

Ionic liquid-based nanofluids (ionanofluids) for thermal applications: an experimental thermophysical characterization

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Conference paper

Kamil Oster*, Christopher Hardacre*, Johan Jacquemin, Ana P. C. Ribeiro and Abdulaziz Elsinawi

Ionic liquid-based nanofluids (ionanofluids) for thermal applications: an experimental thermophysical characterization

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Abstract: Heat transfer fluids materials are manufactured for the purpose of transfer, distribution and storage of heat. Several of their important properties can be listed (for example flash point, thermal expansivity or technical safety). However, to assess the thermal exchange performance of these fluids, a prior knowledge of their heat capacity, density, viscosity and thermal conductivity is obligatory. The most popular heat transfer fluids are based on organic liquids, such as ethylene glycol. However, new technologies and development require more efficient materials. Ionanofluids, mixtures of ionic liquids and nanoparticles, were proposed as a viable replacement for those commonly used fluids due to the properties of ionic liquids (wide liquid range or low vapour pressure and flammability) combined with enhanced thermophysical properties of nanofluids caused by the dispersion of nanoparticles (mainly thermal conductivity and heat capacity). Very few authors reported the extensive analysis of those systems thermophysical properties and impact on the heat exchange efficiency. Moreover, the availability of published data is very limited. The aim of this work is to investigate ionanofluids based on the trihexyl(tetradecyl)phosphonium cation paired with the acetate, butanoate, hexanoate, octanoate or decanoate anion, mixed with carbon nanotubes, boron nitride, graphite or mesoporous carbon as nanoparticles with concentration up to 3 wt %. The density, heat capacity, thermal stability, thermal conductivity and viscosity of selected ionanofluids were determined experimentally as functions of the temperature (up to 363.15 K) and compared with theoretical tools to evaluate the predictive capability. Based on the experimental results, lubrication, heat storage potential and economic analysis were also discussed and compared to commercial heat transfer fluids.

Keywords: heat transfer fluids; ionanofluids; ionic liquids; ISSP-18; nanofluids; nanoparticles; thermophysical properties.

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Johan Jacquemin: Queen's University Belfast, School of Chemistry and Chemical Engineering, Stranmillis Road, BT9 5AG, Belfast, United Kingdom of Great Britain and Northern Ireland; and Université François Rabelais, Laboratoire PCM2E, Parc de Grandmont 37200, Tours, France

Ana P. C. Ribeiro: Universidade de Lisboa, Centro de Química Estrutural, Instituto Superior Técnico, Av. Rovisco Pais 1, 1049-001 Lisbon, Portugal

Abdulaziz Elsinawi: King Fasial University, Materials Engineering Department, College of Engineering, Al-Hasa, 31982, Hofuf, Saudi Arabia

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^{*}Corresponding authors: Kamil Oster and Christopher Hardacre, The University of Manchester, School of Chemical Engineering and Analytical Science, Sackville Street, M13 9PL, Manchester, United Kingdom of Great Britain and Northern Ireland; and Queen's University Belfast, School of Chemistry and Chemical Engineering, Stranmillis Road, BT9 5AG, Belfast, United Kingdom of Great Britain and Northern Ireland, e-mail: kamil.oster@manchester.ac.uk (K. Oster); c.hardacre@manchester.ac.uk (C. Hardacre)

Introduction

Heat transfer fluids (HTFs) have the purpose of transferring, distributing and storing the heat. To describe the thermal performance properly, the most following properties are necessary – thermal conductivity, heat capacity, density and viscosity [1]. Based on these properties, information on the heat transfer rate, the capability to store the energy as the heat or pumpability can be determined [2]. In the case of HTFs, the molecular design is one of the most challenging tasks [3]. It can be performed in two ways - experimentally or theoretically via predictions. Predictions are the most robust, easiest and readily available approach. However, experimental details are still needed to establish the models. Current HTF technologies include organic compounds (glycols, silicones or aromatic compound) and water [1, 2]. In the case of organic compounds, several disadvantages can be listed, for example biotoxicity or flammability. They can be also divided into three main groups depending on the temperature of operation: low temperature (up to 563.15 K), medium temperature (563.15 – 863.15 K) and high temperature (above 863.15 K) [1, 2]. Other properties of HTFs should be also considered during the design process, i.e. corrosivity, flash point, durability, maintenance time or environmental impact [4].

One of the replacement solutions for current HTFs to address these issues is ionic liquids (ILs). These are defined as compounds consisting entirely of ions with melting point below 373.15 K [5]. Due to their ionic structure, strong Coulombic interactions can be distinguished [6]. Moreover, a large variety of hydrogen bonding and van der Waals interactions can be found [7, 8]. These features strongly influence the physical properties of ILs, namely low vapour pressure (and flammability), wide liquid range with low freezing point and relatively high thermal stability, ionic conductivity [9]. ILs have been studied as potential HTFs by many authors, including understanding their advantages and disadvantages [10-14]. For example, Wu et al. reported several ILs consisting of 1-alkyl-3-methylimidazolium cations with various anions in terms of their heat capacity, and further heat storage capacity, structure-property relationship, feasibility and economic analyses [15]. The authors showed that ILs might have very beneficial physical properties as HTFs [15]. Holbrey et al. presented the comparison between 1-alkyl-3-methylimidazolium ILs and commercial HTFs concentrating on their heat capacity [16]. Van Valkenburg et al. investigated 1-alkyl-3-methylimidazolium tetrafluoroborate ILs in terms of their physical properties (thermal stability, heat storage and transfer), and demonstrated that they might be a suitable replacement for conventional HTFs [17]. An extensive review by Chernikova et al. discussed the values of stability associated with the desired low vapour pressure, varied viscosity and corrosivity [18]. The main drawback mentioned was the high cost of production and maintenance. Musiał et al. presented the results of thermophysical properties of pyrrolidinium-based ILs and their comparison to commercial HTFs as a wide range of the temperature and the pressure, and showed that this class of ILs might be a real replacement for currently used HTFs [19]. Zorębski et al. discussed the heat capacity of ILs as the main advantage as HTFs [20]. Our recent work on thermophysical properties of tetraalkylphosphonium carboxylate ILs showed that the viscosity might be a limiting factor to their application [21].

Another solution that has been proposed is mixing liquids with nanoparticles, creating so-called nanofluids [22]. The very first work on nanofluids addressing their potential thermal conductivity, heat transfer coefficient enhancements and pumping power savings caused by dispersion of nanoparticles in liquids was reported by Choi and Eastman [23]. The addition of nanoparticles resulted in enhanced thermal conductivity [24] and heat capacity [25, 26] which are very beneficial for industrial applications. These enhancements were unusual because the mechanism of heat transfer found for liquids nor solids was not able to explain the behaviour in nanofluids. The reasons proposed in the literature are based on Brownian motions of nanoparticles, liquid layering at the liquid/particle interface, nature of heat transport across nanoparticles, nanoparticle clustering. However, it has been shown by several authors based on both theoretical (molecular modelling) [27, 28] and experimental (thermal conductivity measurement) [25, 26] results that interfacial nanolayering seems to be the most influencing factor for the unusual enhancements. Moreover, several potential applications of nanofluids have been indicated, for example exfoliation [29], cold storage systems [30], solar collectors [31].

Further improvements were achieved by mixing ILs with nanoparticles, denoted as ionanofluids [32]. The unique properties of ILs are conserved, and the enhancement of physical properties is also obtained. These offer HTFs with low vapour pressure, low flammability, wide liquid range, enhanced thermal conductivity and heat capacity. The aforementioned enhanced thermophysical properties were repeated in further reports for thermal conductivity [33–35], heat capacity [33, 34, 36], and density [37]. Unfortunately, this area is still very poorly developed, and very few studies have been reported, to date [32, 35, 38, 39]. Therefore, besides the poor understanding of their physical properties behaviour, the prediction of those is also very limited. However, the thermal conductivity was studied by Atashrouz et al. [40, 41], and the density and heat capacity by Oster et al. [42].

The aim of this work is to investigate trihexyl(tetradecyl)phosphonium based ILs together with acetate, $[P_{14,6,6,6}][AcO]$, butanoate, $[P_{14,6,6,6}][ButO]$, hexanoate, $[P_{14,6,6,6}][HexO]$, octanoate, $[P_{14,6,6,6}][OctO]$, or decanoate, [P₁₄₆₆₆][DecO] anion. As shown in our previous work, these ILs were selected based on their very promising properties as heat transfer fluids [21]. To investigate them in further details, ionanofluids with multi-walled carbon nanotubes (MWCNT), boron nitride (BN), graphite (G) and mesoporous carbon (MC), up to 3 wt %, were also chosen. Such concentration of nanoparticles was imposed by the industrial applicability, further addition of nanoparticles can result in sedimentation [43], and increasing cost of materials [44]. The above systems were studied herein in terms of their physical properties behaviour. The physical properties measured and correlated during this work were the density, dynamic viscosity, thermal conductivity, isobaric heat capacity and thermal stability. The evaluation of investigated properties has also been undertaken during this work in order to evaluate predictive tools for such materials. Finally, further derivative features (kinematic viscosity and lubrication properties, volumetric heat capacity and energy storage ability, economic analysis) are also presented and depicted during this work.

Experimental

Materials

Ionic liquids

Trihexyl(tetradecyl)phosphonium carboxylate ionic liquids, [P₁₄₆₆₆][RO], were synthesized in accordance to the procedure in [21]. Generally, the primary ionic liquid, trihexyl(tetradecyl)phosphonium chloride, [P₁₄₆₆₆][Cl] (Cytec Industries Inc., product code: Cyphos IL-101, CAS: 258864-54-9, mass purity: >95%) is mixed with ethanol (Sigma-Aldrich, absolute, CAS: 64-17-5, mass purity: ≥99.8%) at a concentration of 30% (w/w). Thereafter, this mixture is passed through a strongly basic anion exchange resin (Alfa Aesar, product code: IRN-78, CAS: 11128-95-3), at least three times to ensure complete chloride exchange to hydroxide anion which is checked with the silver nitrate test (CAS: 7761-88-8, Sigma-Aldrich, mass purity ≥99.9999%). Finally, the obtained trihexyl(tetradecyl)phosphonium hydroxide, [P_{14,6,6,6}][OH], undergoes the reaction with carboxylic acid to create $[P_{14,6,6,6}][RO]$ (as in Fig. 1), at 328 K on stirring overnight. Trihexyl(tetradecyphosphonium) acetate, (CAS: 1393375-57-9), octanoate, [P_{14,6,6,6}][OctO] (CAS: 1393375-58-0), and decanoate, [P_{14,6,6,6}][DecO] (CAS: 465527-65-5) were obtained. Acetic acid (Sigma-Aldrich, ReagentPlus, CAS: 64-19-7, mass purity: ≥99 %), butanoic acid (Sigma-Aldrich, CAS: 107-92-6, mass purity: ≥99%), hexanoic acid (Sigma-Aldrich, CAS: 142-62-1,

Fig. 1: The general synthesis route for $[P_{14,6,6,6}][RO]$.

mass purity: ≥99.5%), octanoic acid (Sigma-Aldrich, CAS: 124-07-2, mass purity: ≥99%) and decanoic acid (Sigma-Aldrich, CAS: 334-48-5, mass purity: ≥98%) were used in the synthesis in slight molar excess to the amount of chloride salt used in the first step, accordingly. Finally, the product was purified under high vacuum (10⁻³ Pa) at elevated temperature (338 K) for at least 72 h. The summary of all materials used/obtained in this work can be found in Table 1.

1-Octyl-3-methylimidazolium bis[(trifluoromethyl)sulfonyl]imide, [C,C,Im][NTf,] (CAS: 178631-04-4) was also synthesized for the purpose of thermal conductivity calibration following the procedure found in reference [45]. The product was washed with deionised water (18 M Ω cm) 5 times, and the chloride content was checked with the silver nitrate test. The IL was dried under high vacuum (10⁻³ Pa) at 328.15 K for at least 72 h. This IL was accurately described in the literature in terms of the thermal conductivity, therefore, it was selected to check the calibration in this work [46]. The purity of all the investigated ionic liquids was checked with ¹H, ¹³C and ³¹P NMR (B400 Bruker Avance III 400 MHz, ILs dissolved in CDCl., Fig. S1–S3, Supporting Information SI1), Karl Fisher titration (Metrohm 899 Coulometer with 803 Ti Stand and Hydranal Coulomat AG) and CHNS elemental analysis (for carbon and hydrogen content, Perkin Elmer PE2400CHNS). The water content was checked before and after measurements, however, no changes were observed after the experiment. The water content for all samples was <100 ppm. The results of purity control can be found elsewhere [21], however, the final purity was summarized in Table 1.

Nanoparticles

The multiwalled carbon nanotubes (MWCNT) used in this work were Bayer Material Science Baytubes C150 HP (CAS: 308068-56-6, mass purity ≥99 %, 3–15 number of walls, 13–16 nm outer mean diameter, 5–20 nm outer diameter distribution, 4 nm inner mean diameter, 2-6 nm inner diameter distribution, 1-10 mm length, 140-230 kg m⁻³ bulk density; according to the specifications provided by the supplier). Hexagonal boron nitride (BN, CAS: 10043-11-5, mass purity \geq 99.8 %, 70–80 nm size, 2290 kg m⁻³ theoretical true density, <0.03 % Fe₂O₃, <0.002 % CaO, <0.04 % MgO, <0.1 % B₂O₃), graphite (G, CAS: 7782-42-5, ≥99.9 % mass purity, 400–1200 nm size, <0.1% impurities, ~0.2% water) and mesoporous carbon (MC, CAS: 7440-44-0, mass purity: >95%, 20–40 nm average pore size, <5 % water, <2 % ash, 7–10 pH) were obtained from US Research Nanomaterials, Inc. Before preparation of ionanofluids, the boron nitride and graphite were processed by using ball milling to reduce the particle size (Retsch Mixer Mill MM 400, frequency 20 Hz, 3 times for 1800 s). The size of the particles was ascertained with scanning electron microscope (650 FEI Quanta FEG, 15-20, <90, <100, and <50 nm for carbon nanotubes, boron nitride, graphite and mesoporous carbon, respectively, Fig. S4, Supporting Information SI1), and with light scattering (Mastersizer Malvern, dispersion with water, stabilized with sodium laureth sulfate as anionic surfactant, refractive index of 1.330 for water, 1.650 for boron nitride and 2.420 for all carbon-based nanomaterials; 26.60 ± 0.15 nm, 154.5 ± 1.7 nm, 159.1 ± 3.1 nm and 46.23 ± 0.62 nm for carbon nanotubes, boron nitride, graphite and mesoporous carbon, respectively, Fig. S5, Supporting Information SI1). The purity of nanomaterials was checked with X-ray diffractometry (XPert Powder PANalytical, 15–90° scan range, 0.0084° scan step size, Fig. S6, Supporting Information SI1).

