Isoconversional kinetic analysis applied to five phosphonium cation-based ionic liquids

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Abstract

Thermal degradation of five phosphonium cation-based ionic liquids ([P₆₆₆₁₄][BEHP], [P₆₆₆₁₄][(iC₈)₂PO₂], [P₆₆₆₁₄][NTf₂], [P₄₄₄₁₄][DBS] and [P₄₄₄₂][DEP]) was studied using dynamic methodology (25 to 600 °C at 5, 10 and 20 °C/min) in both inert (nitrogen) and reactive (oxygen) atmospheres. In addition, isothermal experiments (90 min at 200, 225 and 250 °C) were carried out with [P₆₆₆₁₄][(iC₈)₂PO₂]. Results indicate that thermal stability is clearly dominated by the coordination ability of the anion, with [P₆₆₆₁₄][NTf₂] outperforming the other ones in both pyrolytic and oxidising conditions. Although the thermal degradation mechanism is affected by atmospheric conditions, the degradation trend remains practically constant. As the dynamic methodology usually overestimates the long-term thermal stability, an isoconversional methodology is better for predicting the long-term thermal stability of these ionic liquids in order to be used as base oil or additive in lubricants formulation. Finally, the model-free methodology can predict at lower costs the ILs performance in isothermal conditions.

Highlights

- Weakly coordinating anions provoke good thermal stability of ILs.
- $[P_{66614}][NTf_2]$ and $[P_{44414}][DBS]$ are the most thermally stable ILs in this study.
- Isoconversional methods are better than dynamic ones for long-term thermal studies.
- Activation energy behavior at low conversions is related to a single reaction.
- Model-free methodology can predict the ILs performance in isothermal conditions.

Keywords: ionic liquids; thermogravimetric analysis; isoconversional method; model-free kinetics

1. Introduction

Ionic liquids (ILs) are salts typically formed by the combination of large organic cations (commonly containing large alkyl chains) and a variety of anions (either inorganic or organic). In the last 15 years, the use of ILs has increased considerably due to their interesting chemical and physical properties, such as negligible vapour pressure, high thermal stability, low melting point, high solvating capacity, high thermal conductivity and wide electrochemical window [1–6]. These properties mean that ILs are excellent candidates for various applications, including their use as environmentally friendly solvents for replacing traditional volatile organic ones [7,8]. Currently, growing interest is focused on the use of ILs for high temperature applications, such as thermal fluids and electrolytes for solar cells and fuel cells [9,10]. Significant research has been focused on the thermophysical properties of imidazolium-based ILs [11,12], but there are only a few studies focused on a novel class of ILs based on phosphorous quaternary cations [13,14]. The phosphorus atom allows for bonding with different substituents, giving a great number of different structures and physicochemical properties [15,16].

One of the most important characteristics of ILs is their thermal stability. An appropriate estimation of this property is a key factor in many engineering applications, including lubrication [17]. Despite the fact that a significant number of works have been focused on the study of thermal degradation of ILs [18–25], only a few of them have used phosphonium based ILs [13,15,26]. In addition, thermal stability is often misleadingly reported to extend up to temperatures of more than 400 °C. This issue occurs if high heating rates (>10 K/min) are used in the temperature ramped thermogravimetric analysis (TGA) studies, overestimating the thermal stability of the analysed samples [22-25]. Due to the fact that most applications require long-term thermal stability, more reliable data is needed in this field. A full evaluation of the long-term thermal stabilities and thermal decomposition is necessary to identify global kinetic models. Nowadays, the knowledge of the kinetic parameters, such as reaction rate and activation energy, is one of the key factors for determining thermal stability. Some research has been focused on the determination of activation energies and kinetic models, describing the thermal degradation of some ILs. Most of these studies performed kinetic analyses based on isothermal experiments. In these cases, some isothermal experiments were measured at low temperatures (below onset temperature) and kinetic parameters were determined assuming zero-order processes [10,27]. Therefore, a prediction of the longterm stability of ILs based on the modelling of non-isothermal TG measurements enables to reduce significantly the time required to carry out the experiment, resulting in a considerable cost reduction [23].

Lubricant oils used in cooling engine parts are exposed to high and long-term thermal stress, which leads to thermal degradation (evaporation of volatiles) and polymerisation, forming a heavier fraction [28]. Volatility is the main parameter linked to oil consumption and, besides the Noack volatility test, the volatility of oils and additives have been measured using TGA [17]. A lot of researches have been published concerning the use of ionic liquids as lubricants since 2001 [29], proving their remarkable potential for this purpose [4,6,30-32]. At the beginning, imidazolium-based ionic liquids with [PF₆] and [BF₄] anions were widely used [33-35]. Despite their good tribological behaviour, the hydrolysis products of those anions are highly corrosive [36]. This fact provoked the introduction of new and more stable ILs based on [FAP] and [NTf₂] anions [37-53], which present enhanced tribological properties due to the formation of fluoride tribofilms, especially at high loads [54]. Recently, phosphonium cation-based ILs have been introduced in this field because of their growing commercial availability and good tribological performance [26,55-62]. Finally, taking into account the importance of thermal stability in the lifetime of lubricants and additives, the determination of thermal models has become an interesting research topic from scientific and technical points of view.

