ABSTRACT

Oxidative stress caused by free radicals has been implicated in several human disorders. Dietary antioxidants can help the body to counteract those reactive species and reduce oxidative stress. Antioxidant activity is one of the multiple health-promoting attributes assigned to bovine whey products. The present study investigated whether this activity was retained during upper gut transit using a static simulated in vitro gastrointestinal digestion (SGID) model. The capacity to scavenge free radicals and reduce ferric ion of whey protein isolate (WPI), individual whey proteins, and hydrolysates pre- and post-SGID were measured and compared using various antioxidant assays. In addition, the free AA released from individual protein fractions in physiological gut conditions were characterized. Our results indicated that the antioxidant activity of WPI after exposure to the harsh conditions of the upper gut significantly increased compared with intact WPI. From an antioxidant bioactivity viewpoint, this exposure negates the need for prior hydrolysis of WPI. The whey protein α-lactalbumin showed the highest antioxidant properties post-SGID (oxygen radical absorbance capacity = 1,825.94 ± 50.21 μmol of Trolox equivalents/g of powder) of the 4 major whey proteins tested with the release of the highest amount of the antioxidant AA tryptophan, 6.955 μmol of tryptophan/g of protein. Therefore, α-lactalbumin should be the preferred whey protein in food formulations to boost antioxidant defenses.

Key words: whey protein hydrolysis, antioxidant activity, simulated gastrointestinal digestion, α-lactalbumin

INTRODUCTION

Antioxidants are widely used in the food industry to prevent food oxidation and provide health benefits to the consumer by combating oxidative stress (Capitani et al., 2009). Oxidative stress occurs due to an imbalance between the elimination and production of reactive oxygen, nitrogen, and sulfur species (Li et al., 2015). Dietary antioxidants, both soluble and insoluble, can reduce oxidative stress through different pathways, including scavenging free radicals by electron or hydrogen atom transfer (Lü et al., 2010). Antioxidants from plants (polyphenols, carotenoids, and vitamins) are widely accepted as natural antioxidants (Fiedor and Burda, 2014). Once ingested, these bioactive compounds or their metabolites survive gut transit and are transported across the intestinal barrier to reach their target organs (Sarmadi and Ismail, 2010; Quirós-Sauceda et al., 2017). Recently, animal-derived proteins from eggs, meat, and milk have been described as a source of antioxidants (Cloetens et al., 2013; Nimalaratne and Wu, 2015; Liu et al., 2016). In particular, products derived from bovine whey, such as whey protein isolate (WPI) and whey protein concentrate (WPC), have been extensively tested for antioxidant potential in vitro (O’Keeffe and FitzGerald, 2014; Önay-Ucar et al., 2014; Torkova et al., 2016). Indeed, recent human intervention studies with whey product supplementation have reported increased levels of different antioxidant biomarkers in plasma, such as glutathione (de Aguilar-Nascimento et al., 2011; Power-Grant et al., 2016).

Dried whey protein-based ingredients differ primarily in the percentage of protein, fat, and lactose, with WPI containing the highest protein concentration at 90 to 95%. The protein component is considered to harbor the antioxidant activity and is composed of β-LG (50–60%), α-LA (15–25%), BSA (6%), lactoferrin (LF; <3%), and immunoglobulins (<1%; Madureira et al., 2007). In an effort to increase the antioxidant potency of commercial whey, different processing methods [enzymatic hydrolysis (Zhidong et al., 2013), fractionation (Hernández-Ledesma et al., 2005), thermal treatment
(Adjonu et al., 2013), pressure treatment (Iskandar et al., 2015), acid treatment (Mohammadian and Madadalou, 2016), and polymerization (Ortega et al., 2015) have been assessed. It is generally agreed that enzymatic hydrolysis and fractionation increase the antioxidant properties of whey (Power et al., 2013). Hydrolysis with the enzyme alcalase (EC 3.4.21.62) commonly delivers the most potent antioxidant fractions (Lin et al., 2012; Zhang et al., 2013). In addition, whey fractions with a molecular weight below 5 kDa consistently exert better antioxidant properties than higher-molecular weight fractions (de Castro and Sato, 2014; Liu et al., 2014; O’Keeffe and FitzGerald, 2014). Recently, Bamdad et al. (2017) identified 8 peptides present in the most antioxidant fraction of β-LG hydrolyzed by alcalase. Once ingested in vivo, whey products are subjected to the chemical (acidic), physical (peristaltic), and enzymatic degradation process of the gastrointestinal tract (Guerra et al., 2012). Given the hydrolytic conditions prevailing in the gastrointestinal tract, it could be argued that enzymatic hydrolysis during whey processing is an unnecessary step to provide the body with antioxidant whey. Indeed, digestion of foods that contain hydrolyzed whey proteins may result in the premature loss of antioxidant activity during gut transit, as hydrolyzed proteins are more easily broken down to individual AA compared with intact protein complexes (Koopman et al., 2009).

