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Abstract

This study assessed the functional profile (foaming, emulsifying and rheological properties), proteomic and metabolomic composition of a naturally foaming and currently unexploited co-product (centrate) from the Quorn fermentation process. Due to the low environmental footprint of this process the centrate is a potential source of sustainable functional ingredients for the food industry. A range of fractions were isolated from the centrate via successive ultrafiltration steps. The retentate 100 (R100) fraction, which was obtained following a 100 kDa ultrafiltration, displayed good foaming, emulsifying and rheological properties. R100 solutions and oil-in-water emulsions displayed high viscosity, while R100 solutions and hydrogels showed high viscoelasticity. R100 foams displayed high stability while oil-in-water R100 emulsions showed small and stable oil droplet size distributions. Large mycelial aggregates were reported in R100 solutions and gels, correlating with their high viscosity and viscoelasticity. A dense mycelial network was observed in R100 foams and contributing to their stability. In parallel tensiometry measurements highlighted the presence of interfacially-active molecules in R100 which formed a rigid film stabilising the oil/water interface. A number of functional metabolites and proteins were identified in the centrate, including a cerato-platanin protein, cell membrane constituents (phospholipids, sterols, glycosphingolipids, sphingomyelins), cell wall constituents (chitin, chitosan, proteins), guanine and guanine-based nucleosides and nucleotides. This study highlighted the potential of functional extracts from the Quorn fermentation process as novel ingredients for the preparation of sustainable food products and the complex and specific nature of the centrate’s functional profile, with contributions reported for both mycelial structures and interfacially-active molecules.

Keywords
Quorn, co-product, centrate, rheology, foam, emulsion
Introduction

Animal-derived functional ingredients including milk and egg proteins play a key role in the food industry due to their ability to form gels and to stabilise oil droplets in emulsions and air bubbles in foams. However, due to their high environmental costs and market volatility, the food industry is looking for sustainable alternatives [1]. One of the strategies employed consists in screening unexploited co-product streams from the food industry for extraction of potential functional alternatives. In this context the production of fungal proteins by Marlow Foods for use in their meat alternative product Quorn is a potential source of sustainable functional ingredients. The term mycoprotein refers to the high-protein biomass produced by fermentation of the fungus *Fusarium venenatum* A3/5 (ATCC PTA-2684) by Marlow Foods, which forms the basis of their Quorn brand products. Mycoprotein contains all essential amino acids [2] and the net protein utilisation value of *Fusarium venenatum* mycoprotein is comparable to that of milk [3]. Moreover, the fermentation of starch into protein by *F. venenatum* results in 90% lower emission of greenhouse gases and benefits in terms of land and water footprints in comparison with beef products [4].

Currently mycoprotein biomass is marketed as the meat substitute Quorn. However, a liquid co-product stream that contains residual hyphal biomass, carbohydrates, nucleotides and proteins as well as the residues of the fermentation feedstock is produced as part of the fermentation process and is currently unexploited. This co-product (centrate) is generated as a result of an RNA-reduction step in which the fermentation broth is first heat-shocked above 68°C for 30 to 45 minutes, then further heated at 90°C and finally centrifuged [5]. The resulting solid deposit (mycoprotein biomass) is processed into a dough ready for conversion to the meat-like texture characteristic of Quorn foods, while centrate is the liquid centrifugation supernatant. This heat shock step carried out on the fermentation broth stops growth, disrupts ribosomes and activates endogenous RNAases which break down cellular RNA to nucleotides [5]. Heat shock also induces diffusion of a fraction of the cell components through the cell wall, including nucleotides and proteinaceous material [5]. The resulting presence of these cell components in the centrate leads to foaming in the centrifuge, indicating the presence of surface-active material.

With only 49 proteins listed on Uniprot (Universal Protein Resource), the *Fusarium venenatum* proteome has so far not been fully characterised [6]. Several protein families produced by filamentous fungi have been reported to be surface-active, including hydrophobins, repellents, rodllins, chaplins, SapB, fungispumins and cerato-platanins [7-9]. One hydrophobin and eight proteins of the cerato-platanin family were reported on Uniprot for *Gibberella zeae* (*Fusarium graminearum* or wheat head blight fungus) [6], which belongs to the same *Fusarium sambucinum* species complex as *Fusarium venenatum*. Filamentous fungi secrete cerato-platanin proteins (CPPs) extracellularly into the culture filtrate, but they have been also reported within the cell wall of fungal hyphae and spores [10-11]. Frischmann et al. [9] reported that the cerato-platanin EPL1 from *Trichoderma atroviride* self-assembles into regularly patterned protein biofilms at air/water interfaces and that EPL1 protein solutions exhibited strong foaming properties.

Denny et al. [2] reported a 6% dietary fibre content for mycoprotein biomass with a predominance of the fungal cell-wall components beta-glucans and N-acetylglucosamine polymers (chitin and chitosan). Beta-glucan extracts from barley and oats are used as functional ingredients in the food industry due to their thickening, emulsifying, water-holding capacity, texturising, stabilising and gelling properties [12]. Similarly chitin, chitosan and their derivatives
are used as functional ingredients in the food industry due to their texture-controlling, emulsifying, thickening, gelling and stabilising properties [13], including those of fungal origin [14]. Nucleotides were reported as breakdown products of RNA during the RNA-reducing heat-shock step [5]. Viscosifying and gelling properties have been reported for guanine-based nucleotides due to the ability of the nucleobase guanine to self-associate [15]. The aim of this study was to assess the proteomic and metabolomic composition of the centrate and the functional profile (foaming, emulsifying and rheological properties) of various fractions derived from the centrate in order to act as a benchmark for the assessment of the exploitation potential of the co-product stream.

Material and methods

Sample preparation

Centrate samples were collected and frozen. Following thawing the centrate underwent a series of ultrafiltration steps using a range of Vivaflow 200 crossflow cassettes (Sartorius, UK) connected to a Masterflex Easy-Load peristaltic pump (Sartorius, UK). Following an initial 100 kDa ultrafiltration, the resulting retentate 100 or R100 (composed of molecules larger than 100 kDa) was collected, while the filtrate 100 underwent further ultrafiltration, using either a 5 kDa membrane or a 50 kDa followed by a 5 kDa membrane. Samples of the centrate and resulting retentates were freeze-dried in a Super Modulyo unit (Edwards, UK).

Nitrogen content

As a wide range of surface-active molecules potentially present in the centrate fractions contain nitrogen (including fungal cell membrane and cell wall constituents such as phospholipids, glycosphingolipids, sphingomyelins, chitin, chitosan and glycoproteins) the Kjeldahl method [16] was used to provide a guideline for preparation of standardised sample quantities for functional tests. 0.1 g of each powder was digested in concentrated sulphuric acid (92%) using a Kjeltec Basic Digestion Unit 20 (Foss, UK) at 440°C in the presence of a selenium catalyst. Distillation of the digested samples into boric acid was carried out using a Tecator Kjeltec 8100 Manual Distillation Unit (Foss, UK). The distilled samples were then titrated using 0.1N hydrochloric acid. The % nitrogen was calculated using the following formula:

\[
Nitrogen = \frac{\text{Titration volume (ml)} \times 14.007}{\text{Sample weight (g)} \times 100}
\]

The % nitrogen obtained was then multiplied by a general conversion factor of 6.25. Measurements were repeated three times.

