allows doping of aerogels with Ln species (Ln = Ce, La, Nd, Pr) through the ion-exchange method.

The mostly mesoporous materials with pore size diameters around 5-15 nm contained up to 12 wt% of metal in organic aerogels and 22 wt% in carbon aerogels. The specific surface area and pore volume of organic aerogels were higher than those of carbon aerogels. The materials were amorphous and no crystal structures were formed during pyrolysis.

In the Michael addition Ln-doped aerogels acted as catalysts. Organic aerogels with different lanthanides catalyze the Michael reaction efficiently. For carbon aerogels the conversion was modest (Pr) to high (Nd).

Acknowledgments

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References


[9] For reviews on lanthanides as catalysts see:


[18] For references on lanthanide grafting on silica aerogels see:


Publication III

Cobalt-Containing Nitrogen-Doped Carbon Aerogels as Efficient Electrocatalysts for the Oxygen Reduction Reaction


The electrocatalysis of the oxygen reduction reaction (ORR) on cobalt-containing nitrogen-doped carbon aerogels (CAs) was studied in alkaline solution. CA-based catalyst materials with varied compositions were prepared through the sol-gel polymerisation of organic precursors (resorcinol derivatives and melamine), followed by insertion of Co by using an ion-exchange process and carbonisation. The concentrations of the precursors had a large effect on the structure and physicochemical properties of the materials, as characterised by using SEM, XRD, XPS, atomic adsorption spectroscopy, and N2-adsorption analysis. The electrocatalytic activity of Co-containing N-doped CAs for the ORR was higher than that of nitrogen-free CAs, and this activity increased with increasing Co content. The most active catalyst materials supported the four-electron reduction of O2 and short-term stability tests indicated their high durability. Co-containing N-doped CAs can be regarded as a promising class of material for the cathode catalysts of alkaline membrane fuel cells.

1. Introduction

Currently, the development of efficient and stable, non-precious-metal catalysts for the oxygen reduction reaction (ORR) is one of the most important issues in optimising the cost and performance of low-temperature fuel cells.[1-8] A promising class of catalysts is transition-metal-containing nitrogen-doped carbon materials (MN/C catalysts) that can be prepared through the pyrolysis of a carbon support in the presence of a nitrogen precursor and a transition-metal (usually Co or Fe) salt.[1,8] Since early studies for the ORR on this type of pyrolysed material by Yager and co-workers in 1989,[9] the electrocatalytic activity of MN/C catalysts has been significantly improved and it currently approaches that of commercial Pt/C catalysts in acidic solution.[10] In alkaline solutions, their activity is even higher and, in some cases, can surpass that of the Pt-based catalysts.[5-7] Still, further enhancement of the volumetric activity of MN/C catalysts remains a great challenge that can be approached by optimising the catalyst structure, increasing the density of the active sites and average active site turnover frequency (TOF).[11] However, rational design of MN/C catalysts is impeded by uncertainties in the exact structures of the active sites for the ORR, the complexity of the reaction mechanism and the uncontrollability of the synthesis processes.[11]

MN/C catalysts can be synthesised from various carbon nanomaterials, such as carbon black, carbon nanotubes, graphene, and so forth.[9] Alternatively, porous N-doped carbon structures can be obtained through the carbonisation of nitrogen-containing organic polymers. For instance, carbon aerogels (CAs) are often prepared through pyrolysis of organic aerogels[9] that can be synthesised by using various methods; the most studied process involves the synthesis of resorcinol-formaldehyde gel by using the sol-gel method and subsequent supercritical drying.[10] To prepare nitrogen-doped CAs or aerogels, pyrolysis in a NH3 atmosphere[10,11] or nitrogen-containing precursors for gel preparation[12-15] can be used. High surface area, low mass density as well as high electrical conductivity and chemical stability make CAs good electrode materials for low-temperature fuel cells, among other applications.[8,16]

Oxygen reduction has been studied on various catalysts based on N-doped CAs or aerogels. For instance, nitrogen doping has been shown to increase the electrocatalytic activity for the ORR with CAs prepared from sustainable precursors through hydrothermal carbonisation.[17-19] However, the ORR activity of these materials in alkaline solution is still considerably lower than that of the Pt/C catalyst.[17-19] Recently, activities similar to that of Pt/C have been reached on metal-free N-doped CAs prepared from ionic liquids.[20] CAs synthesised in the presence of transition metals have shown moderate activities for the ORR in acidic solution,[10,11,12,14,21,22] but, in alkaline solution, activities comparable to that of commercial Pt/C have been reached.[11,22] High ORR activities were also observed on N-doped CAs containing cobalt-oxide nanoparticles that were prepared from commercial melamine sponge.[23] Similarly, high

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activities for different N-doped carbon materials containing nanoparticles of cobalt oxides\textsuperscript{[24-28]} or metallic cobalt\textsuperscript{[31-34]} have been observed in several other studies, and a synergetic effect of N-doped carbon and cobalt or cobalt-oxide nanoparticles on the electrocatalytic activity for ORR has been proposed\textsuperscript{[24-27, 29-31]}. 