Therefore, the objectives of our study were to investigate, under simulated in vitro gastrointestinal digestion (SGID) conditions, whether commercial WPI exposed to the harsh conditions of the upper gut had similar antioxidant activity to WPI hydrolysates prepared with commercial enzymes. We hypothesized that transit of WPI through the upper gut would increase its antioxidant bioactivity to levels comparable to WPI hydrolysates prepared with commercial enzymes. Materials and Methods

Materials

Commercial bovine WPI (Isolac, 91.4% protein content) was purchased from Carbery Food Ingredients (Ballineen, Co. Cork, Ireland). Individual whey proteins were sourced from food industries as representative of commercial products, except BSA (98% protein content) which was purchased from Sigma-Aldrich (Dublin, Ireland). The proteins β-LG (92.1% β-LG content) and α-LA (93% α-LA content) were obtained from Davisco Foods International, Inc. (Le Sueur, MN). The LF (Bioferrin 2000, 95% LF content) was donated by Glanbia Nutritionals, Inc. (Fitchburg, WI). Bromelain (EC 3.4.22.32) was from Kerry Foods Ingredients (Kilnagleyre, Carrigaline, Co. Cork, Ireland), whereas alcalase 2.4 U and neutrase 0.8 L (EC 3.4.24.28) were from Novozymes (Bagsvaerd, Denmark). Reagents were purchased from Sigma-Aldrich unless stated otherwise.

Enzymatic Hydrolysis of WPI

Protein powder (WPI 10%, wt/vol) was hydrated in Milli-Q H$_2$O (300 mL) at 4°C stirring overnight. The protein solution was warmed at 50°C for 15 min before starting the hydrolysis. Hydrolysates were produced using the commercial enzymes alcalase (2.4 Anson units/g), bromelain (600 gelatin digestion units/g), or neutrase (0.8 Anson units/g) at an enzyme-to-substrate ratio of 1:100 (g/g) for 180 min in a water bath at 50°C with continuous stirring. The pH was maintained constant at 8.0 for alcalase or 7.0 for bromelain and neutrase with 2 M solution of NaOH using a Metrohm 842 Titrand dosing unit (Metrohm Ltd., Herisau, Switzerland). After 180 min, the hydrolysis was stopped by inactivating the enzymes in a water bath at 90°C for 10 min. Hydrolysates were freeze-dried and stored at −80°C for further use. Degree of hydrolysis (%DH) of these products was determined according to the following equation (Adler-Nissen, 1986):

\[ \%DH = \frac{B \cdot N_B}{M \cdot h_{TOT} \cdot \alpha} \times 100, \]

where $B$ (mL) is the amount of alkaline solution employed, $N_B$ (mol/L) is the molarity of the NaOH aqueous solution used, $M$ (g) is the protein mass reacting, $h_{TOT}$ (mEq/g) is the number of total peptides bonds contained in the protein, and $\alpha$ is the dissociation degree of α-amino groups that are released during protein hydrolysis. The values $h_{TOT}$ (8.8 mEq/g) and $\alpha$, at pH 7 (0.441) and 8 (0.885), were obtained from Adler-Nissen (1986).

Simulated Gastrointestinal Digestion Static Model

An in vitro SGID was performed according to an established INFOGEST method (Figure 1; Minekus et al., 2014). Oral phase was not performed, as 1 g of protein powder was reconstituted in 5 mL of Milli-Q H$_2$O. Gastric phase was started by mixing 1 g of reconstituted protein powder with 2 mL of simulated gastric
We then added 2.5 mL of a fresh stock solution (8,000 U/mL) of porcine pepsin (EC 3.4.23.1) to reach 2,000 U/mL activity; pH was adjusted to 3.0 manually using HCl (1 M) and volume brought up to 10 mL with Milli-Q H2O. The mixture was incubated for 2 h at 37°C with continuous shaking. Pepsin was inactivated by increasing pH to 6.5 manually using NaOH (1 M) and by diluting 5 parts of gastric chyme with 4 parts of simulated intestinal fluid. After addition of pancreatin, bile extract, and water, the final ratio of gastric chyme to simulated intestinal fluid was 50:50 (vol/vol). Pancreatin (EC 232.468.9) and bile extract prepared in simulated intestinal fluid were added to achieve a final concentration of 200 U/mL and 10 mM, respectively. The pH was adjusted to 7.0, the volume brought up to 20 mL, and the mixture was again incubated for 2 h at 37°C with continuous shaking. After this time, the digestion was stopped by adding the protease inhibitor 4-(2-aminoethyl) benzenesulfonyl fluoride hydrochloride at a final concentration of 1 mM. The final sample concentration after SGID was 50 mg of protein powder/mL. Samples were aliquoted, snap-frozen in liquid nitrogen, and stored at −80°C before analysis. On experimental days, samples were defrosted on ice and diluted, according to the final powder concentration after SGID, to the corresponding assay concentration. The SGID control contained pepsin, pancreatin, and bile salts and it was incubated for the same time as whey test samples. Protease inhibitor was then added and snap frozen.

