Assessing atherosclerotic plaque morphology: comparison of optical coherence tomography and high frequency intravascular ultrasound

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Abstract

Background—OCT can image plaque microstructure at a level of resolution not previously demonstrated with other imaging techniques because it uses infrared light rather than acoustic waves.

Objectives—To compare optical coherence tomography (OCT) and intravascular ultrasound (IVUS) imaging of in vitro atherosclerotic plaques.

Methods—Segments of abdominal aorta were obtained immediately before post-mortem examination. Images of 20 sites from five patients were acquired with OCT (operating at an optical wavelength of 1300 nm which was delivered to the sample through an optical fibre) and a 30 MHz ultrasonic transducer. After imaging, the microstructure of the tissue was assessed by routine histological processing.

Results—OCT yielded superior structural information in all plaques examined. The mean (SEM) axial resolution of OCT and IVUS imaging was 16 (1) and 110 (7), respectively, as determined by the point spread function from a mirror. Furthermore, the dynamic range of OCT was 109 dB compared with 43 dB for IVUS imaging.

Conclusions—OCT represents a promising new technology for intracoronary imaging because of its high resolution, broad dynamic range, and ability to be delivered through intravascular catheters.

Keywords: atherosclerosis; plaque morphology; infrared light; intravascular ultrasound; optical coherence tomography

The need for more detailed assessment of coronary microstructure, to improve risk stratification and guide interventional procedures, has driven the interest in developing new methods of imaging. Recently, high frequency intravascular ultrasound (IVUS) imaging has been used for the in vivo analysis of plaque morphology.1 3 IVUS imaging (20–30 MHz) is superior to conventional angiography in its ability to guide stent deployment, diagnose dissection, assess the effectiveness of coronary intervention, and determine the extent of vessel obstruction.4 6 Clinical trials have been conducted to determine the impact of IVUS imaging on patient management.7 Analysis of plaque microstructure is limited with IVUS imaging, however, because of the relatively low resolution (100–150 μm) and dynamic range (less than 50 dB).8–10 The identification of plaque features predisposed to rupture and progression to acute coronary syndromes in particular has had limited success. Technologies with superior performance may be required for the in vivo diagnosis of plaque instability.

Optical coherence tomography (OCT) is a recently developed imaging technology that shows considerable promise as a method for high resolution intravascular imaging.11 OCT uses infrared light to produce tomographic images on a micrometre scale. OCT imaging is analogous to ultrasound B mode imaging, except that it performs imaging by measuring the intensity of reflected or back scattered light rather than acoustic waves. Tomographic images are produced in a manner similar to that of radar. An optical beam is scanned across the tissue and the reflected or back scattered light is measured as a function of range (depth) and transverse position. The result is a two or three dimensional dataset that represents the intensity of the optical back scattering in a section through the tissue. The intensity of the reflected light is displayed as a false colour or grey scale image.

Although penetration of OCT imaging in non-transparent tissues is limited to a few millimetres, the typical image resolution of OCT is 10–15 μm, almost 10 times greater than ultrasound assessment, magnetic resonance imaging, or computed tomography. OCT imaging with resolution as high as 4 μm has been demonstrated in state of the art systems.12 Furthermore, OCT uses an optical fibre based design that can be readily integrated into catheters and endoscopes. OCT has been investigated, developed, and reduced to clinical practice for imaging transparent tis-
describes the first direct comparison between OCT and IVUS imaging, and assesses the relative performance of both modalities to image in vitro atherosclerotic aortic plaque morphology.

Methods
Aortas were obtained within six hours of post-mortem examination. The samples were stored in physiological saline with 0.1% sodium azide at 0°C. Imaging was performed with OCT and IVUS on segments smaller than 10 × 10 cm with the luminal surface exposed. All IVUS samples and most OCT samples were immersed in saline (OCT does not require saline immersion and imaging can be performed through air). Twenty consecutive sites from five patients were examined by OCT followed by ultrasonic imaging under the direction of a separate, blind, experienced operator. No attempt was made to select for specific plaque morphology due to the preliminary nature of this work. The position of the OCT beam on the sample was monitored with a visible light guiding beam. The peripheral areas of imaged sections were marked with microinjections of dye. The position of the ultrasound beam on the tissue was confirmed through landmarks such as injected dye, abrupt changes in the luminal surface, and metal markers placed in the tissue for orientation. Imaging was performed with specimens at room temperature.

