Spectral analysis of blood stains at the crime scene

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Hyperspectral imaging integrates conventional imaging and spectroscopy, to obtain both spatial and spectral information from a sample. This technique enables investigators to analyze the chemical composition of traces and simultaneously visualize their spatial distribution. Hyperspectral imaging offers significant potential for the detection, visualization, identification and age estimation of forensic traces. The rapid, non-destructive and non-contact features of hyperspectral imaging mark its suitability as an analytical tool for forensic science. This chapter provides an overview of the principles, instrumentation and analytical techniques involved in hyperspectral imaging. We describe recent advances in hyperspectral imaging technology motivating forensic science applications, e.g. the development of portable and fast image acquisition systems. Reported forensic science applications are reviewed. Challenges are addressed, such as the analysis of traces on backgrounds encountered in casework, concluded by a summary of possible future applications.
2.1. INTRODUCTION

The detection and identification of forensic traces are crucial in crime scene investigations. For this purpose a wide range of techniques is available, including chemical enhancement techniques and the use of light sources with 15 to 30 nm bandwidths, which increase the contrast between a trace and its background. Many of these techniques are, however, either destructive or subject to human interpretation. Hyperspectral imaging is suitable for the non-contact identification of evidence, thus minimizing the risk of contamination and destruction of traces. Hyperspectral imaging integrates conventional imaging and spectroscopy to obtain a three dimensional data set containing both spatial and spectral information of a specimen. In addition, analysis of the temporal behaviour of spectra can give insight in the chemical changes within the specimen, which can be used for age estimation purposes. Estimation of the age of forensic traces provides investigators with valuable information, which can assist the reconstruction of the timeline of events.

Hyperspectral imaging was originally developed for remote sensing applications utilizing satellite imaging data of the earth but has since found application in such diverse fields as food science, pharmaceuticals and medical diagnostics. Hyperspectral images are analogous to a stack of images, each acquired at a narrow spectral band. Like spectroscopy, hyperspectral imaging can be applied in different parts of the electromagnetic spectrum, e.g. ultraviolet (UV), visible (Vis), near infrared (NIR), mid infrared (IR) or even the thermal infrared range. In these regions reflectance, transmission, photoluminescence, luminescence or Raman scattering can be recorded by hyperspectral cameras with a spectral resolution similar to miniature spectrographs. Spatial resolutions can be adapted to the application, which range from microscopic to landscapes. Advantages of hyperspectral imaging include speed of data acquisition, reduction of human error, no destruction of traces, no specimen preparation, and the ability to illustrate the results.

Hyperspectral imaging is a powerful emerging tool for the analysis of forensic traces. Latent traces can be detected and visualized by using spectral differences to obtain optimal contrast between a trace and its background.
Individual spectra give information about the chemical composition of the specimen, which is useful for identification and quantification purposes, and the spatial distribution of traces is simultaneously recorded. In the last decade, hyperspectral imaging has proven to be a valuable technique for the imaging of latent fingerprints and the detection of trace materials within these prints. Hyperspectral imaging is also emerging in other fields of forensic science and has shown its value in comparative research of materials including fibres, paint chips, or inks, where the question arises whether two traces share common origin. The possibility of viewing spectral and spatial information side by side is advantageous in these cases.

Recent developments in hyperspectral imaging technology offer added potential for forensic science investigations. Because hyperspectral imaging systems are becoming increasingly portable, they may be used at the scene of investigation, where traces can be viewed and interpreted in the original context. The development of fast scanning systems enables investigators to scan a complete scene, which reduces the workload in forensic laboratories and quickly provides investigators with valuable information which can lead the investigation.

This chapter gives an overview of the principles, instrumentation and analytical techniques involved in hyperspectral imaging, followed by a review of recent forensic science applications. We limited our scope to hyperspectral imaging applications using reflectance, photoluminescence, transmission or Raman scattering. Because forensic traces are typically encountered in many different environmental circumstances, their analysis brings specific challenges, which are also addressed. To conclude, possible future applications are summarized.

2.2. HYPERSPECTRAL IMAGING

2.2.a INTERACTION OF LIGHT AND MATTER

The interaction between light and a specimen is determined by the optical properties of the specimen and the incident light. As hyperspectral imaging
measures such interaction, it may be used to characterize the material. In practice this involves illumination of the object under investigation. Commonly, the first interaction will be on the specimen surface where part of the light will be reflected (Figure 2.1a). This part contains no to little information from within the medium but is governed by the index of refraction difference between media. Upon entering the material, the light can be scattered or absorbed.

![Figure 2.1](image.png)

Figure 2.1. The interaction of light with a specimen may lead to a) specular reflection, b) elastic scattering followed by diffuse reflection, c) inelastic scattering followed by emission of Raman shifted light (dotted lines), d) absorption, and e) absorption followed by photoluminescence emission (dashed lines).

Scattering is the process by which light interacts with structures in a specimen and causes a change in direction of propagation, depending on the wavelength, size of the particle and index of refraction differences (Figure 2.1b). The majority of light is scattered at the identical wavelength of the incident light, a process referred to as elastic scattering. There may also be a small fraction that will be inelastically scattered (Raman scattering) which will cause wavelength shifts corresponding to the vibrational states of the molecules in the specimen (Figure 2.1c). Raman scattering can be measured to chemically analyze the scattering specimen.

