Protein recovery from whisky by-products: a study of using ion exchange chromatography for the recovery of proteins from pot ale

by

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ABSTRACT

Liquid and solid by-products samples from malt whisky (MW), grain whisky (GW) and brewing (B) origin across several Scottish distilleries and breweries were collected and analysed for physical, chemical and nutritional properties. Nutritional properties assessed included protein quantification.

Among the by-products analysed, the focus in this work was placed on pot ale, the liquid by-product from MW processing. Approximately, 2-3 million tonnes of pot ale are generated in Scotland annually, with a protein content of ~1% protein (w/v) or 40% (w/w) on dry matter basis. Current technologies for the recovery of the protein from pot ale, i.e. evaporation, are expensive, require large amounts of energy and produce a low value product called pot ale syrup.

A less energy intensive method with the potential to create a higher value product from pot ale was developed in this work using an ion exchange chromatography (IEC) technique that exploits protein electric charge. Pot ale proteins were found to be positively charged (due to low pH) and cation exchangers were used to bind pot ale proteins. The method was tested and up-scaled from 50 ml to 1400 ml of pot ale at flow rates from 1 ml/min to 30ml/ min.

An economic analysis included in this work showed that using IEC for protein recovery from pot ale can be applied at commercial scale and the protein product used in higher value markets such as aquaculture.
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DEDICATION

To Keara, Lucie, Amy and my parents.
DECLARATION STATEMENT
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CHAPTER 1 – INTRODUCTION

1.1 Background

Brewing and distilling are important economic activities in Scotland, providing more than 10,000 jobs and generating over £3 billion in revenues annually. A high quality environment and raw materials are essential to these industries. Between 2010 and 2014, Heriot-Watt University carried out several projects to address sustainability issues of whisky production.

In this context, the Scottish Founding council (SFC) funded a three year project which started in September 2011. The project was named "Horizon Proteins, Fermentation process co-products: Integrated protein, energy and feedstock recovery".

The overall aim of the project was to design and implement a process to separate and recover protein from by-products which can then be used in aquafeed. The objective was to design an innovative process which had the potential to add-value to distillery by-products, provide a local and sustainable source of protein feed for salmon farmers in Scotland and recover protein for feed purposes which otherwise may be lost from the food chain. The focus was specifically on application of protein from pot ale as an ingredient in salmon feeds.

The vision was to develop a patented process and to have the technical know-how, people and industrial contacts in place after the three years of funding to ensure the commercialisation of the technology. As a first step in commercialisation, the team took part in the Converge Challenge 2013 and made it through to the final. The bid involved writing a comprehensive Business Plan and pitching to a 6-member judging panel.

Following on from this, the team received funding from Scottish Enterprise (SE) through the High Growth Spin-Out Programme (HGSP). This allowed Horizon Proteins to develop into a high-growth spin-out company, aiming revenues for £20 million within five years.
1.2 Thesis objectives

The thesis objectives were aligned with SFC-Horizon Proteins goals explained earlier. But, specifically, for the purpose of this PhD thesis, there were two main areas on which the research was focused:

- Development of a novel and sustainable process for the recovery of whisky pot ale proteins
- Assessment and development of ion exchange chromatography as a technique for protein concentration and separation from pot ale.

As an additional task, the economics behind traditional whisky by-products processing were investigated and compared with the process developed during this work.

It is important to highlight that this work concentrated on whisky pot ale rather than working on all whisky by-products. The degree to which the thesis objectives have been achieved were discussed in the Final Conclusions Chapter.

1.3 Thesis layout

The thesis contains twelve chapters, including the introduction, conclusion and references chapters. The second chapter consists of a literature review about whisky by-products. An understanding of the nutritional, chemical and physical properties of whisky by-products and potential uses for the proteins recovered were studied. This chapter provided the basis of the business plan for Horizon Proteins.

On chapters 3 to 10, more technical and scientific aspects were considered. Protein extraction methods and solid liquid separation studies were presented in Chapter 4 and Chapter 5, respectively. Between Chapter 6 and Chapter 10 the focus was on studying ion exchange chromatography as method for protein concentration and purification.

Finally, in the appendices, the overall process for protein recovery is described and additionally an economical evaluation analysis is included.
CHAPTER 2 - LITERATURE REVIEW

2.1 Whisky and whisky by-products

Whisky spirit uses either malted barley as the sole cereal substrate or a mixture of unmalted cereal grain together with sufficient malted barley to provide the enzymes to convert the cereal starch. It is important to distinguish between the two kinds of Scotch Whisky (i.e. malt and grain) and the cereals used, since the properties of the whisky and its by-products could be affected.

Scotch whisky ingredients for malt whisky production are malt barley, yeast and water. Nothing else is permitted by law. This law was defined in the UK in 1909 and recognised in European (EC) Legislation in 1989. Current UK and Scottish legislation related to Scotch Whisky is the Scotch Whisky Regulations (2009). The term co-products is also used interchangeably with by-products – by-products will be the preferred term in this thesis as it is the legal term used in the Scotch Whisky industry. By-products are clearly defined by European legislation under Article 5 of Directive 2008/98/EC (EU 2008).

A list of the brewing and distilling by-products and their definitions (Crawshaw 2001, Harper 2010) are shown below:

- **Pot ale**: residues from first distillation in malt whisky. Also known as burnt Ale.
- **Spent Wash**: pot ale equivalent from Grain distilling.
- **Draff**: grain solids left after starch and enzyme extraction. Sometimes referred as distillers' grains and used as animal feed or if dried can be used as biomass for heat generation.
- **Dreg**: solid fraction of the spent wash. It contains denatured proteins.
- **Spent Yeast**: post fermentation, may be combined with draff if all-in fermentation
- **Spent Lees**: Residual liquor after second distillation in malt whisky. Mostly water, but also contains some volatile components of the wash other than alcohol. Nutritive value is negligible and normally treated in bioplants.
- **Carbon dioxide (CO₂)**
Figure 2-1 shows a simplified process flow diagram of malt distilleries adapted from Russell (Russell et al. 2003) and Figure 2-2 (also modified from Russell et al) shows a weekly mass balance and process flow diagrams for a mid-sized distillery. A ratio of nearly 11 litres of pot ale for every litre of alcohol produced is in agreement with another source (Mohana et al. 2009). However, another reference (Crawshaw 2001) shows a value about 50% smaller than the figure mentioned previously. From this data, presented Table 2-1, it can be calculated that for every litre of alcohol, on average 5.5 litres of pot ale are produced. More recent figures from the Scotch Whisky Association (SWA 2012) showed that overall whisky production has increase by 49 %, and malt whisky production was up by almost 63% between 2001 and 2011. Assuming a malt whisky production of 255 million lpa, current pot ale generation in Scotland can be estimated between 2-3 million tonnes annually.

<table>
<thead>
<tr>
<th>Year</th>
<th>MW million lpa</th>
<th>PA million tonne</th>
<th>Ratio PA/MW</th>
</tr>
</thead>
<tbody>
<tr>
<td>1970</td>
<td>151</td>
<td>832</td>
<td>5.51</td>
</tr>
<tr>
<td>1975</td>
<td>187</td>
<td>1032</td>
<td>5.52</td>
</tr>
<tr>
<td>1980</td>
<td>164</td>
<td>904</td>
<td>5.51</td>
</tr>
<tr>
<td>1985</td>
<td>102</td>
<td>560</td>
<td>5.49</td>
</tr>
<tr>
<td>1990</td>
<td>182</td>
<td>1000</td>
<td>5.49</td>
</tr>
<tr>
<td>1995</td>
<td>158</td>
<td>872</td>
<td>5.52</td>
</tr>
<tr>
<td>2000</td>
<td>179</td>
<td>984</td>
<td>5.50</td>
</tr>
</tbody>
</table>

Malt whisky (MW) production expressed in lpa (liters of pure alcohol) and density of pot ale (PA) assumed 1 (kg/ L).

Concern for the environment has been and it is a major priority for the whisky industry (SWA 2014). Conventional ways of dealing with pollution have been reconsidered due to social, environmental and economic factors. Several procedures for waste treatment include chemical and/ or biological processes and have reported to have a significant economic impact to the industry (Mohana et al. 2009).
Figure 2-1. By-products from malt distilleries.

Figure 2-2 Weekly material flow for a medium-sized distillery.
2.2 Pot ale

Several texts (Crawshaw 2001, Russell et al. 2003) describe pot ale (PA) as a light brownish turbid liquid, with acidic pH (below 4) with high concentration of organic materials and solids. These solids are mainly intact yeast, yeast residues, soluble protein and carbohydrates and a significant but variable amount of copper. Several sources report different values between 40-140 mg copper/ kg dry matter (DM) (Buxton and Hughes 2013). The origin of this copper is due to the gradual dissolution of copper from the distillation stills leading to the presence of Cu(II) ions in pot ale and spent lees (Lu and Gibb 2008).

Table 2-2 shows a typical composition of pot ale of four different samples from the Hakushu distillery of Suntory in Japan. Table 2-3 presents the analysis of the contents in the separated solids from the same distillery, but presented in another study, which concluded that on a dry basis, 50% of the solids are crude proteins (Kida et al. 1999).

Table 2-2. Typical composition of whisky pot ale from the Hakushu distillery in Suntory, Japan (Kida et al. 1999).

<table>
<thead>
<tr>
<th>Component</th>
<th>mg l⁻¹</th>
</tr>
</thead>
<tbody>
<tr>
<td>Total Organic Carbon (TOC)</td>
<td>15,380 – 17,460</td>
</tr>
<tr>
<td>Suspended Solids (SS)</td>
<td>8,950 – 13,390</td>
</tr>
<tr>
<td>Volatile Suspended Solids (VSS)</td>
<td>8,402 – 12,980</td>
</tr>
<tr>
<td>Volatile Fatty Acids (VFA)</td>
<td>11,411 – 14,821</td>
</tr>
<tr>
<td>Protein</td>
<td>8,392 – 8,980</td>
</tr>
<tr>
<td>NH₄⁺</td>
<td>58 – 80</td>
</tr>
<tr>
<td>K⁺</td>
<td>290 – 971</td>
</tr>
<tr>
<td>Mg²⁺</td>
<td>148 - 277</td>
</tr>
<tr>
<td>Ca²⁺</td>
<td>46 – 58</td>
</tr>
<tr>
<td>NO₃⁻</td>
<td>1.9 - 2.5</td>
</tr>
<tr>
<td>PO₄³⁻</td>
<td>1,560 - 1,580</td>
</tr>
<tr>
<td>SO₄³⁻</td>
<td>223 - 285</td>
</tr>
<tr>
<td>pH</td>
<td>3 - 4</td>
</tr>
</tbody>
</table>

In an earlier work (Tokuda et al. 1998), malt whisky pot ale and grain spirit spent wash without suspended solids were analysed for saccharides and aliphatic acids as shown in Table 2-4. There was, in general, not much difference between them except in the levels
of lactic and propionic acids. However, dextrin contents differed markedly. Based on this work, carbohydrates accounted for 2.51% and 1.41% (w/v) of pot ale and spent wash content, respectively. Dextrin proportion in the carbohydrates was 83.7% (pot ale) and 76.6% (spent wash). In the same work, total organic carbon concentration (TOC) of both pot ale and spent wash of 10,000 to 15,000mg/L, and sometimes as high as 17,000 mg/L was reported.

Table 2-3. Composition of pot ale solids from Hakushu distillery in Suntory, Japan. (Tokuda et al. 1998)

<table>
<thead>
<tr>
<th>Items</th>
<th>Content ratio (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Water</td>
<td>78.23</td>
</tr>
<tr>
<td>Crude protein</td>
<td>11.26</td>
</tr>
<tr>
<td>Crude fat</td>
<td>0.64</td>
</tr>
<tr>
<td>Crude fiber</td>
<td>0.22</td>
</tr>
<tr>
<td>Crude ash</td>
<td>0.91</td>
</tr>
<tr>
<td>Others (non-nitrogen)</td>
<td>99.09</td>
</tr>
</tbody>
</table>

Table 2-4. Organic compounds and organic matter of pot ale and spent wash samples. (Tokuda et al. 1998)

<table>
<thead>
<tr>
<th>Organic compounds and organic matter (% w/v)</th>
<th>Pot ale</th>
<th>Spent wash</th>
</tr>
</thead>
<tbody>
<tr>
<td>Glucose</td>
<td>0.18</td>
<td>0.18</td>
</tr>
<tr>
<td>Fructose</td>
<td>0.08</td>
<td>0.09</td>
</tr>
<tr>
<td>Maltose</td>
<td>0.15</td>
<td>0.06</td>
</tr>
<tr>
<td>Dextrin with oligosaccharide</td>
<td>2.1</td>
<td>1.08</td>
</tr>
<tr>
<td>Lactic acid</td>
<td>0.61</td>
<td>0.42</td>
</tr>
<tr>
<td>Acetic acid</td>
<td>0.06</td>
<td>0.06</td>
</tr>
<tr>
<td>Propionic acid</td>
<td>0.03</td>
<td>0.12</td>
</tr>
</tbody>
</table>

In a more recent study (Mallick et al. 2010) using material from Blair Athol malt whisky distillery (Perthshire, Scotland, UK) different values and parameters for PA are presented in Table 2-5. These data are in agreement with the work presented by Graham et al. 2012 (Graham et al. 2012), where an average BOD value (Biochemical oxygen demand) of 24.9 g/L (with a range of 12.9 – 35.3 g/L) and a COD value (chemical oxygen demand) of 46.8 g/l (with a range of 38.4 -62.9 g/L) from an unnamed distillery were reported. This study revealed significant inconsistencies in distillery pot ale
composition throughout an 8 week sampling period and concluded that compositional variation in pot ale was more due to the inherent differences in pot ale composition, rather than sampling techniques.

Table 2-5. Characteristics of Pot Ale (Blair Athol distillery, Perthshire, Scotland). Adapted from Graham et al. 2012.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Total solids (g/l)</td>
<td>17.0</td>
</tr>
<tr>
<td>Total suspended solids (g/l)</td>
<td>8.3</td>
</tr>
<tr>
<td>Volatile suspended solids (g/l)</td>
<td>8.1</td>
</tr>
<tr>
<td>Total Kjeldahl nitrogen (mg/l)</td>
<td>92.0</td>
</tr>
<tr>
<td>Total COD (g/l)</td>
<td>61.5</td>
</tr>
<tr>
<td>pH</td>
<td>4.1</td>
</tr>
</tbody>
</table>

2.3 Pot ale syrup

In the early years of the 20th century PA was used as animal feed, but due its low solid content, became uneconomical. The solution in those days was to use the PA either as a fertiliser or to dispose the material into the sea. Economic and environmental considerations have made this practice no longer possible and led to the development of Pot Ale Syrup (PAS), which is simply PA concentrated by evaporation. However, using evaporation these days as a method for concentration is no longer an optimal solution from an economic and environmental perspective.

Another important consideration is that protein degradation has been reported during the process of dehydration (Crawshaw 2001). The extent of the protein quality lost seems to be proportional to the temperature and is one of the "most serious problems in the utilization of food waste as animal feed". (Kawashima 2004).

Typical evaporators used in the malt whisky industry can be classified into two types (Piggott et al. 1989): multiple effect (ME) evaporators and mechanical vapour recompression (MVR) evaporators. ME evaporators consists of several evaporators connected in series so that the vapour from the inside of the evaporator tubes serve as a heating medium on the outside of the tubes for the next effect. This configuration improves heat economy (kg steam per kg of water evaporated). ME evaporators with
more than six effects are normally not used, due to capital restrictions. An example of a two-effect evaporator is depicted in Figure 2-3.

![Two-effect evaporator](image)

Figure 2-3. Two-effect evaporator (principle) (Piggott et al. 1989)

The main difference between the ME and the MVR evaporator is that in the latter the vapour is not condensed in a condenser, but it is directed to a vapour compressor, recompressed and directed to the outside of the tubes (Figure 2-4). Efficiencies of MVR are superior to ME evaporators, however, the cost for small distilleries makes them unaffordable.
Figure 2-4. Two-stage MVR evaporator with finisher (Piggott et al. 1989).
The extent of evaporation achieved varies between distilleries. The limiting factor on achieving maximum concentration is the viscosity of the syrup. Typical dry matter content of PAS is between 30-50 per cent. Some of the nutritional benefits of PAS for animal feeding (bovine and pigs) found in the literature (Crawshaw 2001) are summarised below:

- High protein content (34-38% DM)
- High palatability
- Good amino acid balance due to yeast content
- Significant ash content, mainly due to phosphorous content
- Presence of the enzyme phytase from malt and yeast makes this phosphorous highly available to non-ruminant animals
- High digestibility by ruminants. Organic matter digestibility between 89 and 93%
- High Gross Energy (GE) content (20-20.4 MJ/kg DM), which combined with high digestibility results in high Digestibility Energy (DE) and Metabolised Energy (ME) values
- The low pH makes it good for storage
In Table 2-6 the nutritional values of PAS found in literature sources and other commercially available feeds from Whisky origin (Trafford Syrup®, Vitagold® and Spey Syrup®) promoted in the web are compared.

Table 2-6. Nutritional properties and chemical composition of PAS.

<table>
<thead>
<tr>
<th></th>
<th>PAS1</th>
<th>Spey Syrup2</th>
<th>Trafford Syrup2</th>
<th>Vitagold3</th>
</tr>
</thead>
<tbody>
<tr>
<td>Dry matter %</td>
<td>30-50</td>
<td>42</td>
<td>30</td>
<td>35</td>
</tr>
<tr>
<td>Crude Protein %</td>
<td>34-38</td>
<td>32</td>
<td>28</td>
<td>36</td>
</tr>
<tr>
<td>Crude Fibre %</td>
<td>0.20</td>
<td>0.17</td>
<td>1.20</td>
<td>4.69</td>
</tr>
<tr>
<td>Calcium %</td>
<td>0.14-0.20</td>
<td>0.15</td>
<td>0.10</td>
<td>0.08</td>
</tr>
<tr>
<td>Phosphorus %</td>
<td>1.6-2.2</td>
<td>0.21</td>
<td>0.57</td>
<td>0.45</td>
</tr>
<tr>
<td>Magnesium %</td>
<td>0.65</td>
<td>0.60</td>
<td>0.17</td>
<td>0.06</td>
</tr>
<tr>
<td>Sodium %</td>
<td>0.10-0.15</td>
<td>0.10</td>
<td>1.53</td>
<td>0.01</td>
</tr>
<tr>
<td>Potassium %</td>
<td>2.1-2.3</td>
<td>0.22</td>
<td>1.47</td>
<td>0.21</td>
</tr>
<tr>
<td>Copper mg/kg</td>
<td>60-180</td>
<td>40.9</td>
<td>3.5</td>
<td>6.0</td>
</tr>
<tr>
<td>Cystine %</td>
<td>0.7</td>
<td>2.11</td>
<td>1.52</td>
<td>2.01</td>
</tr>
<tr>
<td>Histidine</td>
<td>N/A</td>
<td>3.23</td>
<td>2.06</td>
<td>3.01</td>
</tr>
<tr>
<td>Isoleucine %</td>
<td>1.3</td>
<td>N/A</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Lysine %</td>
<td>2.1</td>
<td>6.47</td>
<td>3.90</td>
<td>3.01</td>
</tr>
<tr>
<td>Methionine %</td>
<td>0.35</td>
<td>1.06</td>
<td>1.41</td>
<td>1.84</td>
</tr>
<tr>
<td>Threonine %</td>
<td>1.9</td>
<td>5.61</td>
<td>3.04</td>
<td>3.60</td>
</tr>
<tr>
<td>pH</td>
<td>3.5 – 3.8</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Notes:
1 (Crawshaw 2001)
2 http://www.kwalternativefeeds.co.uk

Values are expressed as in basis and dry matter basis

N/A: Information Not Available
2.4 Copper content

Copper is an essential element necessary to all animals and humans. It is necessary for the proper growth, development, and maintenance of bone, connective tissue, brain, heart, and many other organs. It has been reported that copper is involved in the formation of red blood cells, the absorption and utilization of iron, the metabolism of cholesterol and glucose, and the synthesis and release of life-sustaining proteins and enzymes. These enzymes in turn produce cellular energy and regulate nerve transmission, blood clotting, and oxygen transport. (Fox 2003)

In animal feeds, copper is incorporated in diets in trace levels. Minimum requirements are recommended and maximum levels are set to avoid or minimise negative effects (i.e. toxicity, anaemia, liver and kidney problems) to humans and animals as well as to the environment (EFSA 2003). Table 2-7 shows the maximum limits of copper content in food diets set by European Regulations (Commission Regulation (EC) No 1334/2003).

<table>
<thead>
<tr>
<th>Animal</th>
<th>Maximum limit (in mg/ kg of the complete feedingstuff)</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Pigs</strong></td>
<td></td>
</tr>
<tr>
<td>Piglets up to 12 weeks</td>
<td>170</td>
</tr>
<tr>
<td>Other pigs</td>
<td>25</td>
</tr>
<tr>
<td><strong>Bovine</strong></td>
<td></td>
</tr>
<tr>
<td>Before the start of rumination</td>
<td>15</td>
</tr>
<tr>
<td>Other bovine</td>
<td>35</td>
</tr>
<tr>
<td><strong>Ovine</strong></td>
<td></td>
</tr>
<tr>
<td></td>
<td>15</td>
</tr>
<tr>
<td><strong>Fish</strong></td>
<td></td>
</tr>
<tr>
<td></td>
<td>25</td>
</tr>
<tr>
<td><strong>Crustaceans</strong></td>
<td>50</td>
</tr>
<tr>
<td><strong>Other species</strong></td>
<td>25</td>
</tr>
</tbody>
</table>

Due to its high copper content PAS has been utilised mainly as cattle and pig feed. Reported copper concentration levels as high of 100 mg/ kg DM in PAS, might constitute a risk if fed to sheep. However, there are some disagreements about this matter (Lewis 2002, Suttle and Underwood 2010).
Based on an American document (Committee on Animal Nutrition 1993) fish appear to have a higher tolerance of copper in diets than of dissolved copper in water. Concentrations of 0.8 to 1.0 ppm of copper sulphate in water are toxic to many fish species, but coho salmon (*Oncorhynchus kisutch*) "resisted up to 1,000 mg copper/kg of copper in the diet with only retarded growth and impaired pigmentation". The same report mentions no harmful effects of feeding diets containing 150 mg copper/ kg rainbow trout (*Oncorhynchus mykiss*) for 20 weeks.

### 2.5 By-products from the Ethanol and Brewing Industry

Detailed nutritional properties of PA/ PAS and economic data (i.e. price, volumes) are difficult to find. However, other by-products from similar industries were researched that might be helpful for future comparisons and references. The by-products of the bioethanol and brewing industries will be described in the following sections.

#### 2.5.1 Bioethanol Industry

Information about the Bioethanol Industry (BEI) by-products, such as nutritional properties (Liu 2011, Belyea et al. 1998, Belyea et al. 2006) and economic data (Cottrill 2007), are abundant and variable. Some of these materials can be found in web sites of (U.S) states agricultural offices, Universities, trade or commodity organizations (University of Minnesota. Department of Animal Science).

By-products from the bioethanol plants include distiller's dried grains (DDG), distiller's dried solubles (DDS), and distiller's dried grains with solubles (DDGS). Additionally, after the fermented mash is distilled, the soluble portion of the remaining residue is condensed by evaporation to produce another by-product called condensed distiller's solubles (CDS). A diagram of a typical bioethanol process is presented in Figure 2-5 (Bothast and Schlicher 2005).

Normally, ethanol plants blend and dry DDS and DDG to produce DDGS, which is the only form available to the feed industry. DDS has a higher concentration of nutrients compared to DDG and DDGS. It is a rich source of vitamins, and is the lowest in fibre and highest in fat, yielding a high DE value (approximately 91% of that found in corn).
Since DDGS is a blend of DDS and DDG, the nutrient composition of DDGS is a mixture between DDS and DDG.

**Figure 2-5.** Process diagram of maize (corn) bio-ethanol (Bothast and Schlicher 2005).
2.5.2 Brewing Industry

Solid by-product streams from breweries include spent grains, trub (also known as hot break), tank bottoms (cold break) and spent/excess yeast. A diagram of the brewing process and the main by-products generated can be observed in Figure 2-6. A summary of typical brewing by-product volumes and protein content is presented in Table 2-8.

![Brewing Process Diagram](image)

**Figure 2-6.** Brewing process in breweries and main by-products generated (Olajire 2012).

<table>
<thead>
<tr>
<th>By-product</th>
<th>Volume (kg of by-product per m³ of beer produced)</th>
<th>Protein content (g of protein per liter of beer produced)</th>
<th>Protein content (DM basis)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Spent-Grain</td>
<td>150</td>
<td>100-200</td>
<td>19-30%</td>
</tr>
<tr>
<td>Spent yeast</td>
<td>2.0</td>
<td>15</td>
<td>40-50%</td>
</tr>
<tr>
<td>Trub</td>
<td>0.8</td>
<td>1-2</td>
<td>40-50%</td>
</tr>
</tbody>
</table>
Spent grains are the leftover solids after milled cereals have been mashed to release carbohydrates and other desirable compounds for use in the fermentation. They vary in composition both across, and within breweries, with average protein levels ranging from 19 to 30% (dry matter basis). These variations are attributed to the raw materials and differences in extraction efficiencies of the individual brewhouses. Some brewers will add cereal adjuncts, which may also influence the levels of protein present in the spent grain. Variety may also be introduced into the final composition of spent grains dependent upon the brewing and milling techniques used (Briggs et al. 2004).

Prior to fermentation, the carbohydrate-rich liquid (the wort) is separated from the spent grains and boiled with hops to release protein and add flavour to the final product. Trub (hot break) is removed in the form of precipitated solids during this boiling step. The protein in trub forms strong hydrogen and hydrophobic bonds with polyphenols, which can be separated. Although trub has a high protein content of around 40-50% dry matter (Hardwick 1994), this protein is not highly digestible (20-30% digestible crude protein) (Hough et al. 2012) which may be presumed to be due to known anti-nutritional compounds such as polyphenols and phytate (Doria et al. 2012, Dai et al. 2007). This sub-optimal protein however only represents a small fraction of the available protein and separation of this protein from anti-nutritional compounds is possible.

When the hot wort is cooled prior to fermentation, or beer cools after fermentation (cold storage/maturation), cold break will be formed, and can continue forming during fermentation. Cold break also consists of protein precipitates, but these are smaller in size than those associated with hot break, and as a result slower to settle, and can require the addition of fining agents to aid their removal. This process is often followed by a filtration step. The quantity of cold break formed is temperature, and thus process dependent, and varies between 0.1-0.7 g/L (Hough et al. 2012). Cold break has been reported to be comprised of up to 70% by weight of protein (South 1996) but the process used to remove cold break will influence the availability of this protein for recovery and reuse.

Excess yeast is removed by brewers at the end of fermentation, and can be pressed or centrifuged to recover the beer, leaving the yeast in the form of pressed cake (Boulton and Quain 2001). This pressed cake is around 30% dry matter (Crawshaw 2001) of which 40-50% is protein. Usually around 20% of the yeast will be kept for re-use in
future fermentations and the remainder is available for protein recovery (Crawshaw 2001). The high protein content in excess/spent yeast makes it a desirable foodstuff and it is commonly sold to farmers as swine feed (Bamforth 2009, Ferreira et al. 2010). In volume terms, Brewer's yeast is the second largest by-product and similar to pot ale. Table 2-9 contrast the main nutritional properties (i.e. amino acids and minerals) of these two compounds and DDGS mentioned earlier.

There can be a large degree of variation between these streams in terms of composition and how they are handled or indeed combined in some cases. However, brewers co-products represent a source of protein from grains and yeast potentially totalling well over 100,000 tonnes per annum in the UK alone, which could be recovered for use in feedstock (Huige 2006).
Table 2-9. Comparison between Distillers Dried Grains with Solubles (DDGS), Brewer’s Dried yeast (BDY) and Pot Ale Syrup (PAS).

<table>
<thead>
<tr>
<th></th>
<th>DDGS$^1$</th>
<th>BDY$^1$</th>
<th>PAS$^2$</th>
</tr>
</thead>
<tbody>
<tr>
<td>DM (%)</td>
<td>92</td>
<td>93</td>
<td>30-50</td>
</tr>
<tr>
<td>CP (%)</td>
<td>29</td>
<td>48</td>
<td>34-38</td>
</tr>
<tr>
<td>Fibre (%)</td>
<td>5</td>
<td>7</td>
<td>0.2</td>
</tr>
<tr>
<td>Cystine %</td>
<td>1.3</td>
<td>1.1</td>
<td>0.7</td>
</tr>
<tr>
<td>Isoleucine %</td>
<td>5.5</td>
<td>4.7</td>
<td>1.3</td>
</tr>
<tr>
<td>Leucine %</td>
<td>8.5</td>
<td></td>
<td>N/A</td>
</tr>
<tr>
<td>Lysine %</td>
<td>2.6</td>
<td>7.0</td>
<td>2.1</td>
</tr>
<tr>
<td>Methionine %</td>
<td>1.8</td>
<td>1.6</td>
<td>0.35</td>
</tr>
<tr>
<td>Threonine %</td>
<td>3.5</td>
<td>4.7</td>
<td>N/A</td>
</tr>
<tr>
<td>Calcium (%)</td>
<td>0.17</td>
<td>0.12</td>
<td>0.14-0.20</td>
</tr>
<tr>
<td>Phosphorous (%)</td>
<td>0.84</td>
<td>1.43</td>
<td>1.6-2.2</td>
</tr>
<tr>
<td>Magnesium (%)</td>
<td>0.21</td>
<td>0.24</td>
<td>0.65</td>
</tr>
<tr>
<td>Sodium (%)</td>
<td>0.04</td>
<td>0.09</td>
<td>0.10-0.15</td>
</tr>
<tr>
<td>Potassium (%)</td>
<td>0.65</td>
<td>1.71</td>
<td>2.1-2.3</td>
</tr>
<tr>
<td>Copper (mg/kg)</td>
<td>59</td>
<td>33.0</td>
<td>60-180</td>
</tr>
<tr>
<td>Manganese (mg/kg)</td>
<td>29</td>
<td>5.7</td>
<td>N/A</td>
</tr>
<tr>
<td>Selenium (mg/kg)</td>
<td>0.38</td>
<td>1.25</td>
<td>N/A</td>
</tr>
</tbody>
</table>

Sources:

$^1$ (Huige 2006)

$^2$ (Crawshaw 2001)
2.6 Market price and prospects for pot ale syrup

Examples of products found in the market were mentioned before (Trafford Syrup®, Vitagold® and Spey Syrup®). All these products are marketed as high protein feeds and compete with other feeding stuffs found in the market (i.e. Soy bean meal, maize, etc.)

The economic value of PAS as animal feed will be discussed later in section 2.8. Typically the price of PAS is in the range of £80-100 per tonne (Scottish Agricultural College 2012). It is worth to mention that protein and energy content are the main parameters when evaluating the price of a feed compound. However, suspended solids and flow ability are factors to take into account since it affects PAS market price.

No historical information about the market prices can be obtained for PAS, but lesson could be learned from the bioethanol industry (i.e. DDGS). In Figure 2-7 from (Wisner 2010) it can be observed that the production of DDGS expanded very rapidly in the last 10 years with the explosive growth in corn processing for ethanol production. In 2005 approximately 20 billion litres of ethanol were produced in the United States (U.S.) and by 2014 this figure more than double to 53 billion litres. Another source indicates that approximately 35 million metric tonnes of DDGS were produced North America in 2011 (University of Minnesota. Department of Animal Science).

Due to this very fast production growth, concerns about the balance between supply and demand of DDGS have been pointed out (Hoffman and Baker 2010). Issues such as the supply of DDGS exceeding the animal feed market raise questions about the economic feasibility of DDGS.

The same scenario could be thought to happen to PAS in the UK. The Scottish whisky and distilling industry is an expanding business and together with new bioethanol plants, would eventually increase the supply of PAS and similar animal feed products with the consequence of bringing the price of PAS down.

One example of bio-ethanol plants in the UK is the £300 million Vivergo plant at Saltend near Hull, which started production in 2012 (FWi 2012) with a capacity of 420 million litres of bioethanol annually and 0.5 million tonnes of animal feed. In
comparison with PAS, it could be calculated that the potential supply of animal feed from PAS in the UK is about 1 M tonne (assuming whisky production volume of 458 million lpa per year, Whisky Production to PAS generation ratio at 10 and PAS density of 1 kg per litre).

![Figure 2-7](image1.png)

**Figure 2-7** Historical and forecasted production of DDGS in the US (Wisner 2010).

![Figure 2-8](image2.png)

**Figure 2-8** Historical price of DDGS as percentage of soy bean meal (SBM). (Wisner 2010).
2.7 Current and future demand for pot ale

Previous sections focused on understanding the supply (i.e. the availability or production) of PA/ PAS and similar by-products (DDGS and BDY). This section will concentrate on the demand side (i.e. the consumption) of PA/ PAS from current and potential customers. Different uses were identified previously by other authors (Russell et al. 2003), including animal feed, human food, fertiliser, fuel and biomass production, all of them with limited viability.

This document will revisit the animal and human markets adding new research material found in the literature. It is important to mention that the work (from Russel et al) did not consider the aquaculture industry as a potential user of PA. The option of this potential market will be analysed in depth in this review.

Other uses rather than food and feed are also important to consider, but due to the limits of this work and two important facts: the growing world population and potential synergies between the Scottish whisky and salmon industry; the focus of this review will stay on food and feed applications.

The world’s population will be around 9 billion people by 2050. A bigger population will demand more animal proteins like beef, poultry and fish. It is crucial that the animal feed industry can meet this challenge in a sustainable and safe way. The second fact is that Scotland is one of the biggest producers of salmon and the whisky industry should take advantage of this.

Both facts raise questions about protein sources and methods used to obtain them, i.e. sustainable means. There is a lot of public debate about this issue (The Guardian 2010) and plenty of material has been written and researched about this topic (Harry 2011, Speedy 2004). Sustainability considerations should be taken for the development of alternative PA protein concentration methods and the usage to protein obtained.
2.7.1 Animal Feed

Global compound feed production is estimated at almost 1 billion tonnes annually with sales equivalent to US$370 billion worldwide (IFIF 2015). European production of animal feed is 150 million tonnes while the UK market is around 15 million tonnes (FEFAC 2013).

To meet this protein demand, it is important to look for alternative protein sources as well as improved efficiencies of protein conversion. Different animals have different protein requirements and different protein conversion ratios.

Approximately 80% of the protein needed for agriculture in the UK is imported (Vivergo Fuels 2012). Proteins contained in PA would help to reduce this dependency. However, the animal feed market could be considered to have limitations, since restricted amounts of PA could be included in the (animal) diets. Moreover, due to the high copper content, it is recommended to avoid the intake for certain animal (i.e. sheep).

Table 2-10 shows a few examples of recommended maximum daily feed rates (per head basis) of commercial PA (Trafford Syrup®, Vitagold® and Spey Syrup®) for several animals. Values vary from 10 to 50% of the animals Dry Matter Intake (DMI).

Table 2-10. Recommended daily feed rates of commercial pot ale syrup.

<table>
<thead>
<tr>
<th></th>
<th>Spey Syrup</th>
<th>Trafford Syrup</th>
<th>Vitagold</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Milking Cows</strong></td>
<td>5</td>
<td>6</td>
<td>15</td>
</tr>
<tr>
<td><strong>Calves (to 12 weeks)</strong></td>
<td>0.75 (10%)</td>
<td>0.75 (10%)</td>
<td>5 (30%)</td>
</tr>
<tr>
<td><strong>Growing Cattle</strong></td>
<td>4 (15%)</td>
<td>5 (15%)</td>
<td><em>ad lib</em> (50%)</td>
</tr>
<tr>
<td><strong>Finishing Cattle</strong></td>
<td>5 (20%)</td>
<td>6 (20%)</td>
<td><em>ad lib</em> (50%)</td>
</tr>
</tbody>
</table>

Note:
Figures are in kg per animal head. Figures in brackets represent the %age of the DMI.

Source: [http://www.kwalternativefeeds.co.uk](http://www.kwalternativefeeds.co.uk)
2.7.2 Aquaculture

In 2012 global fish production (including crustaceans, molluscs and other aquatic animals) reached 158 million tonnes and it is projected to achieve 164 million tonnes by 2020. Current global aquaculture production is 70 million tonnes and is expected to reach 74 million tonnes by 2020. The Fish Farming Industry in the UK produces mainly Atlantic Salmon (*Salmo Salar*). The production volume in 2013 was 163 thousand tonnes with a worldwide retail value worth over £1 billion (The Scottish Government 2013).

Aquaculture feeds (aquafeeds – AQF) have traditionally relied heavily on fishmeal (FM) and fish oil (FO) but the supply of these raw materials is not increasing at the same speed of the demand (EWOS 2010). An expansion in aquaculture capacity implies that the AQF Industry will need to diverge from its dependency on marine resources. This means finding suitable alternative feed sources that promote fish growth and quality, whilst maintaining the human health benefits associated with salmon meat.

The results of a global survey concerning the estimated use of FM and FO within feeds for the major cultivated species groups (includes *Salmo salar*, *Oncorhynchus kisutch* and *O. tshawytscha*) are shown in Table 2-11 (Albert and Marc 2008). Wide variations were observed concerning FM and FO use within and between countries for the same species. For Atlantic salmon, FM usage varies from 20 to 50% and FO from 9 to 35%. To a large extent these variations reflect the differences within and between countries regarding the production systems employed, feed ingredients (FM and FO) availability, quality, cost, legislation, subsidies, incentives and the intended market and market value of the culture species.

