Invasive Plants Are a Valuable Alternate Protein Source and Can Contribute to Meeting Climate Change Targets

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Agriculture has come under pressure to meet global food demands, whilst having to meet economic and ecological targets. This has opened newer avenues for investigation in unconventional protein sources. Current agricultural practises manage marginal lands mostly through animal husbandry, which; although effective in land utilisation for food production, largely contributes to global green-house gas (GHG) emissions. Assessing the revalorisation potential of invasive plant species growing on these lands may help encourage their utilisation as an alternate protein source and partially shift the burden from livestock production; the current dominant source of dietary protein, and offer alternate means of income from such lands. Six globally recognised invasive plant species found extensively on marginal lands; Gorse (*Ulex europaeus*), Vetch (*Vicia sativa*), Broom (*Cytisus scoparius*), Fireweed (*Chamaenerion angustifolium*), Bracken (*Pteridium aquilinum*), and Buddleia (*Buddleja davidii*) were collected and characterised to assess their potential as alternate protein sources. Amino acid profiling revealed appreciable levels of essential amino acids totalling 33.05 ± 0.04 41.43 ± 0.05, 33.05 ± 0.11, 32.63 ± 0.04, 48.71 ± 0.02 and 21.48 ± 0.05 mg/g dry plant mass for Gorse, Vetch, Broom, Fireweed, Bracken, and Buddleia, respectively. The availability of essential amino acids was limited by protein solubility, and Gorse was found to have the highest soluble protein content. It was also high in bioactive phenolic compounds including cinnamic-, phenyl-, pyruvic-, and benzoic acid derivatives. Databases generated using satellite imagery were used to locate the spread of invasive plants. Total biomass was estimated to be roughly 52 Tg with a protein content of 5.2 Tg with a total essential amino acid content of 1.25 Tg (∼24%). Globally, Fabaceae was the second most abundant family of invasive plants. Much of the spread was found within marginal lands and shrublands. Analysis of intrinsic agricultural factors revealed economic status as the emergent factor, driven predominantly by land use allocation, with shrublands playing a pivotal role in the model. Diverting resources from invasive plant removal through herbicides and burning to leaf protein extraction may contribute toward sustainable protein, effective land use, and achieving emission targets, while simultaneously maintaining conservation of native plant species.

Keywords: plant protein, marginal lands, sustainable agriculture, net zero emissions, nutritional characterisation, invasive plants, essential amino acids, cyclic economy
INTRODUCTION

Existing mainstream food supply systems are capable of surpassing global nutritional requirements and yet, we find about 12.3% of the human population living with chronic malnutrition (FAO et al., 2020). Increased atmospheric CO₂ and malnutrition can be causally attributed to inefficiencies along the food supply chain primarily through wastage and poor resource allocation (Bajželj et al., 2014; Sabaté et al., 2015; Xue et al., 2017). The high-energy input required for modern agriculture is mostly sourced from fossil fuels, which directly leads to greenhouse gas emissions. This in turn contributes to climate change, increasing crop vulnerability due to altered pest behaviour, temperature eccentricities, and altered seasonal patterns. Cultivation of crops and plants, which is essentially an anabolic carbon storing process, is rendered ineffective owing to such intensive cultivation practices to meet global food demand. One of the major sources of food-based carbon emission is animal husbandry. Most of the global protein produced is directed toward animal feed. Currently, animal husbandry enjoys the single largest resource allocation; especially land, in global food production and is the dominant source of protein for most people (Day, 2013; FAO, 2019). Justification for the present scale of animal husbandry enumerates reasons such as employment, superior essential amino acid profile of animal products, useful functional properties and gastronomic preferences. However, protein turnover from this enterprise is low, while associated greenhouse gas (GHG) emissions are high, making it the single largest contributing factor to agricultural emissions (Pragna et al., 2018).

In the case of Scotland, large parts of the Scottish Highlands are ill suited for arable farming and are primarily used for rough grazing. Plants such as Gorse (Ulex europaeus), Vetch (Vicia sativa), Broom (Cytisus scoparius), Fireweed (Chamaenerion angustifolium), Bracken (Pteridium aquilinum), and Buddleia (Buddleja davidii) are able to quickly grow in urban as well as marginal lands owing to their robust physiologies and quicker germination times, allowing them to dominate available land and often encroach into farmlands (Eldridgea et al., 2011). Since they can grow in low nutrient conditions, they are known to colonise lands left fallow for rejuvenation. This necessitates the investment of resources to remove such plants. While historically, the excised mass was used for animal fodder during times of famine (Łuczaj, 2010; Pinela et al., 2017), they are currently dried and burnt, dug into the soil for increased organic matter or discarded into the landfill which contributes to carbon emissions. Investigation into their nutritive value and developing efficient protein extraction processes may help solve the problem of resource waste and add potential means of compensation for the investment made toward their active removal. Numerous large-scale leaf protein extraction designs have been proposed, although commercial impetus is lacking owing to efforts being diverted to crop variety selection and design. However, adopting such valorization strategies may mitigate costs generally incurred from raw material sourcing while simultaneously dealing with downstream waste-management. Currently, the resources dedicated to unwanted plant removal is substantial and costs are borne entirely by local farmers, impinging on their income margins. Additionally, these plants pose a threat to local diversity if not kept in check (Parker, 2000; Dassonville et al., 2008; Shiferaw et al., 2019).

Characterisation of these plants as a potential source of nutrition may help provide impetus for their effective utilisation and limit or decrease their spread in their non-native locations or stimulate interest for their conservation in native lands. Extraction methods for obtaining leaf proteins have been well-studied; particularly by Pirie (1977) who pioneered the use of industrial pulping mechanisms to express plant juices. However, contemporary methods for removal of punget and strong-tasting phenolics were not cost effective and subsequent advances made in crop genetics, and development of novel cultivars overshadowed the idea. Recent years, however, have seen an increase in the pursuit of leaf protein recovery, particularly from agricultural waste. Understanding the total protein content and the amino acid composition of native Scottish plants may help open newer avenues for protein sourcing and allow for more robust food systems resilient to global policies and environment.

Improved utilisation and reallocation of land use for alternate and sustainable protein production is required to help reduce dependence on animal husbandry and offset part of the associated emissions. Along with other non-conventional sources such as insects (Henchion et al., 2017), fungi (Bano et al., 1963), and seaweed (Tamayo Tenorio et al., 2018), revalorisation of invasive plants appears to be an abundant and a potentially lucrative route of obtaining protein (Pirie, 1932, 1969a,b; Nagy et al., 1978). The work described here assesses the revalorisation potential of invasive plants through comprehensive nutritional investigation. It considers potential health benefits, and the predicted impact on land resource allocation and associated GHG emissions.

