Recovery of rare-earth elements from NdFeB magnets by zirconium phosphate ion exchangers

Junhua Xu

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I would like to dedicate this thesis for the memory of my father Mr. Qinhai Xu and my supervisor Honorary Professor Risto Harjula.
Abstract

The societal transformation from fossil fuel-based energy sources to ecologically friendly energy sources has sparked the development and utilization of electric (and hybrid) vehicles and electric generators for wind turbines, among others. Permanent magnets are essential components of these technologies.

Over the years, the production of NdFeB permanent magnets has surpassed all other kinds because of their low cost and improved magnetic properties. The rare-earth elements (REEs) Nd and Dy are critical for the production of these magnets, and they come with a significant supply risk. Also since REEs exist simultaneously in minerals, the balance problem has become increasingly evident; Nd and Dy are produced at the cost of overproduction and stockpiling of other REEs. Due to their limited life span, more and more end-of-life (EOL) NdFeB magnets have accumulated as scrap. Recycling Nd and Dy from EOL NdFeB magnets could be a more ecological means to reduce supply chain pressure and to partially solve the balance problem.

The purpose of this thesis is to develop new ion exchangers based on zirconium phosphate (ZrP) for selective recovery of Co, Nd, and Dy from EOL magnets. In general, inorganic ion exchangers, such as ZrPs, are more selective than organic resins because of the ion-sieve functionality originated from rigid structures. Two inorganic ion exchangers, crystalline alpha zirconium phosphate (α-ZrP) and amorphous ZrP (am-ZrP) and one inorganic (am-ZrP)-organic (PAN) ion exchange composite material were synthesized and characterized for their ion exchange properties in this study.

The α-ZrP was synthesized with a lower energy and acid consumption. The ion-exchange capacity from the titration result was 6.65 meq/g. Co was taken up minimally from the Co-Nd-Dy ternary solution in acidic solution (pH 1-3) when compared with the total uptake amount. The am-ZrP was synthesized by using an easy scalability synthesis method at the room temperature. The molecular formula Zr(H2PO4)0.17 (HPO4)1.78 (PO4)0.09 • 0.96H2O was calculated from the results of digestion experiment, 31P NMR, and TG analysis. The molecular formula suggested that the theoretical ion-exchange capacity of am-ZrP was 6.97 meq/g. The column elution study of am-ZrP utilized a stepwise gradient elution; Almost complete metal separation was achieved from a mixed 1.0 mM equimolar solution. These promising results encouraged us to apply am-ZrP to a larger lab-scale study.
To solve the possible operation problems in scale-up column separation, an am-ZrP/polyacrylonitrile composite was synthesized as bead form. X-ray tomography demonstrated a good spatial distribution of ion-exchange active component am-ZrP in the polymer matrix. Column-optimized experiments for the synthesized composite were performed by altering running temperature, speed, and concentration of the elution agent (HNO₃) as well as feed concentration and loading degree. When the column was run at lower speed and at higher temperature, the purity of metal fractions in the effluent was highly enhanced relative to the feed. Gradient elution at 50°C was adopted for metals recovery from the simulated leachate with the concentration 7.6 g/L which in total consisted of 1.4% Co, 9.3% Dy, and 89.3% Nd. Obtaining complete separation was not possible by a single column due to the high Nd concentration in the feed. It is possible to obtain pure Co at the beginning of elution but the separation of Nd and Dy was not possible due to the materials uptake preference for Dy/Nd and their concentration in the feed.
List of original publications


The author contributions to the publications:
The author carried out all the experimental and analyses work together with co-authors. The author drafted all manuscripts I-III.
In article I, the author carried out all experiments and characterizations.
In article II, the author performed all experiments and characterizations, except the digestion experiment and 31P NMR.
In article III, the author completed all the experiments and characterizations, except the SEM and X-ray Tomography imaging, and the intraparticle diffusion modeling.
List of abbreviations

α-ZrP  Alpha-zirconium phosphate
β  The degree of dissociation
am-ZrP  amorphous-zirconium phosphate
$c_{eq}$  The metal concentration in the batch stripping solution at equilibrium
DMF  Dimethylformamide
$E_M$  The cationic equivalent fractions of M in the exchanger phase.
FT-IR  Fourier transform infrared spectra
HDDs  Computer hard disk drives
$K_a$  The acid dissociation constant
$K_d$  The distribution coefficient
$K_1$  The rate constants of the pseudo first-order
$k_2$  The rate constants of Pseudo-second-order kinetic models
MP-AES  Microwave plasma-atomic emission spectrometer
NdFeB  Neodymium-iron-boron
PAN  Polyacrylonitrile
$Q$  The capacity of the ion exchanger
$q_e$  The solute concentration in the sorbent phase at equilibrium
$q_i$  The rate constants of The initial amount of M loaded in the sorbent
$q_m$  The metal M ion equilibrium concentration in sorbent
$q_t$  The solute concentration in the sorbent phase at any given time t
R  Correlation coefficients
REEs  The rare-earth elements
SEM  Scanning electron microscopy
SF  Separation factor
$^{31}P$ MAS NMR  $^{31}P$ magic angle spinning nuclear magnetic resonance
TG  Thermogravimetry
XRD  X-Ray Powder diffraction
$Z_M$  The charge of M ion
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I would like to thank my supervisor Honorary Professor Risto Harjula. I greatly appreciate you accepted me as your PhD student in Belgium after interview. Your trust and guidance encouraged me to be a better researcher and a person with international view. Thank you for your teaching on the chemistry knowledge, experimental skills and paper writing as well as European culture, the travel and social. Your belief motivated me using my knowledge to make the world better. I deeply respect you that you are not only the knowledgeable chemist, but also a kind, optimistic, humorous and friendly person. I will always remember you as a respectable supervisor and friend.

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1 Introduction

Rare-earth elements (REEs) play an essential role in high technology and green industry due to their extensive use in applications such as permanent magnets, catalysts, rechargeable batteries, and lamp phosphors. Recovery and separation of REEs has become an important topic to reduce the pressure of increasing demand for these elements.

In general, NdFeB magnets consist of 30% to 40% REEs, with Nd accounting for the main component (15-30%). In addition, minor elements such as Dy and Co are added whenever special applications are needed. As such, recycling Nd, Dy, and Co from the end-of-life NdFeB permanent magnet is an important supplement for the primary production of REEs. It should be noted that REEs possess similar physical and chemical properties, which leads to difficulty in separating one from another.

Currently, environmentally friendly approaches with low cost and high efficiency are preferable in the metallurgy industry. Ion exchange technology has been extensively utilized in purification, separation and recovery of metals in chemical, food and pharmaceutical industries. Presently, it is still used industrially to produce high purity REEs. Inorganic ion-exchangers are more selective than organic resins because of the ion sieve functionality from rigid structures. So far, inorganic ion-exchangers have been used in large scales in water purification and removal of radionuclides from nuclear waste effluents.

The goal of this thesis was to develop a green separation and recovery process for REEs from NdFeB magnet leachates using zirconium phosphate (ZrP) ion exchangers. ZrPs have been extensively studied due to its high Brønsted acidity, high thermal and chemical stability and good stability under ionizing radiation. The alpha-ZrP (α-ZrP) with a layered structure and amorphous-ZrP (am-ZrP) owning a larger specific surface are worthy to be tested for the metals separation study. For the column separation, the gradient elution process has been proved as an efficient mean for materials separations. For the materials under study, the gradient elution process might be the preferred approach to obtain pure individual Co, Nd, and Dy.
2 Background

2.1 Rare-earth elements

2.1.1 Basic properties of rare-earth elements

The rare-earth elements (REEs) consist of 17 chemically similar elements, namely the lanthanide elements (Z=57-71, La-Lu) plus Sc and Y [1]. As members of the group 3 elements, they share the same typical oxidation state (+3). Certain REEs also present +2 and +4 oxidation states (e.g. Eu²⁺, Ce⁴⁺) due to half or full filling of an electron subshell.

Due to their different atomic numbers, REEs are typically separated into two subgroups, the ‘light’ REEs (lanthanides from La to Sm) and the ‘heavy’ REEs (Gd to Lu as well as Y; Eu can be considered either a light or heavy REE) [2]. The term ‘rare earth’ is derived from the historical difficulties of separation and obtaining high purity rare-earth metals and compounds. In reality, REEs are comparatively abundant in the earth’s upper crust; for example, Ce is as abundant as Cu. Nevertheless, REEs are almost always found together in minerals [3].

Electron configurations are of critical importance and determine the chemical and physical properties of REEs. Lanthanum, cerium, and gadolinium possess [Xe]⁴fⁿ⁶s² electrons, while the remaining lanthanide elements possess electron configurations of [Xe]⁴fⁿ⁻¹⁵d¹⁶s². Scandium and yttrium show chemically similar properties due to their (n-1)d¹⁴ns² configuration for the outermost electron shells, even though they do not have any 4f electrons [4].

In contrast to most other elements, the ionic radii of lanthanide elements continuously decrease with increase in atomic number (Figure 1a). This abnormal phenomenon is called the ‘lanthanide contraction’. This is explained by the imperfect shielding of one electron by another in the same subshell [5]. Compared to the shielding effect of 4f electrons of the lanthanide ions (Ln³⁺), the atomic radius of the lanthanide atoms is not much affected by the lanthanide contraction.
The lanthanide contraction results in regular changes in properties. One of the important properties is basicity (alkalinity), which decreases with increase in atomic number.

\[
\text{La}^{3+} > \text{Ce}^{3+} > \text{Pr}^{3+} > \text{Nd}^{3+} > \text{Pm}^{3+} > \text{Sm}^{3+} > \text{Eu}^{3+} > \text{Gd}^{3+} > \text{Tb}^{3+} > \text{Dy}^{3+} > \text{Ho}^{3+} > \text{Er}^{3+} > \text{Tm}^{3+} > \text{Yb}^{3+} > \text{Lu}^{3+}
\]

The basicity variation of the lanthanide elements provide the possibilities for separating the REEs from each other in hydrometallurgy [6].

