Visible Near-Infrared Hyperspectral Imaging for the Identification and Discrimination of Brown Blotch Disease on Mushroom (Agaricus bisporus) Caps

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Visible Near-Infrared Hyperspectral Imaging for the Identification and Discrimination of Brown Blotch Disease on Mushroom (Agaricus bisporus)

Caps

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Abstract

Brown blotch, caused by pathogenic *Pseudomonas tolaasii* (*P. tolaasii*), is the most problematic bacterial disease in *Agaricus bisporus* mushrooms. Although it does not cause any health problems, it reduces the consumer appeal of mushrooms in the market place, generating important economical losses worldwide. Hyperspectral imaging (HSI) is a non-destructive technique that combines imaging and spectroscopy to obtain information from a sample. The objective of this study was to investigate the use of HSI for brown blotch identification and discrimination from mechanical damage on mushrooms. Hyperspectral images of mushrooms subjected to i) no treatment, ii) mechanical damage or iii) microbiological spoilage were taken during storage and spectra representing each of the classes were selected. Partial least squares- discriminant analysis (PLS-DA) was carried out in two steps: i) discrimination between undamaged and damaged mushrooms and ii) discrimination between damage sources (i.e. mechanical or microbiological). The models were applied at a pixel level and a decision tree was used to classify mushrooms into one of the aforementioned classes. A correct classification of >95% was achieved. Results from this study could be used for the development of a sensor to detect and classify mushroom damage of mechanical and microbial origin, which would facilitate the industry to make rapid and automated decisions to discard produce of poor marketability.

Keywords: mushrooms, *Agaricus bisporus*, brown blotch, *Pseudomonas tolaasii*, mechanical damage, vis-NIR hyperspectral imaging, PLS-DA.
Introduction

Cultivated mushrooms are susceptible to a variety of pests and diseases. *Pseudomonas tolaasii* (*P. tolaasii*) is the causal agent of brown blotch (also known as bacterial blotch) disease$^1$ and the most important pathogenic bacterium of *Agaricus bisporus*$^2$. This disease has been detected and described worldwide and affects not only the button mushroom market but the mushroom market in general$^3$. According to growers, brown blotch is “the worst disease”, because of the large economic losses associated to it. Brown blotch can cause a general loss of crop yield of 10 % and a decrease in quality of another 10 %. The most typical symptoms of brown blotch are pitting and browning of mushroom tissues, induced by the watersoluble toxin tolaasin$^5$. This extracellular toxin is produced by the pathogenic form of *P. tolaasii*$^6$. The colonisation of mushroom caps by *P. tolaasii* results in the appearance of unappealing brown spots on the mushroom cap and stipe$^3$. Lesions are slightly concave blemishes, sometimes small, round or spreading in many directions$^7$. When the damage is more intense, the spots are darker and sunken. Browning affects only the external layers of the cap tissue and is restricted to 2-3 mm below the surface of the cap.

The mushroom industry is in need of objective evaluation methodologies to ensure that only high quality produce reaches the market$^8$. Studies in the field of brown blotch detection include the work of Vízhányó and Felföldi$^9$, who tested the potential of a machine vision system to recognise and identify brown blotch and ginger blotch diseases, both of which cause discolouration in mushroom caps. A vectorial normalisation method was developed to decrease the effect of the natural loss of whiteness of the mushroom surface and increase the differences in the image caused by the disease. The method showed an ability to discriminate between discolouration caused by microbial disease and other sources of discolouration, such as natural senescence. However, no attempt was made to discriminate brown blotch from...
bruises induced by mechanical stress, which is also an important source of discolouration and quality loss in the mushroom industry. Hyperspectral imaging (HSI) is a rapid and non-destructive technology that has recently emerged as a powerful alternative to conventional imaging for food analysis. Hyperspectral images are composed of hundreds of contiguous wavebands for each spatial position of an object. Consequently, each pixel in a hyperspectral image contains the spectrum of that specific position. Hyperspectral images, known as hypercubes, are three-dimensional blocks of data, comprising two spatial and one wavelength dimension. The large quantities of highly correlated data contained in a hypercube are well suited to analysis by dimension reduction approaches such as principal components analysis (PCA) and partial least squares-discriminant analysis (PLS-DA). PLS-DA can also be applied to develop qualitative models for supervised classification between various sample classes.

