Cold enzymatic bleaching of fluid whey

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ABSTRACT

Chemical bleaching of fluid whey and retentate with hydrogen peroxide (HP) alone requires high concentrations (100–500 mg of HP/kg) and recent studies have demonstrated that off-flavors are generated during chemical bleaching that carry through to spray-dried whey proteins. Bleaching of fluid whey and retentate with enzymes such as naturally present lactoperoxidase or an exogenous commercial peroxidase (EP) at cold temperatures (4°C) may be a viable alternative to traditional chemical bleaching of whey. The objective of this study was to determine the optimum level of HP for enzymatic bleaching (both lactoperoxidase and EP) at 4°C and to compare bleaching efficacy and sensory characteristics to HP chemical bleaching at 4°C. Selected treatments were subsequently applied for whey protein concentrate with 80% protein (WPC80) manufacture. Fluid Cheddar whey and retentate (80% protein) were manufactured in triplicate from pasteurized whole milk. The optimum concentration of HP (0 to 250 mg/kg) to activate enzymatic bleaching at 4°C was determined by quantifying the loss of norbixin. In subsequent experiments, bleaching efficacy, descriptive sensory analysis, and volatile compounds were monitored at selected time points. A control with no bleaching was also evaluated. Enzymatic bleaching of fluid whey and retentate at 4°C resulted in faster bleaching and higher bleaching efficacy (color loss) than bleaching with HP alone at 250 mg/kg. Due to concentrated levels of naturally present lactoperoxidase, retentate bleached to completion (>80% norbixin destruction in 30 min) faster than fluid whey at 4°C (>80% norbixin destruction in 12 h). In fluid whey, the addition of EP decreased bleaching time. Spray-dried WPC80 from bleached wheys, regardless of bleaching treatment, were characterized by a lack of sweet aromatic and buttery flavors, and the presence of cardboard flavor concurrent with higher relative abundance of 1-octen-3-ol and 1-octen-3-one. Among enzymatically bleached WPC80, lactoperoxidase-bleached WPC80 contained higher relative abundance of 2,3-octadienone, 2-pentyl furan, and hexanal than those bleached with added EP. Bleach times, bleaching efficacy, and flavor results suggest that enzymatic bleaching may be a viable and desirable alternative to HP bleaching of fluid whey or retentate.

Key words: whey, lactoperoxidase, flavor, bleach

INTRODUCTION

Whey is a by-product of cheese manufacture and is often further processed into value-added products, such as whey protein concentrate 34 or 80% or whey protein isolate (>90% protein). Typical whey processing steps include fat separation, pasteurization, bleaching, UF, diafiltration, and spray drying. The flavor of fluid whey carries through into the final spray-dried products (Croissant et al., 2009), and consumers and product manufacturers demand that dried whey ingredients be colorless with a bland flavor (Kang et al., 2010). The manufacture of Cheddar cheese has continued to increase and Cheddar whey is one of the main sources of cheese whey. Norbixin, a natural orange-colored carotenoid, is added to Cheddar cheese milk to impart the desired orange color and a portion of the norbixin is retained in the fluid whey (Kang et al., 2010) and must be bleached. Off-flavors in dried whey proteins associated with bleaching, either with benzoyl peroxide or hydrogen peroxide (HP), have been well documented in the literature (Croissant et al., 2009; Listiyani et al., 2011, 2012; Jervis et al., 2012). Due to the increased demand for bland, colorless whey ingredients and international concerns with the use of benzoyl peroxide and increasing concerns with HP, chemical bleaching alternatives are desirable (Campbell et al., 2012; Kang et al., 2012). Campbell et al. (2012) recently demonstrated that as little as 10 mg of HP/kg was sufficient for greater than 80% norbixin destruction by lactoperoxidase (LP) in fluid whey at 35°C. Enzymatic bleaching, either using the native LP system or by adding an exogenous peroxidase (EP) has yet to be fully explored.

Lactoperoxidase, a native enzyme found in milk, is often used to increase storage stability and reduce the loss of fresh milk quality due to microbial spoilage. Lactoperoxidase is a member of the peroxidase family and...
when its activators thiocyanate and HP are present, hypothiocyanate, a potent antimicrobial, is produced (Reiter and Harnulv, 1982). In addition to milk preservation, the LP system can be used to bleach whey (Bottomley et al., 1989; Campbell et al., 2012). The strong oxidizing capacity of hypothiocyanate results in the destruction of carotenoid conjugation and subsequent color loss of norbixin in cheese whey. Using the LP system to bleach whey can be highly variable, as levels of LP can vary depending on the lactation cycle of the cow, season, feeding regimen, and breed (Kussendrager and van Hooijdonk, 2000). Similar to LP, thiocyanate concentration in milk and whey can vary widely due to feeding regimen (Seifu et al., 2005). The third component of the LP system, HP, is not normally detected in raw milk and is typically added exogenously. Hydrogen peroxide can be generated endogenously by bacteria, although amounts sufficient to activate the LP system may not be generated (Seifu et al., 2005). Depending on the milk, any 1 of the 3 components that make up the LP system could limit LP activity.

