Horizons in Nutritional Science

Application of advanced synchrotron radiation-based Fourier transform infrared (SR-FTIR) microspectroscopy to animal nutrition and feed science: a novel approach

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Synchrotron radiation-based Fourier transform IR (SR-FTIR) microspectroscopy has been developed as a rapid, direct, non-destructive and bioanalytical technique. This technique, taking advantage of synchrotron light brightness and a small effective source size, is capable of exploring the molecular chemistry within the microstructures of a biological tissue without the destruction of inherent structures at ultra-spatial resolutions within cellular dimensions. This is in contrast to traditional ‘wet’ chemical methods, which, during processing for analysis, often result in the destruction of the intrinsic structures of feeds. To date there has been very little application of this technique to the study of feed materials in relation to animal nutrient utilisation. The present article reviews four applications of the SR-FTIR bioanalytical technique as a novel approach in animal nutrition and feed science research. Application 1 showed that using the SR-FTIR technique, intensities and the distribution of the biological components (such as lignin, protein, lipid, structural and non-structural carbohydrates and their ratios) in the microstructure of plant tissue within cellular dimensions could be imaged. The implication from this study is that we can chemically define the intrinsic feed structure and compare feed tissues according to spectroscopic characteristics, functional groups, spatial distribution and chemical intensity. Application 2 showed that the ultrastructural–chemical makeup and density of yellow- and brown-seeded *Brassica* rape could be explored. This structural–chemical information could be used for the prediction of rapeseed quality and nutritive value for man and animals and for rapeseed breeding programmes for selecting superior varieties for special purposes. More research is required to define the extent of differences that exist between the yellow- and brown-seeded *Brassica* rape. Application 3 showed with the SR-FTIR technique that chemical differences in the ultrastructural matrix of endosperm tissue between Harrington (malt- ing-type) and Valier (feed-type) barley in relation to rumen degradation characteristics could be identified. The results indicated that the greater association of the protein matrix with the starch granules in the endosperm tissue of Valier barley may limit the access of ruminal micro-organisms to the starch granules and thus reduce the rate and extent of rumen degradation relative to that of Harrington barley. It is the first time that the microstructural matrix in the endosperm of barley has been revealed by using the SR-FTIR technique, which makes it possible to link feed intrinsic structures to nutrient utilisation and digestive behaviour in ruminants. Application 4 showed with the SR-FTIR technique that the chemical features of various feed protein (amide I) secondary structures (such as feather, wheat, oats and barley) could be quantified. With a multi-component fitting program (Lorentz function), the results showed feather containing about 88 % β-sheet and 4 % α-helix, barley containing about 17 % β-sheet and 71 % α-helix; oats containing about 2 % β-sheet and 92 % α-helix; and wheat containing about 42 % β-sheet and 50 % α-helix. The relative percentage of the two may influence protein value. A high percentage of β-sheet may reduce the access of gastrointestinal digestive enzymes to the protein structure. Further study is required on feed protein secondary structures in relation to enzyme accessibility and digestibility. In conclusion, the SR-FTIR technique can be used for feed science and animal nutrition research. However, the main disadvantage of this technique is the requirement for a special light source; a synchrotron beam.

Synchrotron: Infrared microspectroscopy: Chemical imaging and mapping: Feed chemistry: Animal nutrition

Traditional ‘wet’ analytical chemistry usually looks for a specific known component through homogenisation of the tissue and separation of the components of interest from the complex matrix. As a result, the information about the spatial origin and distribution of the component of interest is lost and the object of the analysis is destroyed.

Abbreviations: CHO, carbohydrate; FTIR, Fourier transform infrared; NIR, near infrared; NSLS-BNL, National Synchrotron Light Source in Brookhaven National Laboratory; SRC, Synchrotron Radiation Center; SR-FTIR, synchrotron radiation-based Fourier transform infrared.

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(Budevska, 2002). Traditional ‘wet’ chemical analyses rely heavily on the use of harsh chemicals and derivatisation, therefore altering the native feed structures and possibly generating artifacts. The analyses also require reasonable amounts of feed material (g or mg levels), which usually means that they are carried out on composite samples of tissues (Stewart et al. 1995). Also, traditional ‘wet’ chemical analysis is time consuming, expensive and prone to errors within and between laboratories.

