Multi-excitation fluorescence spectroscopy for analysis of non-alcoholic fatty liver disease

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Abstract

Background and objectives: the increasing incidence of non-alcoholic fatty liver diseases (NAFLD) and the consequent progression to cirrhosis is expected to become a major cause of liver transplantation. This will exacerbate the organ donor shortage and mean that ‘marginal’ fatty liver grafts are more frequently used. Autofluorescence spectroscopy is a fast, objective and non-destructive method to detect change in the endogenous fluorophores distribution and could prove to be a valuable tool for NAFLD diagnosis and transplant graft assessment.

Materials and methods: A system was constructed consisting of a fibre probe with two laser diodes that provided excitation light at 375 nm and 405 nm, and an imaging spectrograph system. This was used to distinguish fluorescence spectra acquired from the harvested livers from mice with NAFLD of differing severity (healthy, mild steatosis and steatohepatitic). The fluorescence data was entered into a sparse multiclass probabilistic algorithm for disease classification. Histopathology, TBARS and ALT assays were conducted in addition to the fluorescence measurements.

Results: Thiobarbituric Acid reactive substances (TBARS) and Alanine transaminase (ALT) assays enabled differentiation of the steatohepatitic group from the mild steatosis and control groups (p ≤ 0.028) but failed to separate the mild steatotic group from the control group. The three groups were all clearly differentiated from each other using fluorescence spectroscopy, and classification accuracy was found to be 95%.

Conclusion: Fluorescence spectroscopy appears to be a promising approach for the analysis of diseased liver tissue.
Introduction

Non-Alcoholic Fatty Liver Disease (NAFLD) represents a spectrum of progressive liver disease occurring in the absence of significant alcohol consumption, ranging from steatosis (fatty liver), through steatohepatitis (fat plus inflammation) to liver fibrosis, and ultimately cirrhosis [1,2]. NAFLD is associated with obesity and insulin resistance/diabetes and is characterized by an accumulation of triglycerides (fat) in the liver. Although steatosis has traditionally been considered a relatively benign condition [3], its evolution into more serious progressive fibrosis and cirrhosis of the liver is a concern. NAFLD has become the third most important indication for liver transplantation and is expected to become the leading indication in future decades [4].

Worldwide, the prevalence of NAFLD is estimated to be between 10 and 39% [5]. Its prevalence reaches 57% to 74% among the obese population, 50% among diabetic patients [6] and the overall prevalence is set to rise with the worldwide epidemic of obesity and diabetes (the obese adult population will rise from at least 400 million in 2005 to 700 million by 2013 [7], while the diabetes population is expected to grow from 285 million in 2010 to 438 million in 2030 [8] according to the WHO).

Given the shortage of organ donors, and the high prevalence of NAFLD within the population, the use of steatotic (fatty) ‘marginal’ grafts is increasingly common in liver transplantation [9]. However, severe steatosis and steatohepatitis [10,11] are associated with increased risk of primary graft non-function. Consequently, to aid effective organ selection, there is a need to rapidly and robustly assess donor organs for degree of steatosis and the presence of significant oxidative stress, as these may indicate grafts at greater risk of non-function after transplantation.

At present there is no rapid and accurate test to diagnose NAFLD or non-alcoholic steatohepatitis (NASH). For example, blood biochemistry is insensitive and non-specific, whilst biopsy is time-consuming and not well suited to peri/intraoperative assessment of fatty liver [12]. Several non-invasive imaging techniques have been tested for NAFLD diagnosis. Ultrasound detects steatosis with a limited sensitivity (67%) and specificity (77%) [13], but gives better results when the severity of the steatosis increases [14]. CT imaging enables a sensitivity of 100% and a specificity of 82-93% (10) for steatosis diagnosis. MRI allows the most accurate fat quantification and hence enables the best sensitivity (100 %) and specificity (92 %) for steatosis diagnosis [15]. However, because all of these techniques measure the triglycerides, they can only assess the severity of steatosis (fat content) and are unable to differentiate steatosis from steatohepatitis or cirrhosis since they are characterized by tissue inflammation and fibrosis rather than by triglyceride concentration. In fact, when steatohepatitis and cirrhosis are well established the liver, fat content may decrease, potentially leading to an underestimation of the disease severity [16]. Therefore, a fast and minimally invasive method able to differentiate normal tissue from steatosis and from steatohepatitis would be desirable.