2.1.2 Challenges of REE supply and recycling

Because of their distinctive electron features, REEs possess unique magnetic, electrical, catalytic, and optical properties [3, 7]. These properties make REEs essential components in various applications, such as high-temperature superconductors, secondary batteries, and hybrid cars [8]. REEs currently play a significant role in the transition from traditional to green economy. Consequently, the demand for REEs has significantly increased. The main driving forces behind the demand surge are the applications of REEs in permanent magnets, lamp phosphors, catalysts, and rechargeable batteries [9]. In accordance with the increasing demand for REEs, the yearly global demand for rare-earth metals was estimated to be 210,000 metric tons in 2015. However, the global primary mining production of rare-earth metals was 110,000 metric tons in 2015 [8].

REEs have been classified as the highest supply risk and as the most critical raw materials by the European Commission in both 2010 and 2017 [10]. Due to their applications in green technologies, demand for Nd and Dy has been estimated to increase by 700% and 2600%, respectively, over the next 25 years [11].
To address the supply pressure of REEs, the following three approaches have been proposed: developing less critical metals to substitute for critical REEs and investing in primary mining and recycling of REEs from urban and industrial wastes [9, 12, 13]. There are two kinds of substitution methods, such as substituting REEs with common base metals and substituting critical REEs with less critical ones (for example, using more La and Ce to replace Nd and Dy). This can partially solve the ‘balance problem’ [14]. This refers to the balance between the economic market demand and the natural abundance of REEs in ores [15, 16]. Primary mining is an efficient way to mitigate the supply risk of REEs, but can result in a higher environmental footprint and cause the balance problem [16-18]. Recycling REEs from urban and industrial wastes is one of the strategies encouraged by the green economy towards solving both the supply risk and the balance problem.

Although recycling of REEs has been extensively studied at the laboratory scale, the application of commercial recycling of REEs is insufficient. Regarding urban mining, it is estimated that at most 1% of REEs were recycled in 2011. This was due to inefficient collection, technology obstacles, and lack of motivation [19, 20].

2.1.3 Recycling REEs from permanent magnets

Neodymium-iron-boron alloys (NdFeB magnet) are the most common REE magnets. To suit various applications, the chemical composition has to be tuned by adding minor elements (Table 1). NdFeB magnets are widely used in wind turbines, hybrid and electric vehicles, computer hard disk drives (HDDs), household electrical appliances, and many small consumer electronic devices.

<table>
<thead>
<tr>
<th>Elements</th>
<th>Functions</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>Dy, Tb</td>
<td>Enhances anisotropy, coercivity, and demagnetization temperature</td>
<td>[9, 21]</td>
</tr>
<tr>
<td>Gd</td>
<td>Improves temperature efficiency</td>
<td>[22]</td>
</tr>
<tr>
<td>Nb</td>
<td>Gran refining</td>
<td>[22]</td>
</tr>
<tr>
<td>Co</td>
<td>Improves corrosion resistance</td>
<td>[21]</td>
</tr>
<tr>
<td>Cu, Al</td>
<td>Enhances sintering of the magnet alloy</td>
<td>[22]</td>
</tr>
<tr>
<td>Ga</td>
<td>Improves intrinsic coercivity and high temperature tolerance</td>
<td>[22]</td>
</tr>
</tbody>
</table>
It is estimated that approximately 26,000 metric tons of rare-earth metals have been consumed annually in the production of NdFeB magnets [13]. Due to lifecycle limitations, more and more end-of-life (EOL) magnets have accumulated, pending further treatment and recycling (Table 2). NdFeB magnets contain approximately 31 to 32 wt-% REEs. The main component is Nd, and small mixtures of Pr, Dy, Gd and Tb as well as other REEs are present for different applications. With increasing accumulation, more long-term efforts should be focused on REE recycling from EOL magnets. At the same time, improvements are needed in the development of technology and infrastructure [23]. By the year 2100, supply from recycling is estimated to fulfil half of REE demand [24]. Recycling REEs recycling is increasingly important not only as a supplement for REE demand but also as a more sustainable means of using natural resources [22].

Table 2. Present and future recycling of NdFeB magnets [9]

<table>
<thead>
<tr>
<th>Permanent NdFeB magnets (Nd, Dy, Tb, Pr)</th>
<th>Contribution to recycling (present/future)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Hard disk drives</td>
<td>Decreasing</td>
</tr>
<tr>
<td>Consumer electric and electronic devices</td>
<td>Stable</td>
</tr>
<tr>
<td>Automobiles</td>
<td>Stable</td>
</tr>
<tr>
<td>Electric vehicle and hybrid electric vehicle motors</td>
<td>Increasing</td>
</tr>
<tr>
<td>Wind turbine generators</td>
<td>Increasing</td>
</tr>
</tbody>
</table>

Many methods have been developed for recycling REE magnets, such as direct reuse, reprocessing, hydrometallurgical methods, and pyrometallurgical methods. The advantages and disadvantages of each method are compared and explained below.

Direct reuse in its current form is the most economical means of REE magnet recycling. This is due to low energy input and the fact that chemical consumption is not necessary and no waste is generated. However, direct reuse is only applicable to large and easily accessible magnets, such as wind turbines, large electric motors, and generators in hybrid and electric vehicles [9, 13, 25, 26].

Reprocessing of alloys to magnets after hydrogen decrepitation is particularly suitable for HDDs, as less energy input is required than metallurgical methods and no waste is generated. However, mixed scrap feed and oxidized magnets are not applicable for this method. Sufficient hydrogen access is the key factor for this technology [27, 28].
Hydrometallurgical methods, including leaching, separation and precipitation processes, consist of the same processing steps as the procedures for REE production from minerals. These methods are applicable to all types of magnets. However, these methods require multi-step processing, the consumption of large amounts of chemicals, and generate excessive amounts of waste effluents [8, 9, 22].

Pyrometallurgical methods include liquid-phase processing and gas-phase extraction. These methods consist of directly melting REE magnets to obtain master alloys, which can be generally applied to all types of magnets. Compared with hydrometallurgy methods, no wastewater is generated and fewer processing steps are necessary for pyrometallurgical methods [29, 30].

For liquid-phase processing, the methods require greater energy input and large amounts of solid waste are generated by electrical refining and by the glass slag method. In addition, oxidized magnets are not suitable for direct melting and liquid-metal extraction [9].

Gas-phase extraction can be applicable to non-oxidized and oxidized alloys. Nevertheless, the consumption of chlorine gas and generation of corrosive aluminium chloride are the disadvantages of this method [22].

2.2 Hydrometallurgical method for the recovery of REEs

Hydrometallurgy is a traditional technique in the field of extractive metallurgy and uses aqueous chemistry. It has been extensively adopted industrially for the recovery of metals from ores, concentrates, and residual materials [31, 32]. Hydrometallurgy is also the traditional method for recycling REEs from permanent magnets [33]. The key procedures are leaching and separation (solvent extraction, ion exchange, or precipitation). The final product is obtained by conversion to REE fluorides or oxides (Figure 2) [34, 35].
In commercial separation, mineral acids are applied to dissolve EOL REE magnets to obtain the pregnant leaching solution (containing for example chloride, nitrate, and thiocyanates). Leaching NdFeB magnets is challenging as these magnets contain approximately 72 wt-% iron. Recently, a combined pyrometallurgical and hydrometallurgical method using sulfate or nitration and calcination processes followed by water leaching was developed. More than 95% REEs can be extracted and Fe (less than 1%) and other impurities remains in the solid residue [34, 35]. Electrochemical processes are also useful for selective leaching. More than 95% REEs are extracted followed by membrane electrolysis. All iron was removed by oxidization in the anolyte and subsequently precipitated as Fe(OH)$_3$ [17].

### 2.2.1 Solvent extraction

Solvent extraction is the classic method to separate materials (metal complexes and organic compounds) from the mixture according to the two different immiscible liquids, normally aqueous solution and organic solvent [36]. The leachate is subjected to a solvent extraction process for REE separation. The REEs from an aqueous solution are transferred to the organic phase after the formation of complexes using a selective extractant. Cationic, anionic, and solvating extractants are frequently used, such as di-(2-ethyl-hexyl) phosphoric acid (HDEHP), 2-ethylhexyl phosphoric acid-2-ethylhexyl ester (EHEHPA), 2-Ethylhexyl 2-ethylhexyphosphonic acid (PC88A), bis/2,4,4-trimethylpentyl/phosphinic acid (Cyanex 272), Tri-$n$-butyl phosphate (TBP), and
tricaprylylmethylammonium chloride (Aliquat 336) [37-39]. Representative extractants and extraction mechanisms are shown in Table 3.

<table>
<thead>
<tr>
<th>Type</th>
<th>Representative extractants</th>
<th>Extraction mechanism</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cationic extractants</td>
<td>HDEHP, EHEHPA, PC88A</td>
<td>$\text{Ln}^{3+} + 3\text{HA} \leftrightarrow \text{LnA}_3 + 3\text{H}^+$</td>
<td>[40-42]</td>
</tr>
<tr>
<td>Anionic extractants</td>
<td>$\text{R}_3\text{CH}_3\text{N}^+\text{X}^-$ ($\text{R}$: C8-C12, $\text{X}$: nitrate or thiocyanate)</td>
<td>$[\text{R}_4\text{N\cdot LN(NO}<em>3]\text{]}</em>{\text{org}} + \text{Ln}^{3+} + 3[\text{NO}<em>3^-]</em>{\text{aq}} \leftrightarrow [\text{R}_4\text{N\cdot LN(NO}<em>3]\text{]}</em>{\text{org}}$</td>
<td>[43]</td>
</tr>
<tr>
<td>Solvating extractants</td>
<td>TBP</td>
<td>$3[\text{TBP}]_{\text{org}} + \text{Ln}^{3+} + 3[\text{NO}<em>3^-]</em>{\text{aq}} \leftrightarrow [\text{Ln(TBP)}_3(\text{NO}_3)<em>3]</em>{\text{org}}$</td>
<td>[44]</td>
</tr>
</tbody>
</table>

Due to the very similar physical and chemical properties, the selectivity of adjacent lanthanides is not very satisfactory. At industrial scale, decent separation is achieved often by using hundreds of mixer-settler units and adopted complicated flowsheets with reflux [8]. In addition, solvent extraction often involves the use of toxic volatile organic solvents. Thus, new extractants or new extraction systems are needed for REE recycling [45-47].