The absorption properties of a chemical compound are wavelength dependent. Absorption in the visible wavelength range corresponds to the electronic states of the molecule, while absorption in the NIR and IR is
determined by the vibrational modes. Upon relaxation, return to the ground state, the energy will be released in the form of radiation (heat or photoluminescence) or by transfer to another molecule. So both the spectral absorption and, if present, the induced photoluminescence can be measured to identify the chemical contents of a specimen using hyperspectral cameras in reflectance, or transmission mode (Figure 2.1d/e). Quantitative analysis, however, is complicated because the length of the path travelled by the detected light depends on the optical properties of the specimen.

2.2.b HYPERCUBE FORMATION

Figure 2.2. Hypercube of a blood stain, with two spatial (x,y) and one wavelength ($\lambda$) dimension. From the hypercube an image plane is shown for one wavelength ($\lambda_i$) and a spectrum is obtained from one pixel ($x_j, y_k$).
Hyperspectral images are analogous to a stack of images, each acquired at a narrow spectral band. The resulting data set is a three-dimensional block of data, the so-called hypercube, with two spatial \((x,y)\) dimensions and one wavelength \((\lambda)\) dimension (Figure 2.2). This hypercube provides images for each wavelength \((\lambda_i)\) and a spectrum can be obtained from each individual pixel \((x_j,y_k)\), as depicted in Figure 2.2.

Obtaining information in all three dimensions of a hypercube simultaneously is currently not feasible; hyperspectral instruments can only capture two dimensions at a time. Temporal scanning is needed to create a three-dimensional hypercube by stacking the two-dimensional data in sequence. There are three ways of acquiring a hypercube (Figure 2.3), commonly known as point scanning (or whiskbroom), line scanning (or pushbroom), and area scanning (or stare down). These descriptive names refer to the hardware methodology used to acquire the hypercubes:

- In a point scanning system, a complete spectrum is acquired at a single point. Light originating from this point enters the objective lens and is separated into different wavelengths by a spectrometer and detected by a linear array detector. Once spectral acquisition is completed, the spectrum of another point can be recorded. Scanning has to be performed in both spatial directions to complete the hypercube.

- In the case of line scanning systems, the spectra of all pixels contained in one image line are acquired simultaneously. The light is dispersed onto a two-dimensional charge coupled device (CCD) detector. This way, a two-dimensional data matrix with the spectral dimension and one spatial dimension is acquired. The second spatial dimension of the hypercube is achieved by scanning across the specimen surface in a direction perpendicular to the imaging line. This means that relative movement between the object and detector is necessary, which may be achieved either by moving the specimen (e.g., using a translation stage or a conveyor belt) and keeping the hyperspectral camera in a fixed position or by moving the camera and keeping the specimen fixed.

- An area scanning system also acquires a two-dimensional data matrix but in this case the data represent a more conventional image with two
spatial axes. A complete hypercube is obtained by collecting a sequence of these images for one wavelength band at a time. The wavelength of incoming light in this configuration is typically modulated using a tuneable filter.

Figure 2.3. Methods for acquiring three-dimensional hypercubes: (a) point scanning, (b) line scanning, and (c) area scanning. Hypercubes contain two spatial (x,y) and one spectral (λ) dimension. Blue areas represent data acquired by one scan. Red arrows represent temporal scanning required to complete the hypercube.

2.2.c SYSTEM OPTIMISATION FOR FORENSIC SCIENCE APPLICATIONS

Typical hyperspectral imaging systems contain the following components: objective lens, wavelength modulator, detector, illumination, and acquisition system (Figure 2.4). All these components can be adjusted to the requirements of the application. The forensic environment of analysis may range from laboratory to field conditions, whereas the areas of interest may range from the microscopic to landscapes. As for conventional imaging, different objective lenses can be chosen to obtain the right spatial resolution for each application, e.g. macroscopic lenses, zoom lenses, wide angle lenses etc. For analysis on a microscopic scale, the hyperspectral imaging system can be coupled to a microscope.
Acousto-Optic Tuneable Filters (AOTFs)\(^7\) and Liquid Crystal Tuneable Filters (LCTFs)\(^8\) are the two most common wavelength modulators employed. Major drawbacks of these filters are their size and costs. Recently, hyperspectral imaging systems have been developed and commercialised using Fabry-Pérot filters (Innopharma labs, Ireland). Another recent innovation employs the use of a tuneable laser system based on Optical Parametric Oscillator (OPO) technology\(^9\), which replaces the broadband light source, thus removing the need for a wavelength modulator. The benefits of Fabry-Pérot filters and tuneable lasers compared to tuneable filters are their small size and weight, speed of wavelength tuning and high optical throughput. The development of these new technologies offers potential for low cost, hand held, portable hyperspectral imaging with the desired resolution for trace analysis in forensic science applications.