In this work, an alternative approach using an isoconversional (model-free) methodology is proposed to derive the long-term stabilities of five phosphonium cation-based ILs: trihexyltetradecylphosphonium bis(2-ethylhexyl)phosphate ($[P_{66614}][BEHP]$), trihexyltetradecylphosphonium bis(2,4,4-trimethylpentyl) phosphinate ($[P_{66614}][(iC_8)_2PO_2]$), trihexyltetradecylphosphonium bis(trifluoromethylsulfonyl) imide ($[P_{66614}][NTf_2]$), tributyltetradecylphosphonium dodecylbenzenesulfonate ($[P_{44414}][DBS]$) and tributylethylphosphonium diethylphosphate ($[P_{4442}][DEP]$). Previous papers [27,63] have determined that both model-free and isothermal model-fitting methodologies are in good agreement. Using isoconversional methods, kinetic parameters can be derived without any assumption of the mechanism. Furthermore, an isoconversional rate expression determined using non-isothermal (dynamic) data can predict isothermal reaction data.

2. Experimental

2.1. Ionic liquids

Chemical and structural descriptions of the five ILs used, ([P₆₆₆₁₄][(iC₈)₂PO₂], [P₆₆₆₁₄][BEHP], [P₆₆₆₁₄][NTf₂], [P₄₄₄₁₄][DBS] and [P₄₄₄₂][DEP]), are shown in Table 1. These liquids were provided by Ionic Liquid Technologies GmbH, chosen from an available family of ILs (phosphonium cation-based)

that is currently gaining consideration because of their possible use in lubricant applications (as base fluid or additive).

Table 1. Chemical and structural descriptions of the ILs used in this work.

IUPAC and short name	Empirical formula	Purity (%)	Molecular weight	Chemical structure
$\label{eq:continuous} Trihexyltetradecylphosphonium \\ bis(2,4,4-trimethylpentyl) \\ phosphinate \\ [P_{66614}][(iC8)_2PO_2]$	$C_{48}H_{102}O_2P_2$	95	773.27	
Trihexyltetradecylphosphonium bis(2-ethylhexyl)phosphate [P ₆₆₆₁₄][BEHP]	$C_{48}H_{102}O_4P_2$	98	805.29	
$Trihexyltetradecylphosphonium\\bis(trifluoromethylsulfonyl)\\imide\\[P_{66614}][NTf_2]$	$C_{34}H_{68}F_6NO_4PS_2$	98	764.01	0, 8, 0 0, 0 0
Tributyltetradecylphosphonium dodecylbenzenesulfonate [P ₄₄₄₁₄][DBS]	cylbenzenesulfonate C ₄₄ H ₈₅ O ₃ PS		725.18	
Tributylethylphosphonium diethylphosphate [P ₄₄₄₂][DEP]	$C_{18}H_{42}O_4P_2$	95	384.47	

2.2. Thermal analysis

Thermal analysis was performed in a Mettler Toledo TGA/SDTA 851 thermogravimetric analyser, operating in dynamic and isothermal modes under dry nitrogen and oxygen atmospheres (50 mL/min). Ionic liquid samples (10–12 mg) used as supplied by Ionic Liquid Technologies GmbH were placed in an open aluminium crucible without any previous treatment. Dynamic experiments were performed at

temperatures from 25 to 600 °C at three different heating rates (5, 10 and 20 °C/min). Isothermal experiments were conducted at three different fixed temperatures (200, 225 and 250 °C) during 90 min tests. Results were analysed using Mettler-Toledo STARe version software. The onset temperature (T_{onset}) is the intersection of the baseline weight and the tangent of the weight versus temperature curve as decomposition occurs. In addition, other temperatures were determined at 10% of mass loss, total mass loss, as well as the minimum of differential thermogravimetric (DTG) peaks.

2.3. Kinetic parameters and modelling

The mass loss of an IL under a thermogravimetric experiment may be the result of combining evaporation and/or thermal decomposition kinetics [23,64,65]. The rate of evaporation can be defined using eq. (1), as the mass transfer of the IL into the gas stream flowing over the crucible that contains the sample.

$$\frac{dm}{dt} = M_{IL}BA \frac{p_{vap}(T)}{RT} \tag{1}$$

where M_{IL} is the molar mass of the ionic liquid, B is the mass transfer coefficient, A is the surface area in contact with the gas phase and p_{vap} the vapour pressure. In addition, the rate of heterogeneous thermal decomposition can be described by eq. (2):

$$\frac{d\alpha}{dt} = k(T)f(\alpha) \tag{2}$$

In eq. (2): t is time, k(T) is the temperature-dependent constant and $f(\alpha)$ is a function called the reaction model, which describes the dependence of the reaction rate on the extent of conversion, α .

The rates of evaporation and decomposition increase almost exponentially with temperature, being the decomposition rate more sensitive than the evaporation rate. Thus, at both high heating rates and temperature ranges, thermal decomposition may control the overall mass loss (and vice versa) [64]. Assuming that heating rates used in the TGA experiments carried out in this study can be considered as "high" (>4 K/min), the kinetic process is mainly controlled by thermal decomposition within the whole range of conversion, and eq. (1) could be negligible. In a TGA experiment in which the mass variation versus temperature is obtained, the extent of reaction is calculated using the following equation:

$$\alpha = \frac{m_i - m_t}{m_i - m_f} \tag{3}$$

where m_i and m_f are the initial and final masses, respectively, and m_t is the mass at a specific time, t.

There are several methods for analysing thermal decomposition kinetic data [66]. These methods may be classified according to the experimental conditions selected and the mathematical analysis performed. Experimentally, both isothermal and non-isothermal methods are employed. The mathematical approaches employed can be divided into model-fitting and isoconversional (model-free) methods.