Gel electrophoresis

Sodium dodecyl sulfate polyacrylamide gel electrophoresis (SDS–PAGE) and native PAGE were performed on the centrate fractions using a Mini-Protean Tetra Cell System with TGX 4-20% Tris-glycine gels (Bio-Rad Laboratories
Lt., UK). 1% w/w nitrogen-containing material (NCM) solutions of the centrate fractions were prepared in deionised water and stirred for one hour.

SDS-PAGE samples were prepared by mixing equal proportions of solution and Laemmli 2x concentrate sample buffer (Sigma Aldrich Ltd., UK) followed by heating at 70°C for 10 minutes. Native PAGE samples were prepared by mixing the solution with Novex native TrisGly 2x sample buffer (Life Technologies, UK) in equal proportions. A See Blue Plus2 pre-Stained standard (Life Technologies Ltd., UK) was used as molecular weight marker. The gels were run in Tris/Glycine/SDS buffer (Bio-Rad Laboratories Ltd., UK) at 100 V for 1 hour and subsequently stained in a Coomassie brilliant blue solution (VWR Ltd., UK) and destained in glacial acetic acid/methanol/deionised water 1/4/5. Gels were scanned using a ChemiDoc XRS+ imaging system (Bio-Rad Laboratories Ltd., UK) and analysed using the associated Image Lab software.

Proteomics

Protein digestion was carried out according to Le Bihan et al. [17]. Samples were denatured in 8M urea, reduced by incubation with dithiothreithol prior to cysteine alkylation with iodoacetamide and overnight digestion with 60 μg trypsin. Four μg of peptide samples were acidified with 1% formic acid before centrifugation and vacuum–drying. The samples were analysed by capillary HPLC–MSMS, using 140-minute gradients as described by Martin et al. [18] on a hybrid LTQ–Orbitrap XL instrument (Thermo-Fisher, UK). HPLC quality acetonitrile (Fisher, UK), Suprapure formic acid and sequencing grade trifluoroacetic acid (Merck, Germany) were used. Multicharged (2+, 3+ and 4+) ion intensities were extracted from the LC-MS files and the Mascot Version 2.4 software (Matrix Science Ltd, UK) was used to compare the MSMS data against the NCBI protein database with the following search parameters: maximum missed-cut value of 2, variable oxidation, N–terminal protein acetylation and fixed carbamidomethylation, precursor mass tolerance of 7 ppm and MSMS tolerance of 0.4 Da. A significance threshold of less than 0.05 and a minimum peptide cut-off score of 20 were set. Proteins which were identified based on the Fusarium graminearum proteome and quantified with 2 or more peptide sequences were retained.

Metabolomics

Alcohol insoluble residues of the centrate were produced in 10 ml 70% ethanol. The vials were left overnight at room temperature and the residues were subsequently washed three times with 10 ml of a 10/10/3 chloroform/methanol/water solution by centrifugation at 2,700 x g for 10 minutes. The wash step supernatants were pooled and run in positive (pos) and negative (neg) ion mode on an 6560 Agilent Quadrupole Time of Flight (IM-QTOF) mass spectrometer (MS) (Agilent, UK). Samples were run in both ion mobility QTOF mode and MSMS QTOF mode. A 100mm x 2.1mm, 1.8um ACE C18-PFP column was used. Two solvents were used in separation (solvent A: 50% methanol, solvent B: 95% acetonitrile) using gradient elution at a flow rate of 0.2 ml/min (0 min: 10% B, 5 min: 30% B, 10-12 min: 95% B, 15-16 min: 10% B). The data were analysed using Mass Profiler (MP) and were also extracted using Profinder before uploading to Mass Profiler Professional (MPP) in order to carry out clustering
analysis and visualise feature abundance. Alignment of samples with the Metlin accurate mass metabolite database was carried out with accurate mass matching to 10 ppm.

Rheological properties

Rheological measurements of centrate fraction solutions and oil-in-water emulsions were performed using a Bohlin Gemini controlled stress rheometer (Malvern Instruments, UK) using cone-and-plate geometry. Measurements were repeated three times.

For viscosity measurements 10% w/w NCM solutions of the centrate fractions were prepared in deionised water and stirred for two hours. The commercial whey protein concentrate product Lacprodan 87 (L87, Arla, Denmark) was used as control for all functionality tests. The L87 solution was prepared at a higher protein concentration (15.3%) to match the solid content of a 10% NCM R100 solution (17.65% solids) and assess the possible influence of solid content on viscosity. In order to prepare emulsions for rheology, 15 g solutions of 5% w/w NCM centrate fractions were mixed with 5 g of sunflower oil and homogenised using an Ultra–Turrax high speed homogeniser (IKA–Werke GmbH, Germany) for 1 min. Viscosity measurements were performed using a 4°/40 mm cone (gap 150 μm) at 20°C. The instantaneous viscosity (Pa.s) was measured through a shear rate sweep test ranging from $10^3$ s$^{-1}$ to 50 s$^{-1}$ in up-down mode (15-minute up-sweep, 15-minute down-sweep).

For gelation tests preliminary oscillatory measurements of elastic and viscous moduli (G’ and G’’) were carried out on 10% w/w NCM centrate fraction solutions at 1 Hz over a strain amplitude sweep test ranging from 0.00005 to 50 to determine the linear viscoelastic region of each sample. Small-amplitude oscillatory measurements were then performed using a 2°/40 mm cone (gap 70 μm). G’ and G’’ were measured through a temperature sweep test ranging from 40 to 90°C in up-down mode (15-minute up-sweep, 15-minute down-sweep). The applied strain was chosen from within the linear viscoelastic region for each sample and the oscillation frequency was 1 Hz.

Foaming properties

Two types of foaming methods (gas-sparging and frothing) were carried out. The stability of foams produced by the centrate fraction solutions was first assessed using a gas-sparging test according to Rudin [19] with minor modifications. 40 g solutions of 1% w/w NCM centrate fractions were stirred for one hour. Gas-sparging was carried out at constant flow rate of CO$_2$ (100 cm$^3$/min) until the foam reached the top indicator of the column. The foam stability was quantified as the time needed for the foam to collapse to the second indicator at the bottom of the column. Measurements were repeated three times.

For frothing tests 15 g solutions of 1% w/w NCM centrate fractions were prepared in 50 ml glass beakers. The solutions were frothed for 1 minute using a handheld whisk-type frother (Aerolatte, UK). The height of the resulting foam was measured with a ruler immediately after whisking and every 10 mins until collapse of the foam. The foaming ability was determined as the initial height of the foam, while the foam stability was expressed as the time needed for the foam to fully collapse. Measurements were repeated twice.
Emulsifying properties

Both the EAI/ESI test and oil droplet size distribution were carried out to characterise the oil-in-water emulsions prepared with the different samples. Emulsifying activity index (EAI) and emulsion stability index (ESI) were determined according to Ogunwolu et al. [20] with some modifications. 22.5 g solutions of 1% w/w NCM centrate fractions were mixed with 7.5 g of sunflower oil (to obtain a 3:1 phase ratio) and the mixture was high-speed homogenised for 1 min to produce oil-in-water emulsions. 50 µl of the emulsion were pipetted from the bottom of the vial and suspended in 5 ml of 0.1% (w/v) SDS solution. This step was carried out immediately after emulsification then after 10 min. Absorbance of the diluted emulsions was measured at 500 nm using a Genesys 6 UV/Vis spectrophotometer (Thermo Electron Corporation, USA). The ability of the protein to form an emulsion (emulsifying activity index, EAI) and the emulsion stability index (ESI) were calculated using the following formulae:

\[
EAI \left( m^2/g \right) = \frac{2 \times T \times A_0 \times \text{dilution factor}}{C \times \phi \times 1000}
\]

\[
ESI \left( \text{min} \right) = \frac{A_0}{A_0 - A_{10}} \times \Delta t
\]

where \( T = 2.303 \), \( A_0 = \) apparent absorbance at 0 min, dilution factor = 100, \( C = \) weight per unit volume (g/mL), \( \phi = \) oil volumetric fraction (0.25), \( A_{10} = \) apparent absorbance after 10 min, \( \Delta t = 10 \) min. Measurements were repeated three times.

