

Sunflower meal and spring pea ruminal degradation protection using malic acid or orthophosphoric acid-heat treatments

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Abstract. The effects of solutions of malic or orthophosphoric acids (0.752 Eqg/kg of feed) and heat to protect proteins of sunflower meal (SFM) and spring pea (SP) against ruminal degradation were studied using particle transit, ¹⁵N infusion, *in situ* and electrophoretic techniques. Three wethers fitted with rumen and duodenum cannulae were successively fed three isoproteic diets including SFM and SP, untreated or treated with malic or orthophosphoric acids. Incubations of tested meals were only performed while feeding the respective diet. Estimates of the ruminally undegraded fraction (RU) and its intestinal digestibility of dry matter, organic matter (only for RU), crude protein and starch (only in SP) were obtained considering ruminal microbial contamination and particle comminution and outflow rates. When corrected for microbial contamination, estimates of RU and intestinal digestibility decreased in all tested fractions for both feeds. All RU estimates increased with the protective treatments, whereas intestinal digestibility-dry matter also increased in SFM. Low intestinal digestibility-crude protein values suggested the presence of antitrypsin factors in SP. Protective treatments of both feeds led to consistent increases in the intestinal digested fraction of dry matter and crude protein, being only numerically different for SP-starch (60.5% as average). However, treatments also reduced the organic matter fermentation, which may decrease ruminal microbial protein synthesis. Electrophoretic studies showed albumin disappearance in both SFM and SP, whereas changes in other RU proteins were more pronounced in SP than SFM.

Additional keywords: heat, malic and orthophosphoric acids, protein protection.

Received 2 July 2014, accepted 8 June 2015, published online 27 July 2015

Introduction

Applying protective treatments against ruminal fermentation in high quality proteins is attractive to avoid their microbial degradation, which is usually associated with high ruminal ammonia losses and also with reduced efficiency of microbial protein synthesis (NRC 2001). The inefficiency associated with excessive ruminal protein degradation is important in productive ruminants, whose large amino acid requirements should be supplied mainly by protein concentrates. In particular, this inefficiency is large for concentrates composed of highly degradable proteins such as sunflower or spring pea meals (SFM and SP meals, respectively). In addition, amino acids from these protein concentrates are transformed into compounds such as nucleic bases or amino sugars that are not used for protein synthesis in animal tissues. In SFM, Arroyo *et al.* (2011, 2013) reported a large protection of crude protein (CP) by treating it with malic or orthophosphoric acid solutions and heat. These treatments increased bypass CP by 4.25 times, but the undigested fraction was increased as well, especially when using orthophosphoric acid. This protective effect was suspected to be the result of the high temperature applied (150°C during 6 h),

combined with the high moisture from the acid solution (400 mL/kg). The objective of this study was to determine the effects of these treatments applied at lower temperature and moisture on rumen degradation and intestinal digestion of SFM and SP.

Materials and methods

Meal treatments

Samples of SFM and SP were treated with 4 N solutions (200 mL/kg) of malic acid (268.2 g/L) or orthophosphoric acid (130.6 g/L). Acid doses were equivalent to those (2 N solutions sprayed at 400 mL/kg) used by Arroyo *et al.* (2013). Two fractions of 12.5 kg of each meal were successively sprayed with each acid solution in a concrete mixer using a sprayer. Both fractions were then mixed and allowed to rest for 1 h at room temperature. The blend was then dried in a forced air oven at 120°C for 1 h. Then the oven was turned off and the treated material was left in the oven overnight. During the drying process, the material was stirred every 30 min during the first 2 h and then every 60 min for the subsequent 5 h.

Diet and animals

Diets were isoproteic and included 45% oat (*Avena sativa*) hay and 55% concentrate (fresh weight). The concentrate contained corn grain (30%), barley grain (30%), SFM (15%), SP (22%), and minerals and vitamin (3%). The control diet (UT) included untreated SFM and SP, which were replaced by equivalent amounts of meals treated with malic or orthophosphoric acids in diets including treated meals (MT and PT, respectively). The CP supplied from SFM and SP represented ~55% of total CP in all concentrates. Chemical composition of oat hay, SFM and SP meals and concentrates is shown in Table 1. Ranges of CP, neutral (NDF) and acid (ADF) detergent fibres of diets were (g/kg of DM): 177–181, 323–326 and 150–154, respectively. The DM intake was fixed at 45 g/kg bodyweight^{0.75} and diets were distributed in six equal meals (every 4 h), starting at 0900 hours. The diets UT, PT and MT were offered successively in three experimental periods to three wethers (bodyweight = 57.4 ± 5.29 kg) fitted with rumen cannulae (inside diameter 60 mm) and T-type cannulae (inside diameter 12 mm) in the proximal duodenum. Wethers were pen-housed individually and handled according to animal care principles published in the Spanish Royal Decree 1201/2005 (BOE 2005).

Experimental procedures

Ruminal digestion

The experimental period of 21 days included successively: 10 days of diet adaptation, SFM and SP particle transit study (Days 11–14) and ruminal nylon-bag incubations (Days 15–21). The rumen was emptied at 0900 hours at the end of *in situ* incubations in each period to isolate solid adherent bacteria (SAB) as described by Rodríguez *et al.* (2000). Rumen bacteria were labelled from Days 11 to 21 by continuous infusion (250 mL/day) of a (¹⁵NH₄)₂SO₄ solution (98 atoms% enriched) that provided 25 mg ¹⁵N/day.wether.