Fluorescence spectroscopy has been extensively used for cancer diagnosis in various organs because it provides a fast, objective, minimally invasive (as the instrumentation can be
miniaturised and does not use harmful or ionising radiation) and less traumatic alternative to histopathology [17]. Fluorescence spectroscopy has several potential applications in liver tissue characterisation. In an ex vivo setting it could be an alternative or a complement to histopathology to assess liver biopsies and would also enable non-destructive detection of NAFLD in liver transplant grafts which are more prone to primary non-function. In vivo, it could replace biopsy in patients for whom biopsy would be a high risk procedure. It could also be used as an adjunct in patients already undergoing a surgical procedure (bariatric surgery) or to assess the donor organ at harvest.

Autofluorescence from liver has previously been studied by Croce et al [18-21] in a laboratory microscopy setting. In this work, it was shown that NAD(P)H, vitamin A and flavins were major contributors to liver autofluorescence when excited at 366 nm. In particular, it was shown that accumulation of lipids led to an increased presence of the fluorophore, vitamin A. Also, steatohepatitis can be accompanied by fibrosis, a proliferation of the connective tissue whose main component is fluorescent collagen. Other contributors to the liver autofluorescence are likely to be flavins, riboflavin, arachinoid acid, lipopigments, and porphyrins [22-25], all found to be present in liver and emitting in the visible range [26]. Thus, the construction of a fluorescence spectroscopy probe system seems a promising candidate to non-invasively diagnose and characterise steatosis and steatohepatitis.

In this study a fluorescence spectroscopy probe was constructed that could collect fluorescence emission spectra at two excitation wavelengths. This could excite different subspecies of the endogenous fluorophores present and the emission spectra at different excitation wavelengths could then be used to distinguish different tissue states. A sparse multinomial logistic regression enabled differences between normal, steatotic and steatohepatitic livers to be detected in an animal model.

Material and methods

Multi-excitation fluorescence spectroscopy system

The violet laser fluorescence system combined two laser diodes emitting at 375 nm and 405 nm and a custom made probe (Romack, Inc) consisting of six hexagonally packed fluorescence collection fibres surrounding an excitation fibre that delivered the laser light to the sample (figure 1). The diameter and numerical aperture of the fibres were 200 µm and 0.22 respectively and they were chosen to have low autofluorescence and attenuation in the near ultraviolet spectral region. The probe distal tip was covered by a glass window and the fibres and window were housed in a stainless steel tube with an external diameter of 2 mm. This could enable it to be inserted into endoscope working channels, needles or small orifices. The proximal end was divided into two arms, the first one contained the excitation fibre and was coupled to the laser optics using an SMA connector; the second arm included the six emission fibres arranged in a linear array inside another SMA connector.
The laser light from the two laser diodes was collimated by 4.6 mm focal length lenses and the beams were combined using a dichroic mirror so that both beams could be coupled into the excitation fibre using an 8 mm achromatic lens, with two steering mirrors used to align one of the beams and ensure coaxial propagation. Two beam shutters (one for each laser diode) enabled the selection of the excitation wavelength delivered to the sample. The optical power at the distal end of the probe was 3.5 mW for both lasers.