HSI has been applied at various levels in the assessment of safety and quality of food, including constituent analysis, quality evaluation and detection of contaminants and defects. Additionally, a number of researchers have reported the potential of HSI for identification of microorganisms of concern in food. In the field of mushrooms, HSI has proved useful for the detection of bruise and freeze damage and the prediction of moisture content and enzyme activity, as well as for the evaluation of shelf-life and quality deterioration. Recent advances in the detection of skin damage of other products include work by Ariana et al. with cucumbers and Nicolaï et al. and ElMasry et al. with apples. As regards damage of microbial origin, Gómez-Sanchis et al. proposed a hyperspectral imaging system for the early detection of rot caused by Penicillium digitatum (fungi) in mandarins. This method’s success in classifying rotten fruit was above 91% and it represented an alternative to the operationally inefficient sorting system previously used in the citrus industry. While evidence from the literature points to its feasibility, to the authors’
knowledge, HSI has not been used to detect damage of bacterial origin in horticultural products.

The objective of this study was to investigate the potential application of Vis-NIR HSI for brown blotch identification on mushroom caps and for its discrimination from mechanical damage injuries.

Materials and methods

Mushroom supply and damage

*Agaricus bisporus* mushrooms (strain Sylvan A15, Sylvan Spawn Ltd., Peterbourough, UK) were grown in plastic bags and tunnels in Kinsealy Teagasc Research Centre (Kinsealy, Co. Dublin, Ireland) following common practice in the mushroom industry. Only uniform undamaged closed cap mushrooms from the 1\(^{st}\) and 2\(^{nd}\) flush with a diameter of 3-5 cm were hand-picked in November 2008 (training set) and July 2009 (test set). Samples were placed in a metal grid and carefully delivered to the laboratory in purpose-built containers, to minimise mechanical damage during transport. Mushrooms arrived at the laboratory premises within 1 hour after harvesting and were stored overnight at 4°C.

For each set of mushrooms (\(n_{\text{train}} = 144\) and \(n_{\text{test}} = 108\)), samples were divided in 3 groups (undamaged (U), mechanically damaged (MD) and *P. tolaasii* inoculated mushroom (PT)) of equal size (\(n_{\text{train},i} = 48\) and \(n_{\text{test},i} = 36\), where \(i = \text{U, MD, PT}\)).

Each mushroom class was treated as follows:

U: No treatment.

MD: samples were subjected to vibrational bruising to simulate crop handling and transport. Mushrooms were damaged in batches of 600g (approx) units inside polystyrene plastic boxes. Mechanical damage was induced by using a Gyratory Shaker Model G2
shaking table (New Brunswick scientific Co., Edison, N.J., USA) at 300 rpm amplitude for a
shaking period of 10 min. Samples were stored in an environmental incubator (MLR-350 HT,
SANYO Electric Biomedical Co. Ltd., Japan) at 25°C and 90 % relative humidity (RH) for
24 h prior to imaging.

PT: samples were obtained by inoculating 4 drops of 10 µL/each of a solution of
pathogenic *P. tolaasii* onto each clean cap at $4 \times 10^6$ cfu. Samples were stored in the
incubator for 48 h at 25°C and 90 % RH prior to imaging, to encourage appearance of brown
blotch symptoms on the mushroom caps.

A total number of 252 mushrooms were used in this experiment.

Pathogenic *P. tolaasii* solution

Freeze-dried culture (DMS no. 19342) was purchased from Deutsche Sammlung von
Mikroorganismen und Zellkulturen GmbH (DSMZ), Germany, resuspended in nutrient broth
(NB, Scharlau, Dublin) and incubated at 25°C for 24 h. The pure culture was transferred into
nutrient agar plates (NA, Oxoid, Dublin) and incubated at optimal conditions to obtain
isolated colonies.

“Mushroom tissue block rapid pitting” and “White Line in Agar” (WLA) pathogenicity tests
were carried out following the procedure of Wong and Preece\(^{33}\) to confirm culture
pathogenicity on mushrooms.

- **Mushroom tissue block rapid pitting test**

  The outer skin of a mushroom was peeled off and mushroom cap tissue blocks of approx.
  $15 \times 15 \times 5$ mm were cut. Bacterial isolates were grown on Pseudomonas agar base
  (PAB, Oxoid, Dublin) at 25°C for 24 h and suspended in sterile distilled water ( $10^8$ cfu
  mL\(^{-1}\) approx). Mushroom blocks were placed in duplicate on Petri dishes containing
  sterile water-moistened filter paper. The bacterial solution was inoculated onto the cut
surface of one of the mushroom blocks and incubated at 25°C. Sterile water was inoculated onto the surface of the other mushroom blocks for negative control. Pitting of the cut surface of mushroom tissue revealed pathogenicity of P. tolaasii on the mushroom blocks.

“White Line in Agar” test

Pseudomonas reactans (P. reactans) was streaked out in a line, directly from agar slope culture, across PAB in a Petri dish. This strain had been isolated and provided by Kinsealy Teagasc Research Centre.

