Comparing the accuracy of PCR-capillary electrophoresis and cuticle microhistological analysis for assessing diet composition in ungulates: A case study with Pyrenean chamois

Johan Espunyes, Carme Espunya, Sara Chaves, Juan Antonio Calleja, Jordi Bartolomé, Emmanuel Serrano

Abstract

The study of diet composition is required to understand the interactions between animal and plant ecosystems. Different non-invasive techniques applied on faecal samples have commonly been used for such purposes, with cuticle microhistological analysis (CMA) and emerging DNA-based methods being the most relevant. In this work, we refined and optimized a qualitative DNA-based approach combining PCR amplification of long trnL(UAA) and ITS2 fragments and capillary electrophoresis (PCR-CE), instead of short trnL(UAA) fragments and massive sequencing technologies commonly reported. To do so, we developed a controlled diet assay using a stabled Pyrenean chamois specimen (Rupicapra pyrenaica pyrenaica), which included representative herbaceous and shrubby plant species. We also assessed the impact of sample freshness on the diet determination of this mountain caprinae by exposing faecal samples to the outdoor environment for three weeks. Faecal samples from both experiments were analysed by qualitative PCR-CE and semi-quantitative CMA in order to compare the pros and cons of both approaches. Our results show that all of the offered plant species were detected by both methodologies although CMA overdetected shrubs compared to herbaceous species. At the same time, sample degradation due to sustained climate exposure is a limiting factor for molecular analysis, but not for CMA. Taken all together, our results suggest that the qualitative information obtained by CMA and PCR-CE can be interchangeable when faecal samples are fresh (less than one week after deposition) but, afterwards, molecular analysis underestimates diet composition probably due to DNA degradation. CMA, however, can accurately be used at least three weeks after defecation. Moreover, by combining the results of simultaneous PCR amplification of two complementary genes, this optimized PCR-CE methodology provides a reliable, feasible and more affordable alternative for multiple and routine analyses of complex
samples. Neither CMA nor PCR-CE seems to solve comprehensively the quantification of herbivore diets and thus further research needs to be done.

Introduction

Food selection is a central concept in wildlife nutrition studies [1]. Although food intake evaluation is the starting point for any study of animal nutrition, field ecologists struggle with the challenging task of assessing diet composition. Feeding strategies of free-ranging species are not only influenced by foraging behaviour, food quality and availability but also by the individuals’ requirements for reproduction and maintenance [2]. Large herbivores play a major role in forming the shapes and functions of terrestrial ecosystems as they exert an impact on landscape dynamics and biodiversity [3]. Comprehensive data on their feeding habits are crucial to understand their influence on the ecology of other organisms as they drive the viability of plant communities [4], modify nutrient cycles [5] and regulate carnivore populations [6]. On the other hand, this information is necessary to evaluate their impact on threatened flora [7] and hence to elaborate plans for plant protection [8] or population control [9]. Therefore, knowing the diet composition of herbivores is an essential step towards building an ecological model of plant-herbivore interactions and is basic to many aspects of wildlife and plant research and conservation [10,11]. In fact, modern diet analysis has improved the understanding of dietary niche partitioning and the coexistence of similar herbivores [12,13] as well as enhanced conservation strategies, for example, for endangered woodland caribous [14].

Along these lines, non-invasive techniques for the analysis of herbivore diets (e.g. faeces collection or animal observation) are broadly established since they do not imply animal culling. The simplest procedures are the in situ observation of herbivore ingests of available vegetation, and the time-lapse visualization of chewed plants [15]. However, these approaches are both highly time-consuming because extensive and representative geographical areas need to be monitored in order to obtain conclusive information. Alternatively, information on diet composition can be inferred from the corresponding herbivore faeces through the analysis of plant material remaining after digestion. Two effective faeces-related methodologies should be noted. The first, cuticle microhistological analysis (CMA), is a traditional and well-established method that can provide reliable semi-quantitative data through the identification of plant cell structures, mainly the epidermis and trichomes, visualised under an optical microscope [16–18]. CMA has been extensively applied to different types of complex samples, including faeces, rumen contents and even coprolites [19], due to the strong resistance of plant epidermal cuticles to climate and environmental degradation processes. However, CMA presents important drawbacks that limit its use. Significant expertise in microscopic skills is required [20]. In addition, identification of plant structures is frequently impaired by the strong similarity of anatomic structures between taxonomically related genera to the extent that in some cases only the family level can be determined [18].

The second category of faeces-related methodologies involves the analysis of remaining plant genomic DNA isolated from faeces, also known as DNA barcoding methods [21–23]. DNA barcoding is considered a powerful and accurate alternative to morphological methods and has been applied in many biological studies including the authentication of herbal medicinal products [24,25], ancient permafrost sediment analysis [26–28], and the assessment of herbivore diet composition [29–34]. Next-generation sequencing (NGS) technology, is currently the most used DNA barcoding method and consists of the identification of different taxa using
marker gene sequences, also called barcodes, through amplification by PCR and subsequent sequencing by massive NGS [22,35–38]. In plants, different chloroplast genes have been extensively used—e.g. *matK, trnL(UAA), rbcL* and *trnH-psbA*—either alone or combined with nuclear genes, such as the Internal Transcribed Spacer regions of ribosomal DNA, *ITS1* and *ITS2* [36,39–41]. However, NGS-DNA barcoding implementation has been hampered by the economic cost of sequencing technologies [35,38], whereas an alternative method involving capillary electrophoresis has emerged as a more affordable and suitable option for multiple and routine analysis of complex samples [32,42]. This alternative technique, hereafter called PCR-capillary electrophoresis (PCR-CE), applies capillary electrophoresis to determine the amplicons lengths, instead of DNA-sequencing as in NGS-DNA barcoding. We obtain a migration pattern that allows the detection of several peaks characteristic of multi-species samples. The subsequent botanical adscription of each peak is realized through matching experimental data with those available in the GenBank database [43]. PCR-CE is not commonly used in studies of diet composition despite being fast, inexpensive and frequently applied in the food industry ([44,45], but see [32]). Usually, the resolution of this method is considered low for complex samples because plant species with similar fragment sizes can account for the same peak due to a visual overlap [32]. However, this drawback can be overcome by combining simultaneous PCR amplification of two complementary genes [42].

Here, we wanted to refine and optimize a PCR-CE analysis for a higher resolution in the determination of diet composition. For this purpose, we subjected a stabled chamois specimen to a controlled diet, which included representative herbaceous plants and woody shrubs of the eastern Pyrenees ecosystem. We developed a method combining simultaneous PCR amplification of two genes, *trnL(UAA)* and *ITS2*, and capillary electrophoresis to determine the corresponding fragments length. We expected this PCR-CE alternative method to lead to a higher fragment resolution and more precise plant species identification compared to conventional amplifications methods previously described [46,47] avoiding, at the same time, the high cost and the need for specific expertise on NGS. Simultaneously, cuticle microhistological analysis was also carried on these samples in order to compare both technical approaches.

On the other hand, one of the major concerns in the determination of diet composition is knowing to what extent the freshness of faecal samples may compromise the obtaining of reliable results. In that way, we also aimed at determining if the time-lapse between animal deposition and faecal sampling impacts on CMA and PCR-CE achievements. As a pilot assessment, we exposed a pool of faeces obtained from hunter-harvested Pyrenean chamois (*Rupicapra pyrenaica pyrenaica* Bonaparte) to outdoor environmental conditions for several weeks. We then determined the number of plant species detected by both methodologies every week along the study. Being used in archeobotany [19], we expected CMA results not to be influenced by environmental exposure. On the other hand, due to the high degradability of DNA [48], we expected an effect of the time of exposure on the number of plants detected by the DNA-based method. At the same time, this assessment allowed us to test the efficiency of PCR-CE on a more species-diverse sample.