Ionanofluids

The preparation of (io)nanofluids is a very delicate process, therefore, a systematic procedure is essential and the standard literature protocol was used to prepare the ionanofluids [32, 35, 38, 42]. The proper mass of nanoparticles was weighted and added to ionic liquid with magnetic stirring for 30 min. Finally, the mixture was submitted to the 1 h of ultrasonication which is necessary to break up the nanoparticles agglomeration, stabilisation and preventing creating a so-called bucky-gel. The concentrations of nanoparticles in the investigated ionanofluids were (0.5, 1.0 and 3.0) % by weight. The samples were kept under high vacuum (10⁻³ Pa) to prevent water absorption from the air and removal of trapped air/water in the nanoparticles.

Table 1: Chemicals summary.

Chemical name [CAS]	Supplier	Mass fraction purity	Purification method	Analysis method	Halide/water content (ppm)
Trihexyl(tetradecyl)	Cytec	>0.95	_	_	_
phosphonium chloride	Industries Inc.				
[258864-54-9]					
Ethanol [64-17-5]	Sigma-Aldrich	>0.998	-	-	-
Acetic acid [64-19-7]	Sigma-Aldrich	≥0.99	-	-	-
Butanoic acid [107-92-6]	Sigma-Aldrich	≥0.99	-	-	_
Hexanoic acid [142-62-1]	Sigma-Aldrich	≥0.995	-	-	-
Octanoic acid [124-07-2]	Sigma-Aldrich	≥0.99	_	-	_
Decanoic acid [334-48-5]	Sigma-Aldrich	≥0.98	_	-	_
Trihexyl(tetradecyl)	In house	≥0.98	Washing-extraction	¹ H, ¹³ C and ³¹ P NMR	<5/<100
phosphonium acetate			vacuum	Elemental analysis	
[872700-58-8]				Karl Fisher titration	-1
Trihexyl(tetradecyl)	In house	≥0.98	Washing-extraction	¹ H, ¹³ C and ³¹ P NMR	<5/<100
phosphonium butanoate			vacuum	elemental analysis	
[1393375-56-8]				Karl Fisher titration	-/
Trihexyl(tetradecyl)	In house	≥0.98	Washing-extraction	¹ H, ¹³ C and ³¹ P NMR	<5/<100
phosphonium hexanoate			vacuum	elemental analysis	
[1393375-57-9]	In house	>0.00	Washing autrastian	Karl Fisher titration	-F/-100
Trihexyl(tetradecyl) phosphonium octanoate	In house	≥0.98	Washing-extraction	¹ H, ¹³ C and ³¹ P NMR elemental analysis	<5/<100
[1393375-58-0]			vacuum	Karl Fisher titration	
Trihexyl(tetradecyl)	In house	≥0.98	Washing-extraction	¹ H, ¹³ C and ³¹ P NMR	<5/<100
phosphonium decanoate	III IIouse	≥0.96	vacuum	elemental analysis	<5/<100
[465527-65-5]			vacuum	Karl Fisher titration	
1-octyl-3-methylimidazolium	In house	≥0.98	Washing-extraction	¹ H, ¹³ C and ³¹ P NMR	<5/<100
bis[(trifluoromethyl)sulfonyl]	III IIOU3C	20.70	vacuum	Karl Fisher titration	\J/\100
imide [178631-04-4]			vacaam	narrisher titration	
Toluene [108-88-3]	Sigma-Aldrich	≥0.995	_	_	_
Glycerine [56-81-5]	Sigma-Aldrich	≥0.99	_	_	_
Sodium chloride [7647-14-5]	Sigma-Aldrich	≥0.99	_	_	_
Water [7732-18-5]	Deionised	Ultrapure,	Deionisation	UV, conductivity	_
		type 1		measurement (Merck	
				Millipore Direct-Q	
				3 UV equipment)	
Synthetic sapphire	TA Instruments	Ultrapure (NIST	_	_	_
[1317-82-4]		SRM 720)			
Strongly basic anion resin	Alfa Aesar	≤0.0005 Cl ⁻ ;	-	-	-
IRN-78 [11128-95-3]		≤0.0001 SiO ₂ ;			
		≤0.0006 SO ₄			
Silver nitrate [7761-88-8]	Sigma-Aldrich	≥0.999999	-	-	-
Multi-walled carbon	Bayer Material	≥0.99	-	Supplier: Ashing,	-
nanotubes [308068-56-6]	Science			TEM, SEM, EN ISO 60;	
				This work: XRD,	
				laser diffraction	
				technique	
Boron nitride [10043-11-5]	US Research	≥0.998	_	This work: SEM, XRD	_
	Nanomaterials,				
C	Inc.			TI: C514 VDD	
Graphite [7782-42-5]	US Research	≥0.999	-	This work: SEM, XRD	-
	Nanomaterials,				
	Inc.			TILL I OFFE VES	
Mesoporous carbon [7440-44-0]	US Research	≥0.95	-	This work: SEM, XRD	
	Nanomaterials,				

Techniques

Density

The density, ρ , was measured as a function of temperature, T = (298.15 - 363.15) K, using Anton Paar DMA 4500M densitometer (calibrated on ultrapure degassed water and dry air, relative standard uncertainty of density, u(p) = 0.1%, relative standard uncertainty of temperature, u(T) = 0.01 K, approximate volume of sample 1.5 cm³, 3 independent repeats of measurement, viscosity-induced errors reduced by correction with viscosity), and correlated with temperature with the second-order equation:

$$\rho(T) = \sum_{i=0}^{2} a_i T^i \tag{1}$$

where a_i is the regression parameter determined by the least-square method, along with the standard uncertainty, δa , while the coefficient of determination, R^2 , was used to describe the fit. As a prerequisite for the ionanofluids density calculation, the nanoparticles density was previously determined for multiwalled carbon nanotubes, boron nitride and graphite [42]. However, those for mesoporous carbon have not been reported and, therefore, were determined using the following empirical formulae, in a similar procedure as for carbon nanotubes, boron nitride and graphite [47]:

$$\rho_{NP} = \frac{\rho_{IL+NP} - \rho_{IL} (1 - w_{NP})}{w_{NP}} \tag{2}$$

where w is mass fraction, subscripts IL and NP refer to ionic liquids and nanoparticles, respectively. Equation (2) was also used to calculate the density of ionic liquid + nanoparticles mixtures, as a prediction of the property. To represent the physical properties as a function of the nanoparticles loading, the mass fraction, $w_{_{m\,NP}}$ was recalculated to volume fraction, $\varphi_{_{NP}}$, with the following equation:

$$\varphi_{NP} = \frac{V_{NP}}{V_{NP} + V_{IL}} = \frac{\frac{m_{NP}}{\rho_{NP}}}{\frac{m_{NP}}{\rho_{NP}} + \frac{m_{IL}}{\rho_{IL}}} = \frac{\frac{w_{m,NP}}{\rho_{NP}}}{\frac{w_{m,NP}}{\rho_{NP}} + \frac{1 - w_{m,IL}}{\rho_{IL}}}$$
(3)

where m is a mass, V is a volume, density of nanoparticles was calculated using Eq. (2) for mesoporous carbon, and from the available literature data for the multiwalled carbon nanotubes, boron nitride and graphite. The density of the ionic liquids, ρ_u , was determined experimentally in our previous work [21].

Viscosity

The viscosity, η , was measured as a function of the temperature, T, using TA Instruments AR2000 rheometer (angular displacement resolution 40 nRad, torque resolution 0.1 nN m, temperature standard uncertainty u(T) = 0.01 K) with stainless steel 20 mm parallel plate geometry and shear rate of 10 s⁻¹. The measurement temperature range was (298.15-363.15) K. The viscosity standards Cannon S600 oil (1100 mPa s at 298.15 K), PTB 1000A (780 mPa s at 298.15 K) and ultrapure water (0.88992 mPa s at 298.15 K) were used to ascertain the reported viscosity relative standard uncertainty, $u_{\cdot}(\eta) = 3$ %. The viscosity was correlated with temperature by using Vogel-Fulcher-Tammann (VFT) equation:

$$\eta(T) = \eta_0 \exp\left(\frac{B}{T - T_0}\right) \tag{4}$$

where η_0 , B and T_0 are fitting parameters, determined by the nonlinear curve fitting algorithm in OriginPro 2017 software.

Specific heat capacity

The isobaric heat capacity, c_n , was measured by using differential scanning calorimeter (DSC) equipment, Q100 TA Instruments in the temperature range of (298.15-363.15) K (modulated differential scanning calorimetry technique, MDSC, calibrated on synthetic sapphire (CAS: 1317-82-4, TA Instruments, ultrapure in accordance to the reference standards NIST SRM 720) and checked with 1-hexyl-3-methylimidazolium bis(trifluoromethylsulfonyl)imide as a classified ionic liquid NIST standard [48], under a nitrogen gas flow of $50 \text{ cm}^3 \text{ min}^{-1}$, heating rate $dT/dt = 3 \text{ K min}^{-1}$, amplitude $\pm 0.5 \text{ K}$ and a modulation period 60 s. The results of calibration and post-calibration procedure ascertained that the relative standard uncertainty of measurement as u(c) = 3%, temperature standard uncertainty u(T) = 0.01 K. 3 independent repeats of measurement were performed with 5% repeatability. The specific heat capacity reported in our previous work was measured by DSC equipment Q1000 TA Instruments which enables the automatic heat capacity determination [21, 42]. In this work we used DSC equipment Q100 TA Instruments which does not allow this measurement, however, the results obtained are similar to those reported previously. Therefore, for pure ionic liquids the data from our previous work are used. The specific heat capacity was correlated with temperature with second-order equation:

$$c_{p}(T) = \sum_{i=0}^{2} a_{i} T^{i}$$
 (5)

where a is the regression parameter determined by the least-square method, along with the standard uncertainty, δa , while the coefficient of determination, R^2 , was used to describe the fit. Moreover, the heat capacity of ionanofluids (more specifically the enhancement) was predicted using the following empirical correlations for carbon nanotubes, boron nitride- and graphite-doped ionanofluids [49]:

$$\frac{\lambda_{INF}}{\lambda_{IL}} = (6.00 \pm 0.43) \varphi_{MWCNT} + (32.6 \pm 8.7) \cdot 10^{-4} T - (109 \pm 14) \varphi_{MWCNT}^2 - (5.5 \pm 1.3) \cdot 10^{-6} T^2 + (0.55 \pm 0.14)$$
 (6)

$$\frac{\lambda_{INF}}{\lambda_{IL}} = (28.99 \pm 0.85)\varphi_{BN} + (17.0 \pm 1.5) \cdot 10^{-4}T - (894 \pm 34)\varphi_{BN}^2 - (2.5 \pm 2.1) \cdot 10^{-6}T^2 + (0.70 \pm 0.24)$$
(7)

$$\frac{\lambda_{INF}}{\lambda_{IL}} = (29.5 \pm 1.2)\varphi_G + (23.4 \pm 2.3) \cdot 10^{-4}T - (829 \pm 42)\varphi_G^2 - (2.5 \pm 1.3) \cdot 10^{-6}T^2 + (0.58 \pm 0.38)$$
(8)

Thermal conductivity

The thermal conductivity measurements were performed using a KD2 Pro Thermal Properties Analyzer (Decagon company), based on transient hot-wire principle (KS-1 sensor, 60 mm length, 1.3 mm diameter) with the standard uncertainty of temperature, u(T) = 0.05 K. To ensure that the proper heat transfer occurred between the sensor and investigated material during the measurement, an approximate volume of sample of 30 cm³ was used. The cell was completely closed against the environment, immersed in bath circulator (Grant TC120, 50 % (V/V) ethylene glycol/water, thermal stability ±0.05 K and uniformity ±0.1 K). The time gap between each measurement after the temperature stabilisation was at least 15 min. The temperature range of measurement was (~278 to ~358) K. To obtain accurate results, the calibration procedure, as described in our previous work, was used [21] using glycerine and its water solutions (as a range of viscosity), water (as a range of high thermal conductivity), NaCl water solutions (as electrical conductive materials), toluene (as low viscous material). The calibration constant obtained was 0.9932 ± 0.0075, which results in calibration constant standard uncertainty of 1.50 %, and standard uncertainty of thermal conductivity values of about 3.44% [49]. Finally, the calibration was checked with the well-described (in terms of thermal conductivity) ionic liquid $[C_8C_1Im][NTf_2]$, achieving an excellent repeatability of 0.27 % [21]. The thermal conductivity, λ , was also correlated with temperature by the following linear equation:

$$\lambda(T) = \sum_{i=0}^{1} a_i T^i \tag{9}$$

where a_i is the fitting coefficient. The standard deviation of a_i parameters, δa_i , were calculated based on the least-square approach, and coefficient of determination, R^2 , was used to describe the fit. To predict the thermal conductivity of ionanofluids (more specifically enhancement), the only available model for ionic liquid – based nanofluids, Atashrouz model, was used [40]:

$$\frac{\lambda_{INF}}{\lambda_{IL}} = \left(\frac{\lambda_{NP}}{\lambda_{IL}}\right)^{\varphi_{NP}} \tag{10}$$

where the thermal conductivity coefficients of nanoparticles are (3223.4, 874.1, 35.7 and ~12.0) W m⁻¹ K⁻¹ for carbon nanotubes [50], boron nitride [51], graphite [52], and mesoporous carbon [53], respectively.