MATERIALS AND METHODS

World data on land use and green-house-gas emissions were obtained from FAOSTAT (FAO, 2019).

All chemicals and kits were purchased from Merck (Darmstadt, Germany), and used without further purification unless stated otherwise.

All experiments and measurements were performed in triplicate.

All colorimetric assays were carried out in 96-well-plates and incubated with film cover (Greiner Bio-One, Kremsmuenster, Austria). Absorption values were obtained using SpectraMax 190 (Molecular Devices, San Jose, USA).

Plant Samples

All plant samples were collected between March and July in the North-East of Scotland at the following GPS co-ordinates. Gorse (Ulex europaeus), Broom (Cytisus scoparius), and Bracken (Pteridium aquilinum); 57.257, −2.483, Fireweed (Chamaenerion angustifolium); 57.157, −2.086, Vetch (Vicia sativa); 57.157, −2.139, and Buddleia (Buddleja davidii); 57.158, −2.099. Collection procedure was in accordance to guidelines by the Ministry of Forest, CA (British Columbia Ministry of Forest,
Plant Homogenisation and Extraction

The leaf samples (10 mg) were vortexed in phosphate buffer (10 mL; 10 mM; pH 7.5) at 25°C and kept in an ultrasonic bath (Bransen, B12) for 2 min. The supernatant was separated by centrifugation (4°C; 15 min; 4,000 g). A subsample of the aqueous supernatant extract was used for colorimetric estimation of soluble phenolics, sugar and protein content. Remainder of the supernatant and precipitate were collected and freeze-dried. Freeze-dried precipitate was used to estimate bound proteins.

Total Soluble Phenolics

Fast Blue assay was performed to estimate total soluble phenolics as described by Lester et al. (2012). Sample or Standard (100 µL) was incubated with 2 µL Fast Blue (0.1% w/v) for 30 s on a shaker. Sodium hydroxide (NaOH; 2 µL; 5% w/v) was added and incubated for 90 min in the dark. Absorbance was measured at 420 nm.

Total Soluble Sugars

Lever's assay was used to estimate soluble sugars as previously described by Lever (1973, 1977). Lever reagent comprised of 4-hydroxybenzoic acid hydrazide (PAHBAH, 0.76% w/v), bismuth (III) nitrate pentahydrate (0.48% w/v), potassium sodium tartrate (0.28% w/v), NaOH (2% w/v) in ultrapure Milli-Q water. Sample or standard (5 µL) was incubated with 200 µL of Lever reagent at 70°C for 30 min. The samples were then allowed to cool for 10 min and absorbance was checked at 415 nm.

Protein Estimation

Sample hydrolysis was carried out using HCl (6 N) with phenol (2% w/v) to aid removal of oxygen radicals (Muramoto et al., 1987). Freeze-dried supernatant and retentate (10 mg) were digested in HCl (6 N; 10 mL, prepared as described above) and digested using MARS 6 (CEM, US) microwave digester with a temperature ramp to 150°C for 20 min. The temperature was maintained for 60 min and then cooled overnight.

Protein content was determined by amino acid estimation using the Ninhydrin Assay based on the work described by Harding and MacLean (1916). An aliquot (10 µL) of the supernatant was dried at 60°C in a microtiter plate and resuspended in phosphate buffer (10 mM, 100 µL, pH 7.5) to which ninhydrin reagent (75 µL) was added and incubated at 60°C for 60 min. The absorbance was measured at 570 nm.

Amino Acid Estimation

Hydrolysis for amino acid profiling was performed as previously described for protein estimation with the addition of 10 mg of tryptamine. Essential amino acids were profiled and quantified using Gas Chromatography in tandem with Mass Spectrometry (GC-MS) with U13-C amino acids as internal standards as described previously in Calder et al. (1999).

Phenolic Characterisation

Phenolics were characterised in a three-stage extraction process. Dried sample (5 mg) of freeze-dried, freeze-milled leaf samples were sequentially extracted under acidic, alkali, and finally neutral pH conditions using biphasic solvent extraction with ethyl acetate as the organic layer. The aqueous layer was adjusted to the required pH condition using 1 M HCl or 1 N NaOH.

Quantification was performed using LC-MS as previously described without modifications (Neacsu et al., 2015). Internal standard used were 12C benzoic acid; 2 ng/µL in 0.02% acetic acid in 75% methanol for negative mode MS and 2-amino-3,4,7,8-tetramethylimidazo[4,5-f] quinoxaline; 0.5 ng/µL in acetic acid (0.02% v/v) in methanol (75% v/v) for positive mode MS.

Non-starch Polysaccharide (NSP) Determination

Non-starch polysaccharides were estimated as described by Englyst et al. using GC with inositol as an internal standard (Englyst and John, 1984; Englyst and Hudson, 1996; Bach Knudsen et al., 1997). Briefly, plant samples were hydrolysed in H2SO4 (7 M) at 100°C for 1 h. Monosaccharides were analysed by Gas Chromatography with Flame Ionised Detection (GC-FID) using Inositol as internal standard.

Lignin Content

Lignin content was assayed using the Acetyl Bromide method (Barnes and Anderson, 2017). The protocol was modified to improve carbohydrate removal using enzymes (Hatfield et al., 1999; Fukushima and Hatfield, 2001; Hatfield and Fukushima, 2005; Hojilla-Evangelista et al., 2017). Briefly, the freeze-dried original plant samples (100 mg) were washed with phosphate buffer (10 mM; 10 mL, pH 7.5) at 35°C twice and suspended in citric acid buffer (50 mM, 20 mL, pH 4.9) with 20 units each of cellulase, pectinase and xylanase (all three from Sigma-Aldrich) at 40°C for 4 h. Samples were centrifuged at 4,000 g for 15 min. Retentate was recovered and resuspended in citric acid buffer and incubated with Visozyme® (Novozymes, Denmark) overnight at 32°C. Samples were centrifuged again under the same condition and the supernatant was discarded. Retentate was washed twice with 5 mL ethanol at 30°C, followed by 10 mL of 90% aqueous DMSO at 40°C. Sample was finally washed with 10 mL acetone at 30°C and allowed to dry overnight in vacuo. Dried sample (5 mg) was suspended in acetyl bromide (5% v/v) prepared in glacial acetic acid and incubated for 2 h at 70°C. Samples were left overnight in the dark under ambient temperature. The absorbance of the supernatant was measured at 280 nm. Absorption coefficient was assumed to be 23.6 g/cm/mL.
Statistics and Data Analysis
Statistical analysis was performed using R (Version 4.0.2) and RStudio (Version 1.2.1335). Biological values were tested using one-way (OW)-ANOVA followed by the Tukey-HSD post-hoc test at 99% confidence interval. Significance is expressed as $F$ (degrees of freedom, residuals) = $F$-value, $p$-value as recommended by Field et al. (2012) unless stated otherwise. All graphs and visualisation have been made using ggplot2 (Wickham, 2016).