**Determination of the DH**

The extent of hydrolysis was determined using the 2,4,6-trinitrobenzene 1-sulfonic acid (TNBS) method developed by Fernández and Kelly (2016) based on Adler-Nissen (1979). Samples were prepared at 50 mg/mL and diluted 1:10 in Milli-Q H2O prior to assay. Diluted samples (10 μL) were mixed with 140 μL of SDS (1%) and 1 mL of sodium phosphate buffer (0.2125 M, 7.0).
Free AA Determination

Free AA were determined according to McDermott et al. (2016). Samples after SGID were deproteinated by mixing equal volume of test sample with 24% (wt/vol) trichloroacetic acid. The solution was allowed to stand for 10 min before centrifuging at 14,400 × g (Microcentaur, MSE, London, UK) for 10 min at 4°C. Supernatants were removed and diluted with 0.2 M sodium citrate buffer, pH 2.2. Samples were then diluted 1 in 2 with the internal standard, norleucine, to give a final concentration of 125 nmol/mL. Amino acids were quantified using a Jeol JLC-500/V amino acid analyzer (Jeol Ltd., Garden City, Herts, UK) fitted with a Jeol Na⁺ high-performance cation exchange column.

Reverse-Phase HPLC and SDS-PAGE

Hydrolsates, intact whey proteins, the correspondent SGID samples, and the SGID control were analyzed by reverse-phase HPLC in an Agilent 1200 Series with a binary pump and a diode array detector using ChemStation for LC 3D Systems software (Agilent Technologies). Protein and peptide separation was performed at 35°C using an Agilent ZORBAX StableBond 300SB-C18 column (4.6 × 150 mm, 5 μm) with a pre-column security guard (Agilent Technologies). Samples were prepared at a concentration of 0.2% (wt/vol) in Milli-Q water and 8 μL was injected onto the column after filtration through a 0.45-μm polyethersulfone syringe filter (Sarstedt AG & Co., Nümbrecht, Germany). Mobile phase A was 0.1% trifluoroacetic acid in Milli-Q water and phase B was 0.1% trifluoroacetic acid in HPLC-grade acetonitrile. The flow rate was 0.8 mL/min and the linear gradient elution was 0 to 10 min of 5% B; 10 to 20 min of 10% B; 20 to 22 min of 15% B; 22 to 25 min of 20% B; 25 to 30 min of 30% B; 30 to 50 min of 90% B; and 50 to 54 min of 0% B. The absorbance of the eluent was recorded at 214 nm.

We employed SDS-PAGE to analyze protein profile pre- and post-SGID (Laemmli, 1970). Samples were prepared under nonreducing conditions and run on NuPAGE Novex 4–12% gradient Bis-Tris Mini Gel (Invitrogen, Carlsbad, CA). The Running Buffer MES SDS (Invitrogen) was used with a Xcell SureLock Novex Mini Cell apparatus (Invitrogen). Mark12 Unstained Standard (Invitrogen) was used as the molecular weight standard. Detection of protein bands was performed by gel staining with 0.05% Coo massie blue solution (Fisher Scientific, Hampton, NH).

Measurement of 2,2′-Azinobis(3-Ethylbenzothiazoline-6-Sulfonic Acid) Radical Scavenging Activity

The measurement of the 2,2′-azinobis(3-ethylbenzothiazoline-6-sulfonic acid) (ABTS) radical scavenging activity was performed as described by Re et al. (1999) with some modifications. Potassium phosphate monobasic and dibasic were mixed to prepared a potassium phosphate buffer (PB) at pH 7.0. A 5 mM potassium persulfate solution was prepared in Milli-Q H₂O. The ABTS dianmonium salt radical (7.3 mM) was formed by adding 2 tablets of 10 mg in 5 mM potassium persulfate solution, stirring overnight at 4°C in darkness. Absorbance of diluted ABTS solution (1:10 in PB) at 734 nm was around 0.4. After incubation overnight, the formation of the radical was checked by spectrophotometry (absorbance: 0.8–1.1). The ABTS radical solution was then diluted 1:100 in 10 mM PB (pH 7.0) and, immediately before use, the absorbance at 734 nm was checked (0.8–1.1). This solution was kept on ice in the dark to preserve stability. Protein samples were prepared in Milli-Q H₂O at 15 mg of powder/mL. The incubation was performed in the spectrophotometer sample holder to ensure dark conditions and eliminate time lost due to sample transfer after the start of the reaction. To perform the assay, 5 μL of sample or PB (control vehicle) was added to a cuvette placed in the spectrophotometer, which contained 1 mL of ABTS free radical solution. The mixture was homogenized with a pipette and the absorbance was read after 5 min on a Cary 100 Spectrophotometer (Agilent Technologies) at 734 nm. Antioxidant activity was expressed as percent ABTS radical inhibition using the formula:

\[
\%DH = \frac{(AN2 − AN1/Npb)}{100},
\]