After the light is separated into different wavelengths the detector, e.g. a charge coupled device (CCD), measures the intensity of the collected light. The pixels in the CCD sensor can be arranged in one-dimensional or two-
dimensional arrays, resulting in line detectors and area detectors. Detectors for the mid-infrared region are also available, such as lead selenide (PbSe), indium antimonide (InSb), and mercury cadmium telluride (MCT). To ensure sensitivity of the detector to low light intensities in the infrared regions, the detector may have to be cooled. The CMOS image sensor is another detector that has the potential to compete with CCD. Typical advantages as high speed, low cost, low power consumption, and small size for system integration make them prevail in the consumer electronics market (e.g. low-end camcorders and cell phones). However, the dynamic range and the sensitivity are lower than those of CCD detectors.

The choice of the light properties (broadband vs. monochromatic, specular vs. diffuse, etc.) and consequently the illumination source (halogen, LED, laser, etc.) and lighting arrangement are crucial for the performance and reliability of the system. Halogen lamps, commercially available in various forms, are most common broadband illumination sources used in hyperspectral applications. Halogen lamps can be used directly to illuminate the target (like room lighting) or can be delivered through an optical fibre. Light emitting diode (LED) technology has advanced rapidly during the past few years, and both narrowband and broadband light generators are currently available in the market. This technology is a relatively inexpensive, robust and reliable alternative to halogen lighting, and its use for hyperspectral imaging is likely to expand in the near future, with particular benefits for portable systems. Unlike broadband illumination sources, lasers are powerful directional monochromatic light sources, which make them interesting candidates for photoluminescence and Raman applications. The use of aforementioned tuneable light sources is still limited in hyperspectral imaging applications but they offer promising scope for specific applications including trace analysis. The implementation of digital micro-mirror devices (DMDs) is another recent development in hyperspectral imaging. In this setup, only the region of interest is illuminated. Such systems reduce variations in the spectra arising from scattered light from the background and nearby objects.

Finally, the image acquisition system can be optimized for the application. The desire for on-line monitoring within the process industries has
seen the emergence of real-time online systems typically employing line-scanning approaches. This line scanning setup also offers potential for large-scale forensic science applications, where instead of using a moving stage or conveyor to pass a specimen or product past the detector, the detector itself is moved over a large stationary area of interest such as a wall, a floor or an entire scene of investigation (see Figure 2.5).

![Hyperspectral imaging system at a simulated crime scene.](image)

**Figure 2.5. Hyperspectral imaging system at a simulated crime scene.**

### 2.2.d DATA ANALYSIS

Upon detection, analysis of the data provided in the hypercube is required. Grahn and Geladi detail the types of treatment that can be applied\(^\text{12}\). A summary of processing steps is given below.

**Calibration**

The raw data in a hypercube not only result from the chemical composition of a specimen, but also from the illumination intensity, the sensitivity of the detector and the transmission of the optics\(^\text{13}\). The influence of these factors is a function of wavelength, but may also show spatial variations. Spectral and spatial calibration is required to compensate for this. Calibration measurements
typically performed for reflectance measurements consist of acquisition of the
dark response of the system, measured while covering the lens and dimming
the light source, and the response of a uniform, high reflectance reference.
Using these data, the reflectance (R) can be calculated as follows:

\[ R = \frac{I_{\text{specimen}} - I_{\text{dark}}}{I_{\text{reference}} - I_{\text{dark}}}, \]

where \( I_{\text{specimen}} \) is the intensity of the reflectance measurement of the specimen,
\( I_{\text{dark}} \) the intensity of the dark response, and \( I_{\text{reference}} \) the intensity of the uniform
reference.

It is considered good practice to carry out calibration on a daily basis,
as small changes in electrical power sources, illumination, detector response
and system alignment may result in changes in the detected response. Inclusion
of internal reference standards in each hyperspectral image acquired is also
recommended, which allows monitoring the performance of the system over
time.

**Spectral pre-processing**

Spectral information can be used to gain knowledge about the chemical
composition of a specimen. However, several non-chemical origins cause
systematic variations between spectra, unrelated to the chemical composition
of a specimen, including specular reflections, scattering effects due to surface
inhomogeneities, varying object – illumination distances and random noise. A
number of spectral pre-processing techniques can be applied to reduce these
variations, e.g. smoothing, offset correction, normalization, mean centring,
standard normal variate (SNV) correction\(^1\), multiplicative scatter correction
(MSC)\(^1\), and Savitzky-Golay differentiation\(^16\). The effect of MSC and
differentiation is demonstrated in Figure 2.6; MSC removes variation resulting
from scattering effects, first and second order differentiation eliminate a
constant offset or linear baseline, respectively, and can be used to resolve
overlapping peaks originally appearing as shoulders.
Spectral analysis

Spectral data analysis attempts to address what different components are present in the hypercube, in which concentration and how they are distributed. In some cases the intensity at a single wavelength, the integrated intensity (area) under a spectral peak, or ratios of intensities at different wavelengths can be sufficient for the analysis. However, using these methods, the large amount of spectral information available is not completely exploited. To reduce the amount of variables, while keeping the maximum of variation in the data,
Principal Component Analysis (PCA) can be applied, which is a popular multivariate chemometric method.

