Branched alkylphosphinic and disubstituted phosphinic and phosphonic acids: effective synthesis based on α-olefin dimers and applications in lanthanide extraction and separation†

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A number of mono-alkylphosphinic acids RCH₂CH₂CH(R)CH₂P(O)(H)OH 8–12 were obtained via interaction of α-olefin vinylidene dimers RCH₂CH₂C(R)≡CH₂ 3–7 [R = n-butyl; 3, 8; isobutyl; 4, 9; n-octyl; 5, 10; isopropyl; 6, 11; cyclohexyl; 7, 12] with H₃PO₂ in an isopropanol medium at 90 °C. Hydrophosphinylation of 3 by 8 or PhPO₂H₂ at 140 °C resulted in disubstituted acids 13 and 14. Alkylmethylphosphinic acids 15–19 and functionalized alkylphosphinic acids 20–22 have been synthesized via interaction of silyl ethers of mono-alkylphosphinic acids 8–12 with Mel, [2-chloromethyl]pyridine, acrylic acid or acrylamide. Non-catalytic hydroalumination of 3 with subsequent interaction with PCl₃ and oxidation with SO₂Cl₂ led to the phosphonic acid anhydride, which was further used to obtain alkylphosphonic acids 23 and 24. It has been found that phosphinic acids 8–12 surpass di(2-ethylhexyl)phosphoric acid (extractant P204) in non-selective Ln⁶⁺ extractability (Ln = La, Pr, Nd, Dy and Lu). Significantly higher selectivity for heavy lanthanide extraction (Ln = Dy, Lu), compared to that of P204, is achieved by using a minimal excess of disubstituted phosphonic acids 13–24. Acid 13, which contains two branched substituents, demonstrated unique selectivity in the extraction of Lu in the presence of the other lanthanides. Dialkylphosphinic acids 16–18 and alkylarylphosphinic acid 14 possess a significant potential for the Pr/Nd pair separation. Taking into account the availability of 8–24 and the structural variability of the initial α-olefin dimers 3–7, the newly obtained compounds represent a promising group of rare-earth element extractants.

Introduction

Rare-earth elements (REEs) are widely used in the production of permanent magnets, lamp phosphors, NiMH batteries,¹ in the preparation of effective and irreparable catalysts for hydrocracking, diene polymerization² and in other industrial processes. The extraction and separation of REEs, therefore, is a relevant technical issue.³,⁴ Various extraction methods have been productively used for a long time for this purpose.¹⁵–¹⁸ By now, a great number of compounds have been studied as possible extractants of REEs.⁵ Di(2-ethylhexyl) phosphoric acid (extractant P204, HDEHP or D2EHPA, Scheme 1) is still being used as the primary extractant in industry. However, the main drawbacks of this readily available compound are a low adjacent separation factor, relatively low acidity for REE stripping, as well as low hydrolytic stability. Phosphonic and phosphinic acids are more hydrolytically stable and effective extractants when

Scheme 1 Currently used and prospective REE extractants.
compared with P204. Among these compounds are mono-2-ethylhexyl ester of 2-ethylhexyolphosphonic acid (P507 or PC88A) and bis(2,4,4-trimethylpentyl) phosphonic acid (P272, Scheme 1), which have also been commercially developed.\(^1\) The properties of alkylphosphinic and phosphonic acids R–P(O)(OH)–FG depend on the structure of alkyl substituent R. It has been recently shown that acid 1 which contains β-branched alkyl fragment (Scheme 1), shows the maximum extraction efficiency among isomeric diocytlyphosphinic acids.\(^9\)

Recently\(^1\) we have developed an effective and general method of α-olefin dimerization, which allows to selectively synthesize vinylidene dimers \(\text{RCH}_2\text{CH}_2\text{C}(\text{R})==\text{CH}_2\) containing a linear or branched alkyl R group in high yields. We proposed that a hydrophosphinilation reaction of these compounds can be carried out under controlled conditions with a formation of β-branched mono-alkyl phosphinic acids, which could be used to produce a whole family of new extractants with regulated lipophilicity, namely non-symmetrical disubstituted phosphinic and phosphonic acids (Scheme 1).

