Line-scanning reflectance confocal microscopy of human skin: comparison of full-pupil and divided-pupil configurations

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Line-scanning, with pupil engineering and the use of linear array detectors, may enable simple, small, and low-cost confocal microscopes for clinical imaging of human epithelial tissues. However, a fundamental understanding of line-scanning performance within the highly scattering and aberrating conditions of human tissue is necessary, to translate from benchtop instrumentation to clinical implementation. The results of a preliminary investigation for reflectance imaging in skin are reported. © 2009 Optical Society of America OCIS codes: 040.1240, 180.2520.

Point-scanning confocal microscopes are well proven for optical sectioning and imaging of nuclear and cellular detail in human skin [1]. However, linescanning is fundamentally simpler, and the recent advent of high-quality linear array detectors may enable smaller and lower-cost confocal microscopy [2–4]. Scanning only once, directly behind the pupil of the objective lens, and descanning once, directly onto a linear array detector, allows considerable simplification of the original confocal line-scanning designs.

Of particular clinical importance is imaging performance with reflectance as an endogenous source of contrast, in highly scattering and aberrating human tissues. A fundamental understanding of confocal line-scanning performance in human tissue is therefore a necessary translational bridge from benchtop instrumentation to clinical implementation. For such investigations, the epidermis in human skin is easily accessible and available, and it is an excellent model for the scattering and aberrating properties of epithelial tissue. In this Letter, we present a full-pupil linescanning confocal microscope, and comparison of its performance to a divided-pupil configuration, for reflectance imaging of nuclear and cellular morphology in human epidermis, both *in vivo* and *ex vivo*.

As in the divided-pupil design that we reported earlier [5,6], the full-pupil line-scanning microscope also consists of eight to ten optical components in a simple configuration, with total hardware costs of \$15,000. The electronics is based on field programmable gate array (FPGA) logic. The development of FPGA-based electronics offers the advantages of small footprint, rapid reconfigurability, and complete integration toward a stand-alone system. In the longterm, line-scanning with the use of FPGA-based electronics may enable simple, small, low-cost, and stand-alone confocal microscopes for use at the bedside in diverse healthcare settings worldwide.

Figure 1 shows the full-pupil line-scanning confocal microscope. A collimated beam is incident on a cylindrical lens (L_{cyl}) , which is placed so as to produce a focused line in the back focal plane (BFP) of an objective lens (Obj). The focused line in the BFP is Fourier transformed or refocused to a line that is oriented perpendicular to the axis of the cylindrical lens. This produces a focused line in the object plane (within tissue) that is oriented perpendicular to the plane of this page. The line is scanned by an oscillating galvanometrically driven scan mirror (G). The scan mirror is placed as close as possible to the pupil of the objective lens. No relay telescopes are used such that assembly, alignment, testing, and troubleshooting is reasonably easy. However, the scanning does not occur exactly in the pupil of the objective lens. Thus, the focused line dithers across the pupil. Vignetting at the edges of the scan is minimized by overfilling the pupil. The line is ~ 10 mm in length, with dither

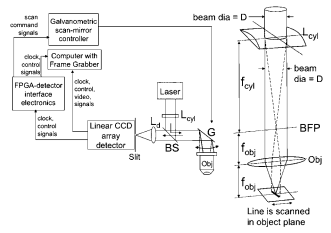


Fig. 1. Full-pupil line-scanning confocal microscope. Illumination is with a collimated beam of diameter ~10 mm from a diode laser at the near-IR wavelength of 830 nm (L4 830S-115-TE/ESYS, Microlaser, Inc.). The illumination path includes a cylindrical lens (L_{cyl}) of focal length f_{cyl} = 100 nm, a 50/50 thin plate beam splitter (BS), a galvanometric scan mirror (G, MiniSAX, GSI Lumonics), and a water immersion objective lens (Obj., Olympus 60×, 0.9 NA) of focal length f_{obj} =4 mm. The detection path includes a spherical detector lens (L_d) of focal length f_d =160 mm and a linear CCD array detector (Reticon LC3022 with photodiode RL1024P Perkin-Elmer). The magnification in the detection path is f_d/f_{obj} =40×.

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of ±0.9 mm, relative to pupil diameter of ~7 mm. The light that is backscattered from the object plane is relayed, descanned, and focused by a spherical lens (Ld) onto a linear CCD array detector. When imaging, the linear array detector, itself, acts as a detection slit aperture, with 14 μ m × 14 μ m pixels that correspond to subdiffraction resolution and almost satisfy Nyquist's sampling condition. The magnification in the detection optics almost satisfies the Nyquist requirement but results in the detected line underfilling the linear array detector by about 11% at the edges. This, along with an estimated 60% illumination falloff due to the Gaussian beam profile causes the edges of the image, in the nonscanned direction, to appear dark.