The study of particle transit in the rumen-reticulum was performed by pulse dosing each animal before the first morning meal (i.e. at 0900 hours). Samples of 50 g of untreated SFM and SP were labelled with europium (Eu) and ytterbium (Yb), respectively, and were consumed completely in 20 min. These meals had previously been washed in an automatic washer to eliminate soluble components, and marked by immersion in

solutions of Eu₂Cl₃ or YbCl₃ (10 mg of Eu or Yb/g of feed) as described by González *et al.* (1998). A total of 22 samples were obtained through the duodenal cannulae, the first sample before supplying the marker and the remaining samples between 1.5 and 96 h post-marker dosing. Samples were oven-dried at 105°C for 48 h, milled to pass a 1-mm screen, and analysed for Eu and Yb. The pattern of Eu or Yb concentrations in the duodenal digesta over time was described for each animal by fitting to the model of Dhanoa *et al.* (1985). Primary and secondary rate constants used in this model were assumed as the rates of outflow (k_p) and of comminution and mixing (k_c) of particles in the reticulo-rumen, respectively, in agreement with Ellis *et al.* (1979) and González *et al.* (2006).

Ruminal incubations were performed using nylon bags (11 by 7-cm internal dimensions) with 46-µm pore size and filled with ~3 g (air-dry basis) of sample (ground to pass a 2-mm screen). Two series of ruminal incubations with 14 bags containing SFM (7) or SP (7) meals were carried out in each diet at 2, 4, 8, 16, 24, 48 and 72 h. The meals incubated in each experimental period were those corresponding to the associated diet. In each incubation, all bags were placed simultaneously in the rumen just before the wethers were offered their first morning meal. Once collected from the rumen, bags were washed with tap water and stored at –20°C. After thawing, bags were washed three times for 5 min in a turbine washing machine (Jata 580; JATA, Abadiano, Bizkaia, Spain). The same washing procedure was applied for two series of two bags for each meal to obtain the 0-h value. Bags were stored at –20°C once again, freeze-dried and immediately weighed to establish degradation kinetics of dry matter (DM) with the exponential model of Ørskov and McDonald (1979):

$$d = a + b (1 - e^{-k_d t})$$

In this model, the constants a and b represent, respectively, the soluble fraction, (which was assumed as the 0-h value) and the nonsoluble but degradable component, which disappears at a constant fractional rate, k_d , per unit time. The undegradable fraction (r) was estimated as $1 - (a + b)$.

Values of ruminally undegraded (RU) DM, organic matter (OM), CP and starch (only in SP) non-corrected or corrected for microbial contamination were determined considering degradation kinetics and both k_p and k_c transit rates with the method proposed by Arroyo and González (2013). This method is

Table 1. Chemical composition (g/kg of dry matter) of untreated (UT), malic acid treated (MT) and orthophosphoric acid treated (PT) of sunflower and spring pea meals, oat hay and used diets

OM, organic matter; NDF, neutral detergent fibre; ADF, acid detergent fibre; CP, crude protein; NDIN and ADIN: insoluble nitrogen in neutral and acid detergent solutions, respectively

Item	Sunflower meal			Spring pea			Hay	Diets		
	UT	MT	PT	UT	MT	PT		UD	MD	PD
OM	932	923	889	963	969	933	840	885	880	881
Starch	–	–	–	368	367	370	–	–	–	–
NDF	309	311	261	201	245	224	459	323	326	324
ADF	177	184	159	85.3	94.5	103	236	154	150	151
Lignin	28.2	44.7	32.5	2.44	1.79	2.38	46.6	26.7	26.9	25.4
CP	344	322	346	221	210	212	189	181	177	178
NDIN (g/kg N)	137	151	103	125	180	150	358	–	–	–
ADIN (g/kg N)	41.9	47.3	38.2	18.8	21.9	23.0	54.2	–	–	–

based on generating composite samples (CS) representative of the chemical composition of the RU feed. For this purpose, the residues obtained at each incubation time were pooled in equal quantities for each animal. The resultant residues for 0, 2, 4, 8, 16, 24, 48 and 72 h of incubation were considered representative of the composition of the rumen outflow of undegraded feed in the intervals 0–1, 1–3, 3–6, 6–12, 12–20, 20–36, 36–60 and 60–84 h, respectively. The proportions in which the different residues were mixed were calculated by the ratio of the flow in each interval and the total flow determined using the feed flow-function proposed by Arroyo and González (2013). The CS were analysed for OM, CP, $^{15}\text{N}/\text{total N}$ and starch (only in SP). The respective RU values (as %) were determined from the concentrations of the tested fraction in the composite samples (Y) and in the whole feed (X) and the RU-DM values, as follows:

$$\text{RU} - \text{MO}, -\text{CP} \text{ or } -\text{starch} = \text{Y} \times \text{RU} - \text{DM}/\text{X}$$

The microbial proportions of N and DM in CS were determined as indicated by Rodríguez and González (2006) using SAB samples as reference; SAB isolates were lyophilised and analysed for DM, OM, N, $^{15}\text{N}/\text{N}$ and total glucose. Microbial proportions of OM and glucose were determined as the microbial DM content of CS samples \times the concentration of these fractions in SAB expressed on DM.

Intestinal digestion

Twenty days after the end of the ruminal studies, wethers were fed again the untreated diet to determine ID-DM, -CP and -starch of untreated and treated meals. Eight subsamples of ~200 mg of each CS were put into round-shaped (approximate diameter 2.5 cm) mobile nylon bags. These bags were inserted randomly through the duodenal cannulae of the respective wether at a rate of one bag every 15 min for 2 h for a total of 8 bags/day.wether, and recovered from the faeces. Bags were then processed by the same methods as those for rumen incubations; bags were then dried at 80°C for 48 h and weighed. The ID-DM was calculated as DM disappearance from the bag during intestinal incubation. Undigested residues obtained in each wether were pooled and analysed for CP, $^{15}\text{N}/\text{N}$ and starch (only in SP). The ID for any fraction was determined as the proportions from the ID-DM value and concentration of this fraction in the CS (Y) and in the intestinal incubated residues (Z):

$$\text{ID} - \text{CP} \text{ or } -\text{starch} = 1 - [\text{Z} \times (1 - \text{ID} - \text{DM})/\text{Y}]$$

The $^{15}\text{N}/\text{N}$ ratio was used to correct for the residual contamination due to adherent rumen microorganisms as previously indicated.