After the solvent extraction process by several stages of mixer-settlers, the low concentration leachate is suitable for recycling by ion-exchange technologies from a recycling efficiency and economic perspective [48, 49].

The problems described above drives the development of environmentally friendly separation methods that eliminate redundant processing units and the use of organic solvents.

2.2.2 Ion exchange

Ion exchange has played a significant role in the progress and development of purification and separation industry. Ion-exchange techniques are not only applied for purification processes, but are also extensively used in separation and extraction processes in the chemical, petrochemical, food, power, and pharmaceutical industries. Ion-exchange techniques in particular are used industrially to produce high-purity REEs [50, 51].
REE separation by ion exchange was initiated to separate fission products from nuclear reactors. With the support of the Manhattan project, ion exchange on organic resins was adopted to separate REEs and actinides [52, 53]. Theoretical analysis of column-separation processes and its pilot-scale separations has been systematically studied at the same time [52-55]. Since then, separation and purification of REEs by ion exchange replaced tedious fractional crystallization [56]. Before the 1960s, ion exchange was the dominant technology for obtaining individual REEs. Even though solvent extraction gradually became the key method for industrial production, ion-exchange technology is still widely implemented industrially to produce high-purity REEs [3, 51].

Inorganic ion exchangers are generally much more selective than organic resins due to the ion sieve functionality from the nanoporous, ordered, and rigid structures [57]. Inorganic ion exchangers are typically hydrous oxides (ZrO$_2$, SnO$_2$, H$_2$SbO$_3$, and Mn$_x$O$_y$), layered compounds (zirconium phosphates and layered double hydroxides), and framework structures that contain cavities or tunnels (zeolites, clays, pharmacosiderites, and ammonium molybdophosphate and sodium titanium silicates) [51, 58-62]. Thus far, inorganic ion exchangers have been used at large scale only for water purification and removal of radionuclides from nuclear waste effluents [63, 64].

2.3 Zirconium phosphate as inorganic ion exchanger

Zirconium phosphates (ZrPs) have received extensive attention because of its unique properties, including high Brønsted acidity, high thermal and chemical stability, and good stability under ionizing radiation. Therefore, ZrPs have found wide applications as catalysts [65, 66], ion exchangers [67-71], acid solids [72], intercalation hosts [73-75].

Alpha-zirconium phosphate, Zr(HPO$_4$)$_2$·H$_2$O (α-ZrP), is one primary crystalline acid salt of zirconium [76]. α-ZrP was first synthesized by Clearfield using a refluxing method and its structure was solved in the 1960s [77]. The compound Zr(HPO$_4$)$_2$·H$_2$O exhibits a layered structure (Figure 3). The layers are constructed by zirconium atoms connected by the oxygen atoms of the phosphate groups. Three oxygen atoms of each phosphate group bond Zr atoms, leaving one –OH group extending into the interlayer space. Adjacent layers are located as the staggered way to form a network resembling a hexagonally shaped cavity. The water molecule is situated in the cavity formed by three P–OH groups. The distances of interlayer space and between P–OH groups are 7.6 Å and 5.3 Å, respectively. The layers are held together by van der Waals forces [69, 78, 79].
Representative of inorganic ion exchangers, the primarily importance of ZrP is acid stability, reasonably high ion-exchange capacity, and selectivity for specific ions [78]. The ion-exchange capacity \( \alpha\text{-ZrP} \) is reported as 6.64 meq/g [69]. The ion-exchange behaviour of ZrP is significantly affected by its degree of crystallinity [80]. Amorphous ZrP (am-ZrP, also called semicrystalline) includes extremely small particles with a layered structure, and is observed often with a weak broad X-ray diffraction [80, 81]. The am-ZrP appears to have large amounts of microspores and possesses a comparatively greater specific surface area than that of crystalline \( \alpha\text{-ZrP} \) [82, 83]. These unique advantages of am-ZrP enhance its function in ion exchangers, catalysts, and adsorbents [82, 84]. However, to the best of our knowledge, extremely limited studies were conducted before this work for the separation of REEs by ZrP materials.

### 2.4 Organic-inorganic ion exchange composite

ZrPs display excellent ion-exchange properties. However, the powdery form of ZrPs easily causes pressure build-up and clogging in fixed bed columns.

To overcome these limitations, a porous composite bead has been developed by embedding the inorganic ion exchanger into porous granulated carriers [85, 86]. Commonly used porous granulated carriers include mesoporous silica, zeolite, activated carbon, alginate, diatomite, cellulose, and porous polymers [87-95].

---

**Figure 3.** Polyhedral representation of the structure of crystalline \( \alpha\text{-ZrP} \). \([\text{ZrO}_6]\) (green octahedron), \([\text{PO}_4]\) (grey tetrahedron), O (grey sphere) and OH (red dot) are shown.
Polyacrylonitrile (PAN) with a linear formula \((\text{C}_3\text{H}_3\text{N})_n\) is known as a common polymer carrier. PAN possesses excellent physicochemical properties, such as good performance on bead formation, strong adhesive force with inorganic materials, good solubility in organic solvents, high thermal stability, good radiation stability, and good chemical stability in strong acids (dissolves when the concentration exceeds 8 M HNO\(_3\), 5 M H\(_3\)PO\(_4\), or 5 M H\(_2\)SO\(_4\)) [96].

The composite’s hydrophilicity, porosity, and mechanical strength can be modified by using a PAN-based organic binding polymer. For the PAN-inorganic composite, inorganic materials existed as the ion exchange active component. The inorganic materials can be dispersed in the polymer with a very wide range, from 5 to 90 wt-% for the different demands [97]. Due to the advantages of the PAN-inorganic composite, this kind of composite based on PAN has been extensively applied in radiochemistry, heavy metal removal, separation, and other applications [87, 98-101].

### 2.5 Ion-exchange theory

\(\alpha\)-ZrP and Nd\(^{3+}\) are shown as an example to explain the ion-exchange process between metal ions and ion-exchanger ZrPs. The reaction can be expressed as

\[
3 \text{ZrP-OH} + \text{Nd}^{3+} \rightleftharpoons (\text{ZrP-O})_3\text{Nd} + 3\text{H}^+ \tag{1}
\]

The distribution coefficient \((K_d)\) represents the distribution of solute ions (Nd\(^{3+}\)) after the equilibrium between the solution and the ion exchanger (\(\alpha\)-ZrP):

\[
K_d = \frac{[\text{Nd}^{3+}]_{\text{eq}}}{[\text{Nd}^{3+}]_{\text{eq}}} \quad \tag{2}
\]

where \([\text{Nd}^{3+}]_{\text{eq}}\) is the concentration of Nd\(^{3+}\) taken up by \(\alpha\)-ZrP at equilibrium (mmol/g) and \([\text{Nd}^{3+}]_{\text{eq}}\) is concentration of Nd\(^{3+}\) remaining in solution after equilibrium (mmol/L).

In a typical batch ion-exchange experiment, \(\alpha\)-ZrP (mass m) is placed in REE solution (volume \(V\)) and rotated until equilibrium. The \(K_d\) for Nd\(^{3+}\) can be expressed as

\[
K_d = \frac{[\text{Nd}^{3+}]_{\text{eq}}}{[\text{Nd}^{3+}]_{\text{eq}}} = \frac{[\text{Nd}^{3+}]_{\text{eq}} - [\text{Nd}^{3+}]_{\text{eq}}}{[\text{Nd}^{3+}]_{\text{eq}}} \times \frac{V}{m} \tag{3}
\]

where \([\text{Nd}^{3+}]_{\text{eq}}\) is the concentration of Nd\(^{3+}\) remaining in the solution after equilibrium (mmol/L) and \([\text{Nd}^{3+}]_{\text{in}}\) is the initial concentration of Nd\(^{3+}\) (mmol/L).
In equilibrium state, the selectivity coefficient is defined as the ratio of ions in solution to ions on the ion exchanger. For the exchange between the metal ions (Nd\(^{3+}\)) and hydronium ions (H\(^{+}\)), the selectivity coefficient can be calculated by

\[
K_{\text{Nd/H}} = \frac{[\text{Nd}^{3+}]_{\text{eq}} [\text{H}^{+}]^{3}}{[\text{Nd}^{3+}]_{\text{eq}} [\text{H}^{+}]^{7}}
\]  

(4)

When sorption mechanism and system are unclear, simpler terminology has usually been used in practical ion-exchange studies. Thus

\[
[M] = q_{\text{eq}}
\]

(5)

\[
[M] = C_{\text{eq}}
\]

(6)

Here \([M]\) (and \(q_{\text{eq}}\)) and \([M]\) (and \(C_{\text{eq}}\)) are the metal concentrations in the ion exchanger and in the solution at equilibrium state, respectively. The indirect measurement was adopted for the uptake by ion exchanger. The metal concentrations are acquired by means of the changes of solution. Therefore

\[
[M] = (C_{0} - C_{\text{eq}}) \times \frac{\nu}{m}
\]

(7)

\[
[H] = Q - Z_{M}[M]
\]

(8)

where \(Q\) represents the total ion-exchange capacity, or how many cations can be taken up in total (milliequivalent per gram, meq/g). \(Z_{M}\) is the ion charge of \(M\).