Thermogravimetric analysis

The thermogravimetric analysis (TGA) was carried out on TA Instruments TGA 550 equipment with platinum-HT pan (calibrated on Curie point with alumel and nickel, nitrogen gas flow 10 cm³ min⁻¹, heating rate of 5 K min⁻¹, samples mass of approximately 25 mg, weighting standard uncertainty of 0.01%, standard uncertainty of signal resolution 0.01 μ g, standard uncertainty of temperature u(T) = 0.5 K). An important factor influencing the significance of thermogravimetric results is the methodology to determine the onset temperature. Differentiation of the obtained thermogravimetric curve produces a Gaussian-like peak function which is used to determine the onset temperature as the crossing point of baseline and extrapolated linear function, while the percentage of weight loss is an equivalent to curve area. Furthermore, as reported by Maton et al. [54], T_{on} is determined by the derivative function (dm/dT vs. T) which has been recognized as a sufficiently accurate parameter of thermal stability, and this is commonly used to describe the thermal decomposition [55, 56]. Moreover, the overestimation of the onset temperature is reduced. Isothermal thermogravimetric studies were also conducted for [$P_{14,6,6,6}$][DecO] (temperatures of 400 K, 450 K, 500 K, 550 K, for 200 min).

Economic analysis

The information concerning whether the considered heat transfer fluids have sufficiently promising properties is discussed as a combination of all the physical properties measured (thermal conductivity, isobaric heat capacity, density and viscosity). There are a few approaches for such assessment, for example Mouromtseff introduced a factor as an implicit function with some basic limitations [57], and Murakami and Mikic proposed figure-of-merit property [58]. Unfortunately, the latter is not sensitive enough to distinguish particular commercial heat transfer fluids, the ionic liquids or ionanofluids [58]. The most novel approach was proposed by Mendonca et al. [59]. Therein, the authors considered an exemplary application as shell and tube heat transfer unit type, prescribing the external constraints and neglecting the pressure drop across the fluid ducts as it does not affect the performance. Thereafter, the only factor which has the influence is heat transfer area expressed by Newton's law of cooling:

$$Q = U_0 A_0 (\Delta T)_{lm} \tag{11}$$

where Q is rate of heat transfer (Q=1 MW), U_0 is overall heat transfer coefficient, A_0 is heat transfer area, and $(\Delta T)_{lm}$ is logarithmic mean temperature difference between inlet and outlet stream temperatures $((\Delta T)_{lm} = 20 \text{ K})$. In the case of circular tubes, U_0 can be calculated by the following equation:

$$\frac{1}{U_0} = \frac{D_o}{h_i D_i} + \frac{1}{h_o} + r_w + r_o + \frac{r_i D_o}{D_i}$$
(12)

where D_a is outside tube diameter ($D_a = 0.020$ m), h_i is the heat transfer coefficient for the inside film of fluid, D_i is the inside tube diameter (D_i =0.018 m), h_o is heat transfer coefficient for the outside film of fluid $(h_a = 2000 \text{ W m}^{-2} \text{ K}^{-1}), r_w$ is the thermal resistance of the tube wall, r_a is the outside fouling resistance, r_i is the inside fouling resistance. This can be also simplified in the following form:

$$\frac{1}{U_0} = \frac{1}{h_o} + \frac{D_o}{h_i D_i} + R \tag{13}$$

where R stands for all the factors related to the resistance (combined resistance constant, therefore R = 0 indicates an ideal flow by placing baffles in the shell part of the exchanges and controlling the fluid velocity in the tubes) [60]. The major assumption in this methodology is that turbulent flow is required for the efficient and accurate optimization. The most recognized correlation for the heat transfer coefficients for smooth circular tube was proposed by Sieder and Tate [61]:

$$Nu = \frac{h_i D_i}{\lambda} = 0.027 Re^{0.8} Pr^{\frac{1}{3}} \left(\frac{\eta}{\eta_w}\right)^{0.14}$$
 (14)

where Nu is the Nusselt number, λ is the thermal conductivity coefficient, η is the dynamic viscosity, η_w is the dynamic viscosity at the wall temperature (assumption: $\eta_{w} = \eta$), Re is the Reynolds number, Pr is the Prandtl number. As Eq. (13) assumes that turbulent flow is required, the Eq. (14) is applicable for values >100 [62]. Pr number can be calculated by the following equation:

$$Pr = \frac{c_p \eta}{\lambda} \tag{15}$$

where c_n is the specific isobaric heat capacity. Whereas the Reynolds number can be determined as follows:

$$Re = \frac{\rho u D_i}{\eta} \tag{16}$$

where ρ is the density, u is the mean velocity over the tube cross section ($u = 0.5 \text{ m s}^{-1}$). França et al. reported that an efficient way to study the cost of a new design was to divide it into five parts, namely the battery limits, utility, off-site, engineering fees and working capital [60]. The main influence herein is the battery limits investment which is related to the cost of individual parts of the equipment, along with the installation [63]. This cost, C_F , is a function of the size (heat transfer area), material of the construction, design pressure, temperature:

$$C_E = C_B \left(\frac{X}{X_B}\right)^m \tag{17}$$

where C_R is the cost of a reference equipment, X is the capacity of the equipment (in this case heat transfer area, A_0), X_B is the capacity of the reference equipment (heat transfer area of reference) and m is a constant depending on the equipment type (m = 0.68). Following the work of França et al., the reference equipment is carbon steel shell and tube heat exchanger with a reference heat transfer area of 80 m² and cost of (3.28 · 10⁴) US\$ [63]. When the factors of materials type, pressure and temperature are included in Eq. (17), the following equation is obtained:

$$C_E = C_B' \left(\frac{X}{X_B}\right)^m f_M f_P f_T \tag{18}$$

where C_B' is the cost of the above described reference equipment ($C_B' = 3.28 \cdot 10^4 \text{ US}$ \$), f_M is the correction factor for materials type ($f_M = 3.4$), f_P is the correction factor for pressure ($f_P = 1.5$), and f_T is the correction factor for temperature (f_T =1.3) [63]. Finally, the equation for the cost of the heat exchange unit is derived as follows:

$$C_E = 3.28 \cdot 10^4 \left(\frac{A_0}{80}\right)^{0.68} \cdot 3.4 \cdot 1.5 \cdot 1.3 = 2.18 \cdot 10^5 \left(\frac{A_0}{80}\right)^{0.68}$$
 (19)

Statistical analysis

All the experimental and calculated values were compared to each other using the relative deviation, RD, and absolute average relative deviation, AARD values:

$$RD = \frac{100(x_{\text{calc}} - x_{\text{exp}})}{x_{\text{exp}}} \tag{20}$$

$$AARD = \frac{100}{n} \sum_{i=1}^{n} \left| \frac{x_{\text{calc}} - x_{\text{exp}}}{x_{\text{exp}}} \right|$$
 (21)

To ascertain the influence of the thermophysical properties, the uncertainties in the enhancement values, the errors of isobaric heat capacity, density and viscosity enhancements, ε , were calculated by using the chain rule of differentiation:

$$\delta \varepsilon = \left| \frac{\partial \varepsilon}{\partial c_{p,INF}} \right| \delta c_{p,INF} + \left| \frac{\partial \varepsilon}{\partial c_{p,IL}} \right| \delta c_{p,IL}$$
(22)

In the case of the thermal conductivity enhancement, the sample covariance can be neglected (due to low numerical values of thermal conductivity and the standard uncertainty, consequently), therefore, excessive numerical artefacts are not propagated:

$$\delta \varepsilon = \sqrt{\left(\frac{\partial \varepsilon}{\partial \lambda_{INF}}\right)^2 (\delta \lambda_{INF})^2 + \left(\frac{\partial \varepsilon}{\partial \lambda_{IL}}\right)^2 (\delta \lambda_{IL})^2}$$
 (23)

Results and discussion

Density

The density, ρ , was measured as a function of temperature over the range of (298.15–363.15) K. All the results for the pure ionic liquids were previously reported in [21]. The experimental data can be found in Supporting Information SI2, also presented in Fig. 2a. The comparison of values in this work and those in

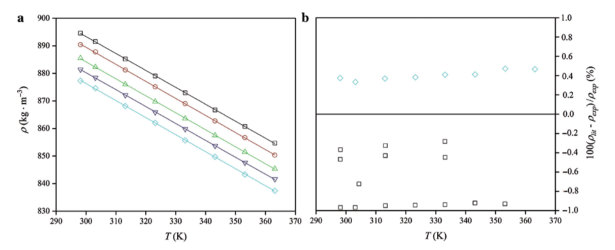


Fig. 2: (a) Experimental data of density, ρ , as a function of the temperature, T, for pure ILs; (b) the relative deviations of density between this work and literature, for $[P_{14,6,6,6}][AcO]$, \Box [64-66]; $[P_{14,6,6,6}][ButO]$, \bigcirc ; $[P_{14,6,6,6}][HexO]$, \triangle ; $[P_{14,6,6,6}][OctO]$, ∇ ; $[P_{14,6,6,6}][ButO]$, \bigcirc $[DecO] \bigcirc [67].$

the literature can be found in Fig. 2b. Relative deviations of (-0.37 [64], -0.45 [65], -0.98 [66])% for $[P_{14,6,6,6}]$ [AcO], and 0.38 % for $[P_{14,6.6.6}]$ [DecO] [67], at 298.15 K, were obtained. The purity of ILs can be a significant factor influencing the thermophysical properties, for example Seddon et al. reported that the residual solvent and halide content affect the values of density, therefore, those should be monitored. Tariq et al. reported the chloride content of $[P_{14,6,6,6}]$ [AcO] in a range of (20–150) ppm, whilst in this work the chloride content was below 5 ppm [64]. Esperança et al. used $[P_{14,6,6,6}]$ [AcO] from the same source as Tariq et al. [65], and it can be seen that the results are consistently similar - negative relative deviations (which indicated higher chloride content of the value from literature, especially by comparing the density of [P_{1/666}][Cl] and those reported in this work) [68]. Similar results can be seen in the work of Fillion et al. [66]. On the other hand, Neves et al. did not report the chloride content [67]. However, the water content reported by Neves et al. is almost 3 times higher than the value reported in this work [67]. This also in a good agreement with the density-water dependence reported by Seddon et al. [69]. As can be seen, the purity of ILs used in this work is very high (low water and chloride contents), in comparison to other reports, therefore, the results presented herein are of high quality.

The comparison of density values to those of commercial heat transfer fluids depends on the type of material, for example the values in this work are similar to synthetic aromatic hydrocarbon mixtures (Therminol ADX10 or Dynalene SF), slightly lower than water and significantly lower than glycol-based HTFs (for example Dowtherm 4000 or Dynalene EG series) [70-72].

The density was also correlated with temperature. The parameters of the second-order equation, $\rho(T)$, can be found in Table S1 (Supporting Information SI3). The density was found to be a decreasing function of temperature as noted for other ILs, while the differences between the lowest (298.15 K) and the highest temperature (363.15 K) are similar for all ILs, (4.67, 4.72, 4.74, 4.73 and 4.76) % for $[P_{14,6,6,6}][AcO]$, $[P_{14,6,6,6}][ButO]$, $[P_{14,6,6,6}][HexO], [P_{14,6,6,6}][OctO]$ and $[P_{14,6,6,6}][DecO],$ respectively.

The density of the ionic liquids containing nanoparticles were also measured (Fig. S7-S11, Supporting Information SI1 and Excel File Supporting Information SI2). The parameters of the $\rho(T)$ equation can be found in Table S1 (Supporting Information SI3). The values of the slope for all systems are very similar for the pure ionic liquids and the ionanofluids.

Solids usually have higher densities than liquids, particularly the investigated nanoparticles (as investigated previously for multiwalled carbon nanotubes, boron nitride and graphite in [42]), therefore, the density of ionic liquid and nanoparticle mixtures have higher values than the pure ionic liquids. This can be observed as enhancement in Fig. 3. It can be seen that the density dependence on the nanoparticles concentration is almost linear for all systems studied.

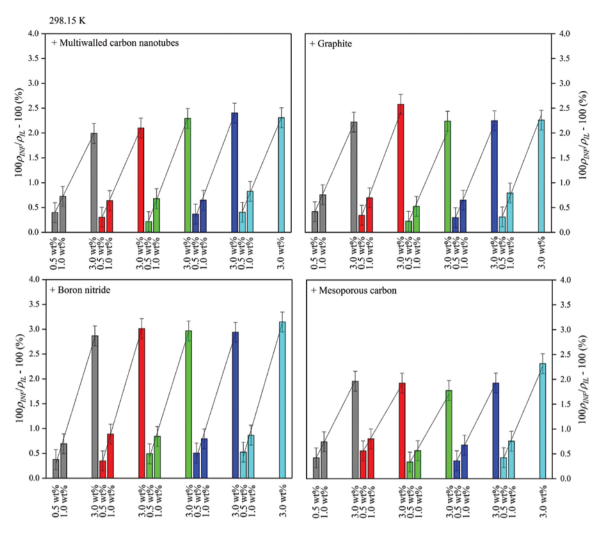


Fig. 3: Density enhancement of ionanofluids, $(100\rho_{NF}/\rho_{1L}-100)$, in comparison to pure ionic liquids, over the mass fraction, for grey $-[P_{14,6,6,6}][AcO]$ INF, red $-[P_{14,6,6,6}][ButO]$ INF, green $-[P_{14,6,6,6}][HexO]$ INF, blue $-[P_{14,6,6,6}][OctO]$ INF, cyan $-[P_{14,6,6,6}][DecO]$ INF, at 298.15 K.

The enhancement values at temperatures 298.15 K and 363.15 K are shown in Fig. 3 and Table 2, respectively. The difference between the enhancements at 298.15 K and 363.15 K is very small and generally only slightly increases with temperature. More specifically the maximum range of enhancement was found as (2.40-1.91, 2.58-2.22, 3.15-2.87 and 2.32-1.77)% for MWCNT, G, BN and MC at 298.15 K, respectively, and (2.44-2.00, 2.70-2.32, 3.33-3.00 and 2.42-1.86)% for MWCNT, G, BN and MC at 363.15 K, respectively while the average enhancements error is $(\pm 0.2\%)$.