Protein electrophoresis

Protein in untreated and treated meals and their 0 h, composite and intestinally undigested samples were extracted and subjected to electrophoresis to determine the digestion site of the different protein fractions.

Chemical and protein electrophoresis analyses

Feed samples were analysed in triplicate using AOAC (2000) procedures for DM (procedure 934.01), ash (procedure 967.05) and CP (6.25 \times Dumas N; procedure 968.06), NDF (Van Soest

et al. 1991) and sequential ADF and acid detergent lignin (Robertson and Van Soest 1981). Analyses of NDF were performed with α -amylase and without sodium sulfite. NDF and ADF were expressed including the residual ash. The insoluble N in neutral and acid detergent solutions was determined by N analysis of the NDF and ADF residues, respectively. Ruminal or intestinal incubated residues were also analysed for N with the Dumas method. Samples of duodenal contents collected for transit studies were analysed by atomic absorption (Yb) or emission (Eu) spectrometry, as described by González *et al.* (1998). N isotopic proportions were performed in an elementary analyser (Flash 1112, Thermo, Bremen, Germany) coupled in continuous flow to an isotope ratio mass spectrometer (Delta V, Thermo).

Homogenate samples of the different protein fractions of SFM and SP were extracted using different solutions: (1) TRIS-HCl pH 6.8 1 M, 10% glycerol, 2.5% SDS and 5% β -mercaptoethanol to obtain total protein (2) 0.4 M NaCl + 3% β -mercaptoethanol to obtain globulins and (3) water to obtain albumins. Solutions were used at 750 μL per mg of N. The extracts were agitated in a vortex and left overnight with gentle stirring at 4°C. After centrifugation (12 500g for 45 min at 4°C) and pellet removal, the supernatant was mixed with a reducing buffer (1:1, v/v) and boiled for 10 min in a water bath. Electrophoresis was carried out using polyacrylamide slab gels [sodium dodecyl sulfate–polyacrylamide gel electrophoresis (SDS-PAGE); 160 \times 155 \times 1 mm] following the Laemmli (1970) protocol modified by Sánchez-Yélamo (in Vázquez *et al.* 2000). For total proteins and globulins, 10 μL per sample were applied, whereas for albumins, 20, 30 and 40 μL per well were tested because fewer proteins were detected in the gels. In all cases the electrode buffer was cold Tris-glycine (pH 8.3) and bromophenol blue was used as front-dye marker. The molecular weights (MW) of the protein samples were determined using a standard protein marker (Invitrogen Mark 12) consisting of myosin (MW 200 KD), β -galactosidase (MW 116.3 KD), phosphorylase b (MW 97.4 KD), serum albumin (MW 66.3 KD), glutamic dehydrogenase (MW 55.4 KD), lactate dehydrogenase (MW 36.5 KD), carbonic anhydrase (MW 31.0 KD), trypsin inhibitor (MW 21.5 KD), lysozyme (MW 14.4 KD) and aprotinin (MW 6.0 KD). After electrophoresis (~4.5 h), gels were stained overnight at room temperature with Coomassie Brilliant blue R-250 followed by de-staining twice in acetic acid: 2-propanol 10% (v/v) for 4 h. At least three repetitions were made to check for the reproducibility of the electrophoretic patterns. In each band pattern, bands were numbered in ascending order from cathode to anode starting with band '1'. The identity of each band was estimated by its relative mobility on the gel, and the molecular masses were estimated using a calibration curve developed with the marker standard polypeptides. Several specific proteins were identified by comparison with data from the literature.

Statistical methods

Degradation and transit kinetics were fitted by non-linear regression. Effects of protective treatments (*t*) on apparent parameters of DM degradation kinetics and RU-DM were

studied by variance analysis considering these factors and the wethers (w) in the model ($y_{ij} = \mu + t_i + w_j + \varepsilon_{ij}$). These same effects and those of the correction of microbial contamination taking place in the rumen (c) of RU, ID and the intestinal absorbable dietary fraction (IADF = RU \times ID) values were studied by variance analyses in a split-plot arrangement of treatments. In this design, the treatment was the whole-plot, tested against the wether \times treatment interaction as the error term, and microbial correction and its interaction with treatment were the subplot treatments ($y_{ijk} = \mu + t_i + w_j + c_k + t_i \times c_k + t_i \times w_j + \varepsilon_{ijk}$). These means were examined by orthogonal contrasts for treatment effects (untreated vs treated) and of acid used (malic vs orthophosphoric acids). This same design substituting microbial correction by the transit model considered (k_p alone compared with k_p and k_c) was also used to examine the differences in RU estimates of the insoluble but potentially degradable fraction of DM. As the animals were adult, maintained in a controlled environment, the period effect was assumed to be negligible in all variance analysis. Effects were declared significant at $P < 0.05$. All statistical analyses were performed using SAS software, version 8.0 (SAS 1999).

Results

Feed digestion

There were no differences among treatments or between the two meals (SFM, SP) in the k_p or k_c values: k_p : 6.36%/h versus 5.53%/h; s.e.m. = 0.295; $P = 0.227$; k_c : 77.7%/h versus 56.2%/h; s.e.m. = 7.05; $P = 0.153$. Therefore, mean values for each feed were used to calculate RU values in untreated and treated meals.