The fluorescence emission from the six fibres was focussed onto the 80 µm wide input slit of an imaging spectrograph (Specim, Ltd) using an achromatic lens doublet (60 mm and 30 mm). A 430 nm long-pass filter was inserted between these two lenses to block the light of the laser diodes reflected by the sample. The light dispersed by the prism-grating-prism element of the spectrograph was then acquired with a sensitive cooled CCD camera (Retiga EXI, Qimaging, 1392 x 1040 pixels). All the optics were assembled together using a cage system and a breadboard so that the system was robust and portable. A LabVIEW program controlled the exposure time (100 ms) and beam shutters and enabled acquisition of fluorescent signals from the laser diodes sequentially with an adjustable number of measurements.

**Study design and animal husbandry**

Research was approved by the local ethics committee and performed in accordance with the UK Animal (Scientific Procedures) Act 1986. Mice were housed under standard conditions and provided with a commercial chow (#801722, SDS, UK) and free access to water. The study was conducted to investigate the autofluorescence spectroscopy of liver tissue from mice with varying degrees of NAFLD and used mouse models with steatosis and steatohepatitis induced by a combination of genetic and dietary modulation [1].

8-week old male non-obese heterozygote Db/m control mice were fed with a control diet (BKS-Cg-m +/- Lepr^db/J, Charles River, Belgium - ‘control group’). This was compared to genetically obese Db/Db mice (BKS-Cg-m +/- Lepr^db/J, Charles River, Belgium) fed either with a nutritionally replete control diet (#A02082003B - ‘mild steatosis group’) or with a choline deficient diet (#A02082002B, Research Diets, USA - ‘steatohepatitis group’). There were initially five animals per group although during the study two of the mice from the mild steatosis group became unwell and had to be culled early and were excluded from the study.

At the end of the study, animals were culled by exsanguination under general anaesthesia and liver tissue was quickly harvested. The liver was bisected and half was immediately snap-frozen in liquid nitrogen for subsequent biochemical analysis. Blood was also obtained for biochemical analysis. Without any treatment or staining, liver tissues were submitted to measurements immediately after culling to minimize air-exposure effects, cleaned of blood to minimize haemoglobin absorption and brought from the operating table to the optical bench. Emission spectra were recorded and the livers were then fixed in formalin solution for further histopathological study.
Data was acquired at between three and five different sites on each liver. At each site six measurements were recorded from the six emission fibres in the probe, probing an overlapping but slightly different tissue volume. This leads to a total of 114 measurements for the normal group, 72 measurements from the steatotic group, and 114 measurements for the steatohepatitic group (note that one measurement consists of two emission spectra – one for 375 nm and one for 405 nm excitation).

**Data processing and analysis**

The six emission spectra recorded for each excitation wavelength were extracted from the CCD images and the intensities at different spatial locations on the image were converted into spectra using calibration data obtained with a number of specific laser wavelengths. The spectrum of a tungsten-halogen calibrated light source (Ocean Optics Inc.) was also recorded at the start of the experiment to take into account the system spectral response and the ratio of this spectrum over the real spectrum was used to correct the raw spectra. Spectra were normalized by the area under the curve and truncated between 450 and 750 nm. The spectra from the two excitation wavelengths were concatenated prior to processing for each measurement.

Sparse multinomial logistic regression (SMLR), as previously developed by Krishnapuram et al. [27], was used to separate the spectral features into three classes: control, mild steatosis and steatohepatitis with an open source software package [28]. SMLR belongs to the family of supervised learning algorithms; it proposes a true multiclass classification of the spectra using classifiers built as a linear combination of the input features, i.e the 106 fluorescence intensities at the 106 wavelengths sampled between 450 nm and 750 nm comprising each spectrum. SMLR has previously been demonstrated to be highly effective in differentiating autofluorescence and diffuse reflectance spectra from healthy, marginal, and tumourous brain tissue [29,30]. The SMLR classification was performed here with a K-fold cross validation rather than with a leave-one-out cross validation since the six spectra from one acquisition cannot be considered to be totally independent. In this approach, the sample pool of 300 spectra was divided into 50 subsets (K=50) each containing the six spectra from the same acquisition so that during the classifier learning groups of six were held out at a time.