The influence of the ionic liquid structure on the intermolecular interactions has been widely investigated in the literature [21, 73–75]. A significant change in the predominant intermolecular interaction occurs at hydrocarbon chain length of ~6 where the pure Coulombic interactions become significantly affected by van der Waals interactions, and/or the volume of anions starts to be an influencing factor. The effect of anion on the enhancement can be studied in this work, as several anions with various chain lengths were studied. França et al. studied the interactions between ions in ILs and nanoparticles (single- and multi-walled carbon nanotubes and graphene), where the driving force for the enhancement was showed to be anion dependent, which they are adsorbed on the surface of nanoparticles more easily (related to polarizability of nanoparticles and charge distribution of ions) [76]. In relation to results observed in this work, it was shown that the enhancement decreases with anion chain length to hexanoate anion (negligible effect of anion chain length

Table 2: The enhancements of density, ρ , dynamic viscosity, η , thermal conductivity, λ , isobaric heat capacity, c_{v} , for ionanofluids in comparison to pure ionic liquids.

Enhnancement (%)	Density (±0.20 %)		Viscosity (0.5 wt % = \pm 2.4 %, 1.0 wt % = \pm 9.0 %, 3.0 wt % = \pm 13.3 %)		Thermal conductivity (±4.5 %)		Heat capacity (± 5.3 %)	
	298.15 K	363.15 K	298.15 K	363.15 K	278.15 K	358.15 K	298.15 K	363.15 K
[P _{14,6,6,6}][AcO] + 0.5 wt % MWCNT	0.40	0.45	19.1	27.9	2.8	0.8	6.1	3.6
[P _{14.6.6.6}][AcO] + 1.0 wt % MWCNT	0.72	0.72	41.3	41.5	6.3	3.2	7.1	4.7
[P _{14,6,6,6}][AcO] + 3.0 wt % MWCNT	1.99	2.05	116.2	135.4	19.9	18.0	11.7	9.3
[P _{14,6,6,6}][AcO] + 0.5 wt % G	0.42	0.48	23.0	28.0	2.0	1.4	11.0	14.5
[P _{14,6,6,6}][AcO] + 1.0 wt % G	0.76	0.83	39.7	50.4	4.4	4.5	17.1	21.9
[P _{14,6,6,6}][AcO] + 3.0 wt % G	2.22	2.36	121.9	155.9	11.3	12.3	29.2	34.5
[P _{14,6,6,6}][AcO] + 0.5 wt % BN	0.38	0.42	12.8	14.7	2.0	1.3	3.9	4.5
[P _{14,6,6,6}][AcO] + 1.0 wt % BN	0.69	0.75	28.5	31.0	4.4	3.9	10.9	11.5
[P _{14,6,6,6}][AcO] + 3.0 wt % BN	2.87	3.00	119.5	93.4	14.9	13.0	21.6	21.5
[P _{14,6,6,6}][AcO] + 0.5 wt % MC	0.42	0.39	26.5	17.2	2.0	0.6	8.8	11.6
[P _{14,6,6,6}][AcO] + 1.0 wt % MC	0.74	0.73	51.4	40.7	3.8	3.2	16.7	20.8
$[P_{14,6,6,6}][AcO] + 3.0 \text{ wt }\% \text{ MC}$	1.96	2.06	117.5	94.2	8.8	7.2	25.1	28.1
$[P_{14,6,6,6}][ButO] + 0.5 \text{ wt }\% \text{ MWCNT}$	0.30	0.27	21.8	23.0	3.8	4.6	5.2	3.1
[P _{14,6,6,6}][ButO] + 1.0 wt % MWCNT	0.64	0.62	50.5	52.0	5.7	6.6	7.5	3.6
[P _{14,6,6,6}][ButO]+3.0 wt % MWCNT	2.10	2.14	156.7	154.0	22.6	21.4	10.3	7.0
[P _{14,6,6,6}][ButO]+0.5 wt % G	0.34	0.34	16.4	20.1	2.6	3.9	12.1	15.9
[P _{14,6,6,6}][ButO]+1.0 wt % G	0.70	0.71	29.5	38.9	5.1	5.9	18.3	22.4
[P _{14,6,6,6}][ButO]+3.0 wt % G	2.58	2.70	130.2	137.9	9.5	10.7	31.5	36.1
[P _{14,6,6,6}][ButO]+0.5 wt % BN	0.35	0.35	16.1	20.5	1.9	1.9	3.5	4.1
[P _{14,6,6,6}][ButO]+1.0 wt % BN	0.89	0.90	42.1	58.3	5.1	5.3	9.6	10.3
[P _{14,6,6,6}][ButO]+3.0 wt % BN	3.01	3.11	102.0	119.5	15.1	16.0	20.6	20.8
[P _{14,6,6,6}][ButO]+0.5 wt % MC	0.56	0.56	15.9	21.6	2.6	1.8	10.1	13.3
[P _{14,6,6,6}][ButO] + 1.0 wt % MC	0.80	0.81	39.2	53.3	5.1	3.5	15.0	18.7
[P _{14,6,6,6}][ButO] + 3.0 wt % MC	1.92	1.98	136.3	135.5	11.4	9.3	27.4	30.9
[P _{14,6,6,6}][HexO] + 0.5 wt % MWCNT	0.22	0.23	28.8	31.5	3.8	2.3	5.7	3.3
[P _{14,6,6,6}][HexO] + 1.0 wt % MWCNT	0.68	0.71	58.4	66.9	5.7	5.0	7.3	3.5
[P _{14,6,6,6}][HexO]+3.0 wt % MWCNT	2.29	2.29	133.1	138.2	21.4	20.0	11.4	8.1
[P _{14,6,6,6}][HexO]+0.5 wt % G	0.22	0.20	19.5	23.6	3.1	3.7	11.5	15.2
[P _{14,6,6,6}][HexO] + 1.0 wt % G	0.52	0.51	50.6	63.6	5.0	6.4	18.7	23.1
[P _{14,6,6,6}][HexO] + 3.0 wt % G	2.24	2.32	143.6	145.4	10.1	11.2	30.3	35.3
[P _{14,6,6,6}][HexO] + 0.5 wt % BN	0.49	0.52	25.9	24.7	3.8	5.0	3.7	4.3
[P _{14,6,6,6}][HexO] + 1.0 wt % BN	0.84	0.88	47.2	53.0	5.0	5.7	11.5	12.0
[P _{14,6,6,6}][HexO] + 3.0 wt % BN	2.96	3.08	99.1	102.5	15.1	17.3	20.3	20.6
[P _{16,6,6,6}][HexO] + 0.5 wt % MC	0.33	0.36	21.0	25.7	1.9	2.3	9.5	12.4
[P _{14,6,6,6}][HexO] + 1.0 wt % MC	0.56	0.60	44.4	54.0	3.2	3.0	15.9	19.8
[P _{14,6,6,6}][HexO] + 3.0 wt % MC	1.56	1.86	97.4	108.4	10.1	10.3	27.0	30.2
$[P_{14,6,6,6}][OctO] + 0.5 wt % MWCNT$	0.37	0.36	17.8	20.2	1.9	1.6	5.1	2.8
[P _{14,6,6,6}][OctO] + 1.0 wt % MWCNT	0.65	0.69	36.8	40.1	7.4	7.1	6.2	4.6
[P _{14,6,6,6}][OctO] + 3.0 wt % MWCNT	2.40	2.45	92.9	98.5	19.8	19.1	11.5	10.0
[P _{14,6,6,6}][OctO]+0.5 wt % G	0.29	0.34	17.2	31.3	1.9	1.6	9.8	14.0
[P _{14,6,6,6}][OctO]+1.0 wt % G	0.65	0.71	43.6	59.1	4.7	3.6	15.8	19.9
$[P_{1/(6,6,6)}][OctO] + 3.0 \text{ wt } \% \text{ G}$	2.25	2.38	121.0	159.3	12.0	11.6	28.7	32.7
[P _{14,6,6,6}][OctO] + 0.5 wt % BN	0.51	0.50	24.1	27.7	2.5	3.6	3.5	4.5
[P _{14,6,6,6}][OctO] + 1.0 wt % BN	0.79	0.78	38.3	46.1	5.6	5.6	1.4	12.6
[P _{14,6,6,6}][OctO]+3.0 wt % BN	2.94	3.08	78.2	94.5	14.9	15.6	20.3	20.8
$[P_{14,6,6,6}][OctO] + 0.5 \text{ wt }\% \text{ MC}$	0.36	0.36	13.5	28.5	1.8	1.4	10.0	12.7
[P _{14,6,6,6}][OctO] + 1.0 wt % MC	0.68	0.80	33.1	55.7	3.8	3.2	16.2	18.3
[P _{14,6,6,6}][OctO] + 3.0 wt % MC	1.93	2.08	97.3	92.6	7.6	9.5	25.2	27.5
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[P _{14,6,6,6}][DecO] + 0.5 wt % MWCNT	0.40	0.41	27.6	25.3	3.3	2.5	5.2	3.3

Table 2 (continued)

Enhnancement (%)	Density (±0.20 %)		Viscosity (0.5 wt % = \pm 2.4 %, 1.0 wt % = \pm 9.0 %, 3.0 wt % = \pm 13.3 %)		Thermal conductivity (±4.5 %)		Heat capacity (± 5.3 %)	
[P _{14,6,6,6}][DecO] + 3.0 wt % MWCNT	2.31	2.44	137.8	144.9	22.0	20.6	11.6	8.9
$[P_{14,6,6,6}][DecO] + 0.5 \text{ wt } \% G$	0.31	0.37	20.6	23.6	3.4	3.3	9.8	14.2
[P _{14,6,6,6}][DecO] + 1.0 wt % G	0.79	0.88	50.9	57.0	5.9	6.4	18.7	22.1
$[P_{14,6,6,6}][DecO] + 3.0 \text{ wt } \% \text{ G}$	2.26	2.41	121.0	135.0	20.6	8.8	29.0	32.7
$[P_{14,6,6,6}][DecO] + 0.5 \text{ wt } \% \text{ BN}$	0.52	0.58	26.1	38.8	1.9	2.5	4.3	4.2
[P _{14,6,6,6}][DecO] + 1.0 wt % BN	0.87	0.94	52.0	60.7	6.2	7.0	11.4	12.2
[P _{14,6,6,6}][DecO] + 3.0 wt % BN	3.15	3.33	106.4	117.2	13.2	13.5	20.5	20.7
[P _{14,6,6,6}][DecO] + 0.5 wt % MC	0.42	0.51	32.6	38.9	1.7	0.8	8.8	11.7
[P _{14,6,6,6}][DecO] + 1.0 wt % MC	0.75	0.80	57.9	69.0	4.3	2.9	15.4	18.8
$[P_{14,6,6,6}][DecO] + 3.0 \text{ wt } \% \text{ MC}$	2.32	2.42	144.0	153.1	9.2	7.3	26.4	29.4

size but caused by the bulkier properties so as a result less favoured adsorption on the surface of nanoparticles), and then increasing again (driven by decreasing the electrostatic forces between cations and anions, therefore, more favoured adsorption onto the surface of nanoparticles).

The density of ionanofluids was previously investigated for a range of ionic liquids, 1-butyl-3-methyl-imidazolium dicyanamide, 1-butyl-3-methylimidazolium bis[(trifluoromethyl)sulfonyl]imide, 1-butyl-1-methylpyrrolidinium bis[(trifluoromethyl)sulfonyl]imide, 1-hexyl-3-methylimidazolium hexafluorophosphate, 1-ethyl-3-methylimidazolium ethylsulfate, with multiwalled carbon nanotubes, boron nitride and graphite [42]. Similar results were found therein namely that the anion size of the ionic liquid was the influencing factor for the enhancement values. The enhancement, as presented above, was also found to increase by increasing the nanoparticles size as follows: MWCNT < MC < BN < G.

The highest value of density repoted in this work is 920.24 kg m $^{-3}$ for $[P_{14,6,6,6}][AcO] + 3$ wt % BN at 298.15 K compared with 894.6 kg m $^{-3}$ of the pure ionic. Even though the addition of nanoparticles to ionic liquids increases the density, the enhancement is still not a vast improvement when compared to commercial heat transfer fluids.

The density of ionanofluids was also calculated by using the additivity model Eq. (2). Unfortunately, the data for pure mesoporous carbon were not available, therefore, similar calculations to those in the work of Oster et al. were needed to be performed [42] which are reported in Supporting Information SI2, and the density for mesoporous carbon is 1477.50 kg m⁻³ at 298.15 K. It was found that the density of carbonaceous nanoparticles is similar, to remind, the density values for investigated carbon nanotubes and graphite are (1626.34 and 1711.99) kg m⁻³ at 298.15 K. It can be seen that the density of mesoporous carbon is very similar to those previously studied, whilst the slight decrease is caused by the high porosity. The deviations between experimental and calculated values can be found in Fig. 4. It should be noted that the data of those ionanofluids containing mesoporous carbon were calculated based on the determined density in this work.

Average absolute relative deviations of (0.21, 0.31, 0.32 and 0.10) % were found for MWCNT, G, BN and MC, respectively and maximum deviations of (0.66, 0.67, 0.58 and 0.31) % were found for $[P_{14,6,6,6}][HexO] + 3$ wt % MWCNT at 363.15 K, $[P_{14,6,6,6}][HexO] + 3$ wt % G at 363.15 K, $[P_{14,6,6,6}][OctO] + 3$ wt % BN at 358.15 K and $[P_{14,6,6,6}][DecO] + 3$ wt % MC at 348.15 K, respectively. Interestingly, all relative deviations between experimental and calculated values are positive. The reason for this might originate from the existing interfacial nanolayering of higher density values (which shifts the calculated results to higher values) and is in a good agreement with work of Shin et al. [25].

Such deviations illustrate that we are able to calculate the density of ionanofluids with high accuracy, with an error of <1% which is good considering that it is a complex system of two phases mixture (nano-solid+liquid). It should be also noted that the calculations for the pure ionic liquids densities with the most accurate approach, i.e. the fluctuation theory-based Tait-like equation of state (FT-EoS), have deviations of up to 6% [77].

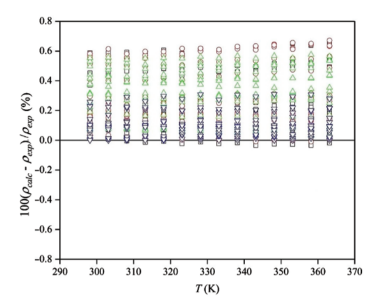


Fig. 4: Experimental vs. calculated density of ionanofluids as relative deviations against the temperature, for $[P_{14,6,6,6}][RO]$ + MWCNT, \square ; $[P_{14,6,6,6}][RO] + G$, \bigcirc ; $[P_{14,6,6,6}][RO] + BN$, \triangle ; $[P_{14,6,6,6}][RO] + MC <math>\nabla$.