The P. tolaasii isolate to be tested was streaked immediately after on to the plates at right angles to the reacting organism (i.e. P. reactans). Plates were incubated at 25°C for 24 h for the production of a white line with the reacting organism.

White line production in the agar between P. reactans and P. tolaasii was interpreted as positive interaction between colonies and a positive response for pathogenicity test.

A loopfull of pathogenicity confirmed working culture was transferred to a sterile 0.8% saline solution (Sigma, Dublin). 4 droplets of 10 µL/each of a 10^8 cfu mL⁻¹ solution were inoculated onto the cap, resulting in inoculation concentration of 4 × 10^6 cfu.

Hyperspectral imaging

Hyperspectral images were obtained using a pushbroom line-scanning HSI instrument (DV Optics Ltd, Padua, Italy). The instrument comprised a moving table, illumination source (150 W halogen lamp source attached to a fibre optic line light positioned parallel to the moving table), mirror, objective lens (25 mm focal length), Specim V10E spectrograph (Spectral Imaging Ltd, Oulu, Finland) operating in the wavelength range of 400-1000 nm (spectroscopic resolution of 5 nm), CCD camera (Basler A312f, effective resolution of 580 × 580 pixels by 12 bits), acquisition software (SpectralScanner, DV Optics, Padua, Italy) and
PC. A cylindrical diffuser was placed in front of the fibre optic line light to produce a diffuse light source. In this study, only spectral data within the wavelength range of 445-945 nm were used, as beyond this range the noise level of the camera was high and the signal efficiency of the light source was low.

Reflectance calibration

Reflectance calibration was carried out prior to mushroom image acquisition in order to account for the background spectral response of the instrument and the “dark” camera response. The bright response (“W”) was obtained by collecting a hypercube from a uniform white ceramic tile; the dark response (“dark”) was acquired by turning off the light source, completely covering the lens with its cap and recording the camera response. The corrected reflectance value (“R”) was calculated from the measured signal (“I”) on a pixel-by-pixel basis as shown by:

\[ R_i = \frac{(I_i - \text{dark}_i)}{(W_i - \text{dark}_i)} \]

where \( i \) is the pixel index, i.e. \( i=1,2,3,\ldots,n \) and \( n \) is the total number of pixels.

HSI images of U mushrooms were acquired on day 0 of the experiment. MD mushrooms were scanned after 24 h of storage. PT mushroom images were taken after 48 h of storage. Data were recorded in units of reflectance and saved in ENVI header format using the acquisition software.

Confirmation of \( P. \ tolaasii \)

After image acquisition of PT mushrooms on day 2 of storage, 0.5 g of the outer skin of 10 % of each mushroom class were extracted with a sharp sterile knife. Skins were suspended separately in 10 mL of a 0.8 % saline solution. Samples were homogenised in a stomacher.
(Seward BA 7020, Seward, UK) for 60 s at high intensity. Serial dilutions of each suspension were prepared and transferred onto NA and PAB plates, which were incubated for 48 h at 25°C to obtain isolated colonies. The same procedure had been carried out after image acquisition of U mushrooms on day 0 and resulting colonies were used as negative controls. Serial dilutions of the pathogenic *P. tolaasii* solution that had been used to inoculate PT mushrooms were also prepared and transferred onto NA and PAB plates; resulting colonies were used as positive controls. Mushroom tissue block rapid pitting and WLA tests were carried out to test for pathogenicity of isolated colonies.

**Image processing**

For each mushroom hyperspectral image, 175 characteristic (i.e. U, MD or PT, depending on mushroom class) regions of interest (ROI) were selected by using an interactive selection tool (“ROI tool”) available in the acquisition software. The ROI’s were 3 × 3 pixels in size and were selected from the central region of the mushroom cap, where possible. Selecting spectra from analogous surface areas in all the mushrooms aimed at minimising the scaling differences caused by mushroom surface curvature. The average reflectance spectrum (“R”) of each ROI was obtained by averaging the pixel spectra of the region. Spectral data of each mushroom set were used to build two-dimensional matrices, where each row represented the spectrum of one ROI.

Prior to the development of multivariate models for damage class prediction, spectra were pre-processed using the Standard Normal Variate (SNV) transformation to reduce spectral variability due to non-chemical biases.