In treated SFM and SP, the soluble fraction of DM (a) decreased and the potentially degradable fraction (b) increased; however, the undegradable fraction (r) was increased except when SFM was treated with orthophosphoric acid (Table 2). In addition, both treatments led to reductions of k_d in SFM whereas reductions in k_d for SP were only numerical. As a result of these changes, treatments increased the apparent RU-DM corresponding to both the b fraction and total feed in both SFM and SP. There was no clear evidence of differences in the protective effect between both acids, especially for SP. The omission of k_c from the model resulted in mean values (\pm s.e.) of the RU-DM of the b fraction for UT, MT and PT meals of: 16.2 (\pm 2.14), 22.2 (\pm 1.19) and 22.1 (\pm 1.67) % in SFM and 18.4 (\pm 1.48), 26.5 (\pm 1.86) and 24.3 (\pm 1.80) % in SP, respectively. These values were higher ($P < 0.001$) than those generated when k_p and k_c were included in the calculations which are indicated in Table 2. Therefore, omitting k_c overestimated the apparent RU- b of DM of UT, MT and PT meals, respectively, by 16.8, 11.0 and 12.4% in SFM, and by 20.1, 15.7 and 18.2% in SP (data not shown).

The microbial contamination with DM or N in CS samples of both feeds showed large variability, and treatment effects were undetected. The large variability observed for the disappearance of DM biomass during the intestinal incubation in SP also prevented detecting effects, which on the contrary were shown for SFM (Table 3). Not correcting ruminal microbial contamination led to consistent overestimations in RU, ID and IADF values of all tested variables in both tested feeds

Table 2. Effects of protective treatments on apparent rumen degradation kinetics and ruminal undegraded fraction (RU) of dry matter of sunflower and spring pea meals (values are expressed as %)

	UT ^A	Meals		Probability of contrasts ^B		
		MT	PT	s.e.m.	C1	C2
<i>Sunflower meal</i>						
a^C	31.6	26.3	26.2	–	–	–
b	50.2	52.0	55.6	4.65	0.003	0.006
r	18.3	21.6	18.2	4.65	0.045	0.006
k_d (%/h)	14.0	8.64	9.94	1.13	0.027	0.464
RU	32.2	41.6	37.9	0.72	0.001	0.021
RU- b	13.9	20.0	19.7	1.03	0.009	0.844
<i>Spring pea</i>						
a	38.9	23.2	21.6	–	–	–
b	55.8	68.2	68.8	1.27	0.001	0.764
r	5.29	8.56	9.60	1.27	0.071	0.592
k_d (%/h)	11.7	8.95	10.5	0.92	0.153	0.311
RU	20.6	31.4	30.2	1.04	0.001	0.427
RU- b	15.3	22.9	20.6	1.38	0.019	0.300

^AUT, MT and PT: meals untreated or treated with malic or orthophosphoric acids, respectively.

^BC1: UT versus treated meals; C2: MT versus PT.

^C a , b , and r represent soluble, non-soluble degradable and undegradable fractions, respectively. k_d : fractional degradation rate of fraction b . RU: ruminally undegraded fraction. RU- b : RU corresponding to the b fraction, calculated using ruminal rates of particle comminution (k_c) and outflow (k_p) as: $RU = r + RU-b$ and $RU-b = bk_c k_p / [(k_d + k_p)(k_d + k_c)]$.

Table 3. Microbial contamination of ruminal undegraded composite samples of sunflower meal (SFM) and spring pea (SP)

	UT ^A	Meals		Contrast probability ^B		
		MT	PT	s.e.m.	C1	C2
<i>DM (g/kg residual DM)</i>						
SFM	39.4	30.5	46.3	6.99	0.91	0.185
SP	30.9	25	25.5	3.8	0.295	0.924
<i>N (g/kg residual N)</i>						
SFM	69.8	37.8	61.8	10.72	0.202	0.19
SP	91	53.2	56.2	13.46	0.093	0.882
<i>Intestinal disappearance of ruminal biomass (% of DM)</i>						
SFM	86.9	77.8	88.4	0.69	0.011	<0.001
SP	65.9	85.6	77.8	11.09	0.307	0.645

^AUT, MT and PT: meals untreated or treated with malic or orthophosphoric acids, respectively.

^BC1: UT versus treated meals; C2: MT versus PT.

(Tables 4, 5). As previously indicated for DM, protective treatments consistently increased RU in the remainder tested fractions in both feeds. Also, positive effects were shown in the ID-DM in SFM, but not for ID-CP in this meal or for the different SP-tested fractions. As a result, treatments also increased the IADF-DM and IADF-CP in both tested meals (Tables 4, 5). Despite the fact that treatments increased the IADF-starch by 60.5% on average, the results were not statistically significant, probably due to the high variability observed (Table 5). Regression analysis showed strong correlations between microbial-corrected values of both

Table 4. Effects of protective treatments and of correcting the microbial contamination taking place in the rumen on *in situ* estimates of ruminally undegraded fraction (RU), intestinal digestibility (ID) and intestinal absorbable dietary fraction (IADF: RU × ID) of sunflower meal

	Meals ^A						s.e.m.	Effects and contrasts ^B			
	UT		MT		PT			Treatments		Correction	
	NC ^C	C	NC	C	NC	C		C1	C2	s.e.m.	P
<i>RU</i>											
Dry matter	32.2	30.9	41.6	40.3	37.9	36.1	0.61	<0.001	0.010	0.17	<0.001
Organic matter	32.8	31.8	43.4	42.3	40.5	39.0	0.61	<0.001	0.022	0.14	<0.001
Crude protein	21.4	19.9	42.2	40.6	33.0	30.9	0.61	<0.001	<0.001	0.19	<0.001
<i>ID</i>											
Dry matter	30.0	27.7	38.1	35.8	40.1	37.4	2.07	0.024	0.573	0.38	0.004
Crude protein	80.1	75.0	83.4	79.5	85.4	78.0	2.92	0.320	0.952	0.87	0.004
<i>IADF</i>											
Dry matter	9.71	8.58	16.0	14.5	15.3	13.6	1.15	0.012	0.628	0.24	0.005
Crude protein	17.5	15.2	35.5	32.5	28.6	24.3	1.24	<0.001	0.013	0.51	0.004

^AUT, MT and PT: meals untreated or treated with malic or orthophosphoric acids, respectively.