The current work is aimed at developing synthetic approaches to mono-alkylphosphinic and disubstituted phosphonic acids prepared from α-olefin dimers \(\text{RCH}_2\text{CH}_2\text{C}(\text{R})==\text{CH}_2\), as well as the study of the REE extraction efficiency of the newly-obtained compounds.

**Experimental**

**Materials and methods**

All of the synthetic experiments were conducted under an argon atmosphere. Solvents used in synthetic procedures were distilled, stored under argon, and degassed under reduced pressure with stirring in the reaction vessels prior to adding reagents (for details on reagent and solvent preparation, see the ESI†). CDCl\(_3\) (Cambridge Isotope Laboratories, Inc., D 99.8%) was used as purchased. The \(^1\)H and \(^13\)C NMR spectra were recorded on a Bruker AVANCE 400 spectrometer (400 MHz) at 20 °C. The chemical shifts are reported in ppm relative to the solvent residual peaks. Elemental analysis (C, H) was performed on a Perkin Elmer Series II CHNS/O Analyser 2400. pH values of the aqueous phase were measured with a digital pH meter calibrated daily with 4.01 and 7.00 standard buffers. The concentration of rare earth ions in the water phase was determined by inductively coupled plasma atomic emission spectroscopy (ICP-AES) with the Agilent 7500C G 315SB instrument after proper dilution. When determined concentrations of lanthanides in water phase were very low, the ICP-AES analyses were repeated without dilution.

**Synthetic procedures**

**α-Olefine dimerization**

*General procedure.* Trisobutylaluminium solution (1 M TIBA in hexane, 5 mL, 5 mmol) and [O(SiMe\(_2\)C\(_p\))\(_2\)]ZrCl\(_2\) (106 mg, 0.25 mmol) were added at 60 °C to liquid monomer (1 mol). After 20 min of stirring, Et\(_2\)AlCl (1 M in hexane, 0.5 mL, 0.25 mmol) and MMAO-12 (1.05 M in toluene, 2.4 mL, 2.5 mmol) were added. After 6–8 h of stirring, the mixture was cooled to room temperature, 5 mL of methanol and 2 mL of water were added. After 10 min of stirring, Na\(_2\)SO\(_4\) (5 g) was added and the mixture was stirred for an additional 10 min. After that, the mixture was filtered through a layer of silica and distilled under reduced pressure. Product yields and characteristics, including \(^1\)H and \(^13\)C NMR spectra, elemental analysis and GC data, are given in the ESL†.

**Hydrophosphorylation of α-olefin dimers: preparation of 8–12**

*General procedure.* A mixture of α-olefin dimer 3–7 (66 mmol), 50% aq. \(\text{H}_2\text{PO}_4\) (14.5 mL, 264 mmol) and i-PrOH (80 mL) was degassed under reduced pressure, filled with argon and heated to 90 °C. AIBN was added with vigorous stirring in 7 portions by 100 mg at 30 min intervals. The mixture was cooled to room temperature and diluted by pentane (100 mL). The organic layer was washed by water (3 × 100 mL), dried over Na\(_2\)SO\(_4\) and evaporated. The residue was purified by gradient column chromatography (silica 60–200, por. 40) using CH\(_2\)Cl\(_2\)–MeOH mixture (from 100 : 1 to 1 : 1 by volume). Product yields and characteristics, including \(^1\)H and \(^13\)C NMR spectra, elemental analysis and GC data, are given in ESL†.