For example, the United Kingdom reported the highest usage of FM and FO within salmon feeds in 2006 (36 and 28%, respectively). This is primarily due to the restrictive demands of the resident national salmon farming associations, major salmon retailers and supermarket chain retailers within the UK concerning the use of FM and FO replacers. This includes the prohibition of some terrestrial animal by-products (i.e. poultry) and genetically modified feed ingredient sources within compound feeds.
The same work (Albert and Marc 2008) agrees that FM and FO usage is expected to decrease in the long term due to decreasing market availability from capture fisheries, increasing market cost and an increased global use of cheaper plant and animal alternative protein and lipid sources.

Some plant proteins, such as soy beans (i.e. SBM) have been used to replaced FM and FO. They are considered an economical alternative with good nutritional properties (e.g. high protein content, good amino acid profile), however certain nutritional characteristics and the presence of several antinutritional factors deserve further analysis. Among the undesirable components in the feeds a few including protease inhibitors, phytate (or phytic acid), saponinins, tannins and phytoestrogens are cited (Chiesa and Gnansounou 2011). From the same report, negative and positive effects are presented, but a further analysis and the impact to aquaculture feeds is beyond the scope of this review.

To be a viable alternative feedstuff to FM, a candidate ingredient must possess certain characteristics, including “wide availability, competitive price, plus ease of handling, shipping, storage and use in feed production” (Gatlin et al. 2007). From a nutritional perspective the same source adds that these alternative feeds must have “low levels of fibre, starch, especially nonsoluble carbohydrates and anti-nutritional compound, plus
have a relatively high protein content, favourable amino acid profile, high nutrient digestibility and reasonable palatability”. From the same document Table 2-12 summarizes the nutrients found in FM (expressed on an as-fed basis), and the range of nutrient concentrations that alternative ingredients (i.e. SBM, DDGS, PAS, etc.) should have to be viable alternatives to FM. Erasmus and other authors recognize that "the primary driver for the use of a particular feed ingredient is its cost, followed by purity, nutrient availability and digestibility. The cost of the nutrient is not only defined in terms of price per tonne, but also in terms of price per kg of live fish weight generated over the entire growth cycle of the fish". (Erasmus 2009)

Some authors conclude that a combination of plant-derived feed ingredients will be required to replace fish meal (Eldar Åsgard et al. 2011, Hardy 2010). Additionally supplements, such as amino acids, flavourings and possibly exogenous enzymes (i.e. phytase), will be needed to produce aquafeeds without or minimal FM that support growth rates necessary for the economic production of farmed fish (Hardy 2000).

Table 2-12 Nutrient content (as-fed basis) of fish meal and targeted ranges in alternative ingredients derived from grains and oilseeds (Gatlin et al. 2007).

<table>
<thead>
<tr>
<th>Category/ Nutrient</th>
<th>Fish Meal</th>
<th>Target Range for alternative ingredients</th>
</tr>
</thead>
<tbody>
<tr>
<td>Crude protein (%)</td>
<td>65-72</td>
<td>48-80</td>
</tr>
<tr>
<td>Crude lipid (%)</td>
<td>5-8</td>
<td>2-20</td>
</tr>
<tr>
<td>Fibre (%)</td>
<td>&lt;2</td>
<td>&lt;6</td>
</tr>
<tr>
<td>Ash (%)</td>
<td>7-15</td>
<td>4-8</td>
</tr>
<tr>
<td>Arginine (%)</td>
<td>3.75</td>
<td>&gt;3.0</td>
</tr>
<tr>
<td>Lysine (%)</td>
<td>4.72</td>
<td>&gt;3.5</td>
</tr>
<tr>
<td>Methionine (%)</td>
<td>1.75</td>
<td>&gt;1.5</td>
</tr>
<tr>
<td>Threonine (%)</td>
<td>2.5</td>
<td>&gt;2.2</td>
</tr>
</tbody>
</table>

Potential substitution of vegetable meals and oils in place of animal-derived ingredients (i.e. FM and FO) offers significant opportunities to decrease environmental pressure of Aquaculture Industries (i.e. Salmon) and the improvement of resource efficiencies.
2.7.3 Experience of Distiller's by-products in Aquaculture

Limited research has been carried out regarding the usage of DDGS in Aquaculture. Gatlin et al (Gatlin et al. 2007) stated that the "relatively high fibre content" (5% against less than 0.2% of FM) limits the use of DDGS in AQF. Based on this fact PA could have a good chance to be incorporated in AQF (fibre content around 0.2%). Another review is provided in the DDGS - University of Minnesota website, where maximum inclusion rates of 8% for salmon and trout are suggested (Shurson 2004).

2.7.4 Experience of brewer's by-products in Aquaculture

In Ferreira's review of brewer's by-products (Ferreira et al. 2010) several publications name brewer's yeast as a potential substitute for live food in the production of certain fish species (including rainbow trout and salmon) or as a replacement for FM. The same work reports up to 50% replacement of FM with no negative effect in fish performance and up to a 30% inclusion of BDY in fish diets improved feed efficiency. An important point made here is in regards to protein digestibility. From the same work, lessons can also be learned about BDY used in rainbow trout. The disruption of cell wall has been suggested to increase protein digestibility. This is in line with previous research carried out by Heriot-Watt (Traub 2011).

2.7.5 Food applications

Based on figures from 2000, the Food and Agriculture Organization of the United Nations (FAO) estimated a protein supply for human consumption from animal origin (Meat: 233 million tonnes, Milk: 568 million tonnes, and Eggs: 55 million tonnes) in 65 million tonnes. That figure of Global meat production and consumption (233 million tonnes) increased in the year 2008 to 281 million tonnes and it is forecast to expand to 300 million tonnes by the year 2020. (Speedy 2004)

Traditional protein sources might not be the enough to this satisfy the increased demand. Alternative protein together with improved production efficiencies (i.e. less waste) will be needed (Harry 2011).
Among the alternative protein sources food yeast or fungal food has been named a good candidate. PA, due to its yeast and protein content, could have a place in this market, too. The work of Russell (Russell et al. 2003) did not forecast a big opportunity for whisky by-products in the food market due to high fibre and copper content; however, specific analysis about PA (with lower fibre content) was not included.

Food yeast is defined by the European Association for Specialty Yeast Products (EURASYP) as Food "a nutritional ingredient consisting of baker's, brewer's or lactic yeast which has been inactivated by heat and consequently has no more fermentative capability." (EURASYP)

A recent review (Ghorai et al. 2009), confirms that over the recent years, the consumption of fungal food has augmented, due to an increased awareness about dietary and health issues among the consumers. The same work revisits the uses and varieties of fungi in food and feed including the strain *Fusarium venenatum*, used to produce myco-protein "Quorn". The nutritional properties of Quorn® are summarised below:

- good protein content having all the essential amino acids (specially enriched with lysine)
- chitinous wall act as a source of dietary fibre
- high vitamin B content
- low in fat
- virtually free of cholesterol

However, there are problems associated with the intake of fungal food. More research is needed and particularly the use of PA in humans will need to be investigated. An important fact mentioned in the literature is the nucleic acid content in fungal food products. The amount of nucleic acid, primarily ribonucleic acid (RNA), that is present in yeast, can lead to gout (Huige 2006). However, techniques and reagents are used to isolate yeast protein from low RNA (Ferreira et al. 2010).

Finally, consumer acceptance is another factor to be considered. An example of this is provided in a review (Siro et al. 2008), where it is affirmed that the consumer acceptance is "widely recognised as a key success factor for market orientation,
consumer led product development, and successfully negotiating marketing opportunities."

2.8 Proteins Economics

2.8.1 Calculation of the economic value

The economic value of any feed is determined by the nutrients contents (i.e. protein, energy, mineral content). Feeds that compete with a by-product depend on the primary nutrient(s) contributed. Protein requirement for beef cattle for example, the feed values are normally compared to soybean meal (SBM).

There are several ways of estimating the value of any feedstuff, including by-products. These range from simple calculations based on the value of the nutrient in one common feedstuff to very specific ration analyses and comparison. The simpler methods may help determine if a feedstuff is generally priced so that it may be a competitive alternative (Iowa Beef Center. Iowa State University 2007).

Comparisons should be made on a dry basis. The effect of moisture might also have a negative effect on the price by increasing storage and handling costs and storage losses. These factors need to be considered when determining the value of high moisture or not easy to handle material.

A price for PAS (P_{PAS}) can be calculated using equations (E. 1) and (E. 2), assuming the following data:

- Soybean meal price (P_{SBM}) assumed at £230 per tonne
- 88% dry matter content for soy bean meal (DM_{SB}) and between 30 to 50% for PAS (DM_{PAS})
- 48% Crude protein content for soybean meal (CP_{SBM}) and between 34-38% for PAS (CP_{PAS})
The calculations show a price for PAS between £ 56 – 103 per tonne, and a bulk protein price \( (P_{BP-SBM}) \) of £545 per tonne. A similar analysis could be made using fishmeal (FM), another important protein source for animal feeding. The parameters to calculate the bulk protein price for FM are presented below:

- Price (\( P_{FM} \)): £832 per ton
- Crude protein (\( CP_{FM} \)): 65%
- Dry Matter (\( DM_{FM} \)): 92%

With the above numbers the bulk protein price would be £1,391 per tonne. A summary of these results are presented in Table 2-13. It must be mentioned that the maximum and minimum price of SBM in the last five years were £111.44 (April 2007) and £270.1 (June 2009) respectively, while the minimum price for FM was £497.96 (November 2007) and the maximum was £1,303.92 (May 2010). A chart of the prices over 2003-2014 of both protein sources (FM and SBM) are presented in Figure 2-9 (Source: http://www.indexmundi.com/commodities/).

### Table 2-13. Price calculation of PAS based on SBM.  

<table>
<thead>
<tr>
<th></th>
<th>FM</th>
<th>SBM</th>
<th>PAS</th>
</tr>
</thead>
<tbody>
<tr>
<td>Average Price (£/ ton)</td>
<td>832</td>
<td>230</td>
<td>56 – 103</td>
</tr>
<tr>
<td>Dry Matter %</td>
<td>92%</td>
<td>88%</td>
<td>30-50%</td>
</tr>
<tr>
<td>Crude protein %</td>
<td>65%</td>
<td>48%</td>
<td>34-38%</td>
</tr>
<tr>
<td>Bulk Protein Price (£/ ton)</td>
<td>1391</td>
<td>545</td>
<td>545</td>
</tr>
</tbody>
</table>
2.8.2 Price comparison between protein sources and grades

Toride (Toride 2004) defined the animal industry as “an industry producing proteins of higher value (meat, milk) from less expensive protein sources (vegetable proteins, i.e. soybean meal).”

In Table 2-14 the price of different protein sources are compared. It is important to highlight that food grade protein (i.e. for human consumption) is approximately 30 times more expensive than feed grade (i.e. animal consumption). Quorn® is considerably more expensive than the other protein sources. This fact might be explained because of Quorn® is a value added product, while beef and fish are commodities.
Table 2-14. Protein prices comparison.

<table>
<thead>
<tr>
<th></th>
<th>Feed Grade proteins</th>
<th>Food Grade Proteins</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>SBM</td>
<td>FM</td>
</tr>
<tr>
<td><strong>Average Price</strong></td>
<td>£/ tonne</td>
<td></td>
</tr>
<tr>
<td>DM</td>
<td>%</td>
<td></td>
</tr>
<tr>
<td><strong>CP</strong></td>
<td>%</td>
<td></td>
</tr>
<tr>
<td><strong>Protein Price</strong></td>
<td>£/ tonne</td>
<td></td>
</tr>
<tr>
<td><strong>Ratio to SBM</strong></td>
<td>1.0</td>
<td>2.1</td>
</tr>
</tbody>
</table>
2.9 Conclusions

PAS might lose commercial attractiveness as animal feed due to economic and environmental factors. With the start-up of new bioethanol plants, concerns about over supply of animal feed products have been expressed. Additionally protein damage has been reported during the traditional protein concentration methods (evaporation). To remain as a competitive product for the animal feed market a more sustainable method to concentrate and purify the proteins from PA must be developed.

PA competes with other by-products from the food and beverages industry (including brewing and distilling) in the animal feed market. The economic value of PA is driven by its protein content, moisture content and the price of soy bean meal. A value between £56 and £103 per tonne was estimated for PAS based on these three variables. Other nutritional properties such as amino acid profile, mineral (i.e. P, K, Ca) and fibre content also influence the price. The relatively high copper content in PA is something that needs to be carefully considered in regards to different animal markets.

Not much detail is available about the nutritional properties of PA. Differences in composition are common between commercially available PAS. These variations seem to happen in other by-products (i.e. DDGS) as well. A good understanding of the nutritional properties of PA such as amino acid profile and mineral content are crucial to establish a market and a price for PA.

Alternative markets to animal feeding (i.e. poultry, pigs, cows and sheep) were also reviewed. Aquaculture is an interesting and logical option to explore, since Scotland is one of the biggest producers of Salmon in the world. Economic and environmental benefits for both industries (synergies) could be obtained. The salmon industry - and particularly the UK - relies heavily on fish meal and fish oil, non-renewable resources, to satisfy its requirements.

Another option investigated for PA was human food. This alternative has some potential, but its viability is uncertain. Unknowns about health issues, cost considerations, added complexity to the process and consumer acceptance were highlighted.
Sustainability issues should be taken into account for process and product development. There is increasing concern about future protein supply for both human and animal needs. This fact could play in favour to both whisky and salmon industry.

Proteins were the main product investigated, but other components of PA such as chitin and glucans (e.g. yeast wall components) should be investigated too. These substances might have higher value than proteins, thus offering good opportunities for further research and commercial applications.

In the next chapter of this thesis, characterisation of brewing and distilling by-products collected during the course of the project will be presented and compared with the information already discussed in this chapter.
CHAPTER 3 - BREWING AND DISTILLING BY-PRODUCTS
CHARACTERISATION

Abstract

Methods for the characterisation of malt distillery by-products were developed in this chapter. Additionally, grain whisky and brewing by-products were analysed. Some of the methods developed were solids content, pH analysis, total nitrogen (crude protein content), soluble protein and metal analysis.

Total protein content in pot ale was calculated at 15 g/L, of which approximately 70% is found in the liquid fraction. The solid fraction was also found to have ~70% of the copper. This fact, that the liquid component of pot ale contains most of the protein and a fraction of copper, led to the decision of focusing future work on the liquid component of pot ale.
3.1 Introduction

In this Chapter, the methods for the characterisation of distillery by-products will be presented. These methods were used through the whole project.

In addition to malt whisky by-products, grain whisky and brewing by-products were also analysed. As discussed in the previous chapter, these by-products have similar properties and future work could contemplate the recovery of proteins from these by-products.

Pot ale from only two sources were analysed in this chapter (Glenkinchie and Speyside). However, during the course of the project other sources were used for protein concentration experiment and a full characterisation of the by-product was conducted. This information is however, not included in this thesis chapter.
3.2 Materials and Methods

3.2.1 By-product sourcing, type and storage

Liquid and solid by-products were obtained from several sources including Breweries and Distilleries located in Scotland. Additionally a by-product from a whisky produced in International Centre for Brewing and Distilling (ICBD) of Heriot-Watt University was analysed.

Brewery sources included Lager (Wellpark - WP) and Ale (Caledonian Brewery – CB). By-products analysed include Spent Grains (SG), Spent Yeast (SY), Trub (TR) and Spent Hops (SH).

Malt Whisky sources included Glenkinchie (GK) Distillery and a Speyside\(^1\) (SS) Distillery. The type of by-products received included Pot Ale (PA) and Draff (equivalent to SG and classified as SG for the purpose of this work). Spent Wash (SW) was the only Grain Whisky by-product and it was sourced from North British (NB) Distillery.

Heriot-Watt University (HW) “whisky” was made utilising molasses and yeast (Saccharomyces cerevisiae) as the main ingredients. The by-product generated was of similar appearance of Pot Ale and hence it was classified as Pot Ale for this work.

Liquid samples (PA, SW, SY and TR) were stored immediately once received at 4°C for a maximum of one week and analyses were performed in this period of time. The residual samples were stored at -15°C for subsequent analysis.

\(^1\) Due to confidentially issues the distillery could not be named.
Solid by-products (SH and SG) were oven dried at 60°C for 2 to 3 days to remove moisture content and then they were placed in sealed plastic bags to avoid samples absorbing ambient moisture.

A brief description of the by-products is presented in Table 3-1 and a summary of the samples collected including source, origin (brewery or distillery) and type is offered in Table 3-2.

**Table 3-1. Brief description of brewing and distilling by-products.**

<table>
<thead>
<tr>
<th>By-product</th>
<th>Abbreviation</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Draff</td>
<td>DR</td>
<td>Solid material with moisture content composed mainly by barley.</td>
</tr>
<tr>
<td>Pot Ale</td>
<td>PA</td>
<td>Liquid material with solids content - mainly yeast cells.</td>
</tr>
<tr>
<td>Spent Grains</td>
<td>SG</td>
<td>Solid material with moisture content composed mainly by wheat.</td>
</tr>
<tr>
<td>Spent Hops</td>
<td>SH</td>
<td>Solid material composed mainly by hops with moisture content.</td>
</tr>
<tr>
<td>Spent Wash</td>
<td>SW</td>
<td>Solid material composed mainly by grains (other than barley) with moisture content.</td>
</tr>
<tr>
<td>Spent Yeast</td>
<td>SY</td>
<td>Liquid material composed primarily of (dead) yeast cells</td>
</tr>
<tr>
<td>Trub</td>
<td>TR</td>
<td>Liquid material composed primarily of spent hops</td>
</tr>
</tbody>
</table>

**Table 3-2. Matrix of by-product sources, origin and types.**

<table>
<thead>
<tr>
<th>Source</th>
<th>Brewery (B)</th>
<th>Distillery (D)</th>
<th>PA</th>
<th>SG</th>
<th>SH</th>
<th>SW</th>
<th>SY</th>
<th>TR</th>
</tr>
</thead>
<tbody>
<tr>
<td>Caledonian Brewery (CB)</td>
<td>B</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Glenkinchie (GK)</td>
<td>D</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Heriot-Watt (HW)</td>
<td>D</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>North British (NB)</td>
<td>D</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Speyside (SS)</td>
<td>D</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Wellpark (WP)</td>
<td>B</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
3.2.2 **Solids content (liquid by-product samples)**

3.2.2.1 **Totals solids content**

Total solids (TS) correspond to the dry weight of the whole liquid sample after oven drying it at 105°C for 24 hours. The analyses were performed in triplicate, the weights recorded in grams (g) and volumes in millilitre (mL).

An aluminium dish (Fisher, FB71085) was pre-dried by placing it in an oven at 105°C for a minimum of 4 hours. The dish was cooled in a desiccator and the weight was recorded in grams. The liquid sample was thoroughly mixed and 10 ml was added to each dish. The weight of each sample - including the dish - was recorded. The aluminium dish containing the sample was put in an oven at 105°C for 24 h and then the sample was removed and placed in a desiccator to cool. The dry weight of the dish containing the oven-dried sample was recorded.

Total Solids on a weight basis (TS\(_{105^\circ C, w/w}\)) was calculated using Equation 1 (E.1.).

Total solids on a volume basis (TS\(_{105^\circ C, w/v}\)) was obtained with E.2. Total Solids in grams per litre of by-product (TS\(_{105^\circ C, g/L}\)) was calculated with E.3.

\[
E.1. \quad TS_{105^\circ C, w/w} = \frac{W_{DD+DS} - W_{DD}}{W_{DD+WS} - W_{DD}} \times 100 \quad (g/ g)
\]

\[
E.2. \quad TS_{105^\circ C, w/v} = \frac{W_{DD+DS} - W_{DD}}{10} \times 1000 \quad (g/ L)
\]

\[
E.3. \quad TS_{105^\circ C, g/L} = \frac{W_{DD+DS} - W_{DD}}{10} \times 1000 \quad (g/ L)
\]

Where:

\(W_{DD}\): Weight of the dry dish (g)

\(W_{DD+WS}\): Weight of the wet sample including the weight of the dry dish (g)

\(W_{DD+DS}\): weight of the dry sample including the weight of the dry dish (g)
3.2.2.2 Total centrifuged solids

The total centrifuged solids correspond to the dry weight of washed solids pellet after centrifugation.

10 ml of the liquid by-product were added to a 15 ml (Anachem, ABCT-15) tube and then centrifuged at 4000 rpm (Heraeus Multifuge 3SR) for 10 minutes. The supernatant was transferred to a dried, pre-weighed aluminium dish and the weight was recorded. The solid content on a weight basis ($\%TS_{w/w}$) was calculated using E.1 and solids content in grams per litre (g/L) of by-product ($TS_{g/L}$) was obtained with E.3.

The remaining pellet after the centrifugation step was resuspended in 5 ml of ddH$_2$O and then centrifuged again at 4000 rpm for 10 min. The supernatant was removed and discarded. This step was repeated one more time and the pellet was resuspended in 10 ml of ddH$_2$O. The pellet was transferred to a dried, pre-weighed aluminium dish. The samples were dried at 105°C for 24 h. Similarly to the calculations of the supernatant solid content, solids content on a weight basis and in g/L was obtained by using E.1 and E.3 respectively.

3.2.3 Dry matter content (solid by-product samples)

Dry matter content of moist solid samples correspond to the dry weight of the whole liquid sample after drying it at 105°C for 24 hours.

This analysis was performed using 6 samples of solid by-products (i.e. DR, SG and TR). Between 5-30 grams of the samples were added to a dried, pre-weighed aluminium dish (Fisher, FB71085) and the weight was recorded. The dry matter content on a weight basis ($\%DM_{w/w}$) was calculated using E.4 and the dry matter content in grams of dried by-product per kilogram of wet by-product ($DM_{g/\text{kg}}$) was calculated with E.5. Symbols of the equations are the same used in section 3.2.2.1 explained above.
3.2.4 Densities (liquid by-product samples)

Similar to the procedure explained above in section 3.2.2, the density ($\rho$) of the liquid samples, expressed in g/L was calculated using E.6. Symbols of the equations are the same used in section 3.2.2.1 explained above.

\[ \rho = \frac{W_{WS}-W_{DD}}{10} \times 1000 \text{ (g/L)} \]

3.2.5 Cell count

The concentration of intact yeast cells in liquid by-product samples was determined by direct counting using a haemocytometer. Samples were gently agitated to ensure yeast cells were resuspended and then the samples were diluted with distilled-deionised water (ddH$_2$O) in a 1 in 10 proportion. Finally, the diluted samples were thoroughly mixed and the cells were counted and the results reported in number of cells per mL of sample.

3.2.6 pH analysis

Prior to pH analysis, the pH meter (Hanna Instruments, pH 201 microprocessor pH Meter) was calibrated with pH 4 and pH 7 buffers. The electrode probe was rinsed in distilled ddH$_2$O and then the probe was inserted into the sample and gently agitated. The reading was allowed to settle and then result was recorded.

3.2.7 Freeze drying

Total or fractionated (centrifuged supernatant and solids) samples were freeze-dried (Edwards Freeze Dryer Super Modulyo) for Total Nitrogen (TN) analysis (section 3.2.8) as the samples were too dilute for the TN detection.
A known volume of sample ($V_s$ in mL) was added to 100 ml pre-weighed boats. Samples were first frozen at -40°C and then left in the freeze-dryer (-60°C, $10^{-1}$ mbar) for at least 2 days or until most of the moisture content was removed. Afterwards, the samples were re-weighed, the solids content of the freeze dried samples in g/ L ($TS_{FD, \ g/L}$) was calculated with and the dry matter content of the freeze dried sample ($DM_{FD}$) was determined with E.8., by reference to the solid content of similar samples oven dried at 105°C (E.3.). Dried samples were ground with a mortar and pestle and used directly for TN Analysis. The freeze-dried samples were stored in a dessicator to avoid the samples absorbing atmospheric moisture and too keep them for long term storage. Symbols of the equations used are the same as in section 3.2.2.1 and other symbols were explained earlier in this paragraph.

\[
TS_{FD, \ g/L} = \frac{W_{DD+DS} - W_{DD}}{V_s} \times 1000 \quad (g/L)
\]

E.7.

\[
DM_{FD} = \frac{TS_{105^\circ C, g/L}}{TS_{FD, g/L}} \times 100 \quad (g/g)
\]

E.8.

3.2.8 Total Nitrogen Content (Kjeldahl Method)

Total Nitrogen (TN) content was determined by the Kjeldahl method using an automated system (Foss Tecator Kjeldahl). This method (Kjeldahl 1883) is suitable for solid samples containing 2-12% protein. Freeze-dried samples with known dry matter content were weighed (0.4-0.6 g) onto nitrogen-free paper (Pergamyn paper) and added to digestion tubes. Digestion, distillation and titration were performed according to the standard protocol. The samples were digested in concentrated $H_2SO_4$ with selenium catalyst. After cooling, the acid solution was diluted with water and made alkaline by addition of 10 M $NaOH$. The liberated ammonia was distilled into an excess of boric acid solution and titrated against 0.1 M $HCl$.

For liquid by-product samples, Total nitrogen content (TN $g/g$) was calculated using E.9. (and reported in g of Nitrogen per grams of dry sample), where T is the volume (reported in mL) of the 0.1 M $HCl$ utilised to titrate the distilled sample, $DM_{FD}$ the dry matter content of the Freeze dried sample (in reference to E.8 and reported in g/g) and $w_s$ is the weight of the freeze dried sample (reported in g).
Crude Protein content of liquid by-products (CP - reported in g of Protein per g of dried sample) was obtained by multiplying TN \( g/g \) by a factor (KF) as shown in E.10. KF values used for individual by-product samples are tabulated in Table 3-3.

TN and CP values in a g/ L basis (grams of nitrogen or protein per Litre of by-product) were obtained using E.11 and E.12, respectively. Where TS \( 105^\circ C, g/ L \) was defined and calculated previously in E.3.

TN and CP values for solid by-product samples were obtained in a similar way to liquid by-product samples. E.13 corresponds to Total nitrogen Content (gram per gram basis), E.14 is TN on g per kg of wet product basis and E.15 corresponds to the CP on a g per kg of wet product basis.

\[
\text{E.9. } \quad TN_{g/g} = \frac{T \times 1.4007}{w_s \times DM_{FD}} \times 100 \quad (g/g)
\]

\[
\text{E.10. } \quad CP_{g/g} = TN_{g/g} \times KF \quad (g/g)
\]

\[
\text{E.11. } \quad TN_{g/L} = TN_{g/g} \times TS_{105^\circ C, g/L} \quad (g/L)
\]

\[
\text{E.12. } \quad CP_{g/L} = CP_{g/g} \times TS_{105^\circ C, g/L} \quad (g/L)
\]

\[
\text{E.13. } \quad TN_{g/g} = \frac{T \times 1.4007 \times 100}{w_s} \quad (g/g)
\]

\[
\text{E.14. } \quad TN_{g/kg} = TN_{g/g} \times DM_{105^\circ C, g/kg} \quad (g/kg)
\]

\[
\text{E.15. } \quad CP_{g/kg} = CP_{g/g} \times TS_{105^\circ C, g/L} \quad (g/kg)
\]
Table 3-3. Kjeldahl Factors (KF) used for Crude Protein (CP) content calculations.

<table>
<thead>
<tr>
<th>By-product</th>
<th>KF</th>
</tr>
</thead>
<tbody>
<tr>
<td>CB-SG</td>
<td>6.25</td>
</tr>
<tr>
<td>CB-SH</td>
<td>6.25</td>
</tr>
<tr>
<td>CB-SY</td>
<td>6.25</td>
</tr>
<tr>
<td>GK-PA</td>
<td>6.25</td>
</tr>
<tr>
<td>GK-SG</td>
<td>6.25</td>
</tr>
<tr>
<td>HW-PA</td>
<td>6.25</td>
</tr>
<tr>
<td>NB-SW</td>
<td>5.75</td>
</tr>
<tr>
<td>SS-PA</td>
<td>6.25</td>
</tr>
<tr>
<td>WP-SG</td>
<td>5.75</td>
</tr>
<tr>
<td>WP-SY</td>
<td>6.25</td>
</tr>
<tr>
<td>WP-TR</td>
<td>6.25</td>
</tr>
</tbody>
</table>

3.2.9 Soluble Protein Content (Bradford Assay)

Bradford reagent was prepared by dissolving 100 mg of Coomassie Brilliant Blue G-250 (Fisher, BP100-25) in 50 ml 95% ethanol in a 1 L volumetric flask and then 100 ml of 85% phosphoric acid was added and finally distilled deionised water (ddH₂O) was supplemented to make up to 1 L. The reagent was be stored in the fridge and protected from light. Bovine Serum Albumin (BSA, 200 mg/ml protein standard; Fluka, P5369) was used for preparing the calibration standards. Diluted BSA (2 mg/ml in 0.15 M NaCl) was stored as 1 ml aliquots in eppendorfs in the freezer and diluted to 1 mg/ml with 0.15 M NaCl before preparing 100 µl standards as follows:

Samples and standards were analysed by adding 30 µl to a 2 ml eppendorf and adding 1.5 ml Bradford reagent, mixed by vortexing and then the absorbance was read at 595 nm between 2 min and 1 hour. All samples and standards were analysed in triplicate. Samples out of the range of the calibration standards were diluted and then re-analysed for the Bradford assay (Bradford 1976).
Table 2: BSA calibration standards for the Bradford assay.

<table>
<thead>
<tr>
<th>Standard BSA mg/l</th>
<th>BSA stock µl</th>
<th>Water µl</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>0</td>
<td>100</td>
</tr>
<tr>
<td>100</td>
<td>10</td>
<td>90</td>
</tr>
<tr>
<td>250</td>
<td>25</td>
<td>75</td>
</tr>
<tr>
<td>500</td>
<td>50</td>
<td>50</td>
</tr>
<tr>
<td>750</td>
<td>75</td>
<td>25</td>
</tr>
<tr>
<td>1000</td>
<td>100</td>
<td>0</td>
</tr>
</tbody>
</table>

3.2.10 Polyphenols content

Total polyphenols in the supernatant fraction was analysed according to the ASBC Method Beer-35 for total Polyphenol analysis of beer (ASBC 1976). Polyphenols react with ferric iron in alkaline solution and the red colour produced is measured at 600 nm.

The carboxymethylcellulose (CMC/EDTA) reagent (1% solution of CMC in 0.2% EDTA) was prepared by adding 10 g CMC (low viscosity, sodium salt of carboxymethylcellulose, Sigma C5678) and 2 g EDTA (disodium salt, Fisher, BPE120) to approximately 500 ml of distilled deionised water with stirring and left to mix for at least 3 hours until the CMC was completely dissolved. The solution was transferred to a 1 L volumetric flask and the remaining volume was made up with ddH$_2$O.

The Ferric reagent (3.5% green ammonium ferric citrate) was prepared by adding 3.5 g of Ammonium Iron(III) citrate (green, 16% Fe, Sigma 09713) to a 100 ml volumetric flask and made up to volume with ddH$_2$O and mixed. It was made sure that the solution was completely clear and a used within a week.

Samples were analysed in a fume hood. 10 ml of the sample were added to 8 ml CMC/EDTA reagent into a 25 ml volumetric flask and well mixed. Then 0.5 ml ferric reagent was added and again the solution was well mixed. Afterwards 0.5 ml ammonia reagent (Ammonium hydroxide solution, Sigma 320145) was added and the solution mixed thoroughly. Finally the volume of the solution was made up to 25 ml using ddH$_2$O and mixed once more. After 10 min the absorbance was measured in a 10 mm cuvette at 600 nm against a blank in a spectrophotometer (Jenway, Model 7305).
The blank was prepared by mixing 10 ml sample and 8 ml CMC/EDTA reagent in a 25 ml volumetric. Ammonia reagent was added (0.5 ml), mixed and the solution was made up to 25 ml with ddH₂O. The solution was allowed to stand for at least 10 min before analysing the blank in the spectrophotometer.

The Polyphenols concentration of the liquid by-product supernatant ([PP] reported in mg/ L) was calculated using E.16 where OD₆₀₀ is the Optical Density of the sample analysed.

\[
[\text{PP}] = \text{OD}_{600} \times 820 \quad \text{(mg/ L)}
\]

E.16.

3.2.11 Metal Content (Cu, Fe Zn, Mn)

Flame Atomic Absorption Spectroscopy (FAAS) was used to determine the concentration of Copper (Cu), Iron (Fe), Manganese (Mn) and Zinc (Zn). A Perkin Elmer AAnalyst 200 with an air/nitrous oxide flame and a multi-element (Cu, Fe, Mn and Zn) hollow cathode lamp were used for analysis (Perkin Elmer, NS3050212). This method determines the soluble metal concentration and samples must be free of solids. All samples were processed in triplicate. All glass and plastic ware were washed in acid (2% HNO₃) and rinsed three times with ddH₂O prior to its use.

3.2.11.1 Procedure for metal content analysis in liquid and solid phases

10 ml of the liquid samples were centrifuged at 4000 rpm for 10 min (Heraeus Multifuge 3SR). The Supernatant (SN) was removed and retained. This (liquid) fraction corresponds to the soluble metal in the by-product under analysis. The remaining material or pellet (P) was washed and resuspended in 5 ml of ddH₂O. Then the material was centrifuged (4000 rpm, 10 mins) and the liquid fraction was removed and retained. The washing step was repeated one more time and the liquid fraction of the washed material were combined. These combined washed fractions correspond to the loosely bound material (LB). The remaining pellet was resuspended in 1 ml of 6 M HNO₃ and transferred to a boiling tube were it was heated at 105°C for 1 hour on a heating block (Hanna Instruments HI 39800 COD Reactor).
The samples were allowed to cool and 6 ml of ddH₂O were added and mixed thoroughly. This material corresponds to the solid-bound (SB) fraction. Solids still present in the sample were centrifuged and only the supernatant was analysed to avoid blockage of the FAAS aspirator. To the liquid (SN) and wash fractions (LB) 200 µl of 69% HNO₃ (15.7M) were added and mixed thoroughly. The samples were stored in the fridge (4°C) for up to two weeks prior to analysis.

Standards for each element were prepared using 1000 mg/ L stock solutions of Cu (Fluka 38995), Fe (Fluka 16596), Mn (Fluka 77036) and Zn (Fluka 18827). Cu and Fe standards were in the range 0 to 5 mg/ L. Mn and Zn standards were in the range 0 to 2.5 mg/ L.

The metal concentration in each sample was determined directly using the method loaded on the Perkin Elmer A Analyst 200. A blank (ddH₂O) and the standards were first analysed, then individual samples, with a mean of three determinations reported in mg/l. Samples out of the range of the standards were diluted (with ddH₂O) and a dilution factor was included in the calculations.

The concentration of metal in the soluble liquid fraction ([SN]) was determined by FAAS with correction for sample acidification as required (i.e. samples diluted with 200 µl HNO₃). The concentration of metals in the loosely-bound ([LB] - reported in mg of metal per litre of by-product) and solid-bound ([P] - reported in mg of metal per litre of by-product) fractions were calculated using \[ E.17 \] and \[ E.18 \], respectively. In these equations A is the absorption reading from the FAAS (reported in mg/L), \( V_0 \) is the original sample volume (in mL), \( V_a \) is the diluted acid volume (in mL) and \( V_W \) is the combined volume of washed material (in mL).

\[
E.17. \quad [LB] = \frac{A \times V_w}{V_0} \quad (mg/ L)
\]

\[
E.18. \quad [P] = \frac{A \times V_a}{V_0} \quad (mg/ L)
\]
3.2.11.2 Total metal content analysis

The total metal concentration was calculated ([T]_c) as the sum of the concentration in the fractions detailed in section 3.2.11.1 above and explained in E.19. At the same time samples were analysed directly following acid digest of a sample. However in some occasions (samples with low metal concentration), the limit of detection of the FAAS was below the equipment capability. In this case the [T]_c value was preferred.

