

INSTITUTO SUPERIOR DE CIÊNCIAS DA SAÚDE EGAS MONIZ

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MAPPING PORTUGUESE SOILS USING SPECTROSCOPIC

TECHNIQUES WITH A MACHINE LEARNING APPROACH

Work Submitted by

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To my parents, Pete Stafford and Lynn Marshall, to my brother Chris Stafford, and to all my friends for their continued support.

Certificate of Originality

This is to certify that I am responsible for the work submitted in this thesis and that the work is original and not copied or plagiarised from any other source, except as specified in the acknowledgements and in references. Neither the thesis nor the original work contained therein has been previously submitted to any other institute for a degree.

Signature:

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Abstract

Soil analysis is an important part of forensic science as it can provide vital links between a suspect and a crime scene based on its characteristics. The use of soil in a forensic context can be characterised into two categories: intelligence purposes or court purposes. The core basis of the comparison of sites to determine the provenance is that soil composition, type etc. vary from one place to another. The aim of this project is to 'map' soils and predict the location of a sample of unknown origin based on the chemometric profiles of Fourier transform infrared (FTIR) spectra, micro x-ray fluorescence profiles and visible spectra. Thirty one samples were collected in triplicate from Monsanto Park in Lisbon for each predetermined collection point on a defined grid. Full FTIR spectra (400-4000cm⁻¹), Visible (1100-401cm⁻¹) spectra, UV $(400-200 \text{ cm}^{-1})$ spectra and μ XRF profiles were collected for all samples. A subset of 43 discriminant features was selected from a total of 1430 using the Boruta feature selection algorithm from the FTIR, µXRF and visible spectra. These discriminant features acted as input data that was used to create a neural network which allowed the prediction of Cartesian co-ordinates (or location) of the samples with a high degree of accuracy (86%) and has shown to be a very useful approach to predict soil location.

Key Words: Artificial neural network; Fourier transform infrared spectroscopy; Micro x-ray fluorescence spectroscopy; Soil analysis; UV-Visible spectroscopy.

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List of Abbreviations

AAS	Atomic Absorption Spectroscopy
ANN	Artificial Neural Network
ATR	Attenuated Total Reflection
EDXRF	Energy Dispersive X-ray Fluorescence Spectroscopy
ETA	Electrothermal Atomisation
FAA	Flame Atomic Absorption
FFT-LW	Fast Fourier Transform Local Weighted
FTIR	Fourier Transform Infrared Spectroscopy
GPS	Global Positioning System
HCl	Hydrochloric Acid
ICP	Inductively Coupled Plasma
ICP-AES	Inductively Coupled Plasma Atomic Emission Spectrometry
ICP-MS	Inductively Coupled Plasma Mass Spectrometry
ICP-OES	Inductively Coupled Plasma Optical Emission Spectrometry
IR	Infrared Spectroscopy
КОН	Potassium Hydroxide
LA-ICP-MS	Laser Ablation Inductively Coupled Plasma Mass Spectrometry
LDA	Linear Discriminant Analysis
LIBS	Laser-Induced Breakdown Spectroscopy
MSE	Mean Squared Error
NIPALS	Nonlinear Iterative Partial Least Squares
NIR	Near Infrared
NKj	Nitrogen Kjeldahl
PCA	Principle Component Analysis
PCR	Principle Component Regression
PLS	Partial Least Squares
PLS-LDA	Partial Least Squares Linear Discriminant Analysis
PSVM	Penalised Support Vector Machine
RMSE	Route Mean Squared Error
RSD	Relative Standard Deviation
SDA	Shrinkage Discriminant Analysis
UV-VIS	Ultraviolet-Visible Spectroscopy
WDXRF	Wavelength Dispersive X-ray Fluorescence Spectroscopy
XRD	X-ray Diffraction
XRF	X-ray Fluorescence Spectroscopy
μXRF	Micro X-ray Fluorescence

1 Introduction

Soil is a complex mixture consisting of crystalline and amorphous minerals, oxides, decomposing organic matter, plants, pollen, microbial residues along with other compounds produced during the formation process (Sugita & Marumo. 1996; Horswell, et al. 2002). Forensic geoscience (analysis of soils and sediment) has been performed for many years for a wide variety of purposes; examples include differentiating between different land use types (Baron et al. 2011), sediment content (Guedes et al. 2009) and the analysis of soil pollutants (Mostert et al. 2010). Due to soils persistence, transferability and Locard's exchange principle 'every contact leaves a trace' (Nickolls, 1956), soil analysis can supply the essential link from the crime scene in question to the suspect and therefore it can be a vital tool. At present, by using a wide range of analytical techniques, crime scene soil samples can be matched to a soil sample taken from a suspect and link them to a particular scene. However, discriminating and characterising soils for intelligence purposes can be much more complex (Baron et al. 2011) due to the enormous variety in composition which is dependent on the location, the type of soil, climate and human activities (Horswell, et al 2002; Pye et al. 2007; Reidy et al. 2013).

The building of soil profile databases using both physical and chemical properties can be rather problematic due to the complex mixtures of material, organic matter and minerals, for example leaves, twigs, rocks and water and even man-made objects such as concrete, glass and ceramics (Pye & Croft, 2004) and thus can be very costly and time consuming (Gogé *et al.* 2014). It has recently been suggested that using national databases of soil properties combined with locally collected samples can aid in predicting the properties of samples (Guerrero *et al.* 2010; Gogé *et al.* 2014), however Ge *et al.* (2011) recognised that creating a soil spectral library with a robust calibration using only one instrumental technique demanded several thousand samples, which is not practicable to be replicated on each different instrument used. Questions arose from this, such as if the instrument is replaced by a new one, can the library still be used with soils scanned with the newer instrument? Also, if a global library was created using different instruments or the same instruments with different conditions, how useful would this be? Soil composition, type etc. varies from one place to another which can create immense problems when using soil comparisons in legal cases, for the reason that the variation can occur equally within a particular site as much as between sites, and the degree of this is still unknown (Baron *et al.* 2011). Due to this, it has been documented that is it more straightforward to eliminate soils based on their profiles and compositions than it is to 'match' an unknown sample to a known, taking into account that it is not possible to provide probabilities that another locality may or may not possess the same or very similar characteristics (Morgan & Bull, 2007; Pirrie *et al.* 2014). One can simply conclude that two samples either do not share a common source or that they are similar in all analytical aspects and therefore cannot be excluded. Forensic soil analysis has already been used in criminal investigations and provided essential information in criminal cases (Dawson *et al.* 2008; Fitzpatrick & Raven, 2012).