In this study, N-doped CAs were prepared through the pyrolysis of organic aerogels synthesised from 3-methylresorcinol (MR), which is a locally available by-product from the oil-shale industry, and its derivative, 2,6-dihydroxy-4-methylbenzoic acid (dHMBB).\textsuperscript{[35]} Melamine (Mel) was used as a nitrogen-containing precursor and the ion-exchange method was employed for metal doping.\textsuperscript{[36, 37]} Considering that Co- and Fe-containing N-doped CAs previously prepared by our group showed high activity for the ORR,\textsuperscript{[38]} we were interested in exploring how the concentrations of the organic precursors affected the physical properties and the ORR activities of the resulting CAs.

2. Results and Discussion

2.1. Physicochemical Characterisation of CA-based Catalysts

The composition of CAs was varied during the preparation process, using different amounts of melamine, MR and dHMBB to study the effects on the properties of CAs. Table 1 shows the molar percentages of monomers that were used during the preparation of different samples and the measured elemental compositions. Co-containing N-doped CAs prepared by using different amounts of monomers are denoted Co-NCA1, Co-NCA2, and so forth (Table 1). For comparison, nitrogen-free Co-containing CA (Co-CA) and a cobalt-free material (NCA) were also prepared; in the latter case, the ion-exchange procedure was not performed.

The molar percentage (mol%) of dHMBB was varied from 10 to 25. It can be noted that, with the increase in the amount of dHMBB in the composition of the aerogel, the Co content also increased (Table 1). This is attributed to dHMBB having the ion-exchange moiety (the carboxylic group) that binds Co\textsuperscript{2+} during immersion in cobalt-nitrate solution. As previously concluded, the higher amount of dHMBB in the sample increased the weight percent of metal in the aerogel.\textsuperscript{[39]}

The melamine content was also varied during gel preparation. The nitrogen content in cobalt-containing samples was changed from 1.6 to 8.4 wt% by increasing the amount of melamine (Table 1). The carbon content in the samples was also affected by the increasing amount of melamine, decreasing from 87 to 69 wt\% by increasing the amount of melamine.

Brunauer–Emmett–Teller (BET) analysis was used to determine the specific surface areas and pore volumes for the CAs (Table 1). Samples Co-NCA1 to Co-NCA4 had high BET surface areas ($S_{\text{BET}} = 314–461 \text{ m}^2\text{g}^{-1}$) and total pore volumes ($V_{\text{TOT}}$) from 228 to 399 mm\textsuperscript{3}g\textsuperscript{-1}. Figure 1 shows the pore-size distribution of sample Co-NCA2, indicating that this material has mostly pores with the diameters ($D_{p}$) of 2 and 4 nm.

Co-NCA5 had significantly lower $S_{\text{BET}}$ and $V_{\text{TOT}}$ values. During preparation, Co-NCA5 had the highest melamine content and no MR was added. The $S_{\text{BET}}$ and $V_{\text{TOT}}$ values for the organic aerogel precursor of Co-NCA5 were also measured and the respective results were 868 m\textsuperscript{2}g\textsuperscript{-1} and 1490 mm\textsuperscript{3}g\textsuperscript{-1}. This shows that a drastic reduction in specific surface area and pore volume occurred during pyrolysis.

Thermogravimetric analysis (TGA) results in Figure 2 show the mass-loss curves of organic aerogels (precursors of Co-NCA1–Co-NCA5) during pyrolysis in an Ar atmosphere. The mass loss was between 56 and 80%, with the highest being for Co-NCA1. Samples that were prepared by using higher melamine contents (Co-NCA3 to Co-NCA5) had similar mass losses (60–65 wt%). The mass-loss curve is smooth in shape, without abrupt changes for Co-NCA1 and Co-NCA2. However, for Co-

\begin{table}[h]
\centering
\caption{Elemental composition, N$_2$-adsorption analysis results and ORR mass activity for CAs.}
\begin{tabular}{|c|c|c|c|c|c|c|c|c|c|}
\hline
Sample & Monomer [mol\%] & Elemental composition [wt\%] & $S_{\text{BET}}$ [m\textsuperscript{2}g\textsuperscript{-1}] & $V_{\text{TOT}}$ [mm\textsuperscript{3}g\textsuperscript{-1}] & $V_{\text{PC}}$ [mm\textsuperscript{3}g\textsuperscript{-1}] & MA$^{[4]}$ [	extmu A g\textsuperscript{-1}] \\
\hline
 & Mel & MR & dHMBB & Co & N & C & H & & \\
\hline
Co-NCA1 & 30 & 60 & 10 & 2.0 & 1.6 & 87.0 & 1.3 & 370 & 330 & 85 & 2.8 \\
Co-NCA2 & 30 & 45 & 25 & 4.3 & 2.5 & 82.2 & 1.0 & 314 & 230 & 70 & 3.2 \\
Co-NCA3 & 40 & 40 & 20 & 4.1 & 2.9 & 80.2 & 1.4 & 461 & 399 & 95 & 3.4 \\
Co-NCA4 & 60 & 15 & 25 & 5.3 & 2.6 & 81.0 & 0.7 & 316 & 228 & 54 & 14.4 \\
Co-NCA5 & 75 & 0 & 25 & 5.6 & 8.4 & 68.9 & 1.2 & 118 & 93 & 1.5 & 12.5 \\
Co-CA$^{[35]}$ & 0 & 75 & 25 & 2.6 & 0.03 & n.d & n.d & n.d & n.d & 2.3 \\
NCA$^{[37]}$ & 60 & 15 & 25 & 0 & 14.3 & 65.2 & 1.3 & n.d & n.d & 0.23 \\
\hline
\end{tabular}
\footnotesize{(a) At –0.2 V. (b) n.d. = not determined.}
\end{table}

\begin{figure}[h]
\centering
\includegraphics[width=\textwidth]{figure1.png}
\caption{Pore-size distribution analysis results for Co-NCA2.}
\end{figure}
NCA4 and Co-NCA5, there is a steep change at 340–370 °C. Melamine resins have also shown large mass loss, owing to the decomposition and evaporation of the products generated in that temperature range. The decomposition of melamine and, therefore, the structure of the material could explain the smaller surface area and porosity of CAs prepared with higher melamine contents.

