Spectrally encoded fiber-based structured lighting probe for intraoperative 3D imaging

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Abstract: Three dimensional quantification of organ shape and structure during minimally invasive surgery (MIS) could enhance precision by allowing the registration of multi-modal or pre-operative image data (US/MRI/CT) with the live optical image. Structured illumination is one technique to obtain 3D information through the projection of a known pattern onto the tissue, although currently these systems tend to be used only for macroscopic imaging or open procedures rather than in endoscopy. To account for occlusions, where a projected feature may be hidden from view and/or confused with a neighboring point, a flexible multispectral structured illumination probe has been developed that labels each projected point with a specific wavelength using a supercontinuum laser. When imaged by a standard endoscope camera they can then be segmented using their RGB values, and their 3D coordinates calculated after camera calibration. The probe itself is sufficiently small (1.7 mm diameter) to allow it to be used in the biopsy channel of commonly used medical endoscopes. Surgical robots could therefore also employ this technology to solve navigation and visualization problems in MIS, and help to develop advanced surgical procedures such as natural orifice translumenal endoscopic surgery.

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1. Introduction

Computer vision techniques have recently been applied in minimally invasive surgery (MIS) to provide more information to the surgeon from the white light endoscopic view and to compensate for the natural cues that are sacrificed when moving from the open surgery platform, for instance the loss of haptic feedback or stereo vision. These techniques may be used to track the surgical instruments, predict the motion of the tissue or detect the tissue surface profile [1,2], but they also have the potential to improve surgical accuracy and reliability by providing augmented views of the tissue and enabling diagnostic assistance.

The detection of tissue surface information in three dimensions has many possible applications in MIS including the registration of multimodal images or pre-operative image data, the improved navigation of robotic instruments and the measurement or characterization of tissue morphology for diagnostic purposes [3,4]. It is commonplace for preoperative MRI, CT or ultrasound scans of a patient to be used for surgical planning to locate and navigate to paronychial tumors. However during the surgery itself surgeons often rely on the optical identification of anatomical landmarks and navigate based on the relative positions between the static pre-operative images and the intraoperative view to make a judgment on how to dissect the tissue. A longstanding aim of the field of computer aided surgery is to help guide the surgery using an augmented reality view of the tissue revealing hidden structures that are present in the preoperative images but not in the live endoscopic view [5]. This could also allow the current finite element tissue models to be fitted to the pre operative data to allow the augmented view to deform and shift as the tissue is manipulated under known boundary constraints [6]. An example of the use of tissue surface morphology is the evaluation of the shape of colonic polyps. These have been shown to be related to the status of the lesion itself and have been proposed as a method of guiding or even replacing biopsies [3,4]. The accuracy of this type of technique is still down to the experience and judgment of the observer.
Quantification of polyp surface shape could be used as an objective measure of these polyps and compared against a library of known cases to guide biopsy selection.

Due to the reasons mentioned above, it is a widely pursued goal of the surgical imaging community to obtain an intraoperative measurement of tissue surface geometry. This has been previously attempted to allow multimodal registration in cadaver measurements [7], microscopic imaging [5] and radiotherapy guidance [8], although MIS applications remain relatively unexplored. Arguably the most successful method of 3D surface measurement during MIS has been stereo reconstruction, where a rigid endoscope with two imaging channels simultaneously acquires corresponding ‘left’ and ‘right’ views of the scene and uses a mathematical model based on a pinhole camera assumption to triangulate the position of salient features (areas of high contrast, such as the intersection of two blood vessels) found in both channels [9–11]. However, this technique is not suited to homogenous or featureless tissues. In this case, structured lighting must be used to project artificial features onto the tissue surface. A number of techniques for extracting depth information using patterned light exist and involve projection of lines, grids or dots [12]. However, the brightness required and the size of the optics needed to provide the pattern has, thus far, mostly limited their use to biomedical applications outside of the endoscopic regime due to the difficulties in miniaturizing projective imaging systems [13,14]. Previous attempts at endoscopic implementation have only been suitable for rigid endoscopy and were not compatible with pattern encoding techniques which are more immune to tissue occlusions [15].

To achieve dense surface reconstruction, a number of projected features is required, which introduces computational problems in the identification of specific features in the presence of occlusions where parts of the tissue may obscure part of the pattern. This results in discontinuities and errors in the 3D reconstruction. To overcome this, the pattern must be coded so that each projected feature can be identified, thereby minimizing the risk that a feature is confused with a ‘missing’ one and causing an error in the reconstruction. In other applications, this has been done spatially (using binary words in the pattern for example) or by introducing a number of different colors into the pattern using digital projection technology [16–18]. However, these techniques were applied to non-endoscopic and in most cases non-biological situations.