where AN1 and AN2 are the amino nitrogen content (mg/g of protein) of the protein substrate before and after digestion, respectively, and Npb is the nitrogen content of the peptide bonds in the protein substrate (mg/g protein). This value is 123.3 mg/g of protein for whey protein (Adler-Nissen, 1979).
% radical inhibition = \[ \frac{(\text{Abs}_{734} \text{ control vehicle} - \text{Abs}_{734} \text{ sample})}{\text{Abs}_{734} \text{ control vehicle}} \times 100, \]

where \( \text{Abs}_{734} \) control vehicle is the absorbance measured at 734 nm of PB with ABTS radical solution after 5 min of incubation. \( \text{Abs}_{734} \) sample is the absorbance of samples reacting with the radical solution after 5 min. A control for the gastrointestinal fluids, including ions, enzymes, and bile extract, was run and the antioxidant value obtained was subtracted from each SGID sample result.

**Ferric-Reducing Antioxidant Power Assay**

Ferric-reducing antioxidant power (FRAP) protocol was performed according to Benzie and Strain (1996) with some modifications to achieve optimum conditions for dairy protein tests. The traditional acetate buffer has a pH of 3.6, close to the whey acid dissociation constant value (4.6), resulting in protein precipitation. As an alternative, 0.2 M HCl/KCl buffer (pH = 2.2) was used. A 10 mM 2,4,6-tris(2-pyridyl)-s-triazine (TPTZ) solution was prepared in 40 mM HCl. A 20 mM FeCl₃ solution was prepared using Milli-Q H₂O. All stock solutions were prepared fresh on the experiment day and kept in darkness to avoid photo-oxidation. A working FRAP solution was prepared at low pH by mixing 0.2 M HCl-KCl (pH 2.2) with 10 mM TPTZ-HCl and 20 mM FeCl₃ at a ratio of 10:1:1. This solution was protected from light and heated to 37°C for at least 1 h in a water bath. Test samples were prepared in Milli-Q H₂O at 15 mg of powder/mL. Trolox (a synthetic analog of vitamin E) was initially dissolved in 200 μL of pure methanol. Then a stock solution was prepared and serial dilutions (800–25 μM) in Milli-Q H₂O were used to generate a standard curve. The reaction was initiated by adding 75 μL of sample to 1.425 mL of FRAP working solution in black microcentrifuge tubes. The mixture was vortexed and allowed to incubate for 1 h at room temperature in complete darkness. After incubation, tubes were vortexed and the absorbance was read at 593 nm on a Cary 100 Spectrophotometer (Agilent Technologies). Results were expressed as micromoles of Trolox equivalents (TE) per gram of powder. The antioxidant value for the gastrointestinal fluids was subtracted from SGID sample results.

**Oxygen Radical Absorbance Capacity Assay**

The oxygen radical absorbance capacity (ORAC) assay protocol was adapted from Zulueta et al. (2009). A potassium PB at 75 mM, pH 7.2, was prepared. Digested assays were run using 0.05 mg/mL of test samples and then expressed per gram of powder. Intact assays were run using 0.1 mg/mL of test samples and expressed as above. Samples, standards, or vehicle control (PB; 20 μL) were pipetted into a black 96-well microtiter plate followed by 120 μL of 0.117 μM fluorescein solution, freshly prepared in PB. The plate was incubated at 37°C for 15 min. During this period, a 40 mM 2,2'-azobis(2-methylpropionamidine) dihydrochloride (APPH) radical solution was prepared in preheated PB and kept at 37°C. After incubation, 60 μL of APPH solution were added to each well. The fluorescence was immediately read at 90-s intervals for 2 h in a Synergy HT BioTek micro plate reader (Winooski, VT). A standard curve was performed with Trolox at 5 different concentrations ranging from 80 to 5 μM prepared in 75 mM PB. Trolox was initially dissolved in 200 μL of methanol, as described previously. Fluorescence data were normalized and area under the curve was calculated. The antioxidant reaction was considered to be finished if the final fluorescence was less than 5% of the initial value. Antioxidant activity was expressed as micromoles of TE per gram of powder. The antioxidant value for the gastrointestinal fluids was subtracted from SGID sample results.

**Statistical Analysis**

Results were compared using one-way ANOVA followed by Bonferroni’s multiple comparison post-hoc test using PASW Statistics 18 software; a P-value less than 0.05 was considered to indicate a statistically significant difference. Results were expressed as mean ± standard deviation. Each experiment was repeated at least in duplicate and on different days.