After imaging, the specimens underwent routine histological processing, being fixed in formalin for 24 hours and subsequently decalcified in a standard Cal Ex solution (Fisher Inc, St Louis, Missouri, USA) for 10 hours. The arteries were processed with routine paraffin embedding. Sections of 5 μm thickness were cut at marked imaging sites, then stained with haematoxylin and eosin, and trichrome blue to identify different components of the vascular wall. Stained histological sections were compared with IVUS and OCT images to provide a better qualitative understanding of the reflectance properties of these tissue structures.

The principles behind OCT imaging have been previously described. OCT functions analogously to ultrasound imaging except that it uses light rather than sound waves. Ultrasound imaging is performed by measuring the delay time (echo delay time) for an incident ultrasonic pulse to be reflected back from different internal structures within the tissue. Because the velocity of sound is relatively slow, this delay time may be measured electronically. OCT performs imaging by measuring the echo delay time of light reflected from internal tissue structures. In contrast to sound, the velocity of light is extremely high, and thus direct measurement of optical echoes cannot be performed electronically. OCT measures the echo delay time of light by interferometry.

Figure 1 represents a schematic diagram of the OCT device. OCT imaging is performed using "low coherence" light generated by a compact, infrared diode source which is cou-
that they travel are matched to within a pulse duration.

The use of low coherence interferometry permits the echo delay time (and the optical path length) of the light reflected from the tissue sample to be measured with extremely high accuracy. In addition, the magnitude of optical interference is a measurement of the intensity of optical backscatter or reflection from the tissue. When the reference mirror is scanned, the backscattered or reflected light from different depths within the tissue sample is measured. The result is measurement of optical backscatter or reflection versus axial range and is analogous to ultrasound A mode ranging.

The optical beam is scanned across the sample and sequential axial measurements are taken at different transverse positions to construct a two-dimensional cross-sectional image of the tissue specimen. The resultant two-dimensional dataset represents the optical backscatter or reflection within a cross-section of the tissue specimen. This dataset can be displayed as a two-dimensional grey scale or false colour image. The magnitude of the optical backscatter is represented as a grey or colour scale to enhance differentiation of the tissue structures.

A superluminescent diode with a 1300 nm wavelength and a 50 nm band width (spectral width) was used as the light source for OCT imaging in this study. The spectral band width (axial resolution is inversely proportional to the spectral band width) yielded an axial spatial resolution of 16 (1) μm. The resolution was verified by measuring the point spread function using a mirror. The lateral or transverse resolution of 30 μm was measured with a Standard Air Force Resolution Chart. Unlike conventional microscopy, the axial resolution of OCT is determined by the coherence length of the light source and not by the focusing properties of the optical beam. Thus higher axial resolutions can be achieved by a shorter coherence length (broader bandwidth) light source.

The signal to noise ratio (SNR) was 109 dB, using an intensity of 160 μW at the sample. The SNR was determined by measuring the maximum detected signal when the optical beam is reflected from a mirror divided by the variance of the background noise level of the instrument. Images of backscatter intensity versus distance were displayed in grey scale or false colour. The axial dimension of the images correspond to 10 μm/pixel. The acquisition times ranged from 20 to 45 seconds, depending upon the size of the image acquired.

Ultrasonic imaging was performed with a 30 MHz, 2·9 French Microview ultrasonic transducer (Cardiovascular Instrument Systems, Sunnyvale, California), which has a mirror speed of 18 000 rpm. The dynamic range was 43 dB (Cardiovascular Instrument Systems, personal communication, 1995) and is similar to other IVUS transducers.10 34 Information was processed and displayed with an Insight III ultrasound system (Cardiovascular Instrument Systems). The image was...
spread functions for IVUS imaging and OCT were measured using a mirror. Digital data were converted to one dimensional data with image processing software (IPLab). Sampling was performed in the centre of the image and 1.5 mm in the transverse (lateral) direction to confirm that the resolution did not deteriorate because of the circular imaging field of the IVUS catheter. Widths were calculated at the full width half maximum. Values represent means (SE) of seven measurements.

