Spectral confocal reflection microscopy using a white light source

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We present a reflection confocal microscope incorporating a white light supercontinuum source and spectral detection. The microscope provides images resolved spatially in three-dimensions, in addition to spectral resolution covering the wavelength range 450-650 nm. Images and reflection spectra of artificial and natural specimens are presented, showing features that are not normally revealed in conventional microscopes or confocal microscopes using discrete-line lasers. The specimens include thin film structures on semiconductor chips, iridescent structures in Papilio blumei butterfly scales, nacre from abalone shells and opal gemstones. Quantitative size and refractive index measurements of transparent beads are derived from spectral interference bands. [DOI: 10.2971/jeos.2008.08026]

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1 INTRODUCTION

The confocal microscope enables the imaging of volume specimens with three-dimensional resolution [1]. For biological imaging, it is most commonly used in fluorescence mode, in conjunction with appropriate fluorescent markers, to reveal structural and functional information about a specimen. The choice of illumination wavelengths had originally been restricted to the available laser lines, but the recent introduction of white light supercontinuum (WLS) sources has provided much greater freedom [2]-[4]. Whereas previously a confocal microscope could in practice use only a small number of laser wavelengths, a WLS source can now provide excitation at any wavelength in a range from around 400 nm to 2000 nm. The combination of the WLS source with spectral detection has provided further versatility for confocal fluorescence microscopy [5, 6]. The versatility of the WLS source in fluorescence microscopy has thus been widely demonstrated and well documented. However, it is also well suited for application to reflection mode confocal microscopy, such as the method already used to characterise the scattering from gold beads [7]. In this paper, we demonstrate the use of a WLS source in a reflection mode confocal microscope incorporating spectral detection. This microscope combines the three-dimensional spatial resolution of the confocal microscope with the spectral resolution of a spectrometer, providing image information that cannot be obtained using either conventional white light microscopy or confocal microscopy using discrete line laser sources. We present images and analysis spectra from a range of specimens that illustrate the capabilities of this method.

2 MICROSCOPE IMPLEMENTATION

The microscope configuration is illustrated in Figure 1(a). Broadband illumination (450-2000 nm) was provided by a Fianium SC450-2 WLS source. The output light was passed through a filter, removing all wavelengths longer than 680 nm and the overall intensity was reduced by a factor of 1000 using a neutral density filter. The resulting beam was expanded then reflected by a 50:50 beam splitter into the back aperture of the objective lens (Olympus MPlanApo, 50×, 0.95 NA, dry). The specimen was mounted on a three-dimensional piezo translation stage (Physik Instrumente NanoCube). Light scattered by the specimen was collected using the same objective lens and was focused by a 200 mm focal length lens onto the 20 μm diameter confocal pinhole. The light passing through the pinhole was collimated by a 160 mm focal length lens before falling upon a blazed diffraction grating (600 line pairs/mm, blazed for 500 nm) placed a distance of 210 mm away from the lens. The spectrally separated beam was passed via a scanning mirror and a 50 mm focal length lens onto the sensor chip of a CCD camera (Andor iXon EMCCD, 512×512 pixels, used in conventional detection mode). The distance between the grating and the mirror was 130 mm and that between the mirror and lens was 100 mm. The resulting line focus on the CCD consisted of the reflection spectrum from a particular specimen point. As the specimen was scanned in the x direction using a piezo translation stage, the galvanometric scan mirror would synchronously sweep the spectral line across the camera. This was achieved by driving both the stage and the mirror with the same scanning signal. By synchronising the camera exposure so that the shutter was open during a single scan, it was possible to retrieve a complete x-λ “image”
in a single camera frame (Figure 1(c)). The microscope also included a second CCD camera, a removable mirror and conventional illumination to facilitate specimen location.

3 SPECTRAL CALIBRATION AND PSEUDO REAL-COLOUR IMAGES

In addition to spectral variations in the source and the wavelength dependent sensitivity of the CCD, various parts of the microscope could introduce chromatic effects, for example through absorption, reflection or diffraction. Figure 1(b) shows the spectra measured at three locations in the system. Spectrum A is the filtered output of the WLS at the input to the microscope, where the spectral variation is predominantly due to the source. Spectrum B was measured after the confocal pinhole, whilst using an aluminium mirror as the microscope specimen. This spectrum includes the effects of the losses in the objective lens and the dependence on wavelength of the power transmitted through the pinhole. Both A and B were measured using a calibrated Ocean Optics USB2000 spectrometer.

Spectrum C was measured using the CCD camera. The correspondence between wavelength and pixels was determined using the band edges produced by a set of bandpass filters at 483, 518, 568, 628, and 638 nm. It was found that the spectrum was uniformly distributed across the pixels, such that the pixel pitch corresponded to a wavelength step of 0.50 nm. This relationship was linear, it was appropriate to consider the signal measured by the CCD pixels as a wavelength spectral density. In the results presented in this paper, the intensity spectra can therefore be regarded as equivalent to a spectral density (in arbitrary units).

For quantitative spectral reflectivity measurements, it was necessary to compensate for the non-uniform illumination spectrum, spectrally dependent losses in the optical system and detector sensitivity. For this reason, all measured pixel values, \( p(\lambda) \), were normalised with reference to spectrum C, to give the renormalised reflection spectrum \( r(\lambda) = p(\lambda)/c(\lambda) \). The calibration function \( c(\lambda) \) was a smoothed version of spectrum C obtained by taking a moving average of ten adjacent pixel values.