In this paper, an endoscopic fiber-optic structured lighting probe for intraoperative 3D imaging is proposed to overcome the limitations of the technologies described above. It uses a broadband laser to encode a dense pattern of spots with a unique color for each, and a segmentation algorithm that recovers the spectral signature of each dot from standard RGB color CCD images. The probe is highly flexible, has a small diameter and produces a pattern of high brightness making it suitable for use with biological tissue and low light collection efficiency endoscope systems. The optical set-up and spot identification algorithm are described, along with initial results characterizing the performance and limitations of the system and 3D reconstructions of ex vivo tissue.

2. Materials and methods

2.1. Probe design

The optical set-up in Fig. 1 shows how the spectrally encoded pattern is generated. The collimated output of a 4 W supercontinuum laser (SC400-4, Fianium Ltd., Southampton, UK) is dispersed using a prism (SF-11 glass) and focused into a thin line to couple it into the 127 fibers of the array end of a custom-made line-to-spot converter (FiberTech Optica, Inc., Canada). The distances between the prism and the coupling lens, and the coupling lens and the fiber array are set to the lens focal length so that the spectral components are sharply focused at the linear array. At the distal end of the probe, a short focal length aspheric lens (f = 4.51 mm) is used to form an image of the end face of the probe. Since the fibers are incoherently bundled, the result is that a random pattern of colored dots is projected onto the screen or
sample. The brass ferrule at the array end of the probe is 12.5 mm in diameter, but the rest of the probe has a maximum outer diameter of 1.7 mm making it compatible with biopsy ports of commonly used flexible endoscopes.

Fig. 1. (a) Broadband laser light from the supercontinuum is dispersed by an SF-11 prism, which is then coupled into the fibers (50 μm core) at the array end of the probe. The projected pattern is a magnified image of the end face of the bundle formed by the projection lens. (b) Emission spectrum of the supercontinuum laser source, with the wavelength range used by the probe indicated between the dashed lines.

Since the light source is a broadband laser and the fibers are closely spaced at the input end (average separation of 68 ± 3 μm), each projected spot effectively contains a unique low bandwidth spectral feature. It should be noted that although the laser is nominally 4 W, it emits over a 420-2200 nm range meaning that most of that power is contained in the near infrared, which is filtered out prior to reaching the prism. Only visible wavelengths are coupled into the fiber bundle so that the final output power at the probe tip after coupling losses is approximately 110 mW, well below the tissue damage threshold.

2.2. Wavelength segmentation and centroiding

The identification and segmentation is based on recovery of the peak wavelength of each spot using knowledge of the relationship between the RGB colorspace and the wavelength of the light. A schematic is shown in Fig. 2 to illustrate the transformation between RGB and wavelength for each pixel and is further described in the text below.

The CIE 1931 xy colorspace is used to represent the colors of visible light in an area defined by wavelengths visible to the human eye [19]. The xy chromaticity coordinates of pure wavelengths form a curve called the spectrum locus at the boundary of the colorspace. The RGB values as recorded by a standard color digital camera however, are only capable of detecting a triangular subset of these colors defined by the RGB response of the camera indicated in Fig. 2 (b). When an RGB image of the spot pattern is recorded by a camera, the position of each pixel within this triangular space can be calculated with knowledge of the RGB system used. This is achieved for each pixel by converting the RGB triplet into the
Fig. 2. Spot segmentation and 3D calibration. (a) Cartoon image showing three projected spots, having different RGB values. (b) Each RGB triplet is converted to \(xy\) coordinates. A line projected through these coordinates from a reference white spot intersects the spectrum locus (dashed) at the dominant wavelength of the pixel. (c) RGB pixels are replaced by the calculated wavelength to form a greyscale ‘\(\lambda\)‘-space’ image which can be thresholded to find the centroids of spots of a particular wavelength. (d) Epipolar geometry showing different positions of a calibration object (c1-c3) and triangulation of points using spot centroids.

The tristimulus values \(XYZ\), and then into the chromaticity coordinates \(xy\) by multiplication by the \(3 \times 3\) transformation matrix \(M\). The elements of M are calculated based on the color coordinates of the vertices of the RGB triangle and the reference white [20].