This project aims to develop a method that can be used for intelligence purposes and for that reason this aspect will be the main focus herein. In soil analysis for intelligence purposes, there are two main aspects to be considered in pursuance of excluding samples due to the dissimilarity or including them because they are very similar and these are the physical properties and the chemical properties of the soil samples.

1.1 Physical Properties

1.1.1 Colour Analysis

It has been previously established that colour is an important aspect of soil analysis and by using Munsell® Soil Colour Charts this can be achieved *in situ* and produce a visual representation of the soil's colour. Sample preparation must be standardised to enable comparison and it was identified that air drying, moistening, iron oxide removal and organic matter decomposition can affect the colour measurements (Sugita & Marumo, 1996). It is sometimes obvious to the naked eye that samples possess different colours while others may be too similar to distinguish. Using Munsell Indices allows values to be given to the various aspects of colour; hue (primary or secondary colour), Value (lightness or darkness) and chroma (intensity or saturation) or L*a*b* indices that expresses colour in relation to the position on a 3D coloured sphere (Croft & Pye, 2004). L* is related to lightness (black ranging to white), a* relates to a green-red continuum whilst b* relates to a blue-yellow continuum (Guedes *et al.* 2009). Cox *et al.* (2000) used the Munsell Colour Chart to assign Munsell values to soil samples collected from Oregon (USA) before and after pyrolysis. It was found that all the samples had the same pre pyrolysis colour and so pyrolysis was carried out. Even after pyrolysis, some samples still had the same post pyrolysis colour and thus it was determined that another technique must be used in order to differentiate them. Guedes *et al.* (2009; 2011) demonstrated that measured L*a*b* values were better for discrimination when applied to dried, un-sieved bulk samples as opposed to pre-treated samples, whereas Croft & Pye (2004) suggested removal of organic matter or analysis of each size fractions before and after heat treatment would provide an in-depth analysis.

1.1.2 Granulometry

It has been demonstrated by many authors that granulometry is very useful in soil analysis. Chazottes *et al.* (2004) found that size distribution of the soil considerably affected results. Particle sizes of 2mm-63 μ m (unimodal distribution) was found to be very representative of the 'original' soil sample, whereas bimodal distribution (soils dominated by the extreme particles, those bigger than 4mm and smaller than 20 μ m, were not very representative. It was suggested than any significant differences in the range of 1mm to 63 μ m must be considered indicative of dissimilarity between samples. These results are probable due to the fact that bigger particles are more likely to detach from material than smaller particles, leading to a different distribution to control samples.

Many authors have differing opinions of which size fractions provide the best discrimination. Pye *et al.* (2007) found bulk samples had the greatest variability and 63-150 μ m has the least, and so less discrimination power when analysis soils using inductively coupled plasma – mass spectrometry (ICP-MS) and inductively coupled plasma – atomic emission spectrometry (ICP-AES). Guedes *et al.* (2009) also found bulk and fractions smaller than 150 μ m gave the best discrimination whilst carrying out colour analysis.

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1.1.3 Palynology

Palynology is pollen and spore science (Hyde & Williams, 1944). Simple palynology consists of pollen and/or spore identification along with pollen and/or spore counting (counting how many times each species occurs within a sample). This allows the creation of a pollen assemblage (profile) which can then help to identify similarities and dissimilarities between samples. It can be a very helpful tool due to the immense variety in the exines (outer shells) making each species unique and the resistive and persistent nature allowing them to survive in particular conditions for thousands of years. It can become complicated if one is not a trained palynologist or someone with little experience due to complexities of pollen identification for example several types of pollen grains can be produced by a single species or grains that look visually similar under a standard microscope come from unrelated plants (Erdtman, 1966).

Mildenhall (1990) recognised there were issues with the use of palynology in forensic science and highlighted these. The problems that were pointed out were the nature of the scene and the uniqueness of assemblages. This evidence is always circumstantial, in the fact that it can connect a person to a particular place but it cannot tell if the person has committed a crime and with assemblages, it is always possible that another location may possess a similar profile. It was also recognised that due to the destructive nature of pollen preparation, other techniques must be carried out before this and so can open the exhibits up to contamination. Mildenhall (2006) demonstrated how important palynological evidence can be in the case of an assault on a young woman. Pollen found on her body and clothing was identical to that found at the scene (with fungal hyphae growth) and it is unlikely she would have had these pollens on her body beforehand. This provided very strong evidence of the scene of the attack and allowed the conviction of the offender. This paper showed the importance of pollen analysis as well the collection of samples from different areas of clothing and that even common pollen can provide vital information in forensic cases and can secure convictions.

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1.2 Chemical Properties

1.2.1 Infrared (IR) Spectroscopy

Infrared spectroscopy is a well known and used technique in forensic soil analysis as it allows one to analyse the organic and inorganic composition of soils. The IR region of the electromagnetic spectrum ranges from 14000 to 10cm⁻¹ and can be subdivided in to three regions: near, mid and far infrared. Near-infrared ranges from wave numbers from 14000-4000 cm⁻¹, mid-infrared ranges from 4000-400 cm⁻¹ and far-infrared ranges from 4000-10 cm⁻¹ (Figure 1) (Smith, 1999; Larkin, 2011).



Figure 1. Infrared region of the spectrum consisting of 3 sub regions: near, min and far infrared (adapted from Viscarra Rossel *et al.* 2006).

Fourier transform infrared spectroscopy (FTIR) is a non-destructive technique that requires little sample preparation however the sample must be dried to remove all water as this interferes with the spectrum and samples that are thick allow less infrared light to pass through the sample resulting in a poor signal to noise ratio. IR spectroscopic techniques are extremely sensitive to the organic and inorganic phases of soil (Viscarra Rossel *et al.* 2006). IR causes vibrations within molecule and these vibrations within particular bonds will only occur at specific wavelengths. This makes it possible to identify functional groups in a molecule and thus the chemical structure and identity of the compound can be confirmed (Haberhauer *et al.* 1998; Larkin, 2011).

Baron *et al.* (2011) collected 60 soil samples from 3 different areas in Lincoln, UK. Samples were taken from 4 flower beds, 4 river banks and from 4 woodland sites (each with 5 replicates). Attenuated total reflectance - Fourier transform infrared spectroscopy (ATR-FTIR) was used to collect full spectra from 4000-400 cm⁻¹ using 128 scans with 4cm⁻¹ resolution. The data analysis consisted of nonlinear iterative partial least squares with linear discriminant analysis (NIPALS-LDA) and partial least squares discriminant analysis (PLS-DA). It was found that samples could be completely separated by the land type although it was more difficult to separate the different sites (flower beds, riverbeds and woodland), even after removing regions of the spectra that had poor signal-to-noise ratio. It was concluded that NIPALS-LDA was successful in modelling the 60 spectra into the 3 land-use types although PLS-DA was poor. It was also found that the NIPALS-LDA tool offered a more straightforward and successful approach for modelling but the authors suggest further work is needed as well as more samples, sites and models at different levels to propose a more methodical approach to dataset increases.