**Materials and methods**

**Study area and sampling procedure**

Fresh plant samples for the controlled diet tests were collected in June 2015 in the Freser-Setcases National Game Reserve (FSNGR), Eastern Pyrenees, Spain (42˚ 22' N, 2˚ 08' E). The FSNGR is a typical sub-humid subalpine and alpine bioclimatic region with a noticeable Mediterranean climatic influence. It covers an area of 20,200 ha, ranging from 1,800 to 2,910 m.a.s.l. and hosts a population of about 3,000 Pyrenean chamois [49]. The fresh faecal samples used
for the time-course degradation study were collected from 12 hunter-harvested Pyrenean chamois during the hunting season of 2014–2015 in the same area. Faecal and plant samples were refrigerated at 4°C during transport to our facilities and stored frozen at -20°C until processing.

**Controlled diet test**

A twelve-year-old female Pyrenean chamois was fed on several diets that overall included 13 vascular plant species, comprising herbaceous plants and woody shrubs representative of the subalpine and alpine grassland ecosystem of the FSNGR. Three different plant mixtures were consecutively offered and intercalated with a four-day exclusive *Medicago sativa* (L.) diet, as shown in Table 1. Fresh faeces were collected before beginning the controlled diet (day 7) and the day after the corresponding diet intake, and were immediately stored at -20°C until processing. The chamois used in this study was observed with ectopia lentis—an ocular condition—in the central Catalan Pyrenees (Pallars Sobirà; 42° 58’ N, 1° 22’ E). The animal was captured by hand by a team of rangers and sent to the wildlife facilities of the Universitat Autònoma de Barcelona for veterinary inspection. The chamois was maintained in an isolated box (6m x 3m) with natural light for six months, pending translocation. Even though the animal showed some degree of visual impairment, it was able to feed selectively on the offered diet (personal observations).

**Ethics statement**

The approval to keep a captive Pyrenean chamois was supported by the Departament d’Agricultura, Ramaderia, Pesca, Alimentació i Medi Natural—Generalitat de Catalunya.

Table 1. Schedule of the controlled diet offered to a stabled specimen of Pyrenean chamois to compare the effectiveness of cuticle microhistological analysis (CMA) and PCR amplification combined with capillary electrophoresis (PCR-CE) in assessing diet composition of herbivores through faecal samples.

| Days    | Diet components | Weight (g) | Percentage (%) |
|---------|-----------------|------------|----------------|
| 1 to 7” | *Medicago sativa* (Ms) | ad libitum |                |
| 8       | Diet 1:         |            |                |
|         | *Calluna vulgaris* ( Cv) | 28.5       | 39.8           |
|         | *Carex caryophyllea* (Cc) | 13.5       | 18.8           |
|         | *Festuca spp.* | 29.7       | 41.4           |
| 9” to 12| *Medicago sativa* | ad libitum |                |
| 13      | Diet 2:         |            |                |
|         | *Helianthemum nummularium* (Hn) | 17.3       | 18.5           |
|         | *Juncus trifidus* (Jt) | 6.7        | 7.2            |
|         | *Nardus stricta* (Ns) | 47.0       | 50.3           |
|         | *Trifolium alpinum* (Ta) | 22.5       | 24.1           |
| 14” to 17| *Medicago sativa* | ad libitum |                |
| 18      | Diet 3:         |            |                |
|         | *Arrhenatherum elatius* (Ae) | 76.4       | 32.3           |
|         | *Hypochaeris radicata* (Hr) | 68.7       | 29.2           |
|         | *Vaccinium uliginosum* (Vu) | 89.4       | 38.5           |
| 19”     | *Medicago sativa* | ad libitum |                |

Composition of the plant mixtures is indicated and the weight of each plant species in the mixture is given in grams (g) and in percentage (%). Plant species initials are listed between brackets.

* Days of faeces collection.

1 Includes a mixture of *F. eska* (Fe), *F. gautieri* (Fg), *F. glauca* (Fgl), and *F. violacea* (Fv).

https://doi.org/10.1371/journal.pone.0216345.t001
(DARPAMN—the Regional authority in charge of livestock and wildlife management). In this work we did not require the approval of the Ethical and Animal Welfare Committee of the Universitat Autònoma de Barcelona. In fact, the Directive 2010/63/EU of the European Parliament indicates: Specific protocols shall not apply to practices not likely to cause pain, suffering, distress or lasting harm equivalent to, or higher than, that caused by the introduction of a needle in accordance with good veterinary practice.

Time-course degradation

Fresh faeces from the hunter-harvested Pyrenean chamois were randomly mixed in 4 pools and left outdoor at the Universitat Autònoma de Barcelona campus (Bellaterra, Spain) for three weeks starting in January 2016. This experiment was set up as a pilot assessment to evaluate the connection among the time-lapse between chamois deposition and faecal sampling—a potential degree of sample degradation—and the performance of both CMA and PCR-CE methods. In that period, temperatures ranged from 2.5˚C to 21.7˚C (mean = 11.1˚C) with short periods of rain between weeks 2 and 3 (S1 Fig). Pellets from the four different mixtures were collected every week and kept at -20˚C until processing. After three weeks of harvesting, they were subsequently tested as described below and the number of identified plant species was monitored using the two methods.

Genomic DNA extraction and multiplex-PCR DNA amplification

Genomic DNA from plants and chamois faeces was extracted from 200 mg of raw material with the DNeasy Plant Mini Kit and the QIAamp DNA Stool Kit (Qiagen, Germany), respectively, following manufacturer’s instructions. Genomic DNA was eluted in 100 μl of milli-Q water and kept at -20˚C. DNA concentration was measured with a NanoDrop 1000 Spectrophotometer (ThermoScientific, USA) and DNA quality was analysed on a 0.8% agarose gel electrophoresis in 1x TBE buffer.

Multiplex-PCR reactions for specific amplification of chloroplastic trnL(UAA) and nuclear ITS2 sequences in plants and faeces were carried out in a total volume of 20 μL containing 1x MyTaq Reaction Buffer (Bioline Reagents Ltd, United Kingdom), 0.4 μM each oligonucleotide (Stab Vida, Portugal; see S1 Table for detailed oligonucleotide sequences), 1 U MyTaq HS DNA polymerase and 20 ng of genomic DNA template. Oligonucleotides trnL-G and trnL-D were used to amplify the trnL(UAA) region [47], while oligonucleotides S2F and S3R amplified the ITS2 region [40]. Oligonucleotides trnL-D and S2F were fluorescence-labelled with 6-FAM (carboxyfluorescein) and HEX (hexachloro-6-carboxyfluorescein) fluorochromes, respectively, to allow simultaneous fluorescent detection by capillary electrophoresis.

Cycling conditions were optimized for multiplex DNA amplification and comprised an initial denaturation for 1 min at 95˚C, followed by 35 cycles of denaturation for 15 s at 95˚C, annealing for 15 s at 56˚C and extension for 45 s at 72˚C, and a final extension step for 3 min at 72˚C. PCR reactions were run on a MJ Research PTC-100 Thermal Cycler (MJ Research Inc., Canada) and amplicons were visualised on a 1.5% agarose gel electrophoresis in 1x TBE buffer.

DNA sizing

Amplicon lengths were determined by automated capillary electrophoresis with a Genetic Analyzer 3130xl system (Applied Biosystems, USA) using LizS600 as a size standard. Electropherograms were in silico analysed with provided Peak Scanner Software version 2.0 (Applied Biosystems, USA). Plant genomic sequences of the trnL(UAA) and ITS2 regions were retrieved...
from the GenBank database using the nucleotide BLASTN tool (e-value < e-04) [50]. Sequence lengths were calculated by flanking corresponding primers to each plant sequence.