Near infrared, Raman and Fourier transform infrared spectroscopic techniques for feed assessment

In feed analysis and quality assessment, spectroscopic techniques including near IR (NIR), Raman and globar-sourced Fourier transform IR (FTIR) spectroscopy exist. The NIR technique is commonly used. The NIR method is a computerised optical technique, in which absorption of near IR radiation is measured on a set of ground samples and a mathematical relationship is established between the optical data and chemical composition obtained by other methods. The NIR technique has been applied in a variety of fields including ruminant nutrition (Marten, 1989). The first use of the NIR method for the analysis of forage quality was reported by Norris & Barnes (1976). Since then, considerable research has been directed towards using the NIR technique to predict the composition of feedstuffs and faeces, in vivo digestibility, intake etc. The advantages of the NIR method, described by Norris (1989), are speed in analysis, simplicity of operation, and the multiplicity of NIR analyses that can be performed on a given sample. The fatal disadvantages are dependent on an appropriate calibration to the direct measurement of the variable for which the NIR method predicts a value, the lack of adequate theory to support the measurement (Norris, 1988), and measuring a set of ground samples. The measuring of ground samples means that the feed sample’s inherent structures are destroyed and that the spatial origin and distribution of the components of interest are lost. The other two IR spectroscopic techniques (Raman and FTIR spectroscopy) are not common in the feed industry. FTIR spectroscopy has limited flexibility for the analysis of solids. Raman spectroscopy instruments suitable for routine use are only recently becoming available and their practical utility requires study. Again, the fatal disadvantage of the two techniques is that the sample needs to be ground. Therefore NIR, Raman and FTIR spectroscopy are not able to link a feed’s inherent structural information to digestive behaviour and nutrient utilisation or reveal chemical features at cellular and subcellular levels.

Synchrotron-based bioanalytical technique

When a globar-sourced FTIR spectroscopy is combined with a microscope, the technique is called ‘FTIR microspectroscopy’. This technique is able to identify molecular constituents in biological samples from their vibrational spectra in the mid-IR region (Marinkovic et al. 2002). A drawback of FTIR microspectroscopy is resulting diffraction effects if the aperture is decreased to limit the field of view to a small region of interest. At the same time, less light overall reaches the detectors, and hence the signal:noise ratio decreases (Wetzel et al. 1998; Miller et al. 2000; Raab & Martin, 2001; Wetzel, 2001; Bonetta et al. 2002; Miller, 2002). In order to study the structural—chemical features of plant feeds, spectral data at the diffraction limit or a few microns in each spatial dimension are collected; synchrotrons and free electron lasers can achieve this. In contrast, the brightness of conventional bench-top IR sources is simply too low by two to three orders of magnitude (Raab & Martin, 2001; Bonetta et al. 2002).

A synchrotron is a giant particle accelerator that turns electrons into light (Synchrotron facts, 2004). Synchrotron light is extremely bright (millions of times brighter than sunlight). The beam is non-divergent, intense and extremely fine (Wetzel et al. 1998; Miller et al. 2000; Raab & Martin, 2001; Miller, 2002).

When a synchrotron light (IR) source, FTIR spectroscopy and microscopy are combined together, the technique is called ‘synchrotron radiation-based FTIR (SR-FTIR) microspectroscopy’. Recently, the SR-FTIR method has been developed as a rapid, direct, non-destructive and non-invasive bioanalytical technique. This technique, taking advantages of synchrotron light brightness (which is usually 100—1000 times brighter than a conventional globar source and has a small effective source size), is capable of exploring the molecular chemistry within microstructures of biological samples with a high signal:noise ratio at ultraspatial resolutions as fine as 3—10 μm (Miller et al. 1998, 2000; Wetzel et al. 1998; Raab & Martin, 2001; Wetzel, 2001; Bonetta et al. 2002; Marinkovic et al. 2002; Miller, 2002; Vogel et al. 2002; Yu et al. 2003b,c, 2004c). This technique is able to provide information relating to the quantity, composition, structure and distribution of chemical constituents and functional groups in a tissue and can encompass a wider spectral range so that more detailed structural information can be extracted. The technique can be used to increase the fundamental understanding of feed structures at the cellular level and bring a new level of understanding of analytical information (Budevska, 2002; Yu et al. 2003b,c, 2004c).