The average oil droplet size distribution of the centrate fraction emulsions (D[3,2] or surface weighted mean) were measured using a Mastersizer 2000 (Malvern Instruments Ltd., UK). The refractive index of oil droplets was set at 1.474 (corresponding to sunflower oil). The laser obscuration was adjusted to 10% obscuration and triplicate measurements were carried out.

Tensiometry

Both surface tension and oil/water interfacial tension were determined for the different samples. For surface tension measurements 1% w/w NCM centrate fraction solutions were serially diluted to obtain 0.5%, 0.25% and 0.1% w/w NCM solutions. The surface tension of the solutions was measured using the Du Nouy method with an EZ-Pi Plus tensiometer (Kibron, UK). 1.5ml of solution was placed into a sterile cup holder. Triplicate measurements were carried out at 20°C.

For oil/water interfacial tension measurements 1% w/w NCM centrate fraction solutions were serially diluted to obtain 0.1%, 0.01%, 0.001% and 0.0001% w/w NCM solutions. A sessile sunflower oil droplet was ejected from a 1.067 mm diameter needle at a rate of 400 µl/min into a glass sample cell containing the centrate fraction solution. The evolution of the oil droplet shape was captured over 60 minutes using a Krüss Easydrop tensiometer (Krüss GmbH, Germany). A 3,000-frame video of the droplet was captured at a frame rate of 50 frames per second. The interfacial tension at the oil/water interface was calculated frame by frame using the associated Drop Shape Analysis software. Measurements were repeated four times. Retraction tests were performed on the aged droplet by removing oil from the droplet at a rate of 100 µl/min with visual observation of the resulting oil/water interface.
Confocal microscopy

Microscopic images of centrate fraction solutions, foams, oil-in-water emulsions and hydrogels were obtained using a Leica TCS2 confocal laser scanning microscope (Leica Microsystems, Germany). Images were recorded at a resolution of 512 x 512 pixels and analysed using the associated DM SDK version 4.2.1 software.

For the imaging of solutions, foams and hydrogels 1% and 10% w/w NCM solutions of the centrate fractions were supplemented with the fluorescent dye Rhodamine B. The dye was excited at 514nm, the collection range was 600-700 nm and a 10x/0.25 dry lens was used. Foams and hydrogels were prepared following the procedures described in the foaming properties and rheological properties sections.

The emulsions were prepared following the procedure described in the emulsifying properties section. Traces of Nile Red dye (Sigma Aldrich Co., UK) were added to the oil prior to mixing with the centrate fraction solution in order to stain the resulting emulsion oil droplets. The dye was excited at 549 nm and fluoresced light was collected at the emission maximum of 628 nm. A 63x/1.40-0.60 oil lens was used.

Statistical analysis

Statistical comparison of the data obtained with the different samples for a given test was carried using the SPSS Statistics 23.0 software (IBM, USA) with a one-way ANOVA test followed by a post hoc Tukey’s HSD (Honestly Significant Difference) test for pair-wise multiple comparison. A p-value of 0.05 was used as cut-off for significance.

Human/animal rights

This article does not contain any studies with human or animal subjects.

Results and discussion

Centrate fraction characterisation

The rationale behind using the whole centrate stream for ultrafiltration instead of its centrifugation supernatant was based on an initial functional comparison between the two samples, during which the whole centrate showed higher foaming and emulsifying properties than its supernatant (results not shown). The pH of all the centrate fractions was more acidic (4.2 to 4.3) than the fermentation broth (6.0) due to the consumption of ammonia added as source of nitrogen for growth of the organism and as pH regulator (Table 1).

Fungal cell and mycelial debris were observed in the centrate and R100+5 solutions (Figures 1a and 1c), with the characteristic mycelial network of the Quorn fermentation broth centrifugation deposit chosen as reference (Figure 1d). Much larger structures were present in R100 (Figure 1b), resulting from the aggregation of the mycelial debris observed in the centrate during their retention on the 100 kDa ultrafiltration membrane. Fungal aggregate formation
has been attributed to a combination of electrostatic interactions, hydrophobic interactions and specific interactions between cell wall components including proteins and polysaccharides [21] and the influence of temperature on mycelial aggregation was previously reported for the filamentous fungus Rhizopus sp [22]. The aggregation process observed could thus have resulted from the pressure exerted on the ultrafiltration membrane-retained material and/or from the temperature increase occurring during ultrafiltration.

The successive ultrafiltration steps resulted in an increase in nitrogen-containing material (NCM): 47.3% (centrate), 56.7% (R100) then 69.8 to 71.6% (further retentates) (Table 1). Following the 100 kDa ultrafiltration step the retention of the aggregated mycelial debris in R100 (observed in Figure 1b) concentrated nitrogen-containing cell wall and cell membrane constituents (including phospholipids, sterols, glycosphingolipids, sphingomyelins, chitin, chitosan and glycoproteins) and contributed to the NCM increase in this sample.

A fraction of the R100 sample did not migrate on the Native-PAGE gel (band g), indicating that material larger than the higher protein marker band size (200 kDa) was present (Figure 2b). Faint bands g (corresponding to non-migrating material) were also reported on the Native-PAGE profiles of the centrate and Quorn fermentation broth centrifugation deposit. Concentrated and faint bands g (non-migrating material) were also respectively observed on the SDS-PAGE profiles of R100 and centrate, however several specific concentrated bands were also present for R100 below 100 kDa in the 10-15 kDa (band b), 20-30 kDa (c and d) and 30-40 kDa (e and f) regions (Figure 2a). These results highlighted the presence of R100 protein aggregates in band g and their breakdown into oligomers or monomers (bands b to f) in the presence of SDS and β-mercaptoethanol. The electrophoretic profiles of R100 also indicated that, in addition to the mycelial aggregates, the concentration of protein aggregates composed of monomers and/or oligomers between 10 and 40 kDa in this fraction also contributed to the increase of NCM level following the 100 kDa ultrafiltration. The Native and SDS-PAGE profiles of the R100+5, R100+50 and R100+50+5 fractions proved similar, with a main band (a) observed below 10 kDa corresponding to protein monomers concentrated as a result of the successive ultrafiltration steps, which contributed to the further increase of NCM concentration in these samples.