In model fitting methodologies, the term $f(\alpha)$ is determined by adjusting a reaction model to experimental data. Subsequently, k(T) can be evaluated by the form of $f(\alpha)$ chosen. This is a limitation because the calculated kinetic parameters are very dependent upon the chosen kinetic model [67]. For zero-order processes, the degradation rate can be represented in integrate form as:

$$\alpha = k(T) \cdot t + C \tag{4}$$

Thus, using isothermal scans, it is possible to obtain k (T) from a linear fitting of α versus time, where C is a constant. Then, when k (T) is known, the Arrhenius equation can be applied:

$$k = Ae^{-E/RT} (5)$$

$$ln k = ln A - \frac{E}{RT}$$
(6)

In Eqs. (5) and (6), A is the pre-exponential factor, E is the activation energy and E is the gas constant. Therefore, a plot of $\ln k(T)$ versus (1/T) will allow us to obtain E from the slope. The major problem of this isothermal methodology is that the sample requires some time to reach the experimental temperature. Model-free (isoconversional) methods allow estimation of the activation energy as a function of α without choosing the reaction model. The basic assumption of these methods is that the reaction rate for a constant extent of conversion depends only on temperature. Considering non-isothermal experiments that are performed at constant heating rate (β) , Eq. (2) can be written as follows:

$$\frac{d\alpha}{dT} = \frac{A}{\beta} e^{-E/RT} f(\alpha) \tag{7}$$

An integration function is shown below, where $g(\alpha)$ is the integrated kinetic function or integral reaction model.

$$\int_0^\alpha \frac{d\alpha}{f(\alpha)} = g(\alpha) = \frac{A}{\beta} \int_{T_0}^T e^{-E/RT} dT$$
(8)

The use of several heating rates enables the application of model-free (isoconversional) methods. These methods make the assumption that the parameters of the model are identical for measurements at all heating rates. It allows for a direct fit of the model to the experimental data without any transformation

[10]. An additional advantage lies in the fact that there are no limitations with respect to the complexity of the model; consequently, it is reliable for solving applied kinetic problems [68-71].

The integral on the right side of Eq. (8) has no analytical solution and some approximation methodology is necessary. The Kissinger-Akahaira-Sunose (KAS) method [72] is based on the following equation:

$$\ln \frac{\beta}{T_a^2} = \ln \left[\frac{RA}{E_a g(\alpha)} \right] - \frac{E_a}{R} \frac{1}{T_a}$$
 (9)

To apply the method, it is necessary to obtain at least three different heating rates (β), with the respective conversion curves being subsequently evaluated from the measured TGA curves [73]. For each conversion (α), $\ln \frac{\beta}{T_{\alpha}^2}$ plotted against $1/T_{\alpha}$, gives a straight line with slope -E α /R and thus, the activation energy is obtained as a function of the conversion. The evaluation of E α dependence is enough to predict the isothermal kinetics from non-isothermal data, as in Eq. (7):

$$t_{\alpha} = \frac{\int_{0}^{T_{\alpha}} e^{-E_{\alpha}/RT} dT}{\beta e^{-E_{\alpha}/RT_{0}}}$$
(10)

The time (t_{α}) , computed by Eq. (10), will be reached at an arbitrary temperature (T_{θ}) with a given conversion (α) .

3. Results and discussion

3.1. Thermal decomposition

Fig. 1 shows the TG and DTG curves obtained under inert (nitrogen) atmosphere for the five samples, with a very large range of thermal stability under these pyrolytic conditions (absence of oxygen). Most of the samples have an onset decomposition temperature over 300 °C, with a couple of T_{onset} values even higher than 400 °C. By analysing the obtained data, the most stable IL is $[P_{66614}][NTf_2]$. The stabilities decrease in the order of $[P_{66614}][NTf_2] > [P_{44414}][DBS] > [P_{66614}][(iC_8)_2PO_2] > [P_{4442}][DEP] > [P_{66614}][(iC_8)_2PO_2]$ samples exhibited only one resolved peak centred at 456, 453 and 376 °C, respectively (minimum in DTG curves, T_{peak1}), whereas the shape of the DTG curves obtained for the samples $[P_{66614}][BEHP]$ and $[P_{4442}][DEP]$ present a more complex mechanism. The latter ILs have, at this heating

rate, two well-resolved peaks (T_{peak1} and T_{peak2}) at 331/467 °C ($[P_{66614}][BEHP]$) and 342/530 °C ($[P_{4442}][DEP]$). This behaviour is possibly related to the influence of a more reactive phosphate type anion [13], causing alkyl phosphates to pyrolyze readily at temperatures of 300 °C and above [74] and thus resulting in an extra peak. Early weight losses in all samples are probably related to the impurities (between 2–5% in weight according to Table 1) present in the ionic liquids used in this study [15]; although they seem more pronounced with phosphate anion ILs (Fig. 1b), probably due to their water content. In view of these results, it can be concluded that anions play an important role in the mechanism of thermal decomposition, because of the similarity of all the cations used in this study. A previous research based on the thermal decomposition of amino- and hydroxyl-functionalised ILs composed of different cations showed that the stability of ILs is generally controlled by the anion [75].

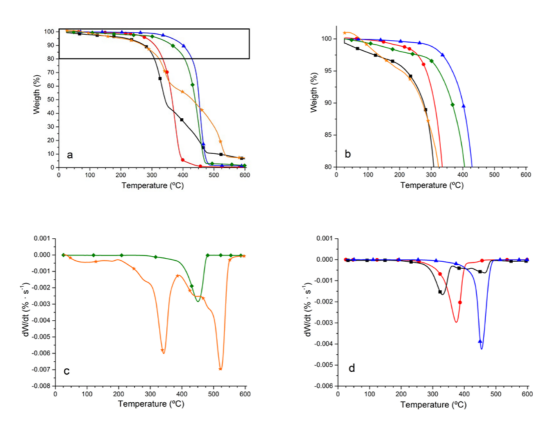


Fig. 1. (a) Full scale TGA, (b) TGA at square region zoom and (c, d) DTG curves for the five ILs obtained in an inert (nitrogen) atmosphere at a heating rate of 10 °C/min: (●) $[P_{66614}][(iC_8)_2PO_2]$, (■) $[P_{66614}][BEHP]$, (♠) $[P_{66614}][NTf2]$, (♦) $[P_{44414}][DBS]$ and (★) $[P_{4442}][DEP]$.