The probability that a spectrum belonged to a certain class is assumed to follow the multinomial logistic regression model where the regression coefficients are the classifier weight vectors of the corresponding class. The spectrum was assumed to belong to the class whose weight vector maximised this probability. The estimation of the weight matrix (106 lines, three columns) for the three different classes (in other words the result of the classifier learning process), was calculated from a maximum *a posteriori* estimate, with the spectra as input training data. This approach contains a number of advantages over traditional maximum likelihood estimation, as described in [27,31].
**Histopathology**

Formalin fixed liver tissue was processed into paraffin wax and sections were stained with H&E. H&E stained sections were assessed using the validated NASH Clinical Research Network histological classification system. Steatosis was scored 0-3 where score 0 = <5% steatosis, 1 = 5-33% steatosis, 2 = 33-66% steatosis, and 3 = >66% steatosis. Lobular inflammation was scored 0-3 where score 0 = none, 1 = <2 foci per x200 field, 2 = 2-4 foci per x200 field, and 3 = >4 foci per x200 field. Finally hepatocyte ballooning was scored 0-2 where score 0 = none, 1 = few ballooned cells, and 2 = prominent ballooning. These scores were then added together to give a NAFLD activity score (NAS) which decides whether a histological diagnosis of NASH is made (≥5 is definite NASH) [32].

**TBARS and ALT biochemistry**

The MCD mouse model of NASH is characterised by increased reactive oxygen species production. To quantify this, a TBARS assay was performed. Free radical damage ("tissue oxidative stress") is a major driver of disease pathogenesis, inflammation and progression in NASH. TBARS is not used in standard practice but is an accepted research tool to determine lipid peroxidation as a surrogate measure for free radical damage within the liver. Inclusion in this study is to provide a comparator directly related to disease pathogenesis beyond that which would be available in routine practice. In practice, the clinical diagnosis of NASH is based on the identification of raised alanine transaminase levels (indicating inflammation) in patients with features of the metabolic syndrome (e.g. obesity, diabetes and dyslipidaemia) with supporting evidence of fatty liver on ultrasound imaging. Approximately 100 mg of frozen liver was placed into a ceramic beads tube. Then 300µl of RIPA buffer was added and the tubes placed into a Precellys rotor at 6500 rpm for 1 cycle of 50 seconds to homogenise the samples. The supernatant was then kept on ice and used in the assay. 100 µl of the lysate was placed into a 1.5 ml microcentrifuge tube and 200 µl of ice cold 10% Trichloracetic acid was added. The sample was then incubated for 15 minutes on ice followed by centrifugation at 3000 g for 10 minutes. Standards were prepared using distilled water and 500 µM Malondialdehyde (MDA) to give known MDA concentrations from 0 µM to 50 µM. 300 µl of the samples and standards were then mixed with 300 µl of 0.67% thiobarbituric acid and incubated at 95 °C for 10 minutes. The samples were then allowed to chill.

Using a 96 well plate 150 µl of each standard and sample was placed into a well, performed in duplicate. Spectrophotometry at 532 nm was performed using a plate reader. Using the absorbance values and standard curve the MDA concentration in µM per 100 mg of liver was calculated for each sample.

The blood obtained when the mice were culled was placed in a plasma collection tube which when centrifuged at 3000 rpm for 10 minutes separated the plasma from the erythrocytes. The
supernatant plasma was removed from the blood and stored at -80°C until analysed in an automated biochemical analyser to give ALT measurements in IU/L.

RESULTS

Histopathology

H&E stained sections showed that the three groups could be clearly distinguished. The non-obese Db/m mice fed chow ('control group') had normal liver histology. The Db/Db mice fed chow ('mild steatosis group') exhibited features of isolated steatosis without lobular inflammation and the Db/Db mice fed a MCD diet ('steatohepatitis group') had NASH with marked steatosis, moderate lobular inflammation and many ballooned cells (see Table 1 and Figure 2).