GPS location was downloaded from the GBIF database. Identifier of the dataset used to generate maps and analysis can be found at https://doi.org/10.15468/dl.tgk8j9. Packages used in R were: "dplyr" (Wickham et al., 2020), "ggspatial" (Dunnington, 2020), "ggrepel" (Slowikowski, 2020), "sf" (Pebesma, 2018), and "maturaleza" (South, 2017).

Principal Component Analysis (PCA) of the phenolic composition was performed on logarithmically transformed data using packages factoextra (Kassambara and Mundt, 2019) and FactoMineR (Lê et al., 2008). Diagnostics of loadings and contribution to the PCA analysis was performed with the help of the package corrplot (Wei and Simko, 2017).

The database on world agricultural values, obtained from FAOSTAT was subjected to exploratory factor analysis using the package psych (Revelle, 2020). Further confirmatory analysis was performed using lavaan (Rosseel, 2012) and lavaanPlot (Lishinski, 2018). Univariate PCA was performed using factoextra, to visualise overarching trends across countries under reduced dimensions. From the database, general regions such as “OECD,” “South-east Asia,” and similar classifications in the database were removed. Finally, all measures of land use were made intrinsic for a country by calculating its percentage coverage relative to its geopolitical land area. A naïve approach was adopted toward factor selection where percentage deviation was used as the metric of choice which showed high variation across individual countries. Factors with a percentage deviation value >100 were chosen for further exploratory factor analysis using maximum likelihood (ML) factor reduction to understand correlation to the latent factors. Finally, a System Equation Mode (SEM) was constructed based on the correlation using confirmatory analysis. Income group classification was performed according to data from World Bank (2020).

RESULTS
Nutritional Characterisation of Six Invasive Plant Species
Six invasive plants (Gorse, Vetch, Broom, Fireweed, Bracken, and Buddleia) were characterised for total protein content, amino acid profile, non-starch polysaccharide (NSP) composition, and phenolic composition to assess their potential for revalorisation.

Protein and Amino Acids
Across the plant samples, Bracken was found to have total protein content significantly greater than other plant samples as shown in Table 1. A one way (OW)-ANOVA test at a 99% confidence interval across plant samples for total protein content was found to be significant $F_{(5,12)} = 725.3; p = 1.88 \times 10^3$, but similarities were revealed using Tukey-HSD post-hoc test between Gorse and Broom ($p = 0.10$), and Vetch and Fireweed ($p = 0.69$). Soluble protein represents the protein fraction extractable using mechanical grinding and buffer solubilisation which ANOVA deemed to have significant overall distinction $F_{(5,12)} = 1,208; p = 8.9 \times 10^{-16}$. A post-hoc test, however, could not distinguish between Gorse and Bracken at a 99% confidence interval ($p = 0.03$). Gorse had 47.3 ± 1.2 mg/g soluble protein (40.8% total protein), while Bracken had 45.2 ± 0.5 mg/g soluble protein (31.6% of total protein). The insoluble protein indicates proteins which remain bound to the cell wall and require further effort for extraction to efficiently use the biomass. ANOVA test found significant difference across the plants $F_{(5,12)} = 1,256; p = 4.45 \times 10^{-16}$, although a post-hoc test could not distinguish between Buddleia and Fireweed ($p = 0.96$).

In Table 2, the amino acid composition of the plants is shown. The difference in amino acid profile was found to be statistically significant $F_{(5,318)} = 4.645; p = 4.17 \times 10^{-4}$ across plants, but a post-hoc test found Buddleia as the influencing component with a difference which was significant at a 99% confidence interval from Broom ($p = 0.005$) and Bracken ($p = 2.1 \times 10^{-5}$) and at a 95% confidence interval from Fireweed ($p = 0.040$) and Gorse ($p = 0.010$). Statistical difference was insignificant between the other five plants. Similarities in individual essential amino acid content at 99% confidence interval is also shown in Table 2. Total essential amino acid content of the leaf mass was found to be similar between Gorse and Broom ($p = 0.99$).

The total essential amino acid content of Gorse, Vetch, Broom, Fireweed, Bracken and Buddleia are 33.05 ± 0.04, 41.43 ± 0.05, 33.05 ± 0.11, 32.63 ± 0.04, 48.71 ± 0.02, and 21.48 ± 0.05 mg/g dry plant mass. When this is presented for % of total protein, the essential amino acid content was the highest for Vetch and appeared to be unique across the plants $F_{(5,12)} = 259.1; p = 8.67 \times 10^{-12}$ using OW-ANOVA. No statistical difference was found between Bracken and Buddleia ($p = 0.015$), and Gorse and Bracken ($p = 0.013$) at 99% confidence interval.

Non-protein Components
Additional plant components such as carbohydrate and non-nutrient plant bio-actives have potential use as food or feed and could provide health benefits or wider uses as

| Table 1 | Total, soluble, and bound protein in Gorse, Vetch, Broom, Fireweed, Bracken, and Buddleia. |
|---------|---------------------------------|
| Plant   | Total  | Soluble  | Insoluble |
| Gorse   | 115.9 ± 0.8^a | 47.3 ± 1.2^b | 63.3 ± 0.3 |
| Vetch   | 106.7 ± 0.3^a | 16.9 ± 0.8 | 99.1 ± 0.4 |
| Broom   | 112.3 ± 1.5^a | 40.3 ± 0.1 | 90.5 ± 0.8 |
| Fireweed| 105.2 ± 1.4^a | 24.0 ± 1.0 | 78.9 ± 1.1^b |
| Bracken | 142.8 ± 0.9 | 45.2 ± 0.5^a | 107.9 ± 0.6 |
| Buddleia| 81.4 ± 0.5 | 7.1 ± 1.1 | 78.1 ± 1.4^a |