**Slide preparation for CMA analysis**

Fresh faeces were lightly ground by hand in a mortar to separate out the epidermal fragments [51]. From each sample, 0.5 g of ground faeces were introduced into a test tube with 3 ml of concentrated HNO$_3$ to allow non-epidermal tissue digestion. The test tubes were placed in a water bath at 80˚C for 2 minutes and the samples were then diluted with 200 ml of distilled water. This suspension was successively sieved through 1 mm and 0.25 mm filters. The 0.25 to 1 mm fraction was dispersed in a 50% (v/v) aqueous solution of glycerine. Samples of the suspensions were spread on glass microscope slides at a density that prevented any significant overlapping of fragments. Cover slips (24 x 60 mm) were then fixed to the slides with DPX varnish (Herter Instruments, Spain). Three slides were prepared from each sub-sample. A reference epidermis collection of the different plant species included in the study was set up in parallel using two different methods. First, fresh plant material was coarsely ground by hand and then processed in the same way as the faeces samples in order to simulate the fragmentation of the epidermis during ingestion and digestion. Second, fresh abaxial and adaxial leaf surfaces were obtained with a scalpel and mounted on a glass microscope slide allowing a more accurate identification of the main morphological features of the different species.

**Fragment identification**

Each slide was examined under a Motic BA210 optical microscope (MoticEurope SLU, Spain) at 400x magnification. Three traverses were scanned, each one 2 mm wide and 60 mm long with 3 mm between traverses. Up to 200 epidermis fragments were identified per sample using the reference collection of plant species, based on the shape of epidermal cells, trichomes and stomata surrounding cells. The data were then pooled and converted to percentages. Reference collection images were captured with a Moticam 2300 camera using provided Motic Images Plus 2.0 software.

**Statistical analysis**

We used a linear model (LM) to explore the effects of the elapsed time between faecal deposition and sampling on the number of plant species identified (response variable) by CMA and PCR-CE. The LM included the sampling period (fixed categorical factor with four categories: Initial and 1st, 2nd, and 3rd week) and the identification technique (CMA and PCR-CE) as fixed explanatory factors, and the number of identified plant species as a response variable. Due to our moderate sample size (n = 32, two techniques x four determinations per week), model parameter uncertainty was estimated by residual-based bootstrap methods using the Boot package 1.3–20 version [52]. LM assumptions were evaluated by exploring the residual pattern (homoscedasticity, linearity and normality) [53]. Weekly differences in the number of plants detected by both methodologies were tested by a Tukey’s HSD [54]. All statistical analyses were performed in R version 3.4.3 [55].

**Results**

**DNA fragments length and CMA reference databases**

We established a database of different plants, 13 of them representative of the eastern Pyrenees ecosystem and *Medicago sativa*, based on the size of multiplex-PCR amplified *trnL(UAA)* and *ITS2* genomic sequences with primers *trnL-G/D* and *S2F/S3R*, respectively. Both genes were
successfully amplified in all the plants tested, confirming the universality of the oligonucleo-
tides previously designed for plant DNA barcoding studies (S2 Fig) [40,47].

The size of the fluorescently labelled PCR products determined after capillary electrophore-
sis ranged from 303 to 593 bp for \textit{trnL(UAA)} amplicons and from 463 to 500 bp for \textit{ITS2}
amplicons, which strongly match the lengths calculated from retrieved GenBank sequences
(S3 Fig and Table 2). Despite the fact that \textit{trnL(UAA)} exhibits higher resolution than
\textit{ITS2}, since the corresponding PCR products show a wider range of sizes, the data obtained from the
two genomic sequence amplifications is complementary. Thus, \textit{Calluna vulgaris} (L.) Hull,
\textit{Carex caryophyllea} Latourr. and \textit{Vaccinium uliginosum} L. showed similar
\textit{ITS2} fragment sizes (488, 490 and 492 bp, respectively), but they could be clearly discriminated through their \textit{trnL}
(UAA) fragments (492, 593 and 464 bp, respectively). The species of the genus \textit{Festuca}
were almost indistinguishable by their \textit{ITS2} fragments (472–475 bp) but exhibited distinct \textit{trnL}
(UAA) fragment sizes (504–517 bp), with the exception of \textit{F. eskia} Ramond ex DC. and \textit{F.
gautieri} (Hack.) K.Richt., which were identical. Furthermore, \textit{Nardus stricta} L. and \textit{Trifolium alpinum}
L. could be clearly differentiated by their \textit{ITS2} fragment (471 and 480 bp, respectively),
but they could be clearly discriminated through their \textit{trnL}
(UAA) fragments (518 and 516 bp). Three species, \textit{Arrhenatherum elatius}
(L.) P. Beauv. ex J. Presl & C. Presl, \textit{Medicago sativa} and \textit{Juncus trifidus} L., exhibited
a shorter \textit{trnL(UAA)} fragment, which sharply differed from the rest of the plant species.

Concerning CMA, all plant epidermis fragments were easily distinguishable at the genus
level based on their anatomical structures [56]. In summary, graminoids presented rectangular
eridermic cells but differences between the species in terms of cell size, cell wall shape and the
presence and shape of trichomes and stomata (Fig 1A). \textit{C. caryophyllea} fragments presented
square to rectangular epidermic cells with wavy cell walls, prickle-shape trichomes and diacytic
stomata. \textit{J. trifidus} also presented rectangular elongated epidermal cells but the wall was
smoother. \textit{A. elatius} displayed rectangular epidermic cells with wavy walls intercalated with
short cells. \textit{N. stricta} also had rectangular epidermis cells with wavy walls but were longer than

### Table 2. DNA fragment lengths reference database for the plant species included in the controlled diet test.

| Plant species              | \textit{trnL(UAA)} | \textit{ITS2} |
|----------------------------|-------------------|--------------|
|                            | amplicon size (bp) | fragment sequence length (bp) | GenBank accession | amplicon size (bp) | fragment sequence length (bp) | GenBank accession |
| \textit{Arrhenatherum elatius} | 305               | 310          | EU434100 | 471           | NA            | -          |
| \textit{Calluna vulgaris}   | 492               | 495          | KP737377 | 488           | 497           | KP737514  |
| \textit{Carex caryophyllea} | 593               | 593          | EU288467 | 490           | 499           | HG915830  |
| \textit{Festuca eskia}      | 517               | 519          | AF478508 | 473           | 484           | KP97144   |
| \textit{Festuca gautieri}   | 517               | 519          | KP699267 | 472           | 482           | AF303414  |
| \textit{Festuca glauca}     | 504               | 506          | JX871940 | 475           | 485           | AY327792  |
| \textit{Festuca violacea}   | 508               | 510          | EF993012 | 472           | 484           | EF584979  |
| \textit{Helianthemum nummularium} | 491          | NA            | -          | 463           | 479           | GU327669  |
| \textit{Hypochaeris radicata} | 530             | 532          | AY504774 | 483           | 485           | AF528461  |
| \textit{Juncus trifidus}    | 343               | 345          | AY437971 | 484           | 495           | AY973508  |
| \textit{Medicago sativa}    | 326               | 327          | GQ488614 | 484           | 486           | AF053142  |
| \textit{Nardus stricta}     | 518               | 520          | EU434097 | 471           | 479           | KJ77049   |
| \textit{Trifolium alpinum}  | 516               | 522          | DQ311725 | 480           | 484           | DQ311995  |
| \textit{Vaccinium uliginosum}| 464               | 466          | DQ286840 | 492           | 498           | DQ217769  |

Amplicon sizes were determined by capillary electrophoresis in this study. Fragment sequence lengths were calculated from retrieved GenBank sequences. (bp), base pairs. NA, not available.
Arrhenatherum elatius and some prickle-shaped trichomes could also be observed. *Festuca* spp. showed long epidermic cells with wavy walls separated by C-shaped short cells, paracytic stomata and spindle-shaped trichomes. Depending on the *Festuca* species, cell walls displayed variable waviness, yet fragments could not be characterised at the species level due to the extraordinary morphological similarity exhibited by this genus.