TBARS Assay

There were significantly raised MDA concentrations in the livers of Db/Db mice on MCD diet (steatohepatitis group) compared to the two other study groups, as expected - p=0.028 with Db/Db mice on standard chow (control group) and p=0.011 with Db/m mice on standard chow (mild steatosis group), see Figure 3. The mean MDA concentrations (µM per 100 mg of liver) were 0.26±0.02 for the Db/m mice on standard chow, 0.28±0.02 for the Db/Db mice on standard chow and 1.79±0.46 for the Db/Db mice on MCD diet (Table 2). These results confirm that there was increased oxidative stress within the liver of the Db/Db mice on the MCD diet. No significant difference was detected between the chow fed Db/Db mice with isolated steatosis or the phenotypically normal Db/m mice fed chow (Figure 3).

ALT Biochemistry

There were significantly increased ALT levels in the plasma of the Db/Db mice fed MCD diet compared to the two other study groups, as expected (p=<0.001 for Db/Db mice on standard chow and Db/m mice on standard chow - see Figure 4). The mean ALT levels in the plasma (IU/L) were 26.40±4.07 for the Db/m mice on standard chow, 91.77±4.78 for the Db/Db mice on standard chow and 307.34±33.38 for the Db/Db mice on MCD diet (table 3). These results confirm that there was biochemical hepatitis within the liver of the Db/Db mice on the MCD diet which is in keeping with the lobular inflammation and ballooning identified on H&E stain. No significant difference was detected between the chow fed Db/Db mice with isolated steatosis or the phenotypically normal Db/m mice fed chow although there does appear to be a trend towards increased ALT in the steatotic Db/Db mice.

Fluorescence spectroscopy results

Mean spectra at 375 nm and 405 nm excitation and error bars representing data standard deviation are presented in Figure 5. At 375 nm excitation, liver autofluorescence peaked at around 475 nm which is attributed to the combined fluorescence of NAD(P)H (emission at around
475 nm) and vitamin A (emission at around 490 nm) [20]. Mild steatotic tissue showed relatively higher 475 nm intensity than healthy tissue, with intensity increasing in conjunction with the severity of the steatosis in these two groups. In steatohepatitis the tissue exhibited lower peak fluorescence than healthy and steatotic tissue. The 405 nm excitation spectra revealed a peak intensity at 500 nm. A secondary peak at 610 nm can be attributed to porphyrins, which has an excitation peak near 405 nm and an emission peak at 610 nm and is synthesised in liver (24). There was a weak 575 nm peak for both excitation wavelengths which could come from lipofuscin, a lipopigment present in the liver [25] which has a maximum fluorescence emission around 575 nm [26] and a concentration that increases with age.

At 405 nm, spectral differences between disease states seem visually less apparent than at 375 nm excitation, although note that the difference is still statistically significant, as shown in the following sections. For this excitation wavelength, vitamin A and NAD(P)H are less strongly excited so their contributions to the fluorescence spectra will appear slightly less important than at 375 nm excitation. The spectral shape also results from the contributions of the other fluorophores present in the liver including flavins, [22], which have an emission band between 500 nm and 600nm [26], and arachidonic acid, which is a free fatty acid metabolized by the liver [23] that emits fluorescence between 400 and 550 nm [18].

**Data processing and analysis**

The results of the SMLR classification performed with a leave-one-out cross validation are presented in the confusion matrices in table 4 for the concatenated dataset. The overall classification accuracy obtained reached 95%.

This table can be reinterpreted in terms of diagnostically more meaningful parameters by considering the sensitivity, specificity, positive and negative predictive values (PPV, NPV) of one group against the two others. The normal tissue (control group) could be differentiated from the mild steatotic and steatohepatitic tissue with 95.6% sensitivity, 96.8% specificity, 94.8% PPV and 96.8% NPV. The steatotic tissue was discriminated from the normal and steatohepatitic tissue with 91.7% sensitivity, 96% specificity, 88% PPV, and 97.3% NPV. Steatohepatitic tissue was separated from the two other groups with 96.5% sensitivity, 100% specificity, 100% PPV and 97.9% NPV.