In general, spectra are compared to other spectra in the hypercube or to reference spectra from an external library using a similarity measure, e.g. the Euclidean distance, Pearson’s correlation coefficient or the spectral angle\textsuperscript{17}. Spectral unmixing can be applied to decompose a measured spectrum into a collection of constituent spectra\textsuperscript{18,20}. Clustering and classification techniques use the spectral information contained in the hypercube and identify regions with similar spectral characteristics. Clustering techniques are unsupervised methods, e.g. k-nearest neighbors\textsuperscript{21}, which require no a priori information about the dataset to achieve clustering. Supervised classification methods, including partial least squares discriminant analysis\textsuperscript{22}, and spectral angle mapping\textsuperscript{23}, require the selection of well-defined and representative calibration and training sets for classifier optimization. On the other hand, hyperspectral image regression enables the prediction of constituent concentrations at the pixel level, thus enabling their spatial distribution in a specimen to be visualized. Numerous approaches are available for the development of regression models (e.g. partial least squares regression, principal components regression\textsuperscript{15}).

### Image processing

Image processing is performed to convert the contrast developed by the previous steps into a picture showing the component distribution. Additionally, a single wavelength image can be selected showing the highest contrast between different components. Greyscale or colour mapping with intensity scaling is frequently used to display compositional contrast between pixels in an image. False colour mapping, in which two or more images at different wavebands are combined to form a new colour image may be employed to enhance apparent contrast between distinct regions of a specimen.

An interesting other approach for presenting the results was demonstrated by Alsberg et al\textsuperscript{24}, who projected the results of hyperspectral
imaging analysis back onto the scene to highlight chemical differences otherwise invisible to the naked eye. This non-destructive technique provides information similarly to traditional forensic methods, e.g. the use of luminol to highlight blood stains at a scene of investigation, and can be useful to guide investigators in their search for traces.

2.3. FORENSIC SCIENCE APPLICATIONS

Although hyperspectral imaging has mainly been used for the analysis of fingermarks, studies are also reported on several other forensic traces, including drugs, hair, dentin, bruises, blood stains, condoms, inks, tapes, firearm propellants, paints and fibres. These applications are described in more detail below.

2.3.a FINGERMARKS

Latent fingermarks are a complex mixture of eccrine deposits from the finger and sebaceous deposits resulting from touching other body parts, such as the face. Eccrine deposits mainly consist of amino acids, inorganic compounds, and proteins, while sebaceous material consists primarily of fatty acid esters. The chemistry of these residues varies among individuals and it shows increasing amounts of sebaceous deposits with age. Fingermark detection techniques aim to create contrast between the ridge details of a latent fingermark and the background on which it is located.

Detection and enhancement of untreated fingermarks

Several authors recently evaluated the possibility of detecting untreated latent fingermarks using hyperspectral imaging. Exline et al. used visible reflectance and photoluminescence hyperspectral imaging to detect untreated latent fingermarks on plastic and paper. Resulting images were compared to images created with a conventional forensic imaging system, in which different excitation and observation wavelengths could be chosen. While both methods succeeded in visualizing latent fingermarks on plastic, hyperspectral imaging
showed enhanced contrast on paper surfaces. Processing tools used included background division, offset correction, normalization and PCA. In a further study Payne et al optimized this visualization technique by using different processing tools to achieve an improved image\textsuperscript{30}.

Unlike visible hyperspectral imaging, NIR and IR hyperspectral imaging yield information about the vibrational modes of a molecule, and thus give additional information about the chemical composition of the material being studied. Bartick et al were the first to show the application of NIR and IR hyperspectral imaging for imaging latent fingermarks, using spectral bands indicative of the chemical components of the deposited material\textsuperscript{27}. They successfully visualized fingermarks deposited on aluminium coated microscope slides.

Crane et al demonstrated the ability of IR hyperspectral imaging to detect latent untreated fingermarks on various porous backgrounds (copier paper, cigarette butt paper, U.S. dollar bill, postcard) and non-porous backgrounds (trash bags, a soda can, tape)\textsuperscript{28}. Fingermarks on the soda can and a black trash bag were clearly visible when viewing the intensity band image at 9842 nm (asymmetric O-C-C stretch ester) (see Figure 2.7). On other backgrounds, more complicated processing tools were required, like PCA, and second derivatives. Processing with these tools rendered most prints clearly visible, even on paper-based porous surfaces. However, the position of the fingermarks was known before collecting the images.
Figure 2.7. Cut and flattened Dr. Pepper’s soda can with fingerprint deposit. (A) Soda can imaged by a document scanner. B) Infrared image of the outlined area obtained by plotting the band intensity at 9842 nm (1016 cm⁻¹). Reprinted from Journal of Forensic Sciences, 52/1, Nicole J. Crane, Edward G. Bartick, Rebecca Schwartz Perlman, Scott Huffman, Infrared Spectroscopic Imaging for Noninvasive Detection of Latent Fingerprints, 48-53, Copyright (2012), with permission from John Wiley and Sons.

In two papers, Tahtouh et al also described the application of infrared hyperspectral imaging to the visualization of untreated fingermarks. Results indicated that the infrared spectra of many untreated fingerprints show peaks due to C-H stretching vibrations around 3333 nm, mainly due to fatty acid residues. These peaks are common to most organic compounds, but they can
be used to visualize fingermarks against some backgrounds, like metals, minerals, and ceramics, that do not contain C-H bonds. For fingermarks on other backgrounds, they stated that some type of chemical enhancement technique is required prior to hyperspectral imaging.