The dominant wavelength detected at the pixel of interest is found by projecting a line from a reference white point, through the pixel’s \(xy\) coordinates, to the point of intersection with the spectrum locus. In practice this is achieved computationally by creating a look-up table of color coordinates representing the mapping of the spectrum locus onto the triangular RGB gamut. For each pixel in an image, the intersection point of a line defined by the reference white and pixel \(xy\), and the nearest side of the RGB gamut is calculated. The closest point to this intersection is then located in the look-up table and its corresponding wavelength returned (Fig. 2 (b)). The result is a greyscale ‘\(\lambda\)‘-space’ image where each pixel value is the calculated wavelength of the light detected.

Once the ‘\(\lambda\)‘-image’ is acquired, spots of a particular wavelength can be isolated by thresholding by wavelength and finding their centroids. The finite bandwidth of each projected spot, sensor noise and uncertainty in locating an RGB triplet’s dominant wavelength means that each spot consists of a narrow distribution of wavelength values in \(\lambda\)-space. Therefore a simple thresholding of the \(\lambda\)-image at a wavelength of interest results in a cluster of pixels scattered within the region of the spot rather than a continuous area. A region-growing algorithm searches for these clusters by examining the nearest neighbors of each pixel in the thresholded image, and those with values within ± 1 nm of the wavelength of interest are deemed suitable for inclusion in the region. In this way, clusters of pixels at the location of a projected spot are ‘grown’ into each other to form a single identifiable region. As a final step, a median filter is applied to remove any erroneously detected pixels outside of the spot. The centroid of the detected region is then recorded along with the histogram of wavelength values within it. The peak value of this distribution is recorded as the ‘label’ for that particular spot and the process is repeated for multiple spots within the image.

2.3. 3D reconstruction

Triangulation of the 3D location of the imaged spots is accomplished through a calibration based on the pinhole camera model and methods previously developed for stereo reconstruction [9]. A calibration object (a plane with a checkerboard pattern of known dimensions) was imaged at 12 different positions in the camera’s field of view. White light and patterned illumination images were recorded for each position of the object to acquire calibration data for calculating the intrinsic geometrical parameters of the camera and also for estimating the 3D path of light rays projected from the probe.

Using the intrinsic results, it was possible to estimate the 3D position of the calibration object in metric space, so with the knowledge that the corresponding images of patterned...
illumination must show the spots lying on these planes, their 3D locations could be calculated. For each calibration image of the projected spots, the centroid of each spot was located using the algorithm described in Section 2.2. Backprojecting each spot’s rays to find the intersection with the calibration plane provided a set of 3D positions that the structured light ray passed through. Lines representing light rays were fitted to each spot’s 3D data, composed of 12 calibration positions, giving the origin of the projection and its position relative to the camera (Fig. 2 (d)). With the projection path of the rays mapped out, the 3D position of an individual spot reflected from an object of unknown shape could be determined by finding its position on the camera’s image plane and projecting a ray through the ‘pinhole’ to intersect with its known projection path. Due to noise in the image the intersection is not guaranteed and the solution is taken as the midpoint of the shortest line connecting the two rays [9].

2.4 Characterization and ex vivo testing

To test the effectiveness of the algorithm in recovering the wavelength of each spot, the pattern was projected onto a white screen and RGB images were recorded using a color CCD camera (DCU 223C, Thorlabs Ltd., UK). The white screen was then removed and spectra of individual spots were recorded using an optical fiber probe placed in the pattern’s plane and connected to a spectrometer (HR 4000, Ocean Optics, Inc., USA).

Since the optical properties of tissue within the body may vary significantly, the effect of background reflectivity on the detected spot wavelength was examined by imaging the pattern on different colored backgrounds. The performance of the system was then evaluated in biological tissue of varying optical properties using ex vivo kidney (ovine) and intestinal (porcine) tissue. Finally, the 3D reconstruction capability of the system was demonstrated by calibrating the projection probe and camera, and determining the shape of two validation objects (plane and cylinder) as well as ex vivo tissue of varying type, color and shape (kidney, fat and liver).

3. Results

3.1 Characterization: algorithm testing

Figure 3 (a) shows the raw RGB image acquired by the camera when the pattern was incident on a plane white screen, while Fig. 3 (b) shows the same pattern after conversion to $\lambda$-space.

Fig. 3. (a) RGB image of pattern recorded by camera. (b) $\lambda$-space image with centroids of spots. (c) Plot showing spot wavelength as calculated by the segmentation algorithm against the wavelength measured using a spectrometer. The transmission response of the camera’s filters (normalized to 1) is overlaid and the identity line is shown in black. The error bars indicate ± 1 standard deviation.
The wavelengths of the spots in the $\lambda$-space image after processing are plotted against the wavelengths measured with the spectrometer in Fig. 3 (c). In order to distinguish closely spaced wavelengths reliably, the algorithm requires a signal in at least two out of the three channels (red, green and blue), which define the working space of the camera’s RGB system. The filter response of the camera used, as measured in the lab using a reflectance standard and tunable filter, is also plotted in Fig. 3 (c) and shows that the areas of the plot with the strongest correlation between calculated and measured wavelength correspond to spectral regions with strong overlap between adjacent channels.

