Ionic liquid pre-treatment of microalgae and extraction of biomolecules

Rupali K. Desai
Thesis committee

Promotors
Prof. Dr M.H.M. Eppink
Special Professor Biorefinery with Focus on Mild Separation Technologies of Complex Biomolecules
Wageningen University

Prof. Dr R.H. Wijffels
Professor of Bioprocess Engineering
Wageningen University

Other members
Prof. Dr A.B. de Haan, Delft University of Technology
Prof. Dr W.J.H. van Berkel, Wageningen University
Dr M.G. Freire, University of Aveiro, Portugal
Dr K. Goiris, KU Leuven, Belgium

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Dedicated to my family
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Chapter 1

General introduction and thesis outline
Chapter 1

Introduction

In the bioprocess and chemical industry, many separation technologies are already highly developed and more than one technology option is available for most processes. However, these separation technologies could be improved in terms of energy efficiency, raw materials used, or cost effectiveness. In addition, changing technologies and changing customer demands continually create new needs for the chemical and biochemical industry. Liquid–liquid extractions (LLE) is an interesting purification alternative and is widely used in the chemical industry due to its simplicity, and ease of scale up [1]. Liquid–liquid extraction is the transfer of certain components from one phase to another when immiscible or partially soluble liquid phases are brought into contact with each other. The conventional process employed in this field consist of water–organic solvent two-phase systems and, therefore, are generally not suitable for separation of biomolecules (e.g. proteins, carbohydrates) due to problems such as protein denaturation [2]. Moreover, with increasing safety concerns for personnel and consumers, extraction by organic solvents is a challenge due to volatility, flammability and toxicity of the solvents [3]. In this respect, complex fluids have been studied for liquid–liquid extraction of biomolecules by means of aqueous two-phase systems (ATPS) that provide a bio-compatible environment. ATPS are formed by mixing polymers (e.g. Polyethylene Glycol (PEG), Polypropylene Glycol (PPG)) with inorganic salt (e.g. phosphate, citrate) or another polymer beyond a certain critical concentration (see Figure 1.1). These systems are used for separation, concentration and fractionation of fragile biomolecules [4]. Polymer-polymer based systems usually display two hydrophobic phases and the difference in polarities depends essentially on the amount of water in each phase. On the other hand, polymer-salt Aqueous Biphasic System (ABS) has a hydrophobic phase constituted by the polymer and a hydrophilic (and more ionic) phase, typically formed by high charge-density salts. The restricted difference in polarities between the two phases prevents vast use of polymer-based ABS for extraction purposes. By virtue of their tunability, ionic liquids (IL) (see below) can ideally cover the whole hydrophilicity–hydrophobicity range [5]. This necessitates the need to develop separation processes with alternative solvents such as IL, supercritical fluids. Microemulsions, such as reverse micelles which are also a two phase system have also been studied for separation. This system suffers from poor back extraction efficiency and denaturation of protein [6]. Therefore the goal of this thesis is to evaluate the feasibility of using IL based system for extraction of value added components, both hydrophilic and hydrophobic components from complex biomass such as microalgae.
Ionic liquids

Ionic liquids are molten salts at a temperature below 100°C. They consist of a large organic cation and a small organic or inorganic anion. The large dimension of its ions does not allow the organization to a crystalline structure and hence are liquids at lower temperature in comparison to conventional salts. The interest in ILs as separation media can be attributed to its interesting chemical and physical properties such as low vapor pressure, high thermal and chemical stability, non-flammability and good solvation properties [8-10]. These are also referred to as designer solvents [11], as their properties such as polarity, viscosity and hydrophobicity can be tuned by combination of cations and anions. These properties make them a desirable class of solvents for liquid-liquid extraction. Most commonly used anions and cations are depicted in Figure 1.2. In the past years ILs have been used for extraction of value added components from biomass [10, 12] and in synthesis [13], in material chemistry as electrolytes for the electrochemical industry and liquid crystals. Moreover, ILs also demonstrated a good performance in biocatalysis [14], while providing a non-denaturing environment for biomolecules and maintaining the protein structure and enzymatic activity [15]. These liquids are promising alternatives for separation and they need to be investigated with respect recycling so as to decrease cost and enhance sustainability of the process.

Figure 1.1 Schematic picture of a typical aqueous two phase diagram Re-printed from [7] with permission from Elsevier
Microalgae

Microalgae are photosynthetic unicellular microorganisms capable of accumulating lipids and other value added components in its biomass (e.g. proteins, pigments). They are considered as promising feedstock for biofuel production as they do not compete for arable land and do not place additional stress on food production [17]. Microalgae have a rigid cell wall [18] and are composed of a non-hydrolysable biopolymer; algenan. The complexity and filamentous structure of algal cell walls makes them stronger and more difficult to disrupt than cells from other organisms. Furthermore, effective solvent extraction requires that the solvent can firstly penetrate the solid matrix enclosing the lipid, secondly physically contact the lipid and thirdly solvate the lipid [19]. Since most microalgae are protected by a cell wall that limits the solvents access to the lipid, mechanical disruption techniques such as bead milling, homogenisation, etc., are required to break open the cells and release the intracellular components (e.g. proteins, carbohydrates, lipids, pigments). Conventional harvesting and extraction processes are energy intensive [20] and thus are the main barriers in the economical production of algae biofuels on a commercial scale [21]. These processes are thus in need of innovation. Traditionally, lipids are extracted from biomass using a combination of organic solvents such as chloroform, methanol and water i.e. Bligh and Dyer’s method [22]. Although the lipid extraction efficiency is higher, such a process would generate a significant amount of solvent waste on large scale which is costly to recycle. However, to develop a process with biodiesel as the only end product is not sustainable and economically feasible. Additionally, use of solvent denatures the proteins and makes the biomass unsuitable for isolation of valuable products (e.g. proteins). Therefore, biorefining of microalgae must be done so as to fractionate it into valuable components such as proteins and
carbohydrates in addition to lipids [23]. Biorefinery is an integrated facility that combines processes and equipment to produce biofuel, high-value biochemicals and energy from the same biomass. Thus, the primary requirement to develop a microalgal biorefinery is to be able to separate the different fractions such that the components retain their functionality. This necessitates the development of separation technologies that are mild and inexpensive [24]. Although, currently there is no established biorefinery process for microalgae, a typical biorefinery would involve multiple unit operations as shown in Figure 1.3. While each of the unit operations is being studied so as to develop alternatives to the current processes [24], this thesis focuses on alternative extraction processes based on ionic liquids.

![Figure 1.3 Schematic diagram of unit operations involved in biorefinery of microalgae](image)

**Outline of thesis**

In this thesis the possibility of using IL and IL based systems for fractionation of components into hydrophilic and hydrophobic components using IL based systems is investigated. Both known and novel systems for extraction of fragile biomolecules from complex biomass such as microalgae are investigated. The effect of ILs on microalgal biomass pre-treatment and subsequent extraction of biomolecules therein was also explored.

**Section 1: Extraction and separation using IL based two phase system**

In chapter 2 we provide a literature review on extraction of proteins using aqueous two phase systems and also discuss about the stability of proteins in ILs. This chapter describes a general approach to protein purification using IL based ABS and factors that influence the partitioning of the proteins in the system. It also discusses the challenges in developing a successful ABS for extraction of proteins. Stability of proteins in ILs is a prerequisite for its use in extraction of these fragile molecules. In chapter 3, the stability of Rubisco together with two other model proteins BSA and IgG, in aqueous solution of ILs was studied. Furthermore, studies were done to systematically understand the relationship between IL concentration and protein stability with the ATPS system. The influence of different process parameters on extraction was studied and compared with the conventional two phase system (PEG/potassium citrate ATPS). In chapter 4, we studied the feasibility of using a novel system, IL based
emulsion stabilised by microgel, for separation of hydrophilic and hydrophobic components from microalgae biomass while keeping the proteins in their native form. Continuous separation of biomolecules using such a system was also investigated.

**Section 2: Pre-treatment using Ionic liquid**

In **chapter 5**, the effect of IL pre-treatment on microalgal cells was investigated under mild conditions of temperature. The extraction efficiency of pigment after pre-treatment was determined and a hypothesis for the extraction of pigment without cell disruption is also presented. In **chapter 6**, we went a step ahead and studied the fractionation of microalgal biomass after pre-treatment with ILs. Extraction of lipids after pre-treatment with ILs and subsequent cell disruption to yield proteins and carbohydrates were studied. Moreover the stability of extracted protein was also studied.

Finally **chapter 7** discusses the major findings in this thesis and bottlenecks and challenges in using ILs and IL based systems for extraction of biomolecules from complex matrices. More studies would be needed to have better understanding of these system and therefore opportunities for future work is also presented.
References


Section I
Extraction and separation using IL based two phase system
Chapter 2

Extraction of Proteins with ABS

This chapter is published as a book chapter:
Chapter 2

Abstract

Over the past years, there has been an increasing trend in research on the extraction and purification of proteins using aqueous biphasic systems (ABS) formed by polymers e.g. polyethylene glycol, (PEG). In general, when dealing with protein purification processes, it is essential to maintain their native structure and functional stability. In this context, ABS, liquid-liquid systems where both phases are water-rich, provide a biocompatible medium for such attempts. More recently, it was shown that the versatility offered by ABS is further enhanced by the introduction of ionic liquids (ILs) as alternative phase-forming components. This chapter describes and highlights the current progress on the field of proteins extraction and purification using IL-based ABS. The general approach for protein extraction using IL-based ABS and factors influencing the partitioning are discussed. In addition, the challenges to be overcome on the use of IL-based ABS for proteins extraction are also presented.

Keywords: Proteins, ionic liquids, aqueous biphasic system, extraction, purification, stability.
Introduction

Proteins are an integral part of all living systems and have various applications in food and feed (both relatively low-value), and pharmaceuticals (high-value). Purification of proteins involves various unit operations using low to high resolution techniques to obtain proteins with desired purity and quality. Proteins, being fragile molecules, can be easily denatured by acid/base solutions, salts and high temperature. Therefore, mild operation conditions for their recovery and purification are required in order to maintain their nativity and functionality. With the current advances in biotechnology, a large increase in the titers of protein production was already observed; yet, the development of cost-effective purification methods is still required. The high cost of protein purification continues to remain a bottleneck in downstream processing of proteins, and mainly for protein value-added biopharmaceuticals. On the other hand, in the field of food and feed, proteins are obtained from e.g. soya and also there has been a growing interest in 3rd generation biofuels from microalgae. For instance, in fuel production processes, large amount of proteins are generated which could be used for feed and food [1-3]. In fact, to make these processes economically feasible it is necessary to refine other components from biomass. Proteins are a major fraction of algae biomass and are normally denatured by the solvents used for lipid extraction. The main challenge therefore lies in separating the proteins in their native form without affecting their functionality. Thus, depending on the biomass or initial medium, protein purification protocols vary and drive the development of more specific, robust and cost-effective methods [4].

Aqueous biphasic systems (ABS) based on polymers were first proposed by Albertsson [5], who studied their applicability in proteins extraction and purification. ABS allows the integration of concentration and purification processing steps and serve as an alternative approach to the traditional processes. Typical ABS are formed by mixing polymer-polymer and polymer-salt combinations above given concentrations to form two distinct aqueous phases, each one enriched in one of the phase-forming components. Both phases are water-rich (~80-90% w/w), and thus ABS can provide a mild and gentle environment for protein separation without affecting their native structure and stability [6, 7]. In addition to the largely investigated polymer-based ABS, in the last decade, ionic liquids (ILs) were proposed as alternative phase-forming components of ABS [8]. And because of the inherent properties of ILs, this possibility allowed the use of ABS in a new range of applications.

The interest on ILs as extractive solvents increased primarily because of their non-volatile nature, which is the major advantage over traditional organic volatile solvents. In addition to their non-volatility, ILs, being composed of cations and anions, can be more easily tuned to achieve specific properties, such as a tunable polarity, viscosity and solvent miscibility. Their tunable polarity enabled them to be used in biotransformations to increase substrate solubility, to dissolve enzymes and to tailor the reaction rate [9]. Moreover, due to their tailoring ability, ILs are also able to form ABS not only with inorganic salts, but also with polymers [10], carbohydrates [11] and amino acids [12]. The main advantage of IL-based
ABS over the conventional systems comprises their ability to tune and tailor the properties of the coexisting phases by permutation and combination of different cations and anions, thereby improving the selectivity of these systems for a wide variety of solutes [13].

Based on the advantages and large recent interest on IL-based ABS for separation purposes, this chapter describes the general approaches of protein purification described in the literature using IL-based ABS and factors that influence the partitioning of proteins in these systems. The challenges in developing a successful ABS for extraction of proteins are also discussed. Finally, this chapter aims a better understanding on the mechanisms ruling proteins extraction using IL-based ABS. Figure 2.1 depicts a scheme on the approach required to use IL-based ABS for the extraction of proteins.

**Figure 2.1 Approach required for proteins extraction using IL-based ABS**

**Extraction of proteins/enzymes using IL-based ABS**

The extraction of proteins using IL-based ABS has been studied by different research groups, and for which a summary is given in Table 2.1. This table was adapted from [14] and updated to include more recent studies.

As a first point, only water-miscible ILs are able to form ABS since water-immiscible ILs do not form two aqueous-rich phases (see Table 2.2). IL-based ABS are formed by mixing water-miscible ILs with salts, carbohydrates, amino acids and polymers [8, 10, 12, 15]. The ability to form ABS with solutes other than salts has indeed been studied [8, 10, 12, 15], but their efficiency in extracting proteins is however scarcely studied. Although more promising than inorganic salts in what concerns the use of more biocompatible systems, these alternative systems suffer the drawback of only being able to form ABS with a limited number of ILs due to their low salting-out ability (carbohydrates, amino acids and polymers versus salts).
Table 2.1 Investigated IL-based ABS for the extraction of proteins (updated from [14]).

<table>
<thead>
<tr>
<th>Protein</th>
<th>IL-based ABS</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bovine serum albumin (BSA)</td>
<td>[C₄mim]Cl/K₂HPO₄, Ammoeng 110™/K₂HPO₄-KH₂PO₄, [C₄mim]Br(n = 4,6,8)/K₂HPO₄, [C₄mim][N(CN)₂]/K₂HPO₄, Guanidinium-based ILs/K₂HPO₄</td>
<td>[16], [17], [18], [19], [20], [21],[22], [23]</td>
</tr>
<tr>
<td>Ovalbumin</td>
<td>[C₄mim]Cl/K₂HPO₄, ILs⁶/K₂HPO₄, Guanidinium-based ILs/K₂HPO₄</td>
<td>[24],[20], [21], [23]</td>
</tr>
<tr>
<td>Lysozyme</td>
<td>Ammoeng 110™/K₂HPO₄-KH₂PO₄, Guanidinium-based ILs/K₂HPO₄</td>
<td>[17],[23]</td>
</tr>
<tr>
<td>γ-globulin</td>
<td>[C₄mim]Br(n = 4,6,8)/K₂HPO₄</td>
<td>[18]</td>
</tr>
<tr>
<td>Myoglobin</td>
<td>[C₄mim]Cl/K₂HPO₄, Ammoeng 110™/K₂HPO₄-KH₂PO₄</td>
<td>[24],[17]</td>
</tr>
<tr>
<td>Haemoglobin</td>
<td>[C₄mim]Cl/K₂HPO₄, ILs⁶/K₂HPO₄, Hydroxyl ammonium ionic liquids⁵/K₂HPO₄</td>
<td>[24],[20], [21]</td>
</tr>
<tr>
<td>Cytochrome c</td>
<td>[C₄mim]Cl/K₂HPO₄, [C₄mim]Br(n = 4,6,8)/K₂HPO₄, Amino-based ILs⁶/K₂HPO₄, Glycine-based ILs⁶/K₂HPO₄</td>
<td>[24],[18],[25], [26]</td>
</tr>
<tr>
<td>Fungal proteins</td>
<td>[C₄mim]Cl/K₂PO₄</td>
<td>[27]</td>
</tr>
<tr>
<td>Trypsin</td>
<td>Ammoeng 110™/K₂HPO₄-KH₂PO₄</td>
<td>[17], [18], [20], [23]</td>
</tr>
<tr>
<td>Lipase CaL-A</td>
<td>[C₄mim][C₂SO₄]/(NH₄)₂SO₄</td>
<td>[28]</td>
</tr>
<tr>
<td>Lipase CaL-B</td>
<td>Imidazolium-based ILs⁶/K₂HPO₄-KH₂PO₄</td>
<td>[29]</td>
</tr>
<tr>
<td>Thermomyces lanuginosus lipase (TIL)</td>
<td>[C₄mim][C₂SO₄]/K₂CO₃</td>
<td>[30]</td>
</tr>
<tr>
<td>Alcohol dehydrogenases</td>
<td>(Ammoeng 100™/Ammoeng 101™/K₂HPO₄-KH₂PO₄</td>
<td>[13]</td>
</tr>
<tr>
<td>Horseradish peroxidase</td>
<td>[C₄mim]Cl/K₂HPO₄</td>
<td>[31]</td>
</tr>
<tr>
<td>Rubisco</td>
<td>Iolilyte 221PG/KH₂PO₄-Na₂HPO₄</td>
<td>[32]</td>
</tr>
<tr>
<td>Wheat esterase</td>
<td>[C₄mim][BF₄]/NaH₂PO₄</td>
<td>[33]</td>
</tr>
</tbody>
</table>

*a Details of ILs used can be found in the corresponding literature.*
Table 2.2 Commonly used cation/anions combination of water-miscible versus water-immiscible ILs.