The posterior probabilities calculated by SMLR of fluorescence spectra belonging to the correct disease group as classified by histopathology are plotted in figure 6. Fifteen sites were spectrally misclassified with a probability below 50%, with the remainder classified with an average probability of 98.9% indicating a strong confidence in the classification of the tissues.
Discussion

This study aimed to assess fluorescence spectroscopy as a way to differentiate healthy livers from steatotic and steatohepatitic livers and compare its performance with ALT and TBARS assays.

Autofluorescence from liver has previously been studied in a tissue microscopy instrument under 366 nm laser excitation by Croce et al. In that study the analysis of the fluorescence spectra and photobleaching from rat livers enabled the determination of the fluorophores responsible for liver autofluorescence [33,34]. Among these, NAD(P)H, vitamin A and collagen were identified as the primary markers to distinguish healthy, steatotic and fibrotic livers. Whilst this work was decisive for the understanding of liver fluorescence and opened perspectives on the potential of fluorescence spectroscopy as a tool to assess liver state, the instrumentation used in this work required the tissues to be excised, sectioned and prepared onto microscope slides. In another study, De Oliveira et al. acquired spectra in a rat liver model with different degrees of steatosis at 532 nm excitation using an optical fibre probe [34]. The most severe steatosis cases were clearly distinguished from the healthy ones by using the ratio of the fluorescence peak intensity at 605 nm over the backreflected excitation light intensity at 532 nm. Moreover, this factor averaged within each study group is shown to follow linearly the fat concentration. However, consecutive study groups could not be differentiated one from another based on this ratio. The use of an algorithm utilising the whole spectrum, similar to SMLR, together with excitation at a wavelength that will allow autofluorescence from vitamin A, collagen, NADH and porphyrins is expected to improve the classification. This work noticeably showed the potential of fluorescence spectroscopy as a tool for NAFLD detection but did not directly assess its diagnosis performance.

The steatohepatitis group exhibited noticeably increased ALT and TBARS levels compared to the steatosis and control groups, leading to a clear differentiation of the steatohepatitic group with a p-value below 0.001. However, no significant differences in these parameters were found between the control and steatotic groups and it was not possible to differentiate these two groups based on ALT and TBARS levels. Fluorescence emission spectra from different disease classes exhibited differences in their spectral shape as shown in figure 5 (A) and (B). The most visually prominent feature was the difference in fluorescence intensity at the peak (around 475 nm) for 375 nm excitation. The normal, mild steatosis and steatohepatitis groups have steatosis scores of 0, 2 and 3 respectively using NAS (see the “steatosis” column in table 1). The relative increase in intensity at 475 nm for 375 nm excitation from the normal (steatosis score 0) to the mild steatosis group (steatosis score 2, see figure 5A) is consistent with an increase in vitamin A content concomitant with the augmentation of lipids (20) accompanied by an increase in NAD(P)H, which has been shown to soar with steatosis (35).

The steatohepatitis group had a steatosis score of 3 and may therefore have been expected to have a higher intensity at 475 nm than the other groups. However, it actually has a lower intensity (Fig 5A), which may be due to the biochemical and biostructural changes induced by
fibrosis in this model of steatohepatitis. Firstly, collagen which is induced by fibrosis is a well known scatterer and fluorophore which will affect the fluorescence and scattering properties of the liver tissue. Secondly, the contribution of vitamin A in fibrotic liver has been shown to be half that of healthy liver when excited at 366 nm [21]. Therefore, the presence of fibrotic tissue in the fibrosing steatohepatitic livers is likely to decrease the peak intensity in this group. Also, the evolution of NAD(P)H with fibrosing steatohepatitis is unknown. The difference between the spectra excited at 405 nm in the spectral interval from 450 nm to 500 nm, was visually less obvious than for 375 nm excitation, although note that the difference is still significant, and that the results from the classification algorithm account for the response across the whole emission spectrum. Spectral differences between the 405 nm excitation spectra also stem from other liver fluorophores such as flavins and arachidonic acid.