**Functionalization of mono-alkylphosphinic acids**

*Methyl(alkyl)phosphinic acids 15–19, general procedure.* The mixture of mono-alkylphosphinic acid (20 mmol), Me\(_2\)SiCl (5.43 g, 50 mmol) and CH\(_2\)Cl\(_2\) (20 mL) was cooled to 0 °C. Triethylamine (5.06 g, 50 mmol) was added. The mixture was stirred for 2 h. After that, the mixture was cooled to 0 °C and methyl iodide (5.75 g, 50 mmol) was added. After 16 h at room temperature, the mixture was diluted by pentane (30 mL). Organic layer was washed by water (3 × 100 mL), dried over Na\(_2\)SO\(_4\) and evaporated. The residue was purified by gradient column chromatography (silica 60–200, por. 40) using CH\(_2\)Cl\(_2\)–MeOH mixture (from 100 : 1 to 1 : 1 by volume). Product yields and characteristics, including \(^1\)H and \(^13\)C NMR spectra, are given in ESL†.

**Extraction experiments**

The lanthanide mixture solutions were prepared from LnCl\(_3\) (1 mol). Two phases were vigorously stirred for 12 hours. The
Results and discussion

Synthesis of alkylphosphinic and alkylphosphonic acids

Compounds 3–7 were obtained via selective catalytic dimerization of α-olefins using our recently developed procedure using the zirconocene pre-catalyst 2 (Scheme 2) activated by TIBA and MAO.10,11 Compounds 3–5 and 7 are isomerically pure vinylidene olefins. Hydrocarbon 6 contains insignificant amounts (~2%) of reduction product (for experimental details and product characterization, see the ESI†).

Vinylidene dimers of olefins 3–7 are promising precursors for preparing functional derivatives with regulated lipophilicity.12–16 The idea of vinylidene olefin functionalization relies on the hydrophosphinilation reaction. This reaction is a general method of synthesis of phosphinic acids and their derivatives.17–18 Radical hydrophosphinilation initiated by peroxides19–21 or AIBN,22,23 needs high temperature and long reaction time. As a result, the product is usually a mixture that includes mono- and diphasphinic acids; the process has low regioselectivity. Catalytic hydrophosphinilation,13,24,26 is not very effective in the case of vinylidene olefins. A recently developed method of phosphinic acid synthesis relies on initiation by microwave irradiation. In the absence of a radical initiator, this method demonstrated low efficiency in the synthesis of mono-alkylphosphinic acids  from vinylidene olefins.27

Starting from vinylidene olefins of various structures, a series of symmetric dialkylphosphinic acids, including compound 1, was obtained in the presence of AIBN. Mono-alkylphosphinic acids were synthesized only in moderate yields.9

We proposed that in the case of highly reactive vinylidene olefins 3–7 thermally initiated hydrophosphinilation with H$_2$PO$_2$ can be carried out in two steps: (1) synthesis of monoalkylphosphinic acids under relatively mild conditions, and (2) transformations of mono-alkylphosphinic acids into dialkylphosphinic acids under more rigorous conditions without MW-irradiation. The choice of the solvent is of fundamental importance in this case. It has been determined earlier22,23 that the presence of strong acids significantly accelerates radical hydrophosphinilation. This method has been effectively used in cases of terminal olefins. However, it is not effective for vinylidene dimers 3–5 due to a side cationic oligomerization reaction. We used a series of solvents (DMF, dioxane, DME, isopropanol) and determined that in the protonic solvent, 3PrOH, hydrophosphinilation of 3 proceeds smoothly under mild conditions (90 °C) and allows to obtain mono-alkylphosphinic acid 8 in 80% yield and high regioselectivity. For α-olefin dimers 4–7, a three hour reaction allowed us to obtain over 90% conversion of initial hydrocarbons and 80–90% isolated yields of branched alkyl phosphinic acids 9–12 (Scheme 3).

By using compound 3, we also demonstrated the possibility of carrying out a hydrophosphinilation reaction in two steps. The acid 8, obtained under relatively mild conditions, was reacted with dimer 3 under substantially more harsh conditions at 145 °C with addition of the initiator by small portions, which resulted in formation of dialkylphosphinic acid 13 (Scheme 3). The reaction was carried out with a high regioselectivity and satisfactory isolated yield. In contrast, interaction of 3 with phenylphosphinic acid proceeded with moderate regioselectivity. Product 14 was obtained with a yield of 42%.