<table>
<thead>
<tr>
<th>Water miscible ILs</th>
<th>Cations</th>
<th>Anions</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Imidazolium, Pyridinium, Ammonium, Phosphonium</td>
<td>Chloride, Bromide, Fluoride, Alkylsulphate, Tosylate, Tetrafluoroborate, Dicyanamide</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Water immiscible ILs&lt;sup&gt;a&lt;/sup&gt;</th>
<th>Cations</th>
<th>Anions</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Imidazolium, Pyridinium, Ammonium, Phosphonium</td>
<td>Bistriflamide, Hexafluorophosphate</td>
</tr>
</tbody>
</table>

<sup>a</sup>Water immiscible ILs also contain water (~2-6%)  

In general, most of the studies reported in the literature deal with imidazolium-based ILs composed of halogens or [BF<sub>4</sub>]<sup>-</sup> anions. Recently, ABS based on guanidinium-based ILs have been studied for proteins extraction [22, 23], and where it was shown that model proteins, such as BSA, could be extracted with high efficiency for the IL-rich phase without losing its native structure and maintain its stability. IL-based ABS could thus serve as potential platforms for protein extraction if the stability of proteins at the IL-rich phase is maintained. While most of the studies focused on the extraction efficiency of model proteins/ enzymes, the studies carried out by Dreyer [17] and Pei [18] made an attempt to understand the mechanisms responsible for the high extractions attained.

**Stability of Proteins in IL-rich phases**

Proteins are complex macromolecules and require a gentle environment to maintain their structural and functional integrity. Changes in this environment, such as solvent concentration, pH, ionic strength, and temperature, could result in denaturation of proteins. Thus, the primary criterion for any protein purification process is the ability to maintain the proteins’ structural integrity and functionality. In this context, to use IL-based ABS for the purification of proteins it is necessary to understand their stability in aqueous solutions of ILs. There are some studies carried out to infer on protein-IL interactions [34, 35], and where model proteins have been used, namely BSA, lysozyme and cytochrome c. However, there are other proteins with higher commercial value, such as monoclonal antibodies, Rubisco (Ribulose-1,5-biphosphate Carboxylase Oxygenase), etc., that should be studied in what concerns their stability in aqueous solutions of ILs and their feasibility to be extracted by ABS. Although some hydrophobic ILs are able to stabilize enzymes [36, 37], they are not discussed in this chapter since these do not form ABS.

In the studies regarding the stability of proteins in aqueous solutions of ILs, the techniques employed to monitor the proteins structural and thermal stability include UV Spectroscopy, Fluorescence, Circular Dichroism (CD), Small Angle Neutron Scattering (SANS), Differential Scanning Calorimetry (DSC), Dynamic Light Scattering (DLS) and Size Exclusion Chromatography (SEC).The stability studies were designed to address the factors that influence the formation of ABS and the stability of proteins, such
Extraction of biomolecules using ABS

as: (i) type of IL; (ii) concentration of IL; (iii) other process conditions, such as pH, ionic strength and temperature; and (iv) protein properties, such as size, charge and surface hydrophobicity.

ABS consists of two aqueous-rich phases: an IL-rich phase and a phase rich in salt, polymer, amino acid or carbohydrate. The concentration of IL in the IL-rich phase of ABS can vary from 1.5-3.0 mol/kg [38], and thus it is prudent to study IL–protein interactions in aqueous solutions. Moreover, it was already shown that the concentration of IL has a strong influence on the protein stability [32, 35]. In our recent study, we have shown that the protein's stability in aqueous solutions of ILs is influenced by the concentration of IL, and by the protein properties, such as size and complexity of the molecule [32]. In this study, the stability of BSA, IgG and Rubisco were studied in aqueous solutions of two ILs, Iolilyte 221 PG and Cyphos 108, at different concentrations [32]. It was found that as the concentration of the IL increases (0-50%, v/v) the proteins start forming aggregates. Rubisco (~540 kDa), being a large complex protein/ enzyme that consists of eight large and small subunits, begins to aggregate at lower IL concentrations (~30%, v/v), while BSA (~67 kDa), a smaller protein, forms no aggregates or only negligible aggregates at 50% v/v of IL(Iolilyte 221 PG). IgG (~150 kDa), with an intermediate size, forms aggregates at 50% (v/v) of Iolilyte 221 PG. In this study [32], the aggregates formation was monitored using SEC and DLS studies. In an additional study, SANS results showed that human serum albumin and cytochrome c form aggregates at high concentrations (50%, v/v) of [C₄mim]Cl and retain their high order structure at lower IL concentrations (25%, v/v) [39]. Lysozyme and interleukin-2 (IL-2) showed increased thermal stability in aqueous solutions up to 40% (w/w) of IL, although it is dependent on the pH [40], indicating thus that the charge of the protein also influences its stability in ILs aqueous solutions. Different ILs with varying “kosmotropicity” were also investigated for their effect on proteins structure and long term stability [41]. In this study, cytochrome c showed no significant changes in its structure when dissolved in hydrated choline di-hydrogen phosphate (containing 20% w/w of water). Cytochrome c additionally showed a higher thermal and long term stability, leading the authors to conclude that the “kosmotropicity” of ILs has strong implications on the proteins stability [41].

The influence of inorganic salts and ion-specific-induced precipitation of proteins is well described by the Hofmeister series [42]. Since ILs are also composed of ions, their influence on protein stability in aqueous solutions can also be explained, to some extent, by the Hofmeister series [34, 41, 43, 44]. Ions can be classified as kosmotropes (water-structure makers) which stabilize proteins and chaotropes (water-structure breakers) which destabilize proteins. The rank of kosmotrope-chaotrope ions according to the Hofmeister series is shown in Figure.2.2. The most suitable combination to enhance protein stability comprises a kosmotropic anion and a chaotropic cation [45-47]. Accordingly to the example described above, the cytochrome c stability in hydrated choline dihydrogen phosphate is a result of this type of ions combination. Though the stability of proteins in ILs can be explained by the Hofmeister series, some deviations were also found while following a reverse trend [44]. Large arrays of
Chapter 2

factors are responsible for the proteins stability in aqueous solutions of ILs, such as the ability to establish hydrogen-bond, electrostatic and dispersive interactions and hydrophobicity.

**Figure 2.2 Hofmeister series and protein stability**

**Partitioning Behavior of Proteins in IL-based ABS**

It has been shown that the partitioning of solutes in typical PEG-based ABS is primarily governed by the system properties, such as type and concentration of phase-forming components, pH, temperature, and the solute properties, such as hydrophobicity, charge, molecular weight, etc. [48]. Thus, partition coefficients and selectivity can be tuned by modifying these parameters. The extraction of proteins using IL-based ABS has been studied by several authors (see Table 2.1), revealing that proteins partition preferentially to the IL-rich phase. Most of these studies are however empirical and to be able to use IL-based ABS as a separation tool on a preparative scale, it is mandatory to understand the mechanisms and factors influencing the partitioning of proteins in these systems.

Different authors have studied the influence of the phase-forming components, concentration, pH and temperature on the partitioning of proteins in IL-based ABS [20, 21, 32, 49]. Proteins distribution in ABS depends on their ability to interact with the phase-forming components and extraction conditions, so that the separation could be protein specific. Cao et.al. [31] studied the extraction of horseradish peroxidase in four alkylimidazolium-based ABS. The enzyme partitioned to the IL-rich phase but its activity decreases with the increase in the alkyl side chain length of the IL. In the same study, increasing the IL concentration favors the maintenance of the enzyme activity. Dreyer et al. [13] studied the feasibility of ABS formation with ammonium-based ILs and showed that Ammocen 110 forms ABS more easily than Ammocen 100 and Ammocen 101. Ammocen ILs contain an oligo-ethylene side chain in the cation which was expected to have a stabilizing effect on the enzyme (alcohol dehydrogenase) extracted. A low temperature for ABS formation together with ILs with oligo-ethylene side chains demonstrated to provide a gentle environment for protein extractions [13]. Desai et al. [32] showed that the partition coefficient of Rubisco increases as the IL and salt concentration increases; however, a decrease in the enzyme activity was observed with higher concentrations (>20%, w/w) of IL.
summary, all these results indicate that the chaotropicity of the IL and its concentration influence the stability of the protein to be extracted.

The system parameters (pH and temperature) also influence the partitioning of proteins to the IL-rich phase through the modification on the proteins charge and surface properties. Protein properties contributing to their partitioning in ABS can be summarized as follows [5]:

\[
\log K = \log K_0 + \log K_{el} + \log K_{\text{hphob}} + \log K_{\text{size}} + \log K_{\text{biosp}} + \log K_{\text{conf}}
\]  

(2.1)

where \( K \) is the partition coefficient and \( K_{el} \), \( K_{\text{hphob}} \), \( K_{\text{biosp}} \) and \( K_{\text{conf}} \) are, respectively, electrostatic, hydrophobicity, biospecificity and configuration which contribute to the partition coefficient value, while \( K_0 \) represents additional factors.

Partitioning of proteins is governed, in a large extent, by the pH of the system. Depending on their isoelectric point, proteins carry a net positive or net negative charge at a given pH. The extraction of BSA, myoglobin, lysozyme and trypsin using IL-based ABS at different pH values showed that proteins are preferentially transferred to the IL-rich phase as the pH increases [17]. On the other hand, the molecular weight of the protein also influences its partitioning in the biphasic system. Dreyer et al. [17] showed that larger proteins, such as BSA, are better extracted in the IL-rich phase while smaller proteins, like myoglobin, remain in the salt-buffer-rich phase. In a separate study, Rubisco, which is large protein (540 kDa) is also extracted into the IL-rich phase [32].

Du et al. [16] studied the extraction of BSA from biological fluids using imidazolium-based ABS and observed that the electrostatic interactions and salting-out effect are the driving forces in proteins partitioning. In summary, research groups [16, 17] have shown that there is a strong correlation between the protein charge and its partitioning in IL-based ABS. Thus, indicating electrostatic interaction between the amino acids on the protein surface and IL cations to be the main driving force. On the other hand, Pei et al. [18] have shown that hydrophobic interactions are the main driving force for protein extraction in IL-based ABS. In the same study, the influence of temperature on the extraction of BSA was evaluated demonstrating that higher temperatures favor the partitioning of proteins to the IL-rich phase.

It could be summarized that the partitioning of proteins in IL-based ABS can be tuned by changing the phase-components and their composition, pH and temperature of the system. Nevertheless, partitioning in IL-based ABS is a quite complex phenomenon not influenced by a single factor, yet it is a result of a combined effect of these factors.

**Recovery of Proteins from the IL-rich Phase**

Like conventional ABS, proteins extraction using IL-based ABS involves two main steps: (i) forward extraction, i.e., extraction of the protein from the initial source/matrix into one of the phases (here, IL-rich phase); and (ii) recovery of the (purified) protein from the IL-rich phase.
In conventional ABS, proteins can be recovered by modification of system parameters, such as pH, change in salt concentration or addition of other salts. The main goal is to achieve a high recovery of a protein with a high purity level without affecting the functionality of the protein. This is indeed one of the major lacunas in the literature since there are almost no attempts on the literature to this end. An isolated work was recently published by Pereira et al. [50] where the protein (BSA) was recovered by dialysis from the IL-rich phase, and allowing the further use of the ABS in a new extraction step. The authors [50] demonstrated the recovery of the protein and the IL reusability in three-step consecutive extractions, concluding that IL-based ABS can be adequately reused without losses on their extraction performance.

ILs being salts and proteins being macromolecules, their separation can be achieved by ultrafiltration and/or nanofiltration, induced precipitation and chromatographic techniques, such as size exclusion chromatography and by the use of affinity tags (HisTags) able to help in recovering the protein from the IL-rich phase by Immobilized Metal Affinity Chromatography (IMAC). Protein recovery studies are thus one of the major lacunas in the IL-based ABS field and must be investigated in the near future.

Conclusions and Future Perspectives

IL-based ABS is a promising platform for the extraction and purification of proteins. However, there are still some issues which need to be addressed to be able to use IL-based ABS on a commercial scale, namely.

1. Currently, studies on proteins of commercial importance are scarce; only few studies were performed, for instance for rubisco and alcohol dehydrogenases. Most studies in the literature address model proteins (BSA, lysozyme, etc.).

2. With a plethora of ILs available and the complex and variable nature of proteins it is difficult to generalize or to predict the behavior of proteins in IL-based ABS. However, the setup of a well-defined guideline with respect to some protein classes would be useful. A mechanistic modelling approach still seems to be far off.

3. Stability of proteins in ILs is the prime requirement to guarantee the viability of IL-based ABS for protein separation. Most studies on this line are focused on model proteins, such as BSA and lysozyme. A pragmatic approach would be to create a public and free available (online) database with respect to the functional stability of commercial proteins in IL-based systems.

4. More sophisticated analytical methods to quantify proteins in the IL-rich phase should be attempted to avoid interferences from the IL. Also for preparative chromatography, the stability and functionality of currently available resins needs to be determined.

5. The high costs of ILs are one of the major drawbacks when envisaging the large-scale application of IL-based ABS. The re-use of ILs in large scale applications is essential to guarantee the economic viability.
6. The ILs used for ABS formation are water-soluble and hence can enter into the ecosystem. Thus, toxicity and biodegradation of ILs poses another concern and must be considered while designing protein extraction and separation processes.

All these points require not only extra efforts to study different IL-based ABS, but more focused studies on the use of biodegradable and biocompatible ILs and efficient IL recycling processes. Since polymers, such as PEG, are able to maintain and even increase the stability of some proteins, IL-PEG ABS seems as an interesting option for protein extraction. Progress in IL-based ABS would open up new applications on their use, especially in biorefinery of 3rd generation biomass feedstocks (e.g., microalgae), where proteins could be separated from more hydrophobic components. IL-based ABS are novel systems and their use for proteins extraction is still in an early stage. Thus, there is ample scope for improvement in protein extractions using IL-based ABS and a strong requirement for further in depth investigations.
References

Extraction of biomolecules using ABS


Chapter 2


Chapter 3

Extraction and stability of selected proteins in ionic liquid based aqueous two phase systems

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*Rupali K. Desai, Mathieu Streefland, René H. Wijffels and Michel H. M. Eppink (2014)*

Extraction and stability of selected proteins in ionic liquid based aqueous two phase systems

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Abstract

Ionic liquid-based aqueous two-phase extraction of a plant protein, Rubisco (Ribulose-1, 5-biphosphate carboxylase oxygenase), using Iolilyte 221 PG and sodium potassium phosphate buffer was investigated as a new alternative extraction method and compared with a conventional PEG-based two-phase system. The influence of various factors, such as concentration of phase components, pH and temperature on partitioning of Rubisco, was evaluated by design of experiments. Rubisco partitions to the ionic liquid (IL) phase and the partition coefficient for IL based two-phase system were 3-4 times higher than in a PEG-based system. Additionally, studies were done in aqueous solution of IL with varying concentrations to develop a relationship between IL concentration and protein stability. In addition to Rubisco, the stability of BSA and IgG1 was investigated in aqueous solution of two ionic liquids: Iolilyte 221 PG and Cyphos 108. No fragmentation or aggregation was observed at 10% w/w concentration of the ionic liquid. However, all three proteins studied formed aggregates at 50% w/w concentration of ionic liquid. This indicates a narrow range of IL concentration for their application in protein extraction.

Key Words: Rubisco, Ionic Liquid, ATPS, BSA, IgG1, Bio-refinery
Introduction

Aqueous two phase extraction of biomolecules has been widely studied over the past decades. First used by Albertsson [1] in the 1950's, the technique has gained attention because it combines several early processing steps such as clarification, concentration and primary purification in one step. The technique is used as a mild primary recovery step for reducing the processing volumes [2]. Aqueous two phase systems (ATPS) are formed by combining two aqueous solutions of polymer and polymer, polymer and salt or salt and salt. When these phase components are mixed beyond a certain critical concentration they separate in two distinct aqueous phases. Selective partitioning of the target molecule depends upon the affinity of the molecule for the phase component used (Polymer type, salt) and system parameters such as pH, temperature and tie line length. PEG is known to stabilise proteins and high water content (~80-90%) in the bulk of the two phases provides a gentle environment for the protein. Thus, aqueous two phase extraction is a mild and the most suitable for extracting biologically active molecules [3-6]. This system has further developed by implementing a new class of extractants, "ionic liquids", to form an aqueous biphasic system. ATPS based on ionic liquids are formed by combining an ionic liquid (IL) solution with a salt solution [7], thus replacing the polymer component of the conventional polymer-salt two phase system.

Ionic liquids are salts, composed solely of ions with a melting point below 100°C. These ILs when compared to organic solvents have low vapour pressure, high solvation capacity and better thermal and chemical stability. Ionic liquids are emerging as the new class of solvents with tuneable properties. The physical properties of ionic liquids such as polarity, hydrophobicity, viscosity can be controlled by permutation and combination of anions and cations. This high tunability makes them a desirable class of extractants in liquid–liquid extraction. Ionic liquids are studied in many fields, ranging from inorganic synthesis [8], extraction of metals [9] to biocatalysis [10]. Despite the several interesting features, most ILs suffers the drawback of being expensive and poorly biodegradable. Nevertheless ILs could be attractive if they could be regenerated and reused.

Aqueous two phase extraction studies with ionic liquids normally involve the use of imidazolium ionic liquid as the cation. Coutinho and co-workers [11] have done some ATPS studies based on ionic liquid with phosphonium as the cation. The Kragl group [12] studied an ammonium based ionic liquid – Amмоeng 110 which is effective in forming aqueous two-phase and purifying enzymes (two different alcohol dehydrogenases); this IL can stabilize the enzymes and enhances the solubility of hydrophobic substrates. In another study extraction of proteins from biological fluid has been investigated using IL-based ATPS [13]. IL based ATPS have been mainly studied for extracting small proteins such as BSA, myoglobin, amino acids and small molecules such as caffeine, vanillin, penicillin and testosterone [14-17]. Apart from this, most of the research using ionic liquids focuses on formation and characterisation
of two phase system while a few studies have been done on protein stability in ATPS [18, 19]. Leveraging on the advantages offered by ATPS and coupling the unique features of IL, such as controlled hydrophobicity, polarity and miscibility, could provide selective extraction of proteins from the biomass. Use of IL based ATPS for extracting commercially important proteins is not well explored. It is thus important to study the extraction of commercial proteins and how it affects the stability of the protein after being extracted in the ionic liquid rich phase.