Data is expressed as mean ± standard deviation in mg/g dry plant mass. Values with same superscript were statistically similar for a category across plant species using OW-ANOVA at 99% confidence interval with Tukey-HSD post-hoc test.
TABLE 2 | Amino acid content of Gorse, Vetch, Broom, Fireweed, Bracken, and Buddleia.

|       | Gorse       | Vetch       | Broom       | Fireweed       | Bracken       | Buddleia       |
|-------|-------------|-------------|-------------|----------------|---------------|----------------|
| His   | 1.54 ± 0.12 | 1.85 ± 0.04 | 2.24 ± 0.01 | 1.84 ± 0.02    | 2.64 ± 0.03   | 1.06 ± 0.11    |
| Ile   | 2.64 ± 0.01 | 3.69 ± 0.02 | 1.90 ± 0.01 | 1.61 ± 0.00    | 3.42 ± 0.00   | 1.72 ± 0.01    |
| Leu   | 7.84 ± 0.19  | 10.18 ± 0.01| 8.03 ± 0.01 | 8.48 ± 0.00    | 11.61 ± 0.01  | 5.39 ± 0.00    |
| Lys   | 6.80 ± 0.11  | 6.83 ± 0.02 | 6.97 ± 0.01 | 6.54 ± 0.00    | 9.22 ± 0.01   | 3.96 ± 0.01    |
| Met   | 1.38 ± 0.03  | 1.80 ± 0.02 | 1.14 ± 0.00 | 1.25 ± 0.01    | 2.21 ± 0.02   | 1.09 ± 0.02    |
| Phe   | 4.53 ± 0.02  | 6.10 ± 0.03 | 4.86 ± 0.01 | 5.27 ± 0.00    | 7.02 ± 0.02   | 3.07 ± 0.01    |
| Thr   | 4.17 ± 0.02  | 5.34 ± 0.03 | 3.45 ± 0.01 | 3.28 ± 0.02    | 6.20 ± 0.02   | 2.66 ± 0.01    |
| Trp   | 0.42 ± 0.03  | 0.59 ± 0.15 | 1.71 ± 0.33 | 1.65 ± 0.11    | 1.64 ± 0.02   | 0.21 ± 0.10    |
| Val   | 3.75 ± 0.01  | 5.05 ± 0.00 | 2.73 ± 0.00 | 2.71 ± 0.00    | 4.75 ± 0.00   | 2.32 ± 0.01    |
| Ala   | 7.69 ± 0.02  | 7.98 ± 0.01 | 8.53 ± 0.00 | 9.45 ± 0.01    | 11.19 ± 0.02  | 4.53 ± 0.01    |
| Arg   | 6.13 ± 0.02  | 6.81 ± 0.02 | 5.84 ± 0.02 | 6.27 ± 0.00    | 8.02 ± 0.01   | 3.73 ± 0.07    |
| Asp   | 20.24 ± 01   | 11.04 ± 01  | 21.75 ± 01  | 11.77 ± 05     | 20.21 ± 09    | 7.32 ± 03      |
| Cys   | 0.45 ± 0.13  | 0.24 ± 0.00 | 0.46 ± 0.01 | 0.45 ± 0.01    | 0.54 ± 0.00   | 0.19 ± 0.00    |
| Glu   | 11.54 ± 0.06 | 14.90 ± 0.04| 12.47 ± 0.04| 13.28 ± 0.04   | 18.68 ± 0.02  | 7.67 ± 0.12    |
| Gly   | 8.13 ± 0.08  | 7.66 ± 0.06 | 9.96 ± 0.02 | 10.47 ± 0.01   | 12.36 ± 0.02  | 4.72 ± 0.02    |
| Pro   | 7.57 ± 0.01  | 6.24 ± 0.01 | 5.90 ± 0.00 | 5.50 ± 0.01    | 8.42 ± 0.01   | 3.61 ± 0.01    |
| Ser   | 8.16 ± 0.05  | 6.93 ± 0.10 | 7.82 ± 0.02 | 7.06 ± 0.07    | 10.51 ± 0.03  | 3.85 ± 0.03    |
| Tyr   | 4.55 ± 0.01  | 5.30 ± 0.01 | 4.54 ± 0.01 | 4.48 ± 0.01    | 7.22 ± 0.01   | 2.85 ± 0.01    |
| Total | 107.51 ± 0.21| 108.54 ± 0.21| 110.32 ± 0.36| 101.35 ± 0.15 | 145.86 ± 0.11 | 59.93 ± 0.37   |

Data is expressed as mean ± standard deviation in mg/g dry plant mass. Histidine to Valine represent essential dietary amino acids where values with same superscript were statistically similar using CW-ANOVA at 99% confidence interval with Tukey-HSD post-hoc test. The * mark represents the significantly lower total amino acid value obtained for Buddleia compared to other plants.

TABLE 3 | Summary of proximate values for phenolic and carbohydrate content.

| Plant | Carbohydrate | Phenolic |
|-------|--------------|----------|
|       | Soluble      | NSP      | Soluble | Lignin |
| Gorse | 26.1 ± 0.7c  | 172.3 ± 9.0 | 25.5 ± 6.8 | 83.7 ± 4.5 |
| Vetch | 53.9 ± 2.0d  | 129.5 ± 3.2 | 29.4 ± 4.8 | 11.7 ± 1.8 |
| Broom | 32.5 ± 3.2c  | 163.3 ± 6.5 | 96.8 ± 18.1 | 64.4 ± 1.3 |
| Fireweed | 54.2 ± 2.3d | 82.8 ± 7.5 | 28.9 ± 3.0 | 35.7 ± 1.2 |
| Bracken | 28.4 ± 3.9f | 172.1 ± 6.3 | 26.9 ± 5.1 | 78.8 ± 2.8 |
| Buddleia | 50.0 ± 2.0d | 112.3 ± 5.3 | 54.1 ± 5.7 | 48.1 ± 7.1 |

Results are presented in mg/g dry plant mass. Values with the same superscript are statistically similar using CW-ANOVA at 99% confidence interval with Tukey-HSD post-hoc test. NSP, non-starch polysaccharides.

Biomaterials contributing to an effective cyclic economy. Table 3 summarises proximate values of non-protein components and their distribution in the aqueous soluble and non-soluble fractions measured across the six leaf samples.