Bhargava et al described an approach to use IR hyperspectral imaging to reveal latent fingermarks overlaid on top of one another, each made under different hand washing conditions. Differences observed in the absorbance of the C-H stretching mode and other vibrational modes in the spectra indicated that the two prints had different chemical compositions. Because of this variation, linear unmixing applied to the spectral content of the data could be used to provide images revealing both superimposed fingermarks.

**Detection and enhancement of treated fingermarks**

Conventionally, fingermarks are treated with chemicals to increase sensitivity and/or contrast with the background. On porous surfaces such as paper, ninhydrin and DFO (1,8-diaza-9-fluorenone) are often applied, which both react with amino acids present in fingermark ridges, causing a purple colour or photoluminescence respectively. On non-porous surfaces such as glass and plastic, cyanoacrylate (superglue) is the most widely used method. Cyanoacrylate fumes polymerize in the presence of moisture and greasy component of the fingermark. The contrast of fingermarks treated with cyanoacrylate can be further enhanced by various methods including luminescent staining.

Exline et al and Payne et al investigated the potential of hyperspectral imaging to increase the contrast and visual quality of treated fingermarks compared to traditional methods of detection. Using visible hyperspectral imaging they investigated fingermarks treated with ninhydrin, DFO, cyanoacrylate, and fluorescent dyes. In some cases, hyperspectral imaging showed significant enhancement over the traditional method, which was mainly due to the suppression of a highly fluorescent background or isolation of the developed latent impression. Hence, the minutiae details were visible using hyperspectral imaging where they were not with the traditional
method. Such examples were examined and compared to the original donor’s prints by a certified fingermark examiner. This process verified that the enhanced detail coincides with actual naturally occurring detail. The added information brought by hyperspectral imaging could sometimes be sufficient for exclusion purposes, whereas the traditional examination would lead to an inconclusive result. In a similar study Miskelly and Wagner used hyperspectral imaging to image chemically treated fingermarks deposited on a newsprint and an aluminium soda can. They showed that background correction is an important step in the visualization of fingermarks on difficult backgrounds.

Although visible hyperspectral imaging is an improvement compared to traditional techniques, it is not always possible to obtain acceptable fingermark images, for example when the background is highly fluorescent, coloured or patterned. However, most dyes that absorb or fluoresce strongly in the visible region are reflective in the NIR. This means that background interferences from dyed surfaces should be reduced when working in the NIR, compared to visible imaging. Maynard et al systematically imaged latent fingermarks on porous, non-porous and semi-porous surfaces. Next to a wide variety of conventional chemical and physical treatments, NIR laser dyes were tested for their ability to produce NIR photoluminescence. Both absorption and photoluminescence properties of the treated marks were examined using NIR hyperspectral imaging. The most suitable enhancement techniques depend on the type of surface. Fingermarks on coloured, printed or watermarked surfaces were imaged in the NIR region without interference from the background colour, both in absorbance and in photoluminescence modes. In these cases, imaging in the NIR region provided advantages over imaging in the visible region.

Background interference problems in the visible region can also be solved by showing differences in chemical composition of the print and the background using mid-infrared hyperspectral imaging, as shown by Tahtouh et al. Infrared hyperspectral imaging of chemically treated fingermarks was carried out on several challenging backgrounds. It was found that infrared hyperspectral imaging gives high quality images of cyanoacrylate-fumed fingermarks on polymer banknotes and aluminium drink cans, regardless of the
printed background. Attempts to acquire images of fingermarks on paper-based porous surfaces treated with established reagents such as ninhydrin were all unsuccessful due to the swamping effect of the cellulose constituents of the paper.

In addition to the chemicals commonly used to create contrast between fingermarks and backgrounds, Tahtouh et al tested four novel chemicals which can be visualized with IR hyperspectral imaging. These were chosen for their potential to produce a strong, isolated infrared spectral band. Each chemical polymerized selectively on fingermark ridges and high quality images were obtained of fingermarks on difficult backgrounds.

Detection and identification of trace contaminations in fingermarks

Next to eccrine and sebaceous deposits, fingermarks may be contaminated by exogenous substances of various sources. These substances might include drugs of abuse, traces of explosives or gunshot residue. Traces found at the scene of investigation can be directly related to an individual by its presence in the fingermark. Many of the approaches involved in collecting traces are destructive. For example, swabbing objects destroys fingerprint deposits within the area. Because of its non-destructiveness, hyperspectral imaging can be used to simultaneously image a latent fingerprint and detect trace information contaminating it.

In 2005, Grant et al had volunteers handle a mixture of common materials before giving fingermarks. When looking at the fingermarks under a visual light microscope, it was impossible to distinguish particles from different materials. However, by using infrared hyperspectral imaging, vibrational spectra of individual particles were obtained and identified by comparison with a spectral library.

In a similar study, Bhargava et al examined traces of explosives within a latent print, using an infrared hyperspectral imaging system (Figure 2.8). They used spectral subtraction to eliminate the effects of latent material on traces. Unique spectral features of the traces were used to provide images of the distribution of these traces. From pixels dominated by the material the full
spectrum of the traces was obtained and compared to databases for identification.