To facilitate enzymatic whey bleaching, a commercial EP is available and can be added to fluid whey product in small quantities to help achieve desired and consistent bleaching efficacy. This enzyme, MaxiBright (MB), is derived from a mushroom, Marasmius scorodonius (Zorn et al., 2003). Very little is known about the enzyme mechanism compared with that of LP; however, it is known that both of these enzymes require similar amounts of HP to activate their respective systems (Bottomley et al., 1989; Zorn et al., 2003). Since the original patent was filed in 2006, several studies have addressed the bleaching capacity of MB on β-carotene in model systems (Scheibner et al., 2008; Pühse et al., 2009; Zelena et al., 2009); however, the bleaching efficacy and subsequent effects on the flavor of MB in conjunction with the natural LP system in fluid whey has yet to be investigated. Studies have demonstrated that chemical bleaching at colder temperatures (<10°C) results in less lipid oxidation (Listiyani et al., 2011). Additionally, colder temperatures enhance membrane stability, microbial quality, and protein integrity. As such, cold bleaching is an attractive process. The objective of this study was to optimize enzymatic bleaching of whey and retentates with both LP and EP at 4°C and to evaluate their subsequent effects on the flavor of whey protein concentrate with 80% protein (WPC80).

MATERIALS AND METHODS

Experimental Design Overview

The study had 2 experimental components: liquid whey and retentate trials and the manufacture of WPC80. Optimum HP levels to activate the LP and EP systems were first determined. Liquid whey and retentate trials were then conducted to determine optimum bleach times at 4°C. The liquid whey treatments with the most bleaching and the fastest bleaching times were then selected for WPC80 manufacture. All treatments within each trial were made from the same lot of milk. All experiments were conducted in triplicate.

Production of Liquid Whey

Cheddar whey was manufactured from HTST (15 s at 72°C) pasteurized whole milk (720 kg/h; model T4 RGS-16/2; SPX Flow Technology, Greensboro, NC). The milk was then cooled to 31°C and transferred to a cheese vat (Kusel Equipment Co., Watertown, WI). Colored Cheddar whey manufacture proceeded as described by Campbell et al. (2011). The whey was drained from the curds at pH 6.3 and a sieve was used to remove any remaining particles. The whey was immediately processed with a hot bowl cream separator (model SI600E; Agrilac, Miami, FL) to reduce the fat content. Fat-separated, fluid whey was then HTST pasteurized as described previously. Whey was cooled to 4°C before bleaching experiments.

Production of Retentate

Fat-separated, pasteurized fluid whey was transferred into a 102-L stainless vat (Fermentor; Blichmann Engineering, Lafayette, IN) equipped with a coil heater (1.3-cm outer diameter; PAC Stainless Ltd., Seattle, WA) and heated to 50°C while recirculating using a peristaltic pump (model 77410-10; Millipore Inc., Billerica, MA). Once the desired temperature was reached, UF commenced. The UF system (model Pellicon 2; Millipore Inc.) was equipped with 10 polyethersulfone cartridge membrane filters (model P2B010V05; 10-kDa nominal separation cutoffs, 0.5 m² surface area; Millipore Inc.). Each sample was run through a peristaltic pump (model 77410-10; Millipore Inc.) and the UF assembly using silicone tubing (model 96440-73; Millipore Inc.) was connected to the vat. Pumps, pump heads, and tubing were all obtained from Cole-Parmer (Vernon Hills, IL). Ultrafiltration and diafiltration continued until the retentate reached 80% protein content on a dry basis (wt/wt), confirmed by a Sprint rapid protein analyzer (CEM Corp., Matthews, NC). Retentates were then collected and cooled to 4°C before bleaching experiments.

Activation of the LP or EP System

The optimum level of HP to activate the LP or EP system was determined by adding 0, 5, 10, 15, 20, 25,
30, 35, 40, 45, 50, 55, 60, 65, 70, 75, 80, 85, 90, 95, 100, 150, 200, or 250 mg/kg HP to pasteurized fat-separated liquid Cheddar whey or retentate (80% protein on a dry basis; 10% solids). Bleaching was then carried out as described below. The concentration of HP that resulted in the most bleaching (10 mg/kg in fluid whey and 15 mg/kg in retentate, according to norbixin destruction determined via HPLC) was selected for further trials.

**Optimum Bleaching Time**

Liquid whey or retentate [10% solids (wt/vol); 80% protein on a dry basis (wt/wt)] was placed in amber glass jars in an ice bath and allowed to equilibrate to 4°C. To activate the LP system, 10 mg/kg of HP (liquid whey) or 15 mg/kg of HP [retentate; 35% (wt/vol); Nelson Jameson Inc., Marshfield, WI] was added and allowed to bleach in an ice bath with gentle agitation. To activate the EP system, 2 dairy bleaching units (DBLU) of MB (DSM, Delft, the Netherlands) was added to the liquid whey or retentate and agitated gently. To that, 10 or 15 mg of HP/kg was added and allowed to bleach in the ice bath with gentle agitation. Aliquots of samples were removed at appropriate time points (fluid whey: 0, 0.5, 1, 2, 4, 6, and 24 h; retentate: 0, 3, 5, 7, 10, 15, 20, 30, 40, and 50 min, and 1, 2, 4, 6, and 24 h). Peroxide test strips (EMD Chemicals, VWR International LLC, West Chester, PA) were used to determine if any HP remained after the bleaching treatment. To consume the remaining HP to stop peroxidase activity, catalase (FoodPro CAT; Danisco) was added at a rate of 20 mg/kg. Measurements, including norbixin, volatile compound analyses, and descriptive analysis were performed immediately.