^BC1: UT versus treated meals; C2: MT versus PT.

^CNC and C: no corrected and corrected by the ruminal microbial contamination.

Table 5. Effects of protective treatments and of correcting the microbial contamination taking place in the rumen on *in situ* estimates of ruminally undegraded fraction (RU), intestinal digestibility (ID) and intestinal absorbable dietary fraction (IADF: RU × ID) of spring pea

	Meals ^A						s.e.m.	Effects and contrasts ^B			
	UT		MT		PT			Treatments		Correction	
	NC ^C	C	NC	C	NC	C		C1	C2	s.e.m.	P
<i>RU</i>											
Dry matter	20.6	20.0	31.4	30.7	30.2	29.4	1.06	0.001	0.449	0.09	0.002
Organic matter	21.0	20.5	31.9	31.3	31.4	30.8	1.04	0.001	0.749	0.07	0.001
Crude protein	13.0	11.9	28.2	26.7	26.1	24.7	1.14	<0.001	0.270	0.16	0.001
Starch	15.4	15.4	29.2	29.1	25.9	25.8	2.01	0.008	0.300	0.007	0.003
<i>ID</i>											
Dry matter	32.1	30.8	32.9	31.6	35.9	34.7	1.90	0.373	0.313	0.22	0.006
Crude protein	35.2	24.8	37.5	28.1	38.6	29.8	2.00	0.222	0.659	1.68	0.007
Starch	64.7	64.2	54.5	53.6	62.7	62.1	6.63	0.484	0.427	0.12	0.007
<i>IADF</i>											
Dry matter	6.64	6.17	10.3	9.67	10.8	10.2	0.66	0.009	0.597	0.09	0.003
Crude protein	4.57	2.99	10.6	7.47	10.0	7.43	0.70	0.004	0.782	0.37	0.003
Starch	10.1	9.97	16.0	15.7	16.5	16.3	2.93	0.164	0.890	0.03	0.005

^AUT, MT and PT: meals untreated or treated with malic or orthophosphoric acids, respectively.

^BC1: UT versus treated meals; C2: MT versus PT.

^CNC and C: no corrected and corrected by the ruminal microbial contamination.

RU-DM and ID-DM in SFM ($ID-DM = -0.55 + 0.955 RU-DM$; $n = 9$; $R^2 = 0.654$; $P = 0.008$). Similar relationships were also shown between IADF and RU of all tested parameters in both feeds (Fig. 1). In SFM, malic acid showed higher protective efficiency than orthophosphoric acid for RU values as well as for IADF-CP (Table 4); this was not observed for SP (Table 5).

Protein electrophoresis

Gel electrophoresis of total proteins, globulins and albumin are shown in Fig. 2 (SFM) and Fig. 3 (SP). In both feeds, images of treated meals were less sharp than those observed in untreated meals, especially in those samples not subjected to washing processes before protein extraction and electrophoresis.

Profiles of SFM showed clear bands for polypeptides in an approximate range of MW of 56 to 6 KD, whereas those of SP fell within a range of 97.4–6 KD.

Albumin was not detected in 0-h samples of SFM nor in those of protected SP meals. In addition, they were not detected in both feeds, untreated or treated CS samples. Similarly, the gel of total proteins of CS samples showed lack of some polypeptides other than albumin in untreated or treated SFM (3, 13 and 14) and SP (1, 2, 4, 6, 12, 14 and 21). Furthermore, some polypeptides (10, 13 and 20) were not detected in the gel of globulins in untreated or treated CS samples of SP. Finally, polypeptides were not observed in intestinal-incubated residues of SFM or SP, with the exception of some diffuse shading in areas corresponding to polypeptides of low MW (<10 KD) identified as 15, 16 and 18.