Training set matrices (raw and SNV-corrected) contained 8400 spectra and test set matrices (raw and SNV) contained 6300 spectra.
Partial least squares- discriminant analysis (PLS-DA)

Partial least-squares discriminant analysis was applied to the training set matrices (raw and SNV-corrected, n=8400) using MATLAB 7.0 (The Math Works, Inc. USA). The aim was to build models that would enable maximum separation of sample spectra into different classes depending on their physical condition. A two step model approach was taken for each set of spectra (i.e. raw and SNV): one model (namely “U/Dam” model) was developed to discriminate between undamaged (U) and damaged (Dam) spectra and another model (namely “MD/PT” model) was built to discriminate between the two classes of Dam, i.e. mechanical (MD) and microbiological (PT). Overall, four models were built: U/Dam_raw, U/Dam_SNV, MD/PT_raw and MD/PT_SNV.

For this purpose, a dummy response variable, $Y$, was constructed and assigned to each spectrum. For U/Dam models, $Y = 0$ for U spectra and $Y = 1$ for Dam spectra. For MD/PT models, $Y = 0$ for MD spectra and $Y = 1$ for PT spectra. In both cases, a cut-off value of 0.5 was used to classify spectra, as suggested by Esquerre, Gowen, O’Donnell & Downey\textsuperscript{35}; spectra with a predicted dummy variable <0.5 were identified as belonging to class 0, while those with predicted $Y$-value $\geq 0.5$ were classified as belonging to class 1.

Spectra from the training data set were split into 10 sections and continuous blocks cross-validation was performed. The decision on the number of latent variables (# LV) to select for each model was made based on the root-mean square error of cross-validation (RMSECV), which is the mean of the sum of squared differences between the actual and the predicted value of the dummy variable.

The models were also applied to the test set matrices (raw and SNV-corrected, n=6300), which represented independent sets of sample spectra. Performance of the classification models was evaluated on the basis of their sensitivity (number of spectra of a given type...
correctly classified as that type) and specificity (number of spectra not of a given type
correctly classified as not of that type) on training and test sets.

Prediction maps

An important feature of hyperspectral imaging is the ability to map the distribution of
components/attributes on samples. In this case, developed PLS-DA models could be applied
to entire hypercubes of mushrooms to form two dimensional prediction images where the
damage class of each pixel as predicted by the PLS-DA models would be represented by its
intensity ("I"). With this in mind, the following step-by-step procedure was carried out on all
mushroom hypercubes:

1. **Masking.** This step was performed to separate the mushroom pixels from the
background. The mask was created by thresholding the mushroom image at 940 nm,
where a pixel threshold value of 0.1 was used to segment the mushroom from the
background. All background regions were set to zero and only the non-zero elements
of the image were used in further steps.

2. **Erosion.** As all the spectra collected to build the models corresponded to interactively
selected ROI’s of the central part of the mushrooms, the image outline (i.e. edge) of
the mushrooms was eroded to spectra showing differences due to sample curvature.
This was done by eroding the masks using disk-shaped structuring elements (SE,
whose radii increased from 0 –i.e. no erosion- to 40 pixels, in 10 pixel gaps), prior to
the application of PLS-DA models. The effect that varying the radius of the SE had on
i) the area of the mask, ii) the pixel distribution of the predicted maps and iii) the
performance statistics of the two models at a pixel level, based upon ANOVA results
obtained using R$^{36}$, was studied.
3. **Application of developed PLS-DA models.** The U/Dam model was applied to eroded hypercubes and following this classification, the MD/PT model was applied only to the pixels previously classified as *Dam*. Three binary images (Bin$_{U}$, Bin$_{MD}$ and Bin$_{PT}$, one for each damage class, where 1 indicated class membership and 0 indicated non-membership) were generated after classifying each pixel as “U” (I$_{U/Dam}$<0.5), “MD” (I$_{U/Dam}$>0.5 and I$_{MD/PT}$<0.5) or “PT” (I$_{U/PT}$>0.5 and I$_{MD/PT}$>0.5) tissue.

4. **Closing.** As enzymatic browning was expected to develop uniformly across MD mushroom caps and bacterial lesions were expected to appear as brown spots of visible size, Bin$_{MD}$ images closed in order to avoid noise in the form of isolated PT pixels in such maps. The *Closing* morphological operator performs dilation followed by erosion; this was done using a diamond-shaped SE with a radius of 3 pixels. The effect of omitting/incorporating Step 4 on i) the pixel distribution of the prediction maps and ii) performance statistics, based upon ANOVA results, was investigated.

5. **Concatenation:** the three binary maps (i.e. Bin$_{U}$, Bin$_{MD}$ and Bin$_{PT}$) were concatenated to build false colour maps where *U*, *MD* and *PT* classified pixels were represented in green, red and blue, respectively.