One method for visualising the spectral image data is to convert the data into pseudo real-colour red-green-blue (RGB) images. These images provide an approximation to the visual appearance of the genuine spectrum. Such images have been obtained before by using three discrete laser lines [8], but this method lacks the spectral coverage of the WLS source. We employed a simple method that is common to many colour imaging systems by summing particular ranges of the spectrum to generate each of the RGB values: the wavelength ranges from 450-500 nm, 500-580 nm and 580-650 nm were allocated to blue, green and red, respectively. In order to ensure an appropriate white balance, \( x=\lambda \) data were taken of the reflection from the surface of a microscope slide. As there should be minimal spectral variation caused by this reflection, the in-focus image should represent white and the relative proportions of the RGB components were adjusted to maintain this appearance. Using this calibration, the RGB images of other specimens were scaled accordingly. The \( x=\lambda \) scan of the microscope slide also showed that the axial chromatic aberration was much smaller than the focal depth for this wavelength range.
4 RESULTS

We investigated the capabilities of the white light confocal reflection microscope by imaging a semiconductor chip. Figure 2(a) shows a conventional microscope image of a region of the chip (obtained using a separate microscope), where the colours arise as a result of complex multilayered semiconductor structure and the varying thickness of the top passivation layer. The pseudo real-colour images in Figure 2(b) are from $x$-$z$ confocal sections through the surface of the chip at the various locations A to E, demonstrating a range of colours. For all of these images, the illumination was incident from the top. Figure 2(c) shows the normalised spectra obtained at these points, showing significant spectral variation in the reflectance at the different points. The oscillating nature of the reflectance spectrum is typical of thin-film interference. The density plot of Figure 2(d) shows both the spatial and spectral variation in the region around point D. The apparent shifting of the spectral bands could arise from small variations in the thickness of the transparent passivation layer.

The spectral confocal reflection microscope is particularly useful for imaging naturally occurring iridescent optical structures. Butterfly wings often have brightly coloured scales, whose optical properties arise from structures similar to photonic crystals [9, 10]. Figure 3(a) shows $x$-$z$ sections of three different scales from a *Papilio blumei* butterfly (note that a gamma correction of 0.5 has been applied in order to enhance dim features). The confocal images reveal three-dimensional variations in reflectance of these scales. Note also the secondary foci produced by the curved surfaces of the concavities. Normalised reflection spectra from the points A to D are also presented. The different reflection spectra of the scales are clearly visible. These images are comparable with the optical and electron microscope images of Vukusic et al., who illustrate photonic structures in other *Papilio* species [10]. In particular, we note that the blue-biased spectrum D corresponds to finely structured ridges between the concavities. For all three scales, it can be seen that longer wavelengths appear to be reflected from a layer deeper than that which reflects the shorter wavelengths. This may be a consequence of a difference in optical properties of lower layers, as noted by Vukusic et al.

Iridescent structures are also produced in the shells of abalone [11]. Figure 4(a) shows pseudo real-colour $x$-$z$ and $x$-$y$ images of a fragment of abalone shell obtained using the spectral confocal microscope. The $x$-$y$ image was constructed as the maximum value pixel projection (sometimes referred to as an autofocus image) from four $x$-$y$ sections taken at 1 $\mu$m axial steps. The striking colours are caused by the optical effects of stacks of aragonite platelets [11]. It is likely that the colour variations across the specimen are due to changes in the orientation of the stacks in different regions. For comparison, we also separately obtained conventional micrographs of an adjacent region of the specimen. The images were taken with a colour digital camera, using a 0.7 NA objective lens and halogen lamp illumination with both open and closed illumination aperture stop (Figure 4(a)). We note that the strong colour variation is not apparent in these images, as the cumulative effect of out-of-focus scatter leads to washed-out, predominantly white images.
Figure 4(b) shows a single x-y image section of an opal gemstone. An opal consists of an array of amorphous silica spheres of uniform diameter, which typically lies in the range 150-400 nm, depending upon the type of opal [12]. The spheres are locally arranged in a face centred cubic packing (occasionally hexagonal close packed), although there are domains over which the packing orientation changes. It is this structural variation that leads to the vivid iridescent colours of these gemstones. The resolution of the confocal microscope is insufficient to resolve the three-dimensional arrangement of the spheres. However, chromatic effects created by the structure, in particular bands of different colours, can be seen in Figure 4(b). Similar bands were observed by Akizuki [13], who suggested they were related to the change in orientation of the sphere packing in different domains. Again, it is important to note that this chromatic structure is not clearly revealed in high NA conventional microscope images of this specimen, which show a strong reflection from the surface (Figure 4(b)).

Many of the spectral features visible in these results are related to interference between reflections from different parts of the specimen. This effect is clearly seen in the reflection spectra from transparent spherical beads, when the geometrical focus is coincident with the centre of the sphere (Figure 5). If we consider the geometrical optics approximation of a sphere that is much larger than the wavelength, all of the focused rays passing through the centre are normally incident on both the front and back surfaces of the sphere. The resulting normal reflections at each surface are similar in magnitude and follow a reciprocal path back through the objective lens. This results in a bright point in the confocal image, apparently from the centre of the bead. By considering the interference between the external and internal surface reflections, one easily obtains the following expression relating the spectrum to the properties...
We have shown that detailed spatial and spectral information can be obtained with a spectrally resolved reflection mode confocal microscope. This has enabled the imaging of specimen features that are not apparent in laser illuminated confocal microscopy or conventional microscopy. This microscope design could be further modified to use line, rather than spot, illumination and a slit instead of the pinhole in the detection path [15]. In this way, an entire x-\(\lambda\) frame could be obtained in a single camera frame without the need for scanning in the \(x\) direction, thus significantly improving the overall acquisition times of \(x-y-z-\lambda\) data sets at the expense of slightly reduced axial resolution [16].

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