**RESULTS**

**In Vitro Antioxidant Activity of WPI**

The antioxidant activity of intact WPI and WPI hydrolysates prepared by 3 individual commercial enzymes (bromelain, neutrase, and alcalase) is detailed in Figure 1. Bromelain, a cysteine protease obtained from pineapple (Abadía-García et al., 2016), was selected because it exposes the reactive thiol group of cysteine residues so that they are available to reduce oxidative agents such as peroxyl radicals. Neutrase is a bacterial metalloprotease, commonly used to improve food protein functionality with an affinity for the AA leucine and phenylalanine (Xu et al., 2014). Alcalase is a non-specific endopeptidase purified from *Bacillus licheniformis*. Not surprisingly, commercial enzyme hydrolysates...
showed significantly higher ABTS and FRAP values than intact WPI (P < 0.05). In particular, alcalase hydrolysis of WPI resulted in 76.41 ± 2.67% ABTS radical inhibition, 26.96 ± 1.18 μmol of TE/g of powder by FRAP, and 737.36 ± 67.24 μmol of TE/g of powder by ORAC assay compared with intact WPI (6.98 ± 0.18% ABTS radical inhibition, 3.81 ± 0.46 μmol of TE/g of powder by FRAP, and 128.74 ± 48.36 μmol of TE/g of powder by ORAC; Figure 1A, B, and C). Bromelain and neustrase hydrolysis of WPI resulted in significantly higher antioxidant activity than intact WPI by FRAP and ABTS (P < 0.05), but we observed no difference in TE as determined by ORAC (Figure 1A, B, and C).

The standardized INFOGEST static in vitro digestion model was employed to ascertain if simulated upper gut transit altered the antioxidant bioactivity of these WPI samples (Figure 1). In all 3 assays, the antioxidant activity of WPI was significantly increased post-SGID (P < 0.05). Remarkably, when subjected to the hydrolytic conditions of the gastrointestinal tract, whey hydrolysates produced by alcalase exhibited significantly lower FRAP and ABTS inhibition than the alcalse-hydrolyzed sample without gastric digestion (Figure 1A and B). However, ORAC activity was increased by SGID of the WPI hydrolysate prepared by alcalase (1.144.73 ± 83.86 μmol of TE/g of powder). Reduction in antioxidant activity by FRAP and ABTS inhibition, but not by ORAC, was also observed with the bromelain-hydrolyzed WPI (Figure 1A, B, and C). The WPI hydrolyzed by neutrase and subjected to SGID exhibited significantly (P < 0.05) lower antioxidant activity by ABTS inhibition but significantly higher antioxidant activity by ORAC and FRAP (Figure 1A, B, and C). Interestingly, subjecting commercial WPI to hydrolysis by either bromelain or neutrase before SGID did not enhance its antioxidant activity, 799.93 ± 56.68 and 908.65 ± 126.11 μmol of TE/g of powder, respectively, compared with WPI post-SGID (885.18 ± 121.91 μmol of TE/g of powder) as determined by ORAC. Only alcalase hydrolysis contributed to higher ORAC value post-SGID compared with the results obtained for WPI post-SGID. Hydrolysates generated by alcalase or bromelain had similar FRAP capabilities to WPI post-SGID (Figure 1B). Only neutrase hydrolysis of WPI post-SGID was significantly better at reducing ferric ions than WPI post-SGID. In the case of ABTS inhibition, commercial enzyme hydrolysis of WPI was not significantly different post-SGID compared with WPI post-SGID (Figure 1A). Antioxidant activity measured after SGID of WPI was significantly higher when measured by ORAC compared with the hydrolysates prepared with the commercial enzymes bromelain or neutrase (Figure 1C). The FRAP capability of WPI post-SGID was similar to alcalase- and bromelain-hydrolyzed WPI (Figure 1B). The WPI post-SGID inhibited less ABTS (26.02 ± 1.84%) than alcalase (76.41 ± 2.67%), bromelain (38.33 ± 1.91%), or neutrase (41.69 ± 6.89%) hydrolyzed WPI (Figure 1A).

**DH of WPI**

To compare the extent of hydrolysis, DH values were obtained for whey samples. The WPI post-SGID had a DH of 49.96%, with alcalase hydrolysis of WPI yielding 13.60% followed by neutrase, 5.28%, and bromelain, 4.31%. However, the comparative kinetics have not been investigated here. Degradation of WPI after hydrolysis with commercial enzymes and after SGID was also evaluated by HPLC (Figure 2). The profile of intact WPI revealed clear peaks correspondent to the caseinomacropeptide, BSA, α-LA, and β-LG (retention times of 31–42 min). The use of alcalase, bromelain, and neustrase showed distinct WPI peptide peaks explained by the diverse enzyme specificity. Fractions at lower retention times contained hydrophilic peptides with higher polarity, whereas hydrophobic peptides with less polarity were eluted at higher retention times. Alcalase hydrolysis resulted in a large number of peaks, primarily at 24 to 33 min. However, bromelain hydrolysates, with the lowest DH, had the least number of peptide peaks (18–19 and 29–31 min). The major peptides obtained after neutrase hydrolysis appeared at the highest retention time (24–39 min), which could explain its poor solubility in water. The chromatograms of each hydrolysate showed further degradation of whey peptides after being exposed to gut conditions. After this breakdown, hydrolysates presented similar HPLC profiles to the chromatogram obtained for WPI after SGID. The HPLC profiles of SGID samples revealed several peaks that corresponded to the hydrolytic activity of pancreatin. The HPLC profile of SGID control did not indicate significant autocatalysis.