For total sample digestion, 0.6 ml of the liquid by-product sample and 0.4 ml of 69% HNO₃ was added to a boiling tube and then heated on the heating block at 105°C for 1 hour. Tubes were removed and allowed to cool. 6 ml of ddH₂O water were added and mixed thoroughly. Cu, Fe, Mn and Zn content was analysed by FAAS according to section 3.2.11.1 The concentration of each metal in liquid by-products ([T] - reported in mg per litre of by-product) was calculated using E.20. Symbols were described earlier in section 3.2.11.1

\[
[T]_c = [SN] + [LB] + [P] \quad \text{(mg/L)}
\]

E.19.

\[
[T] = \frac{A \times V_a}{V_0} \quad \text{(mg/L)}
\]

E.20.
3.2.12 Particle size analysis

A particle size analyser (Mastersizer 2000, Malvern Instruments) was used to measure particle size distribution of the Glenkinchie and Speyside pot ale and the spent wash samples. According to the ASTM method (ASTM 2009) and the equipment manufacturers’ guidelines, a few ml of each sample were placed into instrument and diluted with tap water to achieve an obscuration of less than 20%. Definitions of the parameters obtained with this analysis are presented in below:

Table 3-4. Definitions of the parameters for particle size analysis.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>D[4,3]</td>
<td>Mean diameter over volume (also called the DeBroukere mean)</td>
</tr>
<tr>
<td>D[3,2]</td>
<td>Volume/surface mean (also called the Sauter mean)</td>
</tr>
<tr>
<td>D(v, 0.1), D(n, 0.1)</td>
<td>10% of the volume (v) or number (n) distribution is below this value.</td>
</tr>
<tr>
<td>D(v, 0.5), D(n, 0.5)</td>
<td>Median diameter for a volume (v) or number (n) distribution, where 50% of the distribution (by volume or number) is above and 50% is below this value.</td>
</tr>
<tr>
<td>D(v, 0.9)</td>
<td>90% of the volume (v) or number (n) distribution is below this value.</td>
</tr>
</tbody>
</table>
| Span     | \[
\frac{D(v,0.9)-D(v,0.1)}{D(v,0.5)} \quad \text{or} \quad \frac{D(n,0.9)-D(n,0.1)}{D(n,0.5)}
\] |

3.2.13 Microscopic Imaging

Samples were suspended and then diluted in a 1 in 10 ratio with ddH₂O and placed under microscope (Axiophot, Zeiss) and the images processed with the ZEN software.
3.3 Results and Discussion

3.3.1 Solid content (liquid by-product samples)

The results of the solid content from liquid samples (pot ale, spent yeast, spent wash and trub) were summarised in Figure 3-1. The sample with the highest solid content was trub (163 g/L) and the lowest was the spent wash (43 g/L). Solids content in pot ale samples also showed a degree of variation. Pot ale from industrial sources (GK and SS) showed a solid content of around 50 g/L, while the pot ale obtained from Heriot-Watt University was around 100 g/L.

When the fractions were separated (solid and liquid), the liquid fraction showed a higher amount of solids than the liquid fraction in almost all the samples examined (the exception were spent yeast and spent wash). Supernatant from Glenkinchie pot ale contained approximately 80% of the totals solids. Solids from Speyside were difficult to quantify (less than 1 g/L).

![Figure 3-1. Dry weights of liquid by-products samples from breweries and distilleries.](image)

*P = Pellet or solid fraction*

*SN = Supernatant*

*T = Total (SN + P)*

Error bars in the graph describe the standard deviation of the triplicates.
Dry Matter content of solid by-product-samples

Dry matter content of solids by-products samples (spent grains and spent hops), shown in Figure 3-2, were in the order of 10% (hops) and 15-25% (grains). No substantial difference in the moisture (dry matter) content was observed between distilling (GK) and brewing (WP) grains. The lower value for the Caledonian Brewery could be explained by the sampling method and differences in the brewing process. Caledonian brewery uses a mash tun system, while Wellpark Brewery uses mash filter. It has been reported that mash filtration typically produces grains of lower moisture content than mash tun systems (Andrews et al. 2011).

SG = spent grains (draff)
SH = Spent Hops
Error bars in the graph describe the standard deviation of the triplicates.

Figure 3-2. Dry matter content of solid by-products samples from breweries and distilleries.
3.3.3 Densities, pH and cell count (liquid by-product samples)

The results are presented in the table below (Table 3-5). Densities for all by-products were close to 1 g/ ml. pH was not possible to record in all cases, but pH for pot ale sample were close to pH 4, which is in agreement with other literature sources. Same applied to cell count, the value were close to $10^8$ cells per ml.

Table 3-5. Densities, pH and cell count of liquid by-product samples from Breweries and Distilleries

<table>
<thead>
<tr>
<th>Origin</th>
<th>By-product</th>
<th>Average density (g/ml)</th>
<th>pH</th>
<th>Cell Count ($10^8$ cells/ mL)</th>
</tr>
</thead>
<tbody>
<tr>
<td>CB</td>
<td>SY</td>
<td>0.94 ± 2.11E-02</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>GK</td>
<td>PA</td>
<td>1.00 ± 7.45E-03</td>
<td>4.02</td>
<td>3.17</td>
</tr>
<tr>
<td>HW</td>
<td>PA</td>
<td>1.05 ± 7.55E-03</td>
<td>-</td>
<td>2.74</td>
</tr>
<tr>
<td>NB</td>
<td>SW</td>
<td>1.02 ± 1.72E-02</td>
<td>3.69</td>
<td>1.23</td>
</tr>
<tr>
<td>SS</td>
<td>PA</td>
<td>0.98 ± 5.53E-03</td>
<td>4.20</td>
<td>0.85</td>
</tr>
<tr>
<td>WP</td>
<td>SY</td>
<td>1.03 ± 4.49E-04</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>WP</td>
<td>TR</td>
<td>1.04 ± 2.68E-02</td>
<td>-</td>
<td>-</td>
</tr>
</tbody>
</table>
Crude protein content was analysed for the solid (pellet), liquid and the combined fractions (total) and the results were reported on a dry matter basis (Figure 3-3). The crude protein content for the combined fractions (total) was around 30% for the distilling by-products. Heriot-Watt pot ale and trub from Wellpark had a crude protein content of 16% and 14% respectively.

Protein content of the solids fractions (pellet) were around 25%-50%, with the exception of pot ale from Speyside (73%). This result might be misleading since the solid content from this source was lower than the other sources and not enough material was available to repeat the experiment.

Protein content from the liquid fractions (supernatant) was calculated by difference (mass balance) due to problems with some of the samples becoming too sticky after the drying process. This phenomenon might be related to the presences of carbohydrates in the samples. Carbohydrates analysis was not conducted in this chapter; however, in Section 9.3.5 carbohydrate from three pot ale samples were quantified obtaining values between 10-20 g/L, which are in agreement with the work carried out by Tokuda (Tokuda et al. 1998), discussed earlier in this report (Section 2.2).

A comparison of the protein content from the liquid fractions calculated by difference and experimentally is shown in Figure 3-4. Comparison between crude protein content obtained by mass balanced (SN(c)) and by experimentation (SN). No substantial difference can be observed between the methods, so it might be reasonable to use the calculated method (mass balance) to report the protein content of supernatants.
\[ P = \text{Pellet or solid fraction} \]
\[ \text{SN}(c) = \text{Supernatant (calculated). Difference between Total and Pellet fractions.} \]
\[ T = \text{Total (SN +P)} \]
Error bars in the graph describe the standard deviation of the triplicate.

**Figure 3-3.** Crude protein content (dry matter basis) of liquid by-products samples from brewery and distillery sources.

**Figure 3-4.** Comparison between crude protein content obtained by mass balanced (\( \text{SN}(c) \)) and by experimentation (\( \text{SN} \)).
Protein content of solid by-products was around 15-25% for the grains and 27% for the hops (Figure 3-5).

![Graph showing protein content of solid by-products](image)

G= grains, SH= Hops

Error bars in the graph describe the standard deviation of the triplicates.

**Figure 3-5. Crude Protein Content (dry matter basis) of solid by-products samples from breweries and distilleries sources.**

Protein content of the liquid by-products were also reported on an “as is” basis (Figure 3-6). The protein content in the samples varied from 7 g/L (SW-NB) to 40 g/L (SY-WP). Pot ale samples analysed had ~15 g/L of protein.

By analysing solid and liquid fractions separately (Figure 3-7), it was observed that in pot ale samples >70% of the proteins can be found in the liquid fraction. This result was quite important for future considerations regarding process design.
Figure 3-6. Crude Protein Content (as “is” basis) of liquid by-products samples from Breweries and Distilleries sources.

Figure 3-7. Distribution of protein content in solid and liquid fractions of liquid by-products samples from breweries and distilleries sources.
3.3.5  Soluble protein and polyphenols content

Soluble protein content of pot ale samples were in the range of 0.7-1.0 g/L. Spent wash and Heriot-Watt pot ale had lower protein content than Glenkinchie and Speyside pot ale.

It also important to highlight that protein content calculated with the Kjeldahl method was higher than the Bradford. For example, for GK-PA, considering only the liquid fraction, the protein content was 10 g/L, while with the Bradford method only 0.7 g/L were obtained (a ratio of ~14).

This discrepancy between the Kjeldahl and the Bradford methods, is that the latter responds better to proteins greater than 3 kDa and to peptides comprised of aromatic (phenylalanine, tyrosine and tryptophan) and basic (arginine, histidine and lysine) amino acids. The Kjeldahl method assumes that all nitrogen is protein and it is the standard method for protein analysis used in the food, feed and drink industries (Krohn 2001, Möller 2009).

<table>
<thead>
<tr>
<th>By-product</th>
<th>Soluble Protein Content (mg/L)</th>
<th>Polyphenols Content (mg/L)</th>
</tr>
</thead>
<tbody>
<tr>
<td>CB-SY</td>
<td>ND</td>
<td>ND</td>
</tr>
<tr>
<td>GK-PA</td>
<td>771</td>
<td>519</td>
</tr>
<tr>
<td>HW-PA</td>
<td>332</td>
<td>ND</td>
</tr>
<tr>
<td>NB-SW</td>
<td>350</td>
<td>ND</td>
</tr>
<tr>
<td>SS-PA</td>
<td>1,041</td>
<td>252</td>
</tr>
<tr>
<td>WP-SY</td>
<td>ND</td>
<td>ND</td>
</tr>
<tr>
<td>WP-TR</td>
<td>ND</td>
<td>ND</td>
</tr>
</tbody>
</table>
3.3.6 *Particle size analysis*

A full particle analysis of the spent wash and pot ale samples is presented over the following pages (Table 3-7, Figure 3-8 and Figure 3-9). Average particle size of pot ale samples were calculated between 3-5 microns (in agreement with reported values of yeast cells), while spent wash was 23 microns. The microscopic images confirm these results (Figure 3-10).

**Table 3-7. Particle size analysis of pot ale and spent wash samples.**

<table>
<thead>
<tr>
<th></th>
<th>GK</th>
<th>SS</th>
<th>SW</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>VOLUME DISTRIBUTION</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>D(v, 0.1)</td>
<td>µm</td>
<td>3.03</td>
<td>1.24</td>
</tr>
<tr>
<td>D(v, 0.5)</td>
<td>µm</td>
<td>5.34</td>
<td>6.04</td>
</tr>
<tr>
<td>D(v, 0.9)</td>
<td>µm</td>
<td>31.90</td>
<td>41.65</td>
</tr>
<tr>
<td>Span</td>
<td></td>
<td>5.411</td>
<td>6.684</td>
</tr>
<tr>
<td>Uniformity</td>
<td></td>
<td>1.492</td>
<td>2.002</td>
</tr>
<tr>
<td><strong>NUMBER DISTRIBUTION</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>D(n, 0.1)</td>
<td>µm</td>
<td>0.58</td>
<td>0.53</td>
</tr>
<tr>
<td>D(n, 0.5)</td>
<td>µm</td>
<td>0.76</td>
<td>0.69</td>
</tr>
<tr>
<td>D(n, 0.9)</td>
<td>µm</td>
<td>3.41</td>
<td>1.17</td>
</tr>
<tr>
<td>Span</td>
<td></td>
<td>3.47E+00</td>
<td>9.92E-01</td>
</tr>
<tr>
<td>Uniformity</td>
<td></td>
<td>9.47E-01</td>
<td>3.77E-01</td>
</tr>
<tr>
<td><strong>AVERAGES</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>D[3,2]</td>
<td>µm</td>
<td>4.77</td>
<td>3.48</td>
</tr>
<tr>
<td>D[4,3]</td>
<td>µm</td>
<td>11.77</td>
<td>15.35</td>
</tr>
</tbody>
</table>
Left side of the graph indicates the origin and the by-product type (i.e. GK-PA: Glenkinchie, Pot Ale)

Figure 3-8. Particle size distribution (volume) of liquid distilleries by-product samples.
Left side of the graph indicates the origin and the by-product type (i.e. GK-PA: Glenkinchie, Pot Ale)

Figure 3-9. Particle Size distribution (number) of liquid distilleries by-product samples.
GK-PA zoomed 40x (A1) and 100x (A2); SS-PA zoomed 40x (B1) and 100x (B2); and NB-SW zoomed 10x (C1) and 40x (C2).

Figure 3-10. Microscopic images of liquid distilleries by-product samples.
3.3.7  Metal content

The copper, iron, zinc and manganese content of pot ale and spent wash samples are presented in Figure 3-11. Due to the potential toxicity in feedstuffs (discussed in section 2.4), only copper content was considered in this section. Pot ale samples showed a higher copper content (2-2.5 mg/L) compared to spent wash (<1 mg/L). Another, important result to highlight, is that approximately 70% of the copper was found to be bound to the solids (yeast).

![Graphs showing metal content analysis](image)

Error bars in the graph describe the standard deviation of the triplicates.

Figure 3-11. Metal content analysis (Copper, Iron, Zinc and Manganese)
Since, the majority of the copper was bound to the yeast, solids separation can produce
a product a less toxic product and be incorporated in animal diets as discussed earlier in
the previous chapter.

Other minerals were not determined since only one cathode lamp was available.
However, the analysis of other metals such as calcium and sodium might be beneficial
too and could be incorporated in future work.
3.4 Conclusions

This chapter described analytical techniques used during the rest of the thesis and provided with some examples of the characterisation of by-products from brewing and distilling sources.

Of importance to the work developed through this project, protein from pot ale samples was quantified. Total protein content was calculated at 15 g/L, of which approximately 70% is found in the liquid fraction. The solid fraction was also found to have ~70% of the copper. This fact, that the liquid component of pot ale contains most of the protein and a fraction of copper, led to the decision of focusing future work on the liquid component of pot ale.

Additionally, the protein component seems to be large enough to warrant investigation of recovery, suggesting the possibility of PA as worthwhile protein source commercially.
CHAPTER 4 – PROTEIN EXTRACTION FROM YEAST USING
MECHANICAL AND ENZYMATIC METHODS

Abstract

A high pressure homogeniser and enzymatic methods were used to disrupt yeast cells from pot ale for the release of intracellular protein. 76 mg of soluble protein per gram of dry yeast was released at 800 bar after 10 minutes of recirculation. With a combined method (homogeniser + enzyme) 104 mg/ g were registered.

1 Part of this work counted with the collaboration of the MEng Chemical Engineering student Mrs. Barbara Kalek and the MSc student Sara Bages.
4.1 Introduction

From the previous chapter it was understood that pot ale contains approximately 1% (w/w) yeast with at least 50% protein content on a dry matter basis. However, these intracellular proteins need to be extracted in order to be suitable for feeding purposes or for the recovery of individual proteins.

This chapter aimed to develop a method for the release of soluble protein from yeast cells suspended in pot ale. Three different methods were tested: mechanical, enzymatic, and a combination of both methods.

For the mechanical method and the combined method a high pressure homogeniser was used. A theoretical review of large scale cell disruption methods is presented in the following section together with a review of yeast cell wall properties, as it is an important factor in yeast disruption.

It is also important to highlight the differences in the methodology of this work from the experiments carried out previously by the author (Traub 2011). In that work pot ale was passed through a high pressure homogeniser, whereas in this study, yeast was separated from pot ale supernatant, resuspended and then processed with mechanical and/ or enzymatic methods.
4.2 Literature review

4.2.1 Yeast cell wall

Four main functions have been identified for the yeast cell wall (Klis et al. 2006, Kollar et al. 1997): stabilization of internal osmotic conditions, protection against physical stress, maintenance of cell shape and as a scaffold for proteins.

The major components of fungal cell walls (*Saccharomyces cerevisiae*) are mannoproteins, glucans and chitin, in order in which they are found in the cell wall from the outside to the inside. A typical composition of cell wall is presented in Table 4-1 Figure 4-1 below and Figure 4-1 shows a diagram of yeast cell wall (Walker 1998).

Mannoproteins are mannans linked to proteins and its function seems to be as “filler”. Glucans are defined on the basis of the solubility and the three types of glucans reported in literature are presented Table 4-2 below. Chitin is a linear polysaccharide (of repeating N- acetylglucosamine units) and might be not essential for the mechanical strength of the lateral walls.

**Table 4-1. Major components of yeast cell wall**

<table>
<thead>
<tr>
<th>Component</th>
<th>% of wall mass</th>
<th>Function</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mannoproteins</td>
<td>30 – 50</td>
<td>Filler and Porosity of cell wall.</td>
</tr>
<tr>
<td>Glucans</td>
<td>35 – 55</td>
<td>Mechanical (rigidity, shape flexibility)</td>
</tr>
<tr>
<td>Chitin</td>
<td>1.5 – 6</td>
<td>No mechanical purposes</td>
</tr>
</tbody>
</table>

**Table 4-2. Glucan types found in yeast cell wall**

<table>
<thead>
<tr>
<th>Glucan type</th>
<th>Solubility</th>
<th>Function</th>
</tr>
</thead>
<tbody>
<tr>
<td>β (1-3)-glucan</td>
<td>Alkali and acid insoluble</td>
<td>Maintains wall rigidity and shape</td>
</tr>
<tr>
<td>β (1-6)-glucan</td>
<td>Alkali soluble</td>
<td>Links the components of the inner and outer wall</td>
</tr>
<tr>
<td>β(1-3)-glucan with 8-</td>
<td>Alkali soluble</td>
<td>Confers flexibility to the wall</td>
</tr>
<tr>
<td>12% β(1-6)-glucan</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
Figure 4-1. Composition and structure of the envelope of \textit{Saccharomyces cerevisiae} (Walker 1998)

4.2.2 \textit{Cell disruption}

For products such as enzymes and proteins which remain intracellularly (inside of the yeast), cell disruption is performed in order to release the target product. Methods to achieve cell disruption have been categorized as Mechanical and Non Mechanical Methods and summarised in Figure 4-2 below (Middelberg 1995).

Mechanical methods are more commonly used in large-scale operations, but disadvantages of its use include a release of contaminants and micronisation (fine cell debris). Non mechanical methods are generally used at the laboratory scale, because of economical limitations and potential inference in the target product (Middelberg 1995).
4.2.3 High pressure homogenizer

The High Pressure Homogenizer (HPH) is a widely used technique for cell disruption (APV 2008). A basic HPH design consists of a positive displacement pump that forces a cell suspension through the centre of a valve seat and radially across the seat face. Three different homogeniser valve designs are presented in Figure 4-3. The standard or conventional style is used for most emulsion and dispersion applications, and the CR and CD styles are used for cell disruption.

Several studies (Middelberg 1995) have examined the disruption of yeast cells by HPH which included the release of vitamins from beer during fermentation, the establishment of a cheap process for the manufacturing of single cell protein and enzyme release.
4.2.3.1 Mechanism of disruption

There is some controversy in literature about the explanation of microorganism disruption. General consensus is that impingement of cells on the impact ring is the major cause of disruption. Rapid release of pressure does not cause significant disruption of yeast. Normal stresses are only 20% as efficient as impact. Impact is an important factor in yeast disruption using HPH. (Middelberg 1995, E Keshavarz and Dunnill 1990).

Among other mechanism hypothesis cited in literature (Clarke et al. 2010) include: turbulence, cavitation, shear stress.

4.2.3.2 High pressure homogeniser model

Yeast disruption in a Manton-Gaulin homogenizer was modelled for the disruption of *S. cerevisiae* and soluble protein release was described by a kinetic-rate law (Doran 1995):

\[
\ln \left( \frac{R_m}{R_m - R} \right) = kNp^\alpha
\]

In Equation (1), $R_m$ is the maximum protein available for release, $R$ is the amount of protein release after $N$ passes through the homogenizer, $k$ is a temperature dependent rate constant and $p$ is the operating pressure. The exponent $\alpha$ is a measure of the resistance of the cells to disruption. *S. cerevisiae* has a value of 2.9, but this depends on the growth conditions. Cell concentration seems not to affect cell disruption.
Equation 1 suggests that protein release has a strong dependence on pressure. A smaller number of passes is preferred, because every pass increases micronisation. This phenomenon can cause problems in downstream processes. Pressure increment can, however, lead to temperature rise which cause protein deactivation. Typical operating pressures are in the range from 55 to 200 MPa and temperature increments are 2°C per 10 MPa.

Three to five passes are usually required to release more than 90% of the protein but fewer passes are typically used for practical and economical reasons. The incremental protein released by increased pressure or passes may not justify the cost.

4.2.4 Enzymatic Treatment

There are three enzymatic methods used in cell disruption: autolysis (Iida et al. 2008), phage lysis and foreign lytic enzyme. This document will focus on the latter method.

Due to the high specificity of enzymes, different kinds of enzymes are required for yeast cell wall degradation. A good disruption strategy requires degradation of the mannoprotein, glucans and the chitin. Most lytic systems for yeast include a protease and a glucanase (Crépisi et al. 1993, Iida et al. 2008, Salazar and Asenjo 2007).

One the most cited advantages of enzymatic treatment are its specificity and mildness. Enzymatic treatment is an alternative for process scale disruption, however its cost remain a concern. Additionally the high specificity of the enzyme could also mean a disadvantage since the treated yeast could have different properties depending on the growth stage and the yeast strain.

4.2.4.1 Enzymatic lysis model

Several papers prepared by Hunter (Hunter and Asenjo 1985, Hunter and Asenjo 1987a, Hunter and Asenjo 1987b, Hunter and Asenjo 1990, Hunter and Asenjo 1986) suggest two models of enzymatic lysis of yeast cells. A simplified two-step model, accounting for protein release at cell lysis followed by proteolysis, and a more complex mechanistic model which describes the removal of the two layers of yeast cell wall (glucans and mannoproteins) and the extrusion and rupture of the protoplast and organelles.
**Simple model**

The simple model treats cell lysis and proteolysis as single-step reactions in sequence. Both reactions are modelled with Michaelis-Menten kinetics. Yeast, the substrate of the first reaction, is particulate and the proteins are soluble. The different enzymes of the lytic system are grouped together into an all-inclusive single enzyme, bearing both the proteolytic and yeast lytic activities. All of the cell structures are also considered together as a unified yeast cell mass. When a cell is attacked by enzymes it is presumed to dissolve instantaneously, releasing its entire mass as soluble proteins, peptides and carbohydrates.

This model has two independent variables Y and E. The measured variables are P, S and C. Definitions of the variables are presented in Table 4-3 together with the parameters of the model. The values of the parameters can be found in Hunter’s work, but they will need to be checked for the experiments to be carried out if the yeast and enzyme concentration are out of the model’s specified range.

\[
\frac{dY}{dt} = -k_a(Y - Y_\infty) - \frac{k_p E \cdot (Y - Y_\infty)}{(Y - Y_\infty) + K_m} \quad (2)
\]

\[
\frac{dP}{dt} = -f_{py} \left( \frac{dY}{dt} \right) - \frac{k_p E \cdot P}{P + S + K_{mp} \left( 1 + \frac{Y}{K_i} \right)} \quad (3)
\]

\[
\frac{dS}{dt} = -f_{sy} \left( \frac{dY}{dt} \right) + \frac{k_p E \cdot P}{P + S + K_{mp} \left( 1 + \frac{Y}{K_i} \right)} \quad (4)
\]

\[
\frac{dC}{dt} = -f_{cy} \left( \frac{dY}{dt} \right) \quad (5)
\]

\[
\frac{Y_\infty}{Y_0} = a + bE + cY_0 + \frac{d}{Y_0} \quad (6)
\]

On the right hand side of Equation (2), the initial term represents autolysis and the second term, enzymatic lysis. Equation (3) describes protein breakdown by product-degrading proteases. The first term on the right hand side stands for protein released from lysing cells, and the second term, breakdown of the protein already in solution. Equation (4) shows that peptides are released from lysing yeast, but also arise from
breakdown of longer proteins \( (P) \). Since the protease activity against soluble proteins is considered non-specific, both long- and short-chains proteins will be attacked by the enzyme with essentially the same affinity per gram of substrate. Hence, the peptides \( (S) \) will act as a competitive inhibitor of the enzyme against \( P \), where the inhibition constant is equal to the Michaelis constant \( K_{\text{mP}} \). Carbohydrates release is shown in Equation (5).

Table 4-3. Variables and Parameters of the Simple Model

<table>
<thead>
<tr>
<th>Variables</th>
<th>Parameters</th>
</tr>
</thead>
<tbody>
<tr>
<td>( Y ) Yeast, mg/ l</td>
<td>( k_a ) Rate constant for autolysis</td>
</tr>
<tr>
<td>( Y_0 ) Original yeast concentration, mg/ l</td>
<td>( k_r ) Rate constant for lysis</td>
</tr>
<tr>
<td>( Y_\infty ) Ultimate yeast concentration, mg/ l; proportional to residual turbidity</td>
<td>( K_m ) Micheleais constant for lysis</td>
</tr>
<tr>
<td>( P ) Protein (TCA(^1) – insoluble), mg/l</td>
<td>( k_p ) Rate constant for proteolysis</td>
</tr>
<tr>
<td>( S ) Peptides (TCA – soluble), mg/l</td>
<td>( K_{\text{mP}} ) Micheleais constant, proteolysis</td>
</tr>
<tr>
<td>( C ) Carbohydrates, mg/l</td>
<td>( K_i ) Inhibition constant, proteolysis</td>
</tr>
<tr>
<td>( E ) Enzyme, % (v/v) of reaction mixture</td>
<td>( f_{py} ) Fraction of protein in yeast</td>
</tr>
<tr>
<td></td>
<td>( f_{sy} ) Fraction of peptides in yeast</td>
</tr>
<tr>
<td></td>
<td>( f_{cy} ) Fraction of carbohydrates in yeast</td>
</tr>
</tbody>
</table>

\(^1\) Trichloroacetic Acid

a, b, c, d are constants

**Structured Model**

This model considers lysis of the cell from the viewpoint of progressive breakdown of the cell structures, starting from the outer wall layer and progressing to the subcellular structures inside the protoplast. Here the cell is divided into four regions; the outer wall or wall protein; inner wall or wall glucan; the cytosol and the organelles, here grouped together as mitochondria. A schematic of the reaction pathways are shown in Figure 4-4.
4.2.5 Combined Methods

Both mechanical and non-mechanical methods have advantages and disadvantages as previously discussed. It makes sense, then to combine both techniques in order to maximize protein release at a lower cost. Two approaches of combined methods identified in literature (Doran 1995) include:

- Combination of chemical /physical/ enzymatic treatments
- Pre-treatment with a chemical, physical or enzymatic method followed by mechanical disruption

Pre-treatment methods were reviewed and specifically enzymatic pre-treatment followed by HPH of *S. cerevisiae* broths. In one study (Middelberg 1995), 2 hours of Zymolase pretreatment were combined with a Microfluidizer (4 passes at 95 MPa), which resulted in complete disruption. HPH without pretreatment showed 32% disruption and pure enzymatic treatment resulted in negligible protein release (5%).

In another study (Verduyn et al. 1999) the effect of a commercial baker's yeast treated with HPH and the protease papain was analysed. Isolated HPH treatment released a maximum of 30% of the solids and 34% of the total nitrogen (TN). After the papain treatment of the whole homogenized slurry, high yields of solids and TN (up to 81 and 85%, respectively) were obtained.
4.3 Methods and Materials

4.3.1 Pot ale samples and preparation of yeast suspension

Pot ale (Glenkinchie, UK) was centrifuged (Heraeus Multifuge 3SR at 4000 rpm for 10 minutes), the supernatant discarded and then the solids were washed three times in distilled deionised water (to remove extracellular protein) and finally resuspended in an appropriate buffer solution (pH depending on each experiment) to a concentration of approximately 10 g/L to mimic yeast suspension in pot ale and to avoid blockage in the homogeniser. The suspensions were then processed with enzymatic and mechanical methods for protein release.

4.3.2 Analytical methods

Protein release is reported in this document in mg of protein per dry weight of yeast. Exact dry weight concentrations were measured by drying 10 ml of the suspension in an oven at 105 °C for 24 hours and then recording the weight of the dried solids. Protein content analysis was determined by the Bradford microassay. All tests were performed in triplicate.

4.3.3 High pressure homogeniser

For the mechanical treatment experiments, a High Pressure Homogeniser (HPH) - model APV-1000 from APV Systems was used (Figure 4-5). The homogeniser was mounted with a CD (cell disruption) style valve. A diagram of the experiment can be seen in Figure 4-6.

Prior to the experiments, the HPH was flushed and cleaned with hot water. The water was then emptied from the HPH and 300 ml of a yeast suspension (yeast suspended in 0.2M sodium acetate + 0.2 M acetic acid, pH 4.5) was placed in to the HPH feed hopper.

The HPH pump was then started and the yeast suspension was recirculated for a few minutes at 0 bar, to ensure yeast particles were suspended in the buffer and not settled in the feed hopper. Subsequently, pressure was slowly increased (~1-2 minutes) to the specified pressure (200, 600 and 800 bar) and the suspension was recirculated up to ~15
minutes. ~10 ml samples were collected every ~2 minutes, cooled down, centrifuged and the supernatant analysed for protein content.

Figure 4-5. High pressure homogeniser used for the experiments.

Figure 4-6. High pressure homogeniser diagram.
4.3.4 Enzymatic treatment

Three different enzymes - Rohalase BX (AB Enzymes), Beta-glucanase (Bio-Cat Inc.) and Lyticase (Sigma Aldrich) were dissolved in appropriate buffer (recommended by the manufacturer) and mixed with 300 ml of yeast suspension prepared with the same buffer used for the enzyme solution (same pH). Enzyme dosage, temperature and pH were adjusted for each type of enzyme according to the manufacturers’ recommendation (Table 4-4). The experiments were conducted in a 1 L agitated vessel under constant pH and temperature conditions for 2 hours.

Samples were collected every 10 minutes for the first hour and then every 20 minutes until 120 minutes. Subsequently, samples were placed in a water bath at 85°C for minutes (to terminate enzyme activity), then cooled down to ambient temperature and then centrifuged (Heraeus Multifuge 3SR) at 4000 rpm for 10 minutes. The supernatant was assessed for protein content. For consistency of reporting, t=0 was defined as the time when the temperature reached the set point.

Table 4-4. Conditions used for the enzymatic treatment experiments.

<table>
<thead>
<tr>
<th>Enzyme</th>
<th>pH</th>
<th>Buffer</th>
<th>Temperature (°C)</th>
<th>Dose (µg)</th>
<th>Activity</th>
</tr>
</thead>
<tbody>
<tr>
<td>Rohalase BX</td>
<td>4.5</td>
<td>0.2M acetic acid-sodium acetate</td>
<td>45</td>
<td>100-300</td>
<td>At least 2,000 units/ mg of protein, with +20% protein by biuret</td>
</tr>
<tr>
<td>Beta-Glucanase</td>
<td>5.2</td>
<td>0.2M acetic acid-sodium acetate</td>
<td>37</td>
<td>100</td>
<td>3,000 units/g</td>
</tr>
<tr>
<td>Lyticase</td>
<td>7.5</td>
<td>0.2M Potassium phosphate</td>
<td>27</td>
<td>5.6</td>
<td>10,000 units/g</td>
</tr>
</tbody>
</table>
4.3.5 Combined method

Two experiments were carried out using the enzyme Rohalase BX as a pre-treatment before HPH disruption. 200 mg and 300 mg of the enzyme were added to the yeast suspension and mixed for 2 hours under controlled pH (4.5) and temperature (45°C) and then passed through the homogeniser. Protein release was monitored over time under different pressures (200, 600 and 800 bar).
4.4 Results and discussion

4.4.1 High pressure homogeniser experiments

The experiments performed confirmed the seemed theory outlines in Equation 1. The results presented in Figure 4-7, showed that an increase in the operating pressure and the recirculation time (number of passes) resulted in higher protein concentration in the supernatant of the yeast suspension.

In all experiments, it was observed that at \( t = 0 \) (beginning of the experiment), the protein content of the supernatant was well above 0 mg/L. This measurement could be attributed to the initial recirculation of the suspension and the time needed to reach the pressure set-point (1-2 minutes).

The experiment conducted at 200 bar, showed little protein release after 10 minutes (~18 mg of protein per gram of dried yeast). At 600 bar and 800 bar, 61 mg /g and 76 mg/g were achieved, respectively. After 10 minutes of recirculation, in the 800 bar experiment, protein release seemed to be constant.

![Figure 4-7. Protein release over time using a high pressure homogeniser.](image)

Error bars in the graph describe the standard deviation of the triplicates.
4.4.2 Enzymatic treatment experiments

A comparison chart of the protein release using the enzymes Rohalase (300 mg dose), Beta-Glucanase and Lyticase after 2 hours of treatment is presented in Figure 4-8. It is clear from the graph that Lyticase showed the highest amount of protein released (39.6 mg/g), with nearly 10 times more protein released than the other enzymes (3.96 mg/g for Rohalase and 1.85 mg/g for Beta-glucanase). It is also important to highlight that the Lyticase dosage was 2-6% of the other enzymes.

Error bars in the graph describe the standard deviation of the triplicates.

Figure 4-8. Enzymatic treatment experiments: comparison chart of protein release using the enzymes Rohalase (300 mg dose), Beta-Glucanase and Lyticase after 2 hours of treatment.
Based on the literature review previously presented, the factors that might have influenced the amount of protein released are the enzymatic activity and the specificity of the enzyme. In the first case, as reported by the enzyme manufacturer (Table 4-4), the enzymatic activity of Lyticase is about 40 times higher than Rohalase and 200 times higher than Beta-glucanase\(^1\).

The other factor that might have influenced protein release from yeast cells is the specificity of the enzymes. Lyticase and Rohalase are reported to have \(\beta\)-1,3 and \(\beta\)-1,6 glucanase and protease activity, while for the enzyme Beta-glucanase only beta-glucanase activity is reported (it is not specified what type of glucanase). For an effective yeast wall disruption the three types of enzyme activity are needed.

Protein release for each individual enzyme was monitored over time, and the results are presented in the following graphs. For the Rohalase experiments (Figure 4-9), no benefit (in terms of protein release) was found by increasing the enzyme dose (100 to 300 mg) and by extending the experiment to 120 minutes. For the Beta-glucanase experiments (Figure 4-10), a similar trend to the Rohalase experiments was found. Over time no significant increase in protein release was witnessed.

However, for the Lyticase experiments (Figure 4-11), protein release seemed to increase overtime and a maximum value of 40 mg/g was obtained. Experiments using a higher Lyticase dose were not continued due to economically non-viability, which is discussed later in this chapter.

\(^1\) Assuming that Lyticase activity is 2000 units per mg protein with 20% protein content, equivalent to 400 units per mg of product.
Error bars in the graph describe the standard deviation of the triplicates.

**Figure 4-9.** Protein release overtime using 100, 200 and 300 mg of Rohalase BX.

Error bars in the graph describe the standard deviation of the triplicates.

**Figure 4-10.** Protein release over time using 100 mg of Beta-glucanase.
Error bars in the graph describe the standard deviation of the triplicates.

**Figure 4-11.** Protein release over time using enzymatic treatment (Lyticase)

### 4.4.3 Combined method

Only the enzyme Rohalase was used for the combined experiments due to its lower cost compared to Lyticase and the superior results compared to Beta-glucanase obtained in the enzymatic experiments discussed earlier. Additionally Rohalase is extensively used in the wine industry, making it available as a food grade product.

A comparison of the HPH and the combined experiments is shown in Figure 4-12. At lower pressure (200 and 600 bar), an increase in protein release was observed by using a combined method compared to the pure HPH process. However the enzyme dose did not produce a significant difference in the results.

At 800 bar however, both utilisation and dose of enzyme showed higher protein release compared to pure HPH method. A 200 mg and 300 mg of enzyme dose increased the protein release by 22.6% and 37.8% equivalent to 93.2 and 104.4 mg/g, respectively. This result compared favourably to the work carried out previously with a sonicator (Iida et al. 2008), where 46.7 mg/g of protein was attained.
Figure 4-12. Comparison of protein release using a high pressure homogeniser and combined method with pre-enzymatic treatment (Rohalase BX with a 200 and 300 mg dose).

Protein release over time using a combined method (200 mg and 300 mg of Rohalase for 2 hours followed by HPH) are shown in Figure 4-13 and Figure 4-14. With 200 mg of enzyme at 200 and 600 bar, steady state seemed to happen in the first 2 min of HPH treatment, which suggest a more rapid disruption compared to pure HPH treatment (more than 12 minutes, Figure 4-7).

The combined method experiments showed that Rohalase might have a “weakening” effect on the yeast cell wall. The pure application of Rohalase released a small amount of protein, but when the enzyme treatment was followed by HPH, larger amounts of protein were released.
Error bars in the graph describe the standard deviation of the triplicates.

Figure 4-13. Protein release over time using a combined method (200 mg of Rohalase for 2 hours followed by HPH).

Error bars in the graph describe the standard deviation of the triplicates.

Figure 4-14. Protein release over time using a combined method (300 mg of Rohalase for 2 hours followed by HPH).
4.4.4 Economic analysis discussion

Although the exact price for the application of the enzymes on a large scale could not be obtained, suggested prices are presented in Table 4-4 together with the estimated processing cost. These values were obtained by extrapolating the results discussed earlier and range from £10-70 per g protein. Typical feed protein prices are £1-£2 per kg (see Chapter 2), making the enzymatic treatment process unfeasible for the feed market.

Table 4-5. Estimated processing cost using enzymatic treatment

<table>
<thead>
<tr>
<th>Enzyme</th>
<th>Enzyme used (mg)</th>
<th>Protein released (mg)</th>
<th>Enzyme price(^1) (£ per g enzyme)</th>
<th>Cost (£ per g protein)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Rohalase</td>
<td>100</td>
<td>11</td>
<td>1</td>
<td>10</td>
</tr>
<tr>
<td>Beta-glucanase</td>
<td>100</td>
<td>5.6</td>
<td>1</td>
<td>18</td>
</tr>
<tr>
<td>Lyticase</td>
<td>5.6</td>
<td>120</td>
<td>1400</td>
<td>66</td>
</tr>
</tbody>
</table>

The potential economic advantages in the utilisation of a pre-enzymatic treatment can be translated in lower processing costs: reduced recirculation times (number of passes) and working pressure. For downstream processes lower micronisation could lead to the design of a smaller centrifuge.