The surface morphologies of the CAs were evaluated by using scanning electron microscopy (SEM, Figure 3). The morphology of the CA prepared with the highest MR content (Co-NCA1) is similar to that of resorcinol/formaldehyde CAs with a grany structure composed of linked spheres (Figure 3 a and 3 b). The Co-NCA2 aerogel was prepared by using more melamine and less MR than for Co-NCA1. In the structure of the Co-NCA2 (Figure 3 d), separate particles forming the aerogel network are not visible and the morphology is different from that of the resorcinol/formaldehyde CAs. This is also the case in samples prepared with higher melamine contents (Co-NCA4 and Co-NCA5), where the surfaces of the aerogels appear to be composed of particles merged together. This shows that the addition of resorcinol derivatives is necessary to keep the bead-like structure that is associated with resorcinol/formaldehyde CAs. The structure of melamine-rich aerogels will be affected by the sudden mass loss and decomposition of melamine during pyrolysis.

Figure 3 c shows an SEM image of Co-NCA1 taken by using backscattered electron (BSE) imaging, where brighter spots can be assigned to Co particles. These are evenly distributed in the sample and have sizes ranging from 5 to 40 nm (Figures 3 a–c). The cobalt nanoparticles in Co-NCA3 are also evenly distributed (cf. Figure 3 g) and the sizes are larger (10–100 nm) than in Co-NCA1 (Figures 3 e and 3 f). This is probably caused by the increased amount of metal in the aerogel.[36, 37] Samples Co-NCA4 and Co-NCA5 have smaller nanoparticles (10–40 nm) than Co-NCA3, and SEM images show the formation of long cylindrical structures with a diameter of about 20 nm, which appear to be carbon nanotubes (CNTs) or nanoribbons (Figures 3 h, 3 j and 3 k). The BSE image (Figure 3 j) shows brighter spots (Co nanoparticles) at the end of the long structures. Formation of CNTs during pyrolysis in the presence of cobalt is a well-known process, and cobalt nanoparticles are used as a catalyst in the synthesis of CNTs when using chemical vapour deposition.[46] Cobalt has also been shown to induce the formation of nanoribbons in CAs.[41] From the nitrogen-containing precursors, N-doped CNTs can be synthesised. For instance, CNTs have been prepared by using chemical vapour deposition with melamine and iron precursors.[42] In addition, the formation of nitrogen-doped CNTs through the pyrolysis of melamine-formaldehyde resin in the presence of iron has recently been demonstrated.[43] Therefore, it can be assumed that long carbon nanostructures in Co-NCA4 and Co-NCA5 are also nitrogen-doped CNTs.

Figure 4 shows X-ray diffraction (XRD) peaks for crystalline Co in all samples, which is an expected result under these pyrolysis conditions.[37] A small peak attributed to Co3O4 indicates oxidation of the surface of Co nanoparticles. Formation of graphitic structures is clearly visible in samples Co-NCA1 and Co-NCA2. In Co-NCA5, a peak at the same location cannot be identified. This shows that the increasing melamine content inhibits the formation of graphitic structures in this case. In the diffraction patterns shown in Figure 4, there are a few other small peaks visible at various degrees. The identification of these peaks is problematic, owing to the limited amount of data. It is likely that various cobalt–carbon, cobalt–nitrogen or nitrogen–carbon structures have also formed in the CAs.

X-ray photoelectron spectroscopy (XPS) was used to analyse the surface composition of the catalysts. High-resolution XPS spectra for Co-NCA4 and NCA are presented in Figure 5. The N 1s peaks were deconvoluted into four symmetrical peaks, corresponding to different nitrogen species: pyridinic (398.1 eV), pyrrolic (399.9 eV), graphitic (401.0 eV) and pyridine-N-oxide (403.3 eV).[38] Similar nitrogen species are typically found in Mn/N/C catalysts.[21] It can be seen that, in both catalyst materials, the largest peak belongs to pyridinic nitrogen. Pyridinic and graphitic nitrogen are considered to be the most active centres for oxygen electroreduction.[44] It is also evident that the relative amount of pyrrolic and graphitic N is larger for Co-NCA4. It has been shown earlier that the presence of Co increases the percentage of graphitic N centres[44] that play an important role in ORR electrocatalysis.[46] Analysis of the Co 2p peaks suggested that Co may be present as Co(OH)2 on the surface. However, the interpretation of XPS spectra of transition metals is complicated, because of the complexity of their 2p spectra (peak asymmetries, complex multiplet splitting, uncertain and overlapping binding energies, etc.).[46] Therefore, it is also possible that combinations of other oxides, hydroxides and N-coordinated metal ions might result in similar spectra.