**Production of WPC80**

Industrially, due to manufacturing constraints, bleaching most frequently occurs at the fluid whey level. We also observed differences in the amount of time required for LP and EP bleaching in fluid whey. For these reasons, liquid whey was selected as the bleaching point for WPC80 trials. Treatments were selected based on current industrial practices and from the previous liquid whey trials to achieve maximum bleaching at 4°C [LP: 10 mg of HP/kg for 12 h, EP: 2 DBLU of MB and 10 mg of HP/kg for 1 h, HP: 250 mg/kg for 12 h, and control (no bleaching) for 12 h at 4°C for manufacture of WPC80]. Colored fat-separated cheese whey was freshly manufactured as previously described. The pasteurized separated whey was placed in sanitized 38-L milk cans at 4°C. Two of 3 treatments were administered immediately: LP [10 mg/kg of HP (35% wt/vol; Nelson Jameson Inc.]) or HP (250 mg/kg) and allowed to bleach overnight. The following morning, the last treatment (EP) was administered (2 DBLU of MB and 10 mg of HP/kg) and allowed to bleach for 1 h. If any HP remained after treatment, catalase was added at a rate of 20 mg/kg (FoodPro CAT; Danisco) to consume the remaining HP and stop peroxidase activity.

Bleached wheys were transferred into a 102-L stainless-steel vat (Fermenator; Blichmann Engineering) equipped with a coil heater (1.3-cm outer diameter; PAC Stainless Ltd.) and heated to 50°C while recirculating using a peristaltic pump (model 77410-10; Millipore Inc.). Once the desired temperature was reached, UF commenced. The UF system (model Pellicon 2, Millipore Inc.) was equipped with 5 polyethersulfone cartridge membrane filters (model P2B010V05; 10-kDa nominal separation cutoffs, 0.5 m² surface area; Millipore Inc.). Each sample was run through a peristaltic pump (model 77410-10; Millipore Inc.) and the UF assembly using silicone tubing (model 96440-73; Millipore Inc.) that was connected to the vat. Pumps, pump heads, and tubing were all obtained from Cole-Palmer. Ulfiltration and diafiltration continued until the retentate reached 80% protein content on a dry basis (wt/wt), confirmed by a Sprint rapid protein analyzer (CEM Corp.). Retentates were then collected and spray dried (model Lab 1; Anhydro Inc., Soborg, Denmark). The inlet temperature was 200°C and the outlet temperature was 90°C. Total spray-drying time was about 1 h. The powder was collected and stored in Mylar bags at −80°C following production. All treatments were manufactured from the same lot of milk and the experiment was conducted in triplicate.

**Compositional Analysis**

Total solids of defatted liquid whey and WPC80 were determined by air-oven drying (AOAC International, 2000; method number 990.20; 33.2.44). In powdered WPC80, fat was quantified by the Mojonnier method (AOAC International, 2000; method numbers 932.06 and 989.05). In defatted liquid wheys, fat was quantified using the CEM Smart Trac rapid fat analyzer (CEM Corp.). Protein was determined using the Kjeldahl method in powdered whey proteins (AOAC International, 2000; method number 991.20; 33.2.11) or using the Sprint Rapid Protein Analyzer (CEM Corp.) if the whey was liquid. Mineral analysis (phosphorus, calcium, magnesium, potassium, sulfur, sodium, and iron) was done by the North Carolina State University Analytical Services Laboratory (Raleigh, NC) using a standard dry ash method with inductively coupled plasma optical emission spectroscopy (Lloyd et al., 2009). All samples were measured in duplicate.
Hunter L*a*b* Values

The WPC80 were measured in both powder form (10 g) and liquid form (10 mL of 10% [wt/vol] solution). Ten milliliters of the sample [rehydrated at 10% (wt/vol) solids, if necessary] was placed into the bottom of a 60 × 15-mm polystyrene Petri dish (Becton Dickinson, Franklin Lakes, NJ). The color of WPC80 was measured using a Minolta Chroma meter (CR-410; Konica Minolta Sensing Americas Inc., Ramsey, NJ). Each sample was measured in duplicate. Before measurements were taken, a factory-supplied calibration plate was used to calibrate the instrument. The Hunter CIE L*a*b* color scale (where L* is the lightness of the color, a* is its position between red/magenta and green, and b* is its position between yellow and blue) was used. Reflectance values were taken with a white calibration plate as the background.

Norbixin Extraction and Quantification

Norbixin is the primary carotenoid in water-soluble annatto extracts and was extracted and measured to determine percentage annatto destruction and bleaching efficacy (Kang et al., 2010). Norbixin was extracted and quantified using HPLC. To extract defatted fluid wheys, 200 μL was placed into a 2-mL microcentrifuge tube (VWR International LLC). To this, 800 μL of dilution solution [80% acetonitrile/20% water with 0.1% (wt/vol) formic acid; EMD Chemicals, VWR International LLC] was added. The solution was vortexed and centrifuged at 14,000 × g (Microfuge 18 centrifuge; Beckman Coulter, Brea, CA). The supernatant was removed and placed into vials for quantification by HPLC. To extract powders, samples were first reconstituted to 10% solids using deionized water. Then, 100 μL of sample was placed into a 2-mL microcentrifuge tube. To this, 900 μL of dilution solution was added and the sample was vortexed and centrifuged as previously described. The supernatant was removed and placed into vials for quantification by HPLC. The extraction procedure and measurements were performed with premium full-spectrum F885 flat sheet filters covering all lights (Ergomart, Dallas, TX) to minimize norbixin isomerization and degradation (Mercadante, 2008).