Dynamic viscosity

The dynamic viscosity, η , was measured as a function of the temperature over the range (298.15–363.15) K, as reported previously [21]. The experimental data for pure ionic liquids can be found in Supporting Information SI2, and Fig. 5a. The measured viscosity was also compared to the previously reported data (Fig. 5b) for the two studies available for $[P_{14,6,6,6}][AcO]$ and $[P_{14,6,6,6}][DecO]$ [66, 67]. The maximum observed deviations are 106.93% for $[P_{14,6,6,6}]$ [AcO] (196.2 mPa s in this work and 406 mPa s in [66], at 298.15 K) and 15.18 % for $[P_{14,6,6,6}]$ [DecO] (409.8 mPa s in this work and 472 mPa s in [67], at 298.15 K).

In addition, the viscosity values of investigated systems are higher than commercial HTFs for example (16.9, 6.19 or 3.71) mPa s for Dowtherm MX, Dynalene PG60 or Dowtherm A at 298.15 K, respectively [71, 72].

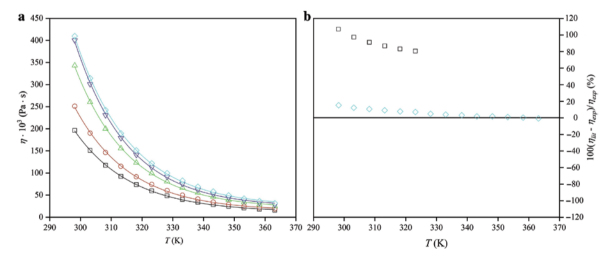


Fig. 5: (a) Experimental data of viscosity, η , as a function of the temperature, T, for pure ILs; (b) the deviations of viscosity between this work and literature for $[P_{14,6,6,6}][AcO]$, \Box [66]; $[P_{14,6,6,6}][ButO]$, \bigcirc ; $[P_{14,6,6,6}][HexO]$, \triangle ; $[P_{14,6,6,6}][OctO]$, ∇ ; $[P_{14,6,6,6}][P_{14,6,6,$ $[DecO] \bigcirc [67].$

Viscosity was found to depend on the anion chain length in the following sequence: $[P_{14,6,6,6}][AcO] < [P_{14,6,6,6}][ButO] < [P_{14,6,6,6}][DecO]$, as expected [78].

The viscosity was correlated with temperature by exponential VFT equation of which parameters can be found in Table S1 (Supporting Information SI3). The effect of temperature on viscosity is enormous in ILs (around 1000 % decrease from 298.15 K to 363.15 K) and, at high temperature the viscosity of ILs and commercial HTFs become similar (at 423.15 K, 0.96 mPa s for Dowtherm MX and 4.65 mPa s for $[P_{14,6,6,6}][AcO]$) [72]. The analysis of each VFT equation leads to the most recognized parameter of so-called glass transition, T_0 , at which a sharp (super-Arrhenius) viscosity occurs when the system is cooled toward the glass transition, T_0 , at which a sharp (super-Arrhenius) viscosity increase can be observed, accompanied with very little structural change, as discussed by Mauro et al. [79]. As a further consequence, below the glass transition temperature, the long structural relaxation times can be caused by a lower mobility of the species. Using the results summarized in Table S2 (Supporting Information SI3), it can be observed that an increase of the anion chain length of ionic liquids results in a decrease of T_0 which reflects to a decrease of their mobility.

The viscosity was also measured for the ionic liquids mixed with nanoparticles (Fig. S12–S16, Supporting Information SI1 and data are reported in Supporting Information SI2). It is well-known that the viscosity increases by adding nanoparticles in ILs [80, 81], as shown herein in Fig. 6 and in Table 2 for the data collected at 298.15 K, and at 298.15 and 363.15 K, respectively. This behaviour can be seen as an enhancement

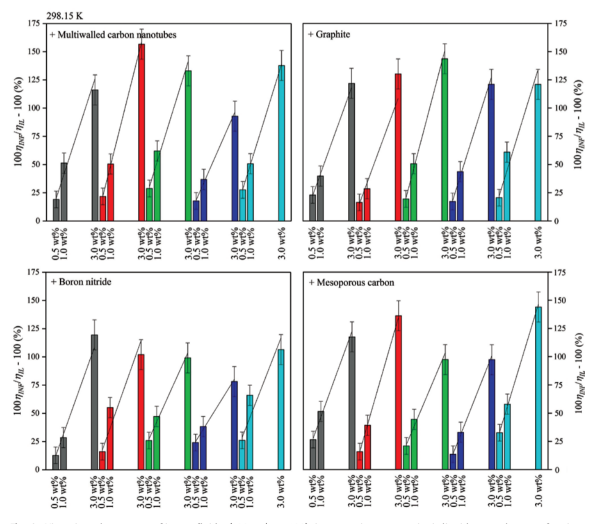


Fig. 6: Viscosity enhancement of ionanofluids, $(100\eta_{_{NP}}/\eta_{_{1L}}-100)$, in comparison to pure ionic liquids, over the mass fraction, for grey $-[P_{_{14,6,6,6}}][AcO]$ INF, red $-[P_{_{14,6,6,6}}][ButO]$ INF, green $-[P_{_{14,6,6,6}}][HexO]$ INF, blue $-[P_{_{14,6,6,6}}][OctO]$ INF, cyan $-[P_{_{14,6,6,6}}][DecO]$ INF, at 298.15 K.

in the ionanofluids viscosity in comparison to the pure ILs. Furthermore, as shown in Fig. 6, in most cases a linear correlation can be used to evaluate accurately the viscosity enhancement caused by the addition of nanoparticles in ILs. In addition, this viscosity enhancement was found to increase with temperature in the ranges of (17.78–28.80, 16.37–22.98, 12.78–26.06, 13.51–32.63) % for 0.5 wt % MWCNT, G, BN and MC at 298.15 K, respectively, and (20.22-31.49, 20.07-31.26, 14.73-38.76, 17.18-38.92) % for 0.5 wt % MWCNT, G, BN and MC at 363.15 K, respectively. For 1 wt % of MWCNT, G, BN and MC, the following enhancements were found (36.79-62.07, 28.52-60.93, 28.51-65.98, 33.07-57.89) % at 298.15 K, respectively, and (40.09-66.91, 38.88-63.63, 30.98-60.71, 40.67–68.97) % at 363.15 K, respectively. The enhancements at a nanoparticle concentration of 3.0 wt % MWCNT, G, BN and MC are much higher (92.88–156.70, 120.95–143.64, 78.15–119.47, 97.35–144.03) % at 298.15 K, respectively, and (98.49-153.96, 135.02-159.27, 93.36-119.49, 92.57-153.10) % at 363.15 K, respectively. It should be noted that the average enhancements errors are (2.4, 9.0, 13.3) % for systems with 0.5 wt %, 1.0 wt % and 3.0 wt % loading of nanoparticles, respectively, as explained in the Experimental part.

The VFT can give an insight into the dependence of the ionic liquid structure onto the viscosity (Table S1, Supporting Information SI3). The influence of pure ionic liquid structure was discussed above, moreover, the addition of the nanoparticles (as well as increasing those concentration) results in a further decrease in the T_0 . Unfortunately, there was no dependence found between the covered nanoparticle types and concentrations and their T_{o} .

Kinematic viscosity

For industrial purposes, the lubrication properties of the HTFs are very important. The selection of lubricating material for a specific application is essential. Materials are classified in several groups, depending on the lubrication abilities. The property on which the classification is performed is called kinematic viscosity, ν , calculated from dynamic viscosity and density ($\nu = \eta/\rho$) at 313.15 K [82–84]. The ISO classification (ISO 3448:1992) includes a wide range of materials from kinematic viscosity of 2.2 mm² s⁻¹ (ISO VG 2) to 1500 mm² s⁻¹ (ISO VG 1500) [85, 86].

The results of kinematic viscosity calculations and assignments to lubrication properties are shown in Fig. 7. The kinematic viscosities for pure ionic liquids were (104.2, 130.5, 177.9, 205.6 and 218.6) mm² s⁻¹ for

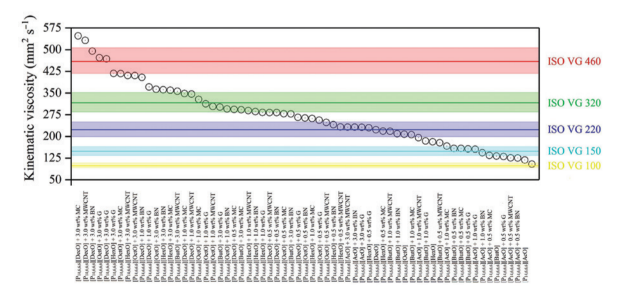


Fig. 7: The ISO lubrication properties classification based on the kinematic viscosity, the bold coloured lines represent the mid-point kinematic viscosity of the class, whereas the borders of coloured field represent the minimum and maximum kinematic viscosity of the class.

 $[P_{14,6,6,6}][AcO]$, $[P_{14,6,6,6}][ButO]$, $[P_{14,6,6,6}][HexO]$, $[P_{14,6,6,6}][OctO]$ and $[P_{14,6,6,6}][DecO]$, respectively. As can be seen, the kinematic viscosity is increasing with the anion chain length which is driven by the dynamic viscosity.

The addition of nanoparticles shifts the kinematic viscosity to higher values, as expected. The maximum kinematic viscosity was found as 547.5 mm² s⁻¹, corresponding to the mixture of $[P_{14,6,6,6}][DecO]+3$ wt % MC. The lubrication properties are also analysed based on the friction force coefficients, however, it does not relate to industrial assignment of material (in accordance to ISO classification). Nevertheless, the previous reports show that ionic liquids have promising lubricative properties (based on the friction coefficients) [87, 88]. In this work, we additionally showed that ionic liquids are very useful as lubricates (due to high variety in ISO classification), moreover, the possibility of doping the ionic liquids with nanoparticle enables the kinematic viscosity to be tailored, starting from relatively low kinematic viscosity (as 104.2 mm² s⁻¹ for $[P_{14,6,6,6}]$ [AcO]) up to higher values (as 547.5 mm² s⁻¹ for $[P_{14,66,6}][DecO]+3$ wt % MC).

Thermal conductivity

The thermal conductivity was measured in the temperature range of (278–358) K. The results for the pure ionic liquids were previously published in [21], also presented in Fig. 8a and collected in Supporting Information SI2. The coefficients of linear equation $\lambda(T)$ are reported in Table S2 (Supporting Information SI3).

To remind, the thermal conductivity decreases by increasing the temperature linearly, while the slopes for all ionic liquids remain similar. The thermal conductivity differences between the lowest and highest temperature investigated herein are (6.71, 7.24, 7.79, 7.62 and 7.23) % for $[P_{14,6,6,6}][AcO]$, $[P_{14,6,6,6}][ButO]$, $[P_{14,6,6,6}][DecO]$, respectively. By increasing the chain length on the anion from acetate to decanoate induces a linear increase on their volume [21]. In fact, a linear decrease in their thermal conductivity could be also expected (based on the approach for simple molecular solvents), however, no dependence was found for investigated ILs.

The difference in thermal conductivity between the largest and smallest anions is ~2.41% which is well below the standard uncertainty of the measurement (3.44%), therefore, it can be assumed that all thermal conductivity values are very similar irrespective of the investigated anion. Thermal conductivity was also measured for the ionanofluids with carbon nanotubes, boron nitride, graphite and mesoporous with a loading up to 3 wt%. The results are presented in Fig. S17–S21, (Supporting Information SI1), reported in Supporting Information SI2, and the coefficients of linear equation as a function of temperature can be found in Table S2 (Supporting Information SI3).

The thermal conductivity was found to linearly decrease with the temperature, and the same dependence was found for the nanoparticle-doped ionic liquids, while the slope was consistently the same, however, the

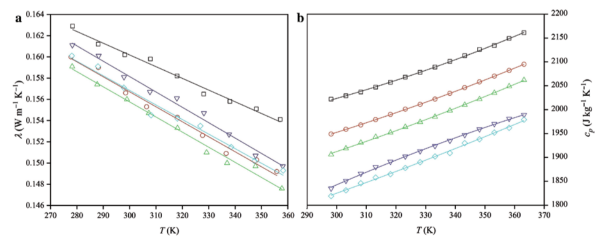


Fig. 8: (a) Thermal conductivity as a function of the temperature; (b) Isobaric heat capacity as a function of the temperature, for $[P_{14,6,6,6}][AcO]$, \Box ; $[P_{14,6,6,6}][ButO]$, \bigcirc ; $[P_{14,6,6,6}][HexO]$, \bigcirc ; $[P_{14,6,6,6}][DecO]$.

values of thermal conductivity were shifted to higher values. This gives, in fact, the opportunity to tune the thermal conductivity profile for a specific application, for example selection of IL influences the temperature profile whilst the addition of nanoparticles shifts the thermal conductivity to higher values.

The enhancements of thermal conductivity values caused by the addition of nanoparticles is shown in Fig. 9 for 278.15 K and Table 2 for 358.15 K. The thermal conductivity of solids is higher than liquids, and for nanoparticles particularly [25]. In the case of the investigated nanoparticles, the thermal conductivity coefficients are (3223.4, 874.1, 35.7 and ~12.0) W m⁻¹ K⁻¹ for carbon nanotubes, boron nitride, graphite and mesoporous carbon, respectively [51–53, 89]. Therefore, the expectation of mixing ionic liquids and nanoparticles is that the thermal conductivity of mixture is higher than for the pure IL. Moreover, increasing the amount of nanoparticle results in higher thermal conductivity. Furthermore, as shown in Fig. 9, this dependence is generally linear.

The range of maximum enhancements observed in this work is (19.81-22.62, 9.52-12.03, 15.12-13.22 and 7.64–11.36)% for MWCNT, G, BN and MC at 278.15 K, respectively, (17.98–21.37, 8.80–12.27, 12.97–17.27 and 7.19-10.27) % for MWCNT, G, BN and MC at 358.15 K, respectively. In general, the changes in the enhancements with temperature are below than the measurement errors (4.5%) and can be neglected. This supports the ascertain that the temperature profile of thermal conductivity originates from the liquid and not the nanoparticles.