### 2.2. Oxygen Reduction on CA-based Catalysts

The electrocatalytic behaviours of various CAs for the ORR were studied by using the rotating disc electrode (RDE) method in 0.1 M KOH solution. A set of RDE polarisation curves for Co-NCA4 in Figure 6 shows that this material is a rather active catalyst for the ORR, as, from the onset potential.
about \(-0.17\) V, the current increases sharply and reaches its diffusion-limited value at approximately \(-0.3\) V.

A comparison of RDE voltammetry curves of O\(_2\) reduction for Co-containing N-doped CAs of different compositions is presented in Figure 7a. It appears that two materials, Co-NCA4 and Co-NCA5, are most active; the half-wave potentials (\(E_{1/2}\)) for these are \(-0.210\) and \(-0.219\) V, respectively, which are only about 20–30 mV more negative than that of the commercial Pt/C catalyst. The other three materials are slightly less active, the \(E_{1/2}\) values being in the range between \(-0.258\) and \(-0.270\) V. The activities of Co-NCA4 and Co-NCA5 are rather similar to that of Co-containing N-doped CAs prepared from commercial melamine sponge\(^{22}\) and slightly lower than that of Fe-containing N-doped CAs synthesised from ionic liquids,\(^{22}\) but considerably higher compared to some metal-free CAs.\(^{16,19}\)

To analyse the RDE data, the Koutecky-Levich equation [Eq. (1)] was used:\(^{24}\)

\[
\frac{1}{j} = \frac{1}{j_k} + \frac{1}{j_d} = \frac{1}{nFkC_{O_2}} + \frac{1}{0.62nF^{1/2}D^{1/2}_{O_2}}V^{1/2}kC_{O_2}^{1/2} \omega^{1/2}
\]  

(1)

where \(j\) is the measured current density, \(j_k\) and \(j_d\) are kinetic and diffusion-limited current densities, respectively, \(n\) is the number of electrons transferred per O\(_2\) molecule, \(k\) is the rate constant for O\(_2\) reduction, \(F\) is the Faraday constant (96,485 C mol\(^{-1}\)), \(\omega\) is the electrode rotation rate (rad s\(^{-1}\)), \(C_{O_2}\) is...
Figure 4. XRD analysis results for the CA materials.

the concentration of oxygen in the bulk \((1.2 \times 10^{-4} \text{ mol cm}^{-3})^{[148]}\) \(D_o\) is the diffusion coefficient of oxygen \((1.9 \times 10^{-5} \text{ cm}^2 \text{s}^{-1})^{[148]}\) and \(\nu\) is the kinematic viscosity of the solution \((0.01 \text{ cm}^2 \text{s}^{-1})^{[148]}\) From the slopes of the Koutecky–Levich plots, the number of electrons transferred per \(O_2\) molecule \(n\) can be calculated by using Equation (1). Figure 7b shows the Koutecky–Levich plots and the values of \(n\) for Co-containing N-doped CAs. It is evident that the ORR yields water as the main product at negative potentials for these materials, as \(3.9 < n < 4.0\), and only Co-NCA2 shows slightly lower \(n\) values. At more positive potentials, the value of \(n\) gradually decreases until \(3.3 < n < 3.7\) at \(-0.5\) V, as a result of increasing production of peroxide. Peroxide production is lower on the most active catalysts, for instance, the peroxide yield on Co-NCA4 is about 15% at low overpotentials and it decreases to zero at \(-1\) V. Low levels of peroxide production on metal-containing N-doped CAs have been observed in earlier studies\(^{[18,22,23]}\) and it is an important pre-requisite for the application of these materials as cathode catalysts on low-temperature fuel cells.\(^{[90]}\)

For a more direct comparison of the electrocatalytic activities, the ORR mass activities (MAs) of the CAs were calculated by using Equation (2):

\[
MA = i_k/m
\]  

(2)

Figure 5. XPS spectra in the N 1s (a, b) and Co 2p (c) regions for Co-NCA4 (a, c) and NCA (b) catalysts.
where \( i_0 \) is the kinetic current at a certain potential and \( m \) is the mass of the catalyst on the electrode surface (Table 1). The values of MA at \(-0.2\) V are highest for Co-NCA4 and Co-NCA5 and about four times lower for the other three materials. According to the results of elemental analysis, these two most active materials also have higher cobalt contents. Increasing ORR activity with increasing Co content has been observed for Co\(_3\)O\(_4\)-containing N-doped graphene, and it has been suggested that one of the active reaction sites in these hybrid materials could be Co-oxide nanoparticles.\(^{24}\) The total nitrogen content is highest for Co-NCA5, but much lower for Co-NCA4, although the activities are similar. This suggests that only a small fraction of nitrogen is composing the active centres for the ORR and the remaining part is either in a passive form or inaccessible to \( O_2 \) molecules. According to the literature, only certain types of nitrogen centres are considered to be involved in ORR electrocatalysis,\(^{21}\) and the activity of the catalysts does not directly depend on the total nitrogen content.\(^{27,40,11}\) The surface area and pore volume do not seem to have a large effect on the ORR activity of the materials, because Co-NCA5, which has a low surface area and porosity, has a similar activity to Co-NCA4, which has much higher \( S_{\text{ BET}} \) and \( V_{\text{ BET}} \) values. However, the nanotubes or nanoribbons observed for Co-NCA4 and Co-NCA5 may contribute to the higher activity of these materials. Both the metal-free\(^{32}\) and transition-metal-containing\(^{32,33}\) N-doped CNTs have been shown to be very active catalysts for the ORR in alkaline solutions. This may arise from improved electrical conductivity of metal/CNT hybrid materials.\(^{24}\) In addition, embedded transition-metal nanoparticles in N-doped CNTs can lower the local work function of the carbon surface and, thereby, enhance the ORR activity.\(^{34}\) Subsurface iron or iron carbide have been suggested to increase the ORR activity of nitride-modified carbon fibres through the stabilisation of the peroxide intermediate.\(^{7}\)