**Centrate fraction composition**

173 proteins were successfully identified based on MOWSE scores of 85 or greater (5% probability of misidentification) and identification with more than one peptide, as defined by Pappin et al. [23]. Out of the 173 identified proteins, the presence of 30 ribosomal proteins and 6 heat-shock proteins (Hsp) resulted from the heat-shock RNA-reduction heating step. Appendix A in the supplementary material highlights the proteins reported in higher relative proportions in R100 in comparison with other centrate fractions. These included a cerato-platanin, HEX-1, the heat shock 70kDa protein 4 and a number of elongation factors and tubulins. The higher proportions of a number of proteins in R100 was due to the presence of the large mycelial aggregates observed in Figure 1b. In particular cerato-platanins have been reported within the cell wall of fungal hyphae and spores [10-11] and members of the Hsp70 (heat-shock protein) family have been shown to be located in the cell wall of Saccharomyces cerevisiae [24] and Candida albicans [25].
A large number of proteins below 100 kDa were reported in the R100 fraction, and the majority of these proteins matched in terms of molecular mass with the concentrated bands (a to g) specifically observed on the SDS-PAGE profile of R100 (Figure 2). Moreover several of these proteins have been characterised by their capacity to self-assemble, including cerato-platanin, HEX-1, translation elongation factors and tubulins, confirming the presence of protein aggregates in this fraction which were reduced to oligomers and monomers under the proteomic sample preparation or SDS-PAGE conditions. In particular cerato-platanins from *Trichoderma* are known to form dimers [26]. The cerato-platanin EPL1 from the fungus *Trichoderma atroviride* was reported to form ordered self-assembled layers at the air/water interface [9] and hydrophobic/hydrophilic interfaces [27]. The HEX-1 protein was also shown to self-assemble into three-dimensional lattices to form Woronin bodies, which are peroxisome-derived organelles that maintain cellular integrity and prevent cytoplasmic bleeding in response to hyphal wounding and cellular damage by sealing septal pores [28]. The heat-shock RNA-reduction process carried out on the Quorn fermentation process has been reported to induce such damage to fungal cells [5], resulting in the possible aggregation and concentration of HEX-1 in the large mycelial structures observed in R100.

Appendix B in the supplementary material lists the metabolites which were identified in centrate and previously reported in the literature for their functional properties. The presence of mycelial debris in the centrate and their aggregation and concentration in the R100 fraction (Figure 1b) implied the presence of fungal cell membrane constituents (phospholipids, sterols, glycosphingolipids and sphingomyelins) and cell wall constituents (chitin, chitosan, beta-glucans and glycoproteins). These materials could also have diffused out of the fungal cells through the damaged cell walls as a result of the RNA-reducing heat-shock process as previously reported [5]. Both α-N-acetylglucosamine, monomer unit of chitin and chitosan, and its derivative α-N-acetylglucosamine-6-phosphate were identified in centrate. Chitin, chitosan and their derivatives are used as functional ingredients in the food industry due to their foaming, emulsifying, thickening and gelling properties [13], including those of fungal origin [14]. A range of fungal cell membrane lipids were reported, including phospholipids, lyso phospholipids, sterols and sterol esters, glycosphingolipids and sphingomyelins. Phospholipids (lecithins) and lysophospholipids are used as emulsifiers [29] while phytosterols and phytosterol esters are known to form oleogels [30]. Gelling properties have been reported for several sphingolipid groups, including ceramides and sphingomyelins [31] and glycosphingolipids [32]. Two functional polysaccharides (inulin and a galactan) were identified. Inulin is used for its viscosifying and gelling properties [33] and was previously identified in the fungus *Aspergillus oryzae* KB [34]. Galactans are constituents of cell walls and used as viscosifying and gelling agents [35]. The production of galactans by fungi has previously been reported in the genus *Pleurotus* [36].

A range of nucleobases, nucleosides and nucleotides (including guanine-based compounds) and their derivatives were reported in the centrate metabolome. The presence of these compounds resulted from the breakdown of cellular RNA releasing nucleotides and the subsequent breakdown of these nucleotides into nucleosides and nucleobases as a result of the RNA-reducing heat-shock treatment. Viscosifying and gelling properties have been reported for the guanine nucleobase and for guanine-based nucleosides and nucleotides due to the ability of guanine to self-associate [15]. A number of monoglycerides and diglycerides and their derivatives, used as foaming and emulsifying ingredients [37], were identified as well as several glycosyldiglycerides, characterised by their foaming and emulsifying properties
A range of triterpenoid and steroidal saponins were identified. These compounds have been characterised by their foaming and emulsifying properties and several Fusarium strains were shown to produce saponins, including the triterpenoid saponin-producing Fusarium avenaceum and Fusarium sp. PN8. Several fatty acyl glycosides, including rhamnolipids, were detected. Fungal rhamnolipids are used as foaming and emulsifying agents. Several sugar alcohols, which are characterised by their viscosifying properties, were identified. A range of fatty amides including fatty acyl ethanolamines was detected. Foaming and emulsifying properties have previously been shown for this class of compounds.

**Rheological properties of centrate fractions**

The functional profile of the R100+5 and R100+50+5 fractions proved very similar so not all the results obtained with R100+5 or R100+50+5 are shown. The yield obtained for the R100+50 sample was insufficient to test its functional profile. The final filtrate obtained (filtrate 100+50+5) did not prove functional (results not shown). 10% w/w NCM R100 solutions showed higher viscosities than similar solutions and emulsions prepared with L87 or other centrate fractions (Figure 3a). The high viscosity reported in R100 solutions resulted from the presence of the large mycelial structures observed in Figure 1b which resisted flow and from the presence of viscosifying compounds (including cell wall and membrane constituents such as chitin, chitosan and galactan). Similarly the high concentration of mycelial aggregates was previously correlated with a decrease of the flow index for fermentation broths of the filamentous fungus Aspergillus terreus. 5% w/w NCM R100 emulsions also showed higher viscosities than similar emulsions prepared with L87 or other centrate fractions (Figure 3b) due to the presence of fungal cells, mycelial debris and viscosifying compounds released from the mycelial aggregates during emulsification.

Unheated 10% w/w NCM R100 solutions showed a higher viscoelasticity than those of L87 and other centrate fractions (Figure 4), resulting from the presence of the mycelial aggregates observed in Figures 1b. R100 displayed a gelation profile at 5% w/w NCM, whereas the centrate equivalent did not (results not shown). Both centrate and R100 fractions gelled at 10% w/w NCM while the R100+5 fraction did not gel at any of the NCM concentrations tested. 10% w/w NCM R100 hydrogels showed a very dense network (Figure 5a) of entangled mycelial aggregates and filaments (Figure 5b). The higher viscoelasticity reported for these gels in comparison with L87 and centrate ones (Figure 4) was due to the presence of these structures and of gelling compounds (including cell wall and membrane constituents such as chitin, chitosan, galactan, phospholipids, sterols, glycosphingolipids and sphingomyelins). The entanglement of mycelial aggregates and filaments reported for R100 gels was reminiscent of the microstructure of Quorn products, which is responsible for their meat-like texture and described as a fibre gel composite composed of an entangled mass of mycoprotein hyphae with gelled albumen protein within the interstitial space. The unique gelation of R100 at the lowest concentration tested (5% w/w NCM) was then due to its high concentration of gel-forming mycelial structures in comparison with the centrate which contained only a limited amount of mycelial debris and thus required a higher NCM concentration (10% w/w) to display a gel-like behaviour. In comparison, the gel obtained using L87 (Figure 5c) showed a more open fractal-like gel network resulting in a lower viscoelasticity.
Foaming properties of centrate fractions

When collected following the 100 kDa filtration, the retentate 100 (R100) foamed strongly, which suggested a further concentration of surface-active material present in the centrate. Foams produced with 1% w/w NCM R100 solutions at the original pH (4.2) by gas-sparging (Figure 6) proved more stable than L87 ones, whereas the other centrate fractions did not yield stable foams. Similarly the native R100 extract displayed a higher foam stability than other centrate fractions and L87 using the frothing assay (Figure 7a). However following frothing R100 showed a lower foaming ability than the other samples, in particular the R100+5 extract (Figure 7a). The high foaming ability measured for the centrate was in line with its ability to naturally foam in the centrifuge during processing of the Quorn fermented broth. Heat treatment applied prior to frothing improved the foaming ability of the R100 and centrate fractions but did not change that of the R100+5 sample (Figure 7b). Gas-sparging profiles showed that the foam stability of R100 was maintained when the pH was adjusted to 7 and was greatly enhanced following heating or in the presence of 100mM NaCl (Figure 6). R100 foams outperformed L87 ones in terms of stability in all conditions.