In this work, studies were done to systematically understand the relationship between IL concentration and protein stability with ATPS system. Thus, extraction of pure Rubisco (purity ~80%) in IL based ATPS (Iolilyte 221 PG/Sodium-potassium phosphate ATPS) was evaluated as it would be the most interesting target for such an application. The influence of different process parameters on extraction were studied and compared with conventional two phase system (PEG/Potassium Citrate ATPS). Additionally the stability of Rubisco, together with two other model proteins BSA and IgG1 in aqueous solution of IL was studied.

The ionic liquid selected on the basis of literature study [17] are: Iolilyte 221 PG-an ammonium based ionic liquid with oligopropyleneglycol unit containing side chain and Cyphos 108 (Tributyl(methyl) phosphonium methylsulfate)- an phosphonium based ionic liquid.

The commercial proteins selected are: Rubisco-a plant protein; Monoclonal antibody (IgG1)-a therapeutic protein; and Bovine Serum Albumin (BSA)-a model protein. BSA and IgG1 were selected for the study in an effort to understand the influence of ionic liquids on proteins with varying size, complexity and isoelectric point (IEP) (Table 1). Rubisco is the most abundant protein found in nature and amounts to nearly 50% of the total protein found in green parts of plants and microalgae. There has been a growing interest in this protein, as it has a high potential to be used as an ingredient in human / animal food [20]. Microalgae have a big potential to be a source for biofuel production. However in order to make this economically feasible, more products from microalgae need to be derived [21]. Thus, purification and efficient separation of microalgal proteins, namely Rubisco is important from the bio-refinery perspective [22].

**Experimental**

**Materials**

Polyethylene glycol (PEG) 400, 1000 and 3350 and potassium citrate tribasic were obtained from Sigma. Sodium phosphate, potassium phosphate and citric acid were purchased from Merck. Ionic liquids, Iolilyte 221 PG and Cyphos 108 were procured from Iolitec. The structure of the ionic liquids used is shown in Figure 3.1 Rubisco and BSA were purchased from Sigma. IgG1 was generously provided by Synthon B.V. Netherlands. The three proteins selected (BSA, IgG1 and Rubisco) widely differ from each other in terms of source, structural complexity, isoelectric point (pl) and molecular weight (Table 3.1).
Table 3.1 Properties of Proteins

<table>
<thead>
<tr>
<th>Proteins</th>
<th>Molecular weight, kDa</th>
<th>pI</th>
<th>No. of subunits</th>
<th>Mol.wt of subunits kDa</th>
<th>Bond between subunits</th>
</tr>
</thead>
<tbody>
<tr>
<td>BSA</td>
<td>67</td>
<td>4.7</td>
<td>Monomer</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>IgG1</td>
<td>150</td>
<td>9.1</td>
<td>Four (L&lt;sub&gt;2&lt;/sub&gt;S&lt;sub&gt;2&lt;/sub&gt;)</td>
<td>Large - 50 Small - 25</td>
<td>Covalently linked subunits</td>
</tr>
<tr>
<td>Rubisco</td>
<td>540</td>
<td>5.5</td>
<td>Eight (L&lt;sub&gt;6&lt;/sub&gt;S&lt;sub&gt;8&lt;/sub&gt;)</td>
<td>Large - 55 Small - 13</td>
<td>Non-Covalently linked subunits</td>
</tr>
</tbody>
</table>

Figure 3.1 Structures of (a) Iolylite 221 PG and (b) Cyphos 108.

Methods

Aqueous two phase system based on polymer (PEG-Salt)

Potassium citrate buffer was prepared by mixing appropriate quantities of 40% w/w citric acid with 40% w/w potassium citrate tribasic to attain the desired pH. Citrate salt was selected due to its biodegradability and low environmental polluting properties. Biphasic system was prepared by mixing appropriate amounts of 50% w/w stock solutions of PEG, 50% w/w stock solution of potassium citrate, water and Rubisco solution (stock solution 2% w/w) to a final weight of 5 g. The final protein concentration in the system was 0.4 mg/g. The systems were mixed using a vortex mixture and incubated at 30°C for 30 min. The system was then centrifuged at 2500 rpm for 10 min to ensure complete separation of phases. The volume of top and bottom phase was measured and samples from respective phases were taken and analysed for the Rubisco content at 280 nm.
Aqueous two phase system based on ionic liquid (IL-Salt)

Two phase system of ionic liquid salt was prepared by mixing appropriate amounts of Iolilyte 221 PG, sodium potassium phosphate buffer, protein solution and water to a final wt. of 5 g. The buffer was prepared by mixing appropriate quantities of 40% w/w di basic potassium phosphate (K₂HPO₄) with 40% w/w mono basic sodium phosphate (NaH₂PO₄) until the desired pH was reached. K₂HPO₄ and NaH₂PO₄ were used to prepare the buffer due to the low solubility of their respective mono and dibasic salts. The system was mixed and incubated for 30 minutes at temperatures selected for the study. Samples from bottom phase were withdrawn and analysed for the Rubisco content at 280 nm. The yield in the IL-rich top phase was calculated by mass balance to avoid possible interference from ionic liquid.

Design of experiments

Modde v.9.1 Design of Experiments (DOE) software (MKS Umetrics, Sweden) was used to study the effect of different factors on the partitioning of Rubisco in PEG-Salt and Ionic liquid-salt two phase systems. A central composite face centred design (CCF) was used for both systems. Four independent variables (factors) at three levels and three replicates at the centre point were studied for the two systems. The coded value for each factor studied for the two systems are shown in Table 3.2 and 3.3.

Table 3.2 Factors and value levels used in CCF design for PEG/Potassium citrate system in % w/w

<table>
<thead>
<tr>
<th>Variables</th>
<th>Factors</th>
<th>Low value (-1)</th>
<th>Centre value (0)</th>
<th>High value (+1)</th>
</tr>
</thead>
<tbody>
<tr>
<td>PEG molecular weight</td>
<td>X₁</td>
<td>400</td>
<td>1000</td>
<td>6000</td>
</tr>
<tr>
<td>PEG concentration % w/w</td>
<td>X₂</td>
<td>24</td>
<td>25.5</td>
<td>27</td>
</tr>
<tr>
<td>Buffer concentration % w/w</td>
<td>X₃</td>
<td>24</td>
<td>26</td>
<td>28</td>
</tr>
<tr>
<td>pH</td>
<td>X₄</td>
<td>6</td>
<td>7</td>
<td>8</td>
</tr>
</tbody>
</table>

Table 3.3 Factors and value levels used in CCF for Iolilyte 221 PG/Sodium -Potassium phosphate system

<table>
<thead>
<tr>
<th>Variables</th>
<th>Factors</th>
<th>Low Value (-1)</th>
<th>Centre Value (0)</th>
<th>High Value (+1)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Iolilyte 221 PG concentration % w/w</td>
<td>X₁</td>
<td>12</td>
<td>16</td>
<td>20</td>
</tr>
<tr>
<td>Buffer concentration % w/w</td>
<td>X₂</td>
<td>20</td>
<td>22.5</td>
<td>25</td>
</tr>
<tr>
<td>pH</td>
<td>X₃</td>
<td>6</td>
<td>7</td>
<td>8</td>
</tr>
<tr>
<td>Temperature °C</td>
<td>X₄</td>
<td>15</td>
<td>25</td>
<td>35</td>
</tr>
</tbody>
</table>

The two response variables studied were partition coefficient (Kₚ) Y₁ and yield (Yₑ) Y₂. The distribution of Rubisco in ATPS was determined by measuring the partition coefficient Kₚ which is calculated as the ratio of Rubisco concentration in the upper phase to that in the lower phase.
Extraction and stability of selected proteins in ionic liquid based aqueous two phase systems

\[ K_p = \frac{C_t}{C_b} \]

The yield \((Y_e)\) is calculated as the percentage of the amount of Rubisco in the top phase to the initial amount for PEG based system and for IL based system it is calculated as:

\[ Y_e = 100 - \frac{C_p \times V_b}{Initial\ Amount} \times 100 \]

Response surface methodology was used to optimise the extraction and the responses were fitted in a quadratic (second order) polynomial regression model to understand the effect of different interactions on the responses. The model’s validity and significance were evaluated using the analysis of variance (ANOVA).

*Stability in ionic liquid solutions*

The effect of ionic liquid on the three proteins was investigated by incubating the proteins in different concentration of aqueous ionic liquid solutions. The samples were prepared by mixing the ionic liquid with water and adding the protein at a concentration of 2 mg/ml. The concentration Iolilyte 221 PG and Cyphos 108 was increased from 10-50% v/v. The stability of proteins was then analysed by gel electrophoresis and size exclusion chromatography (SEC-HPLC). The concentration of ionic liquid in the top phase of ATPS could vary from 20-60% w/w. Therefore the top phase of the ATPS was also analysed for protein stability using gel electrophoresis and size exclusion chromatography.

*Analytical techniques*

**UV Spectroscopy**

The amount of Rubisco in both PEG-Salt and IL-Salt biphasic systems was analysed by measuring the absorbance at 280 nm. All samples were analysed against blanks having the same composition but without protein to avoid interference from the phase components.

**Rubisco Activity**

Rubisco activity is measured spectrophotometrically using NADH-linked enzyme coupled system [23]. The final reaction mixture (3ml) contains 259 mM Tris, 5 mM magnesium chloride, 67 mM potassium bicarbonate, 0.2 mM \(\beta\)-nicotinamide adenine dinucleotide, reduced form, 5 mM adenosine 5'-triphosphate, 5 mM glutathione, reduced form, 0.5 mM D-ribulose 1,5 di-phosphate, 5 units alpha-glycerophosphat dehydrogenase trios phosphate isomerase, and 5 units glyceraldehyde-3-phosphate dehydrogenase/3-phosphoglyceric phosphokinase. Rubisco is added and the oxidation of NADH is measured by monitoring the change in absorbance at 340 nm over a period of 6 minutes. The enzyme activity is then calculated from the rate of NADH oxidation using an extinction coefficient of 6.22 mM⁻¹.

**Size exclusion chromatography (SEC-HPLC)**

The stability of the protein, in terms of fragmentation and/or aggregate formation, was analysed by SEC-HPLC (Thermo Separation Products P4000 pump and AS3000 auto sampler and Ultimate 3000
Diode array detector) using Biosep-SEC-S-3000 column (Phenomenex 300 x 7.8 mm column, 5μ particle size). The mobile phase was 0.1 M sodium phosphate buffer pH 7 and 0.3 M sodium chloride. The samples were centrifuged at 11000 rpm for 1 minute to remove any insoluble particles prior to injection (inj. volume 20 μl). The samples were run in an isocratic mode at a flow rate of 1 ml/min and the protein was detected at 280 nm.

Electrophoresis

To investigate the formation of fragments and aggregates and to confirm that the protein retains its native form, the samples were further analysed by native gel electrophoresis. The samples from the top and bottom phase were diluted with native sample buffer in the ratio 1:2. The samples were then applied on 4-20% Criterion TGX, Tris glycine precast gel and run with 10X Tris glycine native buffer at 125 V for 75 minutes. SDS- gel electrophoresis was performed using precast Criterion XT Bis-Tris gel 12%. The samples were mixed with the sample buffer, reducing agent and heated at 95°C for 5 minutes. The samples were applied on the precast gels and then run using 3-(N-morpholino) propane sulfonic acid (MOPS) buffer at constant voltage of 200 V. The gel (both native and reduced gels) was stained with coomassie brilliant blue R250. All the reagents used for gel electrophoresis were procured

Dynamic light scattering (DLS)

Dynamic light scattering is a non-invasive technique which measures size and size distribution of proteins, particles and other molecules in liquid solution. It measures the fluctuation in light intensity as a function of time. Hydrodynamic radius of proteins in aqueous ionic liquid solution were analysed by Zetasizer Nano S System from Malvern Ltd. The light scattering was measured at a constant temperature of 25°C and scattering angle of 173°. The samples were measured in 40μl disposable cuvettes. The viscosity and refractive index of the dispersant, Iolilyte 221 PG, was set at 1.29 mPa.s and 1.346 for 10% v/v; 2.49 mPa.s and 1.37 for 25% v/v and 9.8478 mPa.s and 1.403 for 50% v/v respectively.

Results and Discussion

In this work, partitioning of Rubisco was studied in two different ATPSs; polymer-salt and IL-salt. Influence of system components: type of polymer, salt and ionic liquid and system parameters: pH, temperature on the extraction of Rubisco was investigated using DoE. Furthermore, the stability of proteins (BSA, IgG1 and Rubisco) in different concentration of aqueous solutions of ionic liquid was investigated using Dynamic Light Scattering (DLS), size exclusion chromatography (SEC-HPLC) and gel electrophoresis (SDS-PAGE) and is discussed in this section.

Partitioning of Rubisco in PEG/salt system

Effect of system components

Different PEG molecular weights were studied to understand its effect on partitioning of Rubisco in aqueous two phase system. The protein partitioned preferentially to the PEG phase (Figure 3.2).
However, with increase in the chain length of PEG the yield decreases due to precipitation of Rubisco at the interface. This could be attributed to the fact that an increase in the chain length of PEG decreases the free volume available in the top phase [24]. Thus there is not enough space to accommodate Rubisco, which is a large protein. Precipitation of Rubisco with increase in the molecular weight is also associated to the increase in hydrophobicity of the top phase. PEG 400 was selected for further studies since it showed good recovery and partitioning with no precipitation of Rubisco.

**Figure 3.2 Effect of molecular weight on partition coefficient $K_p$ and Yield**

Different concentrations of salt and PEG were studied in order to evaluate their effect on partitioning of Rubisco. The salt concentration was increased from 24% w/w to 28% w/w and PEG concentration was increased from 24% w/w to 27% w/w. As shown in Figure 3.3a and 3.3b there is marginal increase in the partition coefficient and recovery of Rubisco with increase in concentrations of phase components.

**Effect of system parameter**

The effect of pH on partitioning of Rubisco was evaluated by varying the pH from 6-8. The net charge on the protein varies with the pH of the ATPS and depending on the pI of the protein it influences the partitioning of protein between the two phases. Rubisco partitioned preferentially to the top phase at all pH studied. The partitioning to the top phase can be explained by the electrostatic interaction between the biomolecule and PEG [25]. From the previous studies by different authors [3, 25, 26], it is observed that negatively charged proteins partition to the PEG phase. Since the isoelectric point of Rubisco is between 5.5-5.7, it is negatively charged at all the pH studied and thus partitions to the PEG phase. The response contour plots (Figure 3.3c and 3.3d) shows that an optimum extraction efficiency of ~93% and partition coefficient of ~12 is achieved at pH close to neutral ~7.

**Partitioning of Rubisco in IL/salt system**

**Effect of system components**

As observed for the PEG based ATPS, partitioning in IL-based ATPS is also influenced by several parameters such as concentration of IL and salt, pH and temperature. The effect of varying
concentration of phase components such as IL and salt on partitioning was investigated. Rubisco is partitioned preferentially to the top phase as the ionic liquid and salt concentration was increased from 12% w/w to 20% w/w and 20% w/w to 25% w/w respectively. The partition coefficient increases from 20 to 45 as the concentration of ionic liquid and salt is increased (Figure 3.4a). A corresponding increase in the yield of Rubisco was observed (Figure 3.4b). This trend of increased partition coefficient and yield is observed at pH 7 and temperature of 25°C. It is interesting to note that the partition coefficient obtained with the ionic liquid based ATPS is of the order 3-4 higher than PEG/salt system. These findings are in agreement with those done by Ruiz-Angel [27]. Ruiz-Angel [27] have reported that the partitioning of myoglobin, cytochrome C, haemoglobin and ovalbumin to the ionic liquid phase is of the magnitude 2-3 higher and is attributed to the vast difference in the polarity of the PEG rich phase and ionic liquid rich phase. The main driving force for partitioning of the protein to the ionic liquid phase is not well understood, though there have been some studies done in this direction [14, 17]. However, these studies suggest that the partitioning of Rubisco to the IL-rich top phase could be attributed to a combined effect of hydrophobic interaction, salting out effect and electrostatic interaction between the positively charged ammonium cation of the ionic liquid and the negatively charged amino acid residues at the surface of the protein.