\(^1\) Enzyme prices were estimated from manufacturer’s websites and interviews.
4.5 Conclusion

HPH is a method that can be used to extract intracellular proteins from yeast suspended in pot ale. Pressure and processing time increase protein release, although beyond 10 minutes of processing, protein release seemed to cease. On the other hand higher pressure could have implications to downstream processes by increasing (yeast) particle micronisation, thus impacting the overall processing costs.

As an alternative to HPH, enzymatic treatment was studied. Only one of the enzymes tested showed (pot ale) yeast wall lytic potential. However, enzyme cost and a lower protein release compared to HPH, might not allow scaling the process up by utilising purely enzymatic treatment. Finally, a combined method (enzyme +HPH) was studied. With this method, at least a 20% increase to pure HPH was achieved.

Although the process economics were not studied in depth, the release and recovery of proteins from pot ale yeast cells seems to be incompatible for the feed market. Perhaps the proteins recovered could be used in higher value protein markets, but more research need to be conducted. In order to minimise cost, an enzyme recovery step could be added, however, the cost effectiveness remains at this stage unknown.

Furthermore, the effect on the enzymes in the final product is also uncertain, like product contamination (i.e. toxicity to salmon) and the effect on protein functionality.
CHAPTER 5- SOLID- LIQUID SEPARATION OF POT ALE: A SCALE-UP ANALYSIS

5.1 Introduction

Protein concentration using ion exchange chromatography (explained later in Section 9) requires solid particiles to be removed from the liquid phase. Generally, centrifuges are used in large scale processes for the separation of suspended solids from liquids.

The Department of Chemical Engineering from the School of Engineering and Physical Sciences of Heriot-Watt University holds a process scale disc stack centrifuge (GEA-Westfalia model SC-6) that potentially could be used for future pilot and/ or commercial trials in an average size malt whisky distillery.

In this Chapter a method to predict the throughput and clarification of pot ale using a large scale disc stack centrifuge was developed. The method is based on scale down models (Boychyn et al. 2000, Tustian et al. 2007).

Additionally disrupted yeast cells in pot ale were assessed for clarification tests in order to understand the effect on particle size and clarification requirements.
5.2 Centrifugation theory

5.2.1 Classification of centrifuges

Centrifuges could be classified into sedimentation and filter centrifuges. In sedimentation centrifuges, solids are transported to the periphery wall of the rotating machine bowl and collected against this surface; liquid is removed from the solids by the close packing of the individual particulates (Brunner and Hemfort 1988, Don and Robert 2008). In filter centrifuges the solids are transported to the surface of a filter element and the solids trapped on this filter, while the liquid drains through the particulates and exits through the filter surface. Another way of classifying centrifuges in based on its operation mode: continuous or batch mode. Generally, continuous centrifuge are often considered for large scale solid-liquid separations. Examples of centrifuge types with approximate capabilities and range of g forces available from each machine type are shown in Figure 5-1 (Beveridge 2000).

\begin{figure}[h]
\centering
\includegraphics[width=\textwidth]{figure5-1.png}
\caption{(A) and (B) Approximate capabilities of various centrifuge forms to sediment/separate particles and levels of suspended solids applicable. NA, Not applicable or not relevant (generally less than 1500 g).}
\end{figure}

\textbf{Figure 5-1.} Centrifuge types with approximate capabilities and range of g forces (Beveridge 2000).
5.2.2 Disc stack centrifuges

Disc-stack centrifuges (DSC) are widely used in the brewing and biotechnology industry for the recovery of cells and cell debris (Boychyn et al. 2000). DSCs key features are a central stack of conical discs, in which most of the separation occurs, and its ability to perform intermittent solids discharge (Figure 5-2). It usually achieves good clarification but relatively poor dewatering compared with solid-bowl centrifuges such as the tubular bowl or multichamber bowl (Boychyn et al. 2004).

Figure 5-2. Bowl section of a self-cleaning disc stack centrifuge (Beveridge 2000).
5.3 Theoretical considerations

Centrifugation performance can be described using the Sigma (Σ) concept of equivalent settling area (Doran 2012) through continuous (Q) or batch flowrate (V/t), and a correction factor (c) to allow for deviations from ideal flow. For a laboratory batch centrifuge with a swing-out rotor, the settling area \( \Sigma_{\text{lab}} \) can be calculated using Eq. 1, where \( V_{\text{lab}} \) is the volume of material in the tube, \( \omega \) is the angular velocity, \( R_1 \) and \( R_2 \) are the inner and outer radii, and \( x \) and \( y \) are the fractional times required for acceleration and deceleration, respectively (Tustian et al. 2007).

$$\Sigma_{\text{lab}} = \frac{V_{\text{lab}} \omega^2 (3-2x-2y)}{6g \ln \left( \frac{2R_2}{R_2+R_1} \right)}$$

Eq. 1

For a disc-stack centrifuge, the settling area \( \Sigma_{\text{ds}} \) can be calculated using Eq. 2 where \( N \) is the number of discs, \( \theta \) is the half disc angle, \( r_1 \) and \( r_2 \) are the respective inner and outer radii of the discs.

$$\Sigma_{\text{ds}} = \frac{2\pi N \omega^2 (r_2^3-r_1^3)}{3g \tan \theta}$$

Eq. 2

A comparison of centrifuges of different designs and sizes is achieved with Eq. 3

$$\frac{Q_{\text{ds}}}{c_{\text{ds}} \Sigma_{\text{ds}}} = \frac{V_{\text{lab}}}{t_{\text{lab}} c_{\text{lab}} \Sigma_{\text{lab}}}$$

Eq. 3

Eq1., Eq2, and Eq.3 assume low particle concentration (no hindered settling), laminar settling of the particle, i.e. \( \text{Re} = \frac{v_s \rho \ d}{\mu} < 0.5 \), where Re is the Reynolds number, \( v_s \) is the particle settling velocity, \( \rho \) is the suspension density, \( d \) is the particle diameter, and \( \mu \) is the suspension viscosity, and particles start at half the settling distance will just be removed from the suspension. Eq. 3 should only be used in the linear region plot of % clarification against \( Q/c\Sigma \) on probability–logarithmic axes.
The particle size settling velocity $v_s$ can also be determined by Stoke’s law (Eq.4), where $g$ is the acceleration gravitational constant ($9.8 \text{ m/s}^2$).

$$v_s = \frac{d^2(\rho_s - \rho_f)g}{18\mu} \quad \text{Eq. 4}$$

5.4 Methods and Materials

Pot ale samples from Glenkinchie (GK), Spey Side (SS) and high pressure homogenised Glenkinchie (GK-HPH) sample were used for the experiments. 10 ml of pot ale was centrifuged (Heraeus Multifuge 3SR) for several minutes and angular velocities (100, 300 and 480 radians per second). The samples were then assessed for clarification using Eq. 5 by optical density (OD) at 670 nm with a spectrophotometer (Jenway 7315 Spectrophotom), where the suffix of OD indicates the feed (f), supernatant (s) and reference (r). OD$_r$ was determined by running the samples in a Minifuge (Eppendorf) centrifuge at maximum speed ($14000$ g) for 10 min.

$$\%C = \left(\frac{\text{OD}_f - \text{OD}_s}{\text{OD}_f - \text{OD}_r}\right) \cdot 100 \quad \text{Eq. 5}$$

A high pressure homogeniser (HPH) - model APV-1000 from APV Systems- was used to disrupt the yeast cells. Approximately 1 L of pot ale was introduced into the homogeniser and recirculated for 12 min at 800 bar to achieve complete cell disruption. The homogenised samples were diluted in a 1:1 proportion using distilled-deionised water to ensure that all readings were in the linear range of the spectrophotometer (0.05 to 1.0). All the experiments were performed in triplicate.

The disc stack centrifuge for the process scale calculations was a GEA-Westfalia model SC6 with nominal capacity of 3 m$^3$/h and capable of centrifugal speeds between 8000 and 12000 rpm. Geometrical properties and other characteristic of this centrifuge are presented in Figure 5-3 and Figure 5-4. For the scale-up calculations the parameters used are in Table 5-1 and Table 5-2.
Table 5-1. Parameters used for scale-up calculations.

<table>
<thead>
<tr>
<th>Number of discs (N)</th>
<th>81</th>
</tr>
</thead>
<tbody>
<tr>
<td>Outer diameter ($r_2$)</td>
<td>124 mm</td>
</tr>
<tr>
<td>Inner diameter ($r_1$)</td>
<td>68 mm</td>
</tr>
<tr>
<td>Half angle ($\theta$)</td>
<td>50°</td>
</tr>
</tbody>
</table>

Table 5-2. Parameters used for lab scale calculations

<table>
<thead>
<tr>
<th>$V_{lab}$</th>
<th>10 ml</th>
</tr>
</thead>
<tbody>
<tr>
<td>$R_2$</td>
<td>192 mm</td>
</tr>
<tr>
<td>$R_1$</td>
<td>110 mm</td>
</tr>
<tr>
<td>Acceleration (x)</td>
<td>45 s</td>
</tr>
<tr>
<td>Deceleration (y)</td>
<td>45 s</td>
</tr>
</tbody>
</table>
Technical data

Bowl
Speed variable 8000 to 12 000 min⁻¹
Total bowl volume 1.8 l
Volume of solids holding space 0.7 l
Max. discharge pressure of centripetal pump approx. 5 bar

Three-phase AC motor
Power rating 5.5 kW
Speed (FC operation) 1800 min⁻¹
Design IM B5
Enclosure IP 55

Weights
Separator complete approx. 230 kg
Bowl approx. 50 kg

Capacity
Rated capacity 3000 l/h

The rated capacity indicates the maximum throughput capacity of the bowl. The effective capacity is normally lower. It depends on the product and the desired clarifying efficiency.

Figure 5-3. Technical data of the disc stack centrifuge (GEA-Westfalia. model SC6) for upscale calculations.

Figure 5-4. Cross section of the GEA-Westfalia model SC6 centrifuge.
5.5 Results and discussion

The results of the clarification tests using a laboratory batch centrifuge were plotted on probability–logarithmic axes as shown in Figure 5-7. The use of these scales on the graph axis allows an easier visualisation of the results and the linearization of the curves.

The correction factor $c_{\text{lab}}$ used was 1.0 (Boychyn et al. 2000). From this graph (Figure 5-7), it can be determined that for the same level of clarification, disrupted samples require a higher area because slower settling rates are achieved (Stoke’s law).

For example, to achieve a 90% clarification, at least one order of magnitude was detected between the GK (4.91x10^{-7} m/s) and the GK-HPH samples (5.38x10^{-8} m/s). GK and SS samples require areas in the same order of magnitude. The settling velocities for a 90% clarification level of SS samples were 2.50 x10^{-7} m/s.

To determine if the process scale centrifuge was adequate to separate yeast particles for a medium size distillery (average pot ale flow rate of 3 m$^3$/h), the $Q_{\text{ds}}$ was calculated using Eq. 3. The right hand side of the equation was obtained using Figure 5-7 for different clarification levels (85%, 90%, 95% and 99%). Since the $\Sigma_{\text{ds}}$ depends on the rotational speed of the centrifuge, $Q_{\text{ds}}$ was plotted against this parameter. The correction factor $c_{\text{ds}}$ used for the calculations was 0.4 (Boychyn et al. 2004)

From the graphs in Figure 5-8 it can observed that the disc stack centrifuge under consideration can theoretically operate at the nominal capacity (3 m$^3$/h) and produce acceptable clarification levels (>90%) for the GK and SS samples. For the GK-HPH sample reduced, clarification levels below 90% can be achieved by operating the centrifuge at lower flowrates and the highest rotational speed. Potential consequences of this operational strategy (low flow rate, high centrifugal speed) would require more energy and increased cooling requirements to avoid overheating of the centrifuge.

Another aspect to consider with the GK-HPH is the cost and sizing of filtration steps downstream of the centrifuge. Conventional large-scale disk-stack centrifuges, as the one reviewed in this Chapter are not effective in the removal of particles less than 1 micron. As particles decrease in size, removing them efficiently decreases
exponentially. A study by Kempken (Kempken et al. 1995) showed that high numbers of small particles (ranging from 1 to 6 microns) were not effectively removed by the centrifugation process. Because of this small-particle loading may have detrimental effects on downstream chromatography or other purification, it is suggested that at least one secondary clarification step might be used after the centrifuge. Centrifuges can be applied to the removal of very high cell densities, but their use as the sole clarification step is often limited (Yavorsky et al. 2003).

Since the GK-HPH sample would have more debris and suspended material than non-disrupted (yeast) pot ale, the replacement requirements of filters need to be increased, thus increasing the cost of the operation.
Figure 5-5. Clarification vs time chart of the GK sample.

Figure 5-6. Clarification vs. $V_{lab}/t_{lab}C_{lab} \Sigma_{lab}$ chart of the GK sample.
Figure 5-7. The probability–log relationship of percent clarification and equivalent flow rate per centrifuge separation area for yeast particles in pot ale samples.
Figure 5-8. Theoretical flowrate of a disc stack centrifuge against rotational speed and clarification level for a Glenkinchie (GK), Speyside (SS) and a high pressure homogenised (GK-HPH) pot ale sample.
5.6 Conclusion

Based on theoretical considerations, the studied disc stack centrifuge (GEA Westfalia SC6) is an adequate choice of equipment, capable of separating yeast cells from pot ale for further processing at the required flow rates supplied by an average size distillery. However, the installation of a downstream filtration step is necessary in order to avoid damaging effects to downstream chromatography or protein purification steps.

The addition of a disruption step upstream of the solid/liquid separation increases the required settling area by a factor of 10. This of course, raises the capital and operational requirements for the separation step. A further economic analysis needs to be carried in order to understand cost/benefits of cell disruption.

In the next chapters of this thesis, the focus will be on the liquid stream of pot ale, i.e. the material that has been centrifuged and/or filtered. From Chapter 6 and Chapter 10 the focus will be on protein concentration and purification methods, particularly ion exchange chromatography. It is also important to highlight that the discarded solids (yeast) from the centrifugation step might have some economic value and further scientific and commercial research is advised in order to maximise the profits of a potential large scale operation for the recovery of protein from pot ale.
CHAPTER 6 – PRELIMINARY STUDIES OF POT ALE PROTEINS CONCENTRATED AND PURIFIED WITH COMMERCIALLY AVAILABLE RESINS USING ION EXCHANGE CHROMATOGRAPHY

Abstract

In this chapter, two sets of experiments were conducted in order to prove that liquid chromatography is a suitable technique to purify and concentrate proteins contained in pot ale. The experiments were carried out using 1 ml columns with commercially available chromatography media, one a cation exchanger (HiTrap Capto S) and the other an anion exchanger (HiTrap Capto Q).

In the first set of experiments, eight trials were conducted. For each column, 50 ml of solids free pot ale was loaded into the column at a loading rate of 1 ml/min. Proteins were then eluted with a 0.5M NaCl gradient over 10 column volumes at four pH conditions (4.5/ 5.8/ 7.2/ 10.1). Elution, washing and equilibration pH conditions were the same for each run. Protein binding was assessed by measuring the areas of the peaks during elution. Maximum protein binding was achieved at a pH value of 4.5 with a HiTrap Capto S column. Overall, the cation exchanger showed higher binding affinity than the anion exchanger and binding capacity decreased with higher pH for the HiTrap Capto S column while the opposite was observed with the HiTrap Capto Q column.

In the second set of experiments, 200 ml of clarified pot ale was loaded into a HiTrap Capto S column and then the proteins were eluted at 4.5 pH using 5 CV steps of 0.2M NaCl increments until reaching 1M NaCl. Three peaks were detected (at 0.2M NaCl, 0.4M and 0.6M NaCl). Properties of the peaks were calculated (area, height, width, resolution and asymmetry) were analysed. Peak area and height decreased with the increased salt concentration while the opposite was observed with the peak width.

An SDS-page analysis was also included, which confirmed the separation of Protein Z and LTP1 from other pot ale components using the HiTrap Capto S column media.
**6.1 Introduction**

Ion Exchange Chromatography (IEC) was evaluated to concentrate protein contained in pot samples. IEC works on the principle of reversible interaction between a charged protein and an oppositely charged chromatography medium (Doran 2012). A theoretical review of this method for protein concentration is covered later in this chapter.

No evidence could be found in literature of using IEC for protein recovery in whisky by-products, specifically pot ale. In the process described in patent GB2094084, pot ale was treated with a caustic material (NaOH, KOH or ammonium hydroxide) to raise the pH causing protein precipitation. The protein precipitate was then allowed to settle and recovered by mechanical means. This method requires a large amount of caustic material which translates into high processing costs. Additionally, a low protein recovery yield was reported (50-60%) (Gilmour et al. 1982).

Ultrafiltration (UF) is another method used for the concentration and purification of proteins (van Reis et al. 2007). There is however, no evidence in literature of UF used in the whisky industry for the recovery of proteins. It has been reported that the Glenllachie Distillery in the North east of Scotland commissioned a waste water system in 2008 compromising membrane bioreactors and cross-flow ultrafiltration technology (Robinson 2009). Another mention of UF technology used in the whisky industry can be found in patent WO 2005/113118 A2 (Peyton et al. 2007). In both cases UF was mentioned as a method for water purification rather than protein recovery.

In the dairy industry, protein recovery from by-products is a well-established process. In a work presented by Daufin et al (Daufin et al. 2001), membrane technology was reported to be a major tool in food processing. This research mentioned that during the 1980’s and 1990’s considerable interest was put on the development of extraction procedures for milk proteins.

Fractionation of the main whey protein was first attempted more than 25 years ago by exploiting the low heat stability of calcium free $\alpha$-lactalbumin. During recent years this process has improved. Removal of calcium from $\alpha$-lactalbumin is obtained by a pH reduction (to about 3.8) or by the addition of a sequestering agent such as citric acid or sodium citrate, resulting in much reduced thermal stability compared to the native
protein. Subsequent heating around 55°C for a limited period of time leads to a reversible and partially denatured form, which undergoes aggregation.

Other processes were also developed from this property of α-lactalbumin. One of these processes uses ultrafiltration, pH adjustment (~4.2) and thermal treatment. This process allows also immunoglobulin and the serum albumin together with α-lactalbumin to precipitate. Further separation is achieved with a clarifier and ultrafiltration.

A recent analysis of the recovery of organic waste in the Spanish wine industry by Ruggieri et al (Ruggieri et al. 2009) find out that some of these wastes were used as by-products whereas the rest were traditionally incinerated or disposed in landfill. This work proposes composting for the recovery of stalk and wastewater sludge to produce sanitised organic amendment for the application of vineyard.

A major disadvantage of UF against IEC is that IEC can selectively separate proteins whereas UF would not be as selective. In UF, depending on the molecular weight cut-off (MWCO) other pot ale components in addition to the proteins will be concentrated as well. This argument, was however challenged in a work from van Reis (van Reis et al. 1997), where UF membranes were used for protein purification using a process known as high performance tangential flow filtration (HPTFF). In bioprocessing HPTFF can potentially be used throughout the purification process to remove specific impurities and/or eliminate protein oligomers or degradation products. In addition, HPTFF can effect simultaneous purification, concentration, and buffer exchange, providing an opportunity to combine several different separation steps into a single scalable unit operation.

Another consideration of UF and precipitation is that these processes are volume dependent separation methods. This means that capital and manufacturing cost are proportional to the volume of solution processed and not to the quantity of product produced (Lightfoot et al. 1987). For dilute protein solutions, such as pot ale (typically ~1% protein content), a large volume must be processed to recover a fixed amount of protein. IEC in contrast is less volume dependent, because its capacity depends mostly on the mass of protein adsorbed, not the volume of liquid (pot ale) processed. In an example found in literature (Etzel 2004), the recovery of 1 kg of lactoferrin from whey
using UF and IEC were compared. In both processes 10,000 L of whey (concentration of lactoferrin in whey is 0.1 g/L) were required. In this comparison, it was highlighted that fluxes of the IEC process were ~300 greater than UF.

In summary, for future work in this thesis and further process development, IEC was chosen as a protein recovery method (purification and concentration). There are technical, economical and other reasons (novelty for the whisky industry) that support this decision in favour to other methods mentioned earlier (UF and precipitation). However, these methods could be integrated in downstream processes and should not be discarded.

An initial step in protein purification using IEC involves the selection of the chromatographic medium. Variables such as surface charge (cationic or anionic), pore size and chemical stability need to be tested in order to achieve maximum protein recovery and throughput. In the experiments presented in this chapter, pot ale samples were loaded into cationic and anionic media and then the proteins bound to the media were eluted at different pH conditions.

Two sets of experiments were conducted. The first experiments aimed to determine which column media (cation or anion exchanger) achieves maximum protein binding and at what pH conditions. The second experiment - based on the results of the first experiment- was designed to identify proteins typically present in pot ale using a SDS-page analysis.
6.2 Theoretical Background

6.2.1 Ion exchange Chromatography

Ion exchange chromatography is based on the binding of charged sample molecules to oppositely charged groups attached to an insoluble matrix. Substances are bound to ion exchangers when they carry a net charge opposite to that of the ion exchanger. This binding is electrostatic and reversible. The pH value at which a biomolecule carries no net charge is called the isoelectric point (pI). When exposed to a pH below its pI, the biomolecule will carry a positive net charge and should bind to a cation exchanger. At pH's above its pI the biomolecule will carry a negative net charge and should bind (under appropriate conditions) to an anion exchanger.

Figure 6-1. Effect of pH on protein net charge (GE-Lifesciences).
6.2.2 Chromatography techniques

A typical chromatography operation involves five consecutive steps assuming: column equilibration, sample loading (adsorption), washing, elution and column regeneration. Typically a down flow pattern is used, at a flowrate appropriate to the column’s hardware specifications (pressure). To minimise peak width, the elution step is carried out lower flowrates. (Willoughby 2002).

In the column equilibration step, a buffer is pumped through the column in order to allow and maximise effective binding to the target molecules, i.e. proteins. Typically salt (NaCl) or acids are used.

The sample is usually pumped (downwards) through the column and the length of the load period will depend on the conditions required. In most cases samples will be loaded until a certain percentage breakthrough (typically 10%) of product is detected.

During the washing step, the same buffer used for the equilibration step is run through the column until all unwanted or loosely bound material is removed from the system.

After washing, during the elution step the required products and any remaining contaminants are removed from the column. In the event of any other contaminant being still present on the column then a second elution or multiple elutions are carried out. Typically pH changes or salt gradients are utilised. To ensure that the column is clean, usually NaOH solution (1M) is used, although attention must be paid to resin or media stability.

If the column will be used for another cycle, the equilibration buffer is pumped through the column once again to regenerate the column. If the column will not be used immediately, the column is rinsed with water and stored in 20% ethanol or sodium azide solution to prevent bacterial growth.
6.2.3 Peak parameters

The definitions of peak parameters (width, width at half height, resolution and asymmetry) are based on Figure 6-2 below.

![Figure 6-2. Typical peak shapes observed in a chromatogram (GE Healthcare Lifesciences).](image)

6.2.3.1 Width

Difference in retention between the peak end and peak start, time or volume base (G-A in the diagram above).

6.2.3.2 Width at half height

Calculated by taking the maximum height of the peak above the baseline, then determining the peak width at half this value above the baseline. Time or volume base. (B-D in the diagram above, where BD bisects CF).

6.2.3.3 Resolution

Resolution can be affected significantly by peak sizes and shapes. The resolution algorithm utilised was:

\[((V_{R2} - V_{R1}) / (2 \times (W_{h2} + W_{h1}))) / 2.354\]
Where \( V_{R1} \) and \( W_{h1} \) are the retention, width, Sigma and width at half height of the previous peak. \( V_{R2} \) and \( W_{h2} \) are the retention and width at half height of the current peak.

### 6.2.3.4 Asymmetry

Peak Asymmetry is simply the ratio of the segments B and A (Asymmetry = \( B / A \)) from Figure 6-3. Where A and B are partial peak width, measured at a 10% the peak height.

![Figure 6-3. Asymmetry ratio (GE Healthcare Lifesciences).](image-url)
6.2.4 Protein profile of pot ale

There is little information about protein profile of pot ale. However, more information can be found about beer. Since similar raw materials are used during the production process of whisky and beer, it can be assumed that pot ale and beer have similar protein profiles. The proteins in beer mainly originate from the barley, which are extracted and undergo modification during malting, mashing, boiling, fermentation and beer clarification steps. Proteins from yeast have been identified although these are present in much lower concentration (Colgrave et al., 2011). The main differences in whisky and beer production is that in brewing, proteins are precipitated from the wort during boiling with hops as the hot break or trub and during cooling as cold break. Many of the large barley storage proteins and in particular hordeins are precipitated and removed during the mashing and boiling stages (Picariello et al., 2011). During wort boiling, proteins are also modified by glycation, acylation and denaturation reactions. This step is omitted in the malt whisky process, although similar high-temperature modifications of soluble proteins may occur during distillation. The boiling step in brewing also deactivates malt enzymes whereas in the whisky process proteolytic enzymes remain active during fermentation allowing continual modification of barley proteins in the wash.

Beer contains a variable amount of protein (20-600 mg/l) with molecular masses ranging from 5 kDa to greater than 100 kDa (Mainente et al., 2011). These proteins are partially glycated through the Maillard reactions that occur during malting and wort production. Protein characterisation has mainly focused on proteins involved in formation of foam and haze. The most important proteins involved in foam formation are the lipid transfer protein LTP1, a 9.7 kDa hydrophobic protein; protein Z a 40 kDa polypeptide; and various hordein-derived polypeptides ranging in size from 10-30 kDa (Evans and Bamforth 2009). LTP1 originates from the barley aleurone and is expressed at the final stages of grain development. It is concentrated in beer foam and constitutes only 1% of the total malt protein (Evans and Hejgaard 1999). Different isoforms with variable molecular weights have been reported (Leiper et al., 2003a, b). Lipid transfer proteins have been isolated from several plants and seem to have several properties in common i.e. molecular weight around 10 kDa, considerable amino acid sequence
homology with four disulphide bridges between 8 conserved cysteines, high isoelectric point and lack of specificity for phospholipids (Sorensen et al., 1993).

Protein Z describes a family of barley Serpins (Serine Proteinase Inhibitors), an albumin-type protein that is estimated to constitute up to 2% of total malt protein (Evans and Hejgaard 1999). It has two isoforms, Z4 and Z7 and it is estimated that it is the most abundant non-dialyzable protein in beer at 10-25% (Evans and Bamforth 2009). In beer, it may be highly glycated with both hexoses and pentoses resulting in increased molecular weights being detected by SDS-PAGE analysis (Leiper et al., 2003a, b). Both LTPI and protein Z are abundant in the barley seed and show amino acid sequence homology to protease inhibitors and this may be the reason why they are not degraded by the proteolytic enzymes present during the malting and brewing processes and survive wort boiling (Sorensen et al., 1993).

Some of the properties of Protein Z were reported by Heejgard (Heejgard 1982) which include foaming properties and was also confirmed later by Evans (Evans et al. 1999). In the food processing, functional properties of proteins in addition to the nutritional value, are important aspects to consider, since they affect product performance. Some of the functional properties cited in literature (Kinsella et al. 1976) include solubility, binding properties, surfactant properties, viscogenic texturizing characteristics and foaming properties.
6.3 Materials and Methods

6.3.1 Pot ale samples and buffer preparation

Pot ale samples were collected from Glenkinchie distillery and kept at 5°C to avoid spoilage. Prior to the chromatography step, suspended solids of the samples were centrifuged for 15 minutes at 5000g.

6.3.2 Pot ale analysis

Soluble proteins were measured with the Bradford assay (Bradford 1976) described earlier in the thesis. Conductivity and pH were measured with the online conductivity and pH monitors incorporated in the liquid chromatography system (Äkta Avant 150 from GE-Healthcare) described below.

Carbohydrates content was measured using a colorimetric assay described in Fournier (Fournier 2001). Briefly, the method involved building a calibration curve using glucose a standard. Samples and standards (5 to 50 μl) were mixed with 500 μl of 4% phenol followed by 2.5 ml 96% sulphuric acid and then absorbance was measured at 490 nm with a spectrophotometer.

6.3.3 Buffers

The buffers used for the experiments were selected accordingly to the pH used i.e. 4.5, 5.8, 7.2 and 10.1. Table 6-1 describes the buffer used for each experiment.
Table 6-1. Buffers utilised for the elution step.

<table>
<thead>
<tr>
<th>pH</th>
<th>Buffer</th>
</tr>
</thead>
<tbody>
<tr>
<td>4.5</td>
<td>0.1 M Sodium Acetate – Acetic Acid</td>
</tr>
<tr>
<td>5.8</td>
<td>0.1 M Potassium Sodium phosphate</td>
</tr>
<tr>
<td>7.2</td>
<td>0.1 M Potassium Sodium phosphate</td>
</tr>
<tr>
<td>10.1</td>
<td>0.1 M Sodium Carbonate – Sodium Bicarbonate</td>
</tr>
</tbody>
</table>

6.3.4 Liquid Chromatography system

The liquid chromatography system used was Äkta Avant 150 from GE-Healthcare (Figure 6-4). The system measures online absorbance, conductivity and pH. Absorbance at 280 nm of the eluted streams was used as protein quantification indicator in addition to the off-line protein assay (Bradford assay).

This equipment uses a fraction collector system and for the experiments every 5 ml of the eluted material were collected in 8 ml tubes (BD Biosciences) for further analysis.

Figure 6-4. Photo of the Äkta Avant - Liquid Chromatography system.
6.3.5 Chromatography media

Commercially available 1 ml columns Hi-Trap Capto S and Hi-Trap Capto Q from GE-Healthcare were used for the experiments (Figure 6-5). The properties of the Capto S and Q columns are presented in Table 6-2.

![Hi-Trap Capto S and Hi-Trap Capto Q columns](image)

Figure 6-5. HiTrap Capto S and HiTrap Capto Q columns utilised during the experiments.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Capto S</th>
<th>Capto Q</th>
</tr>
</thead>
<tbody>
<tr>
<td>Matrix</td>
<td>Agarose with dextran surface extender</td>
<td>Agarose with dextran surface extender</td>
</tr>
<tr>
<td>Ion Exchange type</td>
<td>Strong cation</td>
<td>Strong anion</td>
</tr>
<tr>
<td>Charged Group</td>
<td>$\text{-SO}_3^-$</td>
<td>$\text{-N}^+\text{(CH}_3\text{)}_3$</td>
</tr>
<tr>
<td>Total ionic capacity</td>
<td>0.11 to 0.14 mmol Na+ /ml medium</td>
<td>0.16 to 0.22 mmol Cl-/ml medium</td>
</tr>
<tr>
<td>Particle Size</td>
<td>90 $\mu$m</td>
<td>90 $\mu$m</td>
</tr>
<tr>
<td>Dynamic binding capacity</td>
<td>$&gt;120$ mg lysozyme/ ml medium</td>
<td>$&gt;100$ mg BSA/ ml medium</td>
</tr>
<tr>
<td></td>
<td></td>
<td>$&gt;150$ mg ovalbumin/ ml medium</td>
</tr>
</tbody>
</table>
6.3.6 Chromatography protocols

The chromatography experiments were divided in two (experiment 1 and experiment 2) described in detail later and presented in Table 6-3. Briefly, the protocols consisted in five steps: column equilibration, sample loading, washing (to remove loosely bound proteins and impurities), elution (of proteins) and column cleaning. For this last step, a sequence of distilled water (5 CV), 1M NaOH (5CV or until the absorbance monitor was stable), distilled water and finally 2M NaCl (5CV) were passed through the columns.

6.3.6.1 Experiment 1: Media and pH selection experiments

A total of eight sub-experiments were carried out under four different pH conditions (pH 4.5/ 5.8/7.2/ 10.1) using two column media (Hi-Trap Capto Q and Hi-Trap Capto S). Flowrate (1 ml/ min) and pH were maintained during the equilibration, washing and elution steps. 50 ml of pot ale supernatant was loaded into the column. The elution step consisted of a 0.5M NaCl salt gradient over 10 CV followed by a 1M NaCl step over 5 CV.

6.3.6.2 Experiment 2: Extended sample loading with HiTrap Capto S at pH 4.5

Similar to the Experiment 1, but in this case, 200 ml of pot ale supernatant was loaded to the previously equilibrated (pH 4.5 - 0.1 M Sodium acetate-acetic acid buffer) 1 ml Hi-Trap Capto S column at a flow rate of 2 ml/ min. After the loading step, a 30 CV washing step was included. Elution was performed at a pH of 4.5 (0.1 M Sodium acetate-acetic acid buffer solution) with 5 CV of 20% increments in salt concentration until reaching 1 M NaCl concentration. The elution was followed by a washing step of 1M NaCl over 5 CV.

During the loading, elution and cleaning steps, samples were collected every 5 ml and subsequently analysed for protein content using the Bradford method. These analyses were performed in triplicate.

The steps involved in experiments 1 and 2 are summarised in Summary of the protocols used for experiments 1 and 2.
Table 6-3. Summary of the protocols used for experiments 1 and 2: including buffers, concentrations, pH and volumes used on each step of the chromatography protocol.

<table>
<thead>
<tr>
<th>Step</th>
<th>Experiment 1</th>
<th>Experiment 2</th>
</tr>
</thead>
<tbody>
<tr>
<td>Equilibration</td>
<td>pH 4.5/ 5.8/ 7.2 /10.1 (same as elution)</td>
<td>pH 4.5</td>
</tr>
<tr>
<td>Loading</td>
<td>50 ml (= 50CV)</td>
<td>200 ml (= 200 CV)</td>
</tr>
<tr>
<td>Washing</td>
<td>pH 4.5/ 5.8/ 7.2 /10.1 (same as elution) over 20 CV</td>
<td>pH 4.5 over 30 CV</td>
</tr>
<tr>
<td>Elution</td>
<td>pH 4.5/ 5.8/ 7.2 /10.1 with a 0.5M NaCl gradient over 10 CV followed by a step of 1M NaCl + buffer over 5 CV</td>
<td>pH 4.5 over 5 CV steps with 0.2M NaCl increments to 1M NaCl followed by 1M NaCl over 5 CV.</td>
</tr>
<tr>
<td>Cleaning</td>
<td>pH 4.5/ 5.8/ 7.2 /10.1 (same as elution) + 1M NaCl over 5 CV.</td>
<td>pH 4.5/ 5.8/ 7.2 /10.1 (same as elution) + 1M NaCl over 5 CV.</td>
</tr>
</tbody>
</table>
6.3.7 SDS-page analysis

Proteins were analysed using either Bio-Rad Mini-PROTEAN Tris-Glycine (TGX) or Tris-Tricine gels (Bio-Rad Mini-Protean precast gels, Bio-Rad Laboratories, Herts, UK) along with the corresponding samples and running buffers.

For analysis using the TGX system, samples of known protein concentration, diluted in water where required, were mixed with equal volumes of Laemmli sample buffer (Laemmli 2× concentrate sample buffer, Sigma-Aldrich Ltd., Dorset, England). These were heated at 90°C for 5 min or 70°C for 10 min, centrifuged and loaded on to 4-20% (catalogue #456-1094) or 10% precast polyacrylamide gels (catalogue #456-1034). Gels were run using a Bio-Rad Mini-Protean Tetra Cell System for mini precast gels with Tris-Glycine running buffer. A 10× Tris-Glycine running buffer consisting of (g/l) Tris-Base 30.28; Glycine 144.25; and SDS 10, was diluted to the working volume with water before use. A pre-stained, broad range protein marker (7-175 kD from New England Biolabs Ltd., Herts, UK or 2-250 kD, Precision Plus Protein Dual Extra Standard from Bio-Rad) was run with all gels to estimate protein molecular weight. The exact electrophoresis conditions corresponding to each gel are provided in the results section. After electrophoresis, gels were rinsed with water and stained overnight with a Colloidal Coomassie Blue stain (Pink et al., 2010) consisting of 5% (w/v) aluminium sulphate hydrate (14-18 degree of hydration), 10% (v/v) ethanol, 0.02% (w/v) Coomassie Brilliant blue G-250 and 8% (v/v) orthophosphoric acid. Gels were rinsed and destained with water until the background stain was removed. The SDS-PAGE gels were scanned with a ChemiDoc XRS+ imaging system (Bio-Rad) and images were analysed using GelAnalyzer software (www.gelanalyzer.com) to estimate the molecular weight of protein bands.

Gels performed with the assistance of Dr. Jane White.
For analysis using the Tris-tricine system for peptide analysis, samples were mixed with an equal volume of tricine sample buffer (Catalog #161-0739, Bio-Rad) containing 2% β-mercaptoethanol (catalog #161-0710) and heated at 70°C for 10 min. These were loaded on to 10-20% Mini-PROTEAN Tris-tricine gels (catalog #456-3114) and run using 1× Tris-Tricine running buffer (catalog #161-0744, Bio-Rad) with the electrophoresis conditions as specified in the results below. 15 µl of the Precision Plus Protein Dual Extra Standard was run with all gels to estimate protein molecular weight. After electrophoresis, the gels were rinsed with water and fixed with destain solution for 60 min before staining. The destain contained 2% phosphoric acid and 10% ethanol and colloidal coomassie blue (CCB) stain was as described in Pink et al (Pink et al. 2010). Gels were stained overnight, rinsed with water and incubated with destain for a further 2 hours and rinsed with water until all background stain had been removed. Images were obtained as described above.