Cox *et al.* (2000) also used FTIR to create spectral profiles of the soil samples by collecting the spectra of the samples, then pyrolysis was carried out and the spectra collected again. The spectrum of the pyrolysed sample was then subtracted from the original so only the organic content spectra remained. There was not sufficient information to repeat this experiment but it was shown to be of use when other techniques cannot distinguish between samples.

Gogé *et al.* (2014) used visible and NIR spectroscopy to produce different calibration models built from a local soil database, a national database and a combination of both data sets and to also tested 2 regression algorithms: PLS and fast Fourier transform local weighted (FFT-LW) (a non-linear method) to predict soil properties. The local samples were split into groups and PLS models were built from different numbers of local spectra. PLS models created from the national set were spiked with local spectra and the FFT-LW model was created using 300 neighbours selected from the national data spiked with the local spectra. It was concluded that depending on the property of the soil which is measured, different strategies will perform best and that spiking the national database with locally collected data provided additional value. Guerrero *et al.* (2010) concluded similar results using NIR and nitrogen content values to create PLS calibration models, and were able to accurately predict before and after the models were spiked with locally collected sample (target sites) and the spiking only increased the accuracy. They found that using spectral characteristics was more accurate than using nitrogen values and that using small-sized models integrated the local data more

accurately. Higher accuracies using local models with small sample sizes was observed whereas Gogé *et al.* (2014) suggested a decrease in performance with a decrease in sample size (Stafford, 2013).

1.2.2 X-Ray Fluorescence (XRF) Spectroscopy

X-ray fluorescence (XRF) spectroscopy is a multi-elemental technique that can work with different sample forms, is non-destructive, is able to detect elements with atomic numbers greater than 8 and can be used *in situ*. In an x-ray fluorescence spectrum, the wavelengths present are characteristic of the elements present within the sample. In wavelength dispersive x-ray fluorescence (WDXRF) the samples emitted radiation is diffracted in different directions and a sequential detector moves to detect the x-rays with different wavelengths or a simultaneous detector consisting of fixed single channels to detect specific elements. On the other hand, in energy dispersive XRF (EDXRF) there is only one detector (e.g. (Si(Li))) that is used in combination with a multi-channel analyser according to energies. Although EDXRF is cheaper, WDXRF usually offers greater resolution. Mathematical corrections must be applied to overcome matrix effects that can occur in XRF, which will ensure accurate results are obtained (Levinson, (2001); Krishna *et al.* (2007); Davidson, 2013).

Yu *et al.* (2002) used EDXRF to quantify 19 elements in soil samples and determine the source profiles of these samples. Sixteen samples were collected in total, from 2 different sites in 8 different locations. These 8 different locations possessed different geologies; sedimentary, volcanic or granitic. The authors chose EDXRF over ICP-AES or Atomic absorption spectroscopy (AAS) due to its non destructive nature and the ease of analysing solid samples without the need for digestion.

Singh & Agrawal (2012) acknowledged that X-ray diffraction (XRD) was not a simple technique due to many complexities and so it was combined with EDXRF and AAS for mineral phase identification of soil. Twenty four samples were taken from a 1 hectare field. Five samples were chosen randomly for XRD analysis, with silica removal and grinding with alcohol to avoid matrix effects and peak overlapping problems. EDXRF was found to be very useful in predicting the presence of a chemical/crystalline phase containing specific elements.

Wavelength Dispersive X-ray Fluorescence Spectroscopy (WDXRF) is not a particularly common technique used in soil analysis. Krishna et al. (2007) used sequential WDXRF to determine the levels of 29 major and trace elements (Si, Al, Fe, Mg, Ca, Na, K, Mn, P, Ti, As, Ba, Cd, Co, Cr, Cu, Se, Sr, Mo, Ni, Pb, Rb, S, U, Th, V, Y, Zn, Zr) in agricultural soil samples. Twenty two international reference materials were used to calibrate the spectrophotometer. The samples were not dried prior to analysis as it was recognised this may cause some loss due to evaporation. Matrix effects caused some difficulties but could be corrected using empirical coefficients (alphas) based on count rate, but when there were high concentrations of some elements in certain samples, this was more difficult to correct. Matrix correction of these samples used carried out using empirical formulas based on concentration but if intensity was used, matrix correction was carried out by trial and error. The relative standard deviation (RSD) for most elements was low at less than 5% but for the elements that had higher RSD; this was most likely due to peak suppression and overlapping peaks. Although matrix correction models can produce accurate results this was not the case when there was a high concentration of heavier elements accompanying the lighter elements. This causes a decrease in accuracy unless the standard used was of similar composition to that of the 'unknown' sample.

Despite the fact that this method had low limits of detection (1-2 mg/kg), good precision and accuracy sufficient for use in agricultural monitoring, it may perhaps have potential for use in a forensic context. However, a lower RSD may be required by reducing the peak suppression. Also, 2g of sample was needed to create the pellets used for this analysis and this is considered a bulk sample in a forensic context and this amount of sample will not always be available and so this method will not be of use in trace analysis.

1.2.3 Inductively Coupled Plasma (ICP) Spectroscopy

Inductively coupled plasma spectroscopy can be used with mass spectrometry (ICP-MS), atomic emission spectrometry (ICP-AES) and optical emission spectrometry (ICP-OES). ICP-MS is the fastest growing trace element technique with its most common applications being environmental, geological, biomedical, semiconductor and nuclear fields. ICP is a rapid multielemental technique that can determine concentrations at the ultra trace level. The analysis time is less, detection limits are lower and the fact that it

is multi elemental are just some clear advantages over other atomic spectroscopy techniques such as flame atomic absorption (FAA), electrothermal atomisation (ETA) or ICP-OES (Thomas, 2013).

Pye & Blott (2009) attempted to create a soil database from 1896 soil samples collected in England and Wales from 1999 to 2007 in connection with casework investigations using ICP-MS and ICP-AES. Two laboratories were used to analyse the samples (one third by only ICP-AES and the rest by both techniques) and the data variation was not significant. Methods used to compare soils on the foundation of elemental composition were developed in the author's own laboratory. PCA and Euclidean distances were used to determine the number of elemental concentrations that were indistinguishable for some samples. It was demonstrated that samples that have been taken only a few centimetres apart are likely to be distinguishable based on major and trace elemental concentrations (Stafford, 2013).