Results

Figure 2 compares an OCT image of a haemorrhagic plaque with IVUS imaging and histology. In figure 2A, the IVUS image shows an irregularity in the luminal surface (red arrow). In addition, a hypoechoic area is seen within the vessel wall (green arrow). In figure 2B, the OCT image of the plaque wall shows significantly greater structural detail than the ultrasound image. The plaque and hypoechoic area are shown by OCT. Details that were not apparent with ultrasound assessment, however, become evident with OCT. By comparison with histology examination (fig 2C), the area of low back scattering within the plaque (1) contains large quantities of lipid, matrix, and red blood cells with little cellular structure. In the OCT image, the ringed structure (green arrow) represents an area with a high content of lipid and matrix. The ringed appearance resulted from a continuous line of lipid crescents. The left and right sides of the image appear to be layered where the relatively normal intima abuts a region with a high concentration of lipid and connective tissue.

Figure 3 shows OCT and IVUS images of a large lipid laden plaque. The lesion is also detected by ultrasound assessment (arrow). OCT, however, provides a sharper delineation between intimal wall and plaque. Furthermore, localised collections of lipid are identified with OCT but are not observed with IVUS. The elastic layer is poorly defined with both imaging technologies. Previous work46 has shown that when lipid laden plaque abuts the elastic layer, the optical back scattering contrast between the two layers is relatively small at this wavelength (1300 nm). The use of alternate wavelengths for OCT imaging may lead to improved contrast between layers.

Figure 4 shows a large plaque where layers in the adjacent wall are well demarcated. In figure 5, an intercostal artery emanating from the aorta is imaged with OCT and IVUS. Both imaging modalities demonstrate the presence of the vessel. However, structural detail is superior with OCT, illustrating the interweaving layers of tissue not differentiated by haematoxylin and eosin staining or IVUS but which are seen with trichrome blue staining (fig 5C).

Figure 6 shows the point spread function in the axial direction (which defines the axial resolution). The axial point spread function was 16 (1) μm for OCT and 110 (7) μm for IVUS at the centre of the image. Some 1.5 cm in the lateral (transverse) direction the resolutions

oversampled at 18 μm/pixel. Digital data from OCT and IVUS imaging were converted to analogue with IPLab (Signal Analytics, Vienna, Virginia) on a Power Macintosh 7100/66.

The point spread function was used to compare quantitatively the resolution of the two imaging technologies. Sound and light are strongly reflected at the surface of an interface such as that between air and metal. Thus the measured axial width of the returning signal is a measurement of the resolving power of the instrument. An imaging technology with an infinitely high resolution would resolve the surface as an infinitely sharp line. As any instrument has a finite resolution, however, the surface interface seems to have a finite width in the image. This is the point spread function and defines the axial resolution (the boundary cannot be resolved below the level of resolution of the instrument). The point...
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Figure 5. (A) IVUS and (B) OCT images, and (C) histological specimen showing an intercostal artery emanating from the aorta. Structural detail is superior with OCT, illustrating the intercrossing layers of tissue which are not evident with IVUS imaging. Bar represents 500 μm.

were 15 (1) μm and 107 (7) μm, respectively. Even though the IVUS catheter had a circular field of view, no deterioration in the lateral resolution occurred because the sampling (37 μm/pixel) is much higher than the resolution. In addition, multiple echoes are observed with IVUS, a problem that has been previously described.27 Multiple echoes were not noted with OCT imaging.

Discussion

Structural features such as intimal wall thickness, lipid content, and fissures have been shown by postmortem examination to predispose plaques to rupture and progression to acute coronary syndromes.28-31 Diagnostic technologies superior to those currently available will likely be required to identify these plaque features in vivo as the best current imaging technology, IVUS, does not provide sufficient resolution and contrast.68 Improved identification of plaque microstructure should significantly improve patient risk stratification.

In addition, an intravascular imaging technology with high resolution should improve efficiency and reduce complications associated with invasive catheter based treatment. Interventional procedures to manipulate atherosclerotic plaques could be used microsurgically if the guiding imaging technology could be extended to resolutions on a micrometre scale.