Dicots were also clearly distinguishable by their specific micro-anatomical features, especially trichomes and stomata (Fig 1B). For Ericaceae, *C. vulgaris* presented lobed epidermal cells and anisocytic stomata protected by medium-sized, wavy trichomes while *V. uliginosum* displayed angular to lobed epidermal cells, paracytic stomata and no trichomes. Additionally, veins were easily observable in the *V. uliginosum* fragments. *Helianthemum nummularium* (Cav. Losa & Rivas Goday) showed angular polygonal epidermal cells, cyclocytic stomata and star-shaped trichomes. *Hypochaeris radicata* L. presented irregular rectangular cells with angles often different than 90 degrees, anamocytic stomata and large multicellular hairs. Finally, Fabaceae *T. alpinum* fragments showed angular polygonal epidermal cells, anamocytic stomata and medium-sized trichomes with pointed tips, and *M. sativa* displayed polygonal...
epidermal cells with undulated cell walls and long single-cell trichomes with a reflective dual-cell base.

Analysis of the botanical composition of chamois faeces subjected to a controlled diet

The 13 plant species that were distributed in three different mixture diets and offered to a Pyrenean chamois were completely ingested within the timeframe they were offered. Multiplex PCR amplification of trnL(UAA) and ITS 2 sequences from genomic DNA extracted from chamois faeces yielded products of expected size in all the samples tested (S4 Fig). Analysis of fluorescently labelled PCR products by capillary electrophoresis showed precise and independent peaks of different lengths that could be roughly ascribed to a particular plant species (Fig 2). We were able to identify all the plants included in the controlled diets. For a better interpretation of the electropherograms, the two fluorescence channels were independently preselected. Two common peaks among the three controlled diets of 326 and 484 bp for trnL(UAA) and ITS2 amplicons, respectively, were observed and would correlate with M. sativa background intake.

In diet 1 electropherograms, 6 peaks for trnL(UAA) and 5 peaks for ITS2 amplicons were distinguished, which may correspond to the 7 plant species included in the diet (Fig 2A). In contrast, two species of the genus Festuca, F. eskia and F. gautieri remained undetermined. A wider peak of 474–476 bp in the ITS2 amplification was assigned to the Festuca genus. Diet 2 electropherograms exhibited 7 peaks for trnL(UAA) and 6 peaks for ITS2 amplicons that could be assigned to the 4 diet species (Fig 2B). One trnL(UAA) peak, adjacent to H. nummularium, did not match any plant species but may match a particular amplification profile for this species (S3 Fig). Furthermore, one of the peaks for each gene correlated with C. vulgaris size amplicons suggesting that residual genomic DNA of this species belonging to diet 1 was present in the analysed faeces. Moreover, diet 3 trnL(UAA) electropherograms displayed a total number of 6 peaks: 3 of them corresponded to the diet species, the other 2 to residual H. nummularium and N. stricta from diet 2, and the common M. sativa peak. For the ITS2 channel, results were less resolved as two of the peaks (472 and 483 bp) matched more than one species (Fig 2C).

Regarding CMA analysis, all the species included in the study were identified and quantified in the faeces, with the exception of the species of the genus Festuca, which could not be discriminated due to their almost identical epidermal structures. Results are given as percentages of epidermal fragments quantified in faecal samples (Table 3). Quantification of M. sativa was discarded for better determination of experimental plants. Our results show that epidermal fragments from the different species were preserved in the successive faecal samples throughout the experiment to the extent that species included in diets 1 and 2 were also detected in diet 3.

Furthermore, the CMA approach enables the possibility of linking percentages of epidermal fragments quantified in faeces (% EPF) with those of plant ingested weights (% I). Thus, an EPF/I ratio can be defined for each species in the corresponding diet, providing information about a putative over-detection of a particular species in faeces when EPF/I > 1 (Table 4). For a simplification of the model, only original diet species have been included in ratio calculations.

For example, in diet 1, ingested quantities of C. vulgaris and Festuca spp were equivalent and doubled that of C. caryophylllea ingests (39.7%, 41.4% and 18.8%, respectively) but the first was the predominant species detected in faeces (74.8%, 16.1% and 9.1% resp.). EPF/I ratios suggest an over-detection of C. vulgaris (EPF/I = 1.9) and a significant equivalent infra-
Fig 2. Electropherograms peaks of the three controlled diets offered to the stabled Pyrenean chamois, obtained by capillary electrophoresis of the fluorescence labelled PCR products corresponding to trnL(UAA) (blue) and ITS2 (green) multiplex amplification. For a better interpretation, the two fluorescence channels are shown separately. Plant species initials are as in Table 1.

https://doi.org/10.1371/journal.pone.0216345.g002
detection of *Festuca* spp (EPF/I = 0.4) and *C. caryophyllea* (EPF/I = 0.5). A similar behaviour is observed in diet 2. Despite the fact that *H. nummularium* was not the most ingested species, it was over-detected in the corresponding faeces (EPF/I = 1.5), while *N. stricta*, which was the main species in the diet, was poorly detected, showing a EPF/I ratio of 0.2, whereas for *J. trifidus* and *T. alpinum* the EPF/I was 1.1. The three plant species included in diet 3 were infra-detected in faeces showing EPF/I ratios of 0.2–0.6.

### Time-course degradation analysis

Descriptive statistics for the mean number of plant species found in our pool of faecal samples exposed to outdoor climatic conditions is shown in Table 5. According to our LM, 94.4% of the observed variability in the number of plant species detected was explained by the interaction between the time elapsed since samples were exposed to the environment and the detection technique ($F_{7, 14} = 40.5$, $p < 0.001$). The mean number of plant species detected by

| Diet 1 | EPF (%) ± SE | Diet 3 | EPF (%) ± SE |
|---|---|---|---|
| Calluna vulgaris | 74.8 ± 4.4 | Arrhenatherum elatius | 18.7 ± 4.2 |
| Festuca spp. | 16.1 ± 3.3 | Nardus stricta | 17.6 ± 5.6 |
| Carex caryophyllea | 9.1 ± 0.7 | Festuca spp. | 14.9 ± 3.5 |
| | | Hypochaeris radicata | 10.9 ± 3.1 |
| Diet 2 | EPF (%) ± SE | Diet 3 | EPF (%) ± SE |
| Helianthemum num. | 28.3 ± 2.9 | Vaccinium uliginosum | 9.4 ± 1.9 |
| Trifolium alpinum | 27.8 ± 6.5 | Helianthemum num. | 8.8 ± 1.5 |
| Nardus stricta | 9.7 ± 2.3 | Calluna vulgaris | 7.5 ± 3.3 |
| Festuca spp. | 9.4 ± 1.6 | Carex caryophyllea | 4.5 ± 1.0 |
| Juncus trifidus | 7.9 ± 1.2 | Trifolium alpinum | 2.9 ± 1.7 |
| Carex caryophyllea | 0.3 ± 0.3 | |

(SE), standard error.

https://doi.org/10.1371/journal.pone.0216345.t003

### Table 4. Ratios of epidermal plant fragments (EPF) quantified by cuticle microhistological analysis (CMA) in faecal samples from a stabled specimen of Pyrenean chamois versus ingested plant weights (I) for each plant species in the corresponding diet.

| Plant species | Herbaceous/shrub | % ingest (I) | % EPF (EPF) | EPF/I | Lignin (% of dry matter) | Ref. |
|---|---|---|---|---|---|---|
| Diet 1 | Calluna vulgaris | Shrub | 39.8 | 74.8 | 1.9 | 17.8 | [57] |
| | Carex caryophyllea | Herbaceous | 18.8 | 9.1 | 0.5 | 3.3 | [58] |
| | Festuca spp. | Herbaceous | 41.4 | 16.1 | 1.0 | 5.4 | [14] |
| Diet 2 | Helianthemum num. | Shrub | 18.5 | 28.3 | 1.5 | 7.4 | [59] |
| | Juncus trifidus | Herbaceous | 7.2 | 7.9 | 1.1 | 2.8 | [60] |
| | Nardus stricta | Herbaceous | 50.3 | 9.7 | 0.2 | 5.4 | [58] |
| | Trifolium alpinum | Herbaceous | 24.1 | 27.8 | 1.1 | 6.9 | [58] |
| Diet 3 | Vaccinium uliginosum | Shrub | 38.5 | 9.4 | 0.2 | 7.4 | [61] |
| | Arrhenatherum elatius | Herbaceous | 32.3 | 18.7 | 0.6 | ND | - |
| | Hypochaeris radicata | Herbaceous | 29.2 | 10.9 | 0.4 | 3.1 | [62] |

Overdetecion is reached when EPF/I >1.