After papers demonstrated ICP analysis was a useful technique, Arroyo *et al.* (2009) validated a laser ablation (LA) ICP-MS method for routine soil and sediment analysis. LA-ICP-MS was found comparable to solution ICP-MS and independent proficiency testing using 57 laboratories found the new method was comparable with conventional digestion ICP and AAS methods. With 3 high speed mills a single technician can prepare around 72 samples per day. This method may have been validated and the values for precision, accuracy are given, but nowhere it is stated which validation guidelines were used (Stafford, 2013).

1.3 Statistical Tools

Soil analysis with the use of statistical tools has been carried out for numerous different purposes including environmental, forensic and geological contexts. Guedes *et al.* (2009) investigated the prospect of differentiating between sediment samples collected from beaches and dunes in the north and south of Portugal and Carvalho *et al.* (2013) used a multi-technique approach to distinguish between two river beds over a one year period and investigated if seasonal changes affected the results; cluster analysis was used to determine this. Baron *et al.* (2011) used partial least squares – linear discriminant analysis (PLS-LDA) to determine the possibility of distinguishing different locations within different land use types using FT-IR spectra. Gogé *et al.* (2014) also

used PLS along with principle component analysis (PCA) in an attempt to use visible near infrared (NIR) spectra to create local and national databases and a combination of both to aid in the prediction of soil locations. Guerrero *et al.* (2010) previously explored something similar to Gogé *et al.* (2014), where NIR spectral libraries and models were created using PLS regression and then models were spiked with a few samples from target sites (local samples). Croft & Pye (2004) used 4 different techniques to determine the effectiveness of them on different soil types and 5 footwear types and Cox *et al.* (2000) developed a novel method in which FTIR spectra were collected pre and post pyrolysis which can differentiate samples when other methods cannot.

1.3.1 Artificial Neural Networks

Armenta & de la Guardia (2014) recently reviewed the use of principle component regression (PCR) and PLS, two of the most commonly used techniques for spectral calibration and prediction, and while these are widely used and work well, more sophisticated techniques like artificial neural networks (ANN) have been developed. ANNs are attractive to users because they have incredible information processing characteristics related mainly to nonlinearity, fault and noise tolerance in addition to learning and generalising capabilities (Basheer & Hajmeer, 2000).

Arsoy *et al.* (2013) used a multilayer feed-forward, with back propagation learning ANN to aid in the prediction of soil water content (SWC) using time domain reflectometry (an electromagnetic method) and found the performance of the ANN was better than that of previous calibration models (using unreliable dielectric permittivity of the soil). 50% of the data was used for training the network and 50% for validation, although no testing was carried out. The ANN had an average route mean squared error (RMSE) of 0.009cm³cm⁻³ (for 8 nodes) compared to a range of 0.019-0.033 cm³cm⁻³ for the calibration model.

Most recently, El Haddad *et al.* (2014) applied ANN to process data from an on-site laser-induced breakdown spectroscopy (LIBS) method for soil samples, which were verified by ICP-AES to use as reference values. ANNs were used to determine the relative amounts of silicate, calcareous and ores matrices in the soil samples. Data input into the ANN was not only the raw data but also data with no likeness to the analyte of interest to ensure a good representation of the matrix that allowed correct quantification

of the analyte. Due to the small dataset, the ANN was evaluated by repeating the ANN calculation 5 times, by starting with random weight values to ensure no over fitting occurred. The RMSE was less than 10% when comparing the reference values (from ICP-AES) to the values obtained from the on-site LIBS measurements showing that LIBS is a reliable method.

1.4 Present Work Overview

In this present study, an attempt was made to develop a new method that could reliably and accurately predict the location of known soil samples as well as samples of unknown origin from their FTIR spectra, μ XRF profiles and UV-Visible spectra using feature selection and artificial neural networks. In contrast to previous work, the relationship recognised herein is between the input data and the co-ordinates of the samples locations, as opposed to between the input data and soil classes based on the landscape properties.

A systematic attempt was made based on a machine learning approach with feedforward feature selection and feed-forward neural networks using the values obtained from infrared spectra, μ XRF profiles and UV-Visible spectra as input data. A recursive feature selection using a wrapper method was performed using the Boruta (random forest) classifier algorithm. This was performed in order to reduce the input dimensionality and obtain a much smaller subset of features with the highest possible discrimination whilst maintaining excellent neural network performance. This subset of features was used to train, test and validate a neural network.

The Boruta feature selection method was able to reduce 1430 manually selected initial features to a subset of 43, including features from the FTIR spectra, μ XRF profiles and visible spectra but not from the UV spectra. This new predictor achieved a linear correlation coefficient of 0.86 or 86% accuracy, a great increase from 0.77 when using just FTIR spectra.

2 Materials and Methods

2.1 Sample Collection

Soil samples were collected from Monsanto Park in Lisbon, Portugal, which has an approximate area of 10km². In total, 31 samples were collected in triplicate (total of 93) in 2013 and a further 5 samples were collected in 2014 using a 500 x 500 metre grid of the whole park (Figures 2 and 3). Samples were collected using an in-house built steel core soil sampler with dimensions of 10x3cm, thus samples were collected at a depth of 10cm with a diameter of 3cm and the Global positioning system (GPS) co-ordinates were recorded for each sample site (Table 1) using a Broadcom BCM4750IUB8 GPS receiver.



Figure 2. Map of Monsanto Park, Lisbon, in relation to other cities, showing sample collection sites (created using batchgeo.com). Red = 2013 Samples, Yellow = 2014 Samples.



Figure 3. Satellite image of Monsanto Park, Lisbon, showing sample collection sites (created using batchgeo.com). Red = 2013 Samples, Yellow = 2014 Samples.