Correlation analysis across all measurements given in Tables 1 and 3 revealed a strong relation between soluble carbohydrate and lignin ($r^2 = -0.89$, $p = 5.52 \times 10^{-7}$), soluble protein and lignin ($r^2 = 0.78$, $p = 1.43 \times 10^{-4}$), lignin and NSP ($r^2 = 0.70$, $p = 1.15 \times 10^{-3}$), soluble carbohydrate and NSP ($r^2 = -0.87$, $p = 3.64 \times 10^{-6}$), soluble carbohydrate and soluble protein ($r^2 = -0.89$, $p = 5.57 \times 10^{-7}$), and of particular interest, soluble protein and NSP ($r^2 = 0.77$, $p = 1.58 \times 10^{-7}$). No significant associations were found with insoluble protein bound to the cell wall with any measured proximate components.

Two-way ANOVA test on a model expressing soluble protein content (Table 1) as a function of constituent NSP monomers (Table 4) showed significant association with Arabinose [$F(1,11) = 437.74$, $p = 3.29 \times 10^{-10}$], Fucose [$F(1,22) = 269.72$, $p = 4.37 \times 10^{-9}$], Rhamnose [$F(1,22) = 20.60$, $p = 8.4 \times 10^{-3}$], and Galactose [$F(1,22) = 10.66$, $p = 0.008$]. A multiple regression analysis revealed Xylose and Rhamnose content in the leaf mass could serve as predictors for soluble protein content which was significant at [$F(1,15) = 402.16$, $p = 3.035 \times 10^{-12}$] and [$F(1,15) = 34.69$, $p = 2.97 \times 10^{-5}$], respectively, with values expressed in mg/g dry plant mass in the equation:

$$\text{Soluble Protein} = 0.98 \cdot \text{Xylose} - 3.07 \cdot \text{Rhamnose} + 22.15$$

Goodness of fit of the linear model is shown in Supplementary Figure 1.

From the perspective of process development, such associations are particularly important as plants such as Gorse, Broom and Bracken could yield higher dividends with simple mechanical extraction, leaving behind substrates rich in NSP and Lignin. It further highlights the association between protein and cell wall and the possible treatment; particularly enzymatic, which may be employed to efficiently utilise the biomass. Furthermore, the comparable Lignin and NSP values of Gorse, Broom and Bracken to de-pithed bagasse (Hajiha and Sain, 2015) warrants further investigation into their potential use.
TABLE 4 | Neutral sugar composition of non-starch polysaccharide in Gorse, Broom, Fireweed, and Buddleia. Gorse, Vetch, Broom, Fireweed, Bracken, and Buddleia.

| Plant         | Type        | Rhamnose | Fucose | Arabinose | Xylose | Mannose | Galactose | Glucose |
|---------------|-------------|----------|--------|-----------|--------|---------|-----------|---------|
| Gorse         | Insoluble   | 4.7 ± 0.5a | 1.4 ± 0.1c | 23.0 ± 0.3 | 41.1 ± 0.8 | 4.6 ± 0.4b | 11.7 ± 0.2a | 85.8 ± 1.5 |
|               | Soluble     | –        | 0.1 ± 0.0 | 0.2 ± 0.1 | –       | –       | –         | 0.2 ± 0.1 |
| Vetch         | Insoluble   | 4.6 ± 0.1a | 1.1 ± 0.2 | 10.1 ± 0.1 | 10.0 ± 0.1d | 16.0 ± 0.2 | 16.4 ± 0.2 | 71.3 ± 0.4 |
|               | Soluble     | –        | 0.1 ± 0.0 | 0.1 ± 0.0 | –       | –       | –         | 0.1 ± 0.0 |
| Broom         | Insoluble   | 4.0 ± 0.1b | 1.0 ± 0.0 | 28.4 ± 0.5 | 34.9 ± 1.0 | 5.2 ± 0.1a | 14.4 ± 0.4b | 75.4 ± 2.2 |
|               | Soluble     | –        | 0.1 ± 0.0 | 0.3 ± 0.0 | 0.1 ± 0.0 | –       | 0.3 ± 0.0 | 0.1 ± 0.0 |
| Fireweed      | Insoluble   | 2.6 ± 0.2 | 1.4 ± 0.0d | 11.2 ± 0.3 | 9.8 ± 0.2d | 4.1 ± 0.1d | 17.7 ± 0.4 | 35.8 ± 2.3 |
|               | Soluble     | 0.1 ± 0.0 | 0.1 ± 0.0 | 0.3 ± 0.0 | 0.1 ± 0.0 | 0.1 ± 0.0 | 0.3 ± 0.0 | 0.2 ± 0.0 |
| Bracken       | Insoluble   | 4.2 ± 0.1ab | 1.5 ± 0.0c | 18.3 ± 0.2 | 31.9 ± 0.4 | 6.3 ± 0.1 | 12.4 ± 0.0g | 97.5 ± 2.9 |
|               | Soluble     | –        | 0.1 ± 0.0 | 0.2 ± 0.0 | –       | –       | 0.2 ± 0.1 | 0.1 ± 0.0 |
| Buddleia      | Insoluble   | 7.3 ± 0.2 | 0.7 ± 0.1 | 13.0 ± 0.2 | 7.7 ± 0.2 | 8.7 ± 0.3 | 14.9 ± 0.3b | 60.0 ± 3.0 |
|               | Soluble     | 0.1 ± 0.0 | 0.1 ± 0.0 | 0.3 ± 0.0 | –       | –       | 0.3 ± 0.0 | 0.1 ± 0.0 |

Data is presented at mean ± standard deviation in mg/g dry plant mass. Values with the same superscript are statistically similar using OW-ANOVA at 99% confidence interval with Tukey-HSD post-hoc test. Values with the same superscript indicate similar statistical similarity across corresponding plant samples.

In paper-making or other higher value processing (Rainey and Covey, 2016).

The free sugars measured in the aqueous extract reflect the moieties undergoing translocation in the leaf fated for catabolism or storage (Goldschmidt and Huber, 1992). The soluble phenolics were highest for Broom although, Gorse was found to have the highest overall phenolic content of 109.2 ± 8.1 mg/g dry plant mass with 76.6% of it bound to the plant cell wall components. Syringaresinol was the most abundant lignan across the plants and was particularly high in Gorse (47.4 ± 8.1 mg/Kg), Broom (43.3 ± 6.4 mg/Kg), and Bracken (28.7 ± 2.7 mg/Kg) (Supplementary Table 1). Gorse was rich in chlorogenic acid (1.5 ± 0.1 g/Kg) and in general showed a higher content of all phenolic groups.