For comparison, \( O_2 \) reduction was also studied on cobalt- and nitrogen-free CAs (Figure 8a). In accordance with the earli-

![Figure 7. RDE voltammetry curves of oxygen reduction (a) and dependence of the number of electrons transferred per \( O_2 \) molecule on potential (b) for Co-containing N-doped CAs in \( O_2 \)-saturated 0.1 \( M \) KOH; \( \gamma = 10 \) mV s\(^{-1} \), \( \omega = 1900 \) rpm. Inset shows the Rouseby-Levich plots at \(-0.6 \) V.](image1)

![Figure 8. RDE voltammetry curves of oxygen reduction (a) and dependence of the number of electrons transferred per \( O_2 \) molecule on potential (b) for CA-based catalysts, glassy carbon and Pt/C catalyst in \( O_2 \)-saturated 0.1 \( M \) KOH; \( \gamma = 10 \) mV s\(^{-1} \), \( \omega = 1900 \) rpm. Inset shows the Rouseby-Levich plots at \(-0.6 \) V.](image2)
Higher activities and lower peroxide yields of metal-containing N-doped carbon catalysts, as compared to the metal-free materials, have been observed in several studies,\(^\text{16,35}\) and it has been proposed that the peroxide intermediate is further reduced to OH\(^-\) on metal–nitrogen coordinated centres in alkaline solution.\(^\text{34}\) Interestingly, the Co-containing CA prepared from precursors not containing nitrogen (Co-CA) showed much higher activity for the ORR compared to NCA, but it was lower than that of Co-NCA4 (see Figure 8a and Table 1). Cobalt and cobalt-oxide nanoparticles have been suggested to play an important role in ORR electrocatalysis in alkaline solution. For instance, the N-doped graphene and CoO\(_\text{x}\) nanoparticles supported on undoped graphene showed rather high activities towards the ORR, but the activity was further increased for Co\(_{x}\)O\(_y\)/N-doped graphene, suggesting a synergetic effect between cobalt-oxide nanoparticles and N-containing centres.\(^\text{24,25}\) Similar results have been obtained on CoO-containing N-doped reduced graphene oxide (rGO).\(^\text{26,27}\) In addition, the activity of Co\(_{x}\)O\(_y\)/N-doped mesoporous graphene decreased with decreasing Co\(_{x}\)O\(_y\) loading in the catalyst.\(^\text{28}\) Therefore, the highest activities observed for Co-NCA4 and Co-NCAS can also be attributed to the higher Co content in the catalysts.

There are several explanations in the literature for the synergetic effect of cobalt or cobalt-oxide nanoparticles and N-doped carbon on the ORR kinetics. For instance, it has been attributed to the increase in bonding strength between carbon and the catalyst, owing to nitrogen doping.\(^\text{24,29}\) A dual-site mechanism of the ORR in alkaline solution has been proposed for N-doped carbon catalysts derived from cobalt precursors and polypyrrole, according to which the O\(_2\) molecule is first adsorbed on the N-doped site and then converted to HO\(_2\)\(^-\), which is further reduced on oxide-coated Co nanoparticles.\(^\text{30}\) Alternatively, based on the results obtained on a graphene-based catalyst, it has been suggested that the N-doped graphene domains adjacent to the Co nanoparticles are the main active sites for the ORR, whereas the Co–N–C structures at the interface of Co nanoparticles and N-doped graphene help to activate O\(_2\) molecules for better ORR activity.\(^\text{31}\) Density functional theory (DFT) calculations have indicated that, in N-doped rGO-based catalysts, a stable rGO(N)–Co\(_{x}\)O–Co\(_{x}\)–rGO(N) structure is most likely the active species for the ORR.\(^\text{32}\)

The results obtained in this study also indicate that the most active electrocatalysts contain both cobalt and nitrogen precursors; however, a rather facile 4e\(^-\) reduction of O\(_2\) on nitrogen-free material implies that the main active sites may be cobalt and/or cobalt-oxide nanoparticles. It is also evident that the Co content is higher for the nitrogen-containing CAs compared to Co-CA (Table 1). It has been demonstrated that, in the synthesis of Co-containing N-doped rGO, the presence of nitrogen helps to increase the amount of cobalt.\(^\text{37}\) Although the metal-free nitrogen-containing materials have, in some cases, shown very high activity for the ORR in alkaline solution,\(^\text{31,52,58}\) the activity of Co-free N-doped CA was rather low and there was no obvious dependence of the ORR activity on the nitrogen content for the other CA-based catalysts.