The classification accuracy for single excitation was 80% and 95% for 375 nm and 405 nm excitation respectively. The classification with a combination of spectra at the two excitation wavelengths was found to be 95% and therefore did not improve the result obtained with the 405 nm laser diode only. This suggests that the optical system could be simplified by keeping only the 405 nm laser diode although the combination of the 375 nm and 405 nm might improve the classification performance with other models of NAFLD (data in preparation). Therefore, autofluorescence spectroscopy enabled the differentiation of the three groups with a high classification accuracy that could not be achieved with the ALT and TBARS levels. Moreover, the spectroscopic classification was nearly real time and hence significantly faster than the ALT and TBARS assays (typically 1 day) as well as histopathology (typically at least 2 days). Although in most cases fatty infiltration within the liver is relatively uniform, all diagnostic techniques used in the liver are open to sampling error. Fluorescence spectroscopy should be no more sampling error prone than biopsy, but has the added advantage that multiple readings may be taken without causing tissue damage and so any error may be minimised.

One advantage of the SMLR is that it computes the posterior probability of a spectrum belonging to a particular class, and as pointed out by Majumder [29], this can allow suspicious sites with less than 50% probability of being healthy to be identified and further examined. It could therefore be possible to diagnose NAFLD using fluorescence spectroscopy on freshly excised liver explants or biopsy material. Depending on the posterior probability of classification as healthy, the clinician could then decide on further histological examination. This procedure could help to speed up the diagnostic process as the result from spectroscopy is almost instantaneous, as well as having potential for intraoperative quantitative diagnosis and classification of NAFLD severity in donor liver tissue which may assist in graft selection in the transplant setting.

This result has been obtained in an animal model involving NAFLD mice only, where the contrast between disease states was unambiguous. In practice, NAFLD severity may need to be discriminated from other inflammatory conditions and from varying degrees of fibrosis or cirrhosis. The ability of fluorescence spectroscopy to distinguish NAFLD patients in these different situations remains to be tested.
Conclusion

In summary, we have built a compact dual excitation spectroscopy system using laser diodes that can be used minimally invasively as the optical fibre probe has a 2 mm diameter. Fluorescence spectra at both excitation wavelengths were recorded from healthy, steatotic and steatohepatitic mice and analysed with SMLR. Excellent classification accuracy with strong confidence in the classification was obtained which support the use of fluorescence spectroscopy as a tool to quickly assess liver tissue graft ex vivo or in vivo at harvest.

Acknowledgements

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References

Legends

Figure 1: Experimental set-up of multi excitation wavelength fluorescence spectroscopy probe.

Figure 2: Typical H&E stained liver sections at x 400 magnification showing (A) Db/m mouse liver fed standard Chow., (B) Db/Db mouse liver fed standard Chow showing steatosis only, (C) Db/Db mouse liver fed MCD showing steatosis, lobular inflammation and hepatocyte ballooning.

Figure 3: Mean malondialdehyde concentration in the livers of the mice ±SEM.

Figure 4: Mean ALT levels in the plasma of the mice ±SEM.

Figure 5: Mean spectra at (A) 375 nm and (B) 405 nm excitation and error bars representing standard deviation.

Figure 6: Posterior probability of fluorescence spectra from interrogated tissue sites belonging to their correct disease group (as classified by histopathology): control (114 sites), mild steatosis (72 sites), steatohepatitis (114 sites).
Figure 3: typical H&E stained liver sections at x 400 magnification showing a) Db/Db mouse liver fed MCD showing steatosis, lobular inflammation and hepatocyte ballooning, b) Db/Db mouse liver fed standard Chow showing steatosis only, c) Db/m mouse liver fed standard Chow.