Among the 112 phenolic compounds analysed (Supplementary Table 1), 25 compounds accounted for 90% of total variance observed across all plants and largely influenced their characterisation. Correlation and contribution of each component plant and phenolic compound, respectively, to the composite PCA axes is given in Supplementary Tables 2, 3. The plot (Figure 1) revealed that flavonoids as well as cinnamic- and phenylpyruvic acid derivatives largely influenced the profile diversity. Relative position of plants as function of their phenolic profiles is provided in the Supplementary Figure 2 which shows a loose recapitulation of their phylogenetic grouping. Gorse, Broom, and Vetch, which belong to the Fabaceae family are expected to plot closer while distancing themselves from Bracken, Fireweed, and Buddleia. However, Gorse and Broom appeared to have a rather unique profile while Vetch plots closer to Bracken. Fireweed and Buddleia plot separately, reflecting their distinctive phenolic profiles.

Pearson’s test on cinnamic derivatives against phenylalanine (Table 2) reveals a strong positive correlation to cinnamic acid and 3,4,5-trimethoxycinnamic acid content across the six investigated plants ($R^2 = 0.92$ and 0.94, respectively). Larger samples sizes may be required to corroborate this observation, although established biochemical pathways give some support to this observation (Hyun et al., 2011; Vargas-Tah and Gosset, 2015).

Global Distribution of Investigated Invasive Plants

Based on the hits for GPS points classified under a given invasive plant family, their global and Scottish abundance are shown in Figure 2. In the global list of family abundance, among 635 families identified, Fabaceae featured second while Onagraceae ranked 39, Dennstaedtiaceae ranked 76, and Buddlejaceae ranked 245. The Scottish family abundance identified the presence of 131 families with Fabaceae ranking fourth, Onagraceae ranked 19 Dennstaedtiaceae ranked 52, and Buddlejaceae ranked 62.

The expected resolution for such broad, satellite-based vegetation survey is between 120–250 m. The details are provided in the GPS uncertainty margins available in the GIBF database and an account of uncertainties and challenges in remote measurement is provided by Lawrence et al. (2006); Niphadkar and Nagendra (2016); Baron et al. (2018). At a shrub density between 45 and 70 Kg/m² (Passiourea, 1991), the mean biomass held in invasive plants is roughly 52.2 Tg, which translates to 26.1 Tg of carbon and 5.2 Tg of protein. Globally, annual protein requirement is 215 Tg. This could potentially supply about 2.5% of global annual protein requirements, assuming 1.2 g/Kg body weight of daily protein intake as recommended by the National Institutes of Health (NIH) (Phillips et al., 2016). In Scotland, the biomass held in invasive plants is roughly 448.27 Gg, which translates to 224.14 Gg of carbon and 45 Gg of protein. Scottish annual protein requirements are approximately 118.4 Gg, indicating that invasive plants could supply about 25% of annual protein requirements. Interestingly, in terms of minimum essential amino acids recommended by the World Health Organisation (WHO) (Joint WHO/FAO/UNU...
Expert Consultation, 2007), the estimated protein present in the invasive plant biomass could potentially satisfy about 3.5% of global annual requirements. Comparison of essential amino acids produced from existing agricultural products to the estimated values in the invasive plant mass is given in Figure 3 using global and Scottish datapoints. It was observed that the essential amino acid content of all invasive plants could satisfy partial global and Scottish dietary requirements from locally sourced invasive plants. Estimates of the individual essential amino acid content of commercial agricultural produce as well as the invasive plant
Iyer et al. Invasive Plants as Protein Alternatives

**FIGURE 3** Estimated content of essential amino acid present in invasive plant biomass. (A) Values for global essential amino acid content of agricultural produce and invasive plant mass. (B) Values for Scottish essential amino acid content of agricultural produce and invasive plant mass.

**FIGURE 4** The relative abundance of the families of the plants investigated in this work based on their occurrence in the GiBF database. (A, B) Values represent the protein estimated in the invasive biomass.

Biomass is provided in Supplementary Figure 4. The essential amino acid content in families to which the six plants investigated is shown in Figure 4 and the GPS locations were plotted on a map to understand their spread and identify areas most susceptible to their invasion in Figures 5, 6.

**Implications for Land Allocation**

Allocation of fertile land for food production and forestry has a major impact on global carbon storage (Smith et al., 2008). Of the available fertile land used for economic agriculture and forestry, 25.1% is used exclusively for animal husbandry, predominantly as meadows and pastureland (Figure 7). It is the single largest allocation in agricultural land use surpassing the combined value of all other actively developed land such as irrigated, planted, and organic and permanent crop land.

Despite the landmass available for plant growth shown in Figure 7, emissions associated to high-energy agriculture, potentially offset any anabolic carbon storage. Animal husbandry contributes to 22% of total agricultural emissions through direct enteric fermentation via ruminants and a further 15% through manure management. This involves soil conditioning of crop and pastureland (Figure 8) or disposal, making it the single largest contributor of agricultural emission. Burning of organic matter, either as a means of crop/plant waste disposal or fresh land acquisition through “slash-and-burn” cultivation (Uhl, 1987; Kleinman et al., 1995; Brady, 1996) accounts for 17% of agricultural emissions.

Effect of increased resource allocation to animal husbandry and its use as the dominant source of dietary protein can be observed in the carbon footprint to produce yield. Figure 9 illustrates the carbon-cost of common agricultural raw produce. Cumulative effect of enteric emissions and manure handling on the carbon footprint can be observed for ruminant meat such as cattle, goats, and buffalo, which is about 40 times higher than plant based or non-ruminant chicken produce.

The initial database from FAOSTAT contained factors that were converted to intrinsic percentages where applicable. For example agricultural land area was converted to an intrinsic...
value relative to the individual country size. Each subset of agriculture-associated emissions was converted to a percentage of total agricultural emissions. The initial set of factors were: Bovine Meat (g/day), Livestock (LSU/ha), Pesticide (Kg/ha), Fertiliser (Kg/ha), per capita Emission (t), Agri Employment (% labour force), Enteric Fermentation (% agricultural emission), Crop Residue Burning (% emission), Agricultural Soils (% emission), Soil Manure (% emission), Pasture Manure (% emission), Manure Management (% emission), Synthetic Fertilisers (% emission), Animal Origin Emission (% emission), Grassland (% land area), Agricultural land (% land area), Arable Land (% land area), Cropland (% land area), Forest land (% land area), Permanent Pasture (% land area), Permanent Cropland (% land area), Shrubland (% land area), and Urbanisation (% land area).