Sample Number	Latitude	Longitude	Sample Number	Latitude	Longitude
S01	38° 42' 4"	-9° 13' 27"	S24	38° 43' 23"	-9° 11' 9"
S02	38° 42' 24"	-9° 13' 16"	S25	38° 43' 26"	-9° 10' 50"
S05	38° 42' 49"	-9° 13' 1"	S28	38° 43' 46"	-9° 12' 13"
S06	38° 42' 45"	-9° 12' 52"	S29	38° 43' 46"	-9° 11' 51"
S07	38° 42' 45"	-9° 12' 8"	S32	38° 43' 48"	-9° 10' 51"
S08	38° 42' 51"	-9° 11' 55"	S33	38° 43' 44"	-9° 10' 32"
S11	38° 42' 49"	-9° 10' 50"	S34	38° 44' 7"	-9° 11' 50"
S12	38° 43' 7"	-9° 12' 52"	S36	38° 44' 8"	-9° 11' 10"
S13	38° 43' 5"	-9° 12' 30"	S37	38° 44' 8"	-9° 10' 49"
S14	38° 43' 6"	-9° 12' 5"	S38	38° 44' 5"	-9° 10' 32"
S15	38° 43' 5"	-9° 11' 51"	S39	38° 44' 28"	-9° 11' 50"
S16	38° 43' 8"	-9° 11' 30"	S40	38° 44' 24"	-9° 11' 30"
S17	38° 43' 3"	-9° 11' 6"	S41	38° 44' 29"	-9° 11' 9"
S18	38° 43' 5"	-9° 10' 50"	S42	38° 43' 43"	-9° 11' 47"
S20	38° 43' 26"	-9° 12' 28"	S43	38° 43' 58"	-9° 11' 03"
S21	38° 43' 25"	-9° 12' 3"	S44	38° 43' 42"	-9° 12' 16"
S22	38° 43' 29"	-9° 11' 48"	S45	38° 43' 07"	-9° 11' 55"
S23	38° 43' 27"	-9° 11' 29"	S46	38° 43' 04"	-9° 11' 04"

Table 1. Sample number and location (GPS co-ordinates).

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2.2 Transfer of GPS Co-ordinates to Cartesian Co-ordinates

The input data for the neural network was FTIR spectral data, XRF spectral data and the palynological data with the output being the sample location. This was achieved by converting the GPS co-ordinates to Cartesian Co-ordinates (x and y co-ordinates). This process is necessary due to the curved nature of the earth's surface, so if GPS co-ordinates were used this would give non-linear positioning of the samples. Cartesian co-ordinates are easier for the neural network and it also helps to make the interpretation easier on such a small area. Sample 1 is considered to be the starting point of both the x and y axes and the other samples will be relative to this. The GPS co-ordinates were converted from degrees, minutes and seconds in to decimal degrees using the following equation:

$$c = d + \frac{m}{60} + \frac{s}{3600}$$

Where C is the co-ordinates in decimals, d is the co-ordinates in degrees, m is minutes co-ordinate and s is the seconds co-ordinates. Once converted in to decimal degrees, distances were calculated between samples according to the haversine formula through an in-house built program (table 4).

$$haversin\left(\frac{d}{r}\right) = haversin(\emptyset_2 - \emptyset_1) + \cos(\emptyset_1)\cos(\emptyset_2)haversin(\Lambda_2 - \Lambda_1)$$

Where d is the distance between 2 points (along the great-circle of a sphere), r is the radius of the sphere, $Ø_1$ and $Ø_2$ are the latitude of sample 1 and 2 respectively and Λ_1 and Λ_2 are the longitude or point 1 and point 2 respectively (Shumaker & Sinnott, 1984) with distances in kilometres.

2.3 FTIR Spectroscopy

2.3.1 Sample Preparation

After collection of all samples, the samples were transported to the laboratory in 50mL Falcon[®] flasks. These were then transferred in to separate beakers and dried in an oven at 105°C overnight. Once dried, the samples were sieved through 2mm and 125µm meshes and particles smaller than 125µm were used for the FTIR and UV-Visible analysis.

2.3.2 FTIR Spectroscopy Parameters

The smaller than 125μ m portion of the samples were analysed using a PerkinElmer Spectrum 65 spectrophotometer coupled with an attenuated total reflectance (ATR) accessory. The parameters used were a scan range from 4000-400 cm⁻¹, with a resolution of 4 cm⁻¹, 128 scans and with H₂O/CO₂ correction and the spectra were collected using % transmission. Baseline corrections were carried out manually using the PerkinElmer Version: 10.03.09.0139 software. In between sample application, the ATR crystal was cleaned with 96% ethanol (Purchased from Carlo Erba Reagents) solution and a background scan was performed after every 3 samples.

2.4 µXRF Spectroscopy

2.4.1 Sample Preparation

Sample preparation was carried out using the following procedure before analysis; into a glass beaker, 3.5g of homogenised sample (using a pestle and mortar) was added and placed into an oven at 80-90°C overnight to dry the samples and remove any water. A small portion of the dried samples was then transferred on to an acrylic plate, pressed flat and then placed under the laser beam for analysis.

2.4.2 µXRF Spectroscopy Parameters

A Bruker Artax μ XRF was used for qualitative analysis of the samples. The parameters used were a molybdenum anode, with 25.1 keV of energy. The optic was a capillary 0.060 with an atmosphere of helium. The measurement live time was 120 seconds with a pulse density of 9361cps. The x-ray generator had a high voltage of 50kV and a current of 600 μ A. The X-ray tube used was a MCBM 50-0,6B Mo and the beam size was 17 μ m.

2.5 UV-Visible Spectroscopy

2.5.1 Sample Preparation

Samples were prepared by adding 0.1g of the 125µm portion of each sample to 1mL of deionised water in 2mL eppendorf tubes, vortexed for 20 seconds and centrifuged for 5mins at 10,000rpm.

Some samples were too concentrated and so a dilution was carried out with the following factors:

Sample Dilution Factor		Sample	Dilution Factor		Sample <u>Dilution F</u>		n Factor	
A	UV	VIS	B	UV	VIS	C	UV	VIS
S01	10	0	S01	10	0	S01	10	0
S02	10	0	S02	10	0	S02	10	0
S05	5	0	S05	10	0	S05	10	0
S06	5	0	S06	10	0	S06	10	0
S07	5	0	S07	10	0	S07	10	0
S08	50	5	S08	20	0	S08	20	0
S11	10	0	S11	10	0	S11	10	0
S12	20	0	S12	20	0	S12	20	0
S13	20	0	S13	20	0	S13	40	0
S14	20	0	S14	20	0	S14	40	0
S15	50	2	S15	20	0	S15	100	5
S16	100	0	S16	10	0	S16	20	0
S17	10	0	S17	20	0	S17	20	0
S18	40	0	S18	40	0	S18	50	0
S20	5	0	S20	10	0	S20	20	0
S21	50	0	S21	50	0	S21	100	0
S22	50	0	S22	10	0	S22	10	0
S23	50	0	S23	20	0	S23	20	0
S24	5	0	S24	10	0	S24	10	0
S25	20	0	S25	20	0	S25	20	0
S28	20	0	S28	20	0	S28	20	0
S29	10	0	S29	10	0	S29	10	0
S32	5	0	S32	20	0	S32	10	0
S33	50	0	S33	20	0	S33	40	2
S34	5	0	S34	20	0	S34	10	0
S36	50	0	S36	50	0	S36	50	5
S37	5	0	S37	10	0	S37	20	0
S38	50	0	S38	100	0	S38	50	0
S39	10	0	S39	20	0	S39	20	0
S40	50	0	S40	50	2	S40	50	0
S41	10	0	S41	20	0	S41	20	0

Table 2. Dilution factors of the samples for the UV-Visible spectroscopy.