**In Vitro Antioxidant Activity of Individual Whey Proteins**

To identify which whey protein fraction contributed to greater antioxidant activity, individual whey proteins β-LG, α-LA, BSA, and LF were assessed for their antioxidant activity before and after SGID (Figure 3). Intact α-LA (20.97 ± 1.44%) and LF (18.06 ± 0.44%) were the most effective at inhibiting the radical cation ABTS compared with the other intact proteins (P > 0.05). These samples also exerted the most potent ferric to ferrous ion reduction, tested by FRAP assay (intact α-LA = 8.19 ± 1.19 μmol of TE/g of powder; intact LF = 7.85 ± 0.84 μmol of TE/g of powder). All intact proteins showed peroxyl radical inhibition with values
ranging from 117.65 ± 37.23 μmol of TE/g of powder for LF and 365.14 ± 46.07 μmol of TE/g of powder for α-LA. Antioxidant activity of all proteins appeared significantly enhanced after 240 min of exposure to the pH and enzymes of the stomach and duodenum. Of note was α-LA post-SGID, which exhibited the highest antioxidant activity by all 3 methods (37.29 ± 0.73% ABTS inhibition; FRAP = 15.56 ± 1.21 μmol of TE/g of powder; ORAC = 1,825.94 ± 50.21 μmol of TE/g of powder).

**DH of Individual Whey Proteins by Simulated Upper Gut Transit**

The DH of whey protein samples was determined after SGID by the TNBS method. Values ranged from

![Figure 2](image-url)

*Figure 2*. Reverse-phase HPLC profiles at 214 nm for whey protein isolate (WPI) and whey protein hydrolysates. Samples were prepared at 2 mg/mL and eluted in 0.1% trifluoroacetic acid. (A) Intact WPI and the graph below it simulated gastrointestinal-digested (SGID) WPI, where individual peaks correspond to (1) caseinomacropeptide, (2) BSA, (3) α-LA, and (4) β-LG; (B) WPI hydrolyzed with alcalase for 180 min at 50°C, pH 8.0, with an enzyme-to-substrate ratio of 1:100 and associated SGID sample; (C) WPI hydrolyzed with bromelain for 180 min at 50°C, pH 7.0, with an enzyme-to-substrate ratio of 1:100 and associated SGID sample; and (D) WPI hydrolyzed with neutrase for 180 min at 50°C, pH 7.0, with an enzyme-to-substrate ratio of 1:100 and associated SGID sample. Pancreatin peaks from SGID process are indicated by P. The HPLC profiles of SGID control without WPI were also performed. AU = arbitrary units.
49.96 ± 1.95% for WPI to 65.10 ± 4.61% for BSA (Table 1). The chromatograms of the intact whey proteins (Figure 4) showed distinct peaks (β-LG = 40–42 min, α-LA = 38.5–40 min, BSA = 37.5–39.5 min, LF = 33 and 38.5–39.5 min), indicating a high level of native conformation. After SGID, these peaks disappeared, which indicated native protein degradation. The loss of each intact whey protein after the harsh conditions of the stomach and duodenum was supported by SDS-PAGE (Figure 5).

**Free AA Content in Gastrointestinal-Digested Whey Proteins**

To determine if the high antioxidant activity of α-LA could be explained by the free AA released after SGID, AA analysis was performed on WPI and individual whey protein fractions post-SGID (Table 2). The highest concentration of free AA (50.426 μmol of AA/g of powder) was released from α-LA. Leucine was the predominant AA in SGID of WPI, β-LG, BSA, and LF (6.208–8.154 μmol/g of powder). Lysine was the most abundant (8.154 μmol/g of powder) AA found in α-LA post-SGID. Interestingly, proline was not detected in any of the samples, which can be explained by the lack of specificity of the human gastrointestinal enzymes for this AA. Known antioxidant AA, such as tryptophan, phenylalanine, tyrosine, cysteine, and histidine (Hernández-Ledesma et al., 2005; Power-Grant et al., 2016), were present at relatively high concentrations in all SGID samples compared with other AA. Interestingly, α-LA released the highest amount of tryptophan (6.955 μmol/g of powder).

**DISCUSSION**

Our in vitro studies indicate that the hydrolytic conditions of the gut negate the need for prior hydrolysis of WPI from an antioxidant bioactivity viewpoint. The α-LA, of the 4 major whey protein fractions studied, showed the highest antioxidant properties post-SGID.
when tested by ORAC, ABTS, and FRAP. This appears to accord with a high level of digestion (57.59% DH) and greatest release of the antioxidant AA tryptophan, 6.955 μmol of tryptophan/g of protein.