Objective and applications

To date there has been very little application of this synchrotron-based analytical technique (SR-FTIR) to the study of the structural—chemical features (bonding and functional group characteristics) of feed materials. These structural—chemical features are associated with feed quality and nutritive value in animal diets. The objective of the present article was to review four applications of the synchrotron-based bioanalytical technique (SR-FTIR) for feed science and animal nutrition research. In application 1, using SR-FTIR microspectroscopy, the intensities and distribution of biological components (such as lignin, protein, lipid, structural and non-structural carbohydrates (CHO)) and their ratios in the microstructure of barley tissue within the cellular dimension could be imaged. In application 2, the ultrastructural—chemical makeup and density of yellow- and brown-seeded Brassica rape could be explored and compared. This structural—chemical
information could be used for the prediction of rapeseed quality and nutritive value for man and animals. In application 3, chemical differences in the ultrastructural matrix of endosperm tissue between Harrington (malting-type) and Valier (feed-type) barley in relation to rumen degradation characteristics could be identified. In application 4, chemical features of various feed protein (amide I) secondary structures (such as feather, wheat, oats and barley) could be quantified with a multi-component fitting program (Lorentz function). Finally, the present paper summarises the advantages and disadvantages of the SR-FTIR technique, provides a conclusion and presents implications.

**Principle of synchrotron radiation-based Fourier transform infrared microspectroscopy**

**Advantages of synchrotron radiation-based Fourier transform infrared microspectroscopy**

Using standard globar (conventional thermal IR)-sourced FTIR microspectroscopy cannot reveal the chemical features of micro-biomaterials (<35 to 50 μm, depending on the type of IR microspectrometer). The normal plant feed cell size is about 5–30 μm. With a globar source, a very poor signal:noise ratio within plant cellular dimensions is obtained (Wetzel et al. 1998; Miller et al. 2000; Miller, 2002; Dumas, 2003).

Using SR-FTIR microspectroscopy allows a very small area to be explored; this provides higher accuracy and precision, allows faster data collection, reaches the diffraction limit as a few μm and provides a very good signal:noise ratio within cellular dimensions is obtained (Wetzel et al. 1998; Holman et al. 2002; Marinkovic et al. 2002; Miller, 2002; Dumas, 2003). It can reveal feed structural–chemical features within cellular dimensions (Yu et al. 2003b, 2004b, c, d). Experimentation also shows that the synchrotron IR source is unable to damage any biological tissue.

Detailed comparisons between the conventional globar- and synchrotron-sourced FTIR microspectroscopy have been reported (Wetzel et al. 1998; Miller et al. 2000; Holman et al. 2002; Miller, 2002; Dumas, 2003). Fig. 1 (a) shows that with synchrotron light, a very good signal:noise ratio spectrum at a very small area (5 μm × 5 μm) with high ultra-spatial resolution can be obtained, compared with a conventional globar source (Miller et al. 2000; Miller, 2002). Fig. 1 (c) shows a faster data collection with synchrotron light. It only needed 16 s and thirty-two scans to obtain a very good-quality spectrum at a smaller area (3 μm × 3 μm). However, with a conventional globar source (Fig. 1 (d)), at a larger area of 6 μm × 6 μm, even

**Fig. 1.** (a) Comparison of synchrotron and globar Fourier transform IR (FTIR) microspectroscopy. With synchrotron light, a very good signal:noise ratio spectrum at a very small area (5 μm × 5 μm) with high ultra-spatial resolution is obtained (top) and with a globar source, a very poor signal:noise ratio spectrum is obtained (bottom) (from Miller et al. 2000; Miller, 2002). (b) Visible image. (c and d) Comparison of synchrotron and globar-sourced FTIR microspectroscopy shows a faster data collection with synchrotron light. With synchrotron light (c), only 16 s and thirty-two scans are needed to obtain a very good-quality spectrum at a smaller area (3 μm × 3 μm); with a conventional globar source (d), at a larger area of 6 μm × 6 μm, even when using 1000 scans and 500 s only a very poor quality spectrum can be obtained (from Dumas, 2003).
using 1000 time scans and 500 s, still only a very poor-quality spectrum was obtained (Dumas, 2003).