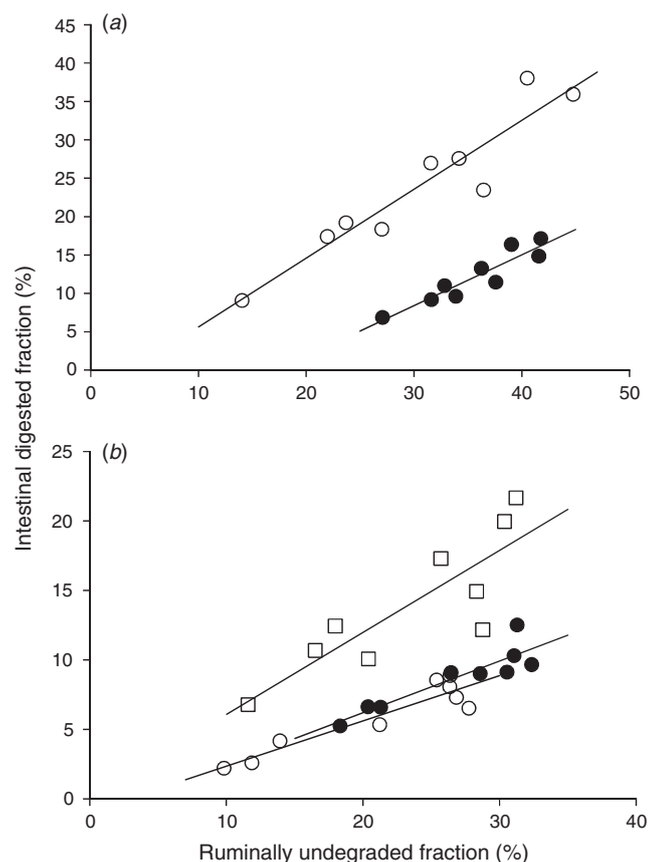


Fig. 1. Relationships between intestinal digested (IADF) and ruminally undegraded (RU) fractions of dry matter (●), crude protein (○) and starch (□) (a) in sunflower meal (SFM) and (b) spring pea (SP). Equations: SFM: DM: $IADF = -11.47 + 0.663 RU$; $R^2 = 0.861$; $P < 0.001$. CP: $IADF = -3.34 + 0.898 RU$; $R^2 = 0.861$; $P < 0.001$. SP: DM: $IADF = -1.23 + 0.371 RU$; $R^2 = 0.813$; $P < 0.001$; CP: $IADF = -0.90 + 0.323 RU$; $R^2 = 0.867$; $P < 0.001$; Starch: $IADF = 0.17 + 0.590 RU$; $R^2 = 0.861$; $P < 0.001$.

Discussion

The chemical composition of the SFM sample may be considered normal except for its low fibre fractions in relation to its CP content. By contrast, the SP sample showed a high fibre content.

Accuracy of *in situ* estimates

The omission of the k_c rate from the calculations of *in situ* degradation may have an effect on the *in situ* digestive estimates through its influence on the digestion site (ARC 1984). In relation to the total mean residence time in the rumen-reticulum, the time associated with the k_c rate was considerable (7.6% and 9.0% as mean in SFM and SP, respectively), supporting the consideration of this rate to improve accuracy of *in situ* estimates, as previously indicated (Arroyo and González 2013). The overestimation of the feed bypass caused by the lack of correction of the microbial contamination of rumen-incubated residues may also lead to overestimations of ID due to the large intestinal digestion of these adherent microorganisms. Both overestimations are directly

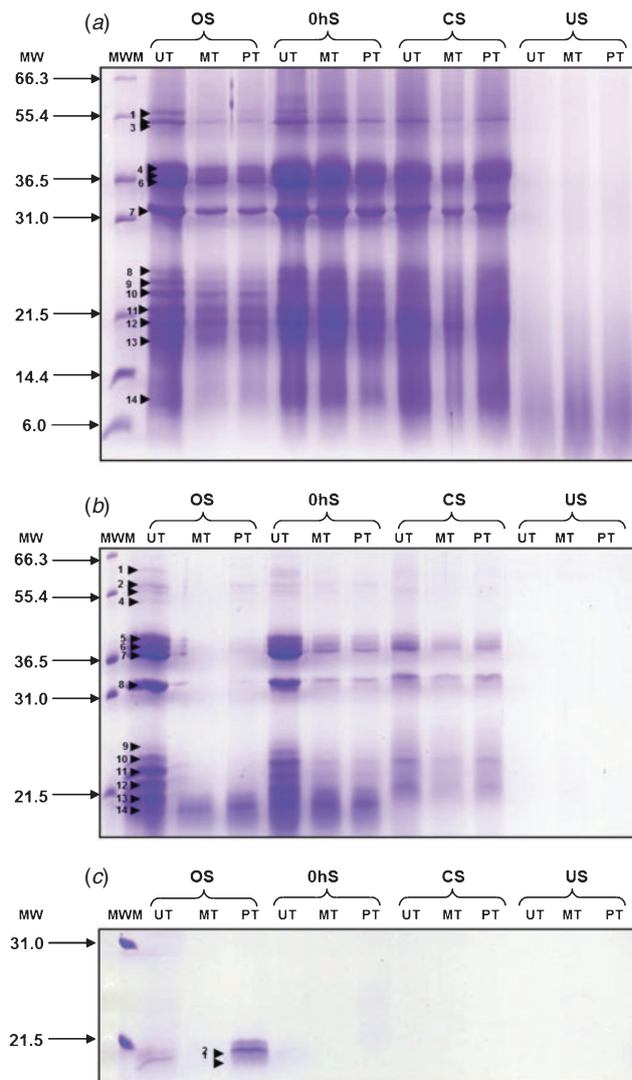


Fig. 2. SDS-gel electrophoresis of (a) total proteins, (b) globulins and (c) albumin of original (OS), 0 h (0hS), ruminally undegraded (CS) and intestinal undigested (US) samples of untreated (UT), malic acid treated (MT) and orthophosphoric acid-treated (PT) sunflower meal.

related to the extent of microbial contamination (Rodríguez and González 2006), whereas the difference in intestinal disappearance between SAB and bypass compounds is also a major factor affecting ID overestimations (Arroyo and González 2013), which is in agreement with the large overestimations shown in ID-CP of SP. Current values for SFM showed higher contaminations (4.0 and 3.1 times as average of the three meals for the DM and CP, respectively), than those observed in another SFM sample of a previous similar experiment carried out with the same doses of both acids (Arroyo *et al.* 2013). As a consequence, RU and ID overestimations in the current sample were also largely higher.