Conditions in the stomach and duodenum (pepsin, trypsin, chymotrypsin, and acidic pH) appear to functionalize WPI in reducing ferric ions, inhibiting ABTS and scavenging peroxyl radicals possibly by the release of bioactive peptides and AA. In agreement, Power-Grant et al. (2015) observed that WPC exposed to simulated SGID had significantly higher ORAC values (36,305 ± 3,390 μmol of Trolox/100 g of powder) than

Figure 4. Reverse-phase HPLC profiles at 214 nm for individual whey proteins at 2 mg/mL and eluted in 0.1% trifluoroacetic acid. (A) Intact β-LG and the graph below it simulated gastrointestinal-digested (SGID) β-LG; (B) intact α-LA and SGID α-LA; (C) intact BSA and SGID BSA; and (D) intact lactoferrin and SGID lactoferrin. P = peaks corresponding to pancreatin; AU = arbitrary units.
intact WPC (13,662 ± 1,018 μmol of Trolox/100 g of powder). Similar to our study, antioxidant activity post-SGID was similar to that obtained with WPC hydrolyzed (32% DH) beforehand (37,391 ± 2,298 μmol of Trolox/100 g of powder). Interestingly, prior hydrolysis of WPC appeared to influence the antioxidant activity after simulated upper gut transit. The WPC with a DH of 45% at processing and then subjected to gastric conditions exhibited 60,613 ± 4,540 μmol of TE/100 g by ORAC assay, whereas a WPC hydrolysate (32% DH) when subjected to gastric conditions had 44,489 ± 2,064 μmol of TE/100 g. Of note was the significant decrease in ORAC values for the WPC hydrolysate (DH 45%) post-SGID compared with its non-SGID equivalent. We found no differences in ORAC values between the WPC hydrolysate DH 32% pre- and post-SGID. In contrast, other forms of processing designed to denature whey proteins, such as high hydrostatic pressure, may increase antioxidant activity even after SGID. Iskandar et al. (2015) observed that hydrostatic pressure at 550 MPa for 1 min at 20°C of WPI can enhance its WPI ferric-reducing activity by 21% after SGID (Iskandar et al., 2015). Pressurized WPI subjected to SGID also protected intestinal cells from oxidative stress after 23 h of treatment (Piccolomini et al., 2012).

The DH for WPI obtained in our study after SGID is in agreement with He et al. (2015), who showed a 49% digestibility by TNBS method after 3 h of digestion. However, these values are higher than the ones obtained by Conway et al. (2013) for WPC (8.2 ± 0.5%) after 2 h of pepsin digestion followed by 3 h of trypsin treatment. The lower protein degradation was also noticeable by SDS-PAGE, where, in Conway et al. (2013), the β-LG band was visible after the digestion process. Adjonu et al. (2013) also obtained DH values ranging between 12 and 13% for WPI hydrolyzed 12 to 24 h by

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<th>Table 2. Free AA released from 1 g of protein powder following simulated gastrointestinal digestion</th>
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1WPI = whey protein isolate, which contains β-LG (50–60%), α-LA (15–25%), BSA (6%), lactoferrin (LF; <3%).
gastrointestinal enzymes individually. This difference supports previous knowledge that DH can be markedly increased by combination of enzymes with different specificity (Lee and Hur, 2017), as in the INFOGEST protocol that more closely resembles in vivo conditions, and that the antioxidant activity is not defined by the DH (Pena-Ramos and Xiong, 2001). It is important to note that SGID samples in our study corresponded to the end of the intestinal phase. However, in vivo, the intestinal lumen will be exposed to these fractions but also to less-hydrolyzed fractions arriving from the stomach (Dupont et al., 2010). Interestingly, Joubran et al. (2015) demonstrated that the antioxidant capacity of α-LA increased in later digestion phases where an in vitro model of the infant gut was employed. The in vitro duodenal samples exhibited higher antioxidant activity than samples from an in vitro model of the adult duodenum. In contrast, in vitro gastric infant samples were less antioxidant than the adult samples. Antioxidant capacity of dairy proteins during gut transit is therefore not only influenced by gut location, but also life stage (Joubran et al. 2015).

The SGID results may indicate that the release of free AA, in addition to individual bioactive peptides, governs antioxidant activity. This is supported by the similar antioxidant activity obtained after whey protein hydrolysis with different enzymes. These samples are abundant in AA such as histidine, phenylalanine, and tryptophan, which can exert their antioxidant activity by hydrogen atom transfer (i.e., ORAC and ABTS) and by electron transfer mechanisms (i.e., FRAP; Elias et al. 2008). Power-Grant et al. (2016) observed that tryptophan has an ORAC value of 1,773,102 ± 25,218 μmol of TE/100 g of AA dry weight, methionine has an ORAC value of 545,413 ± 6,528, histidine has an ORAC value of 191,762 ± 7,308, and cysteine has an ORAC value of 167,600 ± 9,077 μmol of TE/100 g of AA. Therefore, it was proposed that these AA play a major contributing role to milk protein concentrate antioxidant activity. However, it is also noteworthy that several peptides have been identified from whey hydrolysates using Corolase PP [AB Enzymes, Darmstadt, Germany; β-LG: f(15–18) VAGT, f(19–29) WYSL, f(24–26) MAA, f(42–46) YVEEL, f(71–74) IIAE, f(145–149) MHIRL; Hernández-Ledesma et al., 2005; O’Keeffe et al., 2017], thermolysin [EC 3.4.24.27; α-LA: f(101–104) INYW, f(115–118) LDQW; Sadat et al., 2011], and trypsin [β-LG: f(15–20) VAGTWY; Power et al., 2014]. All of these hydrolysates showed antioxidant activity, and the individual peptides, once synthesized, also demonstrated antioxidant properties. Therefore, the antioxidant power of whey hydrolysates is likely to be generated by possible synergy between individual AA and encrypted peptides, which are released during whey protein hydrolysis.