These results are in agreement with others reporting that the thermal stability of ILs is largely determined by the coordinating ability of the anion [76,77]. It is likely that the main goal for achieving weak

coordinating chemistry is a high degree of charge delocalisation. The charge delocalisation should happen over the entire anion, avoiding individual atoms or groups of atoms with high concentrations of charge, such as oxygen or chlorine atoms. This means that anions with accessible oxygen atoms for binding to the cation should have a stronger coordinating ability than others without available atoms with high concentrations of charge. Alternatively, larger anions (containing more atoms) provoke better delocalisation of the charge and weaker coordination [78]. Taking into account all these considerations, ILs containing anions without accessible oxygen for binding to the cation, such as bis(trifluoromethylsulfonyl)imide or with an aromatic group and larger alkyl chains, such as dodecylbenzenesulfonate, provide a better delocalisation of the charge leading to good thermal stability [63,76]. In addition, anion nucleophilicity is another possible approach which should prove useful in predicting quantitative thermal stability trends in ionic liquids [79,80]. The pathway with the lowest activation free energy would be expected to be the most nucleophilic [81].

Regarding the more reactive phosphate type ILs ($[P_{66614}][BEHP]$ and $[P_{4442}][DEP]$), three stages of weight loss up to 600°C were observed when an inert (nitrogen) atmosphere was used: *i*) 25-200 °C, *ii*) 200-500 °C and *iii*) 500-600 °C, with the main weight loss being found in the 200-500 °C case. In this mentioned case, it is possible to find fragments of H_2O , linear hydrocarbons and hydrocarbon arms attached to phosphorus [82]. Since there is no reactive atmosphere, the most abundant fragments should correspond to hydrocarbons with lengths greater than those found in oxidative atmosphere. The absence of a reactive atmosphere makes it impossible for the formation of P_2O_5 , CO_2 and H_3PO_4 .

3.2. Thermo-oxidative decomposition

When analysing the thermal stability of ILs, a question that should be raised is how to unequivocally define its thermal decomposition temperature. The answer is not direct, even if circumscribed to a single technique, such as TGA. Apart from factors related to the sample itself, such as the presence of moisture and/or impurities (which could lead to a significant decrease in the stability), taking into account the operating factor (type of sample, sample size and heating rate) is also required. This coexistence of different factors limits the comparison of data from different sources [15,77], with the experimental atmosphere used in TGA studies being one of the most important factors affecting the results [65,83]. TGA and DTG curves measured in an oxidising atmosphere are shown in Fig. 2. Since the atmosphere

plays a very important role in the thermal degradation of ILs, the shape of these curves is very different from the curves displayed in Fig. 1.

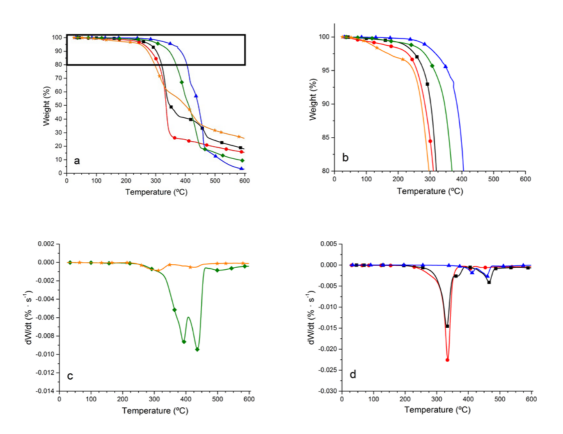


Fig. 2. (a) Full scale TGA, (b) TGA at square region zoom and (c, d) DTG curves for the five ILs obtained in a reactive (oxygen) atmosphere at a heating rate of 10 °C/min: (\bullet)[P₆₆₆₁₄][(iC₈)₂PO₂], (\blacksquare) [P₆₆₆₁₄][BEHP], (\blacktriangle) [P₆₆₆₁₄][NTf2], (\blacklozenge) [P₄₄₄₁₄][DBS] and (\bigstar) [P₄₄₄₂][DEP].

According to Keating et al. [82], it is likely that fragments of H₂O, linear hydrocarbons and hydrocarbon arms attached to phosphorus are found when phosphonium-based ILs react from 250 to 400°C. Thus, linear hydrocarbon fragments of different lengths, such as CH₃, C₂H₅, C₃H₇, C₄H₉, C₅H₁₁, C₆H₁₃, can be distinguished. Besides, the organophosphorus compounds can be result in intact P-C bonds in which the number and length of the hydrocarbon can be diverse, i.e., P(CH₂)₃, P(CH₂)₄, CH₂=P(CH₂CH₂)₂, CH₂=P(CH₂CH₂)₃, P(CH₂CH₂)₄. It is also probable that the longest hydrocarbon arms attached to phosphorus are present in higher amounts between 400 and 500°C. Other mass fragments that can be found are CO₂ and H₃PO₄. The former one results from the oxidation of hydrocarbons, whereas the latter is due to phosphorus oxidation to P₂O₅, which yields phosphoric acid in presence of water. Taking into account that [P₄₄₄₂][DEP] IL has short-length hydrocarbons, the presence of CH₃, C₂H₅, C₃H₇, P(CH₂)₃ and P(CH₂)₄ can be expected. Besides, other fragments such as CO₂ and P₂O₅ can also be expected. In

this case, considering that the hydrocarbon chain of the anion is short, the phosphorus is more easily accessible for oxidation to P_2O_5 . This can explain why the residual content obtained in oxidising atmosphere is very high (25.9%) in comparison with that found in inert atmosphere.