Figure 2.8. Images of a latent fingermark developed by using different vibrational modes to highlight different aspects of the chemical composition of the deposited material. a) Print image developed by the absorbance magnitude at 2,920 cm$^{-1}$. and b) print image developed by absorbance magnitude at 1,568 cm$^{-1}$. c) Example spectra from the oil-rich region (top, dark line) and flake rich region are shown. Reprinted from Analytical and Bioanalytical Chemistry, 394/8, Rohit Bhargava, Non-invasive detection of superimposed latent fingerprints and inter-ridge trace evidence by infrared spectroscopic imaging, 2069–2075, Copyright (2012), with permission from Springer.

Ng et al tested different spectral searching algorithms for their efficacy in finding targeted substances deposited within fingermarks$^{25}$. Out of a range of algorithms which included conventional Euclidean distance searching, the spectral angle mapper and correlation algorithms gave the best results when used with second-derivative image and reference spectra. Aspirin, diazepam,
caffeine, and explosive components were successfully detected and located in a fingermark.

Emmons et al used Raman hyperspectral imaging to examine fingermarks contaminated with traces of explosives (532 nm excitation)\(^40\). To determine if explosives were present in the fingermarks measured, the resulting spectra of each pixel were compared to a spectral library of pure reference specimens of explosives. A false colour picture was created which indicated the possible presence of explosives. Significant differences in the spectra could be used to differentiate between different types of explosives.

Chen et al applied IR hyperspectral imaging and principal component analysis to distinguish between overlapping fingermarks based on exogenous compounds\(^41\). After creating a blank fingermark containing natural secretions, a second finger contaminated with an explosive solution was printed on top of it. Although trace residues of the explosives trapped between the fingermark ridges could be clearly detected, it was not evident which fingermark these chemicals belonged to in cases with overlapped prints.

All the above studies examined fingermarks left on surfaces, which are ideal for infrared reflection analysis. In a forensic science scenario, reflective substrates such as a doorknob, knife blade, or handle should be relatively straightforward for analysis similar to the laboratory situation\(^25\). However, surfaces such as glass, plastic, wood, paper, cloth, etc., will all have their own (sometimes strong) infrared absorptions. These IR absorptions of the underlying surface will mask some parts of the spectrum, rendering these regions unusable for spectral identifications. The effectiveness of finding foreign materials within latent fingermarks in these cases will be dependent on having enough identifying spectral features outside these spectral regions\(^38\).

### 2.3.b Other Traces

Apart from the analysis of fingermarks, the benefits of hyperspectral imaging can be exploited for the analysis of many other traces of importance in forensic science. Latent traces can be detected and visualized by using spectral differences to obtain optimal contrast between a trace and its background.
Individual spectra give information about the chemical composition of the specimen, which is useful for identification, quantification, or age estimation. The possibility of viewing spectral and spatial information side by side is an advantage in comparative research of e.g. fibres, paint chips, or inks, where the question arises whether two traces share common spectral features. Several applications described in literature are reviewed below.

Kalasinsky et al were the first to demonstrate the value of infrared hyperspectral imaging for determining drugs of abuse in hairs. By examining only the interior portion of the hair, drugs exclusively resulting from human ingestion were measured and distinguished from drugs that made contact with the outside of the hair. After microtoming the hair, IR hyperspectral images were obtained of the cortex and the medulla. Drug free hairs of different sources all correlated with standard spectra of proteins. A hair of a chronic drug abuser of hydromorphone was analyzed similarly. Subtraction of the drug free reference spectra yielded a strong indicative band at 5824 nm, which was also present in reference hydromorphone spectra. An intensity band image at 5824 nm showed that the drug was concentrated in the centre of the hair. This way, relative drug concentrations across the hair could be successfully determined and visualized. In a further study, Kalasinsky showed the distribution of drugs in human hairs, which is critical information to validate drug testing data. IR hyperspectral imaging on hairs doped with 6-acetyl morphine (a metabolite of heroin) and cocaine, showed that hydrophobic drugs tend to bind to the medulla of the hair while hydrophilic drugs tend to be spread throughout the cortex of the hair.

Hair colour is basically determined only by eumelanin and phaeomelanin, whose varying ratios produce the observed colour. In forensic science casework, hair colour is normally classified through visual comparison with standardized plates. Birngruber et al investigated the possibility to objectively distinguish hairs from different persons using hyperspectral imaging in the Vis-NIR range. They demonstrated an extreme intra-individual variability in the spectra of single hairs from an individual. Because of this, hairs undistinguishable on the basis of morphology could not be distinguished based on the hyperspectral images.
The chemical composition of the dentinal part of the tooth evolves with increasing age. Tramini et al measured 30 human teeth with Raman hyperspectral imaging (with 632 nm laser excitation) and were able to identify a very small quantity of dental material coming from skeleton debris or biological remains, and determine which kind of tissue it was. They created a PLS regression model to predict the age of an individual based on Raman spectra of his teeth. The model was tested on four teeth, and an age estimation was obtained with a mean error of 5 years.