**Effect of system parameters**

The influence of temperature and pH on formation of aqueous two phase is assessed by different authors [28] [29]. The effect of pH on partitioning of Rubisco was studied in the range 6-8. The studies show that the partitioning of Rubisco increases with the increase in pH (Figure 3.5a, b). Whereas, there is no significant change in the yield when the temperature is increased from 15°C to 35°C (Figure 3.5c, d).
Figure 3.3 Contour plots of PEG and salt concentration effects on a) Partition coefficient, b) Yield of Rubisco and Effect of pH on c) Partition coefficient and d) Yield of Rubisco

Figure 3.4 Contour plots of effect of IL and salt concentration on a) Partition coefficient and b) Yield of Rubisco
Figure 3.5 Contour plots of effect of pH on a) Partition coefficient b) Yield and Effect of Temperature on c) Partition coefficient d) Yield of Rubisco

Model fit and Analysis

A second order polynomial regression model was used to calculate the response contour plot for each response variable. These plots help to visualize the relationship between the variables and the responses [30]. The coefficient of the model’s independent variable describes the effect of the variables on the response. The effects that had less than 95% significance were considered insignificant and discarded. The following reduced regression models were obtained for the responses; partition coefficient and yield for the two systems:

**PEG/Potassium citrate ATPS**

\[
K_p = 0.958 + 0.0315X_1 + 0.022X_2 + 0.0166X_3 + 0.0734X_4 + 0.046X_1X_3 + 0.0464X_2X_4 + 0.11X_1^2 - 0.0743X_2^2 - 0.096X_3^2
\]

\[
R^2 = 0.85
\]

\[
Y_e = -1.365 - 0.175X_1 + 0.053X_2 + 0.0115X_3 - 0.040X_4 - 0.022X_1X_3 - 0.044X_1X_4 - 0.046X_3X_4 + 0.356X_1^2 - 0.12X_2^2 - 0.07X_3^2
\]

\[
R^2 = 0.91
\]
Extraction and stability of selected proteins in ionic liquid based aqueous two phase systems

**Iolilyte 221PG/Sodium Potassium phosphate system**

\[ K_p = -0.263 + 0.188X_1 + 0.085X_2 + 0.307X_3 - 0.028X_4 - 0.080X_1X_2 - 0.095X_1X_3 + 0.096X_1X_4 - 0.11X_2X_4 - 0.11X_3^2 \]

\[ Y_x = 1.98 + 0.011X_1 + 0.0009X_2 + 0.006X_3 - 0.001X_4 - 0.001X_1X_2 - 0.004X_1X_3 + 0.002X_1X_4 - 0.002X_2X_4 - 0.006X_3^2 \]

\[ R^2 = 0.92 \]

The value of \( R^2 \) defines how well the model fits the data. In the model fit, the observed \( R^2 \) value shows a good correlation with the predicted values. The statistical significance of regression model was evaluated by the analysis of variance (ANOVA). The ANOVA analysis (see Appendix, Table 1) performed on the two systems shows that the model is significant. The first F-test, which compares variation in regression model and residuals, is satisfied when \( p < 0.05 \). The second test also known as the lack of fit test, compares the model and replicate errors and is satisfied when \( p > 0.05 \). According to the ANOVA, all parameters, PEG mol. wt., PEG and salt concentration and pH have significant influence on the partitioning (\( K_p \)) of Rubisco in PEG based ATPS. However, the influence of PEG mol. wt. on yield is very prominent. Rubisco precipitates at high mol. wt. PEG and thus low yields. Similarly for the IL based ATPS, as seen in the response contour plots (Figure 3.4 and 3.5) IL, salt and pH substantially influences the partitioning of the Rubisco to the IL phase.

**Stability of the protein**

Proteins are complex macromolecules and retain their structural and functional stability in native environment. Small changes in the protein environment such as temperature, pH and solvent can alter the native fold of the protein. Studies on protein interaction with ionic liquid have been performed by different authors [31-33].

The partition coefficient obtained for the IL/salt ATPS is much higher than the PEG based systems. Nevertheless the stability of the protein in the extracted phase is a prerequisite. The samples of the IL-rich top phase were analysed by gel electrophoresis and SEC-HPLC. In the SEC-chromatogram (Figure 3.6), the band of Rubisco in the top phase is not visible. This result is in accordance with the native gel which shows a very faint band (data not shown). This indicates that Rubisco does not retain its native structure in the IL rich phase.

Although the starting concentration of IL-based ATPS was varied from 12-20% w/w, depending on the partitioning and resulting volumes of each phase, the concentration of ionic liquid in IL rich phase can vary considerably. Therefore, to understand the relationship between ionic liquid concentration and protein stability, pure Rubisco was dissolved in increasing amount of Iolilyte 221 PG and samples were run on SEC-HPLC.

The SEC-HPLC chromatogram (Figure 3.7a) shows a progressive decrease in Rubisco peak intensity with increase in ionic liquid concentration. Additionally, there is no visible fragmentation of Rubisco
observed in the SEC chromatogram. This is further confirmed by gel electrophoresis (Figure 3.7b). However, there is some amount which does not migrate and remains in the well. These studies indicate probably, formation of high molecular weight aggregates with increasing concentration of ionic liquid. The aggregates are too large to pass through the SEC column and hence not observed in SEC chromatogram. Under denaturing conditions (use of SDS and β-mercaptoethanol) these aggregates are reduced to the Rubisco subunits (55 and 13 kDa) and migrate through the gel (Figure 3.7b). However, at higher concentration of the ionic liquid, some amount of protein is still observed in the well under denaturing condition. This indicates that the aggregates so formed are linked by both non-covalent and/or disulphide linkages.

Figure 3.6 Size exclusion chromatography–IL/Salt ATPS top phase

a)
The hypothesis that high molecular weight aggregates were formed was further confirmed by DLS studies (Figure 3.7c). The DLS study shows that the hydrodynamic radius of standard Rubisco and Rubisco in 10% v/v Iolilyte 221 PG is similar. As the concentration is increased to 25% v/v, Rubisco formed an aggregate which is shown by an increase in the hydrodynamic radius. Rubisco activity in...
presence of different concentrations of ionic liquid was also monitored (Figure 3.7d). The study showed a decrease in enzyme activity with increase in ionic liquid concentration. All these results illustrate that the absence of Rubisco peak from the top phase of the aqueous two phase system is associated with the formation of high molecular weight aggregates. The study done by Dreyer [12] on Iolilyte 221 PG/Potassium phosphate aqueous two phase system shows that the concentration of Iolilyte 221 PG in the top phase can be up to 40% w/w or higher depending on the initial ionic liquid concentration in the system. This confirms that high concentration of ionic liquid in the top phase leads to aggregation.

We also studied the behaviour of BSA and IgG1 in aqueous solution containing increasing concentration of the Iolilyte 221 PG. Aqueous two phase extraction of BSA in Iolilyte 221 PG/Phosphate was done by Dreyer [17]. The study showed that BSA was extracted with an efficiency of 85-100% and has improved thermal stability in presence of ionic liquid. Our studies reveal that at high concentration of Iolilyte 221 PG (50% v/v) there is marginal decrease in peak intensity (Figure 3.8a). The SEC results together with DLS study shows aggregation of BSA at 50% v/v of Iolilyte 221 PG (Figure 3.8b). Baker and Heller [34] reported aggregation of human serum albumin at 50% v/v concentration of ionic liquid, 1-butyl-3-methylimidazolium chloride. IgG1 on the other hand too showed aggregate formation at higher concentration (Figure 3.8c). The SEC-HPLC chromatogram shows no aggregation or fragmentation of IgG1 and BSA at 10% v/v. The higher aggregation susceptibility of Rubisco (at 25% IL concentration) compared to BSA (50% IL concentration) could be related to the complexity of the structure, as BSA consist of a single globular subunit and Rubisco consist of 8 large and 8 small subunits linked by non-covalent bonds. So Rubisco is more sensitive for aggregation as different subunit might dissociate from each other and tend to form aggregates. Thus, we hypothesize that aggregation is also influenced by size/complexity (Table 3.1) of the protein.

![Graph](image_url)
Additionally, the behaviour of the proteins was studied in phosphonium based ionic liquid, Cyphos 108. IgG1 and Rubisco precipitated with increasing concentration of ionic liquid whereas BSA remained in solution at all concentrations tested. As the concentration of Cyphos 108 was increased to 20% v/v there is considerable decrease in the BSA peak intensity (Figure 3.9). However, increasing the concentration further from 30% v/v to 50% v/v there is increase in the BSA peak intensity. Simultaneous increase in the peak intensity at lower retention time (~5 min) is also observed. This indicates the formation of high molecular weight species (aggregates). However, these results exhibit a complex behaviour and require advanced analysis for understanding the underlying phenomenon.

Among the two ionic liquids studied the phosphonium based ionic liquid, Cyphos 108 affects the protein structure more than Iolilyte 221 PG. This could be attributed to the acidic nature of aqueous solution of Cyphos 108 which caused unfolding of protein and subsequent aggregation. The acidic nature of aqueous solution of Cyphos 108 is contributed by the anion methyl sulphate which is hydrolysed in water to produce HSO$_4^-$ ion [35]. In a separate study on BSA in polymer based drug delivery system degradation in acidic microenvironment by aggregate formation and hydrolysis is reported [36]. Overall the preliminary studies on the three proteins shows that they retain the native form at low concentration of Iolilyte 221 PG ~10% v/v.

**Figure 3.8** Effect of increasing concentration Iolilyte 221 PG on (a) BSA-SEC HPLC chromatogram (b) BSA-DLS (c) IgG1-SEC-HPLC chromatogram
Conclusions

The purpose of this study was to evaluate the extraction of a large complex protein; Rubisco in IL based ATPS and the stability of proteins (BSA, IgG1 and Rubisco) in aqueous ionic liquid solutions. Aqueous two phase extraction of Rubisco was investigated in PEG/Potassium citrate system and Iolilyte 221 PG/Sodium-Potassium Phosphate system. The results demonstrate that ionic liquid based ATPS showed higher partitioning of Rubisco than PEG based system. However the SEC-HPLC studies shows lack of structural stability of Rubisco. This is further supported by the studies done in aqueous solution of Iolilyte 221 PG. Similar studies on BSA and IgG1 in Iolilyte 221 PG showed formation of aggregates at higher concentration of ionic liquid. BSA in aqueous Cyphos 108 solution formed aggregates and IgG1 and Rubisco were precipitated with increasing concentration.

The empirical findings in the study suggest that higher concentration of the ionic liquids tested, results in protein aggregation. The study also reveals that the size and complexity of the protein influences protein aggregation and subsequent stability in the ionic liquid. From the study done using Cyphos 108 it is evident that the anion also strongly influences the stability of protein. A careful selection of IL with respect to anion can help in designing an IL suitable for extracting protein biomolecules. The current investigation shows that the Rubisco is structurally and functionally stable at ~10% v/v concentration of ionic liquid. Nevertheless it is important to understand that each IL is different and one size fits all does not apply. To summarize, these results help in understanding the behaviour of protein in high concentration of IL.

Acknowledgements

The authors would like to thank Technology Foundation STW (Project No 11410) and Institute for sustainable process technology (ISPT) for the financial support.
Appendix

Table 1 ANOVA table for PEG/Potassium Citrate and Iolilyte/Sodium Potassium Phosphate ATPS

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B. Iolilyte 221 PG/Sodium-Potassium Phosphate buffer system

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References


Chapter 4

Mild separation concept for functional biomolecules using IL based emulsions

Submitted as


Mild separation concept for functional biomolecules using IL based emulsions
Abstract

Fractionation of complex matrices such as biomass (e.g. microalgae) into diverse functional (fragile) biomolecules using a biorefinery concept without disrupting the biomolecules functionalities is a real challenge. Separation of diverse functional (fragile) biomolecules from complex matrices such as biomass (e.g. microalgae) in a biorefinery concept is a real challenge. Known separation processes are designed for single product recovery such as hydrophilic proteins or hydrophobic pigments, discarding the other products as waste. Here we present a novel process for separation of hydrophobic and hydrophilic biomolecules from complex matrices such as microalgae using an ionic liquid-based emulsion stabilised by microgel particles in one step while keeping product integrity. The pigments are extracted in the hydrophobic ionic liquid core and proteins in the aqueous continuous phase. In contrary to most solvent extraction procedures, with this procedure, the proteins retain their nativity opening up an extractive pathway to fully functioning proteins. A perspective towards continuous separation of this novel concept is also presented.
Introduction

Two-phase systems have long been exploited for separation and purification of biomolecules in chemical and biotech industries. Liquid-liquid extraction techniques such as aqueous two-phase systems have been used for primary isolation and separation of water soluble molecules such as proteins [1] whereas, water-organic solvent-based two-phase systems have been used for extraction of hydrophobic molecules such as pigments, lipids, small molecules, etc [2, 3]. Despite advantages, these techniques are focused on recovery of single components. For a truly biobased economy complete utilization of biomass is needed, which is, depending on the type of biomass, a valuable source of phenols, alkaloids, proteins, pigments and lipids. Moreover, product value needs to be protected without degrading fragile biomolecules such as proteins towards low value products. For complete utilization of biomass it is necessary to separate the components within a biorefinery concept into hydrophilic and hydrophobic components before further fractionation [4]. In conventional processes such as biphasic systems using organic solvents, the water-insoluble biomolecules such as pigments, fats and lipids can be recovered. However, these solvents may affect the structure and functionality of water-soluble biomolecules like proteins and enzymes [5, 6]. Therefore, it is of high importance to develop novel techniques and methods that facilitate simultaneous extraction of hydrophobic components (e.g. lipids, pigments) and hydrophilic (fragile) biomolecules from biomass without influencing their functionality and retaining the high value.

Emulsion-based separation systems could provide a solution for simultaneous extraction of biomolecules from biomass. These systems are formed by dispersion of one immiscible liquid in another liquid phase and kinetically stabilised by surfactants [7]. They have been used for both food and cosmetics as well as for separation of biomolecules, in most cases focusing on the isolation of a single product. Reversed micelles, a water-in-organic solvent microemulsion, have been used to extract proteins and enzymes [8]. A disadvantage is the poor back extraction of biomolecules from the hydrophilic core [9] damaging the product as well.

Recently there is a growing interest in ionic liquid (IL) based emulsions and microemulsions [10], due to the unique properties of ILs such as tunability [11] and negligible vapour pressure. ILs are liquid salts at temperatures below 100°C and composed of ions. ILs are potential candidates for extraction [12] and pre-treatment processes [13]. There are very few studies on the use of IL-based microemulsions as extraction media [14, 15]. Chen et al. [14] have studied the extraction of herbal compounds with different polarities using such a microemulsion. The system described however is suitable only for analytical purposes and cannot be used for large scale application. In another study [15], a dual ionic liquid-based microemulsion, wherein one IL is a surfactant and other as a substitute for organic solvent, was used for selective extraction of haemoglobin from whole blood, yet again single product recovery. This article describes the simultaneous separation and extraction of hydrophobic and hydrophilic components from biomass using an ionic liquid emulsion stabilised by polyNIPAm microgel particles.
For this purpose microalgal biomass is used as an example. Apart from being a potential source for biofuels, microalgal biomass contains other value added components such as proteins, pigments and carbohydrates. In this article we show by gel electrophoresis that the major hydrophilic component, i.e. the protein Rubisco retains its native conformation. In addition, the possibility of continuous processing using IL-based emulsions is discussed.

Materials and Methods

Materials

The ionic liquid Cyphos 109 (trihexyl(tetradecyl)phosphonium bistriflamide, purity 98%) was ordered from Iolitec and used without prior treatment. Poly-NIPAM microgel particles were synthetized and characterized as described previously [16]. Spray dried Haematococcus pluvialis cells were provided by Feyecon B.V (Weesp, Netherlands). N-isopropylacrylamide (NIPAm), N,N-methylbisacrylamide (BIS), methacrylic acid (MA), potassium persulfate (KPS), styrene, sodium dodecyl sulphate (SDS) and phosphoric acid were purchased from Sigma Aldrich. Organic solvents such as methanol, tertiary methyl butyl ether and acetone were also purchased from Sigma Aldrich. Standard astaxanthin (~97% purity) was purchased from Ehrenstorfer GmBH. Milli Q water was used for all studies. The material for electrophoresis was bought from Biorad and the staining kit was purchased from Thermo Fisher Scientific.

Methods

Preparation and separation using microgel stabilised emulsion

The microgel-stabilized emulsion is prepared by diluting a stock solution of microgel particles (23 g/l) to the desired concentration with Milli-Q water and an aqueous cell suspension (disrupted cells) and then adding it to the ionic liquid. The cell loading of the aqueous phase was kept constant at 15 g/l for all studies. Two ratios of aqueous phase to IL phase (1:9 and 3:7) and two concentrations of microgel particles in the aqueous phase (0.5 g/l and 1 g/l) were investigated. These ratios were selected as they are within the working window of the emulsion i.e. 1:9 to 4:6. Outside this range the emulsion is not stable. The samples were mixed by using a vortex tube shaker to facilitate separation of proteins and pigments in the aqueous and IL phase respectively. The separation of hydrophilic and hydrophobic components from the cells was determined after different mixing times (1 - 180 minutes). The emulsion was broken by centrifugation at 5000 rpm for 5 minutes. The aqueous phase was analysed for protein content and the IL phase for pigment content. The measurements are done in duplicate and for continuous studies one sample was taken for each time point.

Protein analysis

After breaking the emulsion, the aqueous phase is analysed for total soluble protein content. The supernatant of disrupted H. Pluvialis cells in water was analysed for total soluble protein as control.
Protein content was determined with a commercial assay kit (DC™ Protein assay, Bio-Rad, U.S.) using bovine serum albumin (Sigma–Aldrich A7030) as protein standard. The microplate assay protocol was used and the absorbance was measured at 750 nm using a microplate reader (Infinite M200, Switzerland)). The protein is expressed as the percentage of total protein in the cells:

\[
\% \text{ of Total protein} = \left( \frac{\text{Total soluble protein in aq. phase of emulsion}}{\text{Total soluble protein in cell}} \right) \times 100
\]

**Gel electrophoresis**

The conformation of the proteins after separation using the IL-in-water emulsion was checked by gel electrophoresis. The aqueous phase was diluted 1:1 with native sample buffer. About 25 μl of the diluted sample was then applied on 4–20% Criterion TGX, Tris glycine precast gel and run with 10× Tris glycine native buffer at 125 V for 75 minutes. The gel was stained using a Pierce™ Silver Stain Kit.

**Pigment analysis**

The pigment content in the IL phase was analysed by HPLC using a Thermo Scientific system coupled with DAD-Dionex 3000. Chromeleon software version 7 was used for controlling the system and for data acquisition. The method used is adapted from Fuji Chemical Industry Co., LTD, ([http://www.fujihealthscience.com/assay-method_astareal-l10.pdf](http://www.fujihealthscience.com/assay-method_astareal-l10.pdf)). The samples were analysed using an YMC carotenoid column of 4.6 × 250 mm, 5 μ. A gradient program was run as shown in Table 4.2 using a ternary mobile phase comprising of methanol, tertiary methyl butyl ether and 1% v/v phosphoric acid. The flow rate was set at 1 ml/min and the detection was performed at a λmax of 480 nm. The column temperature was set at 35°C. A calibration curve in the concentration range of 1–11 μg/ml was obtained to determine the amount of astaxanthin in the samples. The standards were run in duplicate. The samples were diluted using acetone.