A high density of air bubbles was observed in centrate foams following frothing (Figure 8a) while only few air bubbles were observed in R100 foams (Figure 8b), which was in agreement with the low foaming ability of R100 (Figure 7a). Micrographs showed that air bubbles in R100 foams were trapped in a dense network of mycelial and cell debris (Figure 8c), which limited the number of air bubbles that could be entrapped thus explaining the low foaming ability of R100. Such mechanism was in agreement with previous findings highlighting that highly viscous fermentation media resulted in issues with gas−liquid mass transfer for the filamentous fungus Aspergillus terreus [45]. The high viscosity reported for R100 solutions (Figure 3a) thus only allowed a limited number of air bubbles to be formed and transported within the liquid, leading to a poor foaming ability, but on the other hand prevented the destabilisation of the air bubbles formed by limiting their movement in the liquid, leading to a high foam stability.

Heating R100 solutions prior to foaming resulted in an increased foam stability following gas-sparging (Figure 6), while in parallel an increased foaming ability but decreased foam stability were observed following frothing (Figure 7b). Extrapolating from the micrographs obtained following heat-based gelation of 10% w/w NCM R100 solutions (Figure 5b), heating the 1% w/w NCM R100 solutions should have resulted in an increased viscoelasticity and denser mycelial network. Such structure would have been maintained during the low-shear gas-sparging process, hence resulting in an enhanced entrapment and stabilisation of the air bubbles. On the other hand the high-shear frothing process would have partially disrupted the mycelial network, hence allowing for a higher number of air bubbles to be cavitated and transported but also limiting their stabilisation due to increased movement, thus leading to the increased foaming ability and decreased foam stability measured.

In parallel the centrate, R100 and R100+50+5 solutions showed significantly lower surface tension values than deionised water (72 mN/m) (Figure 9). The surface tension of the R100 and R100+50+5 fraction solutions proved lower than that of L87 at 1% protein. R100 at 0.5%, 0.25% and 0.1% still reduced the surface tension, equivalent to that obtained with a 1% L87 solution. These results highlighted the presence of molecules with surface activity at the air/water interface in the different centrate fractions.
The combined foaming, microscopy and tensiometry results suggested that small surface-active molecules concentrated in the R100+5 fraction were responsible for foaming ability, while specific material in the R100 fraction contributed to foam stability in addition to the physical stabilisation of air bubbles by the dense mycelial network. Foam-positive material specific to R100 included large mycelial aggregates, molecules larger than 100 kDa, foaming molecules and/or mycelial fragments or molecules released from mycelial aggregates during gas-sparging or frothing (including cell wall and membrane constituents exhibiting foaming properties such as chitin, chitosan and proteins). In particular the higher stability of R100 foams could be due to the concentration of the cerato-platanin in this fraction. Frischmann et al. [9] reported that solutions of cerato-platanin EPL1 from *Trichoderma atroviride* exhibited strong foaming properties. In addition, depending on their hydrophobicity the possible concentration of *Fusarium venenatum* cells and/or spores in the R100 fraction during ultrafiltration could also have contributed to the stabilisation of R100 foams as the shapes and sizes of bacterial cells, viruses and spores fall within the range suitable for Pickering stabilisation of biphasic dispersions including foams and emulsions [47]. Finally, a serpin (SERPB1) was identified in all centrate fractions. Steenbakkers et al. [48] previously reported a fungal serpin (celpin) from *Piromyces* spp. strain E2. Serpins Z4 and Z7 from barley have been characterised as foam-positive proteins in beer [49-50].

The contribution of large mycelial aggregates was further supported by the higher foam stability obtained for the whole centrate (6.49 mins, Figure 6) in comparison with the centrate centrifugation supernatant (3.40 mins, results not shown) in which they were absent. In addition the contribution of mycelial debris released from the breakdown of the large aggregates during frothing was further supported by the higher foam collapse time obtained for the whole centrate (180 mins, Figure 7) in comparison with the centrate centrifugation supernatant (50 mins, results not shown) in which large aggregates were absent.

**Emulsifying properties of centrate fractions**

The centrate and R100 emulsions showed a comparable emulsifying activity index (EAI) to L87 at pH 4.2 and 7 (Figure 10a). The EAI of R100 was greatly increased following prior heating and proved higher than that of L87. R100 emulsions showed a higher emulsifying stability index (ESI) than those of the other centrate fractions and L87 in all the conditions tested (Figure 10b). The stability of R100 emulsions was greatly enhanced by either heating or adjusting the pH to 7. The whole centrate displayed higher EAI and ESI values (13.17 m²/g and 19.72 mins, Figure 10) than the centrate centrifugation supernatant (5.89 m²/g and 17.62 mins, results not shown), highlighting the contribution of large mycelial aggregates to emulsifying properties.

Emulsions prepared with R100 at its original pH (4.2) displayed a smaller mean oil droplet size (D[3,2] surface weighted mean value of 8.8 µm) than L87 (16.1 µm at pH 7) and the other centrate fractions (10.4 µm for centrate and 16.5 µm for R100+5 at pH 4.2) (Figure 11), which was in agreement with their higher EAI and ESI. Smaller oil droplet sizes were also observed for R100 emulsions in comparison with the other samples by confocal microscopy (Figure 12). Small mycelial and cell debris were present in R100 emulsions (Figure 12c), resulting from the breakdown of the large structures observed in R100 solutions (Figure 1b) during emulsification. Similarly to the stabilisation of R100 foams, the presence of mycelium fragments separated and limited the movement of oil droplets, contributing to
emulsion stabilisation. The released mycelial fragments and the remaining mycelial aggregates also resulted in the enhanced viscosity reported for R100 emulsions (Figure 3b), which further limited the movement of oil droplets and stabilised them.

Heating the R100 solutions prior to emulsification resulted in increased EAI and ESI (Figure 10). Extrapolating from the micrographs obtained following heat-based gelation of 10% w/w NCM R100 solutions (Figure 5b), heating the 1% w/w NCM R100 solutions should have resulted in a denser mycelial network. Such structure would have been partially disrupted during the high-shear emulsification process, releasing a number of mycelial fragments contributing to oil droplet formation and stabilisation, while the remaining denser network further stabilised the emulsion, thus leading to increased emulsifying ability and stability.

In parallel oil/water interfacial tension measurements were carried out. A wide range of dilutions (1% to 0.0001% w/w NCM) were screened. Based on the interfacial tension profiles obtained a 0.01% concentration was deemed optimal for comparison between samples. The interfacial tension of a sunflower oil droplet in water was used as control and measured at 32 mN/m. In the presence of 0.1% R100 or R100+5 the oil/water interfacial tension respectively decreased to 15 and 19 mN/m (Figure 13). Following a 1-hour ageing time the interfacial tension dropped to 11 and 14 mN/m respectively. Although these decreases were not as pronounced as with 0.01% L87, these results confirmed the presence of surface-active moieties in the centrate fractions. Following a one-hour aging process the drop surface crumpled or broke upon oil retraction (Figure 14 and Table 2), indicating that the surface was covered with a rigid viscoelastic film that resisted deformation as the drop surface area was reduced. Oil droplets exhibited the strongest crumpling behaviour in R100 solutions and at an R100 concentration as low as 0.0001%. As the mycelial aggregates observed in Figure 1b settled at the base of the cell over time and did not diffuse to the interface, these results confirmed the presence of interfacially-active molecules in the centrate fractions and the concentration and/or specificity of some of this material in R100.