Equivalent fractions or mole fractions are usually used to replace molarities or molalities. For example, the equivalent fraction of \(M^{Z+}\) (\(E_{M}\)) in the sorbent can be calculated from

\[
E_{M} = \frac{Z_{M}q_{M}}{Q}
\]

(9)

\(q_{M}\) is the ion concentration of \(M\) in solid phase (mmol/g) at equilibrium.

For elution studies of sorbent, the elution-% and the distribution coefficient at elution (\(K_{d,\text{elut.}}\)) can be obtained from

\[
\text{Elution - \%} = \left(1 - \frac{q_{\text{eq}}}{q_{i}}\right) \times 100 = \left\{1 - \frac{q_{i} - C_{\text{eq}}(V/m)}{q_{i}}\right\}
\]

(10)

\[
K_{d,\text{elut.}} = \frac{q_{\text{eq}}}{C_{\text{eq}}} = \left\{\frac{q_{i} - C_{\text{eq}}(V/m)}{C_{\text{eq}}}\right\}
\]

(11)
Here $q_i$ is the initial amount of M loaded in the solid phase, $q_{eq}$ is the amount of M in solid phase after stripping, and $C_{eq}$ is the equilibrium concentration of M in the stripping solution.
3 Experimental

3.1 Chemicals and regents

Three ion exchangers were synthesized for this work, namely α-ZrP, am-ZrP, and PAN-encapsulated am-ZrP. The chemicals ZrOCl$_2$·8H$_2$O and ZrCl$_4$ were the Zr sources of α-ZrP and am-ZrP materials, respectively. The chemicals NaH$_2$PO$_4$·H$_2$O and H$_3$PO$_4$ were used as the P source for α-ZrP and am-ZrP materials, respectively. The polymer carrier PAN was used to encapsulate am-ZrP to form the porous beads. The metal salts Co(NO$_3$)$_2$·6H$_2$O, Nd$_3$O$_9$·6H$_2$O and Dy$_3$O$_9$·xH$_2$O were used to prepare the Co-Nd-Dy ternary equimolar solution and simulated leachate. The reagents HNO$_3$, H$_2$SO$_4$, H$_3$PO$_4$, or HCl were employed to study the batch elution and column elution studies.

3.2 Synthesis

3.2.1 Preparation of α-ZrP

α-ZrP was synthesized using a modified recipe from Rajeh and Sziertes [102]. Solutions of NaH$_2$PO$_4$·H$_2$O (828.18 g) in 3 M HCl 600 (mL) and 322.25 g of ZrOCl$_2$·8H$_2$O in 300 mL deionized water were mixed in a 3-L glass Huber reactor (100 rpm). The obtained white homogeneous mixture was then allowed to stand for 24 h at 80°C and for another 24 h in room temperature. Subsequently, 3 L of 2 M HCl and 2 L of 2 M H$_3$PO$_4$ were sequentially used to wash the precipitate to remove unbound Na$^+$ and Cl$^-$ ions. After washed with deionized water to pH 3, the product was dried in an oven at 65°C for 48 h. The dried white product was pretreated using 0.1M HNO$_3$ (solid:liquid ratio 1:10) by a rotating mixer at 23°C for 24 h. The α-ZrP was then rinsed with deionized water to approximately pH 3 and dried in an oven at 65°C for 48 h. The preconditioned product was ground and sieved to desired grain size (200-100 mesh) for further study.
3.2.2 Preparation of am-ZrP

A precipitation method was used for am-ZrP material synthesis according to a previous report [103]. ZrCl₄ (30.7 g) was dissolved in HCl (430 mL, 2 M) and mixed with 400 mL of H₃PO₄ solution (1.25 M). The precipitate obtained was allowed to stand overnight. Subsequently, the white product was washed utilizing deionized water to pH 3. Then the am-ZrP was placed to oven and dried at 60°C for 48 h. Finally the product was ground and sieved to desired grain size (200-100 mesh) for further study.

3.2.3 Preparation of am-ZrP/PAN

The am-ZrP/PAN composite was prepared using methods described previously [99, 100]. Solution A was prepared by mixing am-ZrP (7.2 g), DMF (84 mL), and Tween 80 (2 mL) for 2 h with magnetic stirring at 60°C. PAN (4.8 g) was added to solution A and continued stirring for another 2 h. The composite beads were made by a gelation process where the synthesis mixture was dropwise added to deionized water (2L) using a syringe pump and needle (0.6 mm). The formed beads were aged in deionized water for 24 h. The product was then rinsed with deionized water (2 L). The obtained product was dried by freeze-drying (Christ alpha 1-4 LSC) under 0.570 mbar at -26°C.
3.3 Analytical methods

The structural study for materials used the characterization methods of X-ray powder diffraction (XRD), the characteristic Fourier transform infrared (FT-IR) spectra, thermogravimetry (TG) and solid-state $^{31}$P magic angle spinning nuclear magnetic resonance ($^{31}$P MAS NMR) spectra.

For the morphology and spatial distribution study, scanning electron microscopy (SEM) and X-ray tomography were used.

An Agilent 4200 microwave plasma-atomic emission spectrometer (MP-AES) was used to determine the metal concentrations.

3.4 Experimental plan

$\alpha$-ZrP, am-ZrP, and am-ZrP/PAN were developed for separation of the main components of an NdFeB magnet (Co, Nd and Dy) after a selective leaching procedure.

Due to the layered structure and reported high capacity (6.64 meq/g), $\alpha$-ZrP was chosen as the first ion exchanger to make full use of the ion exchange site’s inner and outside layers. We hoped that the layered structure of $\alpha$-ZrP would bring additional selectivity due to the ion-sieve function of the interlayer spaces. A detailed research experimental design of paper I is presented in Figure 5.

![Figure 5. Flowchart of the research design content of Paper I](image-url)
When compared with $\alpha$-ZrP, am-ZrP has a larger specific surface area and contains numerous microspores and mesopores [104-106]. The experimental design flowchart of paper II is shown in Figure 6.

Figure 6. Flowchart of the research design content of paper II

When comparing the ion exchange results between $\alpha$-ZrP and am-ZrP, am-ZrP showed better ion-exchange behaviour. In addition, column separation for Co, Nd, and Dy were achieved using single-column separation. These promising results led us to apply am-ZrP for the scale-up study for industry.

Am-ZrP in powdery form can easily cause operational problems, such as clogging in the pilot column operations. Thus, am-ZrP was difficult to apply in the pilot-scale test. To overcome this limitation, we employed PAN as the polymer carrier to encapsulate the am-ZrP into composite beads. Moreover, a gradient elution process was utilized for the purpose to achieve well separation. The column experiments were optimized by changing the feed concentration, running speed, operational temperature, and concentration of eluting agent (Figure 7).
Figure 7. Experimental plan for paper III.
4 Results and discussion

4.1 Characterization of synthesis samples

4.1.1 Analysis and structure comparison between $\alpha$-ZrP and am-ZrP

The synthesized $\alpha$-ZrP is a platelet-like highly crystalline material with an interlayer space of 7.6 Å calculated from the (002) diffraction peak (Figure 8a). The SEM image shows regular crystals (Figure 8b). In contrast, the synthesized am-ZrP had weak and broad X-ray diffraction (Figures 8d and 8e). The SEM image showed the amorphous nature of am-ZrP.

For the FT-IR spectrum, the feature band(s) of deformation and vibration of P-OH were observed at 1249, 1069, 1038, and 963 cm$^{-1}$ for $\alpha$-ZrP (Figure 8c) and 987 cm$^{-1}$ for am-ZrP (Figure 8f).

![Figure 8](image)

**Figure 8.** Characterization of synthesized $\alpha$-ZrP. a) XRD pattern, b) SEM image, c) FT-IR spectrum. Characterization of synthesized am-ZrP. d) XRD pattern, e) SEM image, f) FT-IR spectrum.
\( \alpha \)-ZrP was synthesized without hydrofluoric acid (HF). This was beneficial for production of the large crystal size of \( \alpha \)-ZrP (diameter 1-4 \( \mu \)m) in this work synthesis (Table 4). The reaction was completed at lower temperature compared with the previous methods shown in Table 4. Am-ZrP was synthesized using an easily available method by precipitation at room temperature. Both syntheses were in agreement with the aims of green chemistry and were promising for pilot-scale application.

**Table 4.** Different methods for crystalline \( \alpha \)-ZrP preparation

<table>
<thead>
<tr>
<th>Synthesis method</th>
<th>Zr source</th>
<th>P Precursor</th>
<th>Temperature</th>
<th>Reaction time</th>
<th>Average diameter</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>Hydrothermal</td>
<td>ZrOCl₂·8H₂O</td>
<td>3 M H₃PO₄</td>
<td>200°C</td>
<td>24 h</td>
<td>~400 nm</td>
<td>[79]</td>
</tr>
<tr>
<td></td>
<td>ZrOCl₂·8H₂O</td>
<td>12 M H₃PO₄</td>
<td>200°C</td>
<td>24 h</td>
<td>1 ( \mu )m</td>
<td>[79]</td>
</tr>
<tr>
<td></td>
<td>ZrOCl₂·8H₂O</td>
<td>3 M H₃PO₄, 5 M HF</td>
<td>100°C</td>
<td>24 h</td>
<td>1-4 ( \mu )m</td>
<td>[79]</td>
</tr>
<tr>
<td>Refluxing</td>
<td>ZrOCl₂·8H₂O</td>
<td>3 M H₃PO₄</td>
<td>100°C</td>
<td>24 h</td>
<td>~60 nm</td>
<td>[79]</td>
</tr>
<tr>
<td></td>
<td>ZrOCl₂·8H₂O</td>
<td>12 M H₃PO₄</td>
<td>100°C</td>
<td>24 h</td>
<td>~200 nm</td>
<td>[79]</td>
</tr>
<tr>
<td></td>
<td>ZrOCl₂·8H₂O</td>
<td>6 M H₃PO₄</td>
<td>94°C</td>
<td>48 h</td>
<td>~120 nm</td>
<td>[107]</td>
</tr>
<tr>
<td></td>
<td>ZrOCl₂·8H₂O</td>
<td>10 M H₃PO₄</td>
<td>-</td>
<td>-</td>
<td>250 nm</td>
<td>[81]</td>
</tr>
<tr>
<td></td>
<td>ZrOCl₂·8H₂O</td>
<td>NaH₂PO₄·H₂O, 3 M HCl</td>
<td>80°C</td>
<td>24 h</td>
<td>1-4 ( \mu )m</td>
<td>Paper I</td>
</tr>
</tbody>
</table>

### 4.1.2 Determination of molecular formula of am-ZrP

The composition of am-ZrP could be easily changed by adjusting the synthesis conditions. The elemental content was obtained from am-ZrP digestion experiments. The P/Zr ratio was determined to be 2.03. The three peaks of \(^{31}\)P MAS NMR spectrum represented the three different phosphate groups, namely \(-\text{H}_2\text{PO}_4\) (-13.6 ppm), \(-\text{HPO}_4\) (-21.7 ppm), and \(-\text{PO}_4\) (-27.5 ppm) (Figure 9a) [108, 109]. From the peak deconvolution, the ratio of these phosphate groups was estimated to be 9.3:100:4.8.