A data comparison between Figs. 1 and 2 is carried out in Table 2, by collecting several thermogravimetric parameters, such as T_{onset} , $T_{10\%}$, T_{peak1} , T_{peak2} and $Wt_{total\ loss}$ in both inert (nitrogen) and reactive (oxygen) atmospheres. In view of the results, there is a clear influence of the atmosphere on both temperatures of thermal degradation (T_{onset} and $T_{10\%}$) and in the total weight loss ($Wt_{total\ loss}$). These mentioned differences between both atmospheres have also been reflected on the DTG curves (T_{peak1} and T_{peak2}), indicating a mechanism of decomposition/oxidation in two steps for samples [P_{66614}][NTf₂], [P_{44414}][DBS] (Figs. 2c and 2d) against the one-step thermal decomposition process under nitrogen.

Table 2. Thermal results from dynamic scans in both atmospheres at 10 °C/min.

Ionic Liquids	Atmosphere	Tonset (°C)	T _{10%} (°C)	T _{peak1} (°C)	T _{peak2} (°C)	Wt _{total loss*} (%)
[P ₆₆₆₁₄][(iC8) ₂ PO ₂]	N_2	338	313	376		99.6
	O_2	320	286	344	390	84.6
[P ₆₆₆₁₄][BEHP]	N_2	298	276	331	467	93.5
	O_2	296	304	339	470	82.0
[P ₆₆₆₁₄][NTf ₂]	N_2	434	400	456		98.8
	O_2	401	388	422	432	96.7
[P ₄₄₄₁₄][DBS]	N_2	414	367	453		98.5
	O_2	359	353	401	438	89.6
[P ₄₄₄₂][DEP]	N_2	309	272	342	530	92.8
	O_2	265	274	312	423	74.1

^{*}total experiment time: 57.5 min.

Regarding the early weight loss region (Fig.2b), it seems again that the phosphate anion ILs (especially $[P_{4442}][DEP]$) provoke a more pronounced descent, probably related to the water content (analogous to Fig.1b). A comparison of the first the step of weight loss up to 275 °C, depending on the ILs used, using reactive atmosphere (O_2) and inert one (N_2) showed that the initial weight losses and average rates were greater in the inert atmosphere than in the oxidising one for all ionic liquids with the exception of $[P_{66614}][(iC8)_2PO_2]$. The oxygen presence alters the pathway of thermal degradation by initiating reactions of lower activation energy. This fact suggests that thermal degradation starts earlier in this IL in O_2

atmosphere (see Table 3). It should be noted that this process is masked by weight loss processes, especially when heating rates above 1.5°C/min were used, as in agreement with Menczel and Prime [84]. Regarding the step of the greatest weight loss in oxidising atmosphere, it took place between 225-285°C, 200-400°C, 320-500°C, 275-525°C and 200-525°C for [P₆₆₆₁₄][(iC8)₂PO₂], [P₆₆₆₁₄][BEHP], [P₆₆₆₁₄][NTf₂], [P₄₄₄₁₄][DBS] and [P₄₄₄₂][DEP], respectively. However, in the presence of nitrogen, the ranges of temperature in which the greatest weight loss occurred were generally wider, namely 225-475°C, 200-500°C, 200-500°C, 275-525°C and 200-525°C for the same ILs. Therefore, the decomposition temperature and thermal stability of ILs is strongly dependent on the selected anion [16]. In this case, the rate of weight loss does not show a clear tendency, possibly attributed to the different mechanisms for the degradation. Thus, the oxidation tends to show various steps, commonly two, whereas the decomposition exhibits usually one step. Hence, when thermo oxidative processes are involved, decomposition and oxidation take place concurrently. This implies both weight gain associated with the oxidation and weight loss due to the decomposition of the ILs, and this accounts for the declining tendency in mass over this temperature range. However, when nitrogen is employed, the weight is almost completely lost in the previous stage/stages.

Table 3. Mass changes and rates in the first step of weight loss of the different ILs between 25-275°C.

	Reactive atmosphere (O ₂)			Inert Atmosphere (N ₂)		
Ionic Liquid	T ^a Range (°C)	Weight loss (%)	Rate (%•°C ⁻¹)	T ^a range (°C)	Weight loss (%)	Rate (%•°C ⁻¹)
[P ₆₆₆₁₄][(iC ₈) ₂ PO ₂]	25-225	2.31	$1.15 \cdot 10^{-2}$	25-225	1.43	$7.20 \cdot 10^{-3}$
[P ₆₆₆₁₄][BEHP]	25-200	0.78	$4.50 \cdot 10^{-3}$	25-200	4.08	2.33 · 10 ⁻²
[P ₆₆₆₁₄][NTf ₂]	25-200	0.19	1.10.10-3	25-200	0.38	2.10.10-3
[P ₄₄₄₁₄][DBS]	25-275	2.03	8.1.10-3	25-275	2.71	1.08 · 10-2
[P ₄₄₄₂][DEP]	25-200	3.10	1.77 · 10 ⁻²	25-200	5.70	3.26 · 10 ⁻²