The combined EAI/ESI, oil droplet size, microscopy and tensiometry results highlighted the contribution of specific surface-active material (at the oil/water interface) and of viscosifying mycelium fragments (in the bulk emulsion) to the higher emulsifying properties measured for R100. Emulsion-positive material included the released mycelial debris, molecular aggregates larger than 100kDa, emulsifying molecules and/or molecules released from the mycelial structures during emulsification (including cell wall and membrane constituents exhibiting emulsifying properties such as phospholipids, sterols, glycosphingolipids, sphingomyelins, chitin and chitosan). In addition the possible concentration of *Fusarium venenatum* cells and/or spores in R100 following ultrafiltration could have also contributed to the stabilisation of R100 emulsions. Dorobantu et al. [51] demonstrated that cells from four hydrocarbon-degrading bacteria species formed and stabilised oil-in-water and water-in-oil Pickering emulsions depending on their hydrophobicity. A partially hydrophobic moss spore (*Lycopodium clavatum*) also displayed the ability to stabilise oil-in-water emulsions [52-53].

**Conclusion**
The proteomic and metabolomic profiling of the Quorn fermentation process co-product (centrate) coupled with the functional screening of a range of centrate-derived fractions confirmed the presence of functional material in this co-product. In particular the retentate 100 (R100) fraction, obtained following a 100 kDa ultrafiltration, displayed good foaming, emulsifying and rheological properties, highlighting the potential of functional extracts from the Quorn fermentation process as novel ingredients for the preparation of sustainable food products.

The study described the complex and specific nature of the R100’s functional profile, with contributions reported for both mycelial structures and surface-active molecules. Large mycelial aggregates were observed in R100 solutions, contributing to their high viscoelasticity and high foam stability, and formed a fibre gel composite-type structure upon heating, resulting in highly viscoelastic R100 gels. In parallel tensiometry measurements showed the presence of surface-active moieties at the air/water and oil/water interfaces, highlighting the contribution of functional molecules (including cell membrane and cell wall constituents of the mycelial aggregates) and/or cell debris released from mycelial structures during foaming or emulsification to the high foaming and emulsifying properties of R100. Additional studies are required to further understand the contribution of mycelial structures and surface-active molecules to these extracts. Future work will assess the possible applications of functional centrate extracts as novel sustainable ingredients for the food industry. Finally this study highlights the possibility to investigate the functional profile of the other Quorn fermentation process streams (fermentation broth and RNA-reduced broth).

Acknowledgements
The authors wish to thank Dr Tim Finnigan, Dr Muyiwa Akintoye and Dr Sue Gordon from Marlow Foods for their support during this project.

Funding
This work was supported by the Engineering and Physical Sciences Research Council [grant number EP/J501682/1 Foaming and Fat Replacer Ingredients].

Compliance with ethical standards

Conflict of interest The authors declare that they have no conflict of interest.

Compliance with ethics requirements This article does not contain any studies with human or animal subjects.

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Table 1 Characterisation of centrate fractions

| Fraction | Molecule range | Nitrogen-containing material (%) (n=3) | pH in aqueous solution |
|----------|----------------|--------------------------------------|------------------------|
| Centrate | n/a            | 47.3 ± 1.2                           | 4.2                    |
| R100     | > 100 kDa      | 56.7 ± 4.7                           | 4.2                    |
| R100+5   | 5 > kDa < 100  | 69.8 ± 0.3                           | 4.3                    |
| R100+50  | 50 > kDa < 100 | 69.2 ± 1.0                           | 4.3                    |
| R100+50+5| 5 > kDa < 50   | 71.6 ± 2.0                           | 4.3                    |

Table 2 Oil droplet shape profiles upon retraction according to NCM concentration

|                  | 0.0001% | 0.001% | 0.01% | 0.1%   |
|------------------|---------|--------|-------|--------|
| centrate         | no crumpling | crumpling * | crumpling ** | crumpling ** |
| R100             | crumpling * | crumpling ** | crumpling ** | n/a (too turbid) |
| R100+5           | no crumpling | breaking | crumpling ** | crumpling ** |

* major crumpling observed, * minor crumpling observed
Fig. 1 Confocal microscopy of centrate fractions (1% w/w NCM concentration, original pH, Rhodamine, magnification x10 [a,b,c] or x63 [d])
(a) Centrate, (b) R100, (c) R100+5, (d) Quorn fermentation broth centrifugation deposit
Fig. 2 SDS-PAGE (Fig. a) and Native-PAGE (Fig. b) of centrate and broth fractions (1% w/w NCM, original pH)

Fig. a: centrate [lanes 1-2], R100 [3-4], R100+50 [5-6], R100+50+5 [7-8], Quorn fermentation broth centrifugation deposit [9], See Blue reference marker [10]

Fig. b: See Blue marker [1], Centrate [2-3], R100 [4-5], R100+50+5 [6-7], Quorn fermentation broth centrifugation deposit [8-9], See Blue reference marker [10]
Fig. 3 Viscosity profiles of centrate fraction solutions (a) and centrate fraction emulsions (b) during shear sweep

Fig. a: 10% w/w NCM concentration except 15.3% NCM w/w for L87, original pH, n=2

Fig. b: 5% w/w NCM concentration, original pH, n=3
Fig. 4 Gelation profiles (elastic modulus $G'$ and viscous modulus $G''$) of centrate fractions (10% w/w NCM concentration, original pH, $n=3$)

Fig. 5 Confocal microscopy of centrate fraction hydrogels (10% w/w NCM concentration, original pH, Rhodamine, magnification x10 [a,c] and x40 [b])
(a) R100, (b) R100, (c) L87
Fig. 6 Foam stability of the centrate (a, b) and broth (c) fractions as determined by the Rudin method (1% w/w NCM concentration, n=4).

Different letters indicate statistically significant differences (p<0.05)
Fig. 7 Foaming ability and stability profiles of native (a) and heated (b) centrate fractions (1% w/w NCM concentration, frothing assay, original pH, 80°C for 30 mins, n=3)
Fig. 8 Confocal microscopy of centrate fraction interfaces prepared by frothing (1% w/w NCM concentration, original pH, Rhodamine, magnification x10)
(a) air bubbles in centrate foam, (b) air bubbles in R100 foam, (c) mycelial structure in R100 foam

Fig. 9 Surface tension of centrate fractions (0.1 to 1% w/w NCM concentration, n=3)
Fig. 10 Emulsifying ability (a) and stability (b) of the centrate and broth fractions as determined by the EAI/ESI method (1% w/w NCM concentration, n=4)

Different letters indicate statistically significant differences (p<0.05)
Fig. 11: Oil droplet size distribution of emulsions stabilised with centrate and broth fractions (1% w/w NCM concentration, original pH, n=3)

Fig. 12 Confocal microscopy of centrate, broth and whey protein emulsions (1% w/w NCM concentration, original pH, Nile Red, magnification x 63)
(a) Centrate, (b-c) R100, (d) R100+5, (e) L87
Fig. 13: Interfacial tension decrease at the oil/water interface in presence of dispersed centrate and broth fractions (oil droplet in 0.01% w/w NCM aqueous solution, original pH, n=4)