The video timing, control, and synchronization signals are based on an FPGA (Altera-Cyclone). Images are acquired with a frame grabber (Imaq PCI-1410, National Instruments). The linear CCD detector (LC3022, Perkin Elmer) consists of a linear photodiode array (Reticon RL1024P, Perkin Elmer) and operates at a maximum clock frequency of 20 MHz, allowing acquisition of images consisting of 1024 pixels \times 1024 lines with 8 bit gray-scale resolution, at a maximum rate of 20 frames per second. We calculate the sensitivity to be $\sim 10^8$ V/W. The specified dynamic range of ~ 2500 (~ 68 dB) is sufficient for imaging human skin.

With illumination power of $20-100 \ \mu$ W/pixel (i.e., $20-100 \ m$ W/line) on skin, the detected power is estimated to be ~4,000-20,000 photons/pixel in the linear array detector. The calculated rms quantum noise is ~57-179 electrons, while the specified rms read-out noise is ~50 electrons and dark noise is ~52 electrons. Thus, the rms signal-to-noise ratio (SNR) is estimated to be ~35-82. The SNR can be quantum noise-limited when the illumination power is high. However, in practice, dark noise and read-out noise are significant when there is loss of illumination power, especially when imaging deeper in tissue.

The nominal FWHM optical section in line scanning is 20% thicker than that with point scanning [7] and is given by $1.14 n\lambda/NA^2$ (i.e., $1.55 \mu m$) when measured with a plane object, where n is the refractive index (i.e., 1.33) of the immersion medium, λ is the illumination wavelength (i.e., 830 nm), and NA is the numerical aperture (i.e., 0.9) of the objective lens. Optical sectioning was experimentally evaluated in terms of the FWHM of axial line-spread functions (LSFs), using a $\lambda/20$ flat mirror and a 25 μ m slit aperture in front of a photodiode detector. The procedure was described earlier [5,6]. The slit width of $25 \ \mu m$ corresponds to slightly larger than the diffraction-limited lateral resolution. Measurements with the mirror were of the optical section thickness under nominal conditions. To further understand the effects of tissue-induced scattering and aberrations, measurements of LSF were also performed through an overlying layer of full-thickness human epidermis on the mirror surface. Human epidermis specimens, of measured thicknesses $\sim 50-75 \ \mu m$, were harvested from discarded surgical skin excisions under an Internal Revue Board-approved protocol. Five

measurements of optical section thickness were performed.

The nominal FWHM lateral resolution is $0.61\lambda/NA$ (i.e., $0.56 \ \mu m$) in the confocal and $0.79 \ \lambda/NA$ (i.e., $0.73 \ \mu m$) in the nonconfocal direction. Lateral resolution was experimentally evaluated in terms of the FWHM of edge-spread functions (ESFs), from images of a Ronchi ruling target. The FWHM lateral resolution is determined from the measured 10%-90% width of the ESF. The method was detailed earlier [5,6]. The lateral resolution was measured, again, under nominal conditions as well as through an overlying layer of full-thickness human epidermis on the Ronchi ruling target. Five measurements of lateral resolution were performed.

Figure 2 shows examples of measured axial LSFs, lateral ESFs, lateral LSFs obtained as derivatives of the ESFs, and images of an edge in a Ronchi ruling target, both under nominal conditions and through the scattering and aberrating conditions of full-thickness human epidermis. Table 1 shows the measured values for FWHM optical section thickness and lateral resolution for the full-pupil line-scanning confocal microscope. For comparison, also included are previous measurements with our divided-pupil configuration [5,6].

The nominal FWHM optical sectioning, when imaging in the superficial epidermis, is comparable for both pupil configurations. However, through the epidermis, when imaging in the underlying deeper dermis, the degradation is less than twofold for the fullpupil configuration but more than fivefold for the divided-pupil configuration. The higher difference and variability in optical sectioning with the dividedpupil configuration is due to the separate illumination and detection paths (which mimics the well-

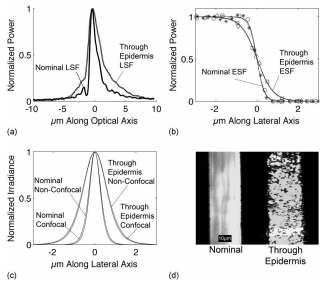


Fig. 2. (a) Experimentally measured axial LSFs, (b) lateral ESFs, and (c) lateral LSFs under both nominal conditions and through full-thickness human epidermis. The 10%–90% width of the ESFs is determined by approximating with sigmoidal fits. Images of an edge in a reflective chrome-on-glass Ronchi ruling target, showing, visually, the nominal resolution and the degradation through (d) epidermis.