Quantification was conducted using HPLC (Waters 1525 Binary Pump; Waters Corp., Milford, MA). Isocratic mobile phase [70% acetonitrile/30% water with 0.1% (wt/vol) formic acid; EMD Chemicals, VWR International LLC] was used at a flow rate of 1 mL/min pumped through a binary pump (Waters 1525; Waters Corp.). Fifty microliters of the sample was injected (Waters 2707 Autosampler; Waters Corp.) onto the column (Phenomenex Kinetex 2.6-μm particle size, 10-cm length, 4.6-mm i.d., 100-A pore size; Phenomenex Inc., Torrance, CA), which was heated to 40°C. The injector temperature was set to 4°C. The sample was sent through a photodiode array detector (Waters 2998; Waters Corp.). A standard curve was created by rehydrating norbixin powder [45% (wt/vol); Chr. Hansen, Milwaukee, WI] in 2.5% (wt/vol) potassium hydroxide (BDH; VWR International LLC) and then diluting in mobile phase. The maxima used for calculation was 482 nm. Norbixin concentration was calculated by TS and correction for dilution during the extraction and solid-phase extraction processes.

Descriptive Sensory Analysis

Sensory analysis was conducted on defatted fluid wheys, retentates, and rehydrated WPC80 (10% wt/vol) using a trained descriptive sensory panel and an established dairy flavor language (Drake et al., 2003, 2009). Panelists (n = 8) each had more than 150 h of previous experience with the sensory analysis of fluid and dried whey products using the Spectrum descriptive analysis method (Meilgaard et al., 2007). All sensory testing was conducted in accordance with the North Carolina State University Institutional Review Board for Human Subjects guidelines.

Defatted fluid wheys, retentates, or reconstituted WPC80 [10% solids (wt/vol)] were evaluated by placing 30 mL in 3-digit-coded 60-mL lidded cups (Solo Cup Co., Champaign, IL). Preparations were conducted with overhead lights off to avoid exposure to light. Samples were evaluated by each panelist in duplicate. Sensory data were collected on paper ballots or using Compusense finite (release 4.8; Compusense Inc., Guelph, ON, Canada).

GC-MS

Selected volatile compounds in defatted fluid wheys and WPC80 powder were extracted by solid-phase microextraction (SPME) using selective ion monitoring. Volatile compounds were selected from previous studies and were compounds that were relevant to flavor or bleaching, or both (Croissant et al., 2009; Campbell et al., 2012; Jervis et al., 2012; Kang et al., 2012; Listiyani et al., 2012). Compounds were then separated and identified by GC-MS using a modified method of Liaw et al. (2010). Liquid samples were tested the day of manufacture and spray-dried powders were reconstituted at 10% solids (wt/vol) and evaluated within 7 d. All samples contained 10% (wt/vol) sodium chloride (Fisher Scientific, Fairlawn, NJ), and 10 μL of internal standard solution (2-methyl-3-heptanone in methanol at 81 mg/kg; Sigma-Aldrich, Milwaukee, WI) in 20-
mL autosampler vials with steel screw tops containing silicone septa faced in Teflon (MicroLiter Analytical Supplies Inc., Suwanee, GA). Samples were injected using a CombiPAL autosampler (CTC Analytics AG, Zwingen, Switzerland) attached to an Agilent 6890N GC with 5973 inert MSD (Agilent Technologies Inc., Santa Clara, CA). Samples were maintained at 5°C before fiber exposure. Samples were equilibrated at 40°C for 25 min before 30-min fiber exposure of a 1-cm divinylbenzene/Carboxen/polydimethylsiloxane (DVB/ CAR/PDMS) fiber (Supelco Inc., Bellefonte, PA) at 31 mm with 4-s pulsed agitation at 250 rpm. Fibers were injected for 5 min at a depth of 50 mm.

The GC method used an initial temperature of 40°C for 3 min, which was increased at 10°C/min to 250°C and then held for 5 min. The SPME fibers were introduced into the split/splitless injector at 250°C. A Zb-5ms column (Zb-5ms 30-m length × 0.25-mm i.d. × 0.25-μm film thickness; Phenomenex Inc.) was used for all analyses at a constant flow rate of 1 mL/min. Purge time was set at 1 min. The MS transfer line was maintained at 250°C, with the quad at 150°C and source at 250°C. Compounds were identified using the NIST 2005 library of spectra (www.nist.gov) and comparison of spectra of authentic standards injected under identical conditions. Relative abundance for each compound was calculated using the calculated recovery of the internal standard concentration to determine relative concentrations of each compound. Retention indices were calculated using an alkane series (Sigma-Aldrich, Milwaukee, WI; van Den Dool and Kratz, 1963).

**Statistical Analysis**

All data was analyzed by a one-way ANOVA using a general linear model with the Fisher least significant difference for means separation (XLSTAT, version 2009.1.02; Addinsoft Inc., New York, NY). Replication was designated as a random effect.