**Mushroom classification**

Based on the percentage of pixels of each damage class on the prediction map, a decision tree (shown in Figure 1) was used to allocate each mushroom to one of the three mushroom classes. As the main objective of this work was to identify *P. tolaasii* inoculated mushrooms, the *PT* pixel percentage of the prediction maps was selected as the discrimination criteria and a cut-off value was established by exploring the pixel histograms of the Bin$_{PT}$ images of U, MD and PT mushrooms (Figure 2). A *PT* pixel percentage of 2 % appeared to be a reasonable cut-off point, as all of the U mushroom predictions and almost all of the MD mushroom predictions exhibited lower values and almost all of the PT mushroom predictions showed...
higher values. After visual inspection of the two PT mushrooms that were below the cut-off value, it was observed that these two mushrooms did not develop any brown blotch on their caps, for which they could be left out for cut-off establishment purposes. As it can be seen in the figure, when the number of PT pixels in the prediction image was higher than 2 %, the mushroom was classified as PT. If the number of PT pixels was lower than 2 % and the amount of MD pixels was higher than the amount of U pixels, the mushroom was classified as MD. Finally, if the number of PT pixels was lower than 2 % and the amount of MD pixels was lower than the amount of U pixels, the mushroom was classified as U.

Sensitivity and specificity of the classification procedure were computed after the application of the decision tree to all of the mushroom hypercubes.

Results and discussion

RGB images

Figure 3 shows representative colour images of the three mushroom classes under investigation in this study. Mushrooms labelled as U (Figure 3a) were white in general appearance, although some of them showed some signs of natural discolouration caused by common picking and transport practice. By day one of storage, MD samples (Figure 3b) exhibited uniform browning over the entire mushroom surface. By day two of storage, P. tolaasii had colonised the cap of most PT mushrooms (Figure 3c), which exhibited slightly concave brown-coloured spots, the typical symptoms of brown blotch disease.

Confirmation of P. tolaasii

The prevalence of Pseudomonas in mushroom surfaces is high, but only pathogenic P. tolaasii is capable of causing brown blotch. Two different types of colonies, whose colours were a) creamy and b) green, were found in PAB plates of PT mushrooms, while only one type appeared in PAB plates of U mushrooms and the P. tolaasii inoculum; the colour of
these colonies was creamy and green, respectively. Isolates of the two types found in PT mushroom plates were obtained by re-streaking representative colonies onto fresh PAB plates. Figure 4 shows growth in PAB plates of the two types of colonies of PT mushrooms (Figures 4a, creamy colonies and 4b, green), the creamy colonies of U mushrooms (Figure 4c) and the green colonies of the *P. tolaasii* inoculum (Figure 4d). The creamy colonies of PT mushrooms (Figure 4a) were found to be similar to those found in U plates (Figure 4c), and both gave a negative response to the mushroom tissue block rapid pitting and WLA tests. The green colonies of PT mushrooms (Figure 4b) were similar to those found in *P. tolaasii* inoculum plates (Figure 4d) and both had a positive response to the two pathogenicity tests. These results confirm that the pitting observed in PT mushrooms was due to pathogenic *P. tolaasii*.

**Spectra**

Mean spectra of the various spectra classes are shown in Figure 5: (a) non-pretreated reflectance spectra and (b) SNV-corrected reflectance spectra. In Figure 5a, signal intensity and shape differences between U and MD spectra were remarkable. The mean MD spectrum exhibited lower reflectance values over the entire spectral region, as expected after bruising had led to loss of whiteness of the caps. The greatest differences in shape between MD and U spectra arose in the 600-800 nm region, where the mean U spectrum exhibited broader features than the mean MD spectrum. Broad spectra in the visible-near infrared wavelength range are characteristic of undamaged mushrooms, corresponding to their white appearance. The spectral differences mentioned above could be related to the formation of brown pigments, mainly melanins, which derive from enzyme-catalysed oxidation products called quinones. The mean PT spectrum appeared to be more similar in shape to the mean MD spectrum, although its slope was not as linear as MD’s was in the 600-800 nm region.
Spectral differences in Figure 5a arose from differences in sample composition, but differences in illumination conditions, sample height and curvature may also have affected the spectral response of the different mushroom classes. Spectra preprocessing methods such as Multiplicative Scatter Correction (MSC) and SNV can be used to compensate for spectral variability caused by these external factors.

The mean spectra of SNV-corrected reflectance spectra of U, MD and PT spectra are shown in Figure 5b. Comparing U and MD spectra, MD exhibited higher SNV-corrected reflectance values in the 450-500 nm region and lower SNV-corrected reflectance in the 500-750 nm region. The oxidation of polyphenolic compounds and subsequent development of brown colour in the MD mushrooms might be partly responsible for this dissimilarity. In the 800-950 nm region, MD spectra showed higher values than U mushrooms. Overall, the shape of the mean spectrum of PT spectra was somewhat intermediate between the mean of U and MD spectra in the wavelength range of study. The visible end of the mean PT spectrum looked more similar to MD than to U, whereas its shape in the >700 nm region was very similar to that of U mushrooms.