Among the available factors, only those with a percentage deviation >100 were chosen as shown in Supplementary Figure 5. Factor analysis is optimal across independent factors with normally distributed data. Within a strict maximum cut-off value of \( R^2 > 0.4 \) and at a 99% confidence interval, correlation analysis found Livestock to have strong and significant associations with Urbanisation and Pesticide use (\( R^2 = 0.79 \) and 0.45, \( p: 7.54 \times 10^{-15} \) and \( 1.88 \times 10^{-4} \), respectively). Since Livestock could be expressed as an equation of Urbanisation and Pesticide, it was filtered out. Lastly, the values for plant protein as a percentage of total protein produced (Plant.Production) failed the Royston test for normality and was filtered out. The final dataset contained six factors, namely, Permanent Cropland (Perm.Cropland), Shrubland, Urbanisation, Labour force in agriculture (Agri.employment), Pesticide use, and Permanent Pasture.

An exploratory factor analysis (EFA) using maximum likelihood (ML) suggested an optimal number of two latent factors (shown in Supplementary Figure 3) with corresponding correlations shown in Supplementary Table 4. The RMSR (root mean square of residuals) was 0.06. The RMSEA (root mean square of approximation) index was found to be 0.055 with a 90% confidence interval ranging from 0 to 0.205 and a BIC at \(-11.85\). The Tucker-Lewis Index (TLI) was 0.88.

Based on the suggestions provided by EFA, a confirmatory analysis (CA) of a system equation model (SEM) was performed where the first latent factor (ML1) was expressed as an additive, non-interactive model comprised of Permanent Cropland, Permanent Pasture, and Shrubland, while the second latent factor was expressed as an additive, non-interactive model comprised of Employment, Pesticide use, and Urbanisation. The resultant model is depicted in Figure 10 below. The RMSEA of the model was 0.064 (10% confidence range: 0.00–0.167) and a comparative fit index (CFI) of 0.925. The standardised root mean square residual was 0.061.

As a complementary assessment, the PCA analysis of the selected high-variant intrinsic agricultural factors is shown in

![Invasive plant distribution in Scotland, identified using GPS-Satellite imagery.](image)

FIGURE 5 | Invasive plant distribution in Scotland, identified using GPS-Satellite imagery.
Figure 11 below. The plot accounts for 56.44% of total observed variance and it is interesting to note that countries, particularly the high and upper-middle income countries plot distinctly compared to the low and lowering middle income groups. This suggests that the emergent factor which can be elucidated over the factor analysis is the overall economic status of a country although no direct economic parameter such as GDP or per-capita income was fed into the factor analysis steps. Correlation and contribution of the factors and variables, respectively, to the composite axes is given in Supplementary Tables 5, 6.
DISCUSSION

The strong association between land use and CO$_2$ emissions was previously demonstrated by Smith et al. (2013) who suggested the reallocation of agricultural resources toward more climate-friendly plant-based cultivation. A detailed account of the carbon flow through the agricultural system was reported by Molotoks et al. (2018), which provides strong quantitative evidence of the contribution of fertiliser use and forest land reclamation toward emissions associated to food production. Complementary to their work, global agricultural data from FAOSTAT was used to identify trends that arise when high variable factors are assessed. The principle underpinning the factor analysis described was that the dynamics of a multicomponent system was a function of its most variable factors. In the principal component analysis (PCA) plot shown in Figure 11, vectors (high variant factors) drove the relative positioning of individual countries; revealing characteristic clustering (the emergent factor), which in this case reflected economic status. Diagonally opposing vectors function antithetically while those of similar orientation amplify effect. A more defined relation between the factors is provided by the structural equation model (SEM) depicted in Figure 10 which, in conjunction with PCA revealed a strong association between the economy and land use and by extension, the emissions associated to agriculture. Shrubland was used a proxy for the marginal lands where native and introduced invasive plants are found. The opposing influence between shrubland and permanent cropland in the PCA plot reveals their contrasting effects in the characterisation of countries and their role in economic output as well as local ecology.

Maps showing the confirmed locations of invasive plants chosen for this study were identified using GPS-satellite imagery as shown in Figures 5 and 6. Among the families of the invasive plants chosen for this study, Fabaceae was highly represented in Scottish and global regions followed by Onagraceae, Dennstaedtiaceae, and Buddlejaceae. These plant families are in abundance in the North-East of Scotland. While satellite-imagery has improved over the decades and is invaluable in identifying endangered plant species and contributing toward conservation efforts, the work toward understanding the spread of invasive species remains relative nascent. The GPS locations from where the plant samples were obtained (see Materials and Methods) were not registered in the year 2020 database despite the plants having a sizeable coverage within those locations. Such errors decrease overall plant count and may underreport the severity of the problem faced on account of their spread. Molotoks et al. (2018) estimated agricultural carbon losses faced by invasive plant encroachment to 1 Pg, while according to estimates here based on satellite surveys, the total biomass present across all invasive plants is about 52 Tg. It is uncertain to what extent the biomass has been underestimated. It may be that recurring loss of agricultural biomass is caused by regular
encroachment of invasive plants from the presently identified locations, which act as their natural reservoirs. Improved data collection is required to ameliorate uncertainties and help give a better estimate of the invasive biomass. Despite the possible underestimation, it was observed that the essential amino acid content of these plants may be sufficient to supplement Scottish and to some extent, global requirements.

Among the invasive plants investigated, Bracken was found to have the highest protein content (Table 1) and Vetch to contain the highest essential amino acid content in its
constituent proteins (Table 2). However, in terms of protein accessibility easily achieved from mechanical grinding, Bracken and Gorse were found to have the highest soluble protein content (Table 1). However, the presence of potent carcinogens, namely, ptaquiloside (Niwa et al., 1983; O’Driscoll et al., 2016) found in Bracken makes it an unsuitable candidate until purification methods capable of removing these anti-nutrients are developed. In Vetch, only 15% of total protein was accessible, which was less than half that of Gorse, Broom, or Bracken. This may require further enzymatic or chemical extraction methods to fully realise the nutritional potential, but processes such as this may result in loss of essential amino acids depending on the nature of treatment. Traditional alkali-based extraction followed by neutralisation can result in severe loss of lysine and tryptophan (Jung et al., 2006) and generate significant effluent volumes, while the enzymatic procedures would substantially increase process cost (Sari et al., 2015).