2.5.2 UV-Visible Spectroscopy Parameters

A PerkinElmer Lambda 25 was used to collect ultraviolet (UV) spectra from 400-200cm⁻¹ and visible spectra (VIS) from 1100-401cm⁻¹ using the coloured liquid. After the spectra were collected, the absorbance values were corrected by multiplying the absorbance by the dilution factors used for each sample.

2.6 Palynological Analysis

Soils were prepared for pollen analysis using the following procedure: a few crystals of each soil were placed in separate glass tubes, 10% potassium hydroxide (KOH) was then added and the tubes placed in a dry bath at 100°C for 20 minutes. The samples were then sieved using a 180-230µm sieve, to remove debris; the samples were washed with 10% KOH, and then centrifuged at 3000 rpm for 5 minutes and the liquid decanted. The samples were mixed with 10% KOH and the tubes were then placed in a dry bath at 100°C for 20 minutes. The samples were then vortexed, centrifuged at 3000 rpm for 5 minutes and the liquid decanted. The samples were then washed with distilled water, vortexed, then centrifuged at 3000 rpm and the liquid decanted (the samples were switched to polypropylene centrifuge tubes with lids). Added to the sample was 37% hydrochloric acid (HCl) (Purchased from Carlo Erba Reagents), the samples were vortexed, and centrifuged at 3000 rpm for 5 minutes and the liquid decanted. Samples were left for 24-48hours after 40% hydrofluoric acid (Purchased from Sigma-Aldrich) was added. The samples were then vortexed, centrifuged at 3000 rpm for 5 minutes and the liquid decanted. The samples were washed with distilled water, vortexed, centrifuged at 3000 rpm for 5 minutes, the liquid was decanted and the samples vortexed again. The samples were washed with glacial acetic acid (Purchased from Carlo Erba Reagents), vortexed, centrifuged at 3000 rpm for 5 minutes and the liquid decanted. The samples were then switched to glass tubes. Acetolysis (9:1 acetic anhydride: sulphuric acid) (Purchased from Panreac) was then carried out by adding the mixture to the samples which were then placed in a dry bath at 98°C for 8 minutes. The samples were again centrifuged at 3000 rpm for 5 minutes and the liquid decanted. The samples were washed with distilled water, vortexed, centrifuged at 3000 rpm for 5 minutes, the liquid was decanted and the sample vortexed again. The samples were then switched to eppendorf tubes. A few drops of glycerine (Purchased from CMD Chemicals) were added to the samples in the eppendorf tubes, the tubes were vortexed, centrifuged at 3000 rpm for 5 minutes and the liquid decanted. A small quantity of the sample was placed on to a microscope slide, which was covered with a cover glass slip and sealed with a small amount of paraffin. Samples were then analysed using an Olympus CX21 biological microscope at 1000X magnification and the pollen grains were identified and counted to a maximum of 100 per slide.

The deionised water was produced in-house with a resistance of $15M\Omega$ using a Helix 10 Millipore and potassium hydroxide was made using KOH pellets purchased from EKA Chemicals with deionised water.

2.7 Input Vectors

Pre-processing of feature vectors were carried out prior to the training of the neural network. All IR spectra were manually baseline corrected, so the baseline was a maximum of 100% transmittance. The UV-Visible spectra were normalised correcting for the dilution factors. Discriminatory peaks present in all the spectra for UV, visible, FTIR and XRF were then manually selected. 20 features were manually selected for visible, 6 for UV and 27 features for both FTIR and μ XRF. The relationship between each of these features was then computed for each method separately.

2.8 Feature Selection

Feature selection was performed on 1430 manually selected features with the purpose of reducing the dimensionality of the input vectors, with the aim of ascertaining a subset of features, with the smallest size which provided the highest possible discrimination between samples. A recursive feature selection wrapper method (Boruta) based on random forest was used in R version 2.15.2 (R Development Core Team, 2008). Input vectors were computed by the feature selection method used and consequently used for the training and selection of one neural network.

2.9 Artificial Neural Network

MATLABs' (The MathWorks, 2011a) Neural Networks Toolbox (The MathWorks, 2011b) was used to develop feed forward fully connected neural networks. The neural networks weights and biases were initialised using the Nguyen-Widrow layer initialisation function, which initialises weights and biases randomly although evenly across all layer's input space. The symmetric sigmoid function was the selected

activation function for the hidden layer and the linear function was chosen for the output layer. The scaled conjugate gradient back propagation (backward propagation of errors) was used as the learning algorithm and the mean absolute error was the performance measure used to stop training. The number of neuron present in the hidden layer was computed based on the number of dimensions of the feature vectors and the number of neurons in the output layer was two, corresponding to x,y co-ordinates of the samples. A general topology of a neural network is shown in figure 4.



Figure 4. General topology of the created neural network, where i relates to the number of the inputs present in the input vector, w relates to the weights, b relates to the bias, h is the number of neurons in the hidden layer and o is the output vector number.

Neural network training was performed after the input vector processing, where the input sequence datasets were divided into three sub-datasets, the training, testing and a validation dataset consisting of 70%, 15% and 15% of the samples respectively which was selected randomly by the neural network. The best neural network was selected for the feature vectors computed through the orthogonal encoding scheme from a total of 1000 trained neural networks. This selection was based on accuracy and standard deviation values of the training, test, validation and overall dataset. The selected neural networks were then externally validated using the external validation dataset.

3 Results and Discussion

3.1 Sample Collection

Some of the planned sites for sampling were not possible to reach due to the land being privately owned and thus no permission to collect and so samples were collected at the nearest achievable location.