**PLS-DA analysis**

Figure 6 shows RMSECV and performance statistics (i.e. sensitivity and specificity) of the four PLS-DA models developed, as a function of the number of latent variables (from 1 to 10). In binary classifications, the sensitivity of a model is a measure of its ability to correctly classify spectra of a given type as being of that type, whereas the specificity is a measure of its ability to correctly classify spectra which are not of a given type as not being of that type. For both U/Dam models (Figs. 6a and 6b), RMSECV exhibited a “corner” (pointed with a red dash circle) at 2 LV. The performance statistics, which were very poor at 1LV, increased at that point and remained at similar levels thereafter, for which 2 was considered to be the optimal # LV for models discriminating between U and Dam spectra. Both MD/PT models
seemed to perform best when 4 LV were selected; RMSECV did not decrease significantly after that and performance statistics remained high.

Numeric values of performance statistics of the selected models are shown in Table 1. When the models built with raw spectra were applied to the training set of spectra, almost perfect classification was achieved in the case of the U/Dam model (sensitivity = 0.997 and specificity = 1.000). The model performed worse when built on SNV-corrected spectra (sensitivity = 0.973 and specificity = 0.999); however differences in sensitivity and specificity were very small. In both cases, almost all of the Dam spectra were classified as such and none or only a few U were misclassified as Dam. When the MD/PT model was applied to the damaged spectra, the sensitivity of the raw model (sensitivity = 0.988) was higher than that of the SNV-corrected model (sensitivity = 0.963), whereas the specificity of the MD/PT_raw model was lower than the MD/PT_SNV model’s (0.983 and 0.998, respectively). These results showed that almost all of the spectra of the mushrooms that had been inoculated with *P. tolaasii* were classified correctly and only a few or none of the spectra of the MD samples were misclassified as PT.

When the models were applied to the test set of spectra, the sensitivity of the U/Dam_raw model was lower (sensitivity = 0.832) but still none of the U spectra were misclassified as Dam (specificity = 1.000). Performance statistics were quite similar for the U/Dam_SNV model (sensitivity = 0.825 and specificity = 0.999). When the MD/PT model was applied to the damaged spectra of the test set, a smaller percentage of raw PT spectra were classified correctly (sensitivity = 0.661) but almost none of the MD spectra were misclassified as PT (specificity = 0.984). As observed for the previous model, the sensitivity of the MD/PT_SNV model was slightly lower (sensitivity = 0.641) and the specificity was higher (specificity = 0.998).
Overall, models built on raw reflectance spectra performed better in this study. However, with a view to generalising the use of U/Dam and MD/PT discrimination models, it might be worthwhile to compromise classification performance in favour of employing more versatile models (e.g. models built on SNV-corrected spectra).

**Prediction maps**

Figure 7 shows examples of prediction maps (with no erosion applied in Step 2) of (a) U, (b) MD and (c) PT mushrooms as a result of the application of raw (top row) and SNV-corrected (bottom row) PLS-DA models to the data hypercubes. Overall, predictions by models built on raw reflectance spectra appeared to be more appropriate than predictions by models built on SNV-corrected reflectance spectra: for each mushroom class, the corresponding pixel class was the main pixel class and pixels were distributed in an even manner. In the example shown (Figure 7), on the top row (i.e. predictions by models built on raw reflectance spectra), neither the map of the U mushroom nor the central region of the prediction of the MD mushroom showed misclassification, whereas most edge pixels of the latter were misclassified as U. Considering that all the spectra selected for model building belonged to central regions of the mushrooms, this misclassification could be related to the inability of the models to account for spectral differences due to mushroom surface curvature. Fewer pixels were misclassified in the prediction map of the PT mushroom, where some pixels were classified as MD. On the bottom row of Figure 7 (i.e. prediction maps by models built on SNV-corrected reflectance spectra), the maps of all mushroom types showed misclassification. For U and PT mushrooms, misclassification happened mainly but not only on the edges, where many U pixels were classified as MD. For MD mushrooms, misclassified pixels were distributed evenly along the mushroom surface. In this case, MD pixels were misclassified as PT.
Considering that the main focus of this work lies in the identification of PT mushrooms, after visual inspection of the prediction maps (Figure 7), PLS-DA models built on non-pretreated spectra were considered more appropriate for this purpose. Models built on SNV-corrected reflectance spectra were therefore discarded and further sections of this paper will focus only on models built on the raw data.