**Table 4.2: HPLC gradient program**

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Chapter 4

Results and Discussion

Separation using emulsion

The potential of the microgel-stabilized IL emulsion to separate the hydrophilic and hydrophobic components from microalgae biomass was investigated. Cell suspension of broken *H. Pluvialis* cells and microgel particles was added to IL and vortexed for different time intervals up to 180 minutes. Samples were taken after different mixing times and analysed for protein and pigment content showing the efficiency of extraction. It was observed that the pigments are extracted in the discontinuous IL phase (Figure 4.1) and subsequent analysis of the aqueous phase confirmed accumulation of proteins in the continuous aqueous phase. These results confirm the separation of hydrophilic and hydrophobic components in a single step.

Some of the key factors influencing the extraction efficiency of these components were investigated: the microgel particle concentration and the ratio of IL to aqueous phase. It is observed that as the relative amount of IL is increased from 1:9 to 3:7, the amount of astaxanthin extracted after 180 minutes increases from 52% to 62% of the total astaxanthin present in the cells (Figure 4.2). In our previous studies [17] we have shown that the microgel particles stabilises the IL water interface. These particles consist of a porous polymer network and allow free transport of molecules across the interface. Increasing the microgel particle concentration from 0.5 g/l to 1 g/l leads to an increase in the extraction efficiency of the pigment up to 120 minutes (Figure 4.3). This can be explained by an increase in the total interface between the water and IL phases, which speeds up the separation process.

Almost all soluble proteins remain in the aqueous phase (Figure 4.4) for all concentration and ratios studied. Additionally, the nativity of the proteins after the separation process was investigated using native gel electrophoresis. As seen in Figure 4.5, the band for the biomarker protein Rubisco (Ribulose-1, 5-bisphosphate carboxylase oxygenase), which is the main protein in the microalgae, is intact and the protein is not broken into its subunits (non-covalently linked 8 large sub units of 55 kDa each and 8 small subunits of 13 kDa each) indicating that the proteins are not denatured in the process of separation.

The extraction efficiency of astaxanthin is lower compared to conventional solvent extractions methods [2] and can be attributed to mass transfer limitation as the pigment has to be extracted from the cells and move across the microgel particles to the IL phase where it is soluble. However, the extraction could be optimised further by modifying biomass loading, using a monodisperse emulsion, use of another IL, etc. The results demonstrate the feasibility of using microgel-stabilised IL emulsions for separating hydrophilic and hydrophobic components from a complex matrix such as microagal biomass and simultaneously retaining the functionality of separated components.
Figure 4.1 Emulsion containing H. Pluvialis cell suspension a) Before mixing b) After mixing the emulsion; aq phase and IL droplets containing proteins and astaxanthin respectively; (inset) is the macroscopic picture of IL droplets formed in the emulsion.

Figure 4.2 Effect of IL- aqueous phase ratio at microgel concentration of 0.5g/l on astaxanthin extraction efficiency.

Figure 4.3 Effect of microgel particle concentration at IL: aqueous ratio of 3:7 on astaxanthin extraction efficiency.
Chapter 4

Figure 4.4 Effect on extraction yield of total soluble protein with IL: aqueous ratios of 1:9 and 3:7 at microgel particle concentration of 0.5 g/L.

Figure 4.5 Gel electrophoresis (native): M- Molecular weight marker, Std- Rubisco standard, CS- control sample supernatant after cell disruption, Aq- aqueous phase of emulsion before adding ionic liquid containing the disrupted cells in the same amount as the samples and 1 min - aqueous phase after 1 min mixing and 180 min - aqueous phase after 180 min mixing.

Continuous extraction process

A continuous process is preferred over a batch system, primarily because it is less time consuming. As a proof of concept for the use of microgel-stabilized IL emulsions in a continuous separation process, a test run was made using the setup depicted in Figure 4.6. The flow rates were set such that the ratio of the IL phase to the water phase in the mixing vessel is 3:7. The cell suspension was mixed with the aqueous microgel solution under magnetic stirring and was equilibrated for initial 30 minutes followed by running in continuous mode until 100 minutes. Samples were taken from the vessel at different time intervals, centrifuged to break the emulsion and finally analysed. It was observed that astaxanthin extraction efficiency was ~45% and that the protein content was ≥85% (see Figure 4.7) after 30 minutes.
and comparable to the batch experiment with similar residence time. However, further improvements in the set-up such as use of different static mixers, would help by increasing the contact time between the phases and thus improving the mass transfer rates. Additionally, the separation between the phases can be enhanced by using temperature controlled settler since the polyNIPAm particles collapse at temperatures higher than 32°C, which promotes the breaking of emulsion [17].

Figure 4.6 Schematic of continuous set up: a) Ionic liquid- Cyphos 109; b) Cell suspension + Microgel particles; c) Peristaltic Pump; d) Mixing vessel; e) Collection Tank

Figure 4.7 Extraction yields of astaxanthin and proteins in continuous set-up

Conclusion

The potential of microgel-stabilised IL-in-water emulsions for simultaneous separation of hydrophilic and hydrophobic components from a complex matrix was investigated in this work using microalgal biomass as an example. The hydrophobic pigment, astaxanthin from *H. Pluvialis* cracked cells is extracted in the IL droplets and the proteins remain in the continuous aqueous phase of the emulsion. The astaxanthin extraction efficiency obtained in the process was ~62% and that of proteins in aqueous phase is ≥ 80%. The proteins recovered in the aqueous phase remained in their native state after separation, which is not the case for other methods such as alkaline or solvent extraction which lead to denaturation. A perspective and approach for continuous processing is also presented. Indeed further optimization studies would be needed to improve process kinetics and economics. The system provides a promising novel alternative not only for separation of diverse biomolecules from complex matrix but
could be extended for analytical applications as well. Future research should thus be concentrated on separation of biomolecules from complex matrix and finding newer application of this system.
Supplementary information

*Characterisation of Core shell microgel particles*

The core as well as the core-shell microgels were characterized by dynamic light scattering:
- Rh (cores) = 60 nm
- Rh (core shell microgels) = 365 nm
The emulsion is polydisperse and the droplets are ~100-200 μm in size
References


Section II

Pre-treatment using ionic liquids
Chapter 5

Novel astaxanthin extraction from *Haematococcus pluvialis* using cell permeabilising ionic liquids

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Chapter 5

Abstract

*Haematococcus pluvialis* (*H. pluvialis*) is a natural source of the food colorant astaxanthin. It is characterised by a thick resistant cell wall composed of a non-hydrolysable biopolymer, sporopollenin. High energy-consuming mechanical disruption is required to improve the extractability of astaxanthin. Opposed to that, this study uses a novel technology with ionic liquids (ILs) to permeabilise the *Haematococcal* cyst cell under mild conditions of temperature and improve the extraction of astaxanthin. The study shows that ≥70% astaxanthin can be extracted from intact *H. pluvialis* cells using 1-ethyl-3methylimidazolium di-butylphosphate (EMIM DBP) at a concentration of 40% w/w in a water solution and temperature of 45°C. Ionic liquids serve as a promising pre-treatment step for extraction of pigments from intact *H. pluvialis* cells. The results obtained opens new avenues for processing of algal biomass.
Introduction

Besides being a promising source for biofuels, microalgae also contain high value products such as pigments, carbohydrates and proteins. *H. pluvialis*, an unicellular green microalgae is a natural source of astaxanthin (3,3’-dihydroxy-β,β’-carotene-4,4’-dione). Astaxanthin, a red–orange ketocarotenoid, is widely used as colour additive in aquaculture and in cosmetics. Astaxanthin also has antioxidant properties which have potential health benefit in animals and humans [1, 2]. All this combined with high market value drives the research in astaxanthin production and particularly from natural sources such as algae, yeast and crustacean by-products [3].

*H. pluvialis* forms non-motile aplanospores or resting cells under nutritional limitations or environmental stress. Astaxanthin accumulates in these cells as a mixture of mono and di-esters under growth limiting conditions [4-6] and can constitute up to 1-5% of the alga’s dry weight [7,8]. The thick cell wall increases the mechanical and chemical resistance of the cell, which necessitates the use of mechanical rupture of the cell to extract the astaxanthin. In aquaculture, *H. pluvialis* is used as feed supplement for improving the colour of salmons and crustacean species to obtain a better market price for these products. Ruptured cells are preferred over intact cells as the latter do not cause pigmentation.

In order to optimally utilize all these algal components, a versatile biorefinery approach for microalgae is urgently needed. One of the major bottlenecks in setting up an algal biorefinery process are the high costs involved due to energy-intensive unit operations [9]. Usually mechanical cell disruption methods such as bead milling or high pressure homogenisation are used. Reducing energy consumption in this step would improve the overall process economics.

In this paper we demonstrate that a possible solution for this is the use of ionic liquids in the extraction process. Ionic liquids (ILs) are salts composed of loosely held anions and cations and, unlike inorganic salts, are liquid over a wide range of temperatures. Ionic liquids have been widely studied as extracting agents and as replacements of organic solvents in various applications such as extraction of metals, alkaloids, proteins, etc. [10-12]. Recently, ILs have also been used to dissolve algae biomass at temperatures above 100°C [13] and together with co-solvents such as methanol [14, 15]. Sun A Choi et al. [16] showed that lipid could be extracted from microalgae using a blend of different ILs. Task specific ILs in combination with ultrasonication has been used to extract astaxanthin from shrimp waste [17]. Del Sesto et al. [18] studied the extraction of botryococcenes and isoprenoids using immiscible ILs as extracting agents. Most of these studies focus on the use of ILs as an extracting agent. Teixeira [13] has nicely depicted complete hydrolysis of *Chlorella* and *Neochloris* algae cells to its constituents using ILs. However, high temperatures above 100°C are needed which could have an effect on protein and carbohydrate recovery. Different chemical, physical and mechanical methods for disruption of *H. pluvialis* have been studied [19]. The studies done by Mendes-Pinto [19] showed that mechanical disruption (e.g. homogenisation) and autoclave treatment are very effective in cell disruption and
subsequent extraction of astaxanthin. Safi and co-workers [20] studied the extraction of protein using different disruption methods from microalgae including *H. pluvialis*. The study also showed that high pressure disruption of the cell yielded most of the proteins from *H. pluvialis*.

Most of the studies mentioned above for extraction of astaxanthin from *H. pluvialis* involve high energy input such as high temperature (above 100°C) and pressure. In this paper we propose the possibility of using ILs as permeabilising agents for thick cell wall *H. pluvialis* under mild conditions. Several ILs were tested for their ability to permeabilise the cells under mild temperatures (below 100°C) and subsequently extract astaxanthin using ethyl acetate (EA). We demonstrate efficient extraction of astaxanthin from *H. pluvialis* without any mechanical cell disruption. This article underlines the potential of using ILs as pre-fractionation of micro algal cells for selective intracellular product recovery.

**Experimental**

**Materials**

Dried *H. pluvialis* intact cells were provided by Feyecon (Weesp, The Netherlands). The cells were spray dried and the astaxanthin content was ~3.2%. The ionic liquids (purity above 95%) used were procured from Iolitec and Sigma and were used without further purification (Table 5.1). Three classes of ILs were used for the study; imidazolium, ammonium and phosphonium. The imidazolium IL were selected such that Sr. No. 1, 3 and 4 (see Table 5.1) varied in the anion but cation was the same and Sr. No. 1 and 2 (see Table 5.1) have the same anion and different cation. Organic solvents and chemicals used in the study, ethyl acetate, methanol, tertiary methyl butyl ether, and phosphoric acid were purchased from Sigma. Standard astaxanthin (~97% purity) was bought from Ehrenstorfer GmBH.

**Table 5.1: List of Ionic liquids used in the study**

<table>
<thead>
<tr>
<th>Sr. No.</th>
<th>Ionic Liquid Names</th>
<th>Abbreviations</th>
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<tbody>
<tr>
<td>1</td>
<td>1-Butyl-3-methylimidazolium dibutylphosphate</td>
<td>BMIM DBP</td>
</tr>
<tr>
<td>2</td>
<td>1-Ethyl-3-methylimidazolium dibutylphosphate</td>
<td>EMIM DBP</td>
</tr>
<tr>
<td>3</td>
<td>1-Butyl-3-methylimidazolium acetate</td>
<td>BMIM Acetate</td>
</tr>
<tr>
<td>4</td>
<td>1-Butyl-3-methylimidazolium Dicynamide</td>
<td>BMIM DCA</td>
</tr>
<tr>
<td>5</td>
<td>Iolilyte 221 PG</td>
<td></td>
</tr>
<tr>
<td>6</td>
<td>Tributylmethylphosphonium methyl sulfate (Cyphos 108)</td>
<td>TBP SO4</td>
</tr>
<tr>
<td>7</td>
<td>Triisobutylmethylphosphonium tosylate (Cyphos 106)</td>
<td>TBP TOS</td>
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**Permeabilisation/Pre-Treatment with Ionic Liquid**

Dried *H. pluvialis* cells (10 mg) were treated with 2.1 ml of IL at different temperatures, 25°C, 45°C, 55°C and 65°C for 90 minutes. The ionic liquid was separated from the cells by centrifuging the cells at 3000 rpm for 10 minutes to ensure complete separation of the ionic liquid. The IL treated cells were further
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treated with 2 ml of ethyl acetate and mixed in a tube rotator for 1 hour. Ethyl acetate was separated and cells were treated with fresh 2 ml of ethyl acetate for an additional 1 hour. The ethyl acetate layer was pooled and evaporated under N₂. The extent of cell permeability was determined by measuring the amount of astaxanthin extracted. The extracted astaxanthin was then re-suspended and diluted with methanol and analysed by HPLC. The IL layer was diluted with methanol and analysed by HPLC for astaxanthin content. Different IL concentrations (IL-water) 80%, 60% and 40% w/w were studied. The control sample was prepared by breaking the cells using bead beater followed by extraction with chloroform – methanol.

HPLC analysis

HPLC analysis was performed using a Thermo Scientific system coupled with DAD (Dionex 3000). Chromeleon software version 7 was used for controlling the system and for data acquisition. The samples were analysed using YMC carotenoid column 4.6 x 250 mm, 5 μ. A ternary mobile phase comprising of methanol, tertiary methyl butyl ether and 1% v/v phosphoric acid was used. The flow rate was set at 1 ml/min and the detection was done at a λ<sub>max</sub> 480 nm. The column temperature was set at 35°C. The gradient program used is described in Table 5.2 (adapted from Fuji Chemical Industry Co., LTD, http://www.fujihealthscience.com/assay-method_astareal-l10.pdf). A calibration curve in concentration range of 1-11 μg/ml was prepared to determine the amount of astaxanthin in the samples. The standards were run in duplicate.

<table>
<thead>
<tr>
<th>Time (min)</th>
<th>Methanol</th>
<th>Tertiary methyl butyl ether</th>
<th>1% v/v Phosphoric acid</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>81</td>
<td>15</td>
<td>4</td>
</tr>
<tr>
<td>15</td>
<td>66</td>
<td>30</td>
<td>4</td>
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<td>23</td>
<td>16</td>
<td>80</td>
<td>4</td>
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<td>16</td>
<td>80</td>
<td>4</td>
</tr>
<tr>
<td>27.1</td>
<td>81</td>
<td>15</td>
<td>4</td>
</tr>
<tr>
<td>35</td>
<td>81</td>
<td>15</td>
<td>4</td>
</tr>
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Light Microscopy and Surface electron microscopy (SEM)

H. pluvialis cells after treatment with IL were observed under the light microscope (Olympus) with 60X magnification. IL treated and non-treated cells were analysed at 2 kV at room temperature in a field emission electron microscope (Magellan 400, FEI, and The Netherlands) and the images were digitally recorded. The samples were glued on sample holders by carbon adhesives tabs and subsequently sputter coated with 10nm Iridium (MED 020, Leica, Austria).

Reusability of ionic liquid

After pre-treatment of the H. Pluvialis cells with ionic liquid, the IL was separated by centrifugation. The IL was then tested for presence of pigment and degradation of IL if any by ¹H NMR (Bruker 400H). The
IL without any treatment was reused with fresh cells and the cells were treated in a manner similar to
the one described above. The amount of pigment extracted subsequently with ethyl acetate was
analysed by HPLC. Reusability of the ionic liquid without treatment was tested three times. The
amount of astaxanthin extracted after each reuse was compared with the cells treated with fresh IL.

Results and discussion

Different ILs (see Table 5.1) were evaluated for their ability to enhance permeability of H. pluvialis and
simultaneous extraction of astaxanthin from H. pluvialis at different temperatures and concentrations.
Samples were investigated using light microscopy to better understand the mechanism of action of ILs
in astaxanthin extraction.