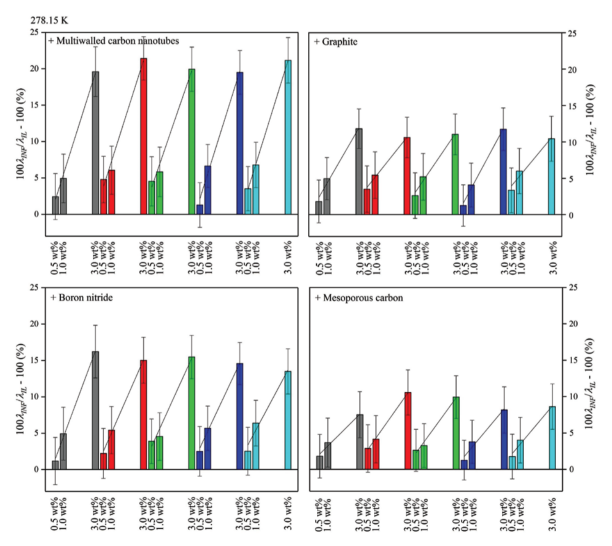


Fig. 9: Thermal conductivity enhancement of ionanofluids, $(100\lambda_{_{NP}}/\lambda_{_{R}}-100)$, in comparison to pure ionic liquids, over the $mass\ fraction,\ for\ grey-[P_{14,6,6,6}][AcO]\ INF,\ red-[P_{14,6,6,6}][ButO]\ INF,\ green-[P_{14,6,6,6}][HexO]\ INF,\ blue-[P_{14,6,6,6}][OctO]\ INF,\ cyan-[P_{14,6,6,6}][NF]\ fraction,\ for\ grey-[P_{14,6,6,6}][AcO]\ INF,\ red-[P_{14,6,6,6}][ButO]\ INF,\ green-[P_{14,6,6,6}][HexO]\ INF,\ blue-[P_{14,6,6,6}][OctO]\ INF,\ cyan-[P_{14,6,6,6}][NF]\ fraction,\ for\ grey-[P_{14,6,6,6}][AcO]\ INF,\ red-[P_{14,6,6,6}][ButO]\ INF,\ green-[P_{14,6,6,6}][HexO]\ INF,\ blue-[P_{14,6,6,6}][OctO]\ INF,\ cyan-[P_{14,6,6,6}][AcO]\ INF,\ cyan-[P_{14$ [P_{14,6,6,6}][DecO] INF, at 298.15 K.

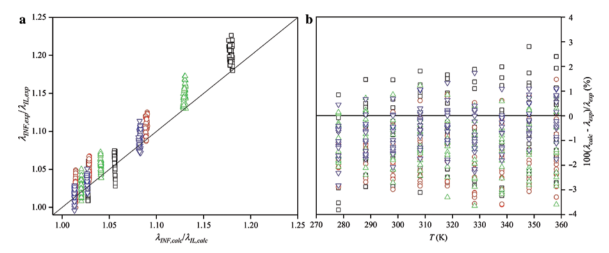


Fig. 10: (a) experimental vs. calculated thermal conductivity of ionanofluids; (b) relative deviations between calculated and experimental values of ionanofluids thermal conductivity against the temperature, for $[P_{14,6,6,6}][RO] + MWCNT$, \Box ; $[P_{14,6,6,6}][RO] + BN$, \triangle ; $[P_{14,6,6,6}][RO] + MC$ ∇ .

The thermal conductivity of ionic liquids was not found to influence the thermal conductivity enhancements as those are very similar for different anions (as shown above in the ranges of enhancements). However, a significant impact can be observed for different types and size of nanoparticles. In this work, 4 different types of nanoparticles were used with a wide range of thermal conductivity (from a very high for carbon nanotubes to relatively low for mesoporous carbon). Carbon nanotubes were found to increase the thermal conductivity the most (as corresponding to their highest thermal conductivity, 3223.4 W m⁻¹ K⁻¹), followed by boron nitride (874.1 W m⁻¹ K⁻¹), graphite (35.7 W m⁻¹ K⁻¹) and mesoporous carbon (~12.0 W m⁻¹ K⁻¹), as expected. Similar results were found for other systems, for example 3.71 % for 1-butyl-1-methylpyrrolidinium dicyanamide + 0.5 wt % MWCNT at 293.1 K (0.84–4.58 % in this work) [38], 4.8 % for 1-hexyl-3-methylimidazolium tetrafluoroborate + 1 wt % MWCNT at 293 K (3.18–7.45 % in this work) [32], or 20.46 % for 1-ethyl-3-methylimidazolium ethylsulfate + 3 wt % MWCNT at 293.77 K (17.98–22.62 % in this work) [35].

To date, only one model was developed to predict the thermal conductivity of ionanofluids (more specifically their enhancement) so-called the Atashrouz model [40]. Oster et al. investigated this model for ionanofluids with several types of ionic liquids [41]. This model is expected to reproduce the thermal conductivity with high accuracy because it is fully based on thermodynamics (modified geometry mean), and the interactions between ionic liquids and nanoparticles are also included in its development [40]. During this work, this approach has been further tested for selected ionanofluids. The results of calculations (experimental enhancement vs. calculated enhancement) are presented in Fig. 10a and collected in Supporting Information SI2.

The average standard uncertainty of the determined enhancements is 4.5 %. All values are well within the linear relation (solid line in Fig. 10a) between the experimental and calculated values of the enhancement. More specifically, in Fig. 10b, the relative deviations are presented. None of the calculated values exceed the standard uncertainty of determined enhancement (all of them are below 4 %). The average absolute relative deviation was found to be (0.66, 1.8, 1.2 and 0.54) % for MWCNT, G, BN and MC, respectively, while the maximum values of relative deviation were -(3.82, 3.61, 3.65 and 2.82) % for $[P_{14,6,6,6}][ButO] + 3$ wt % MWCNT at 278.15 K, $[P_{14,6,6,6}][ButO] + 1$ wt % G at 338.15 K, $[P_{14,6,6,6}][HexO] + 3$ wt % BN at 328.15 K and $[P_{14,6,6,6}][ButO] + 3$ wt % MC at 278.15 K, respectively. Overall, one can appreciate that the Atashrouz model can be used to predict the thermal conductivity of ionanofluids accurately [40].

Isobaric heat capacity

Isobaric heat capacity was measured over the temperature range (298.15–363.15) K. The results for pure ionic liquids are presented in Fig. 8b and reported in Supporting Information SI2. The parameters of linear

equation $c_n(T)$ are collected in Table S2 (Supporting Information SI3). In comparison to commercial heat transfer fluids, the values of heat capacity presented in this work are similar to the commercial HTFs, for example (1.93, 1.90 or 1.63) kJ kg⁻¹ K⁻¹ Therminol ADX10, Dynalene LO-170, Dowtherm Q at 298.15 K, respectively, while the values in this work are (2.02, 1.95, 1.91, 1.84 and 1.82) kJ kg $^{-1}$ K $^{-1}$ for [$P_{14,666}$][AcO], [$P_{14,666}$] [ButO], $[P_{14,6,6,6}]$ [HexO], $[P_{14,6,6,6}]$ [OctO] and $[P_{14,6,6,6}]$ [DecO] at 298.15 K, respectively [70–72].

The heat capacity is found to increase with temperature, as expected. The $c_n(T)$ changes from 298.15 K to 363.15 K are (6.87, 7.49, 8.18, 8.39 and 8.80) % for $[P_{14,6,6,6}][AcO]$, $[P_{14,6,6,6}][ButO]$, $[P_{14,6,6,6}][HexO]$, $[P_{14,6,6,6}][HexO]$, $[P_{14,6,6,6}][HexO]$ and $[P_{14,666}][DecO]$, respectively. In other words, the temperature effect on the Δc_n is increasing with the anion chain length. Heat capacity acts as a representation of the bonds energy in the molecules, therefore, an increase of the anion chain length should result in a larger effect of the temperature on their c_{x} values.

The isobaric heat capacity was also studied for mixtures of the ionic liquids with carbon nanotubes, boron nitride, graphite and mesoporous carbon up to 3 wt % as shown in Fig. S22-S26 (Supporting Information SI1) with data tabulated in Supporting Information SI2. The more insight discussion can be made based on the enhancements of ionanofluids heat capacity in comparison to equivalent ionic liquids heat capacity. In accordance to traditional mixing rules in which the resulting heat capacity is made by the contribution of mixed materials (solids – lower heat capacity, liquids – higher heat capacity), the heat capacity

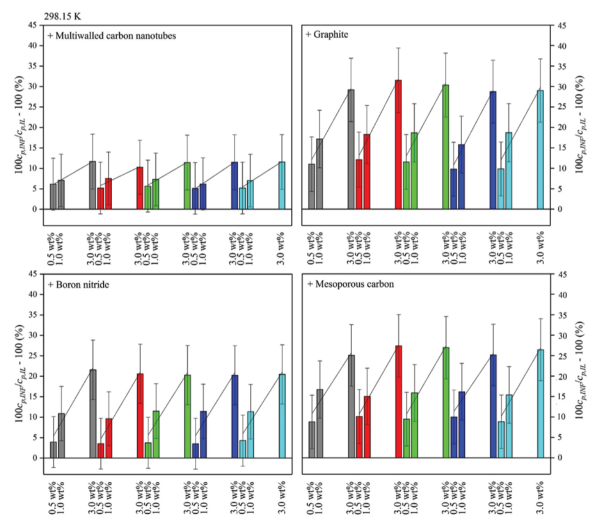


Fig. 11: Isobaric heat capacity enhancement of ionanofluids, $(100c_{p,NH}/c_{p,ll}-100)$, in comparison to pure ionic liquids, over the mass fraction, for grey – $[P_{14,6,6,6}][AcO]$ INF, red – $[P_{14,6,6,6}][ButO]$ INF, green – $[P_{14,6,6,6}][HexO]$ INF, blue – $[P_{14,6,6,6}][OctO]$ INF, cyan – [P_{14,6,6,6}][DecO] INF, at 298.15 K.

of ionanofluids should be lower than that of pure ILs. In reality, for ionic liquid – based nanofluids, the heat capacity increases with the addition of nanoparticle in solution. This behaviour could be associated to the interfacial nanolayers created by adhered liquid molecules on the surface of nanoparticles [42]. The results of these enhancements are presented in Fig. 11 and Table 2. In all cases, a linear correlation could be used again to accurately represent the qualitative dependence of the nanoparticle concentration on the \boldsymbol{c}_p enhancement for each mixture investigated herein.

The maximum c_p enhancements observed were (10.26–11.67, 28.71–31.50, 20.26–21.58 and 25.07–27.40) % for MWCNT, G, BN and MC at 298.15 K, respectively, and (7.00–10.00, 32.67–36.10, 20.61–21.49 and 27.54–30.90) % for MWCNT, G, BN and MC at 363.15 K, respectively. As reported previously [42], the enhancements as a function of temperature depend on the properties of nanoparticles, more specifically, their size. It can be seen that for carbon nanotubes the enhancement decreases with the temperature, which might be related to an increase of the hydrophobicity i.e. less favoured attraction of the liquid molecules [90]. In the case of graphite, the c_p enhancement increases with the temperature due to a hydrophobicity decrease which leads to easier creation of nanolayers [91], whilst with boron nitride it is approximately constant with temperature due to the diversely charged surface of charge distribution between nitrogen and boron [92]. Mesoporous carbon has a relatively high hydrophobicity at room temperature as found with graphite, therefore, it has a similar behaviour to that graphite [93].

However, herein, no dependence of the anion chain length on the c_p enhancement was found. Nevertheless, the type of nanoparticles, or more specifically their size, seems to be an influencing factor to the c_p enhancement. The following dependence of enhancements, MWCNT < BN < MC < G, was observed. In a previous work, it was shown that increasing the size of nanoparticles resulted in increasing the enhancement caused by larger number of liquid molecules adhered on the surface of nanoparticles (investigated for MWCNT, BN and G), which is in a good agreement with results presented in this work [42]. However, in case of mesoporous carbon, the enhancement is higher than for boron nitride but lower than graphite. This change may be the result of the very high porosity of mesoporous carbon (as in the Experimental Section, average pore size 20–40 nm) which influences the adsorption of liquid molecules and, therefore, the enhancement.

The heat capacity of investigated ionanofluids was also calculated by using fully empirical correlations from work of Oster et al. [42]. Unfortunately, the only available models are fully empirical based on data of ionanofluids with carbon nanotubes, boron nitride and graphite. Therefore, only those data were further simulated in this work. The representation of experimental data versus those calculated can be found in Fig. 12a. In addition, the differences can be found in Fig. 12b. The average absolute relative deviations are (0.47, 0.96).

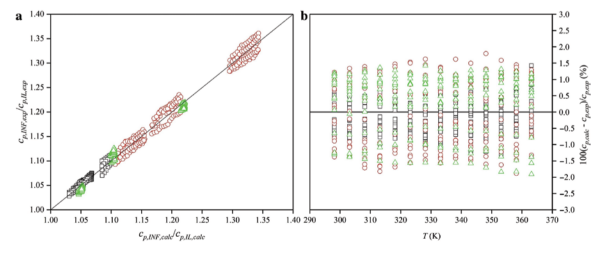


Fig. 12: (a) experimental vs. calculated isobaric heat capacity of ionanofluids; (b) relative deviations between calculated and experimental values of ionanofluids isobaric heat capacity against the temperature, for $[P_{14,6,6,6}][RO] + MWCNT$, \Box ; $[P_{14,6,6,6}][RO] + G$, \bigcirc ; $[P_{14,6,6,6}][RO] + BN <math>\triangle$.

and 0.87) % for ionanofluids with MWCNT, G and BN, respectively, while all the relative deviations were well below 2%. It should be noted that the calculated c_n enhancements are below the heat capacity uncertainty (3%). Thus, it is possible to calculate the heat capacity of ionanofluids with carbon nanotubes, boron nitride and graphite accurately.