Samples were heated with sample buffer at 70°C for 10 min and the electrophoresis run conditions were 150 V for approximately 1 hour.
6.4 Results and discussion

6.4.1 Pot ale sample analysis

Relevant properties of the pot ale supernatant used during the experiments are presented in Table 6-4 below.

Table 6-4. Properties of pot ale used during the experiments.

<table>
<thead>
<tr>
<th>Property</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Soluble protein concentration</td>
<td>414 mg/L</td>
</tr>
<tr>
<td>Conductivity</td>
<td>5.3 mS/cm</td>
</tr>
<tr>
<td>pH</td>
<td>4.0</td>
</tr>
<tr>
<td>Carbohydrate concentration</td>
<td>18.5 g/L</td>
</tr>
</tbody>
</table>

6.4.2 Media selection experiments (Experiment 1)

The absorbance over time (or CV) of the eluted fractions under the conditions described earlier were plotted in the chromatograms shown in Figure 6-7 (HiTrap Capto S) and Figure 6-8 (HiTrap Capto Q). From the chromatograms, different peaks can be observed after 70 CV (during the elution step).

The area and height of these peaks were compared in Figure 6-6. The area of the curve is proportional to the amount of protein in the elution. In all cases the area of HighTrap Capto S experiments are bigger than the areas of the HiTrap Capto Q experiments. This is an indication that Capto S has better affinity for the proteins in pot ale.

The maximum area was found at a pH of 4.5 using HiTrap Capto S (cation exchanger). The area of the peak decreases as the pH increases when the HiTrap Capto S column is used, however the opposite is observed when using the HiTrap Capto Q column. This is due that at lower pH conditions (acid) more protons (H⁺) are available to be exchange with the negatively charged chromatographic medium (HiTrap Capto S) and the proteins are then desorbed from the medium and dissolved in the solution. Similarly, at higher pH conditions (alkali), OH⁻ ions are adsorbed with the positively charged medium and exchanged with the (positively charged) proteins.
The fraction corresponding to the peak height of each experiment (column and pH) were taken for further soluble protein and carbohydrate analysis. The results are presented in Figure 6-9 (protein) and Figure 6-10 (carbohydrates). The soluble protein results correspond to the results presented earlier in Figure 6-6, where a decline in protein concentration was observed with a pH increase, except for the experiment conducted at pH 10.1. However, if the chromatogram in Figure 6-7 is taken into account, the absorbance of the experiments conducted at pH 4.5 and 10.1 have the same magnitude (~ 900 mAU).

Interestingly and not surprisingly, is that carbohydrate concentration at maximum peak height was less than 1.0% of the original sample. This is due that the carbohydrates lack of charge (unlike proteins) flowing through the column media while protein bind to the media.

Another interesting effect observed during the chromatography experiments was the difference in colour on both the eluted samples and the column. Figure 6-11 shows that the elution fraction of the HiTrap Capto S experiment at pH 4.5 had a transparent colour, while the HiTrap Capto Q (pH 10.1) samples had a brown colour similar to the original pot ale. Additionally, after the 4 consecutive experiments, the HiTrap Capto S column remained visually clean (white), while the HiTrap Capto Q column seemed to have accumulated some brown material in the interior (Figure 6-12). There were unsuccessful attempts to clean this material from the column with other solvents apart from the ones mentioned earlier (i.e NaOH, ethanol) such as iso-propanol, but the discussion of this is beyond the scope of this chapter.
Figure 6-6. Comparison of peak and height (a) and area (b) of the Capto S and Capto Q columns.
Figure 6-7. HiTrap Capto S chromatogram for the experiments conducted at pH 4.5 (blue), pH 5.8 (green), pH 7.2 (red) and pH 10.1 (brown).

The absorbance on each experiment was measured at 280 nm and the values were reported in mAU on the primary vertical axis. The secondary vertical axis, represents the values of the salt gradient (black line), where 1M = 100%.
Figure 6-8. HiTrap Capto Q chromatograms for the experiments conducted at pH 4.5 (blue), pH 5.8 (green), pH 7.2 (red) and pH 10.1 (brown).

The absorbance on each experiment was measured at 280 nm and the values were reported in mAU on the primary vertical axis. The secondary vertical axis, represents the values of the salt gradient (black line), where 1M = 100%.
Figure 6-9. Relative soluble protein concentration at maximum peak height to pot ale for the experiments conducted at pH 4.5, 5.8, 7.2 and 10.1 using the Capto S and Capto Q columns.

Figure 6-10. Relative carbohydrate concentration at maximum peak height to pot ale for the experiments conducted at pH 4.5, 5.8, 7.2 and 10.1 using the Capto S and Capto Q columns.
Figure 6-11. Samples eluted at pH 4.5 – using a HiTrap Capto S column - (left) and pH 10.1 – using a HiTrap Capto Q column (right).

Figure 6-12. HiTrap Capto Q (up) and HiTrap Capto S (down) columns after 4 consecutive experiments.
6.4.3 Extended sample loading with Capto S at pH 4.5 (Experiment 2)

In this experiment a 4 fold increase in sample loading allowed the detection of 3 distinct peaks during the elution (Figure 6-13). The peaks (1, 2 and 3, from left to right in the following graphs and tables) appeared at each incremental salt step (0.2, 0.4 and 0.6 M NaCl). After the 0.8M NaCl step, no peaks were detected.

The main peaks parameters (area, width and height) were compared in Figure 6-14. Peak area and specially, peak height decreased as the experiment progressed, but overall the peaks broadened overtime. Additionally, peaks asymmetry and resolution were calculated and the results are presented in Table 6-5. From a peak resolution perspective, it can be said that this was not achieved, but the objective of this experiment was not aimed to produce well resolved peaks.

The conductivity of the mobile phase was monitored continuously, but to simplify the appearance of the results, only the start, highest point and end of the peaks are presented in Figure 6-15. As expected, conductivity increased with salt concentration. Conductivity of the start of peak 3 was seven times higher compared to the start of peak 1, but salt concentration increased only 3 times (from 0.2M NaCl to 0.6M NaCl). Conductivity with 1M NaCl was 87 mS/ cm, about 15 times the conductivity recorded at beginning of the elution, but little protein was unbound from the column, with most of the protein desorption happening at the first elution step (0.2 M NaCl), equivalent to a conductivity to 6-25 mS/ cm.
Figure 6-13 Chromatogram of experiment 2: elution at pH 4.5 with 200 ml of pot ale loaded.

The absorbance was measured at 280 nm (blue line) and 254 nm (red line) and the values were reported in mAU on the primary vertical axis. The secondary vertical axis, represents the values of the salt gradient (green line), where 1M = 100%. Three peaks were indentified and labelled “1”, “2” and “3” in the figure above.
Figure 6-14. Parameters of peak 1, peak 2 and peak 3 from experiment 2 (elution at pH 4.5, 200 ml pot ale loaded) including relative area, height and width of the peaks.

Figure 6-15. Conductivity measurements of peaks 1, 2 and 3 during experiment 2 (elution at pH 4.5, 200 ml pot ale loaded) at start, top and end of the peak.

Table 6-5. Resolution and asymmetry of peaks identified in the chromatogram of experiment 2.

<table>
<thead>
<tr>
<th>Peak No.</th>
<th>Resolution</th>
<th>Asymmetry</th>
</tr>
</thead>
<tbody>
<tr>
<td>Peak 1</td>
<td>-</td>
<td>2.21</td>
</tr>
<tr>
<td>Peak 2</td>
<td>0.55</td>
<td>4.22</td>
</tr>
<tr>
<td>Peak 3</td>
<td>0.54</td>
<td>20.52</td>
</tr>
</tbody>
</table>
Only HiTrap Capto S samples eluted at pH 4.5 using 0.1 M sodium acetate buffer from Experiment 1 (wells A and B in Figure 6-16) and Experiment 2 (wells C-H) were analysed. The images in Figure 6-16 are presented in black and white (a) and colour (b) format to ease the identification of the protein bands in the gel.

Samples in wells A and B were taken from the maximum absorbance values of the 4.5 pH peak collected between 85 to 91 CV (chromatogram in Figure 6-7). Wells C and D correspond to Peak 1 of the chromatogram in Figure 6-13, while wells E and F correlate to Peak 2 and wells G and H to Peak 3. As described earlier, Peaks 1, 2 and 3, correspond to NaCl concentration of 0.2 M, 0.4 M and 0.6 M in the 0.1 M sodium acetate buffer (pH 4.5), respectively. The total µg of protein loaded in each well is shown beneath the labels. A summary of wells and experiments is presented in Table 6-6.

Table 6-6. Correlation between wells and experiments with HiTrap Capto S column.

<table>
<thead>
<tr>
<th>Well</th>
<th>Experiment</th>
<th>Elution pH</th>
<th>NaCl concentration</th>
<th>Peak</th>
</tr>
</thead>
<tbody>
<tr>
<td>A-B</td>
<td>1</td>
<td>4.5</td>
<td>1.0 M</td>
<td>-</td>
</tr>
<tr>
<td>C-D</td>
<td>2</td>
<td>4.5</td>
<td>0.2 M</td>
<td>1</td>
</tr>
<tr>
<td>E-F</td>
<td>2</td>
<td>4.5</td>
<td>0.4 M</td>
<td>2</td>
</tr>
<tr>
<td>G-H</td>
<td>2</td>
<td>4.5</td>
<td>0.6 M</td>
<td>3</td>
</tr>
</tbody>
</table>

From this analysis it could be observed that wells A-B contain low molecular weight proteins (< 7 kDa) while wells C-D indicate the presence of higher molecular weight proteins (30-46 kDa). This band also seems to appear in wells E and F, but it is less evident and not noticeable in wells G and H. The 30-46 kDa band indicates the presence of Protein Z (40 kDa), while LTP1 is around 10 kDa.

The significance of this finding is that individual proteins (Protein Z and LTP1) have been identified in pot ale. Potentially these proteins can be recovered and isolated from other proteins in pot ale. Protein Z, for example, has known functional properties, such as foaming. Functional properties, in addition to the nutritional properties of the proteins might result in a higher value product compared to an ingredient for salmon.
feeding, which value is purely based on nutritional properties (as reviewed earlier in Section 2.8). Yields, quantification and further characterisation is beyond the scope of this thesis and could be proposed for further development.

Figure 6-16. SDS-PAGE (TGX 4-20%) of eluted samples at pH 4.5 using a HiTrap Capto S column.

The figures are presented in black and white (a) and colour (b) format to ease the identification of the protein bands in the gel.
6.5 Conclusions

Proteins in pot ale can be recovered and separated by ion exchange chromatography (IEC). There are other methods available that potentially can be utilised and were discussed in this chapter. Ultrafiltration (UF), high performance tangential flow filtration (HPTFF) and precipitation were suggested as protein recovery methods for pot ale proteins. However, it was concluded that IEC was the method that will be further investigated in this thesis, but the possibility of introducing these unit operations to the overall protein recovery process must be further investigated.

The experiments presented here, demonstrated that a negatively charged column media (HiTrap Capto S) resulted in higher protein capture compared to the positively charged column (HiTrap Capto Q).

The effect of pH on protein recapturing was also studied, showing that the lower pH, the higher the protein recovered. At pH 4.5 maximum recovery was achieved, however when these purified proteins were analysed with SDS-page, no clear protein bands were recognised.

A second experiment with a fourfold increase in the sample loading was conducted (using the HiTrap Capto S column at pH 4.5). This experiment showed 3 distinctive, but not well resolved, peaks during elution. The peaks were analysed with SDS-page, where a clear band around the expected size of Protein Z can be visualised. The identification of Protein Z in pot ale could result in future work where specific proteins can be recovered, offering a product with higher value than nutritional proteins. Protein Z, for example, has been reported to have foaming properties.

Although these experiments were conducted at 1-2 ml/ min, it is possible to increase capacity with larger columns, already available commercially. However, cost of the Capto S media would be a concern. Experiments at higher flowrate, using more economic media will be described in the following chapters.
CHAPTER 7 - POT ALE PROTEIN ADSORPTION USING LOW COST MATERIALS

Abstract

A rapid method to assess pot ale protein adsorption and desorption was developed. The adsorbents used in these experiments had a significantly lower cost compared to the HiTrap Capto S and Q resins (GE Healthcare) described in the earlier chapter. The materials tested included sand, zeolites, glass beads and diatomaceous earth.

For the adsorption experiments 1 ml of clarified pot ale was mixed with a previously treated adsorbent (acid, alkali and water washes) in a 2 ml Eppendorf tube. The mixture of pot ale and adsorbent was centrifuged, and the supernatant was separated again and the soluble protein content was measured and compared with the protein content of pot ale prior to the contact with the adsorbent. Among the adsorbents tested, Zeolite C (zeolite clinoptilolite) and Diaguard (diatomaceous earth) showed the highest protein adsorption results (> 95%).

Desorption experiments focused on Zeolite C and Diaguard, and showed that exposing these materials to high pH conditions (pH>8) resulted in high protein release. A 1 M NaOH solution in contact with the Zeolite or Diaguard resulted in virtually all the protein being desorbed.
7.1 Introduction

In the previous chapter, ion exchange chromatography was utilised as a method for the separation of proteins from pot ale. Commercially available pre-packed columns (HiTrap Capto Q and HiTrap Capto S, from GE Healthcare) were used successfully; however the cost of the media was a concern for upscaling purposes. Typically the cost of Capto S/ Capto Q media is around £5000 per litre (GE Healthcare, Merck-Millipore); whereas the materials used for the experiments presented in this chapter were in the order of £1-£100 per litre (Holistic Valley, Amazon). Other sources indicate a price about US$ 0.03– 0.12 per kg, depending on the quality of the mineral (Babel and Kurniawan 2003).

A quick and simple method was developed to determine the suitability of protein adsorption. Small quantities of pot ale (1 ml) and adsorbent (0.1 mg) were used and the amount of pot ale and adsorbent together with conditions such as salt concentration and pH was evaluated.

A brief literature review is also included which focuses on zeolites, their applications as adsorbent materials and the physicochemical principles behind adsorption.
7.2 Theoretical background

Adsorption is a technique widely used for the removal of pollutants from water such as heavy metals and dyes (Crini 2006). In food processing several materials are used including charcoal, clay, carbon of animal origin, zeolites and in last decades activated carbon, synthetic zeolites and further synthetic resins based in polystyrene, polyacrylic esters have become more popular (Kammerer et al. 2010). However, synthetic resins and activated carbon cannot be suitable for the recovery of bio-compounds, i.e. proteins from food processing streams, due to their relative higher cost compared to natural materials such as zeolites (Crini 2006). Hence, natural zeolite is becoming a popular alternative as adsorbent in several industries as it is described below.

7.2.1 Zeolites

The name “zeolite” was introduced by the Swedish mineralogist Cronstedt in 1756 for certain silicate minerals in allusion to their behaviour on heating in a borax bead (Greek zeo = boil; lithos = stone). More than 40 species of zeolites are available in natural form, with zeolite clinoptilolite being the most abundant and frequently studied (Coombs et al. 1997).

There is not much information available about protein adsorption using zeolite, although some work has been reported on the use of zeolite on waste water treatment and specifically, ammonia removal. Experiments on this subject studied by Cooney (Cooney et al. 1999a), (Cooney et al. 1999b) revealed that the highest ammonium removal efficiency was achieved when the zeolite’s exchange sites were converted to the sodium form. Multicomponent equilibrium experiments were carried out to determine the effects of competing cations on the ammonium-exchange capacity of the zeolite. The laboratory study indicated the zeolite’s selectivity for ammonium ions over other cations typically present in sewage (calcium, magnesium, and potassium), and provided information relevant to the design and operation of a continuous process.

The main components in the natural clinoptilolite structure, are presented in Figure 7-1 (Cooney et al. 1999a), where M1 to M4 describe the molecular bonds between the elements (Na, Ca, K and Mg) and water within the zeolite crystal. The crystal lattice
comprises channels of two different sizes, and each type may display different selectivity behaviour for certain cations.

![Diagram of clinoptilolite structure](image)

**Figure 7.1. Main components of the clinoptilolite structure (Cooney et al. 1999a).**

Apart from economic reasons, other advantages cited about zeolites, include high ion-exchange capacity and relatively high specific surface areas. Another advantage of zeolites over resins is the ion selectivity generated by their rigid porous structures. Cation selectivity is different for each zeolite, and clinoptilolite is highly ammonium-ion selective. Ion selectivity for clinoptilolite follows the order shown below (Cooney et al. 1999a):

\[
    \text{Cs}^+ > \text{Rb}^+ > \text{K}^+ > \text{NH}_4^+ > \text{Ba}^+ > \text{Sr}_2^+ > \text{Na}^+ > \text{Ca}_2^+ > \text{Fe}_3^+ > \text{Al}_3^+ > \text{Mg}_2^+ > \text{Li}^+
\]

This is due to the size and charge of the hydrated cation and specific crystal structure of, and distribution of, the exchange sites in the zeolite. Zeolite-exchange selectivity depends on its particle size, shape, uniformity, purity, and consistency. Due to their regular, uniform structure, zeolites exclude ions by an “ion sieve” action, and the degree of exclusion is specific for each zeolite.
7.2.2 Adsorption mechanism on zeolites

The adsorption mechanism on zeolite particles is complex because of their porous structure, inner and outer charged surfaces, mineralogical heterogeneity and other imperfections on the surface. However, it is recognized that, like clay, the adsorption properties of zeolites result mainly from their ion-exchange capabilities (Duarte-Silva et al. 2014).

Sakaguchi (Sakaguchi et al. 2005) concluded that there are three physicochemical principles that may underlie adsorption: (1) below the pI, mainly Coulomb’s attractions may occur, as in ion-exchange chromatography, (2) at the pI, hydrophobic interactions (a kind of van der Waal’s attraction) plus the three-dimensional mesopore structure are involved, (3) above the pI, the sum of the Coulomb’s repulsions and attractions (such as hydrophobic interactions) and substitution reactions of water on the Al molecule with a protein amino base might be important. At high Si/Al ratios in the presence of a small amount of Al and with mesopores between the zeolite particles, maximal absorption was seen at the pI, suggesting a dependence both on the number of hydrophobic interaction points on the mesopores and on the mesopore morphology (Figure 7-2).

Figure 7-2. Zeolite–protein interactions under different pH conditions (Sakaguchi et al. 2005)
7.2.3 Point of Zero Charge

The point of zero charge (pzc) corresponds to the pH value of the liquid surrounding oxide particles when the sum of surface positive charges balance the sum of surface negative charges. The pzc value characterizes surface acidity: when oxide particles are introduced in an aqueous environment their surface charge is positive if pH(solution)<pzc and is negative if pH(solution)>pzc.

Experimental methods used to determine the pzc of colloidal particles (particle size less than 1 μm) include zeta potential determination and electrophoretic micromobility, are difficult to apply to larger particles (particle size bigger than 20 μm). It can be noticed that zeta potential of coarse particles can be determined from potential flow method.

In the work carried out by Reymond and Kolenda (Reymond and Kolenda 1999), the relationship between silica oxide content of a silica-alumina mixture and pzc is plotted in Figure 7-3. When silica content increases from 0 to 85 wt.% the mixture pzc value varies linearly as a function of silica content. This variation differs from a barycentre law where each oxide behaves independently and contributes to the pzc of the oxide mixture proportionally to its content in the mixture. This suggests that alumina hydrate offers more sites per gram for solid–liquid reactions. When silica content is very high (>90 wt.%) silica surface coverage by alumina particles decreases, silica surface charge becomes predominant. The oxide mixture pzc sharply lowers towards silica pzc.
Variations of the pzc value of a silica–alumina mixture as a function of silica content (Reymond and Kolenda 1999)

7.2.4 Z potential

Electrophoretic light scattering is the most commonly method used for zeta potential measurement due to its sensitivity, accuracy, and versatility. Other methods cited include capillary electrophoresis, acoustic, and electroacoustic.

Theoretical and experimental results have confirmed that zeta potential is affected not only by the suspension conditions such as pH, temperature, ionic strength, and even the types of ions in the suspension, but also by the particle properties such as size and concentration (Wang et al. 2013)
7.3 Methods and materials

7.3.1 Pot ale

Pot ale (collected from Glenkinchie and kept at 4°C) was centrifuged for 15 minutes at 5000g. Solids were then discarded and 1 ml of pot ale supernatant was placed into a 2 ml Eppendorf tube together with the pre-treated adsorbent material as described below. Prior to the adsorption experiments pot ale supernatant was analysed for soluble protein concentration using the Bradford assay (Bradford 1976).

7.3.2 Adsorption and desorption experiments

A list of the adsorbents used for the experiments are presented in Table 7-1 below.

Table 7-1. List of adsorbents used for the pot ale protein adsorption experiments.

<table>
<thead>
<tr>
<th>Adsorbent</th>
<th>Details</th>
<th>Silica oxide content</th>
</tr>
</thead>
<tbody>
<tr>
<td>Diaguard</td>
<td>Diatomaceous earth. JJS Minerals.</td>
<td>94.9%</td>
</tr>
<tr>
<td>Zeolite C</td>
<td>Zeolite clinoptilolite. Holistic Valley. 71.0</td>
<td>71.0%</td>
</tr>
<tr>
<td>Celpure</td>
<td>Diatomite, Amorphous Siliceous Earth. Advanced Minerals.</td>
<td>96-98%</td>
</tr>
<tr>
<td>Glass beads (3 mm)</td>
<td>Sigma Aldrich</td>
<td>-</td>
</tr>
<tr>
<td>Glass beads (0.8 mm)</td>
<td>Sigma Aldrich.</td>
<td>-</td>
</tr>
<tr>
<td>AW Hyflow</td>
<td>Flux Calcined Diatomaceous earth. Advanced Minerals.</td>
<td>Acid washed, up to 44% Crystalline Silica</td>
</tr>
</tbody>
</table>

7.3.3 Pre-treatment of the adsorbents

Prior to exposing the adsorbents to pot ale, the adsorbents were pre-treated with a series of caustic, water and acid washes, to ensure contaminants were removed and the surface of the adsorbent were adequately charge for protein binding.
Approximately 100 mg of adsorbent (the exact weight was recorded) was placed in a 2 ml Eppendorf tube and mixed with 1 ml 1M NaOH. The mixture of adsorbent and NaOH was suspended with a vortex mixer for 20 seconds, left to settle for at least 60 s and the mixed again. This step was repeated 2 more times. The suspension was then centrifuged (Minifuge, Eppendorf) at 14000 rpm for 90 seconds and the liquid was discarded with care of not losing any adsorbent from the tube (a pipette was used in some cases). Subsequently, distilled deionised water and then acetic acid-sodium acetate buffer (0.1 M acetic acid – sodium acetate pH 3.8) was mixed with the adsorbent using the same procedure used with the caustic treatment.

In some experiments the ratio of pot ale to adsorbent was modified (i.e. 100 mg of adsorbent was increased to 1000 mg), and the experiments were conducted in 15 or 50 ml tubes.

7.3.4 Adsorption experiments

For the adsorption experiments, 1 ml of pot ale supernatant was placed into a 2 ml Eppendorf tube together with the pretreated adsorbent material. The mixture of pot ale-adsorbent was suspended with a vortex for 20 s and the tubes were placed in an orbital shaker for 30 minutes at 150 rpm (gentle mixing). After this, the suspension was centrifuged (Minifuge, Eppendorf at 14000 rpm for 90 s), the pot ale separated from the adsorbent (with care of not losing any adsorbent) and analysed for soluble protein content. The adsorbent was kept in the tube for the desorption experiments explained below. The tests were performed in triplicate and the results reported as % Protein Adsorbed (Protein Adsorbed = 1 − c/c₀) where c₀ an c are the soluble protein content before (original pot ale sample) and after the adsorption step, respectively.

7.3.5 Desorption experiments

For the desorption experiments, the adsorbent material from the adsorption experiments kept in the Eppendorf tube was suspended with a vortex mixer in 1 ml of a buffer solution (detailed in the next section) using the same procedure described in the adsorption experiments (20 s vortex, 60 s settle, repeat 2 more times) and then the discarded buffer was analysed for soluble protein content. The tests were performed in triplicate and the results reported as % Protein Desorbed (Protein Desorbed = c/
where $c_0$ and $c$ are the soluble protein content before (original pot ale supernatant) and after the desorption step, respectively.

In some cases, the buffers were added to adsorbent as a sequence of washes and the results are reported as Wash 1 and Wash 2. An example of the procedures used for the experiments is shown in Figure 7-4.

![Figure 7-4. Example of procedure used for adsorption/desorption experiments.](image-url)
7.3.6 **Buffers**

All the buffers were prepared using distilled deionised water and included:

- 0.1 M sodium acetate- acetic acid (NaOAc-HAc) pH 3.8
- 0.1 M sodium acetate- acetic acid (NaOAc-HAc) pH 3.8 + 1M NaCl
- 0.1 M sodium acetate- acetic acid (NaOAc-HAc) pH 5.6
- 0.1 M sodium acetate- acetic acid (NaOAc-HAc) pH 5.6 + 1M NaCl
- 0.1 M potassium phosphate buffer pH 7.0 and 7.8
- 0.1 M sodium carbonate – sodium bicarbonate buffer (pH 8.4-10.8)
- 0.01M, 0.1M and 1M NaOH

7.3.7 **Z-potential analysis**

A ZetaSizer Nano ZS (Malvern Instruments, UK) with a laser Doppler electrophoresis technique was used to analyse the potential electric charge of zeolite. Zeolite samples were separated in fine (<90 micron) and coarse (>90 micron) particles using a 90 micron sieve. Samples of the fine, coarse and the original zeolite as supplied were suspended in 0.1 M sodium carbonate – sodium bicarbonate buffer at a pH 8.4, 10.1 and 11.4. 9 samples were produced in total.

The zetasizer required 2 ml samples, with 50 micrograms of zeolite per ml of sodium carbonate. Accordingly, 20 ml of buffer were thoroughly mixed with 1000 micrograms of zeolite, from which smaller samples were taken for the zetasizer analysis. Before testing, zeolite was resuspended by running each sample for two minutes in an Ultrawave ultrasonic bath mixer. A 2 ml sample was pipetted to an injection cuvette, which was placed into the machine to take triplicate readings.
7.4 Results and discussion

7.4.1 Adsorption experiments

From the adsorption experiments, in Figure 7-5, it can be observed that Diaguard and Zeolite C were the adsorbents with better results in terms of protein adsorption. The Diaguard tests resulted in virtually all protein adsorbed and Zeolite C in 95.9% of protein adsorbed. In third place, with 70.3% protein adsorption was the sand. However, in this case 1000 mg were used, but when 100 mg were used (bar on the far right), only 13.1% of protein adsorption was achieved, suggesting a very low binding capacity.

Similarly, for the glass bead experiments, 1000 mg and 100 mg were used, but only the former amount is reported to simplify the results. With the other adsorbents- Celpure and AW Hyflow - less than 50% adsorption was achieved. This result might be due that AW Hyflow has a lower silica content compared to the other materials as described earlier in Table 7-1.

Based on these results, subsequent experiments - desorption and continuous adsorption – were carried out using Diaguard and Zeolite C as the adsorbents.
Figure 7-5. Relative protein adsorption of 1 ml pot ale supernatant proteins on the materials tested during the experiments.

The quantity of material is indicated in brackets. Typically 100 mg were used, except for sand and glass bead, where 1000 mg were required. The error bars represent the standard deviation of the triplicates of the experiments.
In another experiment, the effect on protein adsorption when the ratio of pot ale to adsorbent was varied is presented in Figure 7-6 and Table 7-2. By increasing the pot ale to adsorbent ratio the amount of protein adsorbed per gram of adsorbent increased almost in the same proportion (left graph in Figure 7-6). However, when the experiments were conducted on the bigger tubes (15 and 50 ml), it was observed that the relative protein adsorbed from the original concentration decreased from ~100% to less than 93% (right graph in Figure 7-6). This phenomenon could be explained to mixing effects or a lower residence time to allow proteins to bind to the adsorbent.

**Figure 7-6.** Effect on protein adsorption when the ratio of pot ale to adsorbent was varied.

**Table 7-2.** Protein adsorption and the variation of pot ale and adsorbent amounts.

<table>
<thead>
<tr>
<th>Pot ale volume (ml)</th>
<th>Adsorbent mass (g)</th>
<th>Pot ale to adsorbent ratio (ml/g)</th>
<th>Protein adsorbed (g)</th>
<th>Protein adsorbed per gram of adsorbent (mg/g)</th>
<th>% of initial protein adsorbed</th>
</tr>
</thead>
<tbody>
<tr>
<td>1.0</td>
<td>0.103</td>
<td>9.73</td>
<td>0.755</td>
<td>7.3</td>
<td>98.4%</td>
</tr>
<tr>
<td>1.0</td>
<td>0.0517</td>
<td>19.35</td>
<td>0.824</td>
<td>16.0</td>
<td>99.7%</td>
</tr>
<tr>
<td>1.5</td>
<td>0.1042</td>
<td>14.40</td>
<td>1.216</td>
<td>11.7</td>
<td>98.2%</td>
</tr>
<tr>
<td>20</td>
<td>1.0264</td>
<td>19.49</td>
<td>15.771</td>
<td>15.4</td>
<td>93.3%</td>
</tr>
<tr>
<td>40</td>
<td>2.0240</td>
<td>19.76</td>
<td>27.214</td>
<td>13.4</td>
<td>90.9%</td>
</tr>
</tbody>
</table>
7.4.2 Effect of pH on protein adsorption

In Figure 7-7, the effect of pH conditioning (prior to the introduction of pot ale) on protein adsorption was studied using Diaguard as the adsorbent material. Acidic pH and below a pH of 8, resulted in more favourable conditions for protein adsorption (close to 100% adsorption). More alkaline conditions (pH > 9), about 5% of the protein remain unbound to the Diaguard particles.

![Figure 7-7. Effect of pH conditioning on protein adsorption using Diaguard particles as the adsorbent materials.](image)

7.4.3 Desorption experiments

From the desorption experiments (Figure 7-8), it can be observed that when the materials were exposed to 1M NaOH, the majority of the protein was detached from the adsorbent. In the case of Diaguard more than 100% of the original protein amount loaded was detected after the introduction of 1M NaOH. These results could be explained due to the presence of impurities that were not removed during the pre-treatment of Diaguard.

For Zeolite C, 76.4% of the original protein was desorbed with the introduction of NaOH and a further 1.9% with the pH 3.8 buffer, another 2.4 % with pH 3.8 and 1 M
NaCl and nothing detected at pH 5.6 + 1 M NaCl. With the other adsorbent the contribution of NaOH in protein desorption became less important.

Figure 7-8. Protein desorbed from the material used during the experiments (Diaguard, Zeolites C, sand, celpure, glass beads and AW Hyflow) under different pH conditions.

The effect of pH on protein desorption using Zeolite C as the adsorbent material is shown in Figure 7-9. At high pH (or higher NaOH concentration), a higher amount of protein desorption was achieved. At pH 13 and 14, protein desorbed was ~80% of the initial protein content added to the zeolite. At pH 12, protein desorption was ~20% and below pH 11, using NaOH and HCl to adjust the pH, little protein was unbound from the Zeolite. However, when the same experiment was conducted using a sodium carbonate - sodium bicarbonate buffer (pH 9 - 11), higher levels of protein desorption were measured (70 - 80%). At pH 8, using a sodium phosphate buffer, 45% of the protein was recovered from the zeolite. It was also observed that samples with darker colour have higher protein content as depicted in Figure 7-10.

From the experiments described earlier in Figure 7-7, protein desorption on Diaguard particles was studied under different pH conditions (Figure 7-11). After the adsorption step, 1 ml of the same buffer (same pH) was added to the diatoms (1st wash) and the
soluble protein was measured. Subsequently, a second wash (the same buffer pH + 1 M NaCl) and finally third with 1M NaOH was added to the solids.

For low pH conditions, negligible protein is released. Most of the protein is unbound with final wash (NaOH wash). At higher pH values (pH>7), some of the protein is desorbed, but again the majority of the protein released with the final wash. The addition of salt (2nd wash) resulted in little protein release compared to the other washes.

The results were in agreement with the theory described earlier in in section 6.2.1, where it was suggested that an increased pH would change the surface charge of the proteins. At low pH, the proteins will be positively charged and at higher pH, protein charge will change from positive to a negative charge, thus detaching from the zeolites or the other materials studied, which are negatively charged.

Another factor that might explain the discrepancy in protein adsorption (and desorption) on the materials studied is the porosity of the materials. Diaguard and zeolite are porous materials, while sand and the glass bead are non-porous. This property was not investigated and discussed in depth in this chapter, but it will be discussed later in Chapter 10.
Figure 7-9. Effect of pH on protein desorption using Zeolite C as the adsorbent material.
Figure 7-10. Colour of the desorbed protein samples under different pH conditions (from pH 8 to pH 14).
Figure 7-11. Protein desorption from Diaguard particles under different pH conditions and subsequent buffer washes.

1st wash= buffer with no NaCl addition, 2nd wash = buffer + 1M NaCl addition, 3rd wash= 1M NaOH wash.
7.4.4 Z-potential analysis

The analysis of the surface charge of zeolite C confirmed that zeolites particles are negatively charged at the pH intended for desorption (8.4-11.4). The results (Figure 7-12) showed that the higher the pH, the higher the electrical potential. However, the electrical potential decreased with particle size, i.e. finer particles (less than 90 microns) have a higher Zeta potential than coarser particles (more than 90 microns).

With the finer particles, at the lower pH (pH 8.4), the Zeta potential increased 14.9% at high pH (pH 11.4). With the coarser particles; a slight decrease (1.8%) was observed when the pH was increased. This result might suggest that the composition of the finer and coarser particles might be slightly different. For example, the coarser particle might content a higher SiO$_2$ content that the finer fraction. However, this would need to be confirmed by a chemical analysis of the fraction, recommended for future work.

![Figure 7-12. Zeta potential analysis of Zeolite C fractions (less than 90 microns, more than 90 microns and “as is” fractions) under different pH conditions.](image-url)
7.5 Conclusions

Diaguard and Zeolite C were the adsorbents that showed better results in terms of protein adsorption. Common sand showed also good protein adsorption properties, but to have comparable results with Diaguard and Zeolite C, the amount of sand utilised was 10 times higher than the amount utilised with Diaguard and Zeolite, suggesting a much lower binding capacity.

All the materials studied have a similar chemical composition, i.e. a high content in silica oxide (more than 70%, except for AW Hyflow with 44% silica content), and the difference in binding capacity might be due to a physical property of the materials: the porosity. More analysis of the material porosity will need to be conducted; however, this aspect will be discussed later in Chapter 10 of this thesis.

From the desorption experiments, it was found that proteins easily desorbed from Diaguard and Zeolite C using alkaline pH (pH>7). Salt addition (ionic effects) does not seem to desorb proteins from Diaguard and Zeolite.

Further studies using continuous adsorption (ion exchange columns) other parameters need to be assessed, such as, particle size. Finer particles may result in higher pressure drops across the ion change column, which translates in lower flowrates.
CHAPTER 8 - PROTEIN CONCENTRATION USING A ZEOLITE PACKED COLUMN: PART I.

Abstract

In this chapter a method for packing zeolite clinoptilolite (zeolite C) into a column (internal diameter 26 mm, height 10 cm) and a method for the concentration of protein contained in pot ale using a liquid chromatography system (Äkta Avant 150) are explained in detailed.

Following the method mentioned above, pot ale samples were loaded into a column at four different flowrates (6, 10, 20 and 30 ml/ min). Breakthrough curves for each case were plotted and at approximately after 5 CV (in all cases), 10% of the initial protein content was measured from the column outlet. The ratio of the protein content in the feed and the column outlet reached a certain level and remained constant for the rest of the experiment, but column saturation ($c/c_0=1$) was not possible to confirm. The level on which the ratio remained constant seemed to increase with the flowrate.

For the elution, it was confirmed (from the batch experiments described in chapter 7) that proteins eluted a high pH (pH>8) and salt (2M NaCl) elution did not seem to release any proteins bound to the zeolite.

A maximum yield of 47.5% was possible to obtain, although the method was not optimised for yield.
8.1 Introduction

In previous chapters it was demonstrated that ion exchange chromatography is a technique that could be used to separate proteins in pot ale. Commercially available media - HiTrap Capto S and Capto Q - were used for this purpose; however, the cost for large scale operations was pointed out as a concern. In chapter Chapter 7, low cost materials were tested for protein adsorption using batch experiments. From these experiments, two materials had potentially showed good results for protein adsorption: Diaguard and Zeolite C.

In this chapter, both materials were packed into a column and tested for protein adsorption using a larger volume of pot ale (1400 ml) under continuous flow. The objectives of this chapter were to achieve flowrates higher than 10 ml/ min (equivalent to 113 cm/ h), depending on column pressure restrictions and to upscale previous experiments conducted with 1 ml columns conducted at 1-2 ml/ min (equivalent to 155-311 cm/h). The maximum recommended pressure for the column used for these experiments (XK-26 columns from GE-Healthcare Life Sciences) was 0.5 MPa.