Lignin content values were taken from the indicated references. ND, data not reported in the bibliography.

https://doi.org/10.1371/journal.pone.0216345.t004
PCR-CE was notably higher than the mean number detected by CMA at the initial starting point of outdoor exposure. However, while the number of species identified by CMA was fairly stable during the trial, the species detected by the molecular method dropped by more than half since the first week. In the faecal material exposed to the environment for three weeks, no species were detected by this technique. According to the Tuckey’s HSD test, the mean number of detected species by both techniques differed significantly in all sampling periods except for the initial sampling \((q = -2.25 [-4.73;0.23], p = 0.09, \text{Table 5})\).

**Discussion**

In this article, we describe and compare two technical approaches for the study of the diet of herbivores (folivores *sensu stricto*) using faecal samples as a source for analysis: a modified PCR-based DNA amplification of marker genes (PCR-CE approach) and a microhistology-based identification of the remaining plant epidermis (CMA approach). In both methodologies, a previous description of the plant species that constitute the available resources for herbivores is an essential prerequisite. The identification of the putative ingested plant species is supported by the collation of the information obtained from the remaining plant epidermis and/or genomic DNA in the faeces with those available in plant reference databases.

Concerning the molecular approach, we have developed an optimized method for multiplex-PCR DNA amplification of two genomic sequences including chloroplast *trnL(UAA)* and nuclear *ITS2* plant genes using total genomic DNA extracted from herbivore faeces as a template. The use of fluorescently labelled oligonucleotides enables the subsequent determination of the amplicon sizes by capillary electrophoresis, which in turn is the basis for plant species determination in diet composition. It has been previously described that DNA barcoding of *trnL(UAA)* using conserved oligonucleotides G and H broadly amplifies a fragment of 10–143 bp covering the P6 loop of the plant *trnL(UAA)* intron, and plant species adscription relies on the application of massive sequencing technologies [30,47]. We show in this work that the amplification of larger *trnL(UAA)* fragments (310–593 bp), including the conserved oligonucleotides G and D and comprising the intron and the exon 2 gene regions, greatly improves the discriminating capacity of the technique due to the hypervariable length of the region and enables a more accurate identification of plant species. Moreover, the Internal Transcribed Spacer 2 (*ITS2*) region, situated between the 5.8S and 26S genes of the nuclear ribosomal cistron, has also been considered an optimal candidate gene marker, due to its appreciable interspecific divergences even in closely related species [39,40]. Since the unique plant structures that can be distinguished in faeces samples are the epidermal cells, which lack chloroplast

### Table 5. Mean ± standard deviation, minimum and maximum number of plant species detected in a pool of Pyrenean chamois faeces exposed to outdoor conditions using cuticle microhistological analysis (CMA) and PCR amplification combined with capillary electrophoresis (PCR-CE).

| Sampling period | CMA       | PCR-CE     |
|-----------------|-----------|------------|
| Initial         | 8.75 ± 0.5 (8–9) | 11 ± 1.83 (9–13) |
| 1st Week        | 8.75 ± 1.5 (7–10) | 4.25 ± 0.71* (3–5) |
| 2nd Week        | 8.75 ± 0.96 (8–10) | 1 ± 0.82* (0–2) |
| 3rd Week        | 8 ± 0.82 (8–10) | 0 ± 0* (--) |

* denotes significant differences between methods, at \(\alpha = 0.05\), Tukey’s HSD test.

https://doi.org/10.1371/journal.pone.0216345.t005
organelles, the concomitant amplification of a nuclear genomic sequence, such as ITS2, serves as a control for DNA template quality. In addition, ITS2 analysis supplements the data obtained from trnL(UAA) concerning plant species identification belonging to a particular genus (i.e., Festuca genus), suggesting that the information obtained from both genes may be complementary. Furthermore, genomic sequences of trnL(UAA) and ITS2 genes from many plant species are freely accessible through public databases, such as GenBank [43], enabling the in silico prediction of gene lengths. It is noteworthy that retrieved GenBank sequences corresponding to both genes are a few base pairs longer compared to the fragment amplicon sizes experimentally determined, probably due to the intrinsic sensitivity of the capillary electrophoresis procedure, as previously described [32]. However, these slight variations do not affect the identification of the plant species since fragment sizes determined by capillary electrophoresis are equivalent in fresh plant tissues and in faeces samples analysis. Although we tested the efficacy of PCR-CE combining two complementary genes on faecal samples from free-ranging chamois, the reliability and resolution of the method should be tested on even more species-diverse diets [14].

CMA provides valuable information about herbivore diet choices through microscopic recognition of digested plant epidermises [16,51]. This method has been widely used for decades due to its versatility and low economic cost, although a high degree of observer expertise is required for accurate microscope interpretation and the level of discrimination between some related plants is relatively low [63]. Microhistological analysis of faeces is more precise than macroscopic identifications of rumen contents [46,64], and may serve to assess diet composition not only in game ungulates but also in species with some degree of protection (e.g., Tatra or Abruzzi chamois among others), thus offering the opportunity to avoid invasive hunting praxis. One of the strong features of CMA is that it offers the possibility to perform semi-quantitative determinations of herbivore diet composition (folivores sensu stricto), through the direct extrapolation of the plant epidermal fragment quantities present in the faecal samples [20]. However, differing degrees of digestibility of plant species that may determine the remaining plant material in the faeces must be considered [18,57,65,66]. To explain this apparent inconsistency, we propose an index of preservation for each plant species, called EPF/I ratio, that would link the relative quantity of one species detected in a faecal sample (EPF) to the corresponding ingested quantity (I). The index would reflect a putative over-detection of a plant species in the faeces when EPF/I > 1, due to a slow or a reduced digestion process of the ingested plant tissues. Considering the plant species included in the study, the shrubby or herbaceous constitution of the plant is most likely the main factor determining the digestibility of a particular plant species. The plant cell wall is composed of polysaccharides and secondary metabolites that assemble to generate complex structures. Shrubs are persistent woody plants with reinforced and thicker cell walls compared to herbaceous plants, partially due to their high lignin content, and therefore are prone to a relatively low digestibility [17,58,60]. Intriguingly, the lignin content values reported in the bibliography for the species included in this study, describe an overall correlation with their structural constitution, where the highest lignin contents are consistent with the shrubby species—C. vulgaris and H. nummularium—and, concomitantly, with their average over-detection in faeces (Table 4). This observation suggests that lignin may exert a protective effect over plant structures throughout the digestion process, which would lead to their preservation, thus triggering the over-detection of shrubby compared to herbaceous species in the faeces. However, lignin content and plant constitution may not be the only explanations for the discrepancies between plant ingests and plant fragments detected in faeces (EPF/I ≠ 1), since in our experimental conditions two herbaceous species, J. trifidus and T. alpinum, showed equivalent EPF/I ratios, but different lignin contents. It is also notable that the concomitant detection of many plant species hinders the observation under
the microscope and debases the relative species quantification, thus hampering the conclusive establishment of EPF/I correlations. This fact occurred in the corresponding faecal samples from diet 3 and especially in the case of the shrubby *V. uliginosum*. We propose that further determinations of individual EPF/I ratios for preferred plant species would provide valuable and more reliable information in CMA-based herbivore diet studies.