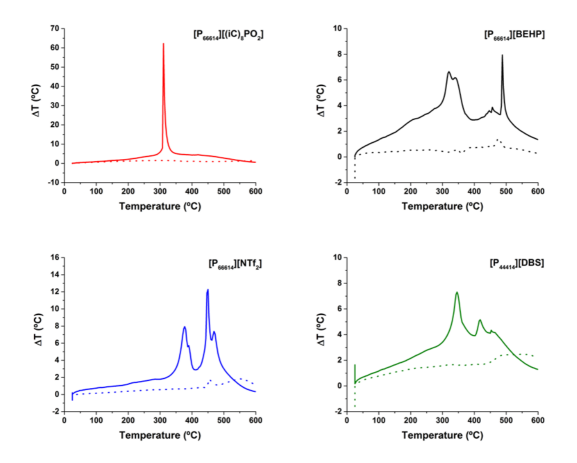
It is also noteworthy that the amount of residue obtained after TGA is larger when reactive atmosphere was employed (see Table 4) and therefore $Wt_{total\ loss}$ is smaller under oxidative conditions. This is explained by the weight gain during the oxidation process. Besides, ILs containing phosphorus in both the cationic and anionic moiety, also presented greater values of residual content. This is due to the oxidation of phosphorus to P_2O_5 , which partly remained in the residue. Such a result is in agreement with the data

reported by Keating et al. [82], who found Phosphorus in the ash of $[P_{66614}][NTf_2]$ and $[P_{66614}][(iC8)_2PO_2]$ after thermogravimetric analysis in the range of 20-740°C.

Table 4. Residual content of different ILs after TGA at 600°C

-	Reactive atmosphere (O ₂)	Inert Atmosphere (N ₂)
Ionic Liquid	Residue at 600°C (%)	Residue at 600°C (%)
[P ₆₆₆₁₄][(iC ₈) ₂ PO ₂]	15.4	0.35
[P ₆₆₆₁₄][BEHP]	18.1	6.5
$[P_{66614}][NTf_2]$	3.3	1.2
$[P_{44414}][DBS]$	10.4	1.5
$[P_{4442}][DEP]$	25.9	7.2

Fig. 3 shows the comparison of SDTA (single differential thermal analysis) curves in both atmospheres for all ILs tested.



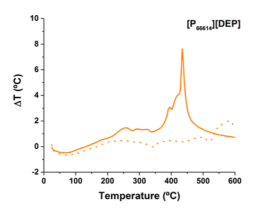


Fig. 3. SDTA for all ILs obtained in both inert (nitrogen, ---) and reactive (oxygen, —) atmospheres at a heating rate of 10 °C/min.

In these curves, the ILs under study and an inert reference are conducted to undergo the same thermal cycles, recording differences in the temperature (ΔT) and identifying physical and chemical phenomena causing heat changes (exothermic or endothermic), such as adsorption (exothermic), crystallisation (exothermic), vaporisation (endothermic), oxidation (exothermic) and so on. The mechanistic differences between thermal and thermo-oxidative degradation is clear in view of these results. The decomposition was determined to be endothermic if the SDTA temperature gradient was negative. Whilst in an inert (nitrogen) atmosphere the thermal decomposition implies a very low signal, exothermic signal measurement in the reactive (oxygen) atmosphere is related with an overlapped oxidation process.

3.3. Kinetic analysis

The above mentioned thermal stabilities have been evaluating using TGA at a single linear heating rate (10 °C/min). The kinetics and mechanism of the thermal decomposition reaction were evaluated from the TG data according to the ICTAC Kinetics Committee recommendations [85,86]. As previously discussed, isothermal studies have shown that ILs exhibit appreciable decomposition at temperatures significantly lower than the values indicated by the onset temperature decomposition calculated from scanning TGA experiments. Therefore, in order to fully evaluate the long-term thermal stabilities and thermal decomposition of the studied ILs, it is necessary to identify global kinetics models.

In this research study, several parameters such as the activation energy (E), the conversion and the degradation decomposition time were estimated as a function of temperature using both TG experimental data and KAS model-free kinetics (in an oxygen atmosphere). As model-free kinetics require at least

three dynamic curves with different heating rates, TGA curves obtained at 5, 10 and 20 °C/min are shown in Fig. 4.

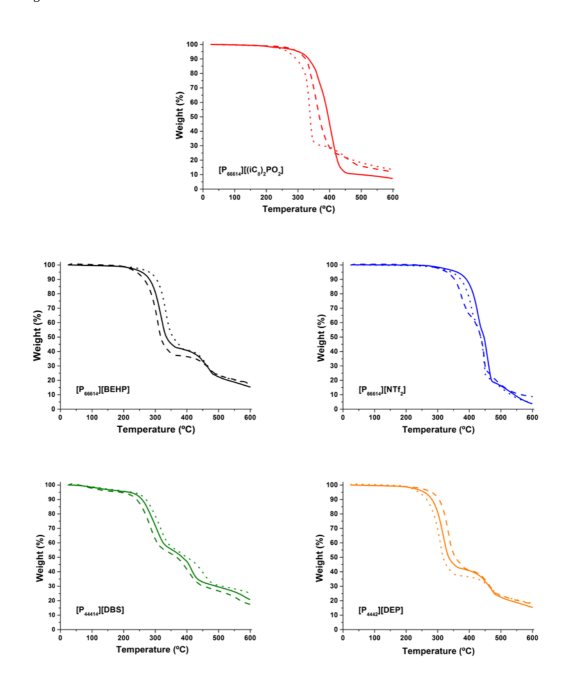


Fig. 4. TGA of all ILs studied in a reactive (oxygen) atmosphere at heating rates of 5 (---), 10 (—) and 20 °C/min (···).