We have recently demonstrated the feasibility of OCT, a new method of micrometre scale imaging, for intravascular diagnostic investigation.16-17 Earlier imaging studies16 of in vitro human aortic and coronary atherosclerotic plaques have shown that fissures, lipid collections, and intimal wall caps can be identified with unprecedented resolution. In the present study, OCT and IVUS were directly compared in terms of their ability to image in vitro aortic plaques.

The most striking advantage of OCT compared with that of ultrasound assessment is the consistently higher resolution and contrast between components of the plaques. The dynamic range and sensitivity of OCT (−109 dB using an incident power of 160 μW) approaches the quantum limit, the theoretical maximum that can be achieved using optical detection. Although light is highly scattered in tissue, this high sensitivity and dynamic range permit imaging of tissue morphology to depths of 1–2 mm, a significant fraction of the vessel wall thickness. In contrast to ultrasound assessment, the contrast mechanism in OCT images arises from differences in the optical scattering properties of tissue rather than acoustic scattering properties. This leads to the dramatic difference in contrast between adjacent areas of distinct tissue composition, which is most prominently seen in figures 2–4. The high resolution, which was evident qualitatively in the images, was confirmed by measuring the point spread function at an interface with both imaging modalities.

The 16 μm resolution of OCT was five times greater than the 110 μm measured for IVUS. Furthermore, the resolution of OCT can be increased to the 4 μm range with the use of broad band width or femtosecond sources.12 Finally, multiple echoes that can lead to image distortion were observed with IVUS.15 Multiple echoes result because sound waves, reflected off the tissues, are also reflected off the surface of the ultrasound transducer (a high acoustical mismatch exists between the transducer and saline/blood). Echo artefacts were not observed with OCT.

The technology on which OCT is based is well suited for clinical use. Because OCT uses optical fibre technology and compact diode laser light sources, it can be engineered into a clinically compatible system. The OCT unit is compact and portable, similar in size to a standard external defibrillator or personal computer, making it attractive for clinical use. OCT is based on fibre optics that are used in optical communications technology, so that an optical fibre (with a standard diameter of
vivo imaging, but imaging time may be reduced significantly. Instrument improvements including increasing source power, scanning speed, and redesigning of electronics are straightforward and should lead to substantial reductions in imaging time. The imaging properties of blood are unknown. Although blood strongly absorbs visible light, it has very low absorption in the infrared region. Optical scattering, however, may reduce the amount of optical signal reflected from the vessel wall. The imaging properties of blood will be the topic of future investigation. If blood leads to a significant reduction in image quality, simultaneous injections of saline may be required for in vivo use similar to that needed in angiography. Unlike IVUS, imaging with OCT can be performed in air or saline without appreciable loss of image quality.

The penetration depths for IVUS versus OCT were not directly compared in this study as the focus of our investigation was to compare resolution. In general, ultrasound assessment is capable of imaging significantly deeper than optical techniques because attenuation and scattering are generally less for acoustic waves at 30 MHz than for light. The results presented in this study, however, suggest that OCT imaging depth will be sufficient for the diagnosis of a range of clinically significant coronary pathologies. OCT with a 1300 nm light source allows imaging through the width of a normal proximal coronary artery (2 mm). In addition, we have previously demonstrated that changing the wavelength of the incident light from 850 nm to 1300 nm results in significant improvements in imaging penetration.

Furthermore, optimisation of the wavelength and increasing the incident power (which is well below the level set by safety standards) will likely result in substantial improvements in OCT imaging penetration.

In conclusion, OCT imaging of plaque microstructure consistently demonstrates superior resolution and contrast when compared with that of IVUS. OCT represents a promising new technology for intravascular imaging due to its high resolution, broad dynamic range, and ability to be adapted for catheter based imaging.

We thank Ms Cindy Kopf for her assistance in the preparation of this manuscript, and Joe Gamba, Kathryn Blackwell, and James Taralli for their technical support. This work was supported in part by National Institute of Health grants NIH-R01-GM35459-09 and NIH-2-R01-EY11289-10; R29-HL55686-01A1, the Office of Naval Research Medical Free Electron Laser Program contract N00014-94-1-0717, and Whittaker Foundation contract 96-0205.

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Heart 1997 77: 397-403
doi: 10.1136/hrt.77.5.397

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