Volumetric heat capacity

The most important property in the case heat transfer fluids is the ability to store the energy (as heat). This can be represented with the volumetric heat capacity, c_v , which is calculated based on isobaric specific heat capacity, c_n , and specific density, ρ , $c_v = c_n \cdot \rho$ [19]. One can describe it as a viability to store the heat in a volume of a material, as a change in temperature. Higher values of volumetric heat capacity result in increased ability to store the heat, therefore, these are favoured. As expected, high isobaric heat capacity and high density contribute to the high volumetric heat capacity.

The ionic liquids investigated in this work have a relatively low density (below 1000 kg m⁻³), however, they have relatively high specific heat capacity (1.82–2.91 kJ kg⁻¹ K⁻¹). The volumetric heat capacity at 298.15 K and 363.15 K are shown in Fig. 13 and Table 3. The volumetric heat capacity found in this work was in the range of $(1.60-2.39 \text{ and } 1.66-2.54) \text{ MJ K}^{-1} \text{ m}^{-3} \text{ at } 298.15 \text{ K and } 363.15 \text{ K, respectively.}$ The maximum was found as 2.39 MJ K^{-1} m⁻³ for $[P_{14,6,6,6}][AcO] + 3.0$ wt % G at 298.15 K and 2.54 MJ K^{-1} m⁻³ for $[P_{14,6,6,6}][AcO] + 3.0$ wt % G at 363.15 K, respectively. While the minimum was as 1.60 MJ K^{-1} m⁻³ for $[P_{14.666}][DecO]$ at 298.15 K and 1.66 MJ K^{-1} m⁻³ for $[P_{14666}]$ [DecO] at 363.15 K, respectively. For the pure ILs, the c_v is increasing with the anion chain length, also driven by the isobaric heat capacity.

The temperature effect was found to be similar for all systems, with an approximate 4% increase in volumetric heat capacity from 298.15 K to 363.15 K (driven by the increase of isobaric heat capacity). As can

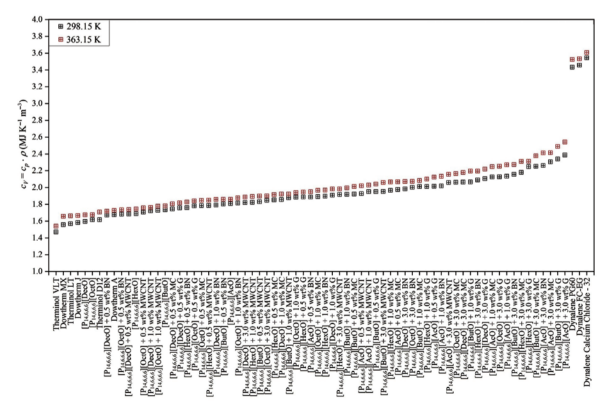


Fig. 13: Volumetric heat capacity, c,, for all investigated systems and some commercially available heat transfer [70-72], for the comparison, at 298.15 K and 363.15 K.

Table 3: A comparison of maximum temperature of operation and volumetric heat capacity, c_{ij} between commercially available heat transfer fluids and fluids in this work.

	Maximum temperature of operation (K)	c, at 298.15/363.15 K (MJ K ⁻¹ m ⁻³)
Dynalene PG-60 [71]	450	3.43/3.53
Dynalene FC-PG [71]	353	3.46/3.52
Dynalene HC-50 [71]	383	3.62/3.70
Dynalene BioGlycol-60 [71]	450	3.54/3.60
Dynalene Calcium Chloride-32 [71]	373	3.55/3.61
Dowtherm SR-1 [72]	393	3.54/3.58
Dowtherm A [72]	673	1.68/1.78
Dowtherm J [72]	588	1.58/1.67
Dowtherm MX [72]	603	1.56/1.68
Therminol VLT [70]	448	1.47/1.54
Therminol LT [70]	453	1.57/1.66
Systems studied in this work	~400	1.60-2.54

be seen the values of volumetric heat capacity of systems in this work are relatively high. The addition of nanoparticles increases the c_v , for example the minimum c_v enhancement at 298.15 K was found as 3.16 % for $[P_{14666}][OctO] + 0.5 \text{ wt} \% \text{ MWCNT, and maximum as } 39.78 \% \text{ for } [P_{14666}][ButO] + 3.0 \text{ wt} \% \text{ G. Generally, the } c_V$ enhancement was observed to be strictly dependent on the isobaric heat capacity. Therefore, the enhancements were found in the following sequence: MWCNT < BN < MC < G.

As a comparison, some commercially available heat transfer fluids were selected. There are several heat transfer fluids whose volumetric heat capacity is lower that the ILs studied in this work. For example, 1.47 MJ K⁻¹ m⁻³ at 298.15 K and 1.54 MJ K⁻¹ m⁻³ at 363.15 for Therminol VLT; values close to 1.56 MJ K⁻¹ m⁻³ at 298.15 K and to 1.68 MJ K^{-1} m⁻³ at 363.15 for Dowtherm MX, and values close to 1.57 MJ K^{-1} m⁻³ at 298.15 K and 1.66 MJ K^{-1} m⁻³ at 363.15 for Therminol LT, 1.58 MJ K⁻¹ m⁻³ at 298.15 K and 1.67 MJ K⁻¹ m⁻³ at 363.15 for Dowtherm J, 1.62 MJ K^{-1} m⁻³ at 298.15 K and 1.71 MJ K^{-1} m⁻³ at 363.15 for Therminol D12, or 1.68 MJ MJ K^{-1} m⁻³ at 298.15 K and 1.78 MJ K^{-1} m⁻³ at 363.15 for Dowtherm A [70, 72]. These heat transfer fluids are mainly based on organic compounds (synthetics, aromatics, silicones and carbohydrates). Obviously, there are some heat transfer fluids that have higher values of volumetric heat capacity, for example 3.43 MJ K⁻¹ m⁻³ at 298.15 K and 3.53 MJ K⁻¹ m⁻³ at 363.15 for Dynalene PG60, 3.46 MJ K^{-1} m⁻³ at 298.15 K and 3.52 MJ K^{-1} m⁻³ at 363.15 for Dynalene FC-EG, 3.55 MJ K^{-1} m⁻³ at 298.15 K and 3.61 MJ K⁻¹ m⁻³ at 363.15 for Dynalene CaCl2-32 [71]. The major impact in their high values is originated from water. Even though water has moderate density (or in comparison to other liquid quite low), the isobaric heat capacity of this material is very high (4.18 kJ kg⁻¹ K⁻¹ at 298.15 K), and all of those heat transfer fluids are based on water.

Thermogravimetric analysis - thermal stability

The thermogravimetric analysis (TGA) was used to investigate the thermal decomposition profile of the pure ILs and mixtures with carbon nanotubes, boron nitride, graphite and mesoporous carbon. The results of thermogravimetric curves can be found in Fig. S28-S87 (Supporting Information SI1). Those figures also include the derivative thermogravimetric curve (dm/dT vs. T). The determined onset temperatures were collected in Table S3 (Supporting Information SI3), and these were also presented against the nanoparticles weight concentration in Fig. 14.

It can be seen (Fig. 14) that the determined T_{on} increases with increasing the number of nanoparticles which is understandable as these nanomaterials have higher thermal stability than the investigated ILs, as can be shown in Fig. S27 (Supporting Information SI1). The nanoparticles were significantly more stable than the investigated ILs (by estimation $T_{on} > 750$ K). Moreover, the T_{on} exhibits an approximately linear dependence with the nanoparticles concentration. The enhancement range of onset temperatures caused by the addition of nanoparticles was (2.02–5.74, 2.71–7.15, 2.49–6.40, 2.38–5.23) % for boron nitride-, carbon

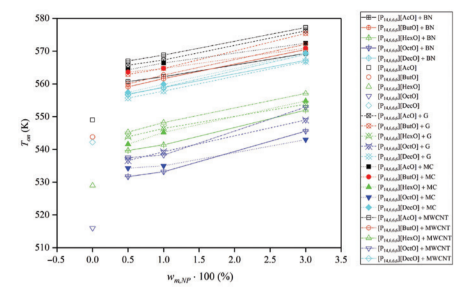


Fig. 14: The onset temperature, T_{on} , determined from thermogravimetric analysis for all investigated ionic liquids and ionanofluids.

nanotubes-, graphite- and mesoporous carbon-doped ionic liquids, respectively, or (2.13-5.14, 2.81-5.81, $2.02-5.31, 3.04-7.15, 2.49-5.02) \% \ for \ [P_{14,6,6,6}][AcO]-, \ [P_{14,6,6,6}][ButO]-, \ [P_{14,6,6,6}][HexO]-, \ [P_{14,6,6,6}][OctO]- \ and \ [P_{14,6,6,6}][P_{14,6,6}][P_{14,6,6,6}][P_{14,6,6}][P_{14,6,6,$ [DecO]-based ionanofluids, respectively. As can be seen, all shifts of T_{on} are similar (nanoparticle- and ionic liquid-dependence) which means that the driving force for the enhancement is purely caused by the addition of more stable species (nanoparticles) into the ionic liquids.

To the best of our knowledge, there are no published data which take into consideration the thermal stability of nanofluids with different type basefluids or nanoparticles, particularly when ionic liquids are used as basefluids. Therefore, the analysis and comparison to previously reported results become more difficult. The linear-like dependence of T_{on} on the nanoparticles concentration shows that the thermal stability is rather dependent on the addition/type of nanoparticles instead of some more sophisticated mechanism. The specific structure of nanofluids, particularly the surface of nanoparticles (i.e. solid-liquid nanolayering) is the cause for unusual behaviour of thermophysical properties of such systems. In case of thermal stability this was not found to be an influencing factor, probably due to not sufficient sensitivity of such measurements.

The thermal decomposition was also found to depend on the anion chain length in the following sequence: [AcO] > [ButO] > [DecO] > [HexO] > [OctO]. $[P_{14,6,6,6}][DecO]$ exhibits a discrepancy from the linear relationship. The possible cause of this behaviour is that decanoic acid is a solid at room temperature, therefore, it influences the molecular structure of [P_{14,6,6,6}][DecO], inducing different molecular recognition than in comparison to other ionic liquids with carboxylate anion. Moreover, similar sequences were found for ionanofluids based on these ionic liquids.

A number of conclusions can be drawn: 1) the onset temperature for all samples is dependent on type of ionic liquid and only the amount of added nanoparticles, 2) the onset temperature is a point at which advanced decomposition can be observed (even up to 20 % of weight change, for example [P14.6.6.6][AcO] + 1.0 wt % G $[P_{_{14,6,6,6}}][OctO] + 0.5 \text{ wt } \% \text{ MWCNT or } [P_{_{14,6,6,6}}][DecO] + 1.0 \text{ wt } \% \text{ MC, thus, the onset temperature cannot be constant.}$ sidered as the descriptor of sample thermal stability, 3) the common feature of all samples is that the decomposition begins to be detectable at about 400 K. The one with the lowest onset temperature was selected, $[P_{1466}][DecO]$ and isothermal TGA measurements were then conducted at 400 K, 450 K, 500 K and 550 K, as shown in Fig. 15.

The weight losses observed are (0.1, 2.9, 6.4 and 11.8) % at 400 K, 450 K, 500 K and 550 K, respectively. It can be seen in Fig. 15 that the $[P_{14,6,6,6}][DecO]$ is stable at 400 K, and the weight loss is probably caused by some water removal (absorbed from the air during sample preparation and transportation). Further increase of the temperature leads to more extensive weight loss due to its decomposition (or evaporation).

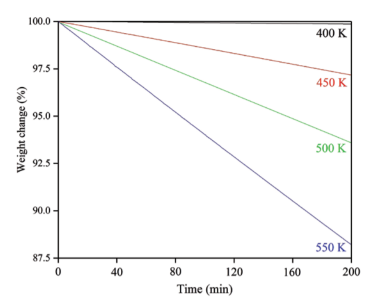


Fig. 15: Isothermal thermogravimetric results as decomposition weight loss over the time at different temperatures for $[P_{14,6,6,6}]$ [DecO].

In terms of practical application of these systems, a consideration as heat transfer fluids is a viable option, as can be seen in Table 3. There are several different types of heat transfer fluids – based on glycols (i.e. Dynalene PG/EG series, Dowcal 100), alkanes (i.e. Therminol VLT), aromatic compounds (i.e. Therminol LT), water (i.e. Dynalene Calcium Chloride series) [70–72]. Some examples of commercially available heat transfer fluids with lower thermal stability can be found (as well as those with higher thermal stability) indicating that the investigated ionic liquids could be successfully considered as replacements. Even though the addition of nanoparticles increases the onset temperature, the maximum temperature of operation is still limited by the thermal stability of ionic liquids.

Ionanofluids economic viability

Physical properties are very important to ascertain whether some material can be used as heat transfer fluid. The most important are thermal conductivity, isobaric heat capacity, density and viscosity. Further investigation of these properties can lead to estimation of the heat exchange reactor.

The comparison of ionic liquids thermal conductivity depends on the composition of commercial fluid, for example water-based materials have higher thermal conductivity (i.e. Dynalene PG series, Dowtherm 4000 or Dynalene EG series), while those based on organic compounds have lower thermal conductivity (i.e. Therminol ADX10 or Dynalene SF) [70–72]. For the density, the comparison also depends on the type of material, for example the values in this work are similar to synthetic aromatic hydrocarbon mixtures (i.e. Therminol ADX10 or Dynalene SF), slightly lower than water, or significantly lower than glycol-based HTFs (for example Dowtherm 4000 or Dynalene EG series) [70–72]. Unfortunately, the viscosity is significantly higher than commercial heat transfer fluids, which is a major drawback.

Based on that qualitative analysis, the price of heat exchange unit for materials in this work should be higher than commercial heat transfer fluids (due to high viscosity). The results can be seen in Fig. 16 and quantitative values in Table 4.