Effect of temperature

The influence of temperature on H. pluvialis cell permeability in presence of 80% w/w ionic liquid-water
solution was investigated at different temperatures: 25°C, 45°C and 55°C. Among the different classes
of ionic liquids tested, intact cells treated with imidazolium based ionic liquid (BMIM DBP, EMIM DBP,
BMIM Acetate, BMIM DCA) were able to extract astaxanthin more efficiently than those treated with
phosphonium (TBP TOS and TBP SO₄) and ammonium (lolilyte 221 PG) based ionic liquids. The amount
of astaxanthin extracted after treatment with BMIM DBP and EMIM DBP was ~higher than BMIM
Acetate and BMIM DCA. Influence of imidazolium based ILs were also studied at an additional
temperature of 65°C. The extraction efficiency of astaxanthin measured for cells treated with
imidazolium ILs was much higher compared to the other ILs and thus to have an upper limit for the
temperature, extraction study was extended to 65°C. The permeability was measured by release of
astaxanthin which was extracted from ionic liquid pre-treated cells using ethyl acetate. As shown in
Figure 5.1, the amount of astaxanthin extracted when intact cells pre-treated with acetone, lolilyte 221
PG and phosphonium IL was ~3-5% at all tested temperatures whereas for BMIM DBP it increased from
~22% to ~65% and for EMIM DBP it increased from 36% to ~70% with temperature increase from 25°C
to 45°C. Increase in temperature from 25°C to 45°C did not increase the extraction of astaxanthin for
BMIM acetate and BMIM DCA. Intact H. pluvialis cells treated with acetone and ethyl acetate under
similar conditions could release only ~5-6% of total astaxanthin. The results show that the anion has an
impact on the cell wall. The long chain of the anion could play a role in dissolving some components of
cysted Haematococcus cell wall. Brandt et al [21] has studied the solubility of wood chips in
imidazolium based ionic liquids with different anions. The study attributed the low solubility of wood
chips treated with BMIM DCA to lower capability of DCA anion to break the hydrogen network in
cellulose and the acetate ions effectively solubilised cellulose. The Haematococcal cell wall is composed
of non-hydrolysable polymer, mainly mannan polymer [22]. Based on these results, we postulate two
hypotheses i) the mannan polymer present in the cell wall has a different solubility than cellulose ii)
complex arrangement of the polymer together with other cell wall component leading to different
solubilisation among imidazolium based ILs. However detailed study with respect to cell wall composition and effect of IL on dissolving the cell wall components is needed before any conclusive decision could be made.

**Figure 5.1** Effect of temperature on algal cell permeability for the different ILs

**Effect of IL concentration**

The pre-treatment of *H. pluvialis* cells was performed using different concentrations of ionic liquid-water solutions ranging from 20% w/w to 80% w/w at 45°C for 90 min. The concentration studies were performed using only imidazolium ILs (BMIM DBP, EMIM DBP, BMIM Acetate, BMIM DCA) since the temperature studies showed that they performed better than the other ILs. As the IL concentration increases from 20% w/w to 80% w/w, the total amount of astaxanthin (astaxanthin in EA layer + IL layer) extracted increases from 6.77 % to 62.25% for BMIM DBP and from 12% to 77.04% for EMIM DBP. The results show that pre-treatment of algal cells with IL concentrations above 40% did not further increase astaxanthin extraction efficiency. On the other hand the amount of astaxanthin lost to the IL increases from ~2% to 18% (Figure 5.2a-b) when the IL concentration increases from 20% to 80%. Pre-treatment with IL apparently weakens the cell wall due to which astaxanthin is leaked out of the cell and in the next step extracted using ethyl acetate after removing the IL from the cells. However, some amount of astaxanthin that is leaked out dissolves in the IL (Figure 5.2b). The amount of astaxanthin that goes to the IL depends on the water content. Water miscible ILs used in this study has low solubility (~120 μg/ml in 80% BMIM DBP and ~66 μg/ml in 80% BMIM DCA) for astaxanthin (data not shown) and solubility further reduces with increase in water content. This explains negligible loss of astaxanthin to the IL phase at low IL concentration and thus eliminating the need to mechanically break the cells to extract astaxanthin. Additionally, the IL could be reused when the concentration of IL used is 40% w/w. We found that EMIM DBP showed maximum extraction efficiency of ~77% among all the ILs tested. This is a good step in overcoming the costs associated with IL usage.
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Figure 5.2 Effect of IL concentration on cell permeability

In order to understand the combined effect of the IL and solvent on the cell surface the cells were studied under microscope with 60X resolution and also by SEM. Light microscopy and SEM studies were done only on cells treated with EMIM DBP, as it gave the best extraction efficiency. Light microscopy shows that the cells were intact and colourless after treatment with IL and subsequent extraction with solvent (Figure 5.3c-d), whereas cells treated with acetone or ethyl acetate under similar conditions remain intact with astaxanthin inside the cell (Figure 5.3a-b). This confirms our idea that IL permeabilises or weakens the cell wall and allows subsequent leaching of astaxanthin by the organic solvent. The SEM studies showed that untreated freeze dried cells (Figure 5.4a) appeared to have a wrinkled surface but did not show any “pitting” or cavities on the cell surface. The ethyl acetate treated cells also showed surface characteristics similar to untreated cells (Figure 5.4b). For the cells treated with IL, it was difficult to observe any changes on the cell surface mainly because of formation of a thin film around the cells which could not be removed by the solvent (Figure 5.4c-d). Although the surface characteristics were not visible, the cells looked intact. Indeed, this film could point at the possible mechanism of action of ILs in the application discussed here.

a) Amount extracted using ethyl acetate (EA) after pre-treatment with IL

b) Amount dissolved in IL after pre-treatment with IL
Novel astaxanthin extraction from *Haematococcus pluvialis* using cell permeabilising ionic liquids

**Figure 5.3** Microscopy of *H. pluvialis* a) Acetone at 45°C; b) Ethyl acetate at 65°C; c) BMIM-DBP at 45°C; d) EMIM-DBP at 45°C

**Figure 5.4** a) Untreated freeze dried cells (Control); b) EA at 45°C; c) EMIM at 45°C; d) EMIM+EA at 45°C
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Hypothesis for mechanism of action

*H. pluvialis* cells undergo considerable changes during morphogenesis from a vegetative flagellate to an aplanospores. The aplanospores have a robust cell wall accounting for 16% of its dry weight [22]. As described by Christoph Hagen [22] *H. pluvialis* cell wall is very robust and is made of highly resistant outer trilaminar sheath and a very thick secondary wall (see Figure 5.5). The trilaminar sheath is made up of sporopollenin-like material (algenan), a nonhydrolyzable aliphatic biopolymer. The secondary wall, just beneath the trilaminar sheath is composed mainly of non-fibrillar mannan polymer [22]. Hydrophilic ILs are known to dissolve cellulose by disrupting the hydrogen bond network of cellulose [23]. Teixeira [13] also showed that algae can be completely hydrolysed into its components by IL in presence of water at high temperatures (above 100°C). We hypothesize that the solubilisation of components of the outer layer of *H. pluvialis* leads to permeabilisation and thus release of astaxanthin. Since the cell wall is composed of mannan polymer, we hypothesize that the polymer structure is partially disrupted by the ionic liquid thus forming tiny holes/passage in the cell wall and paving the way for the solvent to penetrate and extract the astaxanthin. Although, surface characteristics were not clear from SEM studies due to film formation, changes in cell wall morphology could still be confirmed by the intact cells observed under the microscope (Figure 5.3a-d). A possible reason for formation of film around cell treated with ILs could be the release of polysaccharides from the encysted cell or by interaction of the IL with released polysaccharides. Ionic liquids are known to solubilise polysaccharides from lignocellulosic biomass [24, 25]. Polar ionic liquid with low viscosity and high hydrogen bonding capacity are important parameters in dissolution of polysaccharides consisting mainly of cellulose [26-28]. In this article we observed that the extent of permeabilisation varied for cells treated with imidazolium IL with different anions. Abe et al has shown that the anion influences the hydrogen bonding capacity and subsequently polysaccharide dissolution. The study also shows that Iolilyte 221 PG, TBP SO₄, and TBP TOS were less effective than imidazolium based ILs in permeabilising the cell wall which indicates that the cation also plays a role in solubilising the cell wall component. We could thus hypothesize that the permeabilisation of the cell wall is due to possible interaction between the cell wall polysaccharides and IL. Further detailed study on the cell wall composition together with its solubility in IL could explain the underlying principle. This study is beyond the scope of this article. Permeabilisation of algal cell wall using IL is a novel separation tool to obtain pigments from microalgal cell and could be extended to lipid extraction without compromising on fragile water soluble components like proteins [29].
Figure 5.5 Schematic of H. pluvialis cell wall (adapted from Hagen, C.S. et al. [22])

Reusability of Ionic Liquid

The reusability of the ionic liquid is imperative for scale up. To be able to determine this, reusability studies were performed by using the same IL solution for pre-treating four sets of fresh H. Pluvialis cells. The efficiency of the IL was tested by measuring the ability of the IL to permeabilise the cell and thus release of astaxanthin. The amount of astaxanthin that could be extracted using re-used IL was comparable with the control (see Figure 5.6). The IL could be used three times without any treatment and without affecting the efficiency of permeabilisation. A decrease in the efficiency of permeabilisation was observed when used for the fourth time to permeabilise the cells. The ionic liquid layer was tested for presence of pigment after each reuse and it was observed that the amount of pigment in the IL phase after three reuse cycles was negligible (~0.5 µg/ml i.e. 1.68% of total astaxanthin extracted after 4 extractions). The 1H NMR spectra of IL before and after 3 times reuse were comparable (see Figure 5.7).

Figure 5.6 Reusability of Ionic liquid
Conclusions

This paper highlights the potential of using ILs to permeabilise *H. pluvialis*. The haematococcal cyst has a robust cell wall and requires mechanical disruption to extract astaxanthin. The data clearly show that *H. pluvialis* cells are permeabilised by hydrophilic ILs under mild conditions of temperature (below 100°C) and atmospheric pressure. After pre-treatment with IL, astaxanthin is subsequently extracted using ethyl acetate with high efficiency while the cell wall is kept intact. Amongst the different ILs studied, astaxanthin extraction efficiencies obtained with EMIM DBP were the highest (more than 70%). The study shows that the cells can be permeabilised with 40% w/w IL with negligible loss of pigment to the IL phase. In addition the ionic liquid could be used three times without affecting its efficiency to permeabilise the cells. This opens up the possibility of reusing the expensive ILs. Knowing that energy demand drives the overall production cost [13] mechanical cell disruption is the most energy consuming step, and replacing with ionic liquid would have an impact on the overall cost. However, for exact estimations, ionic liquid process should be performed using industrial scale equipment. While the initial findings are promising, further research is necessary to optimise the process parameters in terms of extraction efficiency, use of wet biomass, solid to liquid ratio and to extend the application to other microalgae. A better understanding of the exact mechanism of action would be of great benefit in this. This paper underlines the potential of using ILs in extraction of hydrophobic biomolecules (e.g. pigments) under mild conditions as a replacement of mechanical techniques or harsh conditions. This allows the co-extraction of several classes of hydrophobic biomolecules for different applications from the same cell substrate, whereas the more hydrophilic biomolecules (e.g. proteins) are fractionated more efficiently after cell disruption without compromising the functionality. In this way cell can truly become multi-purpose factories.
Acknowledgements

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References


Chapter 6

Novel selective fractionation technology for fragile biomolecules from *Neochloris oleoabundans* using ionic liquids

To be submitted as:

*Rupali K. Desai, Maria Salvador Fernandez
Mathieu Streefland, René H. Wijffels and Michel H.M. Eppink*

Novel selective fractionation technology for fragile biomolecules from *Neochloris oleoabundans* using ionic liquids
Abstract

Microalgae are a promising source for proteins, lipids and carbohydrates for the food/feed and biofuel industry. To make microalgae production economically feasible it is necessary to optimally use all produced compounds keeping full functionality. Therefore biorefining of microalgae is the key to lower the cost of algal products using mild and effective processing techniques. In this article we have tested the feasibility of few aqueous solutions of imidazolium and phosphonium ionic liquids to selectively fractionate *N. oleoabundans* biomass into a hydrophilic fraction (proteins, carbohydrates) and a hydrophobic fraction (lipids). The results showed that Tributylmethylphosphonium methyl sulfate (Cyphos 108) could permeabilise fresh intact cells of *N. oleoabundans* to yield 68% of total lipids and obtain after cell disruption 80% of total proteins and 77% of total carbohydrates. Moreover, the proteins so obtained kept their native form explaining the novelty of this concept. A perspective on ionic liquid based microalgal biorefinery is also discussed.

Broader Context

Growing energy demands and depleting fossil fuels together with the impact of using these fossil resources on the environment drives the need to develop alternative source for fuel/energy production. Although, microalgae are considered as promising feedstock for biofuel production, the primary bottleneck lies in high cost of production. Apart from lipids, the microalgae also contain other value added components such as carbohydrates and proteins. Thus, using a biorefinery approach would help circumvent the problem of high cost. While the current processes use organic solvent for extraction of proteins from microalgae, use of solvent denatures the protein. In this article we show the feasibility of selectively separating the hydrophobic (lipids) from hydrophilic (proteins, carbohydrates) using ionic liquids without affecting the protein stability. The possible role of ionic liquids in fractionating microalgal biomass could pave way towards development of microalgal biorefinery.
Introduction

Microalgae are promising feedstocks for biofuel production. These photosynthetic microorganisms have high lipid productivity and do not compete for arable land when compared to terrestrial oleaginous crops [1]. Microalgae have a very tough cell wall and thus require energy intensive unit operations to break open the cell and release the intracellular content. Thus, despite the high lipid productivity, the energy input to separate the lipids is much higher than the energy obtained from the biomass indicating the necessity to use less energy intensive unit operations. Apart from lipids, microalgae are also good sources for proteins, carbohydrates and pigments. Utilization of these value added co-products for food, cosmetics, health and chemicals would help in making the process economically feasible [2].

The current processes are mainly focussed on recovery of single component from microalgae mainly lipids for biodiesel production [3]. Most commonly organic solvents are used for extraction of lipids. The Soxhlet [4] method uses hexane as a solvent and the Bligh and Dyer’s [5] method uses chloroform and methanol mixture as a solvent for extraction. As these processes are designed to extract one component, it leaves the biomass unsuitable for recovering other components. Additionally, lipids can also be extracted using sub and supercritical fluids [6], however these methods have high energy requirements and thus impact the overall economics of the process. In a study done by Ursu et al. protein extraction was done using alkaline condition. However, the proteins precipitated under alkaline condition had lower functional properties [7]. It is thus prudent to develop a mild process to fractionate the biomass into its components such that their functional integrity is retained.

Conventional extraction processes based on volatile organic solvents poses safety concerns and is toxic. It is thus necessary to develop newer methods to address these issues. Some of these methods include use of supercritical fluids and recently ionic liquids (IL)

Ionic liquids are salts which are liquid at temperatures below 100°C. They are composed of cations and anions and have negligible vapour pressure. They are known as designer solvents as their properties such as polarity, viscosity can be tailored by using a different combination of cation and anion [8]. This makes IL an attractive solvent for liquid-liquid extraction. ILs were used for lipid extraction from microalgae at elevated temperatures and together with co-solvents such as methanol [9, 10]. Studies using mixtures of ILs have also been performed to extract lipids from algae biomass [11] and dissolution of microalgae in ILs were also demonstrated [12]. Teixeira in his studies have shown energy efficient deconstruction of algae biomass by dissolution and hydrolysis of microalgae in ILs at temperatures above 100°C [13]. Olkiewicz, M., et al.[14] showed ~75% lipid and 93% FAMEs recovery using hydrated phosphonium IL under ambient temperature condition. All these studies have established the potential of ILs to extract lipids from microalgae with high efficiency. While, both Teixeira and Olkiewicz, M., et al have qualitatively demonstrated that all components of microalgae (lipids, proteins and carbohydrates) can be recovered in one process after hydrolysing the microalgae, it does not give any indication about
recovery of proteins and carbohydrates and the stability of the more fragile proteins. Most of these studies address the recovery and extraction efficiency of lipids from microalgae while recovery of other components such as the high value proteins and carbohydrates are not addressed showing the single component isolation strategy. A recent study by Xinhai yu et al. [15] reported energy efficient extraction of lipids from *Chlorella vulgaris* using IL combined with CO₂ capture. The study showed ~75 % lipid (~89% FAMEs) recovery but the proteins were denatured in the process. To be able to recover all components in their full functional state from microalgae biomass it is necessary to use mild techniques. Most of the articles discussed above are focused on lipid extraction from microalgae and not on biorefining of biomass. Additionally, though ILs has been used in the studies the conditions such as temperature are high enough to degrade the protein.

The primary objective of the article is to fractionate the algal biomass into a hydrophilic fraction (proteins, carbohydrates) and a hydrophobic fraction (lipids) using mild pre-treatment with an aqueous solution of ionic liquid. Additionally, the ability of ILs to extract lipids from the cells under different temperature and concentration conditions was studied. The microalgae strain used in this study is *N. oleoabundans* which is a high lipid producing strain [16]. Both fresh and dried algae were studied to understand the influence of ionic liquid pre-treatment on extraction efficiency of individual components and the stability of proteins.

**Materials**

The ILs used in this study was ≥ 95% pure and used without further purification. All the ILs listed in Table 1 was purchased from Iolitec. Chemicals and organic solvents used in the study, ethyl acetate, hexane, methanol sulphuric acid, phenol and fatty acid standards were bought from Sigma.

**Methods**

*Microalgae cultivation*

*N. Oleoabundans* was cultivated in the laboratory in fresh water medium as described by Breuer et al. [17] and the algae were stressed to have a higher lipid content. The microalgae were then harvested by centrifugation (4000 rpm for 10 minutes). The microalgae were freeze dried and used for extraction studies. For the study, using fresh cells the algae were grown at the AlgaeParc pilot facility, Wageningen, The Netherlands. The cell suspension was centrifuged at 4,000 rpm for 10 minutes and used for the study.

*Pre-treatment with Ionic liquid and fractionation of Biomass*

As shown in Figure 6.1 two studies were performed, lipid extraction efficiency of IL from intact microalgae cells at different temperature and IL concentration (A) and IL pre-treatment of microalgae and subsequent fractionation into hydrophilic and hydrophobic component (B).
N. oleoabundans cells ~10 mg of cells (freeze dried and/or fresh cells) were treated with 1.5 ml of aqueous solution of IL under mild temperature conditions for 30 minutes. The studies were conducted with 7 different ILs. The ionic liquids used in the study are listed in Table 6.1. The influence of temperature (35, 45, and 60)°C and concentrations of ILs (40, 80)% w/w on extraction efficiency of lipids was investigated. The amount of lipids extracted in the IL phase was determined by measuring the residual amount of lipids remaining in the cells.