The analysis of bruises, or aging of bruises in particular, can give important evidence in cases of domestic violence or child abuse. Several studies have been performed as initial steps towards the aging of bruises using hyperspectral imaging. A bruise is formed after blunt trauma, which results in blood being present in the skin. In time, haemoglobin in the blood is degraded into other products, including bilirubin. Both haemoglobin and bilirubin have typical spectral features in the visible region. Payne et al showed the possibility to use hyperspectral imaging to differentiate pure blood from blood with bilirubin based on these spectral features. Randeberg et al presented hyperspectral images of bruises on porcine and human skin. They used minimum noise fraction transform, a statistical method similar to PCA, to classify the injuries. Stam et al described how hyperspectral imaging can be used to accurately determine the areas covered by haemoglobin and bilirubin in the bruise, by fitting pixel spectra with a combination of reference spectra of chromophores present in bruises.

Similarly, reflectance spectra of blood stains can be spectrally unmixed to derive the relative amounts of oxyhaemoglobin, methaemoglobin and hemichrome within the blood stains. By comparison of spectra derived from hyperspectral imaging data with a non-linear least squares fit of the theoretical spectra of the haemoglobin derivatives, blood stains were identified in a simulated crime scene and could be distinguished from similarly coloured substances, as demonstrated in Chapter 3. Additionally, the temporal behaviour of the amount of haemoglobin derivatives provided insight in the chemical changes occurring in time, and could be used to estimate the age of
blood stains in Chapter 4\textsuperscript{40}. Figure 2.9 shows a simulated crime scene in which fresh and older blood stains were distinguished using this method.

![Figure 2.9. Simulated crime scene, in which fresh and older blood stains were automatically detected and distinguished (left) based on their reflectance spectra (right).](image)

Important evidence pertaining to sexual assault cases can be provided by the identification of condom lubricant components. In an exploratory study, Wolfe and Exline showed that some of the most common materials found in condom lubricants can be accurately characterized by Raman hyperspectral imaging (with 532 nm laser excitation) without the extensive specimen preparation inherent to other analytical methods\textsuperscript{50}. Using the CH stretching region of the Raman spectrum, they were able to generate contrast based on spectral differences.

To demonstrate the potential of hyperspectral imaging in forensic investigations, Payne et al compared hyperspectral imaging to point measurements performed with traditional spectrometers\textsuperscript{51}. They used Vis/NIR hyperspectral imaging to differentiate between a set of tapes and adhesives, a set of inks (Figure 2.10) and two brands of firearm propellants, based on reflectance and photoluminescence properties. They conclude that hyperspectral imaging offers significant advantages, mainly because a large number of specimens can be analyzed at once. This makes comparisons of different specimens easier and reduces the analysis time.
In the same paper photoluminescence spectra of two multi-layered paint specimens were compared using hyperspectral imaging. Because both spectral and spatial data were gathered, differences in paint layers could easily be highlighted visually, as an alternative to a spectral comparison. This was also shown by Flynn et al who analyzed more multi-layered paint specimens using IR hyperspectral imaging. They presented several ways to display hyperspectral data, which make chemical differences and similarities between heterogeneous specimens easy to visualize and understand for the layperson (such as a juror). The same applies to the visualization of differences in bicomponent fibres, as described in. This study showed that infrared hyperspectral imaging can provide spatially resolved chemical information for those bicomponent fibres where it is possible to detect spectral differences between the two components present. As well as yielding characteristic IR spectra of each component, the technique also provided images clearly illustrating the side-by-side configuration of these components in the fibre. However, in 5 of 11 bicomponent fibres no spectral differences were found.
using integrated peak intensities. Multivariate statistical analysis may improve these results. Markstrom and Mabbott also demonstrated the advantages of using hyperspectral imaging for the comparison of fibers\(^5\) and addressed that the ability to compare fibres simultaneously in one measurement minimizes the chance for errors.

Miskelly and Wagner attempted to improve the visualization of chemically treated soil shoe marks using hyperspectral imaging\(^3\). The standard chemical enhancement technique for such marks is an acidic thiocyanate solution which reacts with iron (III) oxides in the soil to form a coloured iron (III) thiocyanate complex. Unfortunately, this complex has a broad absorption band in the visible spectrum. They point at the necessity of enhancement chemicals with narrow absorption bands, as these can often be readily enhanced relative to the underlying background.

A forensic science application of remote sensing technology was demonstrated by Kalacska et al\(^5\), who used airborne hyperspectral imaging for the detection of mass graves. Analysis of the spectra using Minimum Noise Fraction transform showed a clear separation between an experimental grave, a refilled empty grave, grass and forest. This indicates that airborne hyperspectral imaging can be used to detect the existence of suspicious decomposition properties, i.e. mass graves, which lead to differences in soil chemistry and vegetation.

2.4. TYPICAL CHALLENGES

As demonstrated above, the application of hyperspectral imaging in forensic science casework brings specific challenges. In contrast to pure specimens usually analyzed in the laboratory, traces from casework can consist of complex contaminated mixtures. Next to this, the chemical composition of biological traces may change in time. Although these changes can be used for age estimation purposes\(^1, 45\), the influence of environmental conditions like temperature, humidity, precipitations and light should be studied. For example the influence of temperature and humidity on the aging of blood stains is shown by Bremmer et al\(^56\).
Also, traces are typically not found on ideal neutral reflecting backgrounds used in laboratories (see Figure 2.9). In casework, all possible backgrounds can be encountered (e.g. different materials, porous, non-porous, coloured, patterned, etc.) which may complicate the measurements. Comparison of spectra on different backgrounds typically requires advanced system calibration and data analysis. Miskelly and Wagner and Tahtouh et al examined the spectral properties of enhancement chemicals and experimented with new chemicals for the visualization of shoe marks and fingermarks respectively dedicated for the use of hyperspectral imaging. Although the use of chemicals is not preferred, this may help finding evidence.