Table 3. GPS co-ordinates, decimal degrees and Cartesian co-ordinates of the samples
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Sample	Sample	GPS Co	o-ordinates	ordinates Decimal Degrees		Cartesian Co-ordinates		
Year	Number	Latitude	Longitude	Latitude	Longitude	X / Km	Y / Km	
	S01	38° 42' 4"	-9° 13' 27"	38.7011	-9.2242	0.00	0.00	
	S02	38° 42' 24"	-9° 13' 16"	38.7067	-9.2211	0.27	0.62	
	S05	38° 42' 49"	-9° 13' 1"	38.7136	-9.2169	0.63	1.39	
	S06	38° 42' 45"	-9° 12' 52"	38.7125	-9.2144	0.84	1.27	
	S07	38° 42' 45"	-9° 12' 8"	38.7125	-9.2022	1.90	1.27	
	S08	38° 42' 51"	-9° 11' 55"	38.7142	-9.1986	2.22	1.45	
	S11	38° 42' 49"	-9° 10' 50"	38.7136	-9.1806	3.78	1.39	
	S12	38° 43' 7"	-9° 12' 52"	38.7186	-9.2144	0.84	1.95	
	S13	38° 43' 5"	-9° 12' 30"	38.7181	-9.2083	1.37	1.88	
	S14	38° 43' 6"	-9° 12' 5"	38.7183	-9.2014	1.98	1.92	
	S15	38° 43' 5"	-9° 11' 51"	38.7181	-9.1975	2.31	1.88	
	S16	38° 43' 8"	-9° 11' 30"	38.7189	-9.1917	2.82	1.98	
	S17	38° 43' 3"	-9° 11' 6"	38.7175	-9.1850	3.40	1.82	
	S18	38° 43' 5"	-9° 10' 50"	38.7181	-9.1806	3.78	1.88	
	S20	38° 43' 26"	-9° 12' 28"	38.7239	-9.2078	1.42	2.53	
2013	S21	38° 43' 25"	-9° 12' 3"	38.7236	-9.2008	2.02	2.50	
	S22	38° 43' 29"	-9° 11' 48"	38.7247	-9.1967	2.39	2.63	
	S23	38° 43' 27"	-9° 11' 29"	38.7242	-9.1914	2.84	2.56	
	S24	38° 43' 23"	-9° 11' 9"	38.7231	-9.1858	3.33	2.44	
	S25	38° 43' 26"	-9° 10' 50"	38.7239	-9.1806	3.78	2.53	
	S28	38° 43' 46"	-9° 12' 13"	38.7294	-9.2036	1.78	3.15	
	S29	38° 43' 46"	-9° 11' 51"	38.7294	-9.1975	2.31	3.15	
	S32	38° 43' 48"	-9° 10' 51"	38.7300	-9.1808	3.76	3.21	
	S33	38° 43' 44"	-9° 10' 32"	38.7289	-9.1756	4.22	3.09	
	S34	38° 44' 7"	-9° 11' 50"	38.7353	-9.1972	2.34	3.80	
	S36	38° 44' 8"	-9° 11' 10"	38.7356	-9.1861	3.30	3.83	
	S37	38° 44' 8"	-9° 10' 49"	38.7356	-9.1803	3.81	3.83	
	S38	38° 44' 5"	-9° 10' 32"	38.7347	-9.1756	4.22	3.74	
	S39	38° 44' 28"	-9° 11' 50"	38.7411	-9.1972	2.34	4.45	
	S40	38° 44' 24"	-9° 11' 30"	38.7400	-9.1917	2.82	4.32	
	S41	38° 44' 29"	-9° 11' 9"	38.7414	-9.1858	3.33	4.48	
	S42	38° 43' 43"	-9° 11' 47"	38.7286	-9.1964	2.41	3.06	
	S43	38° 43' 58"	-9° 11' 03"	38.7328	-9.1842	3.47	3.52	
2014	S44	38° 43' 42"	-9° 12' 16"	38.7283	-9.2044	1.71	3.03	
	S45	38° 43' 07"	-9° 11' 55"	38.7186	-9.1986	2.22	1.95	
	S46	38° 43' 04"	-9° 11' 04"	38.7178	-9.1844	3.45	1.85	

3.2 FTIR Spectra

Figures 5 and 6 show the FTIR spectra of the triplicates collected for samples 1 and 2. The spectral profiles both samples are quite similar, including the transmittance of the peaks and this is the same across most of the samples, so to the naked eye it is difficult to differentiate between some samples. Figure 7 shows sample 11 has a very different profile across most of the spectrum with a peak at 2500cm⁻¹ that is not present in figures 5 or 6. This was not surprising as the colour of sample 11 was light beige whereas samples 1 and 2 were dark brown and thus the organic composition differs greatly between the samples. The spectrum from 549-400cm⁻¹ has been removed from all spectra due to the increased noise and low resolution present in this region. All other sample spectra are present on the appendix disk.



Figure 5. FTIR spectra of the triplicate samples collected from location 1.



Figure 6. FTIR spectra of the triplicate samples collected from location 2.



Figure 7. FTIR spectra of the triplicate samples collected from location 11.

3.3 µXRF Profiles

Like with the FTIR spectra the XRF profiles sample 1 and 2 (figures 8 and 9) are very similar although the counts for the different elements differ between samples and the ratios between the different peaks differ also. Figure 10 (Sample 11) has a very different profile to samples 1 and 2 which was not unexpected as the FTIR spectra were very different. All other sample spectra are present on the appendix disk.



Figure 8. µXRF spectra of the triplicate samples collected from location 1.



Figure 9. μ XRF spectra of the triplicate samples collected from location 2.



Figure 10. µXRF spectra of the triplicate samples collected from location 11.

3.4 UV-Visible Spectra

3.4.1 UV-Spectra

Figures 11 and 12 show the corrected UV spectra of samples 1 and 2 have very different profiles across the spectrum but there are some similarities. There are clear differences in the absorbances of each spectrum, within the triplicate samples as well as between the samples. The peak at 260nm and the trough at 220nm is present in almost all samples although the profile to the left and right of the peak and the absorbances differ between samples. Figure 13 shows sample 11B and C have similar profiles to that of samples 1C and 2C. Sample 11A has a very different profile to that of 11B and C as well as samples 1 and 2. All other sample spectra are present on the appendix disk.



Figure 11. UV spectra for the triplicate samples collected from location 1.



Figure 12. UV spectra for the triplicate samples collected from location 2.



Figure 13. UV spectra for the triplicate samples collected from location 11.

3.4.2 Visible Spectra

Figures 14 and 15 show the normalised visible spectra of samples 1 and 2 have similar profiles across the spectrum but there are distinct differences in the absorbance values, where sample 2 has absorbances almost twice that of sample 2. This is the case when looking at the spectra of all the other samples, with very little specific peaks or discrete features present in the visible region. Figure 16 shows sample 11 which has a similar profile to samples 1 and 2 although the B samples have higher absorbance's than the A and C samples. All other sample spectra are present on the appendix disk.