Two weight-loss steps were observed in the TG curve (Figure 9b). The release of physically bound water was suggested for the first weight loss step (7.07%; 25°C—184 °C) and the condensation of \(-\text{H}_2\text{PO}_4\) was suggested for the second weight loss step (4.98%, 184°C—800°C) [110]. From the XRD pattern of the am-ZrP calcined at 800°C (Figure 9c), the substance was identified as ZrP₂O₇ [111], which was consistent with the P/Zr ratio of 2.03 from the digestion analysis.
Finally, after combining the results of the digestion experiment, $^{31}$P MAS NMR spectrum, and TG analysis, the chemical formula of am-ZrP was determined to be $\text{Zr(H}_2\text{PO}_4)_{0.17} (\text{HPO}_4)_{1.78} (\text{PO}_4)_{0.09} \cdot 0.96\text{H}_2\text{O}$. Based on the calculated formulae, theoretical ion-exchange capacity of am-ZrP should be 6.97 meq/g.

**Figure 9.** Characterization of synthesized am-ZrP. a) The deconvolution peaks based on $^{31}$P MAS NMR spectrum, b) TGA curve, c) XRD pattern of am-ZrP calcined at 800°C.

### 4.1.3 Characterizations of synthesized am-ZrP/PAN composite

The regular am-ZrP/PAN spheres are shown in Figure 10a. Their size distribution was evaluated by analyzing a total of 199 particles in the perspective of volume and number distribution. The average bead size was 2 mm in diameter according to the data of equivalent (CE) measurements (Table 5). The circularity value was determined to range from 0.74 to 0.98 (Table 5), indicating a more or less spherical shape. A cross-section of the beads is shown in Figure 10b, which presents the imaged internal porous structure. This is the desired feature for the sorption material (Figure 10c) [94].

X-ray tomography demonstrated the porous feature of the beads and the more or less homogeneous distribution of the inorganic am-ZrP in the polymer matrix (Figure 10d). The porosity ratio of am-ZrP/PAN was determined to be approximately 40%. In addition, the spatial distribution of am-ZrP was characterized along the Z-axis of the bead with XY planes (blue arrow, Figure 10e). The spatial distribution study revealed that am-ZrP was quite evenly distributed in the inner surface and that there was slightly more am-ZrP near the bead surface than elsewhere.
Table 5. Analyzed particle parameters of am-ZrP/PAN beads

<table>
<thead>
<tr>
<th>Name</th>
<th>Volume distribution</th>
<th>Number distribution</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Minimum</td>
<td>Maximum</td>
</tr>
<tr>
<td>CE Diameter (μm)</td>
<td>1640</td>
<td>2341</td>
</tr>
<tr>
<td>Circularity</td>
<td>0.74</td>
<td>0.98</td>
</tr>
</tbody>
</table>

*D [4, 3] is the equivalent volume mean diameter

Figure 10. Synthesized am-ZrP/PAN beads. a) SEM image, b) SEM image of a bead’s cross-section, c) SEM image for the porous structure, d) X-ray tomography image, e) spatial distribution as determined along the blue arrow (Z-axis of the bead) with XY-planes, f) curve of am-ZrP fraction (Z-axis direction).

The XRD pattern of am-ZrP shows a typical feature for amorphous ZrP. As am-ZrP (or semicrystalline ZrP) includes considerably small particles with a layered structure, the layered feature was revealed by weak and broad peaks [78, 80, 81]. The peak shifts were observed from 10.0° to 8.1° (2theta), indicating that the interlayer space was expanded from 9 to 10.8 Å (Figure 11). This phenomenon suggested that DMF was intercalated into the interlayer space. In previous studies, α-ZrP and α-SnP have been studied as the host for DMF intercalation [75, 112]. Double DMF molecules were suggested to be intercalated non-vertically to the interlayer with the hydrogen bond (P)-O-H⋯O-CH-(N), owing to the limited interlayer space (Table 6).
Table 6. DMF intercalation on α-ZrP/DMF, α-SnP/DMF, and am-ZrP/DMF

<table>
<thead>
<tr>
<th>Inorganic material</th>
<th>Basal spacing (Å)</th>
<th>Intercalated basal spacing (Å)</th>
<th>Diameter of DMF (Å)</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>α-SnP</td>
<td>7.8</td>
<td>13.3</td>
<td>3.5</td>
<td>[75]</td>
</tr>
<tr>
<td>α-ZrP</td>
<td>7.6</td>
<td>11.2</td>
<td>3.5</td>
<td>[112]</td>
</tr>
<tr>
<td>am-ZrP</td>
<td>9.0(^a)</td>
<td>10.8</td>
<td>3.5</td>
<td>this work</td>
</tr>
</tbody>
</table>

\(^a\)Semicrystalline am-ZrP material.

Figure 11. Illustration of the intercalation of DMF in semicrystalline ZrP based on XRD patterns.

The bands of the FTIR spectrum of the am-ZrP/PAN composite were consistent with the bands of the PAN beads and am-ZrP (Figure 12a). The strongest bands at 954 cm\(^{-1}\) and 1047 cm\(^{-1}\) are the P-OH deformation and the vibration of the orthophosphate group [113].

For the TG curves of am-ZrP/PAN beads, the elimination of free water molecules contributes to 8% mass loss (<300°C) (Figure 12b). It was suggested that condensation of H\(_2\)PO\(_4\) functional groups of am-ZrP and decomposition of PAN occurs from 300°C to 700°C [87]. The 56.7% am-ZrP content in the am-ZrP/PAN composite could be calculated based on the thermal analysis data from powdery am-ZrP, PAN beads, and am-ZrP/PAN beads.

Figure 12. PAN beads, powdery am-ZrP, and am-ZrP/PAN beads. a) FTIR spectra, b) TGA curves.
4.2 Ion-exchange behaviour study

4.2.1 Potentiometric titration

The P-OH group in α-ZrP can be considered as a weak acid, which undergoes a dissociation reaction:

\[
P{-}OH \leftrightarrow P{-}O + H^+ \quad (12)
\]

The acid dissociation constant \( K_a \) is defined as:

\[
K_a = \frac{[P^{-}][H^+]}{[P{-}OH]} \quad (13)
\]

Here, [POH] is the undissociated phosphate content (mM/g) of the material. [PO\textsuperscript{-}] and [H\textsuperscript{+}] are the concentrations of the dissociated phosphate and hydronium ion inside the pores of α-ZrP.

The degree of dissociation (\( \beta \)) of P-OH can be expressed as

\[
\beta = \frac{[P^{-}]}{[P{-}OH]+[P^{-}]} \quad (14)
\]

After combining equations 13 and 14, the following equation is obtained:

\[
log \frac{1-\beta}{\beta} = pK_a - pH \quad (15)
\]

\( \beta \) can then be calculated from the equation below:

\[
\beta = \frac{1}{1+10^{pK_a-pH}} \quad (16)
\]

Typically, a 1.0 M NaNO\textsubscript{3} solution was used to keep a constant ionic strength. The initial exchange with NaNO\textsubscript{3} is inevitable.

\[
POH + Na^+ \leftrightarrow PONa + H^+ \quad (17)
\]

As a result, the amount of the conversion to the Na form (\( q_{Na1}, \text{mmol/g} \)) from NaNO\textsubscript{3} can be obtained from the following:

\[
q_{Na1} = ([H^+]_{eq} - [H^+]_0) \ (V/m) \quad (18)
\]
Here, $[H^+]_{eq}$ is the concentration of $H^+$ in the solution at equilibrium (mmol/L) and $[H^+]_i$ is the initial concentration of $H^+$ in the solution (mmol/L). $V$ is the solution volume (mL) and $m$ is the material mass (mg).

NaOH is subsequently used for titration. The reaction is described below as:

$$POH + NaOH \leftrightarrow PONa + H_2O$$ \hspace{1cm} (19)

The amount of the conversion to the Na form ($q_{Na2}$, mmol/g) from NaOH can be acquired from the equation:

$$q_{Na2} = ([OH^-]_i - [OH^-]_{eq})(V/m)$$ \hspace{1cm} (20)

where $[OH^-]_i$ and $[OH^-]_{eq}$ is the initial and the equilibrated solution concentrations, respectively.