Physical features differentiating leaves such as their hardness, affect the design used to effectively extract proteins, which is primarily governed by the plant cell-wall composition. It is interesting to note that Gorse, Broom, and Bracken are often subject to herbivory (Tempel, 1981; Broadfield and McHenry, 2019) and values of high lignin and NSP (Tables 3, 4) may reflect investment toward defensive mechanisms by toughening leaf mass or employing anti-nutrients; as in the case of Bracken (Niwa et al., 1983; O’Driscoll et al., 2016). The particularly woody nature of Gorse may be attributed to the high xyllose content capable of producing branched, interlocking structures in conjunction with the lignin (Grantham et al., 2017; Gericke et al., 2018; Wierzbički et al., 2019). When leaf samples are dried, this manifests as a brittle leaf structure making it amenable to crushing and juicing for protein extraction. Bracken and Broom leaf samples on the other hand show similar, but lower xyllose content compared to Gorse, which in conjunction with a relatively lower lignin content and may have resulted in a more flexible leaf structures (Arola et al., 2013; Busse-Wicher et al., 2014).

Gorse has been prolific in more than 16 countries (Broadfield and McHenry, 2019); particularly in Oceania where it was introduced as an ornamental plant as well as to build natural barriers around private properties (Hoffmann and Broadhurst, 2016; Figure 6). Downstream process development for protein extraction using leaves from Gorse appears to be a lucrative solution of deriving nutrition as it had the highest protein recovery using economical methods such as mechanical grinding and buffer extraction. Although leaf proteins have previously faced disregard owing to low essential amino acids such as lysine, tryptophan and methionine, which were lost during traditional alkali extractions (Cuq et al., 1983; Zhang et al., 2017), improvements in throughput and reliability of ultrafiltration technology has since rendered the concern moot, as amino acid preservation is greater given the absence of any chemical treatment. All six plants studied plants exhibit a valuable essential amino acid profile (Table 1) suggesting that the final nutrient yield is a function of protein extractability.

The non-protein residue post-aqueous extraction appears to be rich in polysaccharides and lignin which could potentially be valorised for use in other industries examples being; enzyme assisted ethanol production (Cheng et al., 2017) or use in paper industry. Left over marc has previously been demonstrated as an alternate substrate for edible mushrooms (Mintesnot et al., 2014; Kaszynski, 2016). Overall, the harvest index of such plants is high, as organs such as leaves and parenchyma-rich portions constitute about 48–55% of the plant mass (corresponding to Harvest Index of 0.48–0.55) compared to cereals with indices of about 0.3–0.45 (Singh and Stoskopf, 1971).

Gorse is also rich in soluble phenolics, particularly chlorogenic acid and major bioactive compounds such as phenylpyruvic-, benzoic-, and cinnamic acids derivatives. These compounds are commonly associated with health benefits owing to their anti-inflammatory and free radical-quenching properties (Ozcan et al., 2014; Shahidi and Ambigaipalan, 2015) and their role in maintaining gut health (Ozdal et al., 2016). The nutraceutical market is expected to grow to a value of around $10 billion by the year 2026 (Childs, 1999; Bröring and Cloutier, 2008; Research Markets, 2019) and with growing consumer demands and research investment in the plant-based “well-ness” industry, identification of viable and sustainable sources of bioactives is essential.

Invasive plants tend to have robust physiologies and faster germination times, which increase turnover and reduce income latency. Plants such as Gorse and Bracken grow rapidly during periods of spring, allowing harvest in early summer (Conway, 1957; Bowman et al., 2008). Harvesting and exploiting the nutritional value of such plants may help alleviate some of the burden on the existing agricultural systems to produce protein. Current terrestrial carbon held in vegetation amounts to 302.4 Pg (FAO, 2019) which is almost completely offset by animal husbandry industry. Since agricultural emission is a direct result of land use (Smith et al., 2008; Molotoks et al., 2018; Porter et al., 2019), land improvement and maintenance of pastoral lands adds to the carbon cost of animal products as seen in Figure 8. Although classification of grazing land into rough grazing and maintained meadows is not available for all countries of the world, a significant portion is expected to be repurposed marginal land incapable of high-intensity farming (Asner et al., 2004; Peco et al., 2006). These sites tend to retain a considerable part of plant biodiversity and the native plants found in these sites come under enormous survival pressures from climate change, herbivory, and invasive plants and in many cases have been pushed to extinction (Duncan et al., 2004; Truscott et al., 2008; Lankau, 2012).

Given the average rise in global temperatures was recorded to be 1.96°C (2018–2019) (FAO, 2019; Roe et al., 2019), vulnerable economies, which are primarily agrarian are likely to face greater impact of adverse climatic conditions which could pivot them further into distress. A combination of novel, sustainable nutrition sources introduced through minimum disruption is essential to ensure continued livelihood and nutrition supply. The effect of conscious food choice on global carbon emission has been explored by Bajželj (2014) and Bajželj et al. (2014) and the comparison of scenarios across different food and production choices is quantitatively summarised by the EAT-Lancet Commission (Willett et al., 2019). This study strongly indicates the importance of reduction in the consumption of animal sources, while simultaneously moving away from
intensive cultivation practises and instead focusing on food nutrition quality, rather than production yields. Leaf protein extraction technology could help harness the potential in revalorising the biomass of invasive plants, which is currently disposed of inefficiently and help toward production of alternate and more sustainable protein sources to complement existing food production.

CONCLUSIONS

Agriculture in our current economic system carries the burden of being a critical occupation for human survival, while having to maintain economic viability and now, faces the brunt of being a major contributor of anthropogenic GHG emissions. Based on the relationship between land resource allocation, emission and fiscal patterns observed in global agricultural data, the case is presented for revalorising invasive plants to ameliorate food insecurities. Marginal lands tend to serve as reservoirs for native plant species, which have now come under threat from excessive herbivory from unattended grazing, climate change and added competition from invasive plant species. Using databases cataloguing the location and spread of invasive plants through satellite imagery, Fabaceae species were found to be the second most abundant family globally, with the three most invasive species in Scotland belonging to this family. Among the six invasive plants investigated, Gorse (Ulex europaeus) was identified as a good candidate for implementation owing to its wide geographical spread, high protein extractability and potential of the co-products for further revalorisation efforts.

DATA AVAILABILITY STATEMENT

The raw data supporting the conclusions of this article will be made available by the authors, without undue reservation.

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AUTHOR CONTRIBUTIONS

AI performed the experiments, data collection, and manuscript preparation. SD, CB, and WR supervised the work, helped with data interpretation, and preparation of manuscript. All authors contributed to the article and approved the submitted version.

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SUPPLEMENTARY MATERIAL

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**Conflict of Interest:** The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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