Fig. 14: Evolution of oil droplet shape during retraction after 1 hour ageing in a 0.01% NCM R100 aqueous solution (original pH, n=4)
(a) Full extension, (b1) crumpling during retraction after 1 hour-ageing, (b2) further crumpling with further retraction, (c) droplet breaking during retraction after 1 hour-ageing
### Appendix A: Proteomic markers of the R100 fraction

| Molecular mass (kDa) | Code (Gibberella zeae PH-1) | Identification (Gibberella zeae PH-1) | centrate | R100 | R100+5 | SDS-PAGE band |
|---------------------|-----------------------------|-----------------------------------------|----------|------|--------|---------------|
| 10                  | (translation elongation factor 1-alpha 10kDa monomer?) (Fusarium venenatum) | (18) | (7) | (11) | (a) |
| 15                  | RS12_ERYGR                   | 40S ribosomal protein S12              | 183      | 103  | 421    | a             |
| 17                  | RS15_PODAN                   | 40S ribosomal protein S15 (S12)        | 198      | 76   | 296    | b             |
| 20 (24 in GZ)       | RS5_CICAR                    | 40S ribosomal protein S5               | 134      | 93   | 330    | c             |
| 21                  | FG09866.1                    | 60S ribosomal protein L1B              | 196      | 60   | n/a    | c             |
| 24                  | FG06924.1                    | large subunit ribosomal protein L10Ae | 90       | 53   | n/a    | c             |
| 24                  | FG04471.1                    | cerato-platanin protein                | 56       | 8    | 54     | d             |
| 25                  | FG08737.1                    | HEX-1                                   | 142      | 74   | 360    | d             |
| 25                  | FG01008.1                    | elongation factor 1-beta               | 408      | 73   | 85     | d             |
| 28                  | FG01154.1                    | 60S ribosomal protein L4-B (L9)        | 112      | 79   | 240    | d             |
| 28                  | FG01509.1                    | 40S ribosomal protein S2               | 54       | 43   | 125    | d             |
| 28                  | FG01241.1                    | similar to 14-3-3 protein epsilon      | 117      | 45   | n/a    | d             |
| 29                  | FG06841.1                    | Uncharacterized                        | n/a      | 83   | n/a    | d             |
| 29                  | FG07480.1                    | 40S ribosomal protein S1               | 47       | 33   | 127    | d             |
| 32                  | FG10905.1                    | 40S ribosomal protein S0               | 125      | 91   | 327    | d             |
| 32                  | FG10235.1                    | uncharacterized                        | 510      | 81   | n/a    | d             |
| 33                  | FG00496.1                    | inorganic pyrophophatase               | 104      | 56   | 166    | d             |
| 33                  | FG00602.1                    | uncharacterised                        | 108      | 68   | n/a    | d             |
| 33 (28 in GZ)       | SPEE_NEUCR                   | spermidine synthase (putrescine aminopropyltransferase) (SPDSY) | 208      | 82   | n/a    | d             |
| 33                  | FG05035.1                    | pyridoxine biosynthesis protein PDX1   | 102      | 75   | 83     | d             |
| 34                  | FG01870.1                    | rRNA 2’-O-methyltransferase fibrillarin | 230      | 86   | n/a    | e             |
| 34                  | ADT_NEUCR                    | ADP-ATP carrier protein (ADP/ATP translocase) (adenine nucleotide translocator) (ANT) | 48       | 40   | 343    | e             |
| 35                  | RLS_NEUCR                    | 60S ribosomal protein L5 (CPR4)        | 95       | 84   | 273    | e             |
| (35)                | GBLP_NEUCR                   | guanine nucleotide-binding protein beta subunit-like protein (cross-pathway control WD-repeat protein cpc-2) | 163      | 28   | 126    | e             |
| 36                  | FG05615.1                    | adenosylhomocysteinase                 | 63       | 63   | n/a    | e             |
| 36                  | FG06611.1                    | tubulin beta chain (Fusarium venenatum) | n/a      | 80   | n/a    | e             |
| 36                  | FG02014.1                    | heat shock protein 90                  | 74       | 72   | 149    | e             |
| 38                  | FG07186.1                    | 60S ribosomal protein L4-A             | 27       | 13   | 76     | e             |
| (39)                | NMT1_ASPPA                   | NMT1 protein homolog (thiamine biosynthesis) | 34       | 17   | 173    | f             |
| 40                  | FG02770.1                    | fructose-bisphosphate aldolase         | 36       | 6    | 50     | f             |
| 41                  | FG06675.1                    | 3-isopropylmalate dehydrogenase        | 211      | 26   | 257    | f             |
| 41                  | FG09580.1                    | isocitrate dehydrogenase [NAD] subunit, mitochondrial | 261      | 85   | n/a    | f             |
| (42)                |                               | gamma-actin                            | 40       | 38   | 42     | f             |
| 43                  | FG06668.1                    | (elicitor protein) = SERBP1 (serpine) = plasminogen activator inhibitor 1 RNA-binding protein | 77       | 51   | 52     | f             |
|   |   |   |   |   |   |
|---|---|---|---|---|---|
| 43 | RL3_NEUCR | 60S ribosomal protein L3 | 62 | 46 | n/a | f |
| 43 | FG03816.1 | similar to lactonohydrolase | n/a | 57 | n/a | f |
| 43 | METK_NEUCR | S-adenosylmethionine synthetase = Methionine adenosyltransferase = AdoMet synthetase | n/a | 31 | n/a | f |
| 45 | FG05454.1 | pyruvate dehydrogenase E1 component subunit alpha | 191 | 87 | 336 | f |
| 45 | ILV5_NEUCR | ketol-acid reductoisomerase, mitochondrial precursor (Acetohydroxy-acid reductoisomerase) (Alpha-keto-beta-hydroxylacil reductoisomerase) | 81 | 23 | n/a | f |
| 47 | ENO_ALTAL | enolase = phosphopyruvate hydratase | 2 | 2 | 2 | f |
| 49 | FG07174.1 | acetyltransferase component of pyruvate dehydrogenase complex | 171 | 96 | n/a | f |
| 49 | FG00346.1 | saccharopine dehydrogenase | 89 | 44 | 200 | f |
| 50 | EF1A_TRIRE | elongation factor 1 alpha | 18 | 7 | 11 | f |
| 52 | FG07468.1 | phosphoglycerate dehydrogenase | 23 | 21 | 145 | |
| 53 | FG06039.1 | ATP citrate lyase | 24 | 22 | n/a | |
| 57 | FG08712.1 | Fumarate hydratase | 289 | 94 | 382 | |
| 58 | FG01111.1 | 6-phosphogluconate dehydrogenase | 37 | 12 | n/a | |
| 60 | FG00387.1 | Phosphoglucomutase | 296 | 66 | n/a | |
| 60 | KPYK_TRIRE | Pyruvate kinase | 84 | 52 | n/a | |
| 60 | FG09940.1 | Inositol-3-phosphate synthase isozyme 2 | 70 | 42 | 174 | |
| 63 | FG09834.1 | pyruvate decarboxylase | 5 | 4 | 151 | |
| 64 | FG11326.1 | Glucoamylase | 94 | 47 | 100 | |
| 65 | FG00969.1 | Bifunctional purine biosynthesis protein ADE17 | 148 | 35 | 192 | |
| 66 | FG03993.1 | Homocitrate synthase | 135 | 88 | n/a | |
| 69 | FG10908.1 | Nucleolar protein NOP5 | 263 | 92 | 520 | |
| 69 | FG08875.1 | Sulfate adenylyltransferase | n/a | 20 | 325 | |
| 73 | FG01199.1 | ATP-citrate synthase subunit 1 (Citrate cleavage enzyme subunit 1) | n/a | 18 | n/a | |
| 77 | FG01199.1 | Glucosamine-fructose-6-phosphate aminotransferase | 46 | 30 | n/a | |
| 86 | FG07953.1 | aconitate hydratase | 69 | 16 | n/a | |
| 86 | FG01950.1 | heat shock 70kDa protein 4 | 115 | 27 | 194 | |
| 91 | FG05530.1 | Cell division cycle protein 48 | n/a | 65 | n/a | |
| 117 | FG04181.1 | elongation factor 3 | 29 | 14 | 135 | |
| 198 | FG03631.1 | succinate dehydrogenase (ubiquinone) flavoprotein subunit | 98 | 41 | n/a | |
| 205 | FG05321.1 | Fatty acid synthase subunit alpha reductase | 57 | 25 | 61 | g |
| 234 | FG05322.1 | Fatty acid synthase subunit beta dehydratase | 55 | 34 | 497 | g |
Appendix B: Metabolites identified in centrate and previously reported for their functional properties