The total ion exchange capacity ($Q$) of $\alpha$-ZrP can be calculated from the summation of $q_{Na1}$ and $q_{Na2}$:

$$Q = q_{Na1} + q_{Na2} = ([H^+]_{eq} - [H^+]_i + [OH^-]_i - [OH^-]_{eq})(V/m)$$ \hspace{1cm} (21)

The degree of crystallinity highly affects the titration behaviour. Amorphous materials often show a steady increase in titration curves. Normally, clear inflexion points can be observed when titrating a material with high crystallinity [67, 76]. A total ion-exchange capacity of 6.6 meq/g has been obtained by NaOH titration of $\alpha$-ZrP. The crystalline $\alpha$-ZrP has been identified as a diprotic weakly acidic cation exchanger [68, 69].

In this work, ZrP displayed a diprotic character in titration curves (Figure 13). The ion-exchange capacity of a total of 6.6 meq/g was obtained with the first and second equivalence point at 5 meq/g and at 6.6 meq/g, respectively (Figure 14). To evaluate the $pK_{a1}$ and $pK_{a2}$ for the diprotic character of titration curves, the more acidic sites with the ion-exchange capacity of 5.0 meq/g ($Q_1$) and the weaker acidic sites of 1.65 meq/g ($Q_2$) were distinguished according to the apparent equivalence points in Figure 13. The $pK_a$-value was chosen from the middle point value of the plateaus of the titration curve, in this case the $pK_{a1} = 3.5$ and $pK_{a2} = 6.5$ were obtained.

The degree of dissociation for the more acidic sites and the weaker acidic sites can be calculated using the titration data from Eq. 16 and the equation below.

$$q_{Na} = \beta_1 Q_1 + \beta_2 Q_2$$ \hspace{1cm} (21)
The best fit between Eq. 21 and 22 was then acquired when using \( pK_{a1}' \) (3.3) and \( pK_{a2}' \) (6.3). The \( pK_{a1}' \) and \( pK_{a2}' \) values are very close to the value we obtained from the titration curve.

![Figure 13. Titration curve of α-ZrP with 1.0 M NaOH in a 1.0 M NaNO₃ background.](image)

The titration of am-ZrP was performed using a 1.0 M NaOH solution with 1.0 M NaNO₃ solution as background. The pH of the solution changed from an initial pH 6.5 (1.0 M NaNO₃) to pH 2.6 during equilibrium time, indicating a 2.13 meq/g conversion to the Na form (\( q_{Na1} \)). Adding this conversion value to the titration data, we observed that the ion-exchange capacity was 9.23 meq/g (\( q_{Na1} + q_{Na2} \)) as estimated from the inflection point of the plateau (Figure 14). This ion-exchange capacity value is higher than 6.97 meq/g as calculated from the chemical formula. This deviation might be due to the hydrolysis of material in alkaline solutions [110].

As for the individual \( pK_a \) values in crystalline α-ZrP, we have demonstrated how to determine its values from the titration curve and the relevant equations. These studies were only based on the apparent equivalence points in the titration curves. It was not possible to acquire these \( pK_a \) values from the steadily increasing titration curve of am-ZrP.
The titration of the PAN beads and am-ZrP/PAN beads was studied using 1.0 M NaNO₃ solution as background. The R-OH conversion to Na-form (\(q_{Na1}\)) can be calculated from the difference of the initial and equilibrium pH of the 0.1 M NaNO₃ solution (Eq. 20).

The \(q_{Na1}\) of the pure PAN beads was calculated to be 0.004 meq/g. This value was disregarded from further calculations because it was negligible. The total ion-exchange capacity of pure PAN beads was estimated 0.46 meq/g from the plateau of the titration curve (Figure 15a) [114]. The \(q_{Na1}\) for the am-ZrP/PAN beads was calculated to be 0.53 meq/g. The total ion-exchange capacity (\(q_{Na1} + q_{Na2}\)) of am-ZrP/PAN beads was estimated 4.5 meq/g. (Figure 15b).

The am-ZrP content in the beads was calculated to be 56.7% based from the TG analysis. In addition, the am-ZrP content of am-ZrP/PAN beads can be estimated based on the theoretical capacity (6.97 meq/g) of am-ZrP and the capacity (4.5 meq/g) of am-ZrP/PAN beads. Using the ion-exchange capacities of the am-ZrP/PAN composite, the am-ZrP content in the beads was calculated to be 57.9%, which is consistent with the value from the TG analysis.
4.2.2 Effect of pH on sorption

The effect of pH on metal sorption of α-ZrP, am-ZrP, and am-ZrP/PAN was investigated using 1.0 mM equimolar Co, Nd, and Dy nitrate solution. The uptake amounts for these ion exchangers are shown in Table 7. α-ZrP and am-ZrP showed a similar total metal uptake of 1.6 meq/g at equilibration pH ~3.5. However, am-ZrP had a higher separation factor (SF) based on the $K_d$ value in Figure 16 a, b and c. The SFs of am-ZrP were calculated to be 6.5, 2.1, and 3.2, corresponding to pH 1.0, 1.8, and 2.6, respectively (Figure 16b). Dy was found to be the most favoured element obtained for am-ZrP materials based on the values of $K_d$(Co)=6 mL/g, $K_d$(Nd)=180 mL/g, and $K_d$(Dy)=458 mL/g at pH 1.8. In addition, compared with the total uptake amount, the equivalent-% of Co was not more than 3% at pH below 3, indicating excellent potential separation of Co from Nd and Dy. For am-ZrP/PAN, the strong sorption of Co after pH 4 caused the obviously decrease of the Nd and Dy $K_d$ values (Figure 16c).

For the uptake amount of am-ZrP/PAN, we observed that the metal uptake by am-ZrP increased approximately 50% at pH 3.5 (Table 7) when focused only on the inorganic counterpart. The value was calculated to be 2.43 meq/g for am-ZrP in am-ZrP/PAN beads compared to 1.65 meq/g of pristine am-ZrP. This phenomenon was suggested to result from the DMF intercalation to the layers of ZrP. It is reported that the rather large hydrated metal ions such as REEs are inaccessible in the cavity of the layer due to diffusional resistance [115]. However, with the interlayer space expansion from 9 Å to 10.8 Å after DMF intercalation, the ion-exchange sites became accessible for the hydrated metal ions. Therefore, sorption efficiency was improved and the metal uptake amount increased.
Table 7. Metal uptake data on α-ZrP, am-ZrP, and am-ZrP/PAN in the initial concentration as 1.0 mM equimolar Co, Nd, and Dy.

<table>
<thead>
<tr>
<th>Material</th>
<th>pH&lt;sub&gt;eq&lt;/sub&gt;</th>
<th>Co (meq/g)</th>
<th>Dy (meq/g)</th>
<th>Nd (meq/g)</th>
</tr>
</thead>
<tbody>
<tr>
<td>α-ZrP</td>
<td>3.5</td>
<td>0.44</td>
<td>0.61</td>
<td>0.63</td>
</tr>
<tr>
<td>am-ZrP</td>
<td>3.4</td>
<td>0.43</td>
<td>0.62</td>
<td>0.61</td>
</tr>
<tr>
<td>am-ZrP/PAN</td>
<td>3.5</td>
<td>0.26</td>
<td>0.58</td>
<td>0.59</td>
</tr>
</tbody>
</table>

Figure 16. Distribution coefficients of the metal sorption of 1.0 mM equimolar Co, Nd, and Dy nitrate solution. a) α-ZrP, b) am-ZrP, c) am-ZrP/PAN.

When increasing the metal concentration to 2.0 mM, there is no obvious plateau shown in the uptake curves of α-ZrP (Figure 17a). However, the two separate $K_d$ linear figures were based on the two acid-exchange sites of α-ZrP at pH 1 to 5.3 and pH 5.3 to 6.4 (Figure 17b and 17c). The log $K_d$ versus pH showed a low slope (0.33-0.59) in the first domain and a considerably larger slope (2.25-2.42) in the second domain. This phenomenon indicates that the less acidic exchange site ($pK_{a2}=6.3$) was used for exchanging with the REE and Co ions.

Visual Minteq Software was used to calculate the solubility of Co, Nd, and Dy at the ranges of pH and the metal concentrations [116]. There was no indication of precipitation throughout the study.
4.2.3 Sorption isotherm study

The sorption isotherms of α-ZrP were investigated at approximately pH 2.5 and pH 4.5 at equilibrium state. The metal uptakes of Nd and Dy showed an increasing trend following the equilibrium concentration of the metal, increasing until a plateau emerged approximately at 3.2 mM to 3.6 mM at pH 2.5 (Figure 18a). In contrast, the uptake of Co decreased after the equilibrium concentration of 3 mM. As Nd and Dy have higher affinity than Co, Co was replaced by Dy and Nd as the concentration of Nd and Dy increased. At pH 4.5, we observed that Co and Nd uptake started to decrease after the equilibrium concentration at approximately 2.0 mM (Figure 18b). Therefore, the order of preference for metal adsorption by α-ZrP is Dy>Nd>Co.

Compared to α-ZrP, higher metals uptake was found by am-ZrP at C_{in} 5.0 mM and pH 2.5 (Figure 18c). The total metals uptake of am-ZrP (3.4 meq/g) was approximately five times larger than that of α-ZrP (0.6 meq/g) (Table 8). In addition, α-ZrP and am-ZrP showed the same order of preference Dy>Nd>Co, also Co uptake was rather low on α-ZrP and am-ZrP at pH 2.5 or pH 4.5 (Table 8).
### Table 8. Analyzed data for the sorption isothermal study of $\alpha$-ZrP and am-ZrP

<table>
<thead>
<tr>
<th>Ion exchanger</th>
<th>pH\textsubscript{eq} (equilibrium)</th>
<th>$C_{\text{in.}}$ (initial concentration)</th>
<th>Uptake (meq/g)</th>
<th>Co (Equivalent-%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>$\alpha$-ZrP</td>
<td>2.5</td>
<td>5.0 mM</td>
<td>0.02</td>
<td>0.18</td>
</tr>
<tr>
<td>$\alpha$-ZrP</td>
<td>4.5</td>
<td>5.0 mM</td>
<td>0.20</td>
<td>1.1</td>
</tr>
<tr>
<td>am-ZrP</td>
<td>2.5</td>
<td>5.0 mM</td>
<td>0.18</td>
<td>1.5</td>
</tr>
</tbody>
</table>

Equivalent-% is the percentage calculated from single metal uptake (meq/g) compared with the total metal uptake (meq/g).