Before introducing pot ale into the column, tap water was pumped through the column to test flowrates and pressure drops. Flowrates achieved using Diaguard were lower than the target (~2 ml/ min) at the maximum pressure that the column could withstand (0.5 MPa). Further experiments with Diaguard were then stopped, since at larger scale this process would be become unfeasible. Particle size of Diaguard was 40 microns on average, which might explain the large pressure drop across the column and hence the low flow rates.

Flowrates achieved with Zeolite C, were up to 60 ml/ min, but the experiments reported here were conducted between 6 and 30 ml/ min.

The zeolites particles were classified as 300 microns based on its manufacturing specifications (Holistic Valley), although a large proportion of the particles passed the 90 microns sieve (~75%) as can be observed in Figure 8-1.
Figure 8-1. Particle size distribution of Zeolite C (provided by Holistic Valley).
8.2 Methods and Materials

8.2.1 Experiment description

All the experiments were conducted using the Äkta 150 liquid Chromatography system, an XK 26 column and Zeolite C as the protein binding media as presented in Figure 8-2. The pot ale used for the experiment was collected from Glenkinchie (GK).

A total of four experiments were carried out. The conditions maintained during all the experiment are summarised in Table 8-1. Flowrate and elution conditions, i.e. proportion of NaHCO₃ and Na₂CO₃, pH and salt concentration were changed and are summarised in Table 8-2. To equilibrate the column at least 3 CV of acetic acid-sodium acetate buffer (pH 4.5) were used until the conductivity and pH readings were stable.

![Liquid Chromatography system used for the experiments.](image)
Table 8-1. Conditions maintained during the experiments.

<p>| | |</p>
<table>
<thead>
<tr>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>Sample volume</td>
<td>1400 ml</td>
</tr>
<tr>
<td>Column internal diameter</td>
<td>26 mm</td>
</tr>
<tr>
<td>Column height</td>
<td>10 cm</td>
</tr>
<tr>
<td>Packing media</td>
<td>Zeolite C (Holistic Valley)</td>
</tr>
<tr>
<td>Column type</td>
<td>XK 26 (GE Healthcare)</td>
</tr>
</tbody>
</table>

Table 8-2. Summary of experimental conditions, materials and steps.

<table>
<thead>
<tr>
<th>Experiment</th>
<th>I</th>
<th>II</th>
<th>III</th>
<th>IV</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pot ale source</td>
<td>Glenkinchie (GK)</td>
<td>Glenkinchie (GK)</td>
<td>Glenkinchie (GK)</td>
<td>Glenkinchie (GK)</td>
</tr>
<tr>
<td>Flowrate ml/min (cm/h)</td>
<td>6 (67)</td>
<td>10 (113)</td>
<td>20 (226)</td>
<td>30 (339)</td>
</tr>
<tr>
<td>Elution</td>
<td>2M Salt gradient followed by a pH 10.1 step.</td>
<td>pH 10.1 (Sodium carbonate buffer)</td>
<td>20% incremental steps of Na₂CO₃, pH 8-11.</td>
<td>pH 10.1 (Sodium carbonate buffer)</td>
</tr>
</tbody>
</table>

8.2.2 Sample Loading

In each experiment the chromatography protocol comprised four stages: Sample loading, column washing, protein elution and cleaning. On each stage samples were collected for further analysis. For the loading stage, exactly 1400 ml of clarified pot ale was pumped through the column. For pot ale clarification, approximately 2L of pot ale was transferred into 1L bottles (equal amounts) and centrifuged for 1 hour at 4500 rpm using a centrifuge (AVANTI j-265 p, Beckman Coulter, rotor JLA 8.1). The clarified material was then syphoned into a 2L bottle ensuring no solids were collected. The clarified material pumped through the column was collected in 50 ml samples for further analysis.

8.2.3 Washing

The washing stage consisted of pumping 100 mM acetic acid – sodium acetate buffer (pH 4.5) through the column to remove loosely bound proteins or other contaminants. A maximum of 10 CV (Column Volumes) of acetic acid – sodium acetate buffer (pH 4.5)
or until the absorbance at 280 nm ($A_{280}$) was lower than 100 mAU was used for column washing.

### 8.2.4 Elution

For the elution several conditions were tested. These are summarised in Table 9-2 and explained below:

#### 8.2.4.1 Experiment I

Acetic acid-sodium acetate buffer (pH 4.5) with a 2M NaCl gradient (0-50%) over 10 CV, followed by 10 CV of 2M NaCl and finally 10 CV of 0.1M NaHCO$_3$-Na$_2$CO$_3$ buffer (pH 10.1).

#### 8.2.4.2 Experiment II

10 CV of 0.1M NaHCO$_3$-Na$_2$CO$_3$ buffer (pH 10.1).

#### 8.2.4.3 Experiment III

0.1M NaHCO$_3$-Na$_2$CO$_3$ buffer. With the proportion of Na$_2$CO$_3$ increased by 20% in 5 CV steps. pH of this mixture varied from ~8 to ~11.

#### 8.2.4.4 Experiment IV

Same as experiment II (pH 10.1 elution).

### 8.2.5 Cleaning

Finally, the cleaning stage, 3 CV of 1M NaOH were passed through the column and the material was collected for analysis.
8.2.6 Packing and conditioning of the zeolite in the column

The zeolites were conditioned prior to be introduced into the column to give maximum (protein) adsorption capacity and processing flow rates while maintaining a low pressure drop across the column.

Three sequential steps (Figure 8-3) were necessary in order to achieve the objectives mentioned above (maximum protein adsorption, maximum flowrate and minimum pressure drop). The “Particle Classification” step aimed to remove small size particles (less than 90 micrometres diameter). To achieve this, a 90 microns sieve was used to retain zeolites particles bigger than 90 microns which were subsequently transferred into the column. Approximately 11 to 12 cm (lengthwise) of zeolite were used.

Tap water was pumped at a rate between 10 to 20 ml/ min from the bottom of the column (up flow mode in the Akta) keeping the top end of the column opened allowing smaller zeolite particles to float and escape from the column. This process is also known as elutriation, and its objective was to ensure that all the small particles were removed. This process was carried out for at least 1 hour.

The second step “Bed packing” aimed to maximise the quantity (mass) of the solid phase per unit of volume. To achieve this, water was pumped through the column from the top to the bottom. For the XK-26 column (26 mm internal diameter), typical
flowrates utilised were in the order of 40 – 60 ml/min for at least 2 hours. It was important to ensure that no air was trapped inside of the column and that the column was not pressurising over time. By the end of this step the packing compressed to ~10 cm.

The final step, “Bed equilibration” aimed to remove contaminants attached to the resin and to charge the zeolite particles in order to ensure maximum protein-adsorbent interaction. A solution of 1M sodium hydroxide (NaOH) was used for this step at a flowrate of 10 ml/min (downwards flow) for at least 1 hour.

Subsequently, distilled, deionised water was used to remove any salts at a flowrate of 10 ml/min for at least 1 hour. Finally an acidic buffer solution (i.e. acetate) with a pH range between 4 -6 pH units, was used to ensure that solid particles were charged accordingly in order to ensure maximum attachment of proteins to the zeolites. This step normally was carried out at a flowrate between 20 – 30 ml/min for at least one hour.

8.2.7 Peak areas

The areas of the elution peaks (at absorbance of 280 nm) were calculated using the software Unicorn™ 6.1 incorporated with Äkta system. The calculation is defined as the area between the curve and the baseline, between the peak start and peak end, time or volume base. A zero baseline was chosen for the calculations of this work, i.e. straight line at zero level.

8.2.8 Protein yield and concentration factor determination

During the NaHCO₃-Na₂CO₃ elution, samples with absorbance (280 nm) over 500 mAU were collected, the exact volume recorded and measured for soluble protein content. The yield was calculated as the mass of proteins in the elution divided by the mass of proteins in the feed stream.

The concentration factor was calculated by dividing the protein concentration of the eluted stream by the concentration of the proteins in the feed stream. Protein concentration was determined by the Bradford assay.
8.2.9 Pressure measurements

Prior to the introduction of pot ale into the column the pressure was measured with the online pressure monitors incorporated in the Akta system. Three different fluids (distilled water, sodium hydroxide and a 2M NaCl solution), increasing the flowrate in 5 ml/min steps, until the pressure reading were stable and then it was recorded.

The experiment was conducted by analysing individual components of the system that might increase the pressure such as filters, columns and the packing (zeolite).

8.2.10 SDS-Page analysis

The full procedure was explained earlier in Section 6.3.7. But here, additionally, pot ale samples were passed through ultracentrifugation tubes in order to concentrate the proteins and then compared with the samples passed through the zeolite column.
For the ultra-centrifuged samples, pot ale was centrifuged at 4000 rpm for 5 min and the 10 ml of the supernatant were added to the filter cups (Amicon Ultra-15 Centrifugal Filter Tubes Merck Millipore Ltd) and centrifuged at 4000 rpm, 4°C for approximately 2 hours, until the volume of retentate was reduced by 95%. The filtrate was removed and volume of retentate adjusted to 10 ml with distilled deionised water for dialysis. The retentate was centrifuged as previous and this dialysis step repeated once more. The retentate was transferred to an eppendorf tube, the concentration of protein analysed by Bradford assay. The gel was heated at 70°C for 10 min and the electrophoresis run conditions were 150V for 1 hour.
8.3 Results and discussion

8.3.1 Pressure measurement

In the first part of these measurements (Figure 8-5) it can be observed, as expected that higher flowrates increased the pressure system and the more viscous fluids (1M NaOH and 2M NaCl) registered higher pressures than water for the same flowrate. Since the column was rated to a maximum of 0.5 MPa, the protein separation experiments were conducted to maximum flowrate of 30 ml/min.

In the second part (Figure 8-6), only water was used, but the objective of this exercise was to determine if the packing (zeolites) were causing flow restriction (high pressure). From this experiment, it was concluded that flow resistance due to the zeolites were minimal compared to the system components. It is important to mention that some of the tubing used for these experiments have internal diameter of 0.75 mm. Based on the manufacturers guidance, 100 cm of this tubing generates 0.02 MPa of pressure at flowrate of 10 ml/min.
Figure 8-5. System Pressure using different fluids

Figure 8-6. Pressure contribution of filter, column and packing using distilled water.
8.3.2 Breakthrough curves

The breakthrough curves are presented in Figure 8-7. It can be observed that at approximately 5 CV, ~10% of the soluble protein passed through the column in all the experiments (flowrates). Between ~5 and ~10 CV the ratio \( c/c_0 \) increased rapidly and after ~10 CV the ratio remained constant, but a different value was observed for every experiment. The general trend observed was that the higher the flowrate, the higher the \( c/c_0 \) ratio. This is in agreement with theory, since higher flowrates translates into less residence time.

The kinetics behind protein binding into zeolite was studied in more depth in Chapter 10, where a kinetic model is proposed.

![Figure 8-7. Breakthrough curves at different flowrates (6, 10, 20 and 30 ml/ min).](image)
8.3.3 Chromatograms

The chromatograms presented in the following pages (Figure 8-8 to Figure 8-11) report only the elution and cleaning steps of the experiments. The horizontal axis shows the volume of materials passed through the column (ml), the left vertical axis the absorbance of the material leaving the column (at 280 nm) and the right vertical axis the pH. An additional line (green) shows the concentration of Na$_2$CO$_3$ in the buffer mixture.

Figure 8-8 shows the chromatogram of Experiment I (flowrate 6 ml/ min). In this case, a salt (NaCl) gradient was used and no peaks were detected (between ~2200 -2900 ml), indicating that salt did not desorb the proteins attached to the zeolite. This result is agreement with the batch experiments conducted earlier in this thesis (Figure 7-8 and Figure 7-9).

The absorbance reading under these conditions was ~400 mAU. Fluctuations in the absorbance readings were due to flow problems (due to pressure build up in the column, which was corrected in subsequent experiments). After the NaCl buffer introduction into the column, sodium carbonate – sodium bicarbonate buffer (pH 10.1) was pumped and a first peak was detected (~2800 mAU). The second peak was detected with introduction of NaOH.

An interesting observation detected, was that the absorbance peak followed the increase in pH. The increase in absorbance could be associated with proteins “leaking” out of the column. A possible explanation of this phenomenon (reviewed earlier in section 6.2.1) is that by increasing the pH, the proteins bound to the columns will change their electric charge. At low pH (below 5), the proteins will be positively charged and at higher pH, protein charge will change from positive to a negative charge, thus detaching from the zeolites which are negatively charged.
Figure 8-8. Experiment I (6 ml/min) chromatogram.

The blue line of the chromatogram represents the UV absorption (measured at 280 nm) and is reported in mAU on the primary vertical axis of the chromatogram. The green line represents the salt concentration (0-100%, where 100% = 1M NaCl) and the purple line the pH of the sample reported on the secondary axis.
For Experiment II (10 ml/min) the elution was conducted at pH 10.1 (carbonate buffer). Only two peaks could be detected as shown in Figure 8-9. The first peak at pH 10.1 (from ~2700 ml to 3200 ml) and the second peak (between 3600-3800 ml) eluted with 1M NaOH. From the fraction eluted at pH 10.1 (first peak) a sample was taken for SDS analysis, shown in Figure 8-12.

As well, as in Experiment I, in this experiment, the absorbance increased coincided with the pH increase. It is worth noticing though that the peak in the absorbance when the pH was around 8. Above pH ~8, the absorbance started to decrease, although the slope of the tail is less pronounced than the start of the peak. After 3000 ml this is even more evident, the absorbance trend became almost flat. This suggests that the majority of the proteins detached from the column at pH ~8 and the elution process could be stopped when this pH value has been reached.

![Figure 8-9. Experiment II (10 ml/min) chromatogram.](image)

The blue line of the chromatogram represents the UV absorption (at 280 nm) reported in mAU on the primary vertical axis of the chromatogram. The green line represent the salt concentration (0-100%, where 100% = 1M NaCl) and the purple line the pH of the sample reported on the secondary axis. In this experiment, however, no salt or pH gradient was conducted, i.e. the elution was conducted at constant pH (= 10.8).
For Experiment III (Figure 8-10), increasing amounts of Na$_2$CO$_3$ were introduced into the buffer mixture which translated into higher pH buffer. A total of four peaks were detected, although again, as in Experiment I, fluctuation in the absorbance was observed (due pressure build up in the system) and some of the peaks were difficult to detect. Peak 1 and peak 4 were the highest peaks and with the biggest area. This suggests that elution could be simplified, eluting a lower pH (~pH=8) and using only NaHCO$_3$ in the buffer instead of a combination of NaHCO$_3$ – Na$_2$CO$_3$.

**Figure 8-10. Experiment III (20 ml/min) chromatogram.**

The blue line of the chromatogram represents the UV absorption (measured at 280 nm) and is reported in mAU on the primary vertical axis of the chromatogram. The green line represent the salt concentration (0-100%, where 100%= 1M NaCl) and the purple line describes the pH of the sample reported on the secondary axis. The four peaks detected on the chromatogram were labelled as “Peak 1”, “Peak 2”, “Peak 3” and “Peak 4”.
Experiment IV, performed under similar conditions of Experiment II, but at a higher flowrate 30 ml/ min (Figure 8-11) showed again two peaks, at pH 10.1 and another at pH ~14 (1M NaOH).

Again, as in the previous experiments, it was observed that the absorbance increase was related to the pH surge. In this experiment the absorbance peaked at 2600 mAU when the pH was ~8.

![Figure 8-11. Experiment IV (30 ml/ min) chromatogram.](image)

The blue line of the chromatogram represents the UV absorption (measured at 280 nm) and is reported in mAU on the primary vertical axis of the chromatogram. The green line represents the salt concentration (0-100%, where 100%= 1M NaCl) and the purple line describes the pH of the sample reported on the secondary axis.

Although these experiments were carried out using different elution conditions (buffers), it might be reasonable to conclude that the elution of the proteins (measured by absorbance) coincided when the pH was increased. It was also noticed that the absorbance increased rapidly as the pH increased and then slowly decreased (a long tail in the peak).
This phenomenon could be due to the buffering capacity of the reagents used (carbonate buffer and acetic acid for example). In all the chromatograms it was observed that the absorbance peak occurred before pH reached steady state (i.e. the pH of the elution buffer). On average, it took approximately 200 ml (~4 CV) until the elution pH was stable. Apart from extending the duration of the batch, this could affect the purity and concentration of the product. A theory is that the majority of the proteins elute at the start of the peak (start of the peak the product could be potentially protein rich with low salt/ buffer content) and that the tail of the peak could have less protein and more salt (from the buffer). This needs to be checked and future work will need to analyse the different fractions of the peak (start, top and tail for example) to confirm purity and quantities of the eluted proteins.

A possible way to overcome the long tail of the absorbance peak could be to use a more dilute washing buffer, a different buffer (citric acid) or just water. In these experiments the concentration of the washing buffer (acetic acic- sodium acetate) was 100mM, but this could be reduced to 50 mM or less if possible. On a large scale process, process this would result in a more economic process due to less buffer (volume) required to increase the pH of the elution buffer and less salt required to prepare the buffers.

Another way to improve the purity of the eluted material could be by manipulating the control on the Äkta system. In these experiments, the collection of the product was set over 500 mAU, but this can be easily changed. For example, the Äkta could be programmed to start collecting product when the absorbance is 1000 mAU and stop collecting when the absorbance is 2000 mAU. This would need to be checked in future experiments.

Another suggestion for a potential commercial scale production is that the process could be controlled by pH, rather than absorbance if online absorbance instruments are not suitable or available for larger operations. In addition to simplifying the process, this could translate in lower capital requirements too.
8.3.4 SDS-page analysis

Samples from peaks 1 and peaks 2 from Experiment II (Figure 8-9) were analysed using SDS-page analysis (Figure 8-12). R1 and R2 corresponded to the retentate from pot ale samples passed through the Amicon Ultra-15 centrifugal filter tubes, P1 relates to the first peak of experiment 2 (0.1M NaHCO3-NA2CO3 pH 10 buffer) and P2 corresponds to the second peak of experiment 2 (1M NaOH, pH~14).

From the gel, R1 and R2 wells, the presence of protein Z (40 kDa) are clearly visible, however, the same cannot be confirmed for the samples eluted with zeolite C (P1 and P2). This might be a concentration effect, with sample loading too low to show the presence of this band after staining with coomasie blue.

Figure 8-12. SDS-page analysis of experiment II.

The red oval highlights the presence of protein Z around the 40 kDa mark. The molecular weight of the proteins are placed on the left the figure and the columns correspond to the samples analysed.
8.3.5 Peak areas

The areas of the two biggest peaks (based on height) of the chromatograms presented earlier were compared in the graph below (Figure 8-13). For the first 3 experiments, the results are in agreement with theory. Higher flowrates, imply less residence time (contact of pot ale with zeolite), resulting in a potential smaller proportion of protein bound to the zeolites. However, Experiment IV (highest flowrate) showed the second biggest area.

Experiment II (10 ml/ min) and Experiment IV (30 ml/ min) were conducted at similar conditions (except for the flowrate) and similar areas were obtained. It is also worth to mention that for these experiments, 63% of the total peak area was from the NaHCO$_3$-Na$_2$CO$_3$ peak.

![Figure 8-13. Peak areas from the chromatograms of experiments I, II, III and IV.](image)
8.3.6 Protein yield and concentration factor

In Figure 8-14, the protein concentration factor from the NaHCO$_3$-Na$_2$CO$_3$ and NaOH peaks are presented. In experiment I (6 ml/ min), only the NaHCO$_3$-Na$_2$CO$_3$ peak concentration was measured. In all experiments it can be observed that the NaOH peak had highest concentration. The protein concentration factors for NaHCO$_3$-Na$_2$CO$_3$ peaks of all the experiments varied from 1.5 to 2.7, but no clear trend was possible to detect.

![Figure 8-14. Concentration factor of the eluted proteins from experiments I, II, III and IV.](image-url)
The yield was also calculated in all the experiments and the results are presented in Figure 8-15 for the NaHCO₃-Na₂CO₃ and NaOH peaks. When the proteins from both peaks were combined a maximum of 47.5% yield was obtained. However, when only the NaHCO₃-Na₂CO₃ peak was considered only 37% was recovered. The NaOH contained on average ~10% of the protein.

Possible protein losses could be found in the flow through and wash streams and unbound proteins that remained in the column (attached to the zeolites). A full mass balance was carried out in following chapter.

![Figure 8-15. Protein yield from the eluted proteins from experiments I, II, III and IV from the NaHCO₃-Na₂CO₃ and NaOH peaks.](image)
8.4 Conclusions

The purpose for the experiments explained here was to prove that zeolite c is a suitable adsorbent for protein recovery using ion exchange. The zeolite must however, need to be treated prior to the introduction to the column.

Zeolites need to be classified (particles smaller than 90 microns separated and not used in the column), packed (to minimise column volume) and finally equilibrated (wash at low pH to maximise protein adsorption). After the separation of the small zeolite particles, it was possible to obtain high flowrates with negligible pressure drop.

A second objective of this chapter was to develop a protocol for the concentration of pot ale proteins using the Äkta 150 liquid Chromatography system. A method consisting in sample loading, the washing of loosely bound components, followed by the elution of the proteins using a high pH buffer, and finally a cleaning step of the column (with 1M NaOH) was presented in this chapter.

Although the method was not optimised for yield, concentration or purification, it can be said that on average a ~2X concentration factor was achieved and the protein purity (assessed by SDS-page analysis) remained uncertain, since the bands on the SDS- page analysis were not clearly visible.

During the elution step, it was observed in these experiments that the increase of the absorbance readings (related to protein content) matched the pH increase. However, the maximum absorbance reading did not occur when the maximum or steady state pH was reached. A long tail in the absorbance peaks was observed in all the experiments, suggesting further work to improve this process such as the usage of a more diluted buffer for the washing step which precedes the elution step.

In the following chapter the flowrate and buffers will be kept constant and the column length will be modified (from 10 to 30 cm) to understand the effects on the protein recovery process.
Abstract

Continuing with the zeolite packed column experiments described earlier in this thesis, here, in this chapter the effects of column height on protein yield, concentration and the selective removal of carbohydrate and copper were studied.

Four experiments were carried out maintaining sample loading (1400 ml), flowrate (20 ml/ min), column diameter (26 mm) and media packing (zeolite C). Column height (H=10, 20 and 30 cm), media equilibration conditions (pH 4.5 and pH 7) and elution conditions (proportion of NaHCO$_3$ and Na$_2$CO$_3$) were changed.

The combined areas of the elution peaks from the chromatograms were calculated. A 66% difference in peak areas was observed between the maximum (H=30 cm, pH= 4.5) and minimum (H=10 cm, pH 7.0) areas.

Breakthrough curves were plotted to determine the dynamic binding capacity. The highest capacity was obtained for the largest column equilibrated at pH 4.5 (1.8 mg protein/ ml of zeolite).

The effect on the chemical oxygen demand (COD) based on protein content was analysed on the four experiments. On average a 19% decreased in COD levels was observed by centrifugation and an additional 15% by removing the proteins (flow through of the chromatographic step).

Finally a protein, carbohydrate and copper mass balance was conducted. On average, the combined elution fractions contained 38.7% of the original protein content (feed), while carbohydrates and copper in the eluates contained only 1.3% and 8.8%, of the original content, respectively.
9.1 Introduction

Following the method presented in previous chapter, where a process for the adsorption of pot ale proteins using zeolites as the binding material was developed, in this chapter the focus will be on column height and its influence in the process outputs. Specifically, protein yield, concentration and the selective removal of carbohydrate and copper from proteins were studied.

In the process described in the previous chapter, pot ale was pumped through a column packed with zeolites. In theory, proteins contained in pot ale adsorbed to the zeolites and the rest of the pot ale components flow through. This resultant material (the flow through) becomes then “deproteinated” pot ale.

In the Introductory chapter, it was mentioned that proteins have a detrimental effect on Anaerobic Digestion (AD) a common process used for effluent treatment (i.e. pot ale) in the whisky industry. Some authors recognise the need for a pre-treatment process of pot ale prior to AD. Pre-treatment options are based on enzymatic hydrolysis of yeast cells, solid-liquid separations and pH control (Dionisi et al. 2014, Mallick et al. 2010, Tokuda et al. 1999).

A parallel study was conducted by Dr. Thomas Aspray (School of Life Sciences, Heriot-Watt University) using the deproteinated pot ale obtained out of the experiments presented in this chapter. The deproteinated material was tested for batch biochemical methane potential (BMP) and found to be comparable to untreated pot ale based on comparative chemical oxygen demand loading. These results will be presented and made available in literature (Aspray et al., in preparation), however COD analyses will be presented here.
9.2 Methods and Materials

9.2.1 Pot ale samples and analysis

Pot ale samples were collected from four different distilleries (Glenkinchie-GK, Balblair –BB, Balmenach – BM, Strathairn – SE) and kept at 4°C.

The procedure for determining total solids, soluble protein and carbohydrates was explained earlier in this thesis, but a brief description can be seen below.

Total solids were measured by placing 10 ml of the liquid samples in to an aluminium boat and placed into an oven at 105°C for 24 hours. The analyses were performed in triplicate, the weights recorded in grams (g) and volumes in millilitre (mL).

Soluble proteins were measured with the Bradford assay (Bradford 1976) described earlier in the thesis. Conductivity and pH were measured with the online conductivity and pH monitors incorporated in the liquid chromatography system (Äkta Avant 150 from GE-Healthcare). The analyses were performed in triplicate, and the results are reported in mg of soluble protein per litre of pot ale.

Carbohydrates content was measured using a colorimetric assay described in Fournier (Fournier 2001). Briefly, the method involved building a calibration curve using glucose as a standard. Samples and standards (5 to 50 μl) were mixed with 500 μl of 4% phenol followed by 2.5 ml 96% sulphuric acid and then absorbance was measured at 490 nm with a spectrophotometer.

Copper content was detected by flame atomic absorption spectroscopy (FAAS) and the method was also described earlier in the Thesis.

9.2.2 Chemical Oxygen demand

Chemical oxygen demand (COD) was also measured, but since the procedure has not been explained before a detailed description is provided below. The limits of detection for the procedure is in the range of 9-400 mg/ml. Samples with higher concentration were diluted and the results adjusted accordingly. The procedure was taken from

2 ml of sample was mixed with 0.1 ml silver nitrate (Ag NO₃), placed into a digestion tube, and allowed to stand for two minutes. Subsequently 0.1 ml of Chromium III Potassium Sulphate solution was added to 3.70 ml of COD reagent (oxidising mixture). The mixture was thoroughly mixed allowing any evolved gas to escape by removing the cap.

Four blank solutions prepared in exactly the same way as described earlier, but 2.00 ml of distilled water in place of the sample was used. Samples and blanks were placed in the heating block and allowed to digest at 150°C for 2h. After this time, the digestion tubes were allowed to cool in the air for 5 minutes, and then under running water until the temperature was below 20°C.

Once the tubes were cooled, samples and blanks were transferred to 100 ml conical flasks. The residual dichromate was rinsed into the flask using 10 ml of distilled water. Subsequently, no more than two drops of Ferroin indicator (1,10-Phenanthroline ferrous solution 0.025M from Fisher Scientific, catalogue # 10244490) was added and the residual dichromate was titrated with standardised ferrous ammonium sulphate (FAS). The volume of dichromate used to titrated the samples ($V_s$) and blanks ($V_b$) was recorded and used to calculate the COD values (reported in mg/ml) using the formula below, where M is defined in the paragraph below.

$$\text{COD} = 4000M (V_b - V_s)$$

For the standardisation of FAS, 10 ml of concentrated H₂SO₄ was added to 50 ml of distilled water in a 250 ml flask. The acid water mixture was allowed to cool and exactly 5 ml of 0.02083M potassium dichromate was added with one drop of Ferroin indicator. The solution was titrated with FAS. The volume of FAS was recorded and used to calculate the molarity of the FAS solution (M), using the formula below, where V is the volume (ml) of titrant used.

$$M = \frac{5}{8V}$$
9.2.3 Experiment description

All the experiments were conducted using the Äkta 150 liquid Chromatography system, an XK 26 column and Zeolite C as the protein binding media.

The four experiments (Experiment I, II, III and IV) were carried out maintaining the conditions summarised in Table 9-1. Column height (H=10, 20 and 30 cm), media equilibration conditions (pH) and elution conditions (proportion of NaHCO$_3$ and Na$_2$CO$_3$) were changed and are summarised in Table 9-2. To equilibrate the column at least 3 CV of water or appropriate buffer were used until the conductivity and pH readings were stable.

In each experiment the chromatography protocol comprised four stages: Sample loading, column washing, protein elution and cleaning. On each stage samples were collected for further analysis. For the loading stage, exactly 1400 ml of clarified pot ale was pumped through the column. For pot ale clarification, approximately 2L of pot ale was transferred into 1L bottles (equal amounts) and centrifuged for 1 hour at 4500 rpm using centrifuge (AVANTI j-265 p, Beckman Coulter, rotor JLA 8.1). The clarified material was then syphoned into a 2L bottle ensuring no solids were collected. The clarified material pumped thorough the column was collected in 50 ml samples for further analysis.

The washing stage consisted of pumping either water (pH 7) or an acetic acid – sodium acetate buffer (pH 4.5) through the column to remove loosely bound proteins or other contaminants. A maximum of 10 CV (Column Volumes) of water or buffer or until the absorbance at 280 nm ($A_{280}$) was lower than 100 mAU was used for column washing.

For the elution stage a 0.1M NaHCO$_3$-Na$_2$CO$_3$ buffer was pumped through the column. The proportion of NaHCO$_3$ and Na$_2$CO$_3$ in the mixture was varied in each experiment. For example in Experiment I, the proportion of Na$_2$CO$_3$ was increased in 20% incremental steps. For each incremental step the eluted material was collected for further analysis. Each step consumed 5 CV of buffer and the material collected for analysis was higher than 500 mAU ($A_{280}$). Below this limit the material was discarded.
Finally, the cleaning stage, 3 CV of 1M NaOH were passed through the column and the material was collected for analysis.

**Table 9-1. Conditions maintained during the experiments.**

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<th>Sample volume</th>
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<tr>
<td>Packing media</td>
<td>Zeolite C (Holistic Valley)</td>
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<tr>
<td>Column type</td>
<td>XK 26 (GE Healthcare)</td>
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**Table 9-2. Summary of experimental conditions, materials and steps.**

<table>
<thead>
<tr>
<th>Experiment</th>
<th>I</th>
<th>II</th>
<th>III</th>
<th>IV</th>
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</thead>
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<td>Balmenach (BM)</td>
<td>Glenkinchie (GK)</td>
</tr>
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<td>20</td>
<td>30</td>
<td>30</td>
</tr>
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</tr>
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</tr>
</tbody>
</table>

**9.2.4 Peak areas**

The areas of the elution peaks were calculated using the software Unicorn™ 6.1 incorporated with Äkta system. The calculation is defined as the area between the curve
and the baseline, between the peak start and peak end, time or volume base. A zero baseline was chosen for the calculations of this work, i.e. straight line at zero level.

### 9.2.5 Dynamic binding capacity

The dynamic binding capacity (DBC) was calculated with the formula below (Carta and Jungbauer 2010), where \( C_F \) is the protein concentration in the feed, \( t_b \) and \( V_b \) are the breakthrough time and load volume, respectively. \( Q \) is the volumetric flowrate and \( V_c \) is the column volume. Breakthrough was defined as a value of 10% of the feed concentration.

\[
DBC = \frac{C_F V_b}{V_c} = \frac{C_F Q t_b}{V_c}
\]

### 9.2.6 Mass balance

A protein, carbohydrate and copper mass balance of the chromatographic step was conducted. The inputs and outputs volumes were quantified, and with the concentration determined experimentally (described earlier), the mass of protein, carbohydrate and copper was calculated.

An example of the inputs and output is shown in Figure 9-1. It was assumed that the only in input stream containing protein, carbohydrate or copper was the feed (F) stream. For the output streams 9, 10 and 11, it was assumed that the protein content was low or negligible. These streams correspond to fractions on which the absorbance was below the 500 mAU as described earlier.
Figure 9-1. Process flow diagram of the experiments conducted.


9.3 Results and discussion

9.3.1 Chromatograms

The combined areas of the elution peaks from the chromatograms (Figure 9-4) were calculated and the results are presented in Figure 9-2 and Figure 9-3. From the graphs, it can be noticed that a 66% difference in peak areas was observed between the maximum (H=30 cm, pH= 4.5) and minimum (H=10 cm, pH 7.0) areas. Generally, it was observed that the larger the column, the bigger the area, which is in agreement with theory.

In Experiment I, only 3 peaks were detected during the elution phase, although six incremental steps of Na$_2$CO$_3$ were used for the experiment. This suggested to simplify the elution strategy to have only 3 elution steps (0, 50 and 100% Na$_2$CO$_3$) for subsequent experiments.

Experiments III and IV were carried using the same column height (30 cm), but the pre-equilibration step was changed from pH 7.0 to 4.5, an increment of 10% of the peaks areas was calculated. Another difference with both experiments is that in Experiment III the elution was carried out at pH 10.1 (single step) and in Experiment IV, multiple steps were used. In the final elution step (100% Na$_2$CO$_3$), a small peak was detected, which area was calculated at 12% of the total peak area.

The analysis of the contribution of each peak area is presented in Figure 9-3. For example, in Experiment II (Figure 9-4), four peaks were detected during the elution and cleaning stage. From left to right in the chromatogram, the first peak was detected at ~2600 ml using a buffer mixture of 0.1M- 100% NaHCO$_3$ - 0% Na$_2$CO$_3$, corresponding to the lower pH conditions (pH 8.4). A second and third peak was observed at ~3300 ml (50% NaHCO$_3$/ Na$_2$CO$_3$ mixture, pH 9.7) and at ~3900 ml (0% NaHCO$_3$ - 100% Na$_2$CO$_3$, pH 10.8), respectively. The fourth peak (~4500 ml), corresponds to the elution conditions of 1M NaOH.

Following the example from the above paragraph, the areas of peaks 1, 2, 3 and 4 were compared together with the peak areas of the other experiments (Figure 9-3). In all experiments, it was observed that the biggest areas were obtained with the lower Na$_2$CO$_3$ buffer mixture across all experiments (~pH 8). In Experiment II, for example,
peak 1 represented 37%, peak 2 – 27%, peak 3 – 13% and peak 4 23% of the total area of the peaks.

Figure 9-2: Total peak areas of the chromatograms from Experiments I, II, II and IV.

Figure 9-3. Contribution of individual peaks to total area.
Figure 9-4. Chromatograms of Experiments I, II, III and IV.

The blue lines in the chromatograms represent the absorbance measured at 280 nm and it is reported on the primary axis in mAU. The brown lines represent the conductivity, measured in mS/cm and reported in the secondary axis. Finally the green lines denote the proportion of Na$_2$CO$_3$ in the buffer mixture.
9.3.2 Breakthrough curves

Breakthrough curves of Experiments I, II and IV were plotted in Figure 9-5. It can be observed that these three curves followed the same sigmoidal pattern. Up until ~5 CV, ~20% of the protein were bound to the zeolites. From this time (or CV) a rapid increase in the relative protein concentration was detected.

Unfortunately for all the experiments column saturation (\(c/c_0 = 1\)) was not possible to obtain due to the lack of material. Experiments III and IV concluded at approximately 9 CV. At this point the relative protein concentration of the samples were ~60% of the original concentration. For experiment I, a smaller column was used, the experiment continued until ~27 CV of pot ale were passed through the column. In this case, a slow increase in the relative protein concentration (\(c/c_0\)) from ~0.7 (at ~10 CV) to ~0.9 was measured.

![Figure 9-5. Breakthrough curves of Experiments I, III and IV.](image-url)
9.3.3 Dynamic binding capacity

The dynamic binding capacity (DBC) calculations are presented in Table 9-3. For the smaller column (Experiment I, H= 10 cm), the BC was calculated at 1.40 mg of protein per ml of zeolite. Small improvements (20 -30%) were achieved by increasing the column height and by pH pre-equilibration. The highest capacity was obtained for Experiment IV (1.8 mg protein/ ml of zeolite).

Although the DBC of the commercial resins studied earlier (HiTrap Capto Q and Capto S) were not determined, their manufacturer specifications mentioned a DBC of ~100 mg/ ml. Based on the results presented here, zeolites have a capacity of ~50 times smaller than Capto Q/S. The price per unit of volume of these commercial resins is however, about 300 times more expensive. An economic analysis considering column sizing and other aspects must be carried out, to determine if zeolites were a viable material for the process at a larger scale. However, this assessment is out of the scope this thesis.

Table 9-3. Dynamic binding capacity results.

<table>
<thead>
<tr>
<th>Experiment No.</th>
<th>I</th>
<th>II</th>
<th>III</th>
<th>IV</th>
</tr>
</thead>
<tbody>
<tr>
<td>Column Height (cm)</td>
<td>10</td>
<td>20</td>
<td>30</td>
<td>30</td>
</tr>
<tr>
<td>Equilibration pH</td>
<td>7.0</td>
<td>4.5</td>
<td>7.0</td>
<td>4.5</td>
</tr>
<tr>
<td>V_{b,10%} (ml)</td>
<td>100</td>
<td>ND</td>
<td>400</td>
<td>550</td>
</tr>
<tr>
<td>CV_{b,10%}</td>
<td>1.88</td>
<td>ND</td>
<td>2.51</td>
<td>3.45</td>
</tr>
<tr>
<td>DBC_{10%} (mg/ ml)</td>
<td>1.40</td>
<td>ND</td>
<td>1.7</td>
<td>1.8</td>
</tr>
<tr>
<td>Relative improvement to Experiment I</td>
<td>0%</td>
<td>ND</td>
<td>23%</td>
<td>30%</td>
</tr>
</tbody>
</table>

ND: Not determined
9.3.4 Chemical oxygen demand

The effect on the chemical oxygen demand (COD) based on protein content was analysed on the four experiments and the results are presented in Figure 9-6. On average a 19% reduction in COD levels was observed by centrifugation and an additional 15% by removing the proteins (flow through of the chromatographic step).