The greater resistance to digestion of some shrubby plant species observed by CMA was also perceived at the molecular level since *C. vulgaris* (diet 1) and *H. nummularium* (diet 2) were PCR-amplified in faecal samples belonging to diets 2 and 3, respectively, while none of the herbaceous plants showed this behaviour. Furthermore, PCR amplification is roughly independent of the structural constitution of a plant species but strongly hinges on sample freshness. Thus, while the number of species detected by CMA remained stable in samples kept outdoors for three weeks, the number of species amplified by PCR markedly decayed within the first week. The length of the PCR amplicons, which were 450 bp on average, provides high resolution to discriminate between plant species, but the progressive degradation of the genomic DNA due to deficient preservation conditions is a drawback for amplification success [25,32,48,67]. Although PCR amplification of shorter genomic fragments (70–100 bp in length) from ancient samples containing partially degraded genomic DNA has been previously described in DNA barcoding studies, plant species adscription was performed through massive DNA sequencing instead of amplicon length determination [26–28]. Massive sequencing technologies are powerful tools that provide accurate information but require expensive technology and specific bioinformatic expertise that can serve as a barrier to standard implementation. We propose that determination of amplicon sizes by capillary electrophoresis is a precise, less time-consuming, standardisable and affordable method and that it can replace complex massive sequencing technologies in routine studies of diet composition from fresh faecal samples, as has been previously pointed out [32,42]. As a general guideline, we recommend to start performing a botanical assessment of the study area to not only create a reference collection for CMA but also retrieve the corresponding species genomic sequences from the GenBank database. Then, if researchers are only interested in a qualitative assessment, PCR-CE should be favoured due to its lower time consumption and its higher accuracy to detect plants at the species level (e.g. *Festuca* spp.). On the other hand, CMA must be used when a quick, easy and unexpensive semi-quantitative assessment on field collected faecal samples is required [68].

Conclusions

Taken together, our results suggest that PCR-CE combining the simultaneous amplification of two genes—long *trnL(UAA)* and *ITS2*—is a reliable technique for the determination of diet composition in herbivores. However, while this DNA-based method is qualitative in our experimental conditions, CMA can provide semi-quantitative data, since individual percentages of ingested plant species can be calculated, although a differential index of plant preservation should be taken into account. Additionally, the experimental conditions of outdoor exposure acted concomitantly with proper digestion processes to trigger genomic DNA degradation. Consequently and as expected, faeces freshness and preservation conditions are critical and determinant factors that compromise downstream DNA amplification efficiency but are not limiting in the case of CMA. We conclude that CMA and PCR-CE have both strong and weak features that make the two approaches complementary. Therefore, their simultaneous use can provide more complete, accurate and thorough information in diet composition studies. However, neither CMA nor PCR-CE seems to solve comprehensively the quantification of herbivore diets and thus further research needs to be done.
Supporting information

S1 Dataset. Raw data.
(XLSX)

S1 Fig. (A) Picture of the four pools of faecal samples from the time-course degradation experiment. (B) Environmental conditions while faecal samples were left outdoor (red line: mean daily temperature; red dotted line: minimum daily temperature; red dashed line: maximum daily temperature; blue bars: daily cumulative rainfall).
(TIF)

S2 Fig. Migration patterns on a 1.5% agarose electrophoresis of trnL(UAA) and ITS2 PCR products corresponding to the reference plant database amplifications. Ae: Arrhenatherum elatius, Cv: Calluna vulgaris, Cc: Carex caryophyllea, Fe: Festuca eskia, Fg: Festuca gautieri, Fgl: Festuca glauca, Fv: Festuca violacea, Hn: Helianthemum nummularium, Hr: Hypochaeris radi-cata, Jt: Juncus trifidus, Ms: Medicago sativa, Ns: Nardus stricta, Ta: Trifolium alpinum, Vu: Vaccinium uliginosum. (-): non template negative PCR control. 1Kb ladder is a reference ladder marker.
(TIF)

S3 Fig. Electropherogram peaks of the 14 plant species included in this study, obtained after capillary electrophoresis of the fluoresce labelled PCR products corresponding to trnL(UAA) (blue) and ITS2 (green) multiplex amplification. Peak sizes are summarized in Table 2.
(TIF)

S4 Fig. Migration patterns on a 1.5% agarose electrophoresis of trnL(UAA) and ITS2 PCR products amplified from previous controlled diet faeces genomic DNA (Medicago sativa intake, Ms) and from faeces genomic DNA corresponding to diets 1 to 3. Composition of diets is detailed in Table 1. (-): non template negative PCR control. 1Kb ladder is a reference ladder marker.
(TIF)

S1 Table. Universal oligonucleotide sequences used for specific PCR amplification of chloroplastic trnL(UAA) and nuclear ITS2 regions. Fluorescence-labelling of oligonucleotides TRNL_D and S2F is indicated. Expected amplicon size is given in base pairs (bp).
(DOCX)

Acknowledgments

The authors would like to acknowledge the Servei de Genòmica i Bioinformàtica of the Universitat Autònoma de Barcelona (UAB) for performing capillary electrophoresis analysis and the Servei de Granges i Camps Experimentals of the UAB for the use of their facilities during the animal feeding trial. Special thanks to Carles Conejero for his dedication in caring for “Niebla”, our stabled chamois, and to Miroslaw Ratkiewicz and an anonymous reviewer for their helpful comments on an earlier version of this work.

Author Contributions

Conceptualization: Carme Espunya, Jordi Bartolomé, Emmanuel Serrano.

Data curation: Juan Antonio Calleja.
Formal analysis: Johan Espunyes, Carme Espunya, Sara Chaves, Juan Antonio Calleja, Jordi Bartolomé, Emmanuel Serrano.

Investigation: Jordi Bartolomé.

Methodology: Carme Espunya, Sara Chaves, Jordi Bartolomé.

Supervision: Carme Espunya, Sara Chaves, Juan Antonio Calleja, Jordi Bartolomé, Emmanuel Serrano.

Validation: Carme Espunya, Sara Chaves, Juan Antonio Calleja, Jordi Bartolomé, Emmanuel Serrano.

Visualization: Juan Antonio Calleja.

Writing – original draft: Johan Espunyes.

Writing – review & editing: Johan Espunyes, Carme Espunya, Juan Antonio Calleja, Jordi Bartolomé, Emmanuel Serrano.

References

1. Barboza PS, Parker KL, Hume ID, editors. Integrative wildlife nutrition. Integrative Wildlife Nutrition. Berlin, Heidelberg: Springer Berlin Heidelberg; 2009. https://doi.org/10.1007/978-3-540-87885-8

2. Illius AW. Linking functional responses and foraging behaviour to population dynamics. In: Danell K, Bergstrom R, Duncan P, Pastor J, editors. Large Herbivore Ecology, Ecosystem Dynamics and Conservation. Cambridge: Cambridge University Press; 2006. pp. 71–96. https://doi.org/10.1017/CBO9780511617461.005

3. Danell K, Bergstrom R, Duncan P, Pastor J, editors. Large Herbivore Ecology, Ecosystem Dynamics and Conservation. Cambridge: Cambridge University Press; 2006. https://doi.org/10.1017/CBO9780511617461

4. Virtanen R, Edwards GR, Crawley MJ. Red deer management and vegetation on the Isle of Rum. J Appl Ecol. 2002; 39: 572–583. https://doi.org/10.1046/j.1365-2664.2002.00734.x

5. Pastor J, Cohen Y, Hobbs NT. The roles of large herbivores in ecosystem nutrient cycles. In: Danell K, Bergstrom R, Duncan P, Pastor J, editors. Large Herbivore Ecology, Ecosystem Dynamics and Conservation. Cambridge: Cambridge University Press; 2006. pp. 289–325. https://doi.org/10.1017/CBO9780511617461.012