The mass-transfer-corrected Tafel plots of O\(_2\) reduction on various catalysts were constructed on the basis of the RDE data (Figure 9). The Tafel slope values were in the range between -39 and -47 mV for all Co-containing N-doped materials at low overpotentials, which is consistent with the results obtained for N-doped graphene-containing CoO\(_x\) nanoparticles\(^\text{24,25}\) and suggests that protonation of adsorbed O\(_2\) may be the rate-limiting step.\(^\text{24}\) For Co-CA, the slope was -53 mV, which coincides well with the slope value for Co\(_{x}\)O\(_y\) nanoparticles supported on non-doped graphene.\(^\text{24}\) For the Pt/C catalyst, a slope value of -61 mV was found, as usually observed for Pt catalysts at low overpotentials in alkaline media.\(^\text{39}\) Cobalt-free material showed the highest Tafel slope value (-76 mV).

To evaluate the durability of the most active CA-based catalyst in electrochemical conditions, it was subjected to a short-term stability test (1000 potential cycles between 0 and -0.8 V at 100 mV s\(^{-1}\) in O\(_2\)-saturated 0.1 M KOH). A comparison of RDE results before and after potential cycling showed that this material is very stable, as the E\(_{1/2}\) value shifted by only about 2 mV in the negative direction, and the kinetic current at

![](Image) Figure 9. Mass-transfer-corrected Tafel plots for O\(_2\) reduction on CA-based catalysts in 0.1 M KOH; ω = 1900 rpm.

![](Image) Figure 10. RDE voltammetry curves of oxygen reduction on Co-NCA4 catalyst in O\(_2\)-saturated 0.1 M KOH before and after stability testing; γ = 10 mV s\(^{-1}\); ω = 1900 rpm. Inset shows the change in j\(_k\) values, calculated at -0.2 V during potential cycling.
--0.2 V decreased by about 8% (Figure 10). M/N/C catalysts generally show much higher durability than noble-metal-based catalysts in alkaline and acidic solutions, which can be explained by the incorporation of the active centres into the carbon framework. If degradation of the active centres occurs on the surface, some new centres are exposed to the solution and the ORR activity of the catalyst remains high. It has also been suggested that nitrogen doping of the carbon support increases the strength of the interaction between the cobalt-oxide nanoparticles and carbon, thereby preventing catalyst agglomeration and increasing the stability of the catalyst. Higher durability of N-doped CAs, as compared to Pt/C catalysts, has been observed in both acidic and alkaline solutions. In addition, similarly to the other M/N/C catalysts, these materials show high methanol tolerance and are, thus, promising catalysts for direct methanol fuel cells.

By changing the relative amount of precursors in the synthesis of organic aerogels, the chemical composition of resulting CAs can easily be varied. However, the morphological features of the CAs, most importantly the pore structure and size distribution, also depend on the composition of organic gels and influence the performance of the catalysts. It is important to further optimise the synthesis procedure, to prepare CA-based materials with high density of active sites for ORR and favourable pore structure, which would likely result in even more active electrocatalysts.

3. Conclusions

Cobalt-containing nitrogen-doped CAs were prepared from resorcinol derivatives and melamine, and an ion-exchange process was used to introduce metal into the aerogels. The results showed that, by varying the dhMBA or melamine content, the compositions of the CAs can be altered. CAs showed high surface area and had both meso- and microporosity. The increasing amount of melamine in the compositions of the aerogels decreased the surface areas and pore volumes, owing to the rapid change in mass taking place during pyrolysis, which was demonstrated by using TGA. The SEM images showed the structures of the CAs to be similar to those of resorcinol/formaldehyde aerogels for CAs prepared with low melamine contents, but the structure changed with increasing melamine content. XRD analysis revealed the presence of crystalline structures of cobalt and cobalt oxide and SEM images confirmed the presence of cobalt nanoparticles. In CAs prepared with higher melamine and cobalt contents, CNT structures had also formed.

The electrocatalytic activity of Co-containing N-doped CAs for the ORR was rather high and depended on the composition. The materials with the highest Co content showed the highest ORR activities, which can be associated with the coexistence of cobalt nanoparticles and N-doped carbon structures. The durability of these materials is high in alkaline solution and they mostly support the 4e⁻ pathway of O₂ reduction, thus showing great promise for applications as cathode catalysts in alkaline fuel cells.

**Experimental Section**

**Materials and Preparation of CAs**

5-Methylresorcinol (MR), with a reported purity of >99% was provided by VKG AS (Estonia) and 2,6-dihydroxy-4-methylenzoic acid (dhMBA), with a purity of >99%, was donated by Carboshale OÜ (Estonia). Melamine (Mel), with a purity of >99%, was purchased from Aldrich (USA). Anhydrous sodium carbonate, with a purity of 99.8%, was purchased from Riedel-de Haën (Germany). Formaldehyde (FA) (37 wt% solution in water) was purchased from Sigma-Aldrich. CO₂ (99.8%) was obtained from Eesti AGA AS (Estonia). Cobalt(II) nitrate hexahydrate (>98%) was purchased from Sigma-Aldrich (USA). The Milli-Q water system was used to obtain ultrapure (18.2 MΩ·cm) water.

A supercritical extraction system with a double-clamp autoclave (100 ml in volume) was constructed by NWA Analytische Meßgeräte GmbH (Germany). The pyrolysis oven, type MTF 12/38/400, with a maximum temperature of 1200°C was from Carbolite (United Kingdom).