Figure 14. Visible spectra for the triplicate samples collected from location 1.



Figure 15. Visible spectra for the triplicate samples collected from location 2.



Figure 16. Visible spectra for the triplicate samples collected from location 11.

The actual profile and values of the peaks present in the samples is not of crucial importance as the neural network uses the relationship between the peaks present in the spectra.

3.5 Palynological Analysis

Due to the samples over processing, no surface features were left on the pollen, making it impossible to identify them and thus unfortunately it was not possible to complete the palynological analysis due to the method failing to work as expected.

3.6 Selected Features

Peaks were manually selected from the different spectra. From the visible spectra 5 features, from UV 3 features, from FTIR 27 features and XRF 27 features were identified. The relationship between each feature was computed by dividing each feature by the others. For example, if there are 3 features, A, B and C. The total number of features was calculated by A/B, A/C, B/A, B/C, C/A and C/B, so now there is a total of 6 features selected. This was carried out for each different technique separately. In total, 1430 features were selected for the 4 different methods which were then submitted to Boruta.

3.7 Feature Selection

The Boruta algorithm was used, which selected a total of 43 features from the 1430 total (table 4 in appendix) and these features were then used in the training of the neural network. There were 25 features selected from the FTIR, 15 from μ XRF, and 3 from the visible spectra, however, no features were selected from the UV spectra.

3.8 Neural Network

Initially, 1000 neural networks were created and the best one was chosen. Figure 17 shows the training (figure 17a) had the greatest accuracy of 90%, the accuracy of the testing (figure 17b) had a much lower accuracy of 74% and validation (figure 17c) had an accuracy of 79%. Figure 18 shows that the overall linear correlation factor (accuracy) obtained was 86%. The error histogram that was obtained (figure 19) shows the errors are mainly occurring around zero for the input vectors. Figure 20 shows the MSE for training, testing and validation and the plot shows that the neural network is not to over fitted and the MSE's are very low.



Figure 17. Plot showing the accuracies for the (a) training, (b) testing and (c) validation of the neural network.



Figure 18. Regression plot of classifier output and expected outcomes for the neural network.



Figure 19. Error histogram showing the number of instances per interval of error observed (blue bars for training, red bars for testing and green bars for validation).



Figure 20. Plot of the MSE for the training (blue line), testing (red line) and validation (green line) of the neural network.

4 Conclusion

The neural network was able to produce a correlation factor of 86% using the Boruta (random trees) algorithm, using FTIR, μ XRF and visible spectroscopy data. No features were selected from the UV spectra showing that UV is not a useful technique to use in this method. Only 3 features were manually identified in this region, but these were not discriminatory enough to be used to differentiate the different samples, thus they were not selected during the feature selection process. It was also not possible to collect and therefore use any palynological data, due to the samples being over processed since method failed to work as expected.

The accuracy achieved with this study (86%) is a marked improvement on 77% using just FTIR spectra, so it can be concluded that by adding μ XRF and visible spectroscopy data the accuracy of the prediction is greatly increased. The improved accuracy of this method demonstrates how powerful multiple techniques can be in soil analysis and that this is a strong method that could be widely used.

Taking into consideration that the FTIR crystal showed clear signs of contamination due to soil residues and other samples, it was not in perfect condition and so the obtained results could be somewhat explained by this. This was also the reason that the 2014 samples were not analysed due to the condition of the crystal worsening overtime as a result of high sample throughput. Since the 2014 samples could not be analysed by FTIR, they were not analysed by the other techniques as the data for these samples could not be used in the neural network as the data would be incomplete. A WDXRF instrument was the machine intended to be used but this was not possible so a μ XRF was used which is not as sensitive to trace elements and this could also explain, in part, the correlation coefficient.

5 Recommendations for Further Work

Prospects for the future could be to increase the amount of data input into the neural network by using more techniques which should increase the accuracy of the prediction.

Increase the number of samples by expanding the grid used and reducing the distance between collection sites which will increase the usefulness of this technique.

The samples could be re-analysed on a FTIR with a cleaner or new crystal to reduce noise in the spectra.

Analyse the 2014 samples using FTIR, μ XRF and visible spectroscopy to determine if the method is reproducible across different years.

Also, test different feature selection methods, to see if this increases the number of selected features and improves the overall accuracy of the neural network.

Eventually, create a map the whole of Portugal using this method and possibly expand to other countries for use by police forces, armies and naval fleets.

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Appendix

Feature Number	V1	V 2	Feature Number	V1	V2
6	VIS_980	VIS_905	590	FTIR_700	FTIR_755
10	VIS_905	VIS_980	592	FTIR_700	FTIR_725
15	VIS_800	VIS_905	619	FTIR_671	FTIR_710
54	FTIR_1747	FTIR_1630	683	FTIR_569	FTIR_990
80	FTIR_1630	FTIR_1747	709	FTIR_555	FTIR_990
190	FTIR_990	FTIR_910	723	FTIR_555	FTIR_710
192	FTIR_990	FTIR_870	866	XFR_3.292	XFR_4.4983
216	FTIR_925	FTIR_910	867	XFR_3.292	XFR_4.9193
242	FTIR_910	FTIR_925	942	XFR_4.4983	XFR_3.292
293	FTIR_870	FTIR_990	948	XFR_4.4983	XFR_6.4837
294	FTIR_870	FTIR_925	953	XFR_4.4983	XFR_7.9351
319	FTIR_850	FTIR_990	955	XFR_4.4983	XFR_8.6514
351	FTIR_810	FTIR_800	958	XFR_4.4983	XFR_14.0296
352	FTIR_810	FTIR_785	968	XFR_4.9193	XFR_3.292
377	FTIR_800	FTIR_810	1118	XFR_7.3508	XFR_18.3397
403	FTIR_785	FTIR_810	1140	XFR_7.5832	XFR_14.0296
409	FTIR_785	FTIR_725	1179	XFR_7.9351	XFR_4.4983
435	FTIR_775	FTIR_725	1192	XFR_7.9351	XFR_14.0296
486	FTIR_755	FTIR_737	1231	XFR_8.6514	XFR_4.4983
538	FTIR_725	FTIR_755	1309	XFR_14.0296	XFR_4.4983
541	FTIR_725	FTIR_700	1335	XFR_15.7197	XFR_4.4983
568	FTIR_710	FTIR_671			

 Table 4. Features selected by the Boruta feature selection algorithm, showing the relationship between peaks.

See Appendix Disk for:

FTIR Spectra

UV Spectra

Visible Spectra

XRF Spectra

Neural Network Matlab file.