Protective treatments

The reduction in degradability with the protective treatments in both tested meals was due to a decrease in the soluble fraction

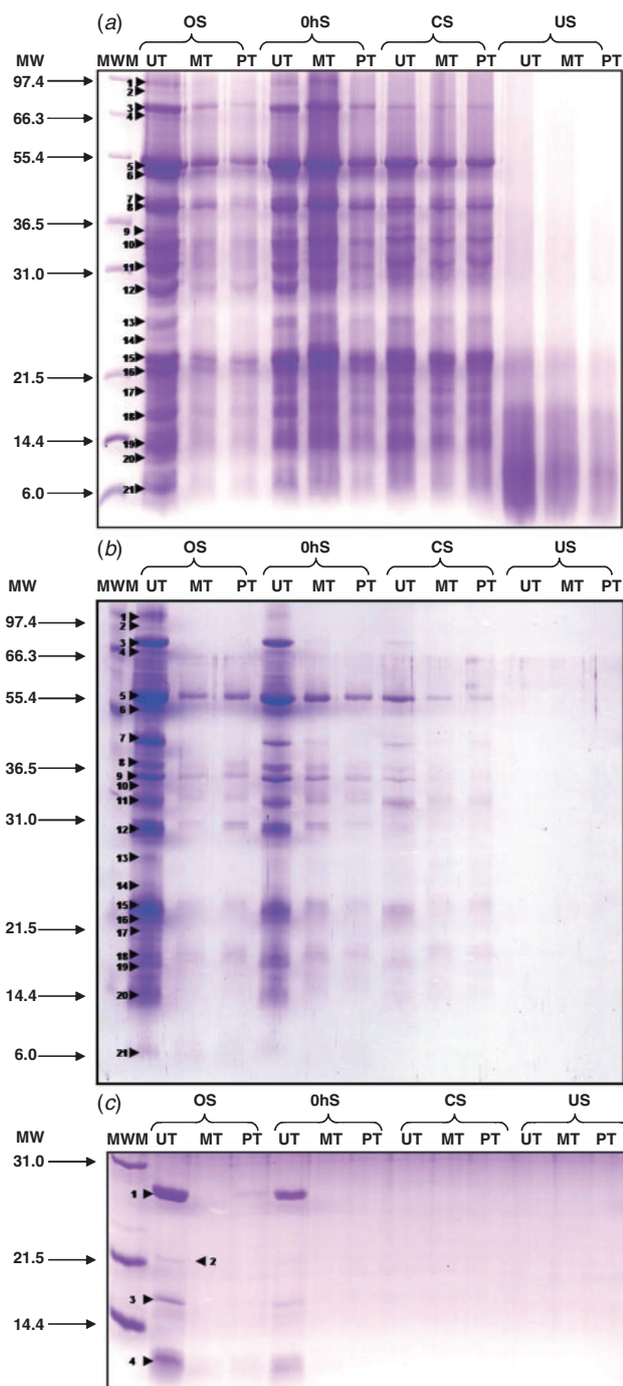


Fig. 3. SDS-gel electrophoresis of (a) total proteins, (b) globulins and (c) albumin of original (OS), 0 h (0hS), ruminally undegraded (CS) and intestinal undigested (US) samples of untreated (UT), malic acid treated (MT) and orthophosphoric acid-treated (PT) spring pea.

and increased *b* fractions, which in addition were degraded more slowly. In consequence, treatments increased the *b*-RU fraction and, therefore, shifted the digestion site to the intestine. However, these effects were lower in SFM compared with previous results by Arroyo *et al.* (2013). The decrease in ruminal DM digestibility in SFM was mainly the result of decreased degradation of CP.

Thus, microbial-corrected values showed that the reduction in degraded CP represented 86.8% (with malic acid) and 70.1% (with orthophosphoric acid) of the reduction in ruminally digested DM. This was not the case in SP, which displayed respective values for this contribution of 35.8% and 31.6%, in agreement with its moderate CP content. Equivalent values for starch were 47.4% and 38.8%, respectively. These results show that degradation of components other than proteins is also affected by these treatments in both feeds.

Observed increases of RU-CP in treated SFM were lower than those reported in Arroyo *et al.* (2013). In addition, in the experiment aforementioned, associated increases of ID-CP were observed, whereas increases seen in the present experiment were not statistically significant. As a result the increments in microbial-corrected values of IADF-CP (113.8% and 59.9% for malic and orthophosphoric acids-treated meals, respectively), were also lower than in Arroyo *et al.* (2013). This lower protection efficiency may be associated with the lower CP degradability of the present untreated SFM sample, but mainly with the lower levels of moisture and heat applied in the present experiment. The similar behaviour observed with treatments in SP also support this limited protection efficiency. This fact was augmented in SP by its extremely low ID-CP, which reduces the benefits associated with changing the digestion site to the intestine. Thus, the contents of feed-undigested CP in the total tract were 8.88% in UT, 19.2% in MT and 17.2% in PT. The low ID-CP observed in SP cannot be associated with a high concentration of indigestible compounds in the RU fraction. Thus, low values were observed both in the untreated meal and in the treated meals, which showed RU fractions 2.15 times higher on average than the untreated meal. On the contrary, it suggests the presence of a non-heat labile antitrypsin factor in SP. Thus, the results of ruminal degradability of DM, CP and starch in the untreated SP sample were close to those proposed by the INRA (2007): 80%, 86% and 79%, respectively, whereas INRA (2007) suggests intestinal digestibility of CP of 91%; 2.5 times higher than the average values for untreated and treated SP meals reported in this study.

Changes with protective treatments in RU and ID demonstrate that the close relationships shown between IADF and RU for both feeds are mainly from the decrease in rumen degradation, although in SFM the increase of ID-DM also contributed to the increased IADF value. Differences in the regression coefficients among chemical fractions also show differential effects of these treatments, which were higher for CP in SFM and for starch in SP.