Effect of varying the radius of the SE in Step 2

Erosion of a binary image is a basic operation to wear the boundaries of regions away. This can be done to overcome the problem introduced by the so-called “edge effect”, by which variability in reflection of light in introduced by spherical surfaces. In this particular case, where PLS-DA models were built on spectra selected from central regions of the mushrooms, it was expected that these models would perform better on central areas of the mushrooms than on edge regions. For this reason, masks defining the mushroom region were eroded using SE’s before the models were applied (see Step 2 in Prediction maps section). This led to a decrease in misclassified pixels (typically belonging to edge regions, as observed in Figure 7b, top row).

Figure 8a shows the effect that increasing the size of the structuring element used in this step (i.e. Step 2) had on the mask (top row) and on the prediction map (bottom row) of a MD class mushroom. As the radius increased (from left to right, from 0 -no erosion- to 40 pixels), the mask became smaller. Consequently, the number of MD class pixels that were misclassified (as U class) decreased progressively. Figure 8b shows the decrease of the average relative area of the mushroom region as a function of the radius of SE, where the relative area of each mask at a certain SE radius value is displayed as a percentage of the area of the mask when the radius was zero (i.e. when no erosion was applied) and the average and standard deviation values were obtained by considering all the mushrooms in the training and test data sets. Figure 8c shows the sensitivity of the PLS-DA models built on raw reflectance spectra
applied at a pixel level as a function of the radius of the SE. The sensitivity of the U/Dam model reached its maximum (sensitivity = 1) at a radius value of 20 pixels when applied to the training set and at a radius value of 40 (sensitivity = 0.972) when applied to the test set. The sensitivity of the MD/PT model was not affected by the radius and remained at its maximum (sensitivity = 1) for the training set, whereas it increased progressively until it reached its maximum (sensitivity = 0.944) at a radius value of 40 pixels. Modifying the radius of the SE did not affect (p>0.05) the specificity of the PLS-DA models (results not shown).

**Effect of omitting/incorporating Step 4**

As the existence of isolated PT pixels in the prediction map had no physical meaning (brown blotch lesion on mushroom caps are detectable by the human eye), a closing step (see Step 4 of *Prediction maps* section) was incorporated to the prediction map routine. This step performed dilation followed by erosion on the BinPT images. Figure 9a shows how the final prediction of a MD class mushroom looked when i) Step 4 was omitted and ii) Step 4 was incorporated in the routine. As it can be observed in the figures, Step 4 removed PT class isolated pixels (blue) in the prediction image and converted them into pixels of class MD (red). The effect that such conversion had on the performance statistics of the models at a pixel level was studied and only the specificity of the MD/PT_raw model on the test set was found to change significantly (p<0.05). Figure 9b shows how specificity changed as the radius of the SE of Step 2 increased, when i) Step 4 was omitted (round marker) and ii) Step 4 was incorporated (square marker) in the routine. The specificity of the MD/PT_raw model improved when this step was incorporated, which means more MD class mushrooms were correctly classified. This figure suggests that adding a *closing* step was important to achieve good levels of classification.
After studying the two effects, a disk radius of 40 pixels was selected for the SE of Step 2 and it was decided to include Step 4 in the generation of prediction maps. Further sections of this paper will only focus on results based on the use of the aforementioned steps.

**Mushroom classification**

The application of PLS-DA models built on raw spectra to the totality of entire hypercubes led to the performance statistics shown in Table 2. For the training set mushrooms, both the sensitivity and the specificity of the U/Dam\_raw model were 1, which means there was no misclassification at all. For the same samples, the sensitivity of the MD/PT\_raw model was 1 and its specificity was 0.98. Only 1 out of 48 MD mushroom was misclassified as a PT mushroom. The models performed quite similarly for the mushroom hypercubes of the test set: for the U/Dam\_raw model, sensitivity = 0.97 and specificity = 1. Only 2 out of 72 Dam mushrooms were misclassified as U, and none of the U was misclassified as Dam. For the MD/PT\_raw model, sensitivity = 0.944 (only 2 out of 36 PT mushrooms were not classified as such) and specificity = 0.97 (only 1 MD mushroom was misclassified as being PT).

These results show the models performed well when applied at a pixel level and could be the first step towards the development of a HSI sensor that would classify independent sets of mushrooms with high levels of accuracy. Overall, the correct classification of the models presented in this paper is higher than the classification of the algorithms by Vízánhý and Felföldi⁹, which correctly classified 81% of the diseased areas of test mushrooms using conventional computer imaging. It should be noted that the procedure described in this paper is longer and more complex than the one presented in that study, and the technology more costly. While the algorithms presented in the aforementioned paper discriminated diseased spots from healthy senescent mushroom parts, the models developed in this paper discriminate microbial spoilage from both undamaged and mechanically damaged samples.
The correct discrimination between PT and MD mushrooms ensure no misclassification of samples whose colour analysis might be similar and hence avoid “false positives”.