![Figure 9-6. Chemical oxygen demand of raw, centrifuged and deproteinated pot ale.](image)

In Experiment I, no reduction in COD was observed from the centrifuged and deproteinated pot ale. This result could be due to an error in the centrifuged pot ale analysis, since in the other experiments (II, III and IV) higher CODs were obtained in the samples analysed.

For Experiments III and IV, COD and soluble protein were monitored during the loading stage. The results are plotted in Figure 9-7 and both COD and soluble protein are reported as the fraction of the initial COD and soluble protein, respectively. The soluble protein curves have an exponential shape (tending to reach saturation point = 100%), while the COD curve shape is logarithmic and tending to reach ~75% of initial COD.
% initial COD

% of initial soluble protein content

Figure 9-7. COD and soluble protein breakthrough curves (experiments III and IV only).
9.3.5 Mass Balance

On each experiment (with the exception of Experiment I, samples were not collected), total solids, carbohydrate, soluble protein and copper content of the fractions (i.e. Flow through, washing, elution and cleaning) obtained during the chromatography experiments were compared with the original material (pot ale supernatant). The results are presented in Figure 9-8 and it can be observed from these graphs that carbohydrate and copper content in the flow through (FT) was higher compared to the other fractions and at similar levels of the original material (SN). Soluble protein content, however, was low in the FT fraction and relatively higher in the elution (E1, E2 an E3) and cleaning (C) fractions. To repeat, E1 corresponded to 1st peak (from left to right in the chromatogram) with the lower pH conditions (high NaHCO₃ concentration).

The concentration of the proteins in the elution fractions were lower than feed. These results indicated that the method was not optimised for protein concentration, but this could be rectified by altering the program in the Äkta system. From the chromatograms (Figure 9-4) it can be observed that elution peaks had long tails. The method was programmed to collect fractions when then absorbance reading were higher than 500 mAU. Increasing this parameter to 1000 mAU might increase the concentration factor, but the expense of protein yield loss.

The volumes of each fraction were taken into account; a mass balance was produced for each experiment. Additionally the results were averaged and presented in Figure 9-9.

On average, the combined elution fractions contained 38.7% (SN), the cleaning (C) fraction 30.4%, the flow through fraction (FT) 9.5% and the washing (W) fraction 3.5% of the original soluble protein mass (SN), respectively. By difference, the losses of the process were estimated at 18.0%, which could be explained by the material that was not collected (i.e. mAU < 500). When the elution fractions were analysed separately, E1 contained on average 24.6% of the original protein, while E2 and E3 captured 9.1% and 5.0% of the original protein. These results were in agreement with the previously discussed results of Figure 9-3, where it was noticed that the bigger areas of the chromatograms corresponded with the lower pH elution conditions (E1).
Similarly, based on the average results, carbohydrates in the FT fraction contained 103.3% of the original content. The figure higher than 100%, could be explained due to the use acetic acid for column equilibration. The procedure for carbohydrate analysis described earlier (Fournier 2001), suggest possible interference with other organic compounds such as organic acids. Apart from the W-fraction, all the other fractions had less or close to 1.0% of the original carbohydrate content. The losses, in this case were -12.5%, which confirmed that carbohydrate might have been “added” to the system.

Finally, the copper content in the elution fraction (on average) was 8.8% of the original content. The FT and the C fractions had 76.5% and 14.5% of the original copper, respectively. The losses were calculated as -3.2%.

The significance of this finding is that the protein product (the elution fraction) contains mostly protein. Carbohydrates and copper are mostly found in the in the flow through fractions suggesting that they did not bind to the zeolite.

On a large scale operation, with a potential incorporation of the flow through into an Anaerobic Digestion (AD) unit it would mean that these carbohydrates could theoretically easily be converted into gas (bio-methane). The influence of copper on AD performance would be hard to predict, but as most of biological systems, copper toxicity could become an issue. But, since the amount of copper introduced into the AD unit would lower than pot ale (assuming that the yeast is also removed from pot ale), it is anticipated that the copper content in the flowtrough would not be an issue for AD and gas generation.
Figure 9-8. Total solids, carbohydrate, soluble protein and copper content of the fractions of Experiments II, III and IV.
Figure 9-9. Mass balance (including carbohydrates, protein and copper) of Experiments II, III and IV.
9.4 Qualitative assessment

Samples were taken during the experiments to assess the colour differences. A few examples are mentioned below.

In Figure 9-10, several samples from the loading phase of Experiment III were compared with the original sample (SN). From this pictured it can be seen that the darkness in the sample increased with the time, e.g. tube 2 (fraction 1 of the loading phase) had a light brownish colour and tube 7 (fraction 28 of the loading phase) had a darker brown colour, similar to the original pot ale sample (SN).

In Figure 9-11, the washing (tube 2), elution (tube 3) and cleaning (tube 4) were compared with the original sample (SN) of Experiment III. Also, in difference in the colours of these fractions can be observed. The darkest colour (almost black) corresponded to the elution fraction. The W fraction was almost transparent, while the C fraction had a lighter brown colour compared to the original SN.

Finally, in Figure 9-12, peak fractions of Experiment IV, were compared and also difference in the colour was observed. The elution material from peak 1 (bottle 2) had a black colour, while peak 2 (bottle 3) had a dark brown colour and peak 3 (bottle 4) and the C- fraction (bottle 5) had light brown colour.

There were other compounds that were not analysed during these experiments such as polyphenols (PP), but PP content in pot ale was briefly mentioned in section 3.2.10. PP or protein-PP complex formation could also explain the difference in colour among the fractions. Protein-PP complex formation has received extensive attention in the beer industry and thus precursors to this complex have become the the target for beer haze treatments (Aron and Shellhammer 2010).

From a process control perspective, this qualitative (or visual) assessment can be used to determine protein content if instrumentation becomes unavailable for economic reasons for example.
Figure 9-10. Breakthrough fractions (experiment III).

Legend:
1=Feed (Supernatant),
2-7=Flow through (fractions 1, 7, 13, 19, 25 and 28 respectively)

Figure 9-11. Elution fractions (experiment III).

Legend:
1=Feed (Supernatant),
2=Wash
3= Elution pH 10.1 (peak 1)
4 = 1M NaOH cleaning fraction

Figure 9-12. Elution fractions (experiment IV).

Legend:
1=Feed (Supernatant),
2= Elution peak 1
3= Elution peak 2
4= Elution peak 3
5=1M NaOH cleaning fraction
9.5 Conclusions

The purpose for the experiments explained here was to study the influence of column height on protein capture. By measuring the area of peaks during the elution, it was observed that by increasing the column volume (height) by a factor of three, a 66% increase in peak areas was measured.

During protein elution, a maximum of three peaks were counted. The elution was conducted with an increasing pH profile (increasing Na₂CO₃ concentration). The first peak (from left to right in the chromatograms) corresponded to lower pH conditions.

The peaks or fractions were analysed for carbohydrate, copper and protein. An important conclusion from these experiments is that on average the combined elution fractions contained 38.7% of the original protein, but virtually all of the carbohydrates (>98%) and copper (>90%) were removed. Although the protein yield was relatively low, it was confirmed that the purity of the protein material was increased (carbohydrates and copper removed).

Another aspect of this work was the analysis of the de-proteinated of pot ale (or flow thorough). It was demonstrated that COD was reduced from the pot ale whilst retaining the residual carbohydrate. This protein poor/ carbohydrate rich material could be integrated to an AD process, thus providing energy (gas) to the whisky process.

The benefits of de-proteinated pot ale are related to ammonia formation (due to possible protein degradation) causing operational problems with the AD process (toxicity to AD microorganism). Reduced COD levels could translate into lower residence times (time necessary to complete the bio-chemical reactions in AD), thus translating in lower capital and operational costs. The effect on copper on AD performance remains unclear and further research would be needed to assess this impact. However, and in spite of the de-proteinated pot ale containing the majority of the copper, the amount incorporated into an AD unit would be less than the original pot ale, since part of the copper remains in the protein product and most of the copper (~70% as reported in section 3.3.7) in pot ale is found in the yeast fraction, suggesting that copper content would not negatively influence AD performance.
CHAPTER 10 - PROTEIN ADSORPTION KINETICS

Abstract

From previous Chapter, 1400 ml of clarified pot ale (from several distilleries across Scotland) was passed through a zeolite packed column at several flowrates (6, 10, 20 and 30 ml per minute). The column length was increased from 10 cm to 30 cm.

Samples were collected every 50 ml and analysed for soluble protein content. The experimental data were fitted to the Linear Driving Force (LDF) and Bohart Adams (BA) models. Other models were also investigated, but both LDF and BA models resulted in fairly good approximations.

The estimation of the adsorption capacity of the pot ale proteins on zeolite using these models were also possible. Although the results differed with each model, the LDF model showed higher adsorption capacities. A trend common to both models was however observed: the higher the flow rate, the lower the adsorption capacity. The maximum capacity (~21.03 mg of protein / ml of zeolite) was obtained when the flowrate was decreased to 6 ml/ min.

Shorter columns showed also higher capacities compared to longer columns. This means that some parts of the larger columns were not in contact with the fluid (pot ale) and potentially (zeolite) packing was poor.
## Nomenclature

<table>
<thead>
<tr>
<th>Symbol</th>
<th>Definition</th>
</tr>
</thead>
<tbody>
<tr>
<td>( C )</td>
<td>Concentration of the protein in the mobile phase</td>
</tr>
<tr>
<td>( C_F )</td>
<td>Concentration of the protein in the feed</td>
</tr>
<tr>
<td>( C_0 )</td>
<td>Initial protein concentration</td>
</tr>
<tr>
<td>( C_s )</td>
<td>Protein concentration in mobile phase at particle surface</td>
</tr>
<tr>
<td>( D_0 )</td>
<td>Protein diffusivity in mobile phase</td>
</tr>
<tr>
<td>( D_e )</td>
<td>Effective pore diffusivity</td>
</tr>
<tr>
<td>( d_p )</td>
<td>Particle diameter</td>
</tr>
<tr>
<td>( H )</td>
<td>Height equivalent to a theoretical plat (HETP)</td>
</tr>
<tr>
<td>( h )</td>
<td>Reduced HETP</td>
</tr>
<tr>
<td>( J )</td>
<td>Mass transfer flux at particle surface</td>
</tr>
<tr>
<td>( K )</td>
<td>Equilibrium constant</td>
</tr>
<tr>
<td>( k_{BA} )</td>
<td>Rate constant for the Bohart Adams model</td>
</tr>
<tr>
<td>( k_f )</td>
<td>Film mass transfer coefficient</td>
</tr>
<tr>
<td>( L )</td>
<td>Column length</td>
</tr>
<tr>
<td>( N )</td>
<td>Plate number</td>
</tr>
<tr>
<td>( \langle \tilde{q}_i \rangle )</td>
<td>Particle average adsorbate concentration</td>
</tr>
<tr>
<td>( q_F )</td>
<td>Adsorbed protein concentration in equilibrium with feed</td>
</tr>
<tr>
<td>( q_m )</td>
<td>Monolayer capacity</td>
</tr>
<tr>
<td>( R )</td>
<td>Separation factor for isotherm parameter</td>
</tr>
</tbody>
</table>
**Re**  Reynolds number

**Sc**  Schmidt number

**Sh**  Sherwood number

**T**  Temperature

**u**  Superficial mobile phase linear velocity

**v**  Interstitial velocity

**v'**  Reduced velocity

**r_{pore}**  Particle’s pore radius

**r_p**  Particle radius

**q_0**  Adsorption capacity per unit volume of fixed bed

**q_t**  Protein concentration in the stationary phase at time $t$
Greek characters

\( \eta \)  
Mobile phase viscosity

\( \varepsilon \)  
Extra particle void fraction

\( \varepsilon_p \)  
Intra-particle porosity factor

\( \tau_p \)  
Intra-particle tortuosity factor

\( \psi_p \)  
Diffusional hindrance coefficient

\( \lambda_m \)  
Protein and pore radii ratio

\( \tau_1 \)  
Dimensionless time

\( \phi \)  
Ratio of stationary and mobile phase volumes in column

\( \psi_p \)  
Diffusional hindrance coefficient
10.1 Introduction

In the previous chapters, a chromatography protocol was developed to concentrate proteins contained in pot ale using zeolite as a capture media. It was demonstrated that proteins can be separated from other components in pot ale (i.e. carbohydrates and copper) on a laboratory scale. In order to up scale the chromatography process, two approaches are normally cited in literature (Harrison 1993). The first approach is more empiric and practical, and focuses in the total cycle, i.e. loading, washing, elution, cleaning. This approach will be reviewed in Appendix 2.

The second approach relies heavily on mathematical analysis involving the solution of partial differential equations. The aim of this approach is to determine the controlling mechanism for each part of the cycle of protein adsorption and construct a valid mathematical model. This approach will be used in this chapter, taking into account some of the models already available in literature and focusing only on the loading step of the ion exchange cycle.

The objective of this chapter is then, to obtain in a kinetic model of protein adsorption onto zeolite. Additionally, and based on the model, the protein adsorption capacity can be calculated.

As presented in earlier chapters, the main proteins found in pot ale were LTP1 and Protein Z (ProZ). The models developed in this chapter, will then assume the properties of these proteins and typical ion exchange chromatography conditions for laboratory scale.

A theoretical background, presenting and describing the equations utilised for the mathematical models are included in the following section. Additionally, the utilisation of zeolites is reviewed.
10.2 Theoretical background and literature review

10.2.1 Zeolite pore size

Of importance for protein adsorption modelling is the knowledge of the pore size and shape. Size and shape of zeolite pores found in literature differ substantially. As an example, in the same paper of Cooney mentioned earlier, the pores are described as elliptical windows having approximate dimensions of 7.9 Å by 3.5 Å; the other channels have a set of windows with dimensions of about 4.4 Å by 3.0 Å. Other papers (Kowalczyk et al. 2006) reported the evaluation of pore size distribution of clinoptilolite using scanning electron microscopy (SEM) techniques showed that the pores of fracture-type clinoptilolite grains range from 25–50 nm to 100 nm in size, as well as pores between crystal aggregates up to 500 nm in size. Other values reported in literature are presented in Table 10-1.

Table 10-1. Zeolite pore size found in literature.

<table>
<thead>
<tr>
<th>Reference</th>
<th>Pore size (nm)</th>
</tr>
</thead>
<tbody>
<tr>
<td>(Nilchi et al. 2006)</td>
<td>0.46–0.76</td>
</tr>
<tr>
<td>(Cooney et al. 1999a)</td>
<td>0.45–0.60</td>
</tr>
<tr>
<td>(Sprynskyy et al. 2010)</td>
<td>4.00–19.7</td>
</tr>
<tr>
<td>(Kowalczyk et al. 2006)</td>
<td>25–50</td>
</tr>
</tbody>
</table>

10.2.2 Adsorption kinetics

The kinetics of protein adsorption on ion exchange media is affected by the nature of the adsorption isotherm. A commonly used expression is the Langmuir isotherm presented in Eq. (6) and rearranged in Eq. (7).

\[
q = \frac{q_mKC}{1 + KC} \tag{6}
\]
\[
\frac{q}{q_{ref}} = \frac{C/C_{ref}}{R + (1 - R)C/C_{ref}}
\]

(7)

Where \( q_m \) is the monolayer capacity, \( K \) equilibrium constant and \( C \) the concentration of the protein in the mobile phase, \( R \) is the separation factor defined in Eq. (8) and \( C_{ref} \) is the reference concentration, normally the feed concentration. The isotherm is approximately linear when \( R \to 1 \) and rectangular when \( R \to 0 \).

\[
R = \frac{1}{1 + KC_{ref}}
\]

(8)

10.2.3 Column efficiency

The efficiency of a chromatographic column can be defined in terms of the height equivalent to a theoretical plate (HETP), \( H \), and the corresponding plate number \( N \), relating to each other as shown in Eq. (9), where \( L \) is the column length.

\[
N = \frac{H}{L}
\]

(9)

The expression for the reduced HETP (\( h \)) is presented in Eq. (10) from the reduced van Deemter curve, plotted in Figure 10-1 (Carta et al. 2005), where \( v' \) is the reduced velocity and \( a, b \) and \( c \) are constants with typical values for \( b=2, a=1 \) and \( c=0.05 \). The dominant contribution to reduced HETP and the approximate range of reduced velocities encountered in practical applications are depicted in Figure 10-1. These curves are used for estimates only.

Eq. (11) defines the reduced velocity \( v' \), as a function of particle diameter \( d_p \), interstitial velocity \( v \) and protein diffusivity \( D_0 \). In Eq. (12) \( u \) is the superficial mobile phase linear velocity and \( \varepsilon \) extra particle void fraction.

\[
h = \frac{H}{d_p} = \frac{b}{v'} + a(v')^{0.33} + c v'
\]

(10)
\( v' = \frac{v_d p}{D_0} \)  

(11)

\( v = \frac{u}{\varepsilon} \)  

(12)

**10.2.4 Mass transfer mechanisms**

For large scale operations, mass transfer controls the overall rate of (mass) protein adsorption (Figure 10-1). The main limitations that protein molecules in the mobile phase found in a typical packed bed of porous adsorbent particles cited in literature (Carta and Jungbauer 2010) include: External Mass transfer, Pore diffusion, Kinetic resistance to binding and Solid or adsorbed phase diffusion.

Figure 10-2 depicts the resistances found in typical chromatography process. For the conditions used in the experiments and for large scale chromatography, only the external mass transfer and pore diffusion resistance will be reviewed.
10.2.5 External mass transfer

The external mass transfer resistance is represented by a film mass transfer coefficient $k_f$, defined in Eq. (13), where $J$ is the mass transfer flux at the particle surface, $C$ is the protein concentration in the mobile phase and $C_s$ is the protein concentration in mobile phase at particle surface.

$$J = k_f (C - C_s)$$  \hspace{1cm} (13)

Engineering correlations for mass transfer coefficients in packed adsorption beds, expressed in terms of the Sherwood ($Sh$), Reynolds ($Re$) and Schmidt ($Sc$) numbers are presented in Eq. (14) to Eq. (17) for protein chromatography in laminar flow conditions ($Re<2000$).

$$Sh = \frac{1.09}{\varepsilon} Re^{0.33} Sc^{0.33}$$  \hspace{1cm} (14)

$$Sh = \frac{k_f d_p}{D_0}$$  \hspace{1cm} (15)
\[ Re = \frac{vd_p}{v} \]  
(16)

\[ Sc = \frac{\eta}{\rho D_0} \]  
(17)

### 10.2.6 Pore diffusion

Pore diffusion occurs in pores that are sufficiently large for the solute to diffuse without interacting with the force field exerted by the pore wall. The molecular diffusion coefficient or diffusivity in solution \( (D_0) \) is a function of the size of the solute, the viscosity of the solution and temperature. In general, proteins diffusivities are in the range of \( 10^{-6} \) to \( 10^{-7} \) cm\(^2\) s\(^{-1}\). Tyn and Gusek (Carta et al. 2005) proposed Eq.(18) for globular proteins where \( \eta \) is the mobile phase viscosity (mPa s), \( T \) the temperature (K) and \( M_r \) is the molecular mass (Da).

\[ \frac{D_0 \eta}{T} = 9.2 \cdot 10^{-8} \frac{1}{(M_r)^{1/3}} \]  
(18)

Pore diffusion is typically expressed in terms of the effective pore diffusivity \( (D_e) \) defined in Eq. (19), where \( \varepsilon_p \) is the intra-particle porosity factor, \( \tau_p \) tortuosity factor and \( \psi_p \) is the diffusional hindrance coefficient. There are some correlations available relating \( \varepsilon_p \) and \( \tau_p \) such as Eq. (20), but in general, \( \tau_p \) increases as \( \varepsilon_p \) decreases. Typical \( \varepsilon_p \) values ranges between 0 (for pellicular stationary phases) to 0.9 (or higher for low density gels).

The diffusional hindrance coefficient \( (\psi_p) \) is generally related to the ratio of protein and pore radii \( \lambda_m = r_m/r_{pore} \). Eq. (21) and Eq. (22) have been suggested to relate \( \lambda_m \) to \( \psi_p \) (Carta et al. 2005), where the protein radius \( (r_m) \) can be estimated from the Stokes-Einstein Equation Eq.(23), where \( k_b \) is the Boltzmann’s constant (= \( 1.38 \cdot 10^{-23} \) Joule/K).

\[ D_e = \frac{\varepsilon_p D_0}{\tau_p \psi_p} \]  
(19)
\[ \tau_p = \varepsilon_p + 1.5(1 - \varepsilon_p) \] (20)

\[ \psi_p = \left( 1 + \frac{9}{8} \lambda_m \ln \lambda_m - 1.539 \lambda_m \right) \] (21)

for \( \lambda_m < 0.2 \)

\[ \psi_p = 0.865(1 - \lambda_m)^2 \left( 1 - 2.1044 \lambda_m + 2.089 \lambda_m^3 - 0.984 \lambda_m^5 \right) \] (22)

for \( \lambda_m > 0.2 \)

\[ r_m = \frac{k_b T}{6\pi \eta D_0} \] (23)

**10.2.7 Mass conservation equations**

A simplified model of chromatography with \( M \) components and for each species \( i \), is presented in Eq. (24) to Eq.(27). This model consists of a mass balance equation, mass transfer rate equations and boundary conditions. It assumes plug flow reactor conditions of the mobile phase thorough a uniformly packed sorption bed. In these equations, \( \langle \tilde{q}_i \rangle \) represents the particle average adsorbate concentration and \( z \) is the column axial coordinate. If axial dispersion is neglected, i.e. \( D_L = 0 \), the right hand side of Eq. (24) becomes 0.

\[ \varepsilon \frac{\partial C_i}{\partial t} + (1 - \varepsilon) \frac{\partial \langle \tilde{q}_i \rangle}{\partial t} + \varepsilon v \frac{\partial C_i}{\partial z} = \varepsilon D_L \frac{\partial^2 C_i}{\partial z^2} \] (24)

\[ \frac{\partial \langle \tilde{q}_i \rangle}{\partial t} = f_i(C_j, q_j^*) \quad j = 1, 2, \ldots, M \] (25)

\[ z = 0: \quad C_i = C_i^F + \frac{D_L}{v} \frac{\partial C_i}{\partial z} \] (26)

\[ z = L: \quad \frac{\partial C_i}{\partial z} = 0 \] (27)
Rate equations for each mechanism are presented in Table 10-2. A simplified rate equation, named the Linear Driving Force (LDF) approximation can be used instead of the External film and Pore diffusion models. The LDF approximation is obtained when the driving force is expressed as a concentration difference. An approximate prediction for the rate parameter $k$ in the LDF model can be taken from Eq. (28).

$$k = \frac{60D_e C_0}{d_p^2 q_F} \tag{28}$$

Analytical solutions of this model for each mechanism are presented in Table 10-3 in a dimensionless form where $n$ is the number of transfer units and $\tau_1$ is the dimensionless time defined in Eq. (29), where $L$ is the column length and $q_F$ is the adsorbed protein concentration in equilibrium with feed. When $\tau_1 = 1$ the column has been supplied with an amount of feed equal to that required to obtain complete saturation of the adsorbent at equilibrium with the feed concentration.

$$\tau_1 = \frac{v t/L - 1}{((1 - \varepsilon)q_F)/\varepsilon C_0} \tag{29}$$

<table>
<thead>
<tr>
<th>Mechanism</th>
<th>Rate Equation</th>
</tr>
</thead>
<tbody>
<tr>
<td>External film</td>
<td>$\frac{d\hat{q}}{dt} = \frac{3k_f}{r_p}(C - C_s)$, $\hat{q} = \hat{q}^*$</td>
</tr>
<tr>
<td>Pore diffusion</td>
<td>$\frac{\partial \hat{q}}{\partial t} = \frac{1}{r^2} \frac{\partial}{\partial r} \left( D_e r^2 \frac{\partial c}{\partial r} \right)$ $\hat{q} = \hat{q}^*$</td>
</tr>
<tr>
<td></td>
<td>$(\frac{\partial c}{\partial r})<em>{r=0} = 0$, $(D_e \frac{\partial c}{\partial r})</em>{r=r_p} = k_f(C - c_{r=r_p})$,</td>
</tr>
<tr>
<td>LDF model</td>
<td>$\frac{d\hat{q}}{dt} = k(\hat{q}^* - \langle \hat{q} \rangle)$,</td>
</tr>
</tbody>
</table>

Table 10-2. Rate equations describing protein adsorption in spherical adsorbent particles.
Table 10-3. Constant pattern expressions for the breakthrough curve with the Langmuir or constant separation factor isotherm with R<1.

<table>
<thead>
<tr>
<th>Mechanism</th>
<th>Number of transfer units (n)</th>
<th>Constant pattern solution</th>
</tr>
</thead>
<tbody>
<tr>
<td>External film</td>
<td>( n = \frac{3\phi k_f L}{\tau_p} \frac{v}{v} )</td>
<td>( n(1-\tau_1) = -1 + \frac{1}{1-R} \ln \left( \frac{1-C/C_0}{C/C_0} \right)^R )</td>
</tr>
<tr>
<td>Pore diffusion</td>
<td>( n = \frac{15\phi D_e L}{r_p^2} \frac{v}{v} )</td>
<td>( n(1-\tau_1) = \frac{15}{2} \ln \left[ 1 + \left( \frac{1-C/C_0}{1-C/C_0} \right)^{1/3} + \left( \frac{1-C/C_0}{1-C/C_0} \right)^{2/3} \right] - \frac{15}{\sqrt{3}} \tan^{-1} \left[ \frac{2}{\sqrt{3}} \left( \frac{1-C/C_0}{1-C/C_0} \right)^{1/3} + \frac{1}{\sqrt{3}} \right] + \frac{5\pi}{2\sqrt{3}} - \frac{5}{2} )</td>
</tr>
<tr>
<td>LDF model</td>
<td>( n = \frac{\phi k q_e L}{c_F} \frac{v}{v} )</td>
<td>( n(1-\tau_1) = 1 + \frac{1}{1-R} \ln \left[ \frac{1-C/C_0}{C/C_0} \right]^R )</td>
</tr>
</tbody>
</table>

10.2.8 Bohart-Adams model for rectangular isotherms

The Bohart–Adams (BA) model (Chu 2010) assumes a rectangular isotherm (R=0) with a quasi-chemical rate expression and ignores dispersive effects (axial dispersion and finite resistance to mass transfer). In this model it is assumed that the sorbate-adsorbent interaction can be represented by a quasi-chemical rate described in Eq. (30), where \( k_{BA} \) is the rate constant for the BA model, \( C \) is the protein concentration in the mobile phase, \( q_0 \) is the adsorption capacity per unit volume of fixed bed and \( q_t \) is protein concentration in the stationary phase at time \( t \).

\[
\frac{\partial q_t}{\partial t} = k_{BA} C (q_0 - q_t)
\]  \( (30) \)

The analytical solution for the previous equation is presented in Eq. (31) to Eq. (33), where \( C_0 \) is the initial protein concentration, \( L \) is the column length, \( v \) the interstitial velocity and \( \varepsilon \) the column void fraction.
\[
\frac{C}{C_0} = \frac{\exp(\alpha)}{\exp(\alpha) + \exp(\beta) + 1}
\]  
(31)

\[
\alpha = k_{BA}C_0 \left( t - \frac{L}{v} \right)
\]  
(32)

\[
\beta = \frac{k_{BA}q_0L}{v} \left( \frac{1 - \varepsilon}{\varepsilon} \right)
\]  
(33)

The above equations may be converted to the commonly quoted form of the Bohart–Adams model (Eq. (34)) using simple algebra and two assumptions. First, the two exponential terms \(\exp(\alpha)\) and \(\exp(\beta)\) are usually much greater than unity, so the third term in the dominator (i.e. 1) on the right-hand side of Eq. (31) can be neglected. And second, because of the adsorption time needed for the adsorbate (i.e. protein) to exit the column is far longer than the time needed for the bulk solution to flow from the column inlet to the outlet, which is given by the expression \(\frac{L}{v}\). Therefore, it can be assumed that \(t \gg \frac{L}{v}\) and disregard the \(\frac{L}{v}\) term in the expression for \(\alpha\). These two approximations simplify Eq. (31) to Eq. (34), which has the form a straight line (i.e. \(y = mx + b\), where \(m\) is the slope and \(b\) the y-axis intercept) and \(u\) is the superficial velocity. The slope and y-axis intercept can be obtained graphically, and the unknowns \(k_{BA}\) and \(q_0\) can be calculated.

\[
\ln \left( \frac{c_0}{c} - 1 \right) = \frac{k_{BA}q_0L}{u} - k_{BA}C_0t
\]  
(34)
10.3 Methods and Materials

10.3.1 Column media and pot ale

A full chromatography protocol including packing, equilibration and elution was described earlier in this thesis. But briefly, zeolite (Zeolite Clinoptilolite - Holistic Valley) was packed in a XK-26 column (GE-Healthcare) and 1400 ml of clarified pot ale (from several distilleries across Scotland) was passed through the column at several flowrates (6, 10, 20 and 30 ml per minute). The column length was increased from 10 cm to 30 cm. The 30 cm experiment was repeated with an pre-equilibration step, which involved of running an acidic buffer (pH 4.5, 0.1M sodium acetate-acetic acid) through the column until the conductivity was stable around ~7 mS/cm.

10.3.2 Breakthrough curves

From the 1400 ml of pot ale passed through the column, 50 ml samples were collected during the protein loading stage of the chromatography protocol. The 50 ml samples were subsequently analysed in triplicate for soluble protein content (Bradford assay) using a 96 multi well plate reader. Briefly, 10 µl of the samples were mixed with 200 µl of the Bradford reagent and then the absorption was read at 595 nm. A 0.15M NaCl solution was used as a blank.

10.3.3 Model fitting

The experimental data of the breakthrough curves were fitted to the LDF and BA models explained earlier in the theoretical background section. All graphs presented are based on Protein Z properties and assumed a value of R=0.5.

For the LDF model, the x-axis is composed of the variables n and τ₁. The n values were calculated using the equations provided in Table 10-3 and τ₁ was calculating using MS-Excel solver by finding the minimum value of the sum of the squares of difference between the experimental method and the LDF model, with qF being the variable to adjust. Similarly with the BA model, Eq. (34) was fitted to a line and the values qo and k_{BA} were obtained by finding the slope and y-axis intercept.
10.4 Results and Discussion

10.4.1 Determination of the rate determining step

The experiments were conducted at flowrates between 6 ml/ min to 30 ml/ min, so the reduced velocity ($v'$) falls in the range of 350 to 3500 and 240 to 2400 (dimensionless units) for protein Z and LTP1, respectively (see Figure 10-4). For this range of $v'$, it can be assumed that the controlling mechanism in mass transfer (Figure 10-1). Physical conditions for the experiments assumed a viscosity of $1 \text{ cP} = 10^{-3} \text{ Pa} \cdot \text{s}$ and a temperature $T=298 \text{ K}$. Other parameters used for the model are presented in Table 10-4 (column and adsorbent properties)

Among the proteins properties presented in Table 10-5, it is worth discussing the protein radius ($r_m$). The radius of Protein Z and LTP1 was calculated at 2.5 nm and 1.7 nm, respectively. Zeolites pore size was assumed at 10 nm. As discussed earlier in the Literature Review, there is controversy about the pore size of zeolites, but for the purpose of this model, a pore size greater than the protein size was chosen, in order to test the pore diffusion model. The hindrance coefficient $\psi_p$ for both proteins is bigger than the recommend value of 0.5 (Carta and Jungbauer 2010) to avoid excessive diffusional hindrance and to achieve this, the pore radius needs to be about eight times the protein radius or $\lambda_m<0.125$. In this work, the values for $\lambda_m$ for Protein Z and LTP1 resulted in at 0.25 and 0.17, respectively.

Table 10-4. Column and adsorbent properties assumed for the model.

<table>
<thead>
<tr>
<th>Particle pore size ($r_{pore}$)</th>
<th>10 nm</th>
</tr>
</thead>
<tbody>
<tr>
<td>Particle size (dp)</td>
<td>100 $\mu$m</td>
</tr>
<tr>
<td>Particle porosity</td>
<td>0.5</td>
</tr>
<tr>
<td>Column voidage</td>
<td>0.4</td>
</tr>
<tr>
<td>Column internal diameter</td>
<td>26 mm</td>
</tr>
</tbody>
</table>
Table 10-5. Calculated properties of Protein Z and LTP1.

<table>
<thead>
<tr>
<th>Protein</th>
<th>$M_r$ (kDa)</th>
<th>$r_m$ ($10^{-9}$ m)</th>
<th>$D_o$ ($10^{-7}$ cm$^2$ s$^{-1}$)</th>
<th>$D_e$ ($10^{-7}$ cm$^2$ s$^{-1}$)</th>
<th>$\psi_p$ ($10^{-7}$ cm$^2$ s$^{-1}$)</th>
<th>$k$ ($10^3$ s$^{-1}$)</th>
<th>$Sc$</th>
</tr>
</thead>
<tbody>
<tr>
<td>Z</td>
<td>30</td>
<td>2.51</td>
<td>8.68</td>
<td>0.84</td>
<td>0.24</td>
<td>1.52</td>
<td>11,527</td>
</tr>
<tr>
<td>LTP1</td>
<td>9.7</td>
<td>1.73</td>
<td>12.6</td>
<td>1.99</td>
<td>0.39</td>
<td>3.58</td>
<td>7,912</td>
</tr>
</tbody>
</table>

The influence of pore radius on the number of transfer units is presented in Figure 10-3 for Protein Z and a 20 cm height column. It can be observed that there is linear relationship between pore size and $n_{pore}$ for $r_{pore}$ values < 20 nm. As an example, assuming a flowrate of 20 ml/ min, an increment of the pore radius from 10 to 20 nm ($\lambda_m$ decreases to ~ 0.125), $n_{pore}$ increased from 10 units to ~20 units.
In the case of larger pore radius ($r_{\text{pore}} > 50 \text{ nm}$), the approximate values for $n_{\text{pore}}$ are 70, 30 and 10 (dimensionless).

![Graph showing relationship between pore radius and number of transfer units ($n_{\text{pore}}$) for a 20 cm column length and Protein Z.](image)

**Figure 10-3.** Relationship between pore radius and number of transfer units ($n_{\text{pore}}$) for a 20 cm column length and Protein Z.

The Sherwood number ($Sh$) can also be obtained using Figure 10-4 (secondary axis) and thus the mass transfer coefficient for the external mass film model $k_f$ can be determined and then plugged into the equation for the number of transfer units ($n_{\text{external}}$) for the external film model (first row Table 10-3). A graphical solution is shown in Figure 10-5 (top row) for different column lengths (10-30 cm) and volumetric flowrates used during the experiments, where it can be observed that $n$ decreases with the volumetric flowrate and increases with column length. As an example, for Protein Z and a 30 cm column length, it can be observed that $n_{\text{external}}$ values range between 300 (5 ml/ min) and 60 units (50 ml/ min).
Figure 10-4. Relationship between flowrate, reduced velocity ($v'$) and Sherwood number (Sh).

The number of transfer units for the pore diffusion model ($n_{pore}$) for the flowrates and column lengths mentioned earlier was also plotted in Figure 10-5 (central row). The equation for the calculation of $n_{pore}$ is presented in Table 10-3 (central row). The main difference with external film model, is that in the pore diffusion model, $n$ is proportional to $1/r_p^2$ instead of $1/r_p$ ($r_p$ = particle or zeolite radius). Therefore values of $n_{pore}$ are expected to be smaller than $n_{external}$. Using the example mentioned in the earlier paragraph, for Protein Z and a 30 cm column length, it can be observed that $n_{pore}$ values vary between 60 (5 ml/ min) and 10 units.

The rate determining step is the one that gives the least number of transfer units when evaluated with the data specific to the situation under study. For both protein Z and protein LTP1, it can be observed that the pore diffusion model gives the least number of transfer unit, so it can be assumed that it is limitation step is pore diffusion.

The number of transfer units was also determined with the LDF model ($n_{LDF}$) using the equation presented in Table 10-3 (last row). In this case, the variables $q_F$ was assumed at 20 mg/ ml and $c_F$ at 0.6 mg/ml and $k$ was calculated using Eq. (28) for each protein (results presented in Table 10-5 last row). The results using the LDF model are very close to the pore diffusion model ($n_{pore}$), suggesting that LDF model is a suitable approximation.
Figure 10-5. Number of transfer units (n) for the external film, pore diffusion and LDF models for different volumetric flowrates (Q), proteins (left: Protein Z and right LTP1 protein) and column length (L).
10.4.2 Constant pattern solutions (LDF model)

The results of the experiments at constant flowrate (Q = 20 ml/ min) with column height increased from 10 cm to 30 cm is shown in Figure 10-6, while the results of the experiments on which the flowrate was altered (Q= 6, 10, 20 and 30 ml/ min) and the height maintained (L=10 cm) is shown in Figure 10-7.