**RESULTS**

**Fluid Whey and Retentate**

**Composition.** Compositionally, samples were not different (P > 0.05). Defatted fluid whey averaged 6.71 ± 0.05% solids, 0.89 ± 0.05% protein, and 0.0 ± 0.01% fat. Retentates (80% protein on a dry basis) averaged 10.6 ± 0.45% solids, 8.80 ± 0.20% protein, and 0.67 ± 0.05% fat. All measurements are reported on a wet-weight basis.

**Norbixin and Color Analysis.** Experiments were conducted to determine the optimum amount of HP needed to activate both the LP and EP systems. In fluid whey at 4°C, 10 mg of HP/kg provided the most bleaching in both LP and EP systems, whereas 15 mg of HP/kg was the most efficient in retentates for both enzymatic systems (results not shown). The optimum level of HP to activate the LP system in fluid whey at 4°C was very narrow (±10 mg/kg), whereas those with EP added displayed a much wider range of HP addition for maximum bleaching activity (Figure 1). The same trend was observed for fluid whey at 20°C and 35°C; the range for HP addition was wider for EP than LP alone (results not shown). The optimum level of HP for retentate bleaching by LP or EP was 15 mg/kg. In retentate, both EP and LP alone exhibited effective bleaching (>80% norbixin destruction) over a wide range of HP addition (15–250 mg/kg) at either cold (4°C) or warm (35°C) temperatures (results not shown).

In fluid whey, bleaching with EP at 2 DBLU of MB with the addition of 10 mg of HP/kg for 1 h at 4°C yielded >80% bleaching. Bleaching using only endogenous enzyme (LP) under the same conditions was slower (P < 0.05), requiring 6 to 24 h to bleach >80% (results not shown) and variability in LP activity and optimum bleach times were observed. In fluid retentate, optimum bleach times at warm temperatures (35°C) for LP and EP were not different (5 min, >80% norbixin destruction; P > 0.05; results not shown), nor were they distinct at 4°C (30 min, >80% norbixin destruction; P > 0.05; results not shown).

**Descriptive Sensory Analysis.** Bleached fluid wheys, regardless of bleaching treatment, had decreased sweet aromatic and cooked/milky flavors (P < 0.05; results not shown). Fluid wheys bleached using HP displayed a distinct sulfur flavor not present in enzymatically (LP or EP) bleached wheys. In retentates, similar to fluid whey, all bleached samples, regardless of bleaching treatment, exhibited decreased sweet aromatic and cooked/milky flavors (P < 0.05). In addition, bleached retentates exhibited increased cardboard flavor intensities compared with control unbleached retentates (P < 0.05). Similar to bleaching in fluid whey, HP bleached retentate exhibited a distinct sulfur note not detected in enzymatically (either HP or LP) bleached retentates.

**GC-MS.** Volatile compound analysis was consistent with descriptive analysis results. In fluid whey, enzymatically bleached wheys (EP or LP) were higher in aldehydes than chemically bleached wheys (HP 250 mg/kg; results not shown). Bleached retentates, regardless of treatment, were higher in aldehydes, including hexanal, heptanal, octanal, nonanal, and decanal, compared with unbleached controls (results not shown). In addition, retentates bleached chemically (250 mg of HP/kg) were higher in dimethyl trisulfide than other treatments (P < 0.05; 0.64 ± 0.05 vs. 0.45 ± 0.02 μg/kg).
Composition. Compositionally, WPC80 were not different ($P > 0.05$). Powdered WPC80 averaged 96.3 ± 0.48% solids, 77.6 ± 2.1% protein, and 3.7 ± 0.4% fat on a wet-weight basis. Phosphorus, potassium, calcium, magnesium, sulfur, and sodium were not different among treatments ($P > 0.05$). Iron was the only distinct mineral; the LP WPC80 was lower in iron than the control WPC80 ($P < 0.05$; Table 1) and this observation has been previously documented in WPC80 from whey chemically bleached with 500 mg of HP/kg (Jervis et al., 2012; Jervis and Drake, 2013).

Norbixin and Color Analysis. Consistent with and as expected from fluid whey results, enzymatic bleaching (LP and EP) removed more norbixin than traditional chemical bleaching (HP; $P < 0.05$; Figure 2). The addition of EP increased the speed of the bleaching process, bleaching 92% in 1 h, whereas LP and HP bleached 97 and 38%, respectively, in 12 h at 4°C. The $L^*a^*b^*$ values were consistent with norbixin values (data not shown).

Descriptive Sensory Analysis. Rehydrated WPC80 that were bleached, regardless of treatment, were characterized by a lack of sweet aromatic and buttery flavors and by increased cardboard flavor compared with the unbleached control (Table 2). Hydrogen peroxide-bleached WPC80 had a distinct oxidized/fatty flavor not detected in the other WPC80. Enzymatically bleached WPC80 with EP displayed a potato/brothy flavor but was also lower in cardboard flavor than enzymatically bleached WPC80 using LP alone (Table 2).

GC-MS. Volatile analysis results were consistent with descriptive analysis results. All 4 WPC80 were distinct in their volatile profiles. Bleached WPC80, regardless of treatment, were higher in 1-octen-3-ol and 1-octen-3-one ($P < 0.05$; Table 3). Among enzymatically bleached samples, LP WPC80 were higher in 2,3-octadienone, 2-pentyl furan, and hexanal than those with added EP ($P < 0.05$; Table 3). Heptanal was higher in EP-bleached WPC80 than the control.