Principle of synchrotron radiation-based Fourier transform infrared analytical technique

As we know, the total energy of a molecule consists of the translational, rotational, vibrational and electronic energies. Different radiation of the electromagnetic spectrum will result in different energy transitions in a molecule. Organic molecules possess bonds and functional groups. These functional groups vibrate independently of each other and weakly interact as entities. Without any electromagnetic radiation effect, these molecules (functional groups) vibrate independently at their equilibrium position. However, when IR radiation (which is part of the electromagnetic spectrum) is applied to organic molecules (functional groups), it breaks down the molecule’s equilibrium (position) stage, and causes two energy transitions in a molecule. It promotes transitions in a molecule between rotational and vibrational energy. When transitions between rotational and vibrational energy levels occur that cause a net change in the dipole moment, the molecule will absorb IR. Therefore an IR absorption profile is unique to a specific molecular vibration frequency (Jackson & Mantsch, 1996). When IR passes through a sample, some IR is absorbed by the sample and some IR is passed through (transmitted). The resulting spectrum represents the molecular absorption—transmission, which creates a molecular fingerprint of the sample. The identification of molecular functional groups is a major application of IR spectrometry (Hergert, 1971; Joe & Roth, 1986; Mathlouthi & Koenig, 1986; Kemp, 1991; Yu et al. 2003c, 2004c).

Unique infrared bands of biological components in feed tissues

Because biological components have unique molecular chemical—structural features, they have their own unique IR spectrum. The characteristic of protein is unique in the peptide bond. The peptide bond contains C=O, C=N and N–H. The amide I bond is primarily C=O stretching vibration (80%) plus C=N stretching vibration. Amide I absorbs at about 1650 per cm. Amide II, which absorbs at 1550 per cm, consists of N–H bending vibrations (60%) coupled with C–N stretching vibrations (40%). Lipid contains both carbonyl C=O ester as well as CH₃ and CH₂ functional groups. The unique IR spectral profile for lipid is: about 1738 per cm (carbonyl C=O), 1470 per cm (CH bending), 2961 per cm (asymmetric stretch CH₃), 2925 per cm (asymmetric stretch CH₂), 2871 per cm (symmetric stretch CH₂), and 2853 per cm (symmetric stretch CH₃) (Jackson & Mantsch, 1996). CHO have lots of sugars, so lots of OH and CO bonds. Depending on bond linkage and type of sugar, the IR position for CHO is between 1180 and 950 per cm (Mathlouthi & Koenig, 1986) and is attributed to C–O stretching vibrations. The bands in this region are very complicated. When studying plant materials, it is customary to look for structural CHO such as hemicellulose and non-structural CHO such as starch (Wetzel et al. 1998). A peak area at the 1420 per cm band can be used to look for a particular type of CHO; β-glucan (Wetzel et al. 1998). In the portion of the spectrum from 1550 to 800 per cm, strong CHO bands are present for both structural and non-structural CHO, particularly in the region 1100–1025 per cm. A major difference between these two forms of CHO is the presence of bands of moderate intensity at approximately 1420, 1370 and 1335 per cm, which indicates the characteristics of structural CHO (Wetzel et al. 1998; Wetzel, 2001). CHO band peaks between 1100 and 1025 per cm depend on whether the CHO is structural or non-structural. A peak at about 1025 per cm indicates non-structural CHO such as starch in the endosperm of cereal grain (Wetzel et al. 1998; Wetzel, 2001). The same principle applies for other biological components, such as lignin at 1510 per cm, hemicellulose at 1246 per cm, cellulase at 1100 per cm and starch at 1025 per cm (Himmelsbach et al. 1998; Miller et al. 1998, 2000; Wetzel et al. 1998; Wetzel, 2001; Marinkovic et al. 2002; Miller, 2002; Yu et al. 2004c). Detailed discussions on functional group assignments in plant materials have been reported by Wetzel et al. (1998), Raab & Martin (2001), Wetzel (2001) and Yu et al. (2004c,d).

Materials and methodology of synchrotron radiation-based Fourier transform infrared technique

Feed preparation and synchrotron transmission and reflection infrared microspectroscopy

The feed samples were frozen at −20°C on objective disks in a cryostatic microtome and then were cut into thin cross-sections (about 5 to 10 μm thickness). The unstained cross-sections of the feed tissues were rapidly transferred to BaF₂ windows (size 13 × 1 mm disk; Spectral Systems, Hopwell Junction, NY, USA) for transmission mode in synchrotron transmission FTIR microspectroscopic work. The unstained cross-sections were dry-mounted onto Low-c-R microscope slides (Kevely Technologies, Chesterland, OH, USA) for