Additional studies were conducted using two ILs, BMIM DBP and TBP SO₄. The cells were pre-treated with 1.5 ml of 40% aqueous solution of IL at 45°C for 30 minutes. The cells, after pre-treatment, were separated from ionic liquids by centrifugation at 3000 rpm for 10 minutes and then contacted with 3 ml of solvent (ethyl acetate/hexane) for 2 hours. The hydrophobic components from microalgae (lipids and pigments) were extracted in the solvent phase. The biomass remaining after pre-treatment containing the hydrophilic components mainly proteins and carbohydrates is suspended in phosphate buffer and then this fraction was analysed for protein and carbohydrate content.

**Table 6.1 Ionic liquids used in the study**

<table>
<thead>
<tr>
<th>Sr. No.</th>
<th>Ionic Liquid Names</th>
<th>Abbreviations</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Iolilyte 221 PG</td>
<td>IO</td>
</tr>
<tr>
<td>2</td>
<td>Tributylmethylphosphonium methyl sulfate (Cyphos 108)</td>
<td>TBP SO₄</td>
</tr>
<tr>
<td>3</td>
<td>Triisobutylmethylphosphonium tosylate (Cyphos 106)</td>
<td>TBP TOS</td>
</tr>
<tr>
<td>4</td>
<td>1-Butyl-3-methylimidazolium dibutylphosphate</td>
<td>BMIM DBP</td>
</tr>
<tr>
<td>5</td>
<td>1-Ethyl-3-methylimidazolium dibutylphosphate</td>
<td>EMIM DBP</td>
</tr>
<tr>
<td>6</td>
<td>1-Butyl-3-methylimidazolium acetate</td>
<td>BMIM Acetate</td>
</tr>
<tr>
<td>7</td>
<td>1-Butyl-3-methylimidazolium Dicyanamide</td>
<td>BMIM DCA</td>
</tr>
</tbody>
</table>
**Fresh/ Freeze dried cells**

**IL Lipid Extraction Efficiency**
- Temperature study 35°C, 45°C, 60°C
- Concentration study 40% w/w, 80% w/w IL

**IL Pre-treatment 45°C 30min**
- Add Solvent (Ethyl acetate/Hexane)
- Separate IL

**Cells resuspended in Buffer**
- Bead beating cells
- Hydrophilic fraction (Proteins and Carbohydrates)

**Solvent Phase**
- Evaporate
- Hydrophobic fraction (Lipids, pigments)

**Separate IL**
- Residual biomass after IL pre-treatment

**CHCl3 - MeOH extraction (Total fatty acids)**

---

**Figure 6.1** Experimental scheme, A) IL lipid extraction efficiency B) IL Pre-treatment and separation of hydrophobic and hydrophilic components

**Fatty acid determination**

The total fatty acids (FA) present in the microalgae is determined by treating the cells with CHCl₃-MeOH as described by [18]. For temperature and concentration studies total amount of fatty acids extracted in the IL phase is determined by measuring residual amount of fatty acids remaining in the cells after treatment with IL and is expressed as follows:

\[
\text{Total FA extracted in IL per mg of biomass} = \text{Total FA in control sample} - \text{Total residual FA in the cells after pretreatment}
\]

For calculating the total FA content of IL and solvent treated cells, the solvent phase (ethyl acetate/hexane) is evaporated under N₂ stream and the residue was analysed for total FA content after transesterification. The samples were analysed in the GC (Agilent 7890A) and the run time was 30 minutes.

**Protein Analysis**

**Protein content**

Protein content was determined with a commercial assay kit (DC™ Protein assay, Bio-Rad, U.S.) using bovine serum albumin (Sigma–Aldrich A7030) as protein standard. The microplate assay protocol was used and the absorbance was measured at 750 nm using a microplate reader (Infinite M200, Switzerland). The cells after pre-treatment with IL were suspended in 1 ml of lysis buffer, 60 mM Tris and 2% SDS, pH 9 in lysing matrix D tubes (6913–500, MP Biomedicals Europe). The sample was bead
beated for 3 cycles of 60s at 6500 RPM with a pause of 120 s between each cycle (Precellys 24, Bertin Technologies). The cell suspension was than heated at 100°C for 30 minutes. The cells were separated by centrifugation and the supernatant was analysed for protein content using the DC™ Protein assay. The protein is expressed as the % of total protein in the cells:

\[
\text{% of Total protein} = \left( \frac{\text{Total protein after pretreatment}}{\text{Total protein in cell}} \right) \times 100
\]

**Gel electrophoresis**

The stability of the proteins after pre-treatment with IL was confirmed by native gel electrophoresis. The cells after pre-treatment are suspended in 50 mM Phosphate buffer and disrupted by bead beating. The supernatant was diluted 1:1 with native sample buffer. The diluted sample ~25 μl was then applied on 4–20% Criterion TGX, Tris glycine precast gel and run with 10X Tris glycine native buffer at 125 V for 75 minutes. The gel was stained with Pierce™ Silver Stain Kit. The material for electrophoresis was bought from Biorad and staining kit was purchased from Thermo Fisher Scientific.

**Carbohydrate analysis**

The total carbohydrate content was determined by acid hydrolysis of IL pre-treated cells, adapted from [19]. The IL- pre-treated cells were suspended in water such that the final cell concentration is 1 mg/ml. To 50 μl of this suspension 450 μl water, 500 μl of 5% phenol solution and 2.5 ml of concentrated sulphuric acid was added. The mixture is incubated at room temperature for 10 minutes and then at 35°C in a water bath for 30 minutes. The carbohydrates react with acidic phenol to give yellow orange colour which is then measured at 483 nm using UV spectrophotometer (Beckman). For control process the cells were directly suspended in water without any pre-treatment. Starch samples were measured as positive controls. The calibration curve was prepared using glucose as the standard. The total carbohydrate content is expressed as:

\[
\text{% Total carbohydrate content} = \left( \frac{\text{Total carbohydrate content after pretreatment}}{\text{Total carbohydrate content in the cell}} \right) \times 100
\]

**Results and Discussion**

Extraction efficiency of different components (e.g. proteins, carbohydrates, lipids) after IL pre-treatment is studied and the protein stability determined by electrophoresis. Aqueous solution of three ionic liquid classes including imidazolium, ammonium and phosphonium ILs were tested at different temperatures and concentration.

**IL lipid extraction efficiency**

Lipid extraction efficiency of aqueous IL solutions after IL pre-treatment at different temperatures (35, 45, 60)°C and concentrations (40, 80)% w/w were studied. In this study lipids represent the fatty acid methyl ester content that could be converted to biodiesel. Three classes of ILs (see table 6.1) were studied. The results show that at 80% w/w concentration of IL, as the temperature increases, the
amount of lipid extracted increases for all ILs (see Figure 6.2). Lipid extraction capacity was higher for BMIM ACE, BMIM DBP, EMIM DBP and TBP TOS, ~8-12% /mg of biomass than for IO, TBPSO₄ and BMIM DCA. Although at 60°C, lipid extraction efficiency was higher, 45°C was selected for concentration studies to maintain mild conditions. Two ILs were selected, BMIM DBP which showed highest extraction efficiency among the ILs tested and TBPSO₄ which showed the least extraction efficiency but could have an impact on the cell wall. The effect of IL concentration on extraction efficiency was tested for BMIM DBP and TBPSO₄ (see Figure 6.3). As the concentration of IL increases from 40% w/w to 80% w/w at 45°C, the amount of lipid extracted increases from 2.61% to 9.89% per mg of biomass for BMIM DBP and from 1.28% to 3.27% per mg of biomass for TBP SO₄. This increase in extraction capacity could be attributed to the hydrophobicity increase of the IL solution.

IL solutions under mild conditions were able to extract lipids from intact microalgae cells; the maximum amount extracted was ~42% of the total fatty acid present in the cells. Based on the results in Figure 6.2 and 6.3, BMIM DBP could permeabilise the cells and extract lipids better than TBP SO₄ indicating that the cation and anion influences the extraction efficiency albeit to a different degree. The results confirm that the lipid solubility in aqueous solution of ILs is low. In other studies by different authors [10, 11, 13, 14] wherein the IL pre-treatment was done at temperatures close to 100°C, lipids released were extracted with organic solvent and extraction efficiency was > 90% of the total fatty acid content. Thus, indicating that temperature is indeed an important factor influencing the extraction efficiency.

![Figure 6.2](image-url)  
**Figure 6.2** Effect of temperature on extraction of lipids using 80% w/w aqueous solution of IL
Figure 6.3 Effect of IL concentration on extraction of lipids at 45°C

IL pre-treatment and extraction with organic solvent

The above studies showed that aqueous solutions could permeabilise the cells as well as extract the lipids without cell disruption. This observation is in accordance with our previously published studies [20] that aqueous IL solutions could permeabilise the intact microalgae cells under mild conditions and release the intracellular hydrophobic components pigments and lipids in this study. Microalgae biomass also contains large amount of proteins and carbohydrates besides lipids. To recover these components in their native form after biomass pre-treatment with ILs, additional studies were performed. Pre-treatment studies were done on both wet and dried N. oleoabundans cells using 40% w/w BMIM DBP and TBP SO4. Despite low lipid extraction efficiency TBP SO4 was selected for further studies to understand the influence of the IL on cell wall permeabilization. Hydrophobic components were subsequently extracted with ethyl acetate and then the cells were mechanically disrupted to recover hydrophilic components. The amount of fatty acid extracted after pre-treatment was compared with Bligh and Dreyer method. The amount of lipid extracted from fresh biomass after pre-treatment was 11.39% and 17.66% for BMIM DBP and TBP SO4 compared to 25.96% using the Bligh and Dreyer method. For the freeze dried biomass, the amount of lipid extracted after pre-treatment was 10.48% and 15.91% for BMIM DBP and TBP SO4 compared to 18.20% using the Bligh and Dreyer method. The results (see Figure 6.4) show that lipid recovery was better with TBP SO4 in comparison to BMIM DBP for both wet and dried cells. This shows that TBP SO4 which has a low lipid extraction capacity, even at 80% w/w concentration, is able to permeabilise the cells and lipids can be subsequently extracted with ethyl acetate. The higher recovery with freeze dried cells could be attributed to effect of drying on the cell walls. The results also show that ethyl acetate alone is not able to permeabilise the cells and extract the lipids. Additionally, studies using hexane instead of ethyl acetate for extracting lipids after IL pre-treatment were performed (not shown). The results showed that no lipids were extracted in the hexane phase. This indicates a possible cooperative role of ethyl acetate together with IL in permeabilising the cell wall.
The hydrophilic components, proteins and carbohydrates after lipid extraction are recovered by cell disruption. The percentage of total protein recovered after pre-treatment using BMIM DBP and TBP SO₄ was 76.81% and 80.29% for fresh cells and for freeze dried cells 33.86% and 62.45% respectively (see Figure 6.5) were observed. The decrease in protein recovery for freeze dried cells could be due to direct contact of IL with proteins in the already compromised cell wall. In a separate study aqueous solution of BMIM DBP (results not shown) and TBP SO₄ [21] in contact with Rubisco (Ribulose-1,5-biphosphate Carboxylase/Oxygenase) causes aggregation/precipitation of the protein molecule. These results thus indicates that TBP SO₄ effectively permeabilises the cell wall such that proteins remain intact inside the cell and can be recovered in their functional state by further cell disruption. The proteins recovered after extraction of lipids were run on native gel and detected using silver stain (see Figure 6.7). Although microalgae contains other proteins, Rubisco is used as the known biomarker protein for microalgae. The native gels shows that Rubisco remains intact and is not dissociated into its subunits indicating that proteins recovered after IL pre-treatment retains it native form. Additionally, the aqueous phase after cell disruption was analysed for carbohydrate content. The percentage of total carbohydrate recovered after pre-treatment using BMIM DBP and TBP SO₄ was 49.06% and 77.10% for fresh cells and 74.62% and 64.81% respectively for freeze dried cells (see Figure 6.6). The results thus show that the microalgae components lipids, proteins and carbohydrates can be selectively fractionated after IL pre-treatment and whereby the proteins retain their full functional composition.

**Table 6.2 Summary of biomass components separated under different IL pre-treatment**

<table>
<thead>
<tr>
<th></th>
<th>Fresh Cells</th>
<th>Freeze dried Cells</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Control</td>
<td>40% BMIM DBP</td>
</tr>
<tr>
<td>Total Fatty Acid/mg of biomass</td>
<td>25.96</td>
<td>11.39</td>
</tr>
</tbody>
</table>
**Figure 6.4** Total lipids extracted with ethyl acetate after IL pre-treatment at 45°C

**Figure 6.5** Total Proteins in the biomass after IL pre-treatment at 45°C

**Figure 6.6** Total carbohydrates in biomass after IL pre-treatment at 45°C
Perspective on IL based microalgal biorefinery

Although microalgae have potential for biodiesel production, the high cost of producing oil is the key bottleneck in its commercialization. Therefore, adoption of an integrated biorefinery approach wherein different products (biofuel, nutraceutical, speciality chemicals) are produced at large scale could improve the economics of oil production. As discussed earlier, current processes are directed towards single product recovery and use of solvents and high temperatures results in denaturation of proteins.

So far studies on the use of ILs in microalgae are focussed on lipid extraction by either hydrolysis of whole cells in IL at high temperature or by use of co-solvents. There are two main approaches to use ILs in microalgal biorefinery mainly:

1. Hydrolysis
2. Pre-treatment

Hydrolysis of microalgae in IL results in separation of oil which could be separated but studies should be done on recovering the proteins and carbohydrates dissolved in IL. Although theoretically dissolved components could be separated using techniques such as chromatography, there is no empirical data to prove it. Hydrolysis of biomass offers the advantage of lower waste generation and reduced number of processing steps. The proteins recovered should also be tested for it functionality as it could be affected by the ILs.

On the other hand pre-treatment studies as described in this article show the novelty of separating lipids without mechanical disruption and subsequent separation of hydrophilic components (proteins,
carbohydrates) in their native form after cell disruption. The process can be optimised further to improve the yields. There are various parameters which influence the efficacy such as biomass loading, time of contact with IL and organic solvent, amount of solvent added and type of IL and should be investigated in detail.

While ILs indeed has a potential role to play in microalgal biorefinery, it can only be realised if they are biocompatible, biodegradable and economical. The ILs must be tested for their reusability and recyclability so as to make the process economically viable. To be able to judge a process superior than other would require a systematic approach and set of criteria on basis of which the process is evaluated. Most of these studies are at their infancy and should be evaluated in terms of energy consumption, efficacy and cost. It is also necessary to have set controls with which it can be compared to.

**Conclusion**

In this article, pre-treatment of *N. oleoabundans* using ILs and subsequent fractionation into hydrophilic and hydrophobic components was studied for both fresh and freeze dried biomass. Additionally, the lipid extraction efficiency of aqueous IL solution under different temperature and concentration conditions were studied. We have demonstrated that aqueous solution of imidazolium and phosphonium based ILs was able to extract lipids from intact microalgae, albeit to a different degree. We have also shown that pre-treatment of microalgae with BMIM DBP and TBP SO₄ at low concentration (40% w/w) results in permeabilisation of cells. The biomass can then be fractionated into hydrophilic and hydrophobic components whereby the proteins were recovered without losing their nativity. The recovery of total fatty acids was ~68% and that of proteins and carbohydrates was ~ 80% and 77% respectively of the total amount present in the cells, after pre- treatment of fresh biomass with TBP SO₄. Most of the current processes that use energy consuming mechanical cell disruption (e.g. bead milling, high pressure homogenisation) [22] and solvents such as methanol/chloroform and hexane [3] are able to recover only lipids and render the proteins unsuitable for use due to denaturation. This article is a step forward in establishing the role of ILs in microalgae biorefinery by developing a novel selective fractionation concept for both hydrophobic compounds (e.g. lipids) and hydrophilic compounds (e.g. proteins, carbohydrates). However, more studies are needed to optimise the process in terms of yields and recycling of ILs. The observations made in this study shows significant potentials in terms of biorefining the algal biomass.

*Note: Fresh and freeze dried cells from different batches. For proteins and carbohydrates average of 2 values are presented and also for temperature and concentration studies values presented is an average of 2.*
References

Chapter 7

General Discussion
Abstract

In the preceding chapters we have shown the feasibility of using IL and IL based systems for extraction of fragile biomolecules from microalgal biomass. We developed two different fractionation techniques; 1) Using ionic liquids and solvents to extract pigments/lipids from intact cells and then further disrupting the cells to obtain carbohydrates and proteins in their native form 2) Using IL based emulsion stabilised by core shell particles to separate hydrophobic from hydrophilic components after cell disruption and keeping the proteins in its native form. There are some challenges in using IL based system such as, recovery from ionic liquid, biodegradation of IL and an outlook on future separations using IL based systems are discussed in this chapter.
Introduction

The need to develop more sustainable and environmentally safe processes drives the development of newer separation and extraction technologies. In addition there is a constant drive to develop energy efficient and cost effective processes. Although the separation technology is product and raw material specific, for most biochemical processes it is possible to develop separation technology principles which could then be tailored to specific needs. ILs has gained considerable attention in the past decade. The role of IL as medium for catalysis [1, 2], in extraction and separation [3, 4], and dissolution of biomaterials [5] has been well demonstrated, the efficiency of IL based systems in extraction of fragile biomolecules such as proteins is scarcely studied. The research in this thesis thus focuses on novel IL based systems for extraction of biomolecules (proteins, pigments, lipids and carbohydrates) from microalgae.