Whey (WPI, WPC, and fresh whey) hydrolysates produced by alcalase have been identified with strong antioxidant activity (Kou et al., 2013; Zhang et al., 2013). We also observed that hydrolysis with alcalase produces hydrolysates with the highest antioxidant activity. Dryáková et al. (2010) observed the highest ABTS inhibition (54%) after hydrolysis of WPC with alcalase. Lin et al. (2012) also described alcalase hydrolysates from WPC as the most potent ferric ion-reducing (0.55 mM FeSO₄ equivalents) agents compared with other commercial enzymes pepsin, trypsin, and flavorzyme. Alcalase and neuramidase hydrolysates from whey proteins also protected lung fibroblast and umbilical vein endothelial cells by boosting intracellular antioxidant defenses (Kong et al., 2012; O’Keeffe and Fitzgerald, 2014). In fact, the hydrolysis of β-LG with alcalase produced the most antioxidant fraction containing the peptides f(27–38) DIQKVAGTWYSL, f(33–38) GTWYSL, f(39–48) AMAASDILL, f(40–48) MAASDILL, f(61–73) ELKPTPEDLEIL, f(87–98) IIAEKTKIPAVF, f(112–121) DTDYKKYLLF, and f(165–172) LSFNPTQL (Bamdad et al., 2017). It is important to note that antioxidant activities from commercial enzyme hydrolysates in our study are likely to be affected by the standard thermal step required for enzyme inactivation, which is known to affect protein denaturation and aggregation (Joyce et al., 2018). In addition, thermal processing influenced the peptide release from milk proteins (Kopf-Bolanz et al., 2014).

The most potent proteins inhibiting ABTS radical and reducing iron were LF and α-LA; the latter was also the more efficient at neutralizing peroxy radicals in ORAC assay. In agreement with our study, Hernández-Ledesma et al. (2005) observed that hydrolysates obtained from α-LA after 24 h of hydrolysis using individual gastric enzymes had significantly higher ORAC values than β-LG hydrolysates (1.065 ± 0.056 and 0.701 ± 0.033 μmol of Trolox/mg of protein, respectively; Hernández-Ledesma et al., 2005). Camel whey protein, which is rich in α-LA (27%) but devoid of β-LG (Hailu et al., 2016), also exerted significantly higher ABTS radical inhibition than its bovine equivalent (Salami et al., 2010). However, Clausen et al. (2009) reported similar ORAC values for 25-μL sample quantities of intact α-LA (0.07 ± 0.04 μM TE) and intact β-LG (0.14 ± 0.09 μM TE) isolated from bovine milk whey by size exclusion chromatography, which is in agreement with our ORAC values for intact α-LA and β-LG. Recently, bovine α-LA (97.2% protein) following an infant in vitro SGID also showed higher DPPH-scavenging potential compared with intact α-LA (Joubran et al., 2017).
Indeed, 4 synthesized dipeptides whose sequences are present in α-LA sequence, EW (25–26 AA), WC (60–61 AA), YW (103–104 AA), and WL (104–105 AA and 118–119 AA), inhibited 50% DPPH radical at concentrations ranging from 0.26 to 3.07 mM (Nongoinnerma and Fitzgerald, 2013). In our study, with a DH value of 57%, α-LA released the highest total concentration (17.441 μmol of AA/g of protein) of the most reactive of 57%, α-LA released the highest total concentration (17.441 μmol of AA/g of protein) of the most reactive amino acids (AA) against peroxyl radicals (tryptophan, methionine, cysteine, tyrosine, and phenylalanine) as described by Hernández-Ledesma et al. (2005) and Power-Grant et al. (2016).

CONCLUSIONS

In conclusion, pre-enzymatic hydrolysis of WPI (based on 3 commercially available proteases tested) does not enhance antioxidant activity during simulated gut transit. This implies that the proteolytic conditions of the gut are capable of generating equivalent antioxidant capacity during SGID of WPI. The α-LA fraction exerted the best antioxidant properties, which are not only maintained after SGID but increased. This suggests that α-LA is the preferred whey protein candidate to supplement food products to counteract free radicals in our body and boost antioxidant defenses.

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