The wavelength plot is marked by two areas of strong correlation separated by flat regions in the red ($\lambda > 600$ nm) and green ($530 < \lambda < 560$ nm), with significant noise in the blue ($\lambda < 490$ nm) due to low signal levels.

Figure 4 shows how the detected spot wavelength varies with background optical properties. When card colored at opposite ends of the spectrum (red and blue) was used as a background (Fig. 4 (a)), a strong correlation between the calculated wavelength of corresponding spots is seen and all values lie along the identity line (with the exception of a number of points at the far end of the red region; $\lambda > 640$ nm).

For ex vivo tissue (Fig. 4 (b)), the wavelength values are again scattered along the identity line showing that the values outputted by the algorithm are in broad agreement in the 450-500 nm and 550-350 nm regions despite the noticeable difference in tissue color (inset). There is a noticeable increase in error however, at the blue end of the spectrum where a number of points deviate from the expected values due to the lower signal levels.

3.2. 3D tissue surface reconstruction

The results of the projector-camera calibration are shown in Fig. 5 (a) and indicate the relationship between the origin and propagation direction of the ‘best fit’ rays emitted from the probe tip and their intersection with the calibration object at different positions. The average error in fitting each ‘ray’ to its set of calibration points was 0.32 mm.

Two opaque validation objects were used to test the system’s 3D reconstruction. These were a flat object (black planar plastic divider, thickness 2 mm) and a brown polymer cylinder (radius 40.5 mm), whose 3D coordinates are shown in Figs. 5 (b) and (c) respectively. A mesh surface was fitted to the 3D point clouds using a least-squares minimization algorithm implemented in Matlab (The MathWorks, Inc., USA). Reprojection errors using the calibrated rays were 1.3 pixels in x and 1.06 pixels in y. The data in Fig. 5 (b) were fit by a planar surface with an error of 0.05 mm. The reconstructed cylindrical object was fitted by a cylinder
Fig. 5. Three-dimensional calibration and validation. (a) Origin and propagation direction of projected spots with respect to the camera (origin) as determined during calibration routine. (b) Planar object. (c) Cylindrical object, diameter = 81 mm. (d) Cross-section of cylindrical object with least-squares fit.

Fig. 6. Three-dimensional reconstruction of ex vivo tissue. (a) Porcine liver, ‘step’. (b) Ovine kidney, convex curve. (c) Porcine liver, convex curve. (d) Porcine tissue, border between fatty tissue and liver. (e) Porcine liver, ‘valley’.

with radius matching that of the validation object (40.5 mm) to sub-millimeter accuracy (rms difference = 0.15 mm) as shown in Figs. 5 (c) and (d).

Following the validation experiments, a number of samples of ex vivo tissue were examined to demonstrate operation of the probe and analysis algorithms over varying background optical properties and curvatures. The results in Fig. 6 show reconstructed 3D data for ovine kidney, porcine liver and fatty tissue, and have varying physical features (convex/concave curve, ‘step’ discontinuity and miscellaneous ‘fine’ structure). The reconstructed surface profiles were observed to match well the observed surface profiles of
the tissues, which can be observed in the color photographs presented. These tests demonstrate that the technique may be applied on \textit{ex vivo} tissue, while the validation with the test objects demonstrates the accuracy that may be obtained.

4. Discussion

The structured lighting probe described in this paper seeks to address the requirements for pattern projection in minimally invasive surgery by providing a pattern of high brightness, high density and employing a codification strategy based on spectral discrimination. The dispersed light from the supercontinuum laser ensures that each projected feature is labeled with a unique color which can then be detected from the camera RGB image using the segmentation algorithm described.

Initial results have shown that by using the segmentation algorithm with knowledge of the camera’s RGB system it is possible to identify the wavelength of individual spots. However, the performance of this algorithm is limited to regions of the spectrum where there is an overlap between the transmission spectra of adjacent filters (blue/green, green/blue). Outside of these regions, the camera only perceives a red, green or blue spot of varying intensity resulting in a flat response below 490 nm, above 600 nm and between 530 and 560 nm.