The above mentioned challenges are not just characteristic for hyperspectral imaging, but are also valid for conventional spectroscopy. The transition from spectroscopy to hyperspectral imaging, however, is not straightforward. While specimen optical properties are independent of the spectral measurement system, the transition from spectroscopy to spectral imaging involves a drastic change in the illumination–collection geometry of the measurement system. Gebhardt et al showed that this change in measurement setup results in a disparity between measured spectra, which is dependent on the optical properties of the specimen and the optical path length. Spectral data bases created for the identification of forensic traces like fibres or printer toners may therefore need to be adapted to hyperspectral imaging applications.

Moving hyperspectral imaging from the laboratory to the scene of investigation brings further complications. Advances in wireless technology and sealed operating units are desirable to prevent contamination. Investigating scenes where chemical, biological, radiological, or nuclear (CBRN) events have occurred poses dangers to investigators. In these cases, a remotely controlled robotic hyperspectral imaging system could provide important information instantly. However, these scenes have even more decontamination requirements.

The complex nature of crime scenes makes for challenging image analysis. Next to this, sunlight, external light sources, shadows and reflections from nearby objects all change the apparent illumination on an object. This
variation can cause large variability in the measured spectra for a fixed object, a problem regularly encountered in remote sensing. Algorithms are needed to distinguish this spectral variability due to nonuniform illumination from spectral variability between objects.

2.5. FUTURE APPLICATIONS

Despite the challenges, hyperspectral imaging offers great potential for providing new, valuable information in forensic science casework. Hyperspectral imaging can be applied to detect traces by optimizing the contrast between a trace and its background, or to differentiate between traces, based on spectral differences. To create contrast between a trace and its background, crime scene investigators traditionally use commercially available light sources with different excitation and barrier filters to isolate regions of the spectrum where traces of interest have high absorption or photoluminescence. This way, e.g., semen and blood traces can be detected (Chapter 6). For laboratory purposes, dedicated forensic imaging systems are available with different light sources, excitation and barrier filters. These systems are mainly used in document analysis, e.g. for differentiating inks, but can also be applied in other fields, e.g. the visualization of gunshot residue patterns on dark clothing. For all these purposes hyperspectral imaging can be used instead. Because hyperspectral imaging systems filter the light in many small bandwidths, maximum spectral differences can be calculated automatically and the choice of one specific filter is no longer necessary, which reduces the risk of human errors.

Many applications currently performed with conventional spectroscopy may also be enhanced with a spatial component using hyperspectral imaging, similar to the way presented for the identification (Chapter 3) and age estimation of blood stains (Chapter 4), as shown in Figure 2.9. In these applications, the added spatial information is crucial, as the blood stain pattern may reveal useful information for the reconstruction of a crime. The spectroscopic identification of other body fluids will also benefit from hyperspectral imaging, as the traces can then be
interpreted in their original context. The spatial distribution of components may be less important in fields like illicit drug analysis. However, using hyperspectral imaging instead of spectroscopic point measurements for the identification of drug components may speed up the process, as many specimens can be imaged at once. The improved speed of hyperspectral imaging compared to spectroscopy, is particularly of advantage in hazardous environments. In explosives investigations especially, the ability to measure specimens without contact or specimen preparation is beneficial, as many accidents occur even when trained personnel handle explosives.

In 2000, Malkoff and Oliver proposed some interesting applications in forensic medicine which are not yet put into practice, like the analysis of patterned injuries (e.g. a tire print on the body of a victim), scanning of body and clothes for toxins, the aging of wounds through the spectral evaluation of local inflammation and repair, and the estimation of the time of death of a victim. They also draw the attention to the need for crime scene reconstruction for operational planning, hazard identification, training etc. Crime scenes are often digitally captured using photography, and panoramic and 3D scanning techniques. These data can be used for a virtual crime scene reconstruction. The addition of hyperspectral imaging data to this reconstruction can give information about the chemical composition of traces and their distribution in the scene. Malkoff and Oliver claimed, this may help in body localization and retrieval, or the characterization of biological or chemical threats. Using hyperspectral imaging, the chemical properties of the scene of investigation can be captured quickly without much disturbance to the scene and analysis can be performed afterwards in the original context.

2.6. CONCLUSION

Recent technological developments in hyperspectral imaging components have opened up the approach to forensic science applications. Fast acquisition, portable, high resolution systems are emerging facilitating the transfer of hyperspectral imaging from the laboratory to the field. Several forensic science applications of hyperspectral imaging were recently explored successfully.
Challenges typically encountered in forensic science casework, e.g. contaminated traces found on non-ideal backgrounds in varying environmental circumstances, emphasize the necessity to modify existing techniques and instrumentation. Key steps in the research process are refining and validating the data to meet the needs of the legal and scientific communities. When introduced in forensic science casework, hyperspectral imaging can help investigators detecting, visualizing and identifying useful traces non-destructively.