Fig.4 shows the behavior of $[P_{44414}][DBS]$ and $[P_{66614}][BEHP]$ samples at lower heating rates, where the temperature values shift lower and consequently the total mass loss increases [87]. On the other hand, $[P_{4442}][DEP]$ behaved different: higher heating rates seem to activate the decomposition reaction and the total mass loss growths. Finally, it is hard to extract conclusions from heating rate variation with

[P₆₆₆₁₄][NTf₂] and [P₆₆₆₁₄][(iC₈)₂PO₂] samples. The model-free approach to kinetic analysis rests upon the isoconversional principle, according to which the reaction rate at a constant extent of conversion is only a function of temperature. This fact implies that curves plotted at different heating rates cannot intersect. If this issue happens, data obtained could lead to unrealistic conclusions [88]. In this research, the crossing curves issue happens at temperatures higher than 420 °C. This fact can be observed in all samples and it is related to the wt.% obtained at the end of pyrolysis (char formation yield). Therefore, an interval of temperatures 200–420 °C was chosen for model-free kinetics calculations. Fig. 5 shows the apparent activation energy versus thermal conversion process obtained by the KAS model-free kinetics methodology.

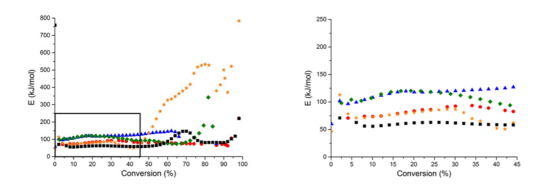


Fig. 5. Apparent activation energy as a function of conversion derived from experiments using 5, 10 and 20 °C/min heating rates for the five ILs: (●) [P₆₆₆₁₄][(iC₈)₂PO₂], (■) [P₆₆₆₁₄][BEHP], (▲) [P₆₆₆₁₄][NTf2],
(◆) [P₄₄₄₁₄][DBS] and (★) [P₄₄₄₂][DEP]. Full scale image (left) and zoom at square region (right).

In general, constant activation energy values should be expected in the case of a single reaction. However, the E profile obtained from Fig. 5 indicates that the decomposition mechanism is a function of the conversion degree. In addition, Fig.5 (right) shows that the activation energy (E) remained practically constant at low conversions (2–40%), with values between 50 and 120 kJ/mol. Table 5 exhibites the average values and their standard deviations. Therefore, the trend of relative nucleophilicities of the anions in this research work is (in decreasing nucleophilic strength): $[P_{66614}][BEHP] > [P_{4442}][DEP] > [P_{66614}][iC_8)_2PO_2] > [P_{44414}][DBS] > [P_{66614}][NTf_2]$, according to the activation energy (E) values (Table 5). These results are in agreement with the delocalisation charge approach, where $[P_{44414}][DBS]$ and $[P_{66614}][NTf_2]$ are the less nucleophilic ILs and therefore the most thermally stable ones.

Table 5. Apparent activation energy values $(0.02 \le \alpha < 0.4)$ for the five ILs.

Ionic liquids	E (kJ/mol)
[P ₆₆₆₁₄][(iC ₈) ₂ PO ₂]	81 ± 7
$[P_{66614}][BEHP]$	60 ± 3
$[P_{66614}][NTf_2]$	115 ± 6
$[P_{44414}][DBS]$	113 ± 8
[P ₄₄₄₂][DEP]	79 ± 5

In order to provide a more realistic description of the thermal stability of these samples, kinetic analysis was performed in oxidising conditions by application of the model-free KAS isoconversional method. This methodology allows one to determine the change in the apparent activation energy during the thermo-oxidative process without choosing the reaction model. Results are shown in Fig. 6.

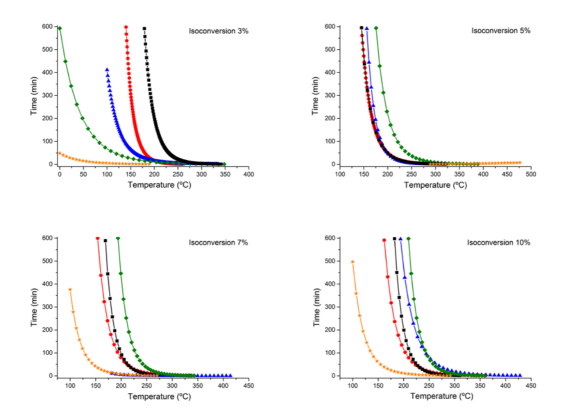


Fig. 6. Estimated isoconversional curves at 3, 5, 7 and 10 % of conversion for the five ILs:

(●) $[P_{66614}][(iC_8)_2PO_2]$, (■) $[P_{66614}][BEHP]$, (▲) $[P_{66614}][NTf2]$, (♦) $[P_{44414}][DBS]$ and (★) $[P_{4442}][DEP]$.