Using flat screens of different colors as objects, the calculated wavelength of individual spots was observed to be constant. This is a feature of this structured lighting probe due to the sharp spectra of the individual spots (FWHM \approx 5 nm), which meant that the observed color did not change and only a variation in spot intensity was observed as the background reflectance spectrum was altered. In biological tissues the same color invariance is observed, particularly in the overlapping spectral regions identified earlier. However, there are noticeable errors in the blue region caused by the diffusion of longer wavelength light from neighboring spots through the tissue between neighboring spots. The intestinal tissue sample was not as strongly absorbing in the 400-500 nm region as the kidney, meaning that adjacent spots were more likely to be influenced by scattered light. For the blue spots this had the effect of biasing the calculated wavelength towards the green. The blue spots were more susceptible to this effect by green rather than red spots due to the relatively lower number of red spots in close proximity. Strong absorption by hemoglobin at wavelengths below 500 nm also led to low reflected intensities at these wavelengths.

For non-biological validation objects, reconstruction of their 3D shape in metric space was demonstrated. High accuracy was achieved, with reprojection errors on the order of one pixel in the calibration and sub-millimeter accuracy in measuring a cylindrical and planar validation objects.

Sections of \textit{ex vivo} tissue were also reconstructed and visually appeared to match the tissue surfaces observed. For each tissue type the morphology could be distinguished, including the convex curves, valley and ‘step change’. However, fine structure in the fatty tissue was not resolved due to undersampling of the surface by the patterned light at that working distance, where the inter-spot spacing was of the order of several millimeters. This means that at long working distances this probe is limited to resolving sparse 3D structure with a resolution of less than a centimeter. However, at short working distances (2 cm or less) the pattern diameter could be reduced to \approx 2 cm and the inter-spot separation to \approx 2 mm or less, which would make more dense reconstruction possible. This would also be compatible with potential applications such as quantification of polyp shape during colonoscopy, where typical polyps can have diameters between 7 and 10 mm [4]. The segmentation and centroiding algorithm worked well in the green-red region of the spectrum, but a number of spots at the blue end could not be segmented and used in the reconstruction, reducing the accuracy of the final surface fit.

More calibration images could improve the accuracy of the calibration and hence the accuracy of reconstruction, however, the main source of error in the 3D reconstruction is in the detection of the spots. Their shape is dependent on distance and surface orientation, and
any errors, from a partial occlusion of a spot for example, means that the centroid does not reflect the correct ray path. The amount by which these image measurement errors are converted into triangulation errors is determined by the ‘baseline’ (distance between the projector and camera). Previous results from stereo endoscopy have already demonstrated that reliable reconstructions can be achieved using a small baseline (≈5 mm) if the working distance is sufficiently short (< 5 cm) [11,21], which suggests that a future application for this device could be a colonoscope.

5. Conclusions

A structured lighting probe has been built that is capable of delivering patterned light to the tissue using a compact fiber probe that is compatible with existing endoscopic devices. The optical set-up addresses the correspondence problem by coding the pattern with different wavelengths, assigning a unique wavelength to each projected spot. This can minimize the problem of occlusion, common to structured lighting set-ups, by tracking exactly which projected features have been blocked from view and avoid the risk of detected spots being matched to the calibrated rays of ‘missing’ spots.

A segmentation algorithm based on the CIE 1931 chromaticity diagram has been shown to successfully recover the wavelengths of individual spots in areas of the spectrum where there is an overlap between the transmission spectra of the camera’s RGB filters. In measurements on colored paper and tissue of varying optical properties it was shown that the calculated wavelength did not change with background reflectivity in the optimum region. This means that it will be possible to use the probe on a variety of tissues or at organ boundaries where optical properties may vary significantly.

Future work will be focused on optimizing the spectral output of the probe so that the number of spots with wavelengths in the optimal blue/green and green/red regions is maximized. A customized filter arrangement with increased spectral overlap will also be used to extend the range of these spectral regions. Spatial constraints will be added to the algorithm to aid discrimination of any spots that appear to have the same wavelength. To overcome some of the limitations imposed by absorption in tissue and to allow for practical clinical use, an interleaved imaging system will also be implemented. This would involve a high speed synchronized camera/shutter system to acquire patterned and white light images alternately in such a ratio as to minimize the visual impact of the pattern and make simultaneous normal white light viewing and 3D data acquisition possible. High-speed acquisition sequences of varying exposure times would also enable the acquisition of composite high dynamic range images where blue spots are made visible in the presence of hemoglobin without saturating those closer to the red end. Furthermore, absolute validations of the instrument accuracy in measurement of tissue will be carried out using ground truth data from CT images in order to better understand and minimize sources of error arising from diffusion and absorption of light. We believe that these modifications will allow a clinical structured lighting system to be constructed.

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