6. Pierce BM, Bleich VC, Bowyer RT. Social organization of mountain lions: does a land-tenure system regulate population size? Ecology. 2000; 81: 1533. https://doi.org/10.2307/177304

7. Velamazán M, San Miguel A, Escribano R, Perea R. Threatened woody flora as an ecological indicator of large herbivore introductions. Biodivers Conserv. 2017; 26: 917–930. https://doi.org/10.1007/s10531-016-1279-3

8. Brookshire JE, Kauffman BJ, Lytjen D, Otting N. Cumulative effects of wild ungulate and livestock herbivory on riparian willows. Oecologia. 2002; 132: 559–566. https://doi.org/10.1007/s00442-002-1007-4 PMID: 28547642

9. Tanentzap AJ, Burrows LE, Lee WG, Nugent G, Maxwell JM, Coomes DA. Landscape-level vegetation recovery from herbivory: progress after four decades of invasive red deer control. J Appl Ecol. 2009; 46: 1064–1072. https://doi.org/10.1111/j.1365-2664.2009.01683.x

10. Mishra C, Van Wieren SE, Ketner P, Heitkönig IMA, Prins HHTT. Trans-Himalaya. J Appl Ecol. 2004; 41: 344–354. https://doi.org/10.1111/j.0021-8901.2004.00885.x

11. Kowalczyk R, Taberlet P, Coissac E, Valentini A, Miquel C, Kamiński T, et al. Influence of management practices on large herbivore diet-Case of European bison in Bialowieza Primeval Forest (Poland). For Ecol Manage. 2011; 261: 821–828. https://doi.org/10.1016/j.foreco.2010.11.026

12. Kartzinel TR, Chen P a., Coverdale TC, Erickson DL, Kress WJ, Kuzmina ML, et al. DNA metabarcoding illuminates dietary niche partitioning by African large herbivores. Proc Natl Acad Sci. 2015; 112: 8019–8024. https://doi.org/10.1073/pnas.1503283112 PMID: 26034267

13. Bison M, Ibanez S, Redjadj C, Boyer F, Coissac E, Miquel C, et al. Upscaling the niche variation hypothesis from the intra- to the inter-specific level. Oecologia. 2015; 179: 835–842. https://doi.org/10.1007/s00442-015-3390-7 PMID: 26198049
14. Newmaster SG, Thompson ID, Steeves RADD, Rodgers AR, Fazekas AJ, Maioles JR, et al. Examination of two new technologies to assess the diet of woodland caribou: video recorders attached to collars and DNA barcoding. Can J For Res. 2013; 43: 897–900.

15. Gordon IJ. Animal-based techniques for grazing ecology research. Small Rumin Res. 1995; 16: 203–214. https://doi.org/10.1016/0921-4488(95)00635-X

16. Stewart DRM. Analysis of plant epidermis in faeces: a technique for studying the food preferences of grazing herbivores. J Appl Ecol. 1967; 4: 83–111. https://doi.org/10.2307/2401411

17. Anthony RG, Smith NS. Comparison of rumen and fecal analysis to describe deer diets. J Wildl Manage. 1974; 38: 535. https://doi.org/10.2307/3800866

18. Holechek JL, Vavra M, Pieper RD. Botanical composition determination of range herbivore diets: a review. J Range Manag. 1982; 35: 309. https://doi.org/10.2307/3898308

19. Rivera L, Baraza E, Alcover JA, Bover P, Rovira CM, Bartolomé J. Stomatal density and stomatal index. Newmaster SG, Thompson ID, Steeves RADD, Rodgers AR, Fazekas AJ, Maioles JR, et al. Examining the impact of climate change on the diet of woodland caribou. Can J For Res. 2013; 43: 897–900.

20. Alipayo D, Valdez R, Holechek JL, Cardenas M. Evaluation of microhistological analysis for determining ruminant diet botanical composition. J Range Manag. 1992; 45: 148. https://doi.org/10.2307/4002773

21. Hajibabaei M, Singer GAC, Hebert PDN, Hickey DA. DNA barcoding: how it complements taxonomy, molecular phylogenetics and population genetics. Trends Genet. 2007; 23: 167–172. https://doi.org/10.1016/j.tig.2007.02.001 PMID: 17316686

22. Angers-Loustau A, Petrillo M, Paracchini V, Kagkli DM, Rischitor PE, Puertas Gallardo A, et al. Towards plant species identification in complex samples: a bioinformatics pipeline for the identification of novel nuclear barcode candidates. PLoS One. 2016; 11: e0147692. https://doi.org/10.1371/journal.pone.0147692 PMID: 26807711

23. Coissac E, Hollingsworth PM, Lavergne S, Taberlet P. From barcodes to genomes: extending the concept of DNA barcoding. Mol Ecol. 2016; 25: 1423–1428. https://doi.org/10.1111/mec.13549 PMID: 26821259

24. Newmaster SG, Grguric M, Shanmughanandhan D, Ramalingam S, Ragupathy S. DNA barcoding detects contamination and substitution in North American herbal products. BMC Med. 2013; 11: 222. https://doi.org/10.1186/1741-7015-11-222 PMID: 24120035

25. Parveen I, Gafner S, de Vries H, Boessenkool S, Coissac E, et al. Fungal paleodiversity revealed using high-throughput metabarcoding of ancient DNA from arctic permafrost. Environ Microbiol. 2013; 15: 1176–1189. https://doi.org/10.1111/1462-2920.12020 PMID: 23171292

26. Hofreiter M, Poinar HN, Spaulding WG, Bauer K, Martin PS, Possnert G, et al. A molecular analysis of ground sloth diet through the last glaciation. Mol Ecol. 2000; 9: 1975–1984. https://doi.org/10.1046/j.1365-294X.2000.01106.x PMID: 11123610

27. Haile J, Holdaway R, Oliver K, Bunce M, Gilbert MTP, Nielsen R, et al. Ancient DNA chronology within sediment deposits: are paleobiological reconstructions possible and is DNA leaching a factor? Mol Biol Evol. 2007; 24: 982–989. https://doi.org/10.1093/molbev/msm106 PMID: 17255121

28. Bellemain E, Davey ML, Kauserud H, Epp LS, Boessenkool S, Coissac E, et al. Fungal paleodiversity revealed using high-throughput metabarcoding of ancient DNA from arctic permafrost. Environ Microbiol. 2013; 15: 1176–1189. https://doi.org/10.1111/1462-2920.12020 PMID: 23171292

29. Soininen EM, Valenti A, Coissac E, Miquel C, Gielly L, Brochmann C, et al. Analysing diet of small herbivores: the efficiency of DNA barcoding coupled with high-throughput pyrosequencing for deciphering the composition of complex plant mixtures. Front Zool. 2009; 6: 16. https://doi.org/10.1186/1746-135X-6-16 PMID: 19695081

30. Valenti A, Miquel C, Nawaz MA, Bellemain E, Coissac E, Pompanon F, et al. New perspectives in diet analysis based on DNA barcoding and parallel pyrosequencing: The trnL approach. Mol Ecol Resour. 2009; 9: 51–60. https://doi.org/10.1111/j.1755-0998.2008.02352.x PMID: 21564566

31. Rayé G, Miquel C, Coissac E, Redjaï C, Loison A, Taberlet P. New insights on diet variability revealed by DNA barcoding and high-throughput pyrosequencing: chamois diet in autumn as a case study. Ecol Res. 2011; 26: 265–276. https://doi.org/10.1007/s11284-010-0780-5

32. Czernik M, Taberlet P, Świslocka M, Czajkowska M, Duda N, Ratkiewicz M. Fast and efficient DNA-based method for winter diet analysis from stools of three cervids: moose, red deer, and roe deer. Acta Theriol. 2013; 58: 379–386. https://doi.org/10.1007/s13364-013-0146-9 PMID: 24244041