**Conclusion**

Results presented in this work show that raw reflectance data of mushroom caps could be used to classify mushrooms according to their damage class (i.e. undamaged, mechanically damaged or brown blotch diseased). PLS-DA models were developed to initially sort mushrooms into undamaged or damaged classes and to further classify the damaged into mechanically damaged or microbiologically diseased classes. The application of the models at a pixel level together with the use of a decision tree allowed for correct classification of >95%. This study demonstrates the potential use of hyperspectral imaging as an automated tool for detection of brown blotched mushrooms and for their discrimination from mechanically damaged mushrooms. Knowledge gained in this research using HSI could be incorporated towards the development of simpler sensors to detect and classify mushroom damage of different sources. Such a system could aid the industry in increasing quality control standards by correctly identifying low quality produce. However, further research and validation at industrial scale are required to facilitate its adoption.

**Acknowledgements**

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**References**


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**Figures**

**Figure 1**

- Prediction map
- Count the number of *U*, *MD* and *PT* pixels in the prediction image
- Is the percentage of *PT* pixels greater than 2%? Yes → Classify the hypercube as *PT*
- No → Is the percentage of *MD* pixels greater than the percentage of *U*? Yes → Classify the hypercube as *MD*
- No → Classify the hypercube as *U*

**Figure 1** Decision tree for mushroom hypercube classification, where *U* = undamaged, *MD* = mechanically damaged and *PT* = *P*. tolaasii inoculated.
Figure 2 Histograms showing number of mushrooms as a function of the percentage of pixels in the BIN\textsubscript{PT} binary images of (a) undamaged (U), (b) mechanically damaged (MD) and (c) P.tolaasii inoculated (PT) mushrooms. All the mushrooms (i.e. training and test set mushrooms) of each class were plotted together, making a total of 86 samples per mushroom class.

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**Figure 3**

(a) (b) (c)

**Figure 3** Representative colour images of (a) undamaged (U), (b) mechanically damaged (MD) and (c) *P. tolaasii* inoculated (PT) mushrooms.

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**Figure 4**

(a) Creamy colonies in PAB plate of PT mushrooms, (b) Green colonies in PAB plate of PT mushrooms, (c) Creamy colonies in PAB plate of U mushrooms and (d) Green colonies in PAB plate of *P. tolaasii* inoculum.

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**Figure 5**

(a) Mean ± standard deviation raw reflectance spectra and (b) mean ± standard deviation SNV-corrected reflectance spectra of selected regions of undamaged (U), mechanically damaged (MD) and P. tolaasii inoculated (PT) mushroom caps. For each mushroom group, the broader line represents mean spectrum and the narrower lines represent ± standard deviation spectra.

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Figure 6

Figure 6 Root-mean square error of cross-validation (RMSECV) and performance statistics (i.e. sensitivity and specificity) of PLS-DA models of (a) U/Dam_raw model, (b) U/Dam_SNV model, (c) MD/PT_raw model and (d) MD/PT_SNV model as a function of the number of latent variables (# LV), where _train = training set and _test = test set.
Figure 7 Prediction images (after no erosion in Step 2) of (a) undamaged (U), (b) mechanically damaged (MD) and (c) *P. tolaasii* inoculated (PT) mushrooms by PLS-DA models built on raw reflectance spectra (top row) and SNV-corrected reflectance spectra (bottom row).

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**Figure 8**

(a) Binary masks (top row) and prediction maps (bottom row) of a mechanically damaged (MD) mushroom, (b) Average relative mask area ± SD and (c) Sensitivity of PLS-DA models, as a function of the radius of the structuring element (SE) used for erosion in Step 2.

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**Figure 9**

Figure 9 Effect of the exclusion/incorporation of Step 4 to the prediction map routine in terms of (a) the pixel distribution in the prediction map of a mechanically damaged (MD) mushroom when (i) Step 4 was omitted and (ii) Step 4 was incorporated and (b) the Specificity of the MD/PT_raw model on the test set as a function of the radius of the structuring element (SE) used for erosion in Step 2.

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### Tables

#### Table 1

Table 1 Performance statistics at spectra level of all the PLS-DA models built on reflectance spectra.

<table>
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<th>Specificity</th>
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<th>Specificity</th>
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<td>0.998</td>
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### Table 2

Performance statistics at a pixel level PLS-DA models built on raw reflectance spectra.

<table>
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