Table 1. Select minerals in powdered whey protein concentrate with 80% protein (WPC80) with different bleaching treatments

<table>
<thead>
<tr>
<th>Bleaching treatment</th>
<th>P (% weight)</th>
<th>K (% weight)</th>
<th>Ca (% weight)</th>
<th>Mg (% weight)</th>
<th>S (% weight)</th>
<th>Fe (mg/kg)</th>
<th>Na (mg/kg)</th>
<th>Ash (% weight)</th>
</tr>
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<tbody>
<tr>
<td>Control</td>
<td>0.37&lt;sup&gt;a&lt;/sup&gt;</td>
<td>0.71&lt;sup&gt;a&lt;/sup&gt;</td>
<td>0.52&lt;sup&gt;a&lt;/sup&gt;</td>
<td>0.06&lt;sup&gt;a&lt;/sup&gt;</td>
<td>0.99&lt;sup&gt;a&lt;/sup&gt;</td>
<td>9.1&lt;sup&gt;a&lt;/sup&gt;</td>
<td>2,100&lt;sup&gt;a&lt;/sup&gt;</td>
<td>3.3&lt;sup&gt;a&lt;/sup&gt;</td>
</tr>
<tr>
<td>HP</td>
<td>0.37&lt;sup&gt;a&lt;/sup&gt;</td>
<td>0.66&lt;sup&gt;a&lt;/sup&gt;</td>
<td>0.55&lt;sup&gt;a&lt;/sup&gt;</td>
<td>0.06&lt;sup&gt;a&lt;/sup&gt;</td>
<td>0.98&lt;sup&gt;a&lt;/sup&gt;</td>
<td>7.7&lt;sup&gt;a&lt;/sup&gt;</td>
<td>1,900&lt;sup&gt;a&lt;/sup&gt;</td>
<td>3.2&lt;sup&gt;a&lt;/sup&gt;</td>
</tr>
<tr>
<td>LP</td>
<td>0.37&lt;sup&gt;a&lt;/sup&gt;</td>
<td>0.72&lt;sup&gt;a&lt;/sup&gt;</td>
<td>0.53&lt;sup&gt;a&lt;/sup&gt;</td>
<td>0.06&lt;sup&gt;a&lt;/sup&gt;</td>
<td>0.94&lt;sup&gt;a&lt;/sup&gt;</td>
<td>7.4&lt;sup&gt;a&lt;/sup&gt;</td>
<td>2,200&lt;sup&gt;a&lt;/sup&gt;</td>
<td>3.5&lt;sup&gt;a&lt;/sup&gt;</td>
</tr>
<tr>
<td>EP</td>
<td>0.38&lt;sup&gt;a&lt;/sup&gt;</td>
<td>0.67&lt;sup&gt;a&lt;/sup&gt;</td>
<td>0.56&lt;sup&gt;a&lt;/sup&gt;</td>
<td>0.06&lt;sup&gt;a&lt;/sup&gt;</td>
<td>0.96&lt;sup&gt;a&lt;/sup&gt;</td>
<td>8.0&lt;sup&gt;a&lt;/sup&gt;</td>
<td>2,000&lt;sup&gt;a&lt;/sup&gt;</td>
<td>3.4&lt;sup&gt;a&lt;/sup&gt;</td>
</tr>
</tbody>
</table>

<sup>1</sup>Means within a column not sharing a common superscript are different ($P < 0.05$).

<sup>2</sup>Control = no bleach; HP = 250 mg of hydrogen peroxide/kg; LP = lactoperoxidase; EP = exogenous peroxidase (MaxiBright, DSM, Delft, the Netherlands).
whereas dimethyl disulfide was higher in LP-bleached WPC80 than the control ($P < 0.05$; Table 3). Hexanal, a key volatile indicative of lipid oxidation, was highest in HP- and LP-treated WPC80.

**DISCUSSION**

Levels of HP needed to activate enzymatic systems at cold temperatures were in the range of values previously reported for other temperatures (Bottomley et al., 1989; Campbell et al., 2012). In agreement with previous research, chemical bleaching using HP (250 mg/kg) at cold temperatures was not very effective in fluid whey but was more effective in retentate (Listiyani et al., 2012; Fox, 2013). At 4°C, enzymatic bleaching of fluid whey was more effective than traditional chemical bleach with HP. Furthermore, the addition of

**Table 2.** Descriptive sensory profiles of whey protein concentrate with 80% protein (WPC80)

<table>
<thead>
<tr>
<th>Sensory attribute</th>
<th>Control</th>
<th>HP</th>
<th>LP</th>
<th>EP</th>
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</thead>
<tbody>
<tr>
<td>Aroma intensity</td>
<td>2.1b</td>
<td>2.5a</td>
<td>2.3a</td>
<td>2.4a</td>
</tr>
<tr>
<td>Sweet aromatic</td>
<td>1.3a</td>
<td>ND</td>
<td>ND</td>
<td>ND</td>
</tr>
<tr>
<td>Cooked/milky</td>
<td>2.4a</td>
<td>2.4a</td>
<td>2.3a</td>
<td>2.3a</td>
</tr>
<tr>
<td>Butter</td>
<td>0.6a</td>
<td>ND</td>
<td>ND</td>
<td>ND</td>
</tr>
<tr>
<td>Cardboard</td>
<td>1.2a</td>
<td>2.5a</td>
<td>2.3a</td>
<td>1.9b</td>
</tr>
<tr>
<td>Potato/brothy</td>
<td>ND</td>
<td>ND</td>
<td>ND</td>
<td>1.1a</td>
</tr>
<tr>
<td>Oxidized/fatty</td>
<td>ND</td>
<td>1.2a</td>
<td>ND</td>
<td>ND</td>
</tr>
<tr>
<td>Astringent</td>
<td>1.3b</td>
<td>1.6a</td>
<td>1.4a</td>
<td>1.4a</td>
</tr>
</tbody>
</table>

a–c Means within a row not sharing a common superscript are different ($P < 0.05$).