The newly obtained mono-alkylphosphinic acids 8–12 were used in the synthesis of asymmetrical phosphinic and phosphonic acids. We used a method developed earlier by Regan and coll. in order to introduce alkyl substitutes and functional groups.28–33 This method includes generation of silyl ethers through interaction of phosphinic acids with TMSCl in the presence of bases. The silyl ethers act as P-nucleophilic particles and promote alkylation or Michael addition in situ. We introduced mono-alkylphosphinic acids 8–12 into a reaction with TMSCl and methyl iodide, which produced a series of methyl[ -]alkylphosphinic acids 15–19 in high yields. The interaction of 8 with 2-(methyl chloride)pyridine resulted in product 22. The applicability of the Michael reaction was demonstrated in the case of interaction of 8 with acrylic acid and acrylamide, providing compounds 20 and 21, respectively (Scheme 3).

Ethers of alkylphosphinic acids 23 and 24 were obtained from 1-hexene dimer 3 through a multistage reaction. At the first stage, via interaction with triisobutylaluminium (TIBA), an organoaluminium derivative was obtained. This reaction is traditionally carried out in the presence of [η$^5$-C$_5$H$_5$ZrCl$_2$], however, in the case of 3, there is no need for a catalytic process to take place: an autooxidation test showed that 3 quantitatively transformed into tris(2-butylcyclohexyl)aluminium after 40 h at 140 °C. The reaction of the organoaluminium derivative with an excess of PCl$_3$ led to compound C$_{12}$H$_{27}$PCl$_3$ of ca. 75% purity. The latter, upon treatment with SO$_2$Cl$_2$, was reacted with 2-ethylhexanol or 2,6-diisopropylphenol. The products of these reactions were compounds 23 and 24, respectively (Scheme 3).
All newly obtained compounds were purified by gradient column chromatography. Their structures and purity were confirmed by spectral methods and elemental analysis (see ESI†) prior to studying them as lanthanide extractants.

Lanthanide extraction

Standard methods of extractability studies of organic compounds are based on determining the element concentrations in organic and aqueous phases after reaching equilibrium. According to published data,1,7,34,35 dialkylphosphoric, dialkylphosphonic and dialkylphosphinic acids exist in hydrocarbon media as dimers (HL)2. The extraction equilibrium can be described by eqn (1).†

\[
\text{Ln}^{3+\text{aq}} + 3(\text{HL})_2\text{org} \rightleftharpoons [\text{Ln}(\text{HL})_2]_{\text{org}} + 3\text{H}^+\text{aq}
\]  

(1)

A number of conventional parameters are used for a qualitative analysis of extractant properties: distribution ratio (D), extraction efficiency (E) and separation factor (β) (See ESI†). The extraction equilibrium constant (Kex) for each lanthanide can be represented by the eqn (2).

\[
K_{\text{ex}} = \frac{[\text{Ln}(\text{HL})_2]_{\text{org}}[\text{H}^+]_{\text{aq}}}{[\text{Ln}^{3+\text{aq}}]([\text{HL}]_2)_{\text{org}}}
\]  

(2)

Rare-earth elements extraction ability of the synthesized phosphinic and phosphonic acids was studied for La, Nd, Pr, Dy, and Lu. Di(2-ethylhexyl)phosphonic acid (P2O4) was used as a benchmark. All extraction experiments were carried out at initial concentration of each lanthanide in aqueous phase of 0.002 M (with the total lanthanide concentration [La]0 = 0.01 M) and at [NaOH] = 1 M at equal volumes of organic and aqueous phases.