**Figure 18.** Isothermal sorption curves of Co, Nd, and Dy nitrate solution. a) $\alpha$-ZrP at approximately pH\textsubscript{eq} 2.5, b) $\alpha$-ZrP at approximately pH\textsubscript{eq} 4.5, c) am-ZrP at approximately pH\textsubscript{eq} 2.5.

#### 4.2.4 Batch elution study

To choose the right mineral acid for the column elution study, batch elution was necessary as pre-test experiments. For this work, four different acids at two concentrations were tested (Table 9). The most efficient total elution (96.7% Dy and 99.1% Nd) was obtained using 0.1 M H\textsubscript{2}SO\textsubscript{4} as eluent for $\alpha$-ZrP. Compared to other acids the difference in elution percentage was significant (>25%) and also a difference in eluent concentration was observed. An increase in H\textsubscript{2}SO\textsubscript{4} concentration decreased the elution. That is unusual and was not seen with the other acids. SF (the ratio of $K_d$) was calculated based on the $K_d$ values. HCl would be a good candidate due to a SF (Dy/Nd) of 3.08 at a concentration of 1.0 M (Table 9). However, HCl was not adopted in our work due to its high corrosiveness.

For am-ZrP, H\textsubscript{2}SO\textsubscript{4} was also the most efficient when compared to the elution results of other acids. We observed that 85% Nd and 83% Dy were removed by 1.0 M H\textsubscript{2}SO\textsubscript{4} (Table 9). The elution efficiencies of acids at 0.1 M was observed as H\textsubscript{2}SO\textsubscript{4}>HNO\textsubscript{3}>HCl>H\textsubscript{3}PO\textsubscript{4}. HCl and H\textsubscript{3}PO\textsubscript{4} showed
less than 30% elution at this concentration. The separation factors calculated from $K_d$ values of Dy and Nd show a decreased order as HCl (1.8)>HNO$_3$ (1.2)>H$_2$SO$_4$ (0.9)>H$_3$PO$_4$ (0.7).

Table 9. Batch elution data of $\alpha$-ZrP and am-ZrP in different mineral acids

<table>
<thead>
<tr>
<th>Stripping agent (Mol/L)</th>
<th>$\alpha$-ZrP</th>
<th>am-ZrP</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Elution (%)</td>
<td>$K_d$ (ml/g)</td>
</tr>
<tr>
<td></td>
<td>Dy</td>
<td>Nd</td>
</tr>
<tr>
<td>HCl 0.1</td>
<td>62</td>
<td>67</td>
</tr>
<tr>
<td></td>
<td>1.0</td>
<td>92</td>
</tr>
<tr>
<td>HNO$_3$ 0.1</td>
<td>70</td>
<td>65</td>
</tr>
<tr>
<td></td>
<td>1.0</td>
<td>86</td>
</tr>
<tr>
<td>H$_3$PO$_4$ 0.1</td>
<td>22</td>
<td>5.7</td>
</tr>
<tr>
<td></td>
<td>1.0</td>
<td>82</td>
</tr>
<tr>
<td>H$_2$SO$_4$ 0.1</td>
<td>96</td>
<td>99</td>
</tr>
<tr>
<td></td>
<td>1.0</td>
<td>85</td>
</tr>
</tbody>
</table>

4.2.5 Sorption kinetics

For the sorption kinetics study, a pseudo-first-order equation, pseudo-second-order equation, and simplified model of resistance to intraparticle diffusion were employed to determine the characteristic sorption constants of $\alpha$-ZrP and am-ZrP [30-32].

The pseudo-first-order equation, the pseudo-second-order equation, and the intraparticle diffusion model were as follows:

$$\log(q_e - q_t) = \log q_e - \left(\frac{k_1}{2.303}\right)t$$  \hspace{1cm} (25)
\[
\frac{t}{q_t} = \frac{1}{k_2 q_e} + \frac{1}{q_e} \quad (26)
\]

\[
q_t = k_{\text{int}} t^{1/2} + C \quad (27)
\]

Here \(q_e\) is the maximum uptake value at equilibrium and \(q_t\) is the uptake value at any time \(t\). The rate constants \(k_1\) and \(k_2\) belong to the pseudo-first-order and the pseudo-second-order models, respectively. The slope \(k_{\text{int}}\) is the intraparticle diffusion constant. If a straight line can be obtained from a plot of \(q_t\) versus \(t^{1/2}\), the adsorption mechanism should follow the process of intraparticle diffusion.

We observed that the metal uptake by \(\alpha\)-ZrP increased with time, increasing until equilibrium was almost reached at 24 h (Figure 19a). A shorter time period of 12 h was observed in reaching equilibrium for the sorption by am-ZrP and am-ZrP (Figure 19b and 19c). At this point (12 h), the total uptake value for Dy and Nd were 1.27 meq/g (am-ZrP) and 1.37 meq/g (am-ZrP/PAN), respectively. All three ion-exchangers followed the pseudo-second-order model. The results suggest that the rate-limiting step for the adsorption of metal ions is the ion-exchange process [118-120].

For the am-ZrP/PAN composite, an intraparticle diffusion model based on Fick’s law was used to determine the diffusion coefficients [117].

The intraparticle diffusion model based on Fick’s law is as follows:

\[
\frac{\partial \hat{q}_j}{\partial t} = \frac{1}{r^2} \frac{\partial}{\partial r} \left( r^2 D_j \frac{\partial \hat{q}_j}{\partial r} \right),
\]

\[
\frac{\partial \hat{q}_j}{\partial r} \bigg|_{r=0} = 0, \quad \hat{q}_j \bigg|_{r=r_p} = f\left(c_j\right) \quad (28)
\]

here \(D_j\), \(\hat{q}_j\), and \(r\) are the intraparticle diffusion coefficient, local concentration, and radial coordinate of species \(j\), respectively.

Metal uptake by the am-ZrP/PAN composite was reasonably fast, even though the material’s particle size was approximately 2 mm in diameter (Fig. 19d). The value of \(D_{\text{Dy}}\) 1.05·10\(^{-13}\) m\(^2\)/s (\(R^2=0.937\)) and \(D_{\text{Nd}}\) 1.02·10\(^{-13}\) m\(^2\)/s (\(R^2=0.967\)) were an order of magnitude lower than that of \(D_{\text{Co}}\) 4.31·10\(^{-12}\) m\(^2\)/s (\(R^2=0.990\)), most likely due to their larger hydrated radii [121, 122].
4.3 Column experiments

4.3.1 Loading and breakthrough

The metal loading on columns of α-ZrP, am-ZrP, and am-ZrP/PAN materials were tested using 1.0 mM equimolar Co, Nd, and Dy.

The breakthrough of Co was observed first and followed by Nd and Dy (Figure 20a, b and c). This suggests that the metals were preferred by these ion-exchange materials in the order Dy>Nd>Co. The loaded amount of Co relative to the total metal loading amount ranged from 0.8 to 3.7 equivalent-% (Table 10). When compared with α-ZrP, am-ZrP displayed a higher total metal loading capacity (2.4 meq/g), less Co, and a larger equivalent Dy/Nd ratio (1.9).

The column (am-ZrP) breakthrough points of Co, Nd, and Dy were observed at 20, 125, and 135 BV, respectively (Table 10, Fig. 20b). The Co desorption from loaded column was indirectly observed by
the Co concentration, which exceeded approximately 20% to that of the initial feed. Thereafter, a desorption of Nd also took place at 190 BV. This was caused by the exchange of Nd to Dy, since Dy has the strongest affinity on am-ZrP.

**Figure 20.** Breakthrough curves of Co, Nd, and Dy with initial concentrations of 1.0 mmol/L equimolar nitrate solution. a) α-ZrP at pH 2.5, b) am-ZrP at pH 1.8, c) am-ZrP/PAN at pH 1.8.

Even though the am-ZrP/PAN composite beads are of relatively large size (approximately 2 mm in diameter), the Co, Nd, and Dy loading equivalent percentages closely resembled these values of equivalent percentages when using the powdery am-ZrP as ion exchanger (Table 10). The higher concentration of Co between 8 to 20 BV and subsequently Dy after 20 BV was observed, indicating the same preferred order (Dy>Nd>Co) as pristine am-ZrP. Moreover, it is worth noting that the value of equivalent ratio (2.0) between the loaded amount of Dy and Nd was better than that of the powdery am-ZrP (1.9).
Table 10. Data of the column loading experiments under the loading solution of 1.0 mmol/L equimolar Co, Dy, and Nd

<table>
<thead>
<tr>
<th>Materials</th>
<th>pH of feed</th>
<th>Breakthrough point (BV)</th>
<th>Loaded amount (meq/g)</th>
<th>Equivalent-%</th>
<th>Equivalent ratio (Dy/Nd)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>Co</td>
<td>Nd</td>
<td>Dy</td>
<td>Co</td>
</tr>
<tr>
<td>α-ZrP</td>
<td>2.5</td>
<td>2</td>
<td>8</td>
<td>8</td>
<td>0.01</td>
</tr>
<tr>
<td>Am-ZrP</td>
<td>1.8</td>
<td>20</td>
<td>125</td>
<td>135</td>
<td>0.02</td>
</tr>
<tr>
<td>Am-ZrP/PAN</td>
<td>1.8</td>
<td>2</td>
<td>20</td>
<td>22</td>
<td>0.03</td>
</tr>
</tbody>
</table>

Equivalent ratio is the ratio of metals uptake (meq/g).

4.3.2 Metal elution

We wanted to have single metal elution from the single-column study. For the α-ZrP packed column, HNO₃ at different concentrations and mixed with H₃PO₄ was tested for elution of loaded metals. With a fully loaded column it was not possible to have separate metal elution bands. To see clear elution bands of a single metal, the degree of metal loading must be considered.