1 Attribute intensities were scored on a 0- to 15-point universal intensity scale (Meilgaard et al., 2007). Most dried ingredient flavors fall between 0 and 4 (Croissant et al., 2009; Listiyani et al., 2011).

2 Control = no bleach; HP = 250 mg of hydrogen peroxide/kg; LP = lactoperoxidase; EP = exogenous peroxidase (MaxiBright, DSM, Delft, the Netherlands).

3 ND = not detected.
EP in fluid whey increased the speed of bleaching at 4°C. In an industrial setting, dosing the correct amount of HP for enzymatic bleaching into a continuous fluid whey system can be difficult to do precisely. Seasonal variations and processing deviations may also occur that alter the amount of HP needed to activate the LP system, further complicating matters. As such, it may be beneficial in industrial settings to add EP to increase the speed of bleaching as well as robustness of the bleaching system.

In retentate, the addition of EP did not increase the speed of enzymatic bleaching ($P > 0.05$) but in fluid whey, the addition of EP greatly increased the speed of bleaching ($P < 0.05$; EP: 1 h, LP: 12 h). The decreased effect of the exogenous enzyme in retentate is likely because native LP is concentrated along with protein during UF and is present at much higher levels in retentate than in fluid whey. As such, enzymatic activity is increased and the speed of bleaching in retentate increases compared with fluid whey. Similar to fluid whey, the LP system in retentate can be permanently inactivated if too much HP is dosed into the system. Fox (2013) demonstrated that in liquid whey protein retentate, 250 mg of HP/kg destroyed more norbixin than 500 mg of HP/kg, suggesting that the LP range for HP is much higher in retentate (up to 250 mg/kg) than in fluid whey (up to 20 mg/kg), which is also consistent with results from the current study. Increases in enzyme levels, whether from membrane filtration or the addition of EP, increase the range at which HP can be dosed without permanently inactivating LP.

As expected, bleached fluid wheys and retentates, regardless of bleaching treatment, had decreased sweet aromatic and cooked/milky flavors and increased cardboard flavor ($P < 0.05$). Increased cardboard flavor as a result of lipid oxidation in fluid whey, 34 and 80% protein retentate, and 34 and 80% protein spray-dried powders have been extensively documented (Croissant et al., 2009; Campbell et al., 2011, 2012; Listiyani et al., 2011, 2012; Jervis et al., 2012; Kang et al., 2012). Fluid wheys and retentates that were chemically bleached with HP contained a distinct sulfur flavor not present in enzymatically (LP or MB) bleached wheys ($P < 0.05$; results not shown). Increased volatile sulfur compounds and distinct sulfur flavor in chemically HP bleached WPC80 have been previously reported (Jervis et al., 2012) and higher concentrations of dimethyl trisulfide were documented in fluid retentates that were bleached with HP in the current study. Previous studies have also demonstrated functional differences between HP-bleached whey protein and unbleached controls, suggesting that HP affects protein integrity (Jervis et al., 2012; Campbell et al., 2013). Volatile sulfur compounds are formed from protein degradation of sulfur-containing amino acids (Wright et al., 2006). Although it does not directly influence cardboard flavor, hexanal is considered a good indicator of lipid oxidation and cardboard flavors (Whitson et al., 2010). Hexanal relative abundance was higher in LP- and HP-bleached samples than unbleached or EP-bleached WPC80. Higher relative abundance of this lipid oxidation compound is consistent with sensory results that

### Table 3. Relative abundance (μg/kg) of select volatile compounds in rehydrated spray-dried whey protein concentrate with 80% protein (WPC80) with different bleaching treatments