In order to determine the conditions that would reveal selectivity of the used extractants, two parameters were varied: initial pH of the aqueous phase and initial extractant concentration in organic phase [HL]0. First, we compared lanthanide extractability of mono-alkyl phosphonic acids 8, 10, and P2O4 at [HL]0 = 0.200 M and at initial pH = 2.00 and 3.00. The extraction efficiency of P2O4 was expectedly lower for lighter lanthanides at pH = 2. The acids 8 and 10 were more effective towards extraction of all lanthanides than P2O4 regardless of initial pH of aqueous phase (Table 1: entries 1–4, 6, 8, see extraction efficiency), not demonstrating any selectivity toward heavier lanthanides. For example, the separation factor values βDy/Nd for P2O4 (βDy/Nd = 225 and 262) and compounds 8, 10 (βDy/Nd = 5.4–15.6) differ by more than an order of magnitude (for details, see ESI†).

Dependency of lanthanide extraction efficiency from [HL]0 was estimated at initial pH = 3 of the aqueous phase, using the more sterically crowded alkylphosphinic acid 8. Expectedly, lanthanide extractability did not change significantly at lowering the phosphinic acid concentration by 29%. Selectivity toward heavier lanthanides (Dy, Lu) increases greatly upon further decrease of [HL]0 from 0.142 M to 0.067 M (Table 1, entries 9 and 10). This behavior correlates with eqn (1)

\[
\begin{align*}
\text{La} & : 100.00 \\
\text{Nd} & : 100.00 \\
\text{Pr} & : 100.00 \\
\text{Dy} & : 100.00 \\
\text{Lu} & : 100.00 \\
\end{align*}
\]

Table 1 Data for lanthanide extraction using P204, 8 and 10 at different pH and extractant concentrations ([La] = 0.01 M)

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a Published data. Distribution ratio and separation factors have been calculated based on extraction efficiency. D = E([Vaq]/[Vorg])/(1 − E), Vaq = Vorg, βLant/Lanet = Dlant/Dlanet. b Unreliable separation factor values have been omitted.
Table 2 Data for lanthanide extraction using P204, B, 10, 13–24. [Ln]₀ = 0.01 M, [HL]₀ = 0.067 M, initial pH = 3.00

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Unreliable separation factor values have been omitted.

Describing the equilibrium of lanthanide complex formation—the Ln³⁺ ion forms a complex with six molecules of hexafluorophosphinate.

In order to compare extraction ability of newly obtained compounds, further experiments were carried out at [HL]₀ of 0.067 M (the ratio of [HL]/[Ln]₀ = 1/6.7), which provided ca. 11% molar excess of the extractant (see eqn (1)). Under these conditions, the extraction characteristics of 8 barely differed from those of 10. Both mono-alkylphosphinic acids demonstrated slightly worse selectivity towards late lanthanides (Dy, Lu) in comparison with P204, β_Dy/Nd = 124 (8) 39.5 (10) and 279 (P204). Therefore, lanthanide extraction with compounds 9, 11 and 12, which are structurally similar to 8 and 10, has not been studied. Comparative experiments regarding extraction of La, Nd, Pr, Dy, and Lu at pH = 3 and initial concentrations [Ln]₀ = 0.01 M and [HL]₀ = 0.067 M were carried out for compounds 13–24. The results are summarized in Table 2.

As it has been noted while discussing preliminary experiments (Table 1), mono-alkylphosphinic acids 8 and 10 are effective non-selective extractants for all examined lanthanides, possessing significantly lower selectivity compared to even P204. Substitution of the hydrogen atom (8, 10) with a methyl group (15, 17), and further, by a 2-butyloctyl fragment (13), abruptly lowers the extraction efficiency (see ESIF) and lowers Ig D of all lighter lanthanides (La, Pr, Nd) (Fig. 1). This allows us to consider phosphinic acids containing branched alkyl and methyl substitutes as selective extractants of heavy lanthanides. The extraction properties of compounds 15–19 are similar to each other (Fig. 2). The substitution of n-alkyl fragment R (15, 17) with an isoalkyl one (16, 18) leads to an increase in separation efficiency for Nd/Pr pair. The best separation of light (La, Nd, Pr) and heavy (Dy, Lu) lanthanides is achieved with 17, containing a more bulky alkyl fragment based on a 1-decene dimer.