The preparation of metal-doped CAs was based on the ion-exchange method. The amounts of substances (in grams) to be used for gel preparation were calculated by using Equation (3):

$$\text{substances} = \frac{\text{monomers} + \text{FA} + \text{Na}_2\text{CO}_3}{\text{water added} + \text{water from FA solution}} = 0.25$$  \hspace{1cm} \text{(3)}$$

The ratio of substances to water was always kept at 0.25. The amounts of the monomers (Mel, MR and dhMBA) used were varied during preparation from 0 to 75 mol% [Eq. (4)].

$$x\%\text{ (Mel)} + y\%\text{ (MR)} + z\%\text{ (dhMBA)} = 100\%\text{ (monomers)}$$  \hspace{1cm} \text{(4)}$$

The amount of sodium carbonate used was calculated by adding the amount needed for converting dhMBA into the sodium salt of dhMBA (molar ratio of dhMBA to Na₂CO₃ = 2) and an additional amount as catalyst (molar ratio of monomers to Na₂CO₃ = 100). The molar ratio of FA to melamine was 3.7 and that of FA to resorcinol derivatives (MR and dhMBA) was 2. At first, two separate solutions were prepared in two vials. In one vial, a mixture of Mel, FA solution and half of the water was prepared. The mixture was kept at 70°C for about 25 min to allow the Mel to dissolve. In the second vial, a mixture of MR, dhMBA, sodium carbonate and half of the water was prepared. Both homogeneous solutions were poured into one vial, mixed, and then poured into a glass test tubes and allowed to gel and cure for 4 days at 85°C.

To obtain Co²⁺-doped gels, the hydrogels were soaked in a 0.1 M solution of cobalt nitrate for 3 days (the solution was changed every 24 h), after which the sodium ions bound to the ion-exchange moiety (the carboxylic group) were exchanged for cobalt ions.

In order to use supercritical CO₂ drying for aerogel preparation, solvent exchange was carried out for acetone by changing the solution every 2 h, starting with a 25:75 mixture of acetone and water and proceeding with 50:50 and 75:25 mixtures and finally with 100% acetone for 2 days. A more detailed description of the method used for supercritical drying has been published elsewhere.
The CAs were prepared through pyrolysis in a N₂ atmosphere by heating at 10 °C/min -1 to 800 °C and maintaining at this temperature for 1 h. Co-containing N-doped aerogels prepared with different amounts of the monomers are denoted as Co-NCA1, Co-NCA2, and so forth (Table 1). For comparison, nitrogen-free Co-containing CA (Co-CA) and cobalt-free (NCA) materials were also prepared; in the latter case, the ion-exchange procedure was not performed.

Physicochemical Characterisation of Catalyst Materials
Elemental analyses were performed by the vario MICRO cube (Elementar, Germany) and the cobalt content was measured by using a Spectra AA 220F flame atomic absorption spectrometer (Varian, USA). SEM images were taken using a ZEISS ULTRA 55 (Germany) microscope. Nitrogen-adsorption analyses were performed by using a KELVIN 1042 Sorptometer built by Costech International (USA). Helium was used as a carrier gas and N₂ as the adsorbive gas. Nitrogen-adsorption data were taken at relative pressures (p/p⁰) from 0 to 1 and at liquid-nitrogen temperature. The specific surface area (S BET) was calculated according to the BET theory. The specific micropore volume (V mic) was determined by using the t-plot, whereas the pore-size distributions were determined by using the Barrett–Joyner–Halenda (BJH) method from the desorption branch. The XRD pattern was recorded with a Rigaku Ultima IV diffractometer by using a D/Max Ultra IId detector. CuKα radiation was used, and a Ni filter to remove Kβ radiation. The recorded diffraction patterns were analysed by using Rigaku HIGHScore. TGA tests were conducted by using Labsys Evo TG-DTA 1600 (Setaram, France) and heating the samples in an Ar atmosphere at 10 °C/min -1 to 800 °C. The XPS measurements were carried out with a SCIENTA SES-100 spectrometer by using a non-monochromated MgKα X-ray source (incident energy = 1253.6 eV), a take-off angle of 90 ° and a source power of 300 W. The pressure in the analysis chamber was below 10⁻⁹ Torr. Raw data were processed by using CasaXPS software, and data processing involved removal of k=0 and k=1 satellites, removal of the background and fitting of the components. Background removal was performed by using a Tougaard background; for fitting components, the Gauss–Lorenz hybrid function was used (GL 30, Gauss 70%, Lorenz 30%) for the best fit.

Determination of Electrocatalytic Activity Towards the ORR
Glassy carbon (GC) discs (GC20-5S, Tokai Carbon) with a geometric area (A) of 0.2 cm² were used as electrode substrates. The GC electrodes were polished to a mirror finish with 1 and 0.3 µm alumina slurries (Buehler) and sonicated in Milli-Q water for 5 min. The CAs were ball-milled to fine powders and the catalyst ink was prepared by mixing the catalyst (4 mg), Nafion solution (40 µL; 5 wt% solution in lower alcohols, Aldrich) and 2-propanol (960 µL). The ink was subjected to ultrasonication for at least 40 min and the GC electrode was drop-coated twice with 5 µL aliquots of the catalyst, resulting in a loading of 200 µg cm⁻², which has been determined as the optimal loading for this type of material. For the 20 wt% Pt catalyst supported on Vulcan carbon XC-72 (ETEK), the same ink formulation and coating procedure were used, resulting in a loading of 40 µg cm⁻² of Pt on the electrode. The catalyst-coated electrodes were dried in an oven at 65 °C for at least 30 min before measurements.