33. Hibert F, Taberlet P, Chave J, Scotti-Saintagne C, Sabatier D, Richard-Hansen C. Unveiling the diet of elusive rainforest herbivores in next generation sequencing era? The tapir as a case study. PLoS One. 2013; 8: e60799. https://doi.org/10.1371/journal.pone.0060799 PMID: 23560107
34. Burkepile DE, Parker JD. Recent advances in plant-herbivore interactions. F1000Research. 2017; 6: 119. https://doi.org/10.12688/f1000research.10313.1 PMID: 28323868

35. Staats M, Arulandhu AJ, Gravendeel B, Holst-Jensen A, Scholtens I, Peelen T, et al. Advances in DNA metabarcoding for food and wildlife forensic species identification. Anal Bioanal Chem. 2016; 408: 4615–4630. https://doi.org/10.1007/s00216-016-9595-8 PMID: 27178552

36. Hollingsworth PM, Graham SW, Little DP. Choosing and using a plant DNA barcode. PLoS One. 2011; 6: e19254. https://doi.org/10.1371/journal.pone.0019254 PMID: 21637336

37. Shokralla S, Spall JL, Gibson JF, Hajibabaei M. Next-generation sequencing technologies for environmental DNA research. Mol Ecol. 2012; 21: 1794–1805. https://doi.org/10.1111/j.1365-294X.2012.05538.x PMID: 22486820

38. Pompon F, Deagle BE, Symondson WOC, Brown DS, Jarman SN, Taberlet P. Who is eating what: diet assessment using next generation sequencing. Mol Ecol. 2011; 6: e19254. https://doi.org/10.1371/journal.pone.0019254 PMID: 21637336

39. Yao H, Song J, Liu C, Luo K, Han J, Li Y, et al. Use of ITS2 region as the universal DNA barcode for plants and animals. PLoS One. 2010; 5: e13102. https://doi.org/10.1371/journal.pone.0013102 PMID: 20957043

40. Chen S, Yao H, Han J, Liu C, Song J, Shi L, et al. Validation of the ITS2 Region as a Novel DNA Barcode for Identifying Medicinal Plant Species. PLoS One. 2010; 5: e8613. https://doi.org/10.1371/journal.pone.0008613 PMID: 20062805

41. Hollingsworth PM, Forrest LL, Spouge JL, Hajibabaei M, Ratnasingham S, van der Bank M, et al. A DNA barcode for land plants. Proc Natl Acad Sci. 2009; 106: 12794–12797. https://doi.org/10.1073/pnas.0905845106 PMID: 19666222

42. Pegard A, Miquel C, Valentini A, Coissac E, Bouvier F, François D, et al. Universal DNA-based methods for assessing the diet of grazing livestock and wildlife from feces. J Agric Food Chem. 2009; 57: 5700–5706. https://doi.org/10.1021/jf803680c PMID: 19566081

43. NCBI. Database resources of the National Center for Biotechnology Information. Nucleic Acids Res. 2016; 44: D7–D19. https://doi.org/10.1093/nar/gkv1290 PMID: 26615191

44. Bazakos C, Dulger AO, Uncu AT, Spaniolas S, Spano T, Kalaitzis P, A SNP-based PCR–RFLP capillary electrophoresis analysis for the identification of the varietal origin of olive oils. Food Chem. 2012; 134: 2411–2418. https://doi.org/10.1016/j.foodchem.2012.04.031 PMID: 23442703

45. Spaniolas S, May ST, Bennett MJ, Tucker GA. Authentication of coffee by means of PCR-RFLP analysis and lab-on-a-chip capillary electrophoresis. J Agric Food Chem. 2006; 54: 7466–7470. https://doi.org/10.1021/jf061164n PMID: 17002409

46. Nichols RV, Åkesson M, Kjellander P. Diet assessment based on rumen contents: a comparison between DNA metabarcoding and macroscopy. PLoS One. 2016; 11: e0157977. https://doi.org/10.1371/journal.pone.0157977 PMID: 27322387

47. Taberlet P, Coissac E, Pompon F, Gielly L, Miquel C, Valentini A, et al. Power and limitations of the chloroplast trnL (UAA) intron for plant DNA barcoding. Nucleic Acids Res. 2007; 35: e14–e14. https://doi.org/10.1093/nar/gkl938 PMID: 17169982

48. Deagle BE, Eveson JP, Jarman SN. Quantification of damage in DNA recovered from highly degraded samples—A case study on DNA in faeces. Front Zool. 2006; 3: 1–10.

51. Zuur AF, Ieno EN, Elphick CS. A protocol for data exploration to avoid common statistical problems. Methods Ecol Evol. 2010; 1: 3–14. https://doi.org/10.1111/j.2041-210X.2009.00001.x

52. Zar J. Biostatistical analysis. 5th ed. Pearson Education Limited; 2010.

53. Johnson MK, Wofford HH, Pearson HA. Microhistological techniques for food habits analyses. Res. Pap. SO-199. USDA For Serv. 1983; 1–40.

54. Bartolome J, Franch J, Plaikats J, Seligman NG. Diet selection by sheep and goats on Mediterranean heath-woodland range. J Range Manag. 1998; 51: 383. https://doi.org/10.2307/4003322
58. Marinas A, García-González R. Preliminary data on nutritional value of abundant species in suprafores-tal pyrenean pastures. Pirineos. 2006; 161: 85–109.

59. Hayes F, Mills G, Jones L, Abbott J, Ashmore M, Barnes J, et al. Consistent ozone-induced decreases in pasture forage quality across several grassland types and consequences for UK lamb production. Sci Total Environ. 2016; 543: 336–346. https://doi.org/10.1016/j.scitotenv.2015.10.128 PMID: 26595401

60. Heggerget TM, Gaare E, Ball JP. Reindeer (Rangifer tarandus) and climate change: Importance of winter forage. Rangifer. 2002; 22: 13. https://doi.org/10.7557/2.22.1.388

61. Asshoff R, Hättenschwiler S. Growth and reproduction of the alpine grasshopper Miramella alpina feeding on CO₂-enriched dwarf shrubs at treeline. Oecologia. 2005; 142: 191–201. https://doi.org/10.1007/s00442-004-1714-0 PMID: 15480804

62. Hendriks RJJ, de Boer NJ, van Groenendael JM. Comparing the preferences of three herbivore species with resistance traits of 15 perennial dicots: the effects of phylogenetic constrains. Plant Ecol. 1999; 143: 141–152. https://doi.org/10.1023/A:1009832621516

63. Carrière S. Photographic key for the microhistological identification of some Arctic vascular plants. Arctic. 2002; 55: 247–268. https://doi.org/10.14430/arctic709

64. Mohammad AG, Pieper RD, Wallace JD, Holechek JL, Murray LW. Comparison of fecal analysis and rumen evacuation techniques for sampling diet botanical composition of grazing cattle. J Range Manag. 1995; 48: 202. https://doi.org/10.2307/4002421

65. Vavra M, Holechek JL. Factors Influencing Microhistological Analysis of Herbivore Diets. J Range Manag. 1980; 33: 371. https://doi.org/10.2307/3897886

66. Leslie DM, Vavra M, Starkey EE, Slater RC. Correcting for differential digestibility in microhistological analyses involving common coastal forages of the Pacific Northwest. J Range Manag. 1983; 36: 730. https://doi.org/10.2307/3898197

67. King RA, Read DS, Traugott M, Symondson WOC. Molecular analysis of predation: a review of best practice for DNA-based approaches. Mol Ecol. 2008; 17: 947–963. PMID: 18208490

68. Espunyes J, Bartolomé J, Garei M, Gálvez-Cerón A, Aguilar XF, Colom-Cadena A, et al. Seasonal diet composition of Pyrenean chamois is mainly shaped by primary production waves. PLoS One. 2019; 14: 1–23. https://doi.org/10.1371/journal.pone.0210819 PMID: 30673575