It should also be noted that the replacement of the methyl substituent with a β-branched alkyl fragment in compound 13 leads to a unique extraction selectivity for the Dy/Lu pair (Fig. 1). This selectivity makes 13 markedly different from compound 1 described earlier. This allows us to examine 13 as a promising selective extractant for Lu and heavier lanthanides from REE mixtures.

The extraction ability of functionalized di-substituted alkylphosphinic acids 20–22, phosphonic acid 23 and alkylarly phosphonic acid 14 is qualitatively close to that of P204 (Fig. 3). At the same time, the characteristics of extractant 24 are similar.
potential for separation of Pr from Nd due to higher extraction efficiency (42.9% for Pr and 55.2% for Nd) at $\beta_{\text{Nd/Pr}} = 1.64$.

Furthermore, 14 possessed the highest value of $\beta_{\text{Dy/Nd}} = 564$ among all studied acids.

Within the series of synthesized phosphinic acids, 14 represents a sole example of an entirely unexplored class of chemical compounds – (branched alkyl)(aryl)phosphonic acids. Our experiments have demonstrated that this extractant class deserves the closest attention and further studies.

By using eqn (2), we calculated the $\lg K_{\text{ex}}$ which lie in the intervals of 3.6 to +1.9 for La and −0.6 to +6.8 for Lu (for diagram, see ESI†). The values of $\lg K_{\text{ex}}$ for most compounds studied herein are higher by 3–6 orders of magnitude than those of dialkylyphosphinic acids of the ($\text{C}_3\text{H}_7$)$_2$POOH type, which explains their high non-selective extraction ability at $[\text{HL}]_0 = 0.2$ M. Compound 13, ($\text{C}_{12}\text{H}_{25}$)$_2$POOH, possessing the worst overall extraction ability, has $\lg K_{\text{ex}}$ values that are similar to those published earlier for dioctylphosphinic acids. The previously published $\lg K_{\text{ex}}$ values for di-[2,3-dimethylbutyl]-phosphinic acid are in good agreement with our data. With the exception of 20 and 24, there is a positive correlation between the $\lg K_{\text{ex}}$ values and the general extraction ability of the studied acids. For compounds 8, 10 and extractant P204, $\lg K_{\text{ex}}$ data that have been obtained at different initial values of $[\text{HL}]_0$ and $pH$ differ insignificantly. The differences can be explained by the fact that the equilibrium shown on eqn (1) does not take into account the formation of higher HL oligomers and other side reactions. However, as it has been shown earlier, these side reactions do not play a fundamental role in comparative studies of lanthanide extraction by organic compounds of the same type from aqueous solutions.

Conclusions

We have proposed herein an effective method of synthesis of branched alklyphosphinic and alklyphosphonic acids based on vinylidene dimers of $\alpha$-olefins. The synthetized phosphinic (8–22) and phosphonic (23, 24) acids have demonstrated a high lanthanide extraction potential. It has been determined that:

- Monosubstituted phosphinic acids 8 and 10 possess a higher non-selective lanthanide extractability in comparison with industrial extractant P204;
- The previously unknown alkylmethylphosphonic acids 15–19 are effective and selective extractants of heavy lanthanides (Dy, Lu).
- Dialklyphosphinic acid 13, containing two bulky branched alkyl substituents and being formally similar in structure to the commercial extractant P272 and to the described earlier dioctylphosphinic acids, demonstrates high selectivity for the “heavy” lanthanide separation within the pair Dy/Lu;
- Extractant 14, bearing phenyl and branched alkyl substituents, demonstrates a high potential for separation of “light” lanthanides from La and for lanthanide separation within the Pr/Nd pair. It significantly surpasses P204 by both parameters.

Therefore, the newly synthesized phosphinic and phosphonic acids may be considered as promising extractants for the separation of rare-earth metals. New structural types of...
asymmetrical acids, especially (branched alkyl)(aryl)phosphinic acids that are derivatives of 14, are being considered for further research.

Acknowledgements

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Notes and references