<table>
<thead>
<tr>
<th>Volatile compound</th>
<th>Control</th>
<th>HP</th>
<th>LP</th>
<th>EP</th>
</tr>
</thead>
<tbody>
<tr>
<td>1-Hexen-3-one</td>
<td>0.367b</td>
<td>1.62a</td>
<td>0.783b</td>
<td>0.431b</td>
</tr>
<tr>
<td>1-Octen-3-ol</td>
<td>ND³</td>
<td>2.96a</td>
<td>3.83⁴</td>
<td>2.95⁵</td>
</tr>
<tr>
<td>1-Octen-3-one</td>
<td>ND⁶</td>
<td>2.95⁶</td>
<td>3.01⁶</td>
<td>1.97⁶</td>
</tr>
<tr>
<td>2,3-Octadienone</td>
<td>0.21³</td>
<td>0.632b</td>
<td>1.507⁷</td>
<td>0.552⁷</td>
</tr>
<tr>
<td>2-Methyl butanal</td>
<td>0.199⁸</td>
<td>0.206⁹</td>
<td>0.211¹</td>
<td>0.231⁸</td>
</tr>
<tr>
<td>2-Pentyl furan</td>
<td>1.467b</td>
<td>3.569a</td>
<td>9.385⁷</td>
<td>3.373⁷</td>
</tr>
<tr>
<td>3-Methyl butanal</td>
<td>0.044³</td>
<td>0.059⁴</td>
<td>0.079⁴</td>
<td>0.061⁴</td>
</tr>
<tr>
<td>Decanal</td>
<td>0.099⁵</td>
<td>0.115⁶</td>
<td>0.128⁶</td>
<td>0.083⁶</td>
</tr>
<tr>
<td>Diacetyl</td>
<td>0.285⁷</td>
<td>0.249⁷</td>
<td>0.253⁷</td>
<td>0.240⁷</td>
</tr>
<tr>
<td>Dimethyl disulfide</td>
<td>0.17⁴</td>
<td>0.705b</td>
<td>1.77²</td>
<td>0.650b</td>
</tr>
<tr>
<td>Dimethyl sulfide</td>
<td>0.64³</td>
<td>0.484⁴</td>
<td>0.498⁴</td>
<td>0.53⁴</td>
</tr>
<tr>
<td>Heptanal</td>
<td>0.052⁶</td>
<td>0.194⁶</td>
<td>0.456⁶</td>
<td>0.63⁴</td>
</tr>
<tr>
<td>Hexanal</td>
<td>1.59⁷</td>
<td>7.69⁷</td>
<td>5.98⁷</td>
<td>3.54b</td>
</tr>
<tr>
<td>Nonanal</td>
<td>0.53³</td>
<td>0.569⁴</td>
<td>0.49²</td>
<td>0.33³</td>
</tr>
<tr>
<td>Octanal</td>
<td>0.55³</td>
<td>0.004⁴</td>
<td>0.365⁴</td>
<td>0.45³</td>
</tr>
<tr>
<td>p-Xylene</td>
<td>0.29⁶</td>
<td>0.401⁴</td>
<td>0.187⁴</td>
<td>0.18⁷</td>
</tr>
<tr>
<td>Toluene</td>
<td>2.91⁴</td>
<td>2.65⁴</td>
<td>2.99⁴</td>
<td>2.4⁹</td>
</tr>
</tbody>
</table>

³ᵃ Means within a column not sharing a common superscript are different ($P < 0.05$).
¹ Control = no bleach; HP = 250 mg of hydrogen peroxide/kg; LP = lactoperoxidase; EP = exogenous peroxidase (2 dairy bleaching units of MaxiBright).
³⁲ ND = not detected.
LP- and HP-bleached WPC80 were higher in cardboard flavor than unbleached or EP-bleached WPC80.

The WPC80 from fluid whey bleached using EP exhibited a distinct potato/brothy flavor. Potato flavor can be caused by a wide array of volatile compounds, but is mainly attributed to methional, an aroma compound formed from the degradation of the amino acid methionine (Jansky, 2010). Methional was not detected by headspace volatile compound analysis in EP WPC80, even though a distinct potato flavor was detected by trained sensory panelists. The threshold for methional is very low (less than 5 μg/kg; Karagül-Yüceer et al., 2004). It is possible that methional was below instrumental headspace detection but still readily detected by trained panelists. It is also possible that another compound is responsible for the potato flavor in EP WPC80 documented by trained panelists. Methional has been detected previously by gas chromatography-olfactometry and by solvent extraction with GC-MS, but not by SPME GC-MS in dried chromatography-olfactometry and by solvent extraction with GC-MS, but not by SPME GC-MS in dried.

Chromatography-olfactometry and by solvent extraction with GC-MS, but not by SPME GC-MS in dried.

It is imperative that dried dairy ingredients be colorless and bland in flavor to increase ingredient applications. Bleaching is required to eliminate color, but also produces undesirable flavors that can carry through into the finished product and influence consumer acceptance. Alternative bleaching agents, such as EP, can bleach effectively and can eliminate more color than traditional chemical bleaching agents in less time. These results demonstrate that off-flavors due to lipid oxidation are still present in enzymatically bleached WPC80 but are the same or lower in intensity than HP-chemically bleached WPC80 and lower in lipid oxidation volatiles that HP-chemically bleached WPC80. By bleaching at cold temperatures, manufacturers can minimize off-flavors and decrease membrane fouling. The addition of exogenous enzyme increased the speed of bleaching at 4°C in fluid whey and also the range of HP allowed for enzymatic bleaching in fluid whey. As such, EP may be beneficial in an industrial setting where continuous dosing of a narrow range of HP is difficult.

CONCLUSIONS

It is imperative that dried dairy ingredients be colorless and bland in flavor to increase ingredient applications. Bleaching is required to eliminate color, but also produces undesirable flavors that can carry through into the finished product and influence consumer acceptance. Alternative bleaching agents, such as EP, can bleach effectively and can eliminate more color than traditional chemical bleaching agents in less time. These results demonstrate that off-flavors due to lipid oxidation are still present in enzymatically bleached WPC80 but are the same or lower in intensity than HP-chemically bleached WPC80 and lower in lipid oxidation volatiles that HP-chemically bleached WPC80. By bleaching at cold temperatures, manufacturers can minimize off-flavors and decrease membrane fouling. The addition of exogenous enzyme increased the speed of bleaching at 4°C in fluid whey and also the range of HP allowed for enzymatic bleaching in fluid whey. As such, EP may be beneficial in an industrial setting where continuous dosing of a narrow range of HP is difficult.

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REFERENCES

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