For the constant flowrate experiments (Q=20 ml/ min), it can be observed that the model is well suited to the experimental data. For the 10 cm column, n was set to 3 units, while for the 30 cm column, n was set to 5 transfer units. The values for q_F are presented and discussed later in section 10.4.4. The saturation of the column (c/ c_0=1) occurs at values of n(τ_1-1) approaching to 10 and 90% of saturation capacity is reached when n(τ_1-1) ~3.5.

Symbol (♦) represents the experimental data and the solid lines the LDF model.

Assumptions for the LDF model are based on protein Z properties and a value of R =0.5

Figure 10-6. Constant pattern solution (LDF model) for Q= 20 ml/ min and H=10 cm and 30 cm.
From the graphs showing the experiments with fixed column length (L=10 cm) and variable flowrate fitted to the LDF model (Figure 10-7), it can be observed that at the lower and higher flowrates i.e. Q= 6 ml/ min and Q= 30 ml/ min, data fitting to the model becomes less evident compared to the middle range flowrates (Q=10 and 20 ml/ min).

Additionally, it can be detected that the column (adsorbent) at Q= 30 ml/ min, nearly reaches saturation. This could be explained due to poor column packing, i.e. channelling and the reduction of the contact time (residence time) between protein and adsorbent.
Symbol (♦) represents the experimental data and the lines the theoretical model.

Assumptions for the LDF model are based on protein Z properties and a value of R = 0.5.

Figure 10-7. Constant pattern solution (LDF model) for H=10 cm and Q=6, 10, 20 and 30 ml/min.
10.4.3 Bohart Adams model (BA model)

Similarly to the LDF model, the experimental data were fitted to the BA model, but in this case a linear regression analysis was possible and the $R^2$ values were presented in the graphs for both constant flowrate experiments (Figure 10-8) and constant column length (Figure 10-9).

For the constant flowrate experiments it can be observed that for the taller column tests ($L=30$ cm) the model fits better the data than the lower column trial ($R^2= 0.88 – 0.93$ compared to 0.73 for the 10 cm column length).

Symbol (♦) represents the experimental data and the lines the theoretical model.

Figure 10-8. Bohart-Adams solution for $Q= 20$ ml/ min and H=10 cm and 30 cm.
Symbol (♦) represents the experimental data and the lines the theoretical model

Figure 10-9. Bohart-Adams solution for Q = 6, 10, 20 and 30 ml/min and H = 10 cm
Also for the constant height experiment, and like the LDF model, a better fit for the BA model at intermediate flowrates can be observed as shown in the linear fit regression analysis of the curves in Figure 10-10.

Figure 10-10. Linearised Bohart-Adams solution for Q= 6, 10, 20 and 30 ml/ min and H=10 cm
10.4.4 Adsorption capacity

The adsorption capacity was calculated with the BA and LDF models, with the latest showing higher capacities. In general, it was observed, that the difference in capacity between the models decreases as the column height and flowrate increases.

For the experiments at constant flow rate (Figure 10-11) using the LDF model, the adsorption capacity was calculated at 13.96, 7.89 and 5.71 mg of protein per ml of zeolite for the 10 cm, 30 cm and 30 cm (pre-eq) experiments, respectively. For the same experiment, but using the BA model, the adsorption capacity was calculated at 5.37, 3.41 and 2.47 mg/ml for the 10 cm, 30 cm and 30 cm (pre-eq) experiments.

![Figure 10-11. Adsorption capacity vs. column length (L) at Q=20 ml/ min calculated with the BA and LDF models.](image)

Similarly, with the experiments at constant height (Figure 10-12), with the LDF model the adsorption capacity was calculated at 21.03 mg/ml at a flowrate of 6 ml/min while a decrease to 8.57 mg/ml was detected at a flowrate of 30 ml/min. The calculations based on the BA model, for the same points mentioned before, a capacity of 7.86 and 2.06 mg/ml was obtained for the low and high flowrates, respectively.
Figure 10-12. Adsorption capacity vs. volumetric flowrate for the BA and LDF models for the experiment using a 10 cm column height.
10.5 Conclusions

For the conditions carried out during the experiments (i.e. flowrate), it was calculated that mass transfer controls the overall rate of protein adsorption. Among the mass transfer mechanism cited in literature, the external film and pore diffusion models were investigated. Further calculations showed that pore diffusion was the controlling step (least number of transfer units).

The pore diffusion model is however difficult to model and analytical solutions are only available for rectangular or irreversible isotherms (R=0). The other complication with this model is the assumption of the zeolite pore size. There is conflict in literature about zeolites pore size and due to the limitations of time and resources, the zeolites pore size could not be determined experimentally. It was assumed a pore size radius of 10 nm. In contrast, Protein Z and LTP1, the main proteins in pot ale, have a radius of 2.5 and 1.7 nm, based on empirical correlations calculated in this chapter.

There are however simpler approximations. The LDF (linear driving force) and BA (Bohart-Adams) kinetic models for protein adsorption were studied and represented a good approximation for the experiments carried out using zeolite packed columns and pot ale proteins.

Both approximations allowed also the estimation of the adsorption capacity of the pot ale proteins on zeolite. The results differed with each model, with the LDF model showing higher adsorption capacities. A trend common to both model was however observed: the higher the flow rate, the lower the adsorption capacity. The maximum capacity (~21.03 mg of protein / ml of zeolite) was obtained when the flowrate was decreased to 6 ml/ min.

Shorter columns demonstrated also higher capacities (per unit of volume) compared to longer columns. This means that some parts of the larger columns were not in contact with the fluid (pot ale) and potentially (zeolite) packing was poor.
CHAPTER 11 –CONCLUSIONS AND FUTURE WORK

11.1 General Conclusions

From the characterisation chapter (Chapter 3) it was concluded that pot ale, the liquid by-product from malt whisky processing, contain important amounts of protein that are currently underutilised. In the literature review chapter (Chapter 2), it was estimated that approximately 2-3 million tonnes of pot ale are generated in Scotland annually. Assuming 1% (w/v) protein content, it can be easily calculated that there are potentially 20-30 thousand tonnes of protein (worth between £40-60 million) available from pot ale.

Current processes to dispose or treat pot ale include: direct disposal to the sea, spreading on land as fertiliser, evaporation to produce pot ale syrup and anaerobic digestion. However, all these methods for disposal or treatment have their limitations and do not fully exploit the value of the proteins in pot ale.

In this thesis, a process was developed to purify and concentrate the proteins found in pot ale. The process includes a solid-liquid separation and a simultaneous concentration and purification step.

On a commercial scale, the solid liquid separation can be achieved by the use of a disc stack centrifuge. A model to upscale the requirements of large scale centrifugation was presented in Chapter 5.

For the protein purification and concentration step, ion exchange chromatography was proved to be successful. This method takes advantage of the electric charge of the proteins. Due to the low pH of pot ale, most of the proteins are positively charge. A commercially available cation exchanger (HiTrap Capto S) was used to confirm that the proteins can bind to the (Capto S) resin and then elute in a purified and concentrated form.

A problem with commercially available resins is, however, the cost. Low cost alternatives were tested on a batch mode to determine the suitability of protein adsorption in Chapter 7. Among the materials experimented, zeolite c was further analysed for continuous adsorption using a packed column (Chapters 8-10). The zeolite
experiments showed that pot ale can be processed at equivalent flow rates used with the Capto S resin (~300 cm/h) and obtain protein recovery yields around 40% (a maximum yield of 47.5% was obtained). The purity of the product (in terms of specific proteins, i.e. protein Z and LTP1) with zeolite c was lower than Capto S. However the zeolites experiments showed that little copper and carbohydrate can be found in elution fractions the (less than 10% and 1% for copper and carbohydrates, respectively).

This technology is not restricted for the production of protein for aquaculture feeds. With further improvements it could result in the manufacture of specific proteins (fractionation of proteins) with functional properties. This could include alternatives to whey and egg proteins, and proteins which have attractive solubility, emulsifying, foaming and gelling properties and could potentially be included in human foods.

11.2 Review of the objectives

The first objective presented earlier in this work: the development of a novel and sustainable process for the recovery of whisky pot ale proteins, it can be concluded that this objective has been achieved.

In terms of the novelty of the work, no evidence was found of the utilisation of ion exchange chromatography as a method for protein recovery from pot ale. This work served as the basis for the patent GB 1411943.2 (Process and protein product), which allowed to carry out further research on protein recovery from distillery by-products at Heriot-Watt University and the incorporation of the University’s spin-out business “Horizon Proteins”.

The sustainability of the process has yet to be analysed. Data from a large scale operation needs to be collected and compared with existing pot ale processing methods. However, it would be fair to mention that this method requires lower amount of energy to process pot ale than evaporation.

The sustainability of the product (protein) also needs to be assessed with real data, but an advantage of this product is the proximity to the customer compared to other protein sources utilised in Scottish aquaculture, i.e. fishmeal and soy bean meal. A life cycle
analysis could be suggested as future work to evaluate the sustainability of both product and process.

In terms of the waste streams generated with the process developed in this work (yeast and de-proteinated pot ale mainly), it would be reasonable to recommend processing methods. The flow through (or de-proteinated pot ale), for instance can be processed in an anaerobic digester, from which methane can be obtained. The methane can be then burned and used as an energy source for the distillery requirements or for drying requirements of the protein product. It this not clear though, if the de-proteination of pot ale has a detriment in gas production yields. This can be then suggested as future work.

The second objective of the thesis, the assessment and development of ion exchange chromatography as a technique for protein concentration and separation from pot ale, was also carried out successfully. The process has yet to be optimised (yield increase), but several modifications to improve protein yields were mentioned in the thesis. These adjustments can be considered for future work.

Finally, the economic analysis in the appendices allowed the understanding of the economics behind pot ale processing. It resulted evident that pot ale syrup processing and the use of pot ale as fertiliser offer little value compared to the recovery of protein with the method developed in this work.

11.3 Future work

For the cell disruption experiments, some of the topics suggested to be included for future works are: economic analysis of high pressure homogenisation, the utilisation of spent lees for yeast suspension and the increment of yeast cell concentration to reduce processing times.

In terms of the final product further research to determine full nutritional properties of the product is needed. It is also very important to test the product as an aquafeed ingredient, from formulation to feed trials.

Further effort would be also needed to determine market opportunities for individual proteins available in pot ale, such as such as Protein Z and LTP1. It was highlighted in this thesis that Protein Z, for example has well known foaming properties. Exploiting
the functional properties of pot ale proteins could lead to new market opportunities and increase the value of the recovered proteins. It is necessary, however, to forecast demand, price and investigate customers’ requirements for the production of these proteins.

From a process perspective, further work needs to be conducted in order to maximise yield. Areas to look at include zeolite conditioning (pH, particle size, buffers) and flowrates (for loading and eluting). Also the work should not be limited to use zeolite c as the sole resin used for protein recovery. Another suggested research area is process cost reduction. Buffer utilisation and concentration are a few examples that were mention in this thesis (Section 8.3.3).

Another aspect that needs to be investigated is the drying of the protein product. This step would require significant energy (however less than pot ale evaporation since a lower volume would be processed due to the concentration step obtained with the chromatography). An integrated process that recovers the energy potential from the flow through material was discussed earlier.
CHAPTER 12 - REFERENCES


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Appendix 1 - Pot ale evaporation economics

Mass Balance

Assuming a single effect evaporator is used to concentrate the solids in pot ale, an overall and solids (sugars, proteins, yeast cells, etc.) mass balance can be conducted as presented in the equations below (Eq. 1 and Eq.2) and shown in Figure 1. F represents the feed (pot ale), V the Vapour stream (or water removal) and P the product stream (pot ale syrup) and $x_{s,F}$, $x_{s,V}$, $x_{s,P}$ are the solid content for the feed, vapour and product streams.

\[ F = V + P \quad \text{Eq. 1} \]
\[ F x_{s,F} = V x_{s,V} + P x_{s,P} \quad \text{Eq. 2} \]

![Figure 1. Process flow diagram of pot ale evaporation.](image)

Typical solid contents of the process streams in the pot ale syrup evaporation process are $x_{s,F} = 5\%$, $x_{s,V} = 0\%$ and $x_{s,P} = 45\%$. The water removal rate (WRR) or V can be established as the calculation basis and for this example, it can be assumed as $V=1 \text{ ton/h}$. From these values, it can be calculated that $F=1.125 \text{ ton/h}$ and $P=0.125 \text{ ton/h}$. 


Energy Balance

The quantity of steam required per hour (S) and the area of heat transfer surface (A) can be calculated by performing an energy balance around the evaporator.

Assuming that the temperature of the feed (T_f) is 45°C, the working pressure in the evaporator is 77 kPa absolute and that pot ale has similar properties of water: boiling point of 91°C at 77 kPa, specific heat (c_p) = 4.186 x 10^3 J kg⁻¹°C⁻¹ and latent heat of vaporization (λ) = 2281 kJ kg⁻¹ at 91°C. Steam is supplied at 300 kPa. The overall heat transfer coefficient (U) is 2500 J m⁻² s⁻¹ °C⁻¹ [3].

From steam tables, at 300 kPa the specific heat of vaporisation λ_s is 2164 kJ kg⁻¹; and the boiling point temperature at 77 kPa (abs) is 91°C and latent heat λ is 2281 kJ kg⁻¹.

In the absence of heat losses, the heat transferred from the steam to the tubes of the evaporator (q_s) equals that transferred from the tubes to the pot ale (q). From Eq.3 and Eq.4, it can be determined that S = 1154 kg/ h.

\[
q_s = S\lambda_s \quad \text{Eq. 3}
\]
\[
q = Fc_p(T-T_f)+V\lambda \quad \text{Eq. 4}
\]

To determine the Heat transfer Area (A) Eq. 5 is used, where ΔT is the overall temperature drop (and equal to T-T_f). Solving Eq. 5 for A, it gives a value of A= 6.03 m².

\[
q = UA\Delta T \quad \text{Eq. 5}
\]

Steam requirement for the evaporation of pot ale decreases proportionally to the number of effects (n). Table 1 shows the steam requirement (kg steam per kg of water removed) based on the number of effects [4]. Naturally, lower steam consumption means lower operational costs. However, by increasing the number of effects capital requirements escalate.

1 Typical overall coefficients for long tube evaporators are in the range of 2000 -5000 J m⁻² s⁻¹ °C⁻¹
Table 1 - Steam economy per number of effects.

<table>
<thead>
<tr>
<th>n</th>
<th>Kg steam/ kg H₂O evaporated</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>1.2</td>
</tr>
<tr>
<td>2</td>
<td>0.6</td>
</tr>
<tr>
<td>3</td>
<td>0.4</td>
</tr>
<tr>
<td>4</td>
<td>0.3</td>
</tr>
<tr>
<td>5</td>
<td>0.25</td>
</tr>
</tbody>
</table>

Processing and capital costs estimation

Medium sized distilleries in Scotland generate on average 2 - 3 tonnes of pot ale per hour. Based, on this flow rate range, the water removal rate to concentrate pot ale can be estimated at 2 tonnes per hour for medium sized distilleries. Large distilleries in Scotland have a capacity of approximately 10 times bigger than medium size distilleries; hence the water removal rate for pot ale evaporation for large scale distilleries can be estimated at 20 tonnes per hour.

The processing cost of four different scenarios was calculated for medium and large scale distilleries (WRR of 2 and 20 tonne per hour, respectively). The scenarios considered were a processing plant with three Multiple Effect Evaporator (3 ME), five Multiple Effect Evaporator (5 ME), five Multiple Effect Evaporator with Thermal Vapour Recompression (5 ME+TVR) and Mechanical Vapour Recompression (MVR). The latest case (MVR) steam consumption is negligible, but electricity consumption is substantially higher than the other configurations.

Cost of steam and a cost of electricity were assumed at £7 per tonne and 7.2 p/ kWh, respectively [5]. Capital costs were estimated by a combination of quick methods defined by Coulson and Richardson and Evaporators suppliers (APV). The market price of the product (pot ale) is typically sold in the range of £60-70 per tonne [6].

Detailed costs are presented below and a summary of the calculated processing costs is presented in Figure 2. It is clear that concentrating pot ale using evaporation methods for medium size distilleries is not an economically viable option. For larger distilleries, it is an option, but a larger capital investment is required.

Total Fixed Capital Cost (TFC) was estimated as the addition of the Capital Cost (CC) and the Physical Plant Costs (PPC)

\[
TFC = CC + PPC
\]
Capital Cost was estimated as the addition of several items including Major Equipment (ME), equipment erection, piping, (all detailed in the table below). All items are a function of the Major Equipment.

**Table 2 – Capital cost estimation of pot ale evaporation**

<table>
<thead>
<tr>
<th>Capital Cost (CC)</th>
<th>% of ME</th>
</tr>
</thead>
<tbody>
<tr>
<td>Major Equipment (ME)</td>
<td>100%</td>
</tr>
<tr>
<td>Equipment erection</td>
<td>45%</td>
</tr>
<tr>
<td>Piping</td>
<td>45%</td>
</tr>
<tr>
<td>Instrumentation</td>
<td>15%</td>
</tr>
<tr>
<td>Electrical</td>
<td>10%</td>
</tr>
<tr>
<td>Building Process</td>
<td>10%</td>
</tr>
<tr>
<td>SUBTOTAL (CC)</td>
<td>225%</td>
</tr>
</tbody>
</table>

Physical Plant Cost was estimated by the addition of items such as Design and Engineering (see Table below).

<table>
<thead>
<tr>
<th>Physical Plant Cost (PPC)</th>
<th>% of CC</th>
</tr>
</thead>
<tbody>
<tr>
<td>Design and Engineering</td>
<td>25%</td>
</tr>
<tr>
<td>Contractor’s fees</td>
<td>5%</td>
</tr>
<tr>
<td>Contingency</td>
<td>10%</td>
</tr>
<tr>
<td>SUBTOTAL (PPC)</td>
<td>40%</td>
</tr>
</tbody>
</table>

The Operational Cost was calculated adding the variable cost (VC) and the fixed cost (FC). The variable cost was assumed to be mainly electricity and steam. The Fixed cost was calculated as a function of the Total Fixed Capital Cost (TFC), detailed in the table below.
The Total Fixed capital Costs (TFC) for a 2 and 20 tph water removal rate evaporator based on 3 multiple effect evaporator (3 ME), 5 multiple effect evaporator (5 ME), 5 multiple effect evaporator + thermal vapour recompression (5 ME + TVR) and Mechanical vapour recompression (MVR) are shown in the Table below. Figures are presented in thousands of pounds.

<table>
<thead>
<tr>
<th>Fixed costs</th>
<th>% of TFC</th>
</tr>
</thead>
<tbody>
<tr>
<td>Maintenance</td>
<td>10%</td>
</tr>
<tr>
<td>Labor</td>
<td>0%</td>
</tr>
<tr>
<td>Depreciation</td>
<td>10%</td>
</tr>
<tr>
<td>Insurance</td>
<td>1%</td>
</tr>
<tr>
<td>Taxes</td>
<td>2%</td>
</tr>
<tr>
<td><strong>Total Fixed Cost (FC)</strong></td>
<td><strong>23%</strong></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th></th>
<th>3 ME</th>
<th>5 ME</th>
<th>5 ME + TVR</th>
<th>MVR</th>
</tr>
</thead>
<tbody>
<tr>
<td>2 tph WRR</td>
<td>473</td>
<td>1,418</td>
<td>1,575</td>
<td>1,733</td>
</tr>
<tr>
<td>20 tph WRR</td>
<td>663</td>
<td>2,215</td>
<td>2,626</td>
<td>2,728</td>
</tr>
</tbody>
</table>

Figure 2. Cost of pot ale evaporation.
Conclusions

The production of pot ale syrup for small to medium sized distilleries is not economically feasible and requires large amount of energy. It is however, an option for larger distilleries, although significant capital is required to build an evaporation plant.

References

Appendix 2 – Scale up of an ion exchange column for protein recovery from pot ale in medium size malt whisky distillery

Theoretical Background

There are two approaches available for scaling up chromatography. The first approach relies on mathematical analysis involving the solution of partial differential equations. This approach was reviewed earlier in Chapter 10. A second approach is more empirical and practical, and involves only simple equations. This method was taken from Simpson [1].

In this model, the protein production rate \( Q \), a known quantity can be expressed by three factors:

\[
Q = N_c P_c C_d
\]  \hspace{1cm} \text{Eq. 1}

Where \( N_c \) is the number of columns, \( P_c \) productivity per cycle per column (kg protein/cycle/column) and \( C_d \) the number of cycles per day (cycles/day). The production rate (kg protein/day) can be calculated from the protein concentration and output of pot ale from the distillery. In this case the protein rate is 43 kg/day (assuming 3000 L/h pot ale flow rate, with 0.6 g of protein concentration per L and 90% protein yield recovery).

To calculate \( P_c \), Eq.2 is used where, \( C \) is the column dynamic binding capacity (kg protein per L of resin), \( D \) is the column diameter and \( h \) column height (expressed in cm).

\[
P_c = \frac{\pi CD^2 h}{4000}
\]  \hspace{1cm} \text{Eq. 2}

In Eq. 3, the number of cycles per day, is dependent on bed height (h cm), linear flowrate (v cm/h), \( B_L \) (Bed volumes of the loading step, dimensionless), \( B_W \) (bed volumes of the Washing step, dimensionless), \( B_E \) (Bed volumes of the Elution step, dimensionless), \( B_R \) (Bed volumes of the Regeneration step, dimensionless) and \( t_H \) , the holding time (hours). \( B_L \) can be calculated using Eq. 4, where \( C_F \) is the protein concentration in the feed (kg protein/L of pot ale).

\[
C_d = \frac{24}{\frac{h}{v} (B_L + B_W + B_E + B_R) + t_H}
\]  \hspace{1cm} \text{Eq. 3}
\[ B_L = \frac{1000C}{c_F} \quad \text{Eq. 4} \]

Previous equations can be combined and the number of columns \( N_c \) expressed as a function of known parameters as shown in Eq. 5

\[ N_c = \frac{53.05Q \left( \frac{h}{V} \left( \frac{1000C}{c_F} + B_W + B_E + B_R \right) + t_H \right)}{CD^2h} \quad \text{Eq. 5} \]

If the holding time \( (t_H) \) is zero or negligible, \( N_c \) becomes independent of bed height as shown in Eq. 6.

\[ N_c = \frac{53.05Q \left( \frac{1000C}{c_F} + B_W + B_E + B_R \right)}{CD^2V} \quad \text{Eq. 6} \]

Another case is when the loading step dominates all others. This case is applicable when the protein concentration in the feed stream is very low, then Eq. 5 becomes Eq. 7. At low feed concentration the number of columns is independent of bed height and capacity. The column’s primary role has changed from separation to concentration

\[ N_c = \frac{53050Q}{D^2vc_F} \quad \text{Eq. 7} \]

**Assumptions**

In Table 1 the process parameters are presented. A medium size distillery with an average pot ale flowrate of 3000 L/h was used for the calculations. In Table 2, the column parameters were chosen based on the experiments conducted earlier in the thesis. Column height and linear velocity were maintained and a column diameter of 50 cm was assumed. Similarly, in Table 3, the ion exchange parameters used previously in the thesis were used to scale up the ion exchange step.
Table 1. Process parameters.

<table>
<thead>
<tr>
<th>Parameters</th>
<th>Value</th>
<th>Units</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pot ale flowrate</td>
<td>3000</td>
<td>L/ h</td>
</tr>
<tr>
<td>Protein concentration in pot ale (cF)</td>
<td>0.6</td>
<td>g/ L</td>
</tr>
<tr>
<td>Protein recovery yield</td>
<td>90%</td>
<td></td>
</tr>
</tbody>
</table>

Table 2. Column and resin parameters.

<table>
<thead>
<tr>
<th>Parameters</th>
<th>Value</th>
<th>Units</th>
</tr>
</thead>
<tbody>
<tr>
<td>Dynamic binding capacity (C)</td>
<td>0.030</td>
<td>Kg protein/ L resins</td>
</tr>
<tr>
<td>Column height (h)</td>
<td>30</td>
<td>cm</td>
</tr>
<tr>
<td>Column diameter (D)</td>
<td>50</td>
<td>cm</td>
</tr>
<tr>
<td>Linear velocity (v)</td>
<td>700</td>
<td>cm/h</td>
</tr>
</tbody>
</table>

Table 3. Ion exchange process parameters.

<table>
<thead>
<tr>
<th>Parameters</th>
<th>Value</th>
<th>Units</th>
</tr>
</thead>
<tbody>
<tr>
<td>Washing step bed volumes (Bw)</td>
<td>3</td>
<td>-</td>
</tr>
<tr>
<td>Elution step bed volumes (Be)</td>
<td>5</td>
<td>-</td>
</tr>
<tr>
<td>Regeneration step bed volumes (Bc)</td>
<td>5</td>
<td>-</td>
</tr>
<tr>
<td>Holding time (tH)</td>
<td>0.25</td>
<td>h</td>
</tr>
</tbody>
</table>

Results and Discussion

The results are presented in Table 4. From the calculations 3 columns are necessary to keep with supply of pot ale (3000 L/h). The columns specified must contain 59 L of resin. Another alternative would be to double the column diameter (D=100 cm), while maintaining column height (h= 30 cm) and linear flowrate (700 cm/h). This change would decrease column requirements four times \( N_c \sim 1/D^2 \) and only 1 column would be necessary.

In large scale protein purification, rather short columns from 15 to 45 cm are used [2],[3], mainly due to adsorption-desorption behaviour of the protein contradicts almost all of the benefit of long columns. Additionally, long columns contribute to pressure drop and resin compression, which must be avoided. [4]
However, with short, wide columns a good flow distribution is needed. Modern ion exchange columns are equipped with flow distributors that direct the flow evenly over the face of the column minimising dead volumes [5],[6].

Table 4. Results of the ion exchange column design for a medium size malt whisky distillery.

<table>
<thead>
<tr>
<th>Parameters</th>
<th>Value</th>
<th>Units</th>
</tr>
</thead>
<tbody>
<tr>
<td>Number of Columns (Nc)</td>
<td>2.92~3</td>
<td>-</td>
</tr>
<tr>
<td>Productivity per column per cycle (Pc)</td>
<td>1.77</td>
<td>kg protein/cycle/column</td>
</tr>
<tr>
<td>Number of cycles per day (Cd)</td>
<td>7.53</td>
<td>(cycles/day)</td>
</tr>
<tr>
<td>Protein production rate (Q)</td>
<td>38.9</td>
<td>h</td>
</tr>
</tbody>
</table>

References

Appendix 3 – Economic analysis of protein recovery using an ion exchange process

A block diagram of the process developed in this work for the recovery of protein from pot ale is presented in Figure 1. The process counts with 4 unit operations: centrifugation, filtration, a simultaneous protein concentration/purification step (Ion Exchange Chromatography) and a final drying step.

![Diagram of the process](image)

**Figure 1. Process block diagram of pot ale protein recovery**

For the economic model it was assumed that pot ale protein content is 10 g of protein / L of pot ale and the final product has a protein content of 70% on a dry matter basis. The market price of the product (pot ale protein) was assumed at 80% of fishmeal protein £1600 per tonne. A large malt whisky distillery was used for the calculations for the purpose of comparison with alternative pot ale processing technologies (to be discussed in Appendix 5). Annual protein production was estimated at 605 tonne. This information can be seen in Table 2.

Capital costs were estimated by a combination of quick methods [2],[3]. Total Fixed Capital Cost (TFC) was estimated as the addition of the Capital Direct Cost (DC) and the Capital Indirect Costs IC). In this case, the total fixed capital cost required amount ~£2.1 million (Tables 3, 4 and 5).

Processing costs were estimated at ~£1400 per tonne of protein (Table 6), of which spray drying costs were estimated at £131/ tonne. For the spray dryer, it was assumed 350 kW of gas 15 kW were needed [1]. Chromatography resins can be used up to 300 cycles.
### Table 1. Distillery parameters.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Units</th>
</tr>
</thead>
<tbody>
<tr>
<td>Distillery capacity</td>
<td>10</td>
<td>Million litre of pure alcohol per annum (m Lpa/ y)</td>
</tr>
<tr>
<td>Distillery utilisation factor</td>
<td>90%</td>
<td>-</td>
</tr>
<tr>
<td>Pot ale factor</td>
<td>10</td>
<td>L of pot ale/ Lpa</td>
</tr>
<tr>
<td>Protein concentration in pot ale</td>
<td>10</td>
<td>g/ L</td>
</tr>
</tbody>
</table>

### Table 2. Protein price.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Source</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fish meal protein (FMp)</td>
<td>£2000</td>
<td><a href="http://www.indexmundii.com">www.indexmundii.com</a></td>
</tr>
<tr>
<td>Pot ale protein price (% of FMp)</td>
<td>80%</td>
<td></td>
</tr>
<tr>
<td>Price inflation</td>
<td>10% pa</td>
<td></td>
</tr>
</tbody>
</table>

### Table 3. Major equipment (ME) costs.

<table>
<thead>
<tr>
<th>Equipment</th>
<th>Value</th>
<th>Source</th>
</tr>
</thead>
<tbody>
<tr>
<td>Centrifuge</td>
<td>£500,000</td>
<td>Interview with suppliers</td>
</tr>
<tr>
<td>IEX columns</td>
<td>£90,000</td>
<td>Interview with suppliers</td>
</tr>
<tr>
<td>Spray drier</td>
<td>£300,000</td>
<td></td>
</tr>
<tr>
<td>Total Major Equipment (TME)</td>
<td>£890,000</td>
<td></td>
</tr>
</tbody>
</table>

### Table 4. Direct Cost (DC)

<table>
<thead>
<tr>
<th>Item</th>
<th>Value</th>
<th>% of TME</th>
</tr>
</thead>
<tbody>
<tr>
<td>Major Equipment</td>
<td>£890,000</td>
<td>100%</td>
</tr>
<tr>
<td>Installation</td>
<td>£347,100</td>
<td>39%</td>
</tr>
<tr>
<td>Piping</td>
<td>£178,000</td>
<td>20%</td>
</tr>
<tr>
<td>Instrumentation</td>
<td>£133,500</td>
<td>15%</td>
</tr>
<tr>
<td>Electrical</td>
<td>£89,000</td>
<td>10%</td>
</tr>
<tr>
<td>Building process</td>
<td>£89,000</td>
<td>10%</td>
</tr>
<tr>
<td>TOTAL Direct Cost (DC)</td>
<td>£1,726,600</td>
<td></td>
</tr>
</tbody>
</table>

### Table 5. Indirect Cost (IC)

<table>
<thead>
<tr>
<th>Item</th>
<th>Value</th>
<th>% of DC</th>
</tr>
</thead>
<tbody>
<tr>
<td>Design and Engineering</td>
<td>£172,660</td>
<td>10%</td>
</tr>
<tr>
<td>Contractor's fees</td>
<td>£86,330</td>
<td>5%</td>
</tr>
<tr>
<td>Contingency</td>
<td>£172,660</td>
<td>10%</td>
</tr>
<tr>
<td>SUBTOTAL</td>
<td>£431,650</td>
<td></td>
</tr>
</tbody>
</table>
Table 6. Processing costs.

<table>
<thead>
<tr>
<th>Item</th>
<th>Factor</th>
<th>Basis</th>
<th>Cost, thousand £/y</th>
<th>Cost, £/ton of protein</th>
</tr>
</thead>
<tbody>
<tr>
<td>Raw materials</td>
<td>0.0</td>
<td>of pot ale disposal cost</td>
<td>0.0</td>
<td>0</td>
</tr>
<tr>
<td>Operating labor</td>
<td>1.0</td>
<td>Operators @ £30k pa</td>
<td>30.0</td>
<td>50</td>
</tr>
<tr>
<td>Operating supervision</td>
<td>0.15</td>
<td>of operating labour</td>
<td>4.5</td>
<td>7</td>
</tr>
<tr>
<td>Utilities</td>
<td></td>
<td>Calculated (Table 7)</td>
<td>89.2</td>
<td>148</td>
</tr>
<tr>
<td>Maintenance and repairs</td>
<td>0.06</td>
<td>of FCI</td>
<td>129.5</td>
<td>214</td>
</tr>
<tr>
<td>Operating supplies</td>
<td>0.15</td>
<td>of maintenance &amp; repair</td>
<td>19.4</td>
<td>32</td>
</tr>
<tr>
<td>Laboratory charges</td>
<td>0.15</td>
<td>of operating labour</td>
<td>0.7</td>
<td>1</td>
</tr>
<tr>
<td>Resins and buffers</td>
<td></td>
<td>Calculated (Table 7)</td>
<td>356.6</td>
<td>590</td>
</tr>
<tr>
<td><strong>Variable cost =</strong></td>
<td></td>
<td></td>
<td><strong>630.0</strong></td>
<td><strong>1,042</strong></td>
</tr>
<tr>
<td>Depreciation</td>
<td>0.1</td>
<td>Of FCI</td>
<td>215.83</td>
<td>357</td>
</tr>
<tr>
<td><strong>TOTAL COST</strong></td>
<td></td>
<td></td>
<td><strong>1,399</strong></td>
<td></td>
</tr>
</tbody>
</table>

Table 7. Cost of utilities.

<table>
<thead>
<tr>
<th></th>
<th>Factor</th>
<th>Basis</th>
<th>£/kWh</th>
<th>£/m³</th>
</tr>
</thead>
<tbody>
<tr>
<td>Gas</td>
<td>0.025</td>
<td></td>
<td>£/kWh</td>
<td></td>
</tr>
<tr>
<td>Electricity</td>
<td>0.12</td>
<td></td>
<td>£/kWh</td>
<td></td>
</tr>
<tr>
<td>Water</td>
<td>0.26</td>
<td>£/m³</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Solvent</td>
<td>0.5</td>
<td>£/m³</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Resin</td>
<td>0.5</td>
<td>£/m³</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Cost of fuel can be offset by anaerobic digestion (AD) of de-proteinated pot ale. An economic analysis of AD is presented in Appendix 4

References

Appendix 4 – Economic analysis of anaerobic digestion of pot ale

**Gas Production**

| Pot ale COD | 0.035 kg COD/ L pot ale | Range 0.030-0.060 |
| Factor | 0.245 Nm3/ kg COD |
| Heat capacity | 8.30 kWh/ Nm3 methane | 6.1-8.3 kWh/ Nm3 (55-75% v/v Methane) |
| Volume | 857,500 Nm3/ y |
| Energy | 7,117,250 kWh/ y |

**Renewable Heat incentive (RHI)**

| RIH tariff | 6.8 p/ kWh |
| Subsidies | £483,973 |

**Heavy fuel Oil**

| Heat value | 11.72 kWh/ L |
| Cost | 4.5 p/ kWh |
| Fuel expenses (offset) | £320,276 |

**Capital Cost AD**

| Boiler | £130,000 |
| AD | £1,200,000 |
| Total | **£1,330,000** |
Appendix 5 – Economical comparison of pot ale processing technologies

As discussed earlier, a comparison of different pot ale processing technologies is presented in this Appendix. The processing technologies available are: (1) fertiliser, (2) anaerobic digestion and (3) protein recovery + anaerobic digestion.

For technology (3) it was assumed that gas production is reduced by 40% due to COD drop (as review in chapter 9). Capital reductions due to lower COD were not incorporated in the model.

For the economic evaluation, the net present value (NPV) was calculated assuming a 10% discount rate over 5 years.
<table>
<thead>
<tr>
<th>Pot ale products</th>
<th>Initial investment (£k)</th>
<th>Yield (per tonne of pot ale)</th>
<th>Units</th>
<th>Market Price per unit of product</th>
<th>Processing Cost per unit of product</th>
<th>Income (loss) per unit of product</th>
<th>Median Size distillery Income (loss) pa (£k)</th>
<th>NPV (£k)</th>
<th>NPV + disposal savings (£k)</th>
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</thead>
<tbody>
<tr>
<td>Fertiliser</td>
<td>£0</td>
<td>1 tonne</td>
<td></td>
<td>£0.0</td>
<td>£12.50</td>
<td>-£12.50</td>
<td>-£1,250</td>
<td>-£4,738</td>
<td>£0</td>
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<tr>
<td>Pot ale syrup</td>
<td>£1</td>
<td>1/9 tonne</td>
<td></td>
<td>£70.0</td>
<td>£48.00</td>
<td>£22.00</td>
<td>£244</td>
<td>£926</td>
<td>£5,664</td>
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<tr>
<td>Biogas (75% methane)</td>
<td>£1,330</td>
<td>8.58 m3</td>
<td></td>
<td>£0.94</td>
<td>£0.04</td>
<td>£0.90</td>
<td>£771</td>
<td>£1,593</td>
<td>£6,331</td>
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<tr>
<td>Gas+proteins</td>
<td>£3,430</td>
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<td>£1,263</td>
<td>£1,356</td>
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<tr>
<td>Gas</td>
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<td>5.15 m3</td>
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<td>£0.94</td>
<td>£0.04</td>
<td>£0.90</td>
<td>£463</td>
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<tr>
<td>Proteins</td>
<td></td>
<td>1.0% tonne</td>
<td></td>
<td>£2,000</td>
<td>£1,400</td>
<td>£600</td>
<td>£600</td>
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<tr>
<td>Yeast</td>
<td></td>
<td>1.00% tonne</td>
<td></td>
<td>£200</td>
<td>£0.00</td>
<td>£200</td>
<td>£200</td>
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