The electrocatalytic activities of the CAs were evaluated by using the RDE method, using an ED1101 rotator and a CTV110 speed control unit (Radiometer). The potential was applied by using an Autolab potentiostat/galvanostat PGSTAT30 (Eco Chemie B.V., The Netherlands). Experiments were conducted in a three-electrode glass cell at room temperature (23 ± 1 °C). A Pt wire served as the counter electrode and a saturated calomel electrode (SCE) was used as the reference; all potentials were referred to this electrode. Electrochemical experiments were carried out in 0.1 M KOH solution, prepared from KOH pellets (p.a. quality, Sigma–Aldrich) and saturated with Ar (99.9999%, AGA) or O₂ (99.999%, AGA). A continuous flow of gas over the solution was maintained during the electrochemical measurements. The RDE polarisation curves at various electrode rotation rates (Ω) between 360 and 3100 rpm were obtained by scanning the potential from 0 to −1.2 V at a scan rate (v) of 10 mV s⁻¹. For data presentation and analysis, the background current registered in O₂-free solution was subtracted from the experimental current and current densities were calculated per geometric area of the electrode (A).

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Keywords: carbon aerogels - electrocatalysis - nitrogen doping - non-precious metal catalysts - oxygen reduction

# Curriculum vitae

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3. **Language competence/skills**

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5. Professional employment

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6. Research activity

01.03.2015-30.06.2015 University of Lisbon (Portugal), the group of Carlos Nieto de Castro. Working with ionanofluids and thermal conductivity measurements.
## Elulookirjeldus

1. **Isikuandmed**

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<td>Sünniaeg ja -koht</td>
<td>02.08.1988, Tallinn, Eesti</td>
</tr>
<tr>
<td>E-posti aadress</td>
<td><a href="mailto:kristiina.kreek@ttu.ee">kristiina.kreek@ttu.ee</a></td>
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2. **Hariduskäik**

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<thead>
<tr>
<th>Õppeasutus</th>
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<tr>
<td>Tallinna Reaalkool</td>
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<td>2010</td>
<td>Rakenduskeemia ja biotehnoloogia/bakalaureusekraad</td>
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<td>Tallinna Tehnikaülikool</td>
<td>2012</td>
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3. **Keelteoskus**

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4. **Täiendusõpe**

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<td>2012-2014</td>
<td>Doktorikool „Funktsionaalsed materjalid ja tehnoloogiad“, Tallinna Tehnikaülikool, Eesti</td>
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<td>2013-2015</td>
<td>POKE training school, Sustainable Chemistry and Process Technology in the Northern Baltic Sea Region</td>
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<td>25.01.2016-29.01.2016</td>
<td>MICACT doktorikool 2: modelleerimine ja ioonsed elektroaktiivsed polümeerid, Tartu Ülikool</td>
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5. Teenistuskäik

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<td>2012-2015</td>
<td>Tallinna Tehnikaülikool</td>
<td>õppetöö läbiviimine analüütilise keemia alal</td>
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6. Teadustegevus

01.03.2015-30.06.2015 Lissaboni Ülikool (Portugal), Carlos Nieto de Castro rühm. Ionanofluidumite uurimine and soojusjuhtivuse mõõtmised.
DISSERTATIONS DEFENDED AT TALLINN UNIVERSITY OF TECHNOLOGY ON
NATURAL AND EXACT SCIENCES

5. **Eola Valdre**. Endothelial-Specific Regulation of Vessel Formation: Role of Receptor Tyrosine Kinases. 2000.
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<th>Page</th>
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<tr>
<td>100</td>
<td>Priit Pruunsild</td>
<td>Neuronal Activity-Dependent Transcription Factors and Regulation of Human BDNF Gene.</td>
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<td>101</td>
<td>Tatjana Knjazeva</td>
<td>New Approaches in Capillary Electrophoresis for Separation and Study of Proteins.</td>
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<td>Atanas Katerski</td>
<td>Chemical Composition of Sprayed Copper Indium Disulfide Films for Nanostructured Solar Cells.</td>
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<td>103</td>
<td>Kristi Timmo</td>
<td>Formation of Properties of CuInSe$_2$ and Cu$_2$ZnSn(S,Se)$_4$ Monograin Powders Synthesized in Molten KI.</td>
<td>2011</td>
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<td>104</td>
<td>Kert Tamm</td>
<td>Wave Propagation and Interaction in Mindlin-Type Microstructured Solids: Numerical Simulation.</td>
<td>2011</td>
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<td>Adrian Popp</td>
<td>Ordovician Proetid Trilobites in Baltoscandia and Germany.</td>
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<td>Sea Ice Deformation Events in the Gulf of Finland and This Impact on Shipping.</td>
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<td>Kairit Zovo</td>
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<td>Evaluation of the Biological Effects of Engineered Nanoparticles on Unicellular Pro- and Eukaryotic Organisms.</td>
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<td>117</td>
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<td>Dispersion Analysis of Wave Motion in Microstructured Solids.</td>
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<td>Liina Kaupmees</td>
<td>Selenization of Molybdenum as Contact Material in Solar Cells.</td>
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<td>Development of Methods for CE Analysis of Plant Phenolics and Vitamins.</td>
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<td>124</td>
<td>Ardo Illaste</td>
<td>Analysis of Molecular Movements in Cardiac Myocytes.</td>
<td>2012</td>
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125. **Indrek Reile.** 3-Alkylcyclopentane-1,2-Diones in Asymmetric Oxidation and Alkylation Reactions. 2012.

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