| Group                          | Subgroup                                      | Potential functionality | Compound               |
|-------------------------------|-----------------------------------------------|-------------------------|------------------------|
| Fatty acid amides             | Fatty acid primary amides                     | Foam, emulsion          | 12 (pos) / 13 (neg)    |
|                               | Fatty acid ethanolamines                      | Foam, emulsion          | 3 (pos) / 1 (neg)      |
|                               | Other fatty acid amides                       | Foam, emulsion          | 3 (pos) / 3 (neg)      |
| Glycerides                    | Monoglycerides (MG)                           | Foam, emulsion          | 7 (pos) / 16 (neg)     |
|                               | Diglycerides (DG)                             | Foam, emulsion          | 9 (pos) / 20 (neg)     |
|                               | Triglycerides (TG)                            | Viscosity               | 3 (pos) / 22 (neg)     |
| Glycosyldiglycerides          | Monogalactosyldiglycerides (MGDG)             | Foam, emulsion          | 7 (pos) / 1 (neg)      |
|                               | Other glycosyldiglycerides                    | Foam, emulsion          | 0 (pos) / 2 (neg)      |
| Phospholipids                 | Phosphoglycerides (PG)                        |                         | 15 PG (pos) / 12 PG (neg) |
|                               | Phosphatidic acids (PA)                       |                         | 4 PA (pos) / 7 PA (neg) |
|                               | Phosphatidylethanolamines (PE)                | Emulsion                | 11 PE (pos) / 5 PE (neg) |
|                               | Phosphatidylinositolis (PI)                   | Emulsion                | 21 PI (pos) / 28 PI (neg) |
|                               | Phosphatidylserines (PS)                     | Emulsion                | 11 PS (pos) / 9 PS (neg) |
| Lysosphospholipids            | LysoPG (LPG)                                  |                         | 8 LPG (pos) / 8 LPG (neg) |
|                               | LysoPA (LPA)                                  |                         | 9 LPA (pos) / 9 LPA (neg) |
|                               | LysoPE (LPE)                                  | Emulsion                | 9 LPE (pos) / 7 LPE (neg) |
|                               | LysoPI (LPi)                                  | Emulsion                | 6 LPI (pos) / 4 LPI (neg) |
|                               | LysoPS (LPS)                                  | Emulsion                | 1 LPS (pos) / 4 LPS (neg) |
| Sphingolipids                 | Ceramides (Cer) and ceramide phosphates (CerP)| Emulsion, gel           | 4 Cer (pos) / 1 Cer (neg) |
|                               | Sphingomyelins                                | Emulsion, gel           | 2 CerP (pos)           |
|                               | Cerebrosides (non sulphated) and derivatives  | Emulsion, gel           | 5 PE-Cer (pos) / 2 PE-Cer (neg) |
|                               | Sulfatides                                    | Emulsion, gel           | 1 PI-Cer (pos) / 1 PI-Cer (neg) |
|                               | Other glycosphingolipids                      | Emulsion, gel           | 6 (pos) / 7 (neg)      |
| Steroids                      | Sterols and derivatives (episteryl oleate)   | Oleogel                 | 9 (pos) / 1 (neg)      |
|                               | Sterol esters                                 | Oleogel                 | 5 (pos) / 1 (neg)      |
| Sugar alcohols                |                                               | Viscosity               | 7 (pos) / 4 (neg)      |
| Class                        | Subclass                        | Function              | (pos) / (neg)         |
|------------------------------|---------------------------------|-----------------------|-----------------------|
| Saponins                     | Triterpenoid saponins           | Foam, emulsion        | 7 (pos) / 7 (neg)     |
| Steroidal saponins           | Foam, emulsion                  | 8 (pos) / 1 (neg)     |
| Other steroidal glycosides   | Foam, emulsion                  | 0 (pos) / 9 (neg)     |
| Fatty acyl glycosides        | Rhamnolipids                    | Foam, emulsion        | 3 (pos) / 1 (neg)     |
| Other fatty acid glycosides  | Foam, emulsion                  | 8 (pos) / 13 (neg)    |
| Polysaccharides              | Chitin (monomer)                | Foam, emulsion, viscosity, gel | a-N-Acetylglucosamine (pos) |
| Chitin derivative            | Foam, emulsion, viscosity, gel  | N-Acetylglucosamine 6-phosphate (neg) |
| Inulin, galactan             | Viscosity, gel                  | 2 (pos) / 1 (neg)     |
| Nucleobases                  | Nucleobases                     | Viscosity, gel        | Guanine (pos/neg)     |
| Derivatives                  | Viscosity, gel                  | 5 (pos) / 8 (neg)     |
| Nucleosides                  | Ribonucleosides                 | Viscosity, gel        | Isoguanosine (pos)    |
| Deoxyribonucleoside          | Viscosity, gel                  | Deoxyguanosine (pos)  |
| Ribonucleoside derivatives   | Viscosity, gel                  | 6 (pos) / 1 (neg)     |
| Deoxyribonucleoside derivatives | Viscosity, gel              | 2 (pos) / 0 (neg)     |
| Ribonucleoside monophosphates | Viscosity, gel                  | 3 (pos) / 1 (neg)     |
| Ribonucleoside monophosphate derivatives | Viscosity, gel | 2 (pos) / 5 (neg)     |
| Ribonucleoside diphosphates  | Viscosity, gel                  | 3 (pos) / 3 (neg)     |
| Ribonucleoside diphosphates glycosides | Viscosity, gel | 1 (pos) / 6 (neg)     |
| Ribonucleoside diphosphate glycosyl aminoacids | Viscosity, gel | 0 (pos) / 1 (neg)     |
| Ribonucleoside diphosphate diglycerides | Viscosity, gel, foam, emulsion | 3 (pos) / 2 (neg)     |
| Ribonucleoside diphosphate fatty acyl glycosides | Viscosity, gel, foam, emulsion | 0 (pos) / 3 (neg)     |
| Other ribonucleoside diphosphate derivatives | Viscosity, gel | 0 (pos) / 2 (neg)     |
| Deoxyribonucleoside diphosphate derivatives | Viscosity, gel | 1 (pos) / 2 (neg)     |

(pos): number of compounds reported in positive ionisation mode, (neg): number of compounds reported in negative ionisation mode
Appendix B Metabolites identified in centrate and previously reported for their functional properties
(pos): number of compounds reported in positive ionisation mode, (neg): number of compounds reported in negative ionisation mode

Graphic abstract