The increased supply of intestinally digested CP from treated meals should also be associated with a decrease in the ruminal microbial protein synthesis owing to their reduced OM degradation. Based on corrected results of IADF-CP and OM degradability (calculated as 100-RU) and the parameters of ruminal synthesis and intestinal availability of microbial protein in the PDI system (INRA 2007), the total supply of intestinal digestible protein from SFM samples was 111.7, 154.3 and 134.9 g/kg DM in UT, MT and PT meals, respectively. Similar values in SP were 78.0, 77.6 and 76.1 g/kg DM. When CP concentrations of the meals were considered, protective treatments using malic or orthophosphoric acids respectively increased the protein value by 47.5% and 20.1% in SFM and

Table 6. Range of estimated molecular mass of polypeptidic bands detected by SDS-PAGE in sunflower meal and spring pea studied samples

Band no.	Range of molecular mass (KD)	Sunflower meal		Band no.	Range of molecular mass (KD)	Spring pea	
		Protein nomenclature	References			Protein nomenclature	References
1	56	Unknown		1 } 2 }	96–91	Unknown	
2	53	Unknown		3 } 4 }	70–68	Convicilin subunits	[1, 2, 3, 4, 5]
3	46	Unknown		5 } 6 }	55–53	Vicilins (50 KD)	[2, 3, 4]
4 } 5 } 6 }	40–30	Helianthinin (larger polipeptides)	[2, 3, 6]	7 } 8 }	40–45	α legumin (40KD)	[2, 3, 4]
7	33	Unknown		9 } 10 } 11 }	36.5–31	Vicilins (30 KD)	[2, 3, 4]
8 } 9 } 10 }	27–23	Helianthininm (smaller polipeptides)	[2, 3, 6]	12 } 13 } 14 } 15 } 16 } 17 }	30–20	β -legumin and vicilins	[2, 3, 4]
11 } 12 } 13 }	21.5–15	Albumins	[7]	18 } 19 } 20 }	20–6	Vicilins (of lower mol. mass) and albumins	[2, 3, 4, 5]
14	12	2S-methionine-rich protein	[8, 9]	21 }			

[1] Croy *et al.* (1980); [2] Derbyshire *et al.* (1976); [3] Casey (1999); [4] Casey and Domoney (1999); [5] O’Kane *et al.* (2004); [6] Allen *et al.* (1985); [7] Kortt and Cadwell (1990); [8] Kortt *et al.* (1991); [9] Shewry and Pandya (1999).

by 4.95% and 1.79% in SP. These increases may be somewhat higher than the above calculated values due to a higher efficiency of microbial synthesis associated with the reduction of the proportion of CP in the fermented OM (NRC 2001). The negative effect of these treatments on ruminal microbial synthesis through the reduction of fermented OM seems to indicate that they are mainly of interest in protein-rich feeds.

Results from the present study of SFM are in agreement with the conclusions of Arroyo *et al.* (2011, 2013) pointing to the higher efficacy of malic acid than orthophosphoric acid to protect proteins. In addition, the inclusion of malate in the diet may provide additional benefits for the ruminal fermentation pattern (Martin and Streeter 1995; Callaway and Martin 1996; Carro *et al.* 1999).

The range of MW in polypeptidic bands detected by SDS-PAGE is in agreement with that observed by Spencer *et al.* (1988) in both feeds. A proposition of band distribution comparing current MW values with literature data is presented in Table 6. An uncertain identification of some polypeptidic bands occurred in SFM, probably because the hexane extraction used in the industrial process may extract cell-wall proteins, which are incorporated to the proteins present in the SDS-PAGE gels, hindering this identification. Albumin disappearance in 0-h samples of SFM agrees with its soluble character. However, residual albumin that showed in the 0-h sample of the UT-SP meal is in agreement with results of Spencer *et al.* (1988). These authors found albumin beyond 4 h of *in vitro* incubation,

indicating that their degradation resistance was almost similar to that of bovine serum albumin, which is usually used as a standard of a protein relatively resistant to rumen degradation. However, the absence of albumin in CS profiles of untreated and treated meals of both feeds show that the practical contribution of albumin to ruminal protein outflow should be near zero even in SP. Bands of CS profiles in both feeds showed lower intensity compared with the original or 0-h samples, indicating quantitative changes resulting from degradation. The disappearance of polypeptidic bands in CS samples supports higher effects of ruminal degradation in SP than in SFM, which agrees with the quantitative CP degradation results. *In vitro* studies of Spencer *et al.* (1988) showed a faster breakdown of major subunits of convicilins, vicilins and α -legumin in peas. In addition, Aufrère *et al.* (1994, 2001) indicated a faster *in situ* degradation of vicilins and convicilins than legumin in peas. Present results provided partial support to these observations because most disappearing bands in SP have MW corresponding to both vicilins and convicilins. The higher degradation resistance of SP-legumin may be associated with the existence of disulfide bridges not present in the other SP-globulins (Casey and Domoney 1999). No polypeptidic bands were detected after intestinal digestion for SFM. Therefore, the undigested CP contents shown by *in situ* microbial-corrected data (4.70%, 8.10% and 6.6% in UT, MT and PT meals, respectively), should correspond mainly to non-protein compounds or too small peptides, which cannot be detected with SDS-PAGE techniques. These kinds of compounds should also

be present in intestinally digested SP samples that also showed poorly resolved bands in the low MW areas, probably corresponding to non-digested peptides. The latter observation is in agreement with the low intestinal digestibility shown *in situ*.

Conclusions

The protective treatments tested in this experiment decreased the ruminal degradation, displacing the digestion site to the intestine. These effects were higher with malic than with orthophosphoric acid. However, a reduction of the moisture and heat compared with previous technical recommendations decreased the treatment efficacy. Also, the treatments' effectiveness decreased with protein concentration due to the reduction of the ruminal-fermented OM and, presumably, of the resulting microbial protein synthesis. However, their effectiveness is drastically reduced when anti-nutritive factors are present in the intestine, as seems to be the case in the SP sample tested. Correction for microbial contamination of ruminal-incubated residues was important for the accuracy of current estimates.

Acknowledgements

This work has been supported by the CICYT-funded Project AGL 2006-08300 and Comunidad de Madrid-funded Projects S2009/AGR-1704 and MEDGAN ABI-2913. Analyses of ^{15}N isotope ratios were performed at the 'Servicio Interdepartamental de Investigación, Universidad Autónoma de Madrid', Spain.

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