Firstly, the prices for all pure ionic liquids are higher than those of selected heat transfer fluids, for example (1130.7, 1245.8, 1385.7, 1460.9 and 1482.4) k\$ for $[P_{14,6,6,6}][AcO]$, $[P_{14,6,6,6}][ButO]$, $[P_{14,6,6,6}][DecO]$ and $[P_{14,6,6,6}][DecO]$ at 298.15 K, and (542.2, 584.1, 643.2, 668.4 and 696.1) k\$ for $[P_{14,6,6,6}][AcO]$, $[P_{14,6,6,6}][ButO]$, $[P_{14,6,6,6}][HexO]$, $[P_{14,6,6,6}][DecO]$ at 363.15 K, respectively. While the

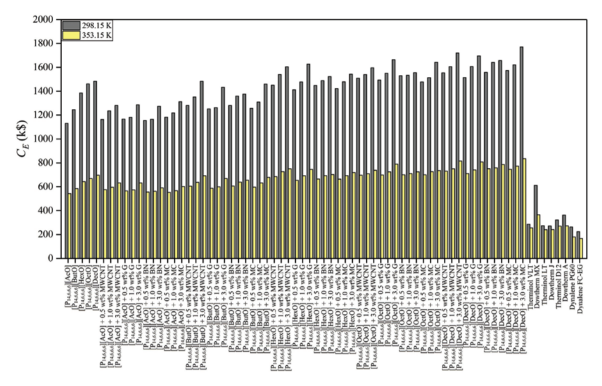


Fig. 16: The results of economic analysis in the meaning of the heat exchange reactor, C_{c1} at 298.15 K and 363.15 K [70–72].

values for selected HTFs are for example (285.4, 612.1 or 362.8) k\$ for Therminol VLT, Dowtherm MX or Dowtherm A at 298.15 K, and (255.6, 365.3 or 272.6) k\$ for Therminol VLT, Dowtherm MX or Dowtherm A at 363.15 K, respectively. The ILs are more expensive by up to 5 times which is also caused by their higher viscosity [70, 72].

On the other hand, the price of the pure ionic liquids decreases with temperature, and over 100 K this drops by over 100 %. The decrease for commercial heat transfer fluids is not that significant as they already have quite low viscosity, and as discussed before, the viscosity in this work is decreasing for about 1000 % within the investigated temperature range. At higher temperatures, the price for ionic liquids and commercial heat transfer fluids would be very similar. Moreover, for industrial purposes the fluids are usually used at high temperature which is more applicable for ionic liquids.

Despite the decreasing viscosity, other properties also contribute to the decrease of the price, for instance the isobaric heat capacity (7–9 % of increase within 100 K temperature change). Unfortunately, thermal conductivity and density counteract the beneficial changes in viscosity and isobaric heat capacity. For exampleapproximately 5% and 7% decreases in the density and thermal conductivity over a 100 K temperature change are observed, respectively.

Despite the high cost of the ILs, they also have other important properties, namely low vapour pressure [94], low freezing point [95], high thermal stability [54], or wide liquid range [96]. Therefore, the application of them as HTFs is beneficial as they can help to overcome many limitations associated with the engineering issues of the heat exchange, for example pressure control.

Furthermore, the addition of nanoparticles causes an increase in the price. It was observed that the enhancement was independent on the type of ionic liquid or nanoparticle, average as (3.1, 6.8 and 13.7) % for ionanofluids containing 0.5 wt %, 1.0 wt % and 3.0 wt % nanoparticles, respectively. This is a very significant shift, for example 13.7 % is at least 155 000 \$. These shifts are mainly caused be the increase in viscosity (of even up to 150%), even though the other properties are improved, for example density (up to ~3%), isobaric heat capacity (up to ~35%) and thermal conductivity (up to ~20%). Unfortunately, as can be seen, the viscosity is still a remaining problem and it influences the thermal performance of ionanofluids, even though other

Table 4: The results of economic analysis in the meaning of the heat exchange reactor, C_E , at 298.15 K and 363.15 K, along with the temperature effect, Eq. (19).

	C _E at 298.15 K (k\$)	C _E at 363.15 K (k\$)	Temperature effect (%)
[P _{14,6,6,6}][AcO] Neat	1130.7	542.2	108.54
$[P_{14,6,6,6}][AcO] + 0.5 \text{ wt } \% \text{ MWCNT}$	1163.0	575.0	102.26
[P _{14,6,6,6}][AcO] + 1.0 wt % MWCNT	1234.5	596.3	107.03
[P _{14,6,6,6}][AcO] + 3.0 wt % MWCNT	1280.6	632.1	102.59
[P _{14,6,6,6}][AcO] + 0.5 wt % G	1166.0	565.2	106.30
$[P_{14,6,6,6}]$ [AcO] + 1.0 wt % G	1180.4	573.0	106.00
$[P_{14,6,6,6}][AcO] + 3.0 \text{ wt } \% \text{ G}$	1286.3	632.2	103.46
$[P_{14,6,6,6}][AcO] + 0.5 \text{ wt } \% \text{ BN}$	1155.2	555.1	108.11
$[P_{14,6,6,6}][AcO] + 1.0 \text{ wt } \% \text{ BN}$	1164.9	562.4	107.13
$[P_{14,6,6,6}]$ [AcO] + 3.0 wt % BN	1273.0	591.2	115.32
$[P_{1/(6.6.6)}]$ [AcO] + 0.5 wt % MC	1181.5	552.6	113.81
[P _{14,6,6,6}][AcO] + 1.0 wt % MC	1217.7	566.2	115.07
$[P_{14,6,6,6}][AcO] + 3.0 \text{ wt } \% \text{ MC}$	1311.9	601.4	118.14
[P _{14,6,6,6}][ButO] Neat	1245.8	584.1	113.29
[P _{14,6,6,6}][ButO] + 0.5 wt % MWCNT	1280.6	604.6	111.81
[P _{14,6,6,6}][ButO] + 1.0 wt % MWCNT	1351.3	636.7	112.23
[P _{14,6,6,6}][ButO] + 3.0 wt % MWCNT	1482.4	691.9	114.25
[P _{14,6,6,6}][ButO] + 0.5 wt % G	1251.5	587.0	113.20
[P _{14,6,6,6}][ButO] + 1.0 wt % G	1262.7	599.8	110.52
[P _{14,6,6,6}][ButO] + 3.0 wt % G	1432.8	668.5	114.33
[P _{14,6,6,6}][ButO] + 0.5 wt % BN	1279.9	606.1	111.17
[P _{14,6,6,6}][ButO] + 1.0 wt % BN	1359.9	638.3	113.05
[P _{14,6,6,6}][ButO] + 3.0 wt % BN	1375.2	654.9	109.99
[P _{14,6,6,6}][ButO] + 0.5 wt % MC	1257.0	596.5	110.73
[P _{14,6,6,6}][ButO] + 1.0 wt % MC	1308.7	631.5	107.24
[P _{14,6,6,6}][ButO] + 3.0 wt % MC	1459.9	678.1	115.29
[P _{14,6,6,6}][HexO] Neat	1385.7	643.2	115.44
[P _{14,6,6,6}][HexO]+0.5 wt % MWCNT	1450.3	685.5	111.57
[P _{14,6,6,6}][HexO] + 1.0 wt % MWCNT	1540.4	726.0	112.18
[P _{14,6,6,6}][HexO]+3.0 wt % MWCNT	1604.2	750.0	113.89
[P _{14,6,6,6}][HexO] + 0.5 wt % G	1411.3	654.0	115.80
[P _{14,6,6,6}][HexO] + 1.0 wt % G	1477.4	692.3	113.40
[P _{14,6,6,6}][HexO]+3.0 wt % G	1626.2	744.9	118.31
[P _{14,6,6,6}][HexO] + 0.5 wt % BN	1447.9	664.9	117.76
[P _{14,6,6,6}][HexO] + 1.0 wt % BN	1488.9	692.9	114.88
[P _{14,6,6,6}][HexO] + 3.0 wt % BN	1521.8	701.8	116.84
[P _{14,6,6,6}][HexO]+0.5 wt % MC	1422.0	663.9	114.19
[P _{14,6,6,6}][HexO] + 1.0 wt % MC	1477.7	693.0	113.23
[P _{14,6,6,6}][HexO] + 3.0 wt % MC	1542.5	719.3	114.44
[P _{14,6,6,6}][OctO] Neat	1460.9	668.4	118.57
[P _{14,6,6,6}][OctO]+0.5 wt % MWCNT	1508.8	696.0	116.78
[P _{14,6,6,6}][OctO] + 1.0 wt % MWCNT	1539.1	708.7	117.17
[P _{14,6,6,6}][OctO] + 3.0 wt % MWCNT	1596.0	737.0	116.55
[P _{14,6,6,6}][OctO]+0.5 wt % G	1492.5	699.1	113.49
[P _{14,6,6,6}][OctO] + 1.0 wt % G	1549.6	724.9	113.77
[P _{14,6,6,6}][OctO]+3.0 wt % G	1663.3	789.4	110.70
[P _{14,6,6,6}][OctO]+0.5 wt % BN	1529.5	700.0	118.50
[P _{14,6,6,6}][OctO] + 1.0 wt % BN	1533.0	710.4	115.79
[P _{14,6,6,6}][OctO] + 3.0 wt % BN	1555.6	724.9	114.60
[P _{14,6,6,6}][OctO]+0.5 wt % MC	1476.7	700.4	110.84
[P _{14,6,6,6}][OctO] + 1.0 wt % MC	1513.8	725.4	108.68
[P _{14,6,6,6}][OctO] + 3.0 wt % MC	1641.8	734.9	123.40
[P _{14,6,6,6}][DecO] Neat	1482.4	696.1	112.96
[P _{14,6,6,6}][DecO] + 0.5 wt % MWCNT	1553.9	730.2	112.80
[P _{14,6,6,6}][DecO]+1.0 wt % MWCNT	1604.5	749.8	113.99
14,6,6,611DCCO] 1 1.0 WE 70 WINDCOM	1004.5	742.0	113.99

Table 4 (continued)

	C _E at 298.15 K (k\$)	C _E at 363.15 K (k\$)	Temperature effect (%)
[P _{14.6.6.6}][DecO]+3.0 wt % MWCNT	1719.0	815.5	110.79
[P _{14,6,6,6}][DecO] + 0.5 wt % G	1514.0	709.7	113.33
[P _{14,6,6,6}][DecO] + 1.0 wt % G	1605.9	740.4	116.90
[P _{14,6,6,6}][DecO] + 3.0 wt % G	1695.6	807.0	110.11
[P _{14,6,6,6}][DecO]+0.5 wt % BN	1556.8	751.1	107.27
[P _{14,6,6,6}][DecO] + 1.0 wt % BN	1641.4	757.1	116.80
[P _{14,6,6,6}][DecO] + 3.0 wt % BN	1657.5	786.3	110.80
[P _{14,6,6,6}][DecO] + 0.5 wt % MC	1572.7	746.0	110.82
[P _{14,6,6,6}][DecO] + 1.0 wt % MC	1620.4	772.8	109.68
[P _{14,6,6,6}][DecO] + 3.0 wt % MC	1769.6	834.8	111.98
Therminol VLT [70]	285.4	255.6	11.66
Dowtherm MX [72]	612.1	365.3	67.56
Therminol LT [70]	273.0	241.0	13.28
Dowtherm J [72]	271.1	241.7	12.16
Therminol D12 [70]	322.1	270.4	19.12
Dowtherm A [72]	362.8	272.6	33.09
Dynalene PG60 [71]	263.7	179.3	47.07
Dynalene FC-EG [71]	224.2	166.6	34.57

properties are improved. On the other hand, the temperature effect on the price (and physical properties, particularly viscosity) is similar to that of the pure ionic liquids.

Unfortunately, based on the conducted economic studies, ionic liquid – based nanofluids do not seem to be promising materials for heat transfer applications. Despite the enhanced thermal conductivity, density and heat capacity, which are favourable, the viscosity is also enhanced which significantly increases the price of the unit. In contrast, the pure ionic liquids are promising materials for heat transfer applications especially at high temperature due to the decrease in viscosity. Additionally, the properties of ionic liquids (low vapour pressure or wide liquid range) makes them even more attractive solution. Oster et al. discussed ionic liquids in mixture with water in terms of application as heat transfer fluids [21]. These mixtures seem to be the most promising as: 1) heat capacity and thermal conductivity are enhanced to very high values and 2) the viscosity is significantly decreased. However, some of the desirable properties of the ionic liquids are lost in this case, for example low vapour pressure. However, reliable data over the entire concentration range are thus a necessary prerequisite for the use of such mixtures. Furthermore, there may be a negative effect in terms of excess properties especially in the water rich region, which would have a negative impact on thermal properties [97, 98].

Conclusions

Comprehensive studies on ionanofluids composed of trihexyl(tetradecyl)phosphonium acetate, butanoate, hexanoate, octanoate and decanoate ionic liquids with multiwalled carbon nanotubes, boron nitride, graphite and mesoporous carbon nanoparticles up to 3 wt % of concentration, has been reported. The experimental data of density, dynamic viscosity, thermal conductivity, isobaric heat capacity, short-range and long-range thermogravimetric analysis were presented, along with calculated kinematic viscosity, volumetric heat capacity and estimated heat exchange unit cost governing studied systems. The prediction of density, thermal conductivity, isobaric heat capacity was also conducted. The discussion presented in this work finalized the impact of ionic liquids and ionic liquid - based nanofluids properties into a heat transfer fluids design where the ionanofluids do not represent better thermophysical properties than the pure ionic liquids. Most importantly, this is the first work of this kind in which ionanofluids are considered for industrial purposes from both molecular and engineering perspective, including a comprehensive analysis and comparison to commercial heat transfer fluids.

As expected, the limiting factor in terms of heat transfer fluid design in the case of ionic liquids is their viscosity whereas many of the other physical properties are positive. The addition of nanoparticles into the ionic liquids results in the beneficial enhancements of heat capacity, thermal conductivity and density, however, such an addition causes a drastic increase of their viscosity, which negatively overrides the other enhancements. Temperature was found to affect the enhancement, as well as type of ionic liquid and nanoparticles/nanoparticles concentration.

It should be noted that discussion on whether a material is good or not is strongly dependent on the specific application for the heat transfer fluid. The recommendation from this study is to use ionic liquids at moderately high temperatures (up to 400 K) in which pressure control is problematic and can be overcome by the very low vapour pressure of the ionic liquids.

The modelling of physical properties of materials considered as heat transfer fluids might result in a significant reduction in time and cost of ionanofluids design. In this work, the presented calculations and the comparison to experimental data (density and thermal conductivity) show that the presented models are capable to predict those physical properties. In terms of heat capacity, empirical equations designed for specific nanomaterials were used successfully, however, this shows that more versatile models are still needed to allow the design of other types of nanomaterials.

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