Table 1. Measured Values of FWHM Optical Section Thickness and Confocal Lateral Resolution for	
Full-Pupil and Divided-Pupil Configurations ^a	

Measured Parameter		Full-Pupil Configuration	Divided-Pupil Configuration
FWHM optical section thickness	Nominal Through epidermis	$1.7 \pm 0.1 \ \mu m$	$1.7 \pm 0.1 \ \mu m$
FWHM confocal	Nominal	$2.8 \pm 0.8 \ \mu \text{m}$ $0.8 \pm 0.1 \ \mu \text{m}$	$9.2 \pm 1.7 \ \mu \text{m}$ $1.0 \pm 0.1 \ \mu \text{m}$
lateral resolution	Through epidermis	$1.6\pm0.3~\mu\mathrm{m}$	$1.7\pm0.1~\mu{ m m}$

^{*a*}For the full-pupil configuration, lateral resolution in the nonconfocal direction is nominally 0.9 μ m and 1.6 μ m through full-thickness epidermis.

known theta microscope geometry [8]). With separate paths, the overlying heterogeneous epidermis and dermo-epidermal junction produces spatially varying mismatches between the illumination and detection LSFs and loss of peak irradiance in the focal plane in the underlying deeper dermis. By comparison, the coaxial illumination and detection paths in the fullpupil configuration result in the LSF being less sensitive to the overlying epidermis and dermoepidermal junction.

The difference and variability in lateral resolution is comparatively smaller. The diffraction-limited FWHM lateral resolution with a confocal microscope is only 30% better than that with a conventional microscope [7]. In practice, this marginal improvement usually degrades owing to detection aperture size, instrumental imperfections, and tissue-induced scattering and aberrations, such that the two pupil configurations offer somewhat comparable lateral resolution.

Figure 3 shows images of human skin, both *in vivo* and *ex vivo*, with both full-pupil and divided-pupil line-scanning confocal microscopes. The images confirm that the measured nominal FWHM optical sectioning and lateral resolution are comparable for the full-pupil and divided-pupil configurations, and ad-

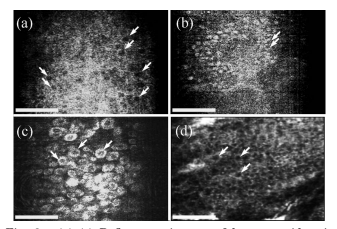


Fig. 3. (a)–(c) Reflectance images of human epidermis with the full-pupil line-scanning confocal microscope. Dark-appearing nuclei are seen within bright and grainy cellular cytoplasm (arrows) in (a) the spinous layers *in vivo* at depths of $\sim 20-50 \ \mu\text{m}$, (b) the basal layer *in vivo* at depths of $\sim 50-100 \ \mu\text{m}$, and (c) the superficial layers of surgically excised skin *ex vivo*. Also shown is an earlier image of the (d) spinous layer *in vivo* with the previously reported divided-pupil configuration. Scale bar, 75 $\ \mu\text{m}$.

equate for visualizing nuclear and cellular detail in the epidermis. Both out-of-focus multiply scattered background light and speckle noise is evident in the images. The visually assessed contrast is lower and the images somewhat more noisy with the full-pupil configuration than that seen earlier [5,6] with our divided-pupil configuration. The rejection of background light in the epidermis is stronger and the resulting image contrast superior with the dividedpupil configuration, owing to the angular separation of the detection path from the illumination path. Within dark nuclei, intranuclear structure is often distinctly noticeable, which is not seen with the fullpupil configuration. However, as noted in Table 1, through epidermis and when deeper in the underlying dermis, the full-pupil configuration provides significantly better optical sectioning with less tissueinduced variability.

In conclusion, line-scanning confocal microscopy may offer a competitive alternative to current pointscanning technology for imaging nuclear detail in epithelial tissues for clinical utility. The full-pupil configuration is easier to implement than the divided-pupil configuration. However, the choice of pupil configuration and the trade-offs in optical sectioning, resolution, and contrast will depend on the scattering and aberrating properties of tissue for any desired clinical application.

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