To achieve better separation, a multistep gradient elution process was employed for the separation study by stepwise variation of the elution solution (HNO₃) concentration [123]. The am-ZrP material, with its more favourable ion-exchange properties, was employed to conduct the column separation for the Co, Nd, and Dy mixture. Decreasing the metal loading% to 8%, three separate elution bands were obtained with only a slight overlap from 25 to 32 BV (Figure 21a). The outlet order of metals (Co>Nd>Dy) was reversed with the adsorption order. Under these conditions, pure Co, Nd, and Dy can be produced (Figure 21a).

To set more realistic separation data, higher metal concentrations were tested using am-ZrP/PAN. A higher concentration solution totalling 1.2 g/L (approximately 10.7 wt% Co, 41.1 wt% Nd, and 48.2 wt% Dy) was used as the feed. Also the effect of higher temperature was tested. Thus, the gradient elution using 0.1, 0.2, 0.5 and 1 M HNO₃ at 50°C was performed with the column loaded at approximately 10%. Compared with the initial concentration, the concentrations of Co, Nd, and Dy
in the outlet were significantly enhanced with the gradient elution and the possibility to enrich Co by this process is seen possible (Figure 21b).

Finally, a simulated magnet leachate was employed as feed to test the suitability of metal separation in a practical setting using the am-ZrP/PAN beads. A total metal concentration of 7.6 g/L (1.4 wt% Co, 9.3 wt% Dy, and 89.3 wt% Nd) was prepared based on previous reports [47, 124]. Although the gradient elution method was utilized to elute metals at 50°C from an approximately 10%-loaded column (Figure 21c), the separation and recovery of Nd and Dy is extremely difficult due to the high Nd concentration in the feed and the higher selectivity of Dy over Nd.

![Figure 21. Gradient elution curves of Co, Nd, and Dy in the initial feed at pH 1.8 in 1 BV/h. a) The am-ZrP (approximately 8% degree loading) in feed at 1.0 mM at 23°C. b) am-ZrP/PAN (approximately 10% degree loading) in 1.2 g/L feed at 50°C. c) The am-ZrP/PAN (approximately 10% degree loading) in simulated magnet leachate (7.6 g/L) at 50°C.]

4.3.3 Stability of the material in regeneration

![Figure 22. XRD pattern of fresh and used am-ZrP/PAN beads for Run 3 and Run 10 in the column.]

**Figure 22.** XRD pattern of fresh and used am-ZrP/PAN beads for Run 3 and Run 10 in the column.
In general, the α-ZrP and am-ZrP materials are stable in acidic solution as they are synthesized from strongly acidic precursors. The decrease of interlayer space 1.0 Å and 1.4 Å was observed from am-ZrP/PAN after Run 3 and Run 10 compared with the unused one (Figure 22). These results suggest that the intercalated DMF in the layers was gradually released during the column process. The XRD pattern displayed the layers was possible to gradually shrink in the using process.
5 Conclusions

In this work, α-ZrP (Zr(HPO₄)₂ ⋅ H₂O) and am-ZrP (Zr(H₂PO₄)₀.₁₇ (HPO₄)₁.₇₈ (PO₄)₀.₀₉ ⋅ 0.₉₆H₂O) were synthesized and tested for the ability to recover and separate Co, Nd, and Dy metal ions. Crystalline α-ZrP showed a low sorption amount, which suggests that the exchanging metals cannot diffuse through the interlayer cavity in large extent. The sorption preference order Dy>Nd>>Co of α-ZrP and am-ZrP was obtained from batch ion-exchange experiments. We found that both materials had reasonable total metal capacity for Co, Nd, and Dy from equimolar solution but Co was not preferred. Am-ZrP had a higher selectivity and a higher sorption capacity when compared with that of α-ZrP or even the commercial classic Chelex 100 resin.

A gradient elution method proved to be an efficient process to achieve almost complete separation of metals using am-ZrP as ion-exchanger. Column loading at low degree has been shown to provide sufficient space for separate metal bands to develop in the elution process. Pure metal effluents were obtained by the gradient elution method using HNO₃ at different concentrations at room temperature. After obtaining promising results, larger lab-scale column separation was tested by using PAN-encapsulated am-ZrP composite in order to avoid possible operational problems associated with powdery am-ZrP.

X-ray tomography demonstrated a good spatial distribution of ion-exchange active component am-ZrP in the polymer matrix. Based on the batch experiments, the am-ZrP/PAN composite showed almost 50% higher sorption when compared to pristine am-ZrP. It appeared that the expansion of interlayer due to DMF intercalation enabled metals entering the cavity of the interlayer, resulting in increased uptake. This work demonstrated that it is possible to obtain fractions of Co with significantly improved purity relative to the simulated NdFeB magnet leachate through the single column form. However, it was not possible to achieve complete Nd and Dy separation due to the high Nd concentration in the simulated leachate solution and the higher Dy selectivity of the material.

Future work should consider the following two perspectives. From a material development perspective, enhanced uptake is suggested due to the intercalation of DMF. This provides a means for future study on intercalation chemistry and inorganic-organic hybrid materials.

From a process design perspective, even though it was difficult to achieve complete separation for the simulated leachate by single column, using multiple columns or a continuous simulated moving bed could be promising for future purification study.
Appendix

1 Methods for structural study

**X-Ray Powder diffraction**

X-Ray Powder diffraction (XRD) is primarily used for identification of crystalline material [125]. From these measurements, α-ZrPs can be identified by its unique diffraction peaks, such as (002), (110), and (112) plane. The interplanar spacing ($d_{hkl}$) can be calculated by Bragg’s law (Equation 13).

$$\lambda = 2d_{hkl} \sin \theta \quad (1)$$

The XRD patterns were obtained by utilizing a Philips PW 3710 X-ray diffractometer operated with Cu-Kα ($\lambda=1.542$ Å) radiation (40 kV and 40 mA). The data were recorded from 5° to 70° in 2θ (step length 0.02°, counting time 0.5 s per step).

**Fourier transform infrared spectra**

Fourier transform infrared (FT-IR) spectra was used to identify the chemical group in the material by means of detecting the infrared absorption caused by certain bond vibrations [126].

Here, FT-IR spectra was employed to identify the POH group from the synthesized and relevant materials. FT-IR spectra were acquired using a Perkin Elmer spectrum one FT-IR spectrometer in ATR (attenuated total reflection) mode. The measurement range is from 600 to 4000 cm\(^{-1}\) at a resolution of 4.0 cm\(^{-1}\).

**Thermogravimetric analysis**

Thermogravimetry (TG) is a technique to measure the changes in mass of a material against temperature changes in a specific atmosphere. The measurements are often simultaneously processing combined thermogravimetry and differential thermal analysis [127].

TG analysis was performed using a Mettler Toledo TG850. Samples were normally heated to 800°C or up to 1000°C when needed. The heating rate was programed as 5°C/min, as the measurement atmosphere depends on the measurement of program setting in nitrogen or air flow.

**Solid-state \(^{31}\)P magic angle spinning nuclear magnetic resonance**

Solid-state \(^{31}\)P magic angle spinning nuclear magnetic resonance (\(^{31}\)P MAS NMR) spectra were used for detecting phosphorous in different chemical environments. The different phosphate groups in am-ZrP can therefore be distinguished. The ratio of these phosphate groups can be obtained based on the
peak deconvolution results. The molecular formula can then be calculated from the combination of the ratio of these phosphate groups and the TG results.

$^{31}$P MAS NMR spectra were recorded from a Bruker Avance III NMR spectrometer (500 MHz, 4 mm H/X/Y MAS probe). The sample was placed into a zirconia rotor (4 mm). The sample of $^{31}$P MAS NMR spectra recorded with a MAS rate (12 KHZ), 90° pulse (77 KHZ RF), and a 100-s recycle delay (64 scans).

2 Methods for morphology and spatial distribution study

Scanning electron microscopy

Scanning electron microscopy (SEM) was utilized to record the surface and the cross-section morphology of the synthesized samples.

The sample images were collected using a Hitachi Hi-Tech S-4800 field-emission scanning electron microscope (FESEM) by sputtering for deposition of a 3 nm-thick layer of Pd-Au.

For the size distribution investigation of the am-ZrP/PAN composite beads, samples without coating were imaged using a Hitachi SU3500 scanning electron microscope. In addition, Malvern Morphologi G3 software was employed to analyse the morphology of 199 beads from an SEM image.

X-ray tomography

X-ray tomography was used to reveal a structure’s internal attributes by generated 3-dimensional imaged volumes from 2-dimensional X-ray image slices [128].

For spatial distribution of am-ZrP in an am-ZrP/PAN bead, the bead was recorded using a GE phoenix v\text{\textregistered}ome|x s 240. An optional nanofocus tube (180 kV/15 W, 90-kV, 300-μA) was used in a 27-W tube power. The sample was obtained with an isotropic 1.33-μm resolution/voxel size. 2700 projections (exposure time: $2 \times 4000$ ms) were performed using a 4000-ms skip at each angle for 9 h total.

3 Method for metals determination

Microwave plasma-atomic emission spectrometer

An Agilent 4200 microwave plasma-atomic emission spectrometer (MP-AES) was utilized to determine the metal concentrations. The OneNeb nebulizer and a double-pass cyclonic spray chamber was equipped for sample introduction.

The analytical cycle was a 30-s rinse with nitric acid (v/v 5%) and a subsequent 30-s sample uptake with pump speed 15 rpm. The internal standard and the buffer ion was used by adding 0.1 mL of La
(1000 mg/L) and 0.1 mL Cs (100 000 mg/L) into the 9.8 mL diluted samples. The detection wavelength of Co, La, Nd, and Dy was set at 340.512, 394.910, 430.358 and 353.171 nm, respectively.