Figure 4: Mean malondialdehyde concentration in the livers of the mice ±SEM
Figure 5. Mean ALT levels in the plasma of the mice ±SEM

Figure 6: Mean spectra at (a) 375 nm and (b) 405 nm excitation and error bars representing standard deviation.
Figure 7: Posterior probability of fluorescence spectra from interrogated tissue sites belonging to their correct disease group (as classified by histopathology): steatohepatitis (114 sites), mild steatosis (72 sites), control (114 sites).

<table>
<thead>
<tr>
<th>Disease group</th>
<th>Number of measurements</th>
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<tbody>
<tr>
<td>Steatohepatitis (A)</td>
<td>114</td>
</tr>
<tr>
<td>Mild steatosis (B)</td>
<td>72</td>
</tr>
<tr>
<td>Control (C)</td>
<td>114</td>
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</tbody>
</table>

Table 1: Number of tissue fluorescence measurements per disease group.

<table>
<thead>
<tr>
<th>Study Group</th>
<th>Steatosis</th>
<th>Lobular inflammation</th>
<th>Hepatocyte ballooning</th>
<th>NAFLD activity score</th>
</tr>
</thead>
<tbody>
<tr>
<td>Steatohepatitis A</td>
<td>3</td>
<td>2</td>
<td>2</td>
<td>7</td>
</tr>
<tr>
<td>Mild steatosis B</td>
<td>2</td>
<td>0</td>
<td>0</td>
<td>2</td>
</tr>
<tr>
<td>Control C</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
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Table 2. Median NAFLD activity score for the mice.
### Table 3: Mean malondialdehyde concentration in the livers of the mice ±SEM

<table>
<thead>
<tr>
<th>Pilot Study Group</th>
<th>Malondialdehyde Concentration (µM per 100mg of liver) ± SEM</th>
</tr>
</thead>
<tbody>
<tr>
<td>Db/Db mice on MCD diet Steatohepatitis group A</td>
<td>1.79±0.46</td>
</tr>
<tr>
<td>Db/Db mice on standard chow Mild steatosis group B</td>
<td>0.28±0.02</td>
</tr>
<tr>
<td>Db/m mice on standard chow Control group C</td>
<td>0.26±0.02</td>
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### Table 4. Mean ALT levels in the plasma of the mice ±SEM.

<table>
<thead>
<tr>
<th>Pilot Study Group</th>
<th>ALT biochemistry (IU/L) ± SEM</th>
</tr>
</thead>
<tbody>
<tr>
<td>Db/Db mice on MCD diet Steatohepatitis group A</td>
<td>307.34±33.38</td>
</tr>
<tr>
<td>Db/Db mice on standard chow Mild steatosis group B</td>
<td>91.77±4.78</td>
</tr>
<tr>
<td>Db/m mice on standard chow Control group C</td>
<td>26.40±4.07</td>
</tr>
</tbody>
</table>

### Table 5: Confusion matrix presenting the ability of fluorescence spectroscopy to distinguish between control, moderate steatosis and steatohepatitis disease groups.

<table>
<thead>
<tr>
<th>Disease group</th>
<th>Fluorescence spectroscopy</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Steatohepatitis</td>
</tr>
<tr>
<td>Histopathology</td>
<td>113</td>
</tr>
<tr>
<td>Mild steatosis</td>
<td>0</td>
</tr>
<tr>
<td>Control</td>
<td>0</td>
</tr>
</tbody>
</table>

**Table 3:** Mean malondialdehyde concentration in the livers of the mice ±SEM

**Table 4:** Mean ALT levels in the plasma of the mice ±SEM.

**Table 5:** Confusion matrix presenting the ability of fluorescence spectroscopy to distinguish between control, moderate steatosis and steatohepatitis disease groups.