From Fig. 6, it is possible to estimate the time required for a fixed mass loss (conversion) of the ionic liquid at a given temperature. If 200° C is used, $[P_{4442}][DEP]$ reached 10% conversion in about 25 min, whereas $[P_{66614}][(iC_8)_2PO_2]$ and $[P_{66614}][BEHP]$ needed more than 100 min. Finally, $[P_{66614}][NTf_2]$ and

[P₄₄₄₁₄][DBS] did not achieve the 10% mark in the total duration of the test (600 min). The analysis would be similar for the others conversion rates (3%, 5% and 7%). In addition, it is possible to predict the sample behaviour at isothermal conditions. The model-free methodology allows one to estimate the sample performance in isothermal conditions. In order to check these results, some isothermal experiments were carried out for the [P₆₆₆₁₄][(iC₈)₂PO₂] sample (Fig. 7). Isothermal experiments indicate that thermo-oxidative decomposition reaches values of 6, 22 and 23% of conversion when the sample is heated at 200, 225 and 250 °C, respectively, for 90 min. In addition, it is possible to predict the sample behaviour in isothermal conditions at a given temperature (T₀) by using Eq. (10) with non-isothermal kinetic data. In order to solve the mentioned equation, the integral is determined using the approximate formula obtained by Wanjun et al. [89]. Thus, the temperature (in °C) at which each conversion value is reached in the selected time could be obtained [66]. The deviations between experimental and model values may be linked to not taking into account the evaporation kinetics. Anyway, From Fig. 6 data, results obtained at low conversions ($\alpha < 0.1$) by the application of the model-free methodology with $T_0 = 200$ °C are in good agreement with those obtained by experimental isothermal analysis. Although this result seems to support the decision concerning evaporation, further research considering this evaporation kinetic should be conducted in all temperature range to validate completely the model [65]. On the other hand, differences between 200 and 225 isotherms prove the previous statement regarding the overestimation of long-term thermal stability of the ILs with conducting standard TGA experiments. If the 320 °C value of T_{onset} calculated from classic dynamic TGA analysis were precise, the 225 and 250 isotherms should be closer to 200 °C one.

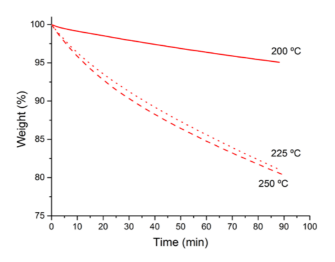


Fig. 7. Isothermal scans for $[P_{66614}][(iC_8)_2PO_2]$ in a reactive (oxygen) atmosphere.

4. Conclusions

The thermal degradation of five phosphonium cation-based ILs, $[P_{66614}][BEHP]$, $[P_{66614}][(iC_8)_2PO_2]$, $[P_{66614}][NTf_2]$, $[P_{44414}][DBS]$ and $[P_{4442}][DEP]$, was studied in both nitrogen (inert) and oxygen (reactive) atmospheres using dynamic and isothermal approaches. From the obtained results, the following conclusions can be drawn:

- Thermal degradation of ILs is clearly dominated by the nature of the anion, endowing the weakly-coordinating-anion ILs (such as [NTf₂] and [DBS] based) with a great thermal stability. On the basis that the pathway with the lowest activation free energy (E) would be expected to be the most nucleophilic, the trend in decreasing nucleophilic strength is as follows: $[P_{66614}][BEHP] > [P_{4442}][DEP] > [P_{66614}][(iC_8)_2PO_2] > [P_{44414}][DBS] > [P_{66614}][NTf_2]$. Therefore, both approaches are in good agreement.
- Although the atmosphere has a clear influence on the thermal degradation mechanism, the trend of the thermal stability among different samples did not change. Both temperatures of thermal degradation (T_{onset} and T_{10%}) presented the [P₆₆₆₁₄][NTf₂] and [P₄₄₄₁₄][DBS] as the most thermal stable ILs in both pyrolytic and oxidising conditions.
- Despite of the fact that the decomposition mechanism is a function of the conversion degree, the
 activation energy (E) remained practically constant at low conversions, with values ranging from
 60–115 kJ/mol for all ILs studied.
- The long-term thermal stability of the ILs is also overestimated with conducting standard TGA
 experiments; so the use of isoconversional methods is more appropriate.
- The results from both the application of the model-free methodology and the isothermal
 experiments are in good agreement especially at low conversions, so the former can be used to
 predict at lower costs the long-term thermal stability.
- The determination of thermal models, especially those which can predict the long-term thermal stability, is an interesting research topic from lubrication science approach in order to predict the lifetime of lubricants and additives.
- The physicochemical properties of phosphonim cation-based ILs (conferred by the size, symmetry and structure of their ions) combined with an excellent tribological behaviour make them excellent candidates for being used as a lubricant additive. The utilization of this kind of

substances as an additive in motor oils should take into account current and future specifications for motor oils (e.g. PC-11, ILSAC GF-5 and ILSAC GF-6) which limit phosphorus concentration under 800 ppm.

5. Nomenclature

A	$(m^{3} \cdot mol^{-1})^{n-1} s^{-1}$	Pre-exponential factor
E	kJ/mol	Activation Energy
$M_{ m IL}$	mol	Molar mass of the ionic liquid
В	m/s	Mass transfer coefficient
A	m^2	Surface area in contact with the gas phase
p_{vap}	Pa	Vapour pressure
f(a)	-	Reaction model
k(T)	J/K	Temperature dependent constant
R	J/K/mol	Gas constant
t	S	Time
T	°C	Temperature
T_{on}	°C	Onset Temperature
T_{10}	°C	Temperature at 10% of mass loss
T_0	°C	Arbitrary Temperature to a given extent of reaction, α
m_i	kg	Initial mass
$\mathbf{m_f}$	kg	Final mass
$\mathbf{m_t}$	kg	Mass at given time, t
α	-	Extent of reaction
β	°C/min	Heating rate

Acknowledgements

The authors would like to thank to the Ministry of Economy and Competitiveness (Spain) and FICYT (Foundation for the Promotion in Asturias of the Applied Scientific Research and Technology) for supporting the research projects STARLUBE (DPI2013-48348-C2-1-R) and GRUPIN14-023, respectively, under the framework this research was developed. The Unit of Thermal Tests and Elemental

Analysis from the Scientific-Technical Services at the University of Oviedo as well as Dr. Paula Oulego Blanco are also acknowledged.

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