Biomass feedstock such as microalgae, apart from being a potential source for biofuels also contains other value added components such as proteins and carbohydrates. The recovery of these components is mostly based on solid-liquid extraction processes. These processes such as one employing volatile organic solvents aim at recovering only one type of biomolecule i.e. hydrophobic lipids. The spent biomass which is a rich source of hydrophilic components, mainly proteins and carbohydrates after removal of lipids is rendered unsuitable for feed and food application. Furthermore, the proteins are denatured by use of organic solvents. This necessitates the need to develop mild separation techniques for complete utilization of biomass. Interest in separation processes based on ILs has gained interest in past couple of years due to higher extraction yields of bioactives from biomass under moderate conditions of temperature and pressure [6]. Thus, the aim of this thesis was to study the feasibility of using novel IL based systems for separation of multiple (fragile) biomolecules and pre-treatment of microalgal biomass. In this chapter a summary of the outcomes of the feasibility studies and bottlenecks obtained in the study are discussed. Additionally an outlook on the future separation studies is presented.

Protein Stability

One of the prime requirements to use IL based systems for separation of proteins from microalgae or other feedstocks is to determine its stability in ionic liquids. In chapter 3, we studied the stability of three proteins (Rubisco, IgG, and BSA) of varying nature in two ILs (Iolityte 221 PG and Cyphos 108). It was observed that the proteins form aggregates at higher concentration of ILs, however, the extent to which the aggregates are formed varied depending on the type of protein. We also observed an unexplained behaviour of BSA in Cyphos 108, wherein as the concentration of Cyphos 108 increased to 20% the BSA peak intensity (see chapter 3, Figure 3.9) decreased and when the concentration was increased to 50% an increase in the BSA peak intensity was observed together with the formation of aggregates (chapter 3). Thus, indicating that the molecular interactions between BSA and Cyphos 108
Chapter 7

promote this behaviour. The number of studies on protein stability in ILs is limited to model proteins, such as BSA, lysozyme etc. Due to the diverse nature of proteins and ILs it is indeed difficult to state general guidelines for protein stability in ILs. However, based on our studies and together with the available literature [7] it could indeed be stated that high concentrations of ILs affect the structural integrity of the proteins such as promoting protein unfolding or induce the formation of aggregates. Additionally, the protein molecular weight too determines the IL threshold concentration. As seen in chapter 3, BSA which is a small sized molecular weight protein (~66 kDa) is stable up to 50% concentration of Iolilyte 221 PG whereas Rubisco, a high molecular weight protein (~540 kDa), forms aggregates at 25% concentration of Iolilyte 221 PG.

Studies on proteins of commercial interest are limited. As a step forward in this direction we looked into the stability of IgG and Rubisco in imidazolium based ILs. The stability of IgG and Rubisco was tested in different concentrations (10% v/v to 40% v/v) of 1-ethyl 3-methyl imidazolium dibutyl phosphate (EMIM DBP). A trend similar to the ILs studied in chapter 3 was observed, higher concentration causes aggregation of IgG and Rubisco. In Figure 7.1 a-b, the stability of IgG in EMIM DBP is shown using high performance size exclusion chromatography (SEC-HPLC). SEC-HPLC studies showed that IgG was stable at 40% IL concentration. Qualitative studies revealed better stability of IgG when compared to Rubisco in imidazolium ILs. As also mentioned in chapter 3, the difference in stability of protein in ILs could be due to difference in size, structure, folding and connections of the subunits of the two molecules, Rubisco ~540 kDa and IgG 150 kDa. Structurally IgG has 2 large and 2 small subunits which are covalently linked unlike Rubisco which has non-covalently linked 8 large and 8 small subunits. We also tested the recovery of IgG from EMIM DBP by using desalting columns and the fractions collected showed an IgG band on native gel. Ionic liquids being salt could be separated from proteins using desalting columns. The proteins being large molecules elutes first and then the IL. However the study is more qualitative and a quantitative study must be done to determine the precise recovery of the protein from IL. While these are preliminary studies, future work should be focussed on recovery of proteins from ILs and investigate the protein-IL interactions explaining the molecular behaviour between ILs and proteins. Recovery of components dissolved in ILs is an important aspect and demands more research on this topic.
In this thesis we studied two novel IL based systems for separation of biomolecules:

1. IL based Aqueous Biphasic System (ABS)
2. IL based emulsion stabilised by microgel particles

As discussed in chapter 2 and 3, IL based aqueous biphasic system are novel systems and have advantage over traditional aqueous biphasic system in terms of better extraction efficiency and selectivity. In chapter 3, we studied the extraction of Rubisco using IL based ABS. We observed higher partition coefficient for Rubisco, a major microalgal protein, in the IL rich phase in comparison to the conventional PEG based ABS. Despite high partitioning, the higher concentration of IL in IL rich phase led to protein instability. As mentioned earlier, protein stability in IL plays an important role in separation processes using IL based system. Rubisco stability studies revealed that when the concentration of IL used, Ioliyte 221 PG, was ~10% v/v then it retains it nativity. Designing ABS such that the concentration of IL in the IL rich phase is ~10% v/v, would help to stabilise the protein. Higher concentration of IL leads to formation of aggregates. This directs towards more systematic studies on phase diagrams which would help in designing ABS system with desired concentrations of phase components.

*Figure 7.1 Effect of increasing concentration of ionic liquid-EMIM DBP on a) IgG and b) Rubisco*
While IL based ABS are widely studied for extraction of biomolecules such as alkaloids, antioxidants [8] using model systems, their potential for extracting biomolecules like antibiotics from complex matrices such as fermentation broth is also established [9]. Additionally, these systems offer the possibility to tailor the polarities of the phases in order to improve efficiency and selectivity [10]. This ability of IL based ABS holds huge prospect in extracting fragile biomolecules from microalgal biomass which contains molecules of different polarity. No work has been done on separating microalgal proteins using IL based ABS except, the studies described in this thesis. Although the feasibility studies show aggregation of Rubisco at concentration studied, additional work is required before this technique is rendered suitable for microalgal protein separation. More extensive studies using different ILs and careful design of IL based ABS should be performed using microalgal biomass.

We also investigated another novel system based on ILs, which is an emulsion stabilised by microgel particles for its feasibility to separate fragile biomolecules of varied polarities simultaneously from microalgae (chapter 4). The emulsion is IL in water type, wherein a hydrophobic IL, cyphos 109, forms the discontinuous phase and water forms the continuous phase. With mixing microalgae biomass (cracked cells), the proteins remain in the aqueous phase and pigment, astaxanthin, is extracted in the hydrophobic core. The interesting part is that the proteins retains its nativity. The microgel particles used to stabilise the emulsion are porous hence allows the transfer of small molecules and prevents direct contact of proteins with the IL, thus assists in retaining nativity of the protein. In chapter 4, we presented a proof of concept for a novel continuous separation. The microgel particles are thermoresponsive, a property of emulsion which was unexplored in this work. As the temperature is raised above 32°C the particles collapse and the emulsion breaks. This property of the emulsion should be explored in the separation process in the near future. The technique is still at its infancy and should be tested with different ILs and smaller IL droplets so as to increase mass transfer. Additionally, although the emulsion facilitates simultaneous separation of components from biomass, the process still needs to be optimised in terms of efficiency and reusability of IL and microgel particles. Future research should focus not only on optimization of process parameters but also on finding newer applications for this emulsion, which provides a complete toolbox for separation and extraction of value added components from different biomass streams.

Pre-treatment studies on microalgal biomass using IL

In addition to IL based systems, we also investigated the influence of IL solutions on microalgal cells and recovery of components thereof. Microalgae are known to have a tough cell wall and require mechanical disruption to be able to release the intracellular components. In chapter 5, we showed that a 40% aqueous solution of EMIM DBP, an imidazolium based IL, was able to permeabilise intact cells of *H. pluvialis* under mild condition and astaxanthin could subsequently be extracted with ethyl acetate. Microscopic study confirmed that the cells were not ruptured. Based on these observations we
postulated a possible mechanism wherein we stated permeabilization of the microalgal cells by ILs. However, it's unclear as to what solubilises in the IL to make the cell wall porous. A similar observation was also made when \textit{N. oleoabundans} cells were treated with IL (chapter 6). In addition to imidazolium IL, a phosphonium based IL, Cyphos 108, permeabilised the cell wall of \textit{N. oleoabundans}. However, this IL did not permeabilise the cell wall of \textit{H. Pluvialis}. This indicates the difference in cell wall composition of different algae. The ability of IL to permeabilise both fresh and freeze dried cells of \textit{N. oleoabundans} rules out the possible role freeze drying of cells on the cell wall permeability, although it would indeed aid in weakening the cell wall. We hypothesize that IL pre-treatment together with solvent ethylacetate partially dissolves some component of the cell wall thereby leaching the lipids/pigments from the intact cells.

In chapter 6, we went a step ahead and showed the feasibility of biomass fractionation. After cell wall permeabilization with IL and de-lipidation with ethyl acetate, the cells were mechanically disrupted to recover the proteins and carbohydrates. The proteins so recovered were in their native form. These results portrays the potential of ILs in fractionation and separation of microalgal biomass components in a biorefinery concept. While the early results are promising, we need more in depth knowledge and understanding of the effect of IL on cell walls. The most logical approach would be to study the cell wall of microalgae and its composition. Together with this we need advanced analytics to analyse components dissolved in IL. These studies would help in selecting and designing a suitable IL for different microalgae.

\textbf{Key outcomes of the thesis}

\textit{Table 7.1 A summary of the main findings of the thesis}

<table>
<thead>
<tr>
<th>Main Finding</th>
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<tr>
<td>• Rubisco is stable at low concentration (~10%) of Ionic liquids</td>
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<tr>
<td>• Extraction of pigments/lipids from intact microalgal cells using ionic liquids and solvents and further fractionation into native proteins and carbohydrates by cell disruption</td>
</tr>
<tr>
<td>• Separation of hydrophobic from hydrophilic components using IL based emulsion stabilised by microgel particles.</td>
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<tr>
<th>Bottleneck</th>
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<tr>
<td>• Analysis of dissolved component in ILs</td>
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<tr>
<td>• Cost of IL</td>
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<tr>
<td>• Biodegradability of ILs</td>
</tr>
<tr>
<td>• Recovery of biomolecules from IL</td>
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\textbf{Ionic liquids in Separation process and Challenges}

ILs has gained popularity as unique solvents in different areas of separation techniques. Owing to tunable properties which can be selected by choosing appropriate cationic or anionic constituents, they
can be applied in chromatographic methods as mobile phase or stationary phase modifiers and electro-chromatographic methods such as capillary electrophoresis or in sample preparations processes as new extraction solvents [11]. Most of the IL based separation techniques are focused towards analytical separation and sample preparation [11]. ILs has been used successfully for the separation and extraction of a wide range of analytes (organic ions, inorganic ions, metal ions and organic compounds) with different properties. On the other hand, more targeted IL-based materials are needed for the extraction and separation of bioactive targets compounds from complex matrices such as plants and different biomass at preparative scale.

Despite the potential shown by ILs in pre-treatment and separation there are bottlenecks which needs to be overcome before successful use of ILs in large scale preparative applications. The cost of using ILs as extractants or for pre-treatment should be considered. The economic use of ILs could be achieved by recycling and reusing the ILs. ILs is considered green primarily due to their non-volatile nature. However, not all ILs can be considered green, as some of them are toxic and are non-biodegradable.

Conclusion and Outlook

In conclusion, this thesis describes two novel IL based systems for extraction of fragile molecules from complex matrices and also shows direction towards continuous processing. It also demonstrates the permeabilization of microalgal cell wall using ILs and subsequent fractionation of biomolecules therein. To develop more sustainable scalable processes, future research should focus on using biodegradable ILs such as cholinium based IL. While high extraction efficiency of biomolecules in ILs is achieved, recovery of biomolecules from ILs must be studied as ILs are non-volatile. The IL based emulsion studied in this thesis has potential and efforts should be made to find newer applications. IL based ABS should be assessed for extracting multiple biomolecules from different biomass streams (e.g. microalgal biomass).

Currently, IL based separation system despite being milder and high in yields have to compete with existing technologies, thus requiring the process to be cost effective. Nevertheless, they hold a good chance in microalgal biorefinery, wherein there is need to develop an efficient process to ensure complete fractionation of the biomass. The role of ILs in fractionation of microalgal biomass is shown in this thesis. With research ongoing within several research groups on reusing/recycling ILs and development of more biodegradable ILs, brings the processes based on these liquids closer to an ideal sustainable process which eventually will be the need as the world moves towards a biobased economy.
References


Summary
Summary

Liquid-liquid extraction (LLE) techniques are widely used in separation primarily due to ease of scale up. Conventional (LLE) systems based on organic solvents are not suitable for extraction of fragile molecules such as proteins as it would result in denaturation. On the other hand Aqueous Biphasic System (ABS) though suitable for extraction of proteins they are restricted by limited polarity range. Ionic liquids have gained interest in extraction over the past years due to its non-volatility and tunable properties. In this thesis we explored the feasibility of using two ionic liquid based systems for extraction: 1) Ionic liquid based aqueous two phase system for extraction microalgae protein and 2) ionic liquid based emulsion for separation of hydrophilic and hydrophobic components from complex biomass such as microalgae. Additionally the influence of IL pre-treatment on microalgae cell wall and subsequent fractionation of its component was also investigated.

In part 1 of the thesis we studied extraction of biomolecules using known and novel IL based system. In Chapter 2, a review on protein extraction using ionic liquid based aqueous two phase system has been discussed. The stability of proteins in ionic liquids and factors influencing the extraction of proteins using ABS are presented. A systematic approach towards use of ionic liquid based system for extraction of proteins together with the major bottlenecks in using such systems is discussed.

The stability of proteins in ionic liquids is crucial to use ionic liquid based extraction system for their extraction. Therefore, in Chapter 3 the stability of commercially important proteins such as Rubisco and IgG, and model protein such as bovine serum Albumin (BSA) was studied in various aqueous solutions of ionic liquids at different concentrations. We observed that protein properties such as size, number of subunits influence its stability in ionic liquid, however the general trend was that the proteins were found to be stable at low concentration of ionic liquids, ~10% w/w. The extraction efficiency of Rubisco in PEG based and ionic liquid based aqueous two phase systems were evaluated. The partition coefficient for Rubisco in ionic liquid rich phase was 3-4 fold higher than in PEG based two phase system.

In Chapter 4, another ionic liquid based system, an ionic liquid in water emulsion stabilised by microgel particles was studied for feasibility of separation of components in microalgal biomass. The system was evaluated for its efficiency to selectively separate hydrophilic components such as proteins from hydrophobic components such as pigments present in lysed microalgal biomass. The system could selectively separate the components and moreover the proteins so obtained retained its nativity. A proof of concept for continuous operation is also presented.

In part 2 of the thesis we explored the possibility of using ionic liquids for pre-treatment of microalgal biomass. Microalgae are known to have a tough cell wall and usually require energy intensive mechanical cell disruption to recover the intracellular components. In Chapter 5 the influence of aqueous solution of different ionic liquids on cell wall of *H. pluvialis* was studied. Aqueous solution of
the ionic liquid, 1-ethyl, 3-methyl imidazolium dibutyl phosphate was found to permeabilise the cell wall of Haematococcus pluvialis under mild condition of temperature below 50°C. The pigment, astaxanthin could then be recovered using ethyl acetate. We hypothesized that ionic liquid was capable of dissolving certain components such as mannan polymer thereby disrupting the polymer structure and forming tiny holes in the cell wall, thus paving the way for solvent to penetrate and extract astaxanthin. Using microscopy we showed that the cells remain intact after ionic liquid and solvent treatment and thus shows that ionic liquid permeabilises the microalgae cell wall.

While we showed that tough cell wall of H. pluvialis could be permeabilised with ionic liquid under mild condition, we extended the study to other microalgae, Neochloris oleoabundans in Chapter 6. Different ionic liquids were tested for their ability to permeabilise and subsequently extract all hydrophobic components (e.g. lipids) from the cell without affecting the functionality of fragile molecules such as proteins. The results showed that microalgae could be permeabilised with the ionic liquid, Tributylmethylphosphonium methyl sulfate in addition to 1-ethyl, 3-methyl imidazolium dibutyl phosphate used for H. Pluvialis, indicating possible difference in the cell wall composition of two microalgae strains. The lipids are extracted using ethyl acetate and the cells were further disrupted by mechanical cell disruption to separate the proteins and carbohydrates. The protein analysis showed that they were native. While these were all feasibility studies, the process can optimised to improve efficiency.

Finally in Chapter 7, we summarise and highlight the main finding in this thesis together with the implication of overall results of the thesis for future research is discussed. The results in this thesis highlight the potential of Ionic liquids and ionic liquids based extraction system for extraction of biomolecules from complex biomass such as microalgae.
Acknowledgement

Finally the most read part of the thesis, acknowledgement. PhD is not only about science but as the name suggest, it’s a Philosophy. There are times during your period of PhD when you are at the extremes of emotions. It’s then the people around you, who support and be a part of this journey. I want to take this opportunity to thank all those who have been a part of my PhD journey and make it memorable.

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About The Author

Rupali K. Desai was born on 10th July, 1981 in Mumbai, India. After completing her Junior college in Mumbai in 1999, she started her Bachelors in Pharmacy at Bharati Vidyapeeth College of Pharmacy, Mumbai University. In 2003, she started working as a trainee in a Pharmaceutical Company, Cipla Ltd. and at the same time started preparing for entrance exams for further studies. In 2004, she started her Masters in Bioprocess Technology from the Institute of Chemical technology, Mumbai. After completion of Masters, she was working as a Process development Scientist in Biocon, Bangalore, India. In 2010, she moved to the Netherlands. She was working for DSM in Delft before she joined Bioprocess Engineering group of Wageningen University on an STW project under the supervision of Prof. Dr M.H.M Eppink and Prof. Dr Rene Wijffels and Dr Mathieu Streefland. Since July 2016, she works as a Scientist, downstream processing, at uniQure, Amsterdam, The Netherlands.
List of Publications


# Overview of completed training activities

## Discipline specific

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<td>Advanced Downstream processing course</td>
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## General

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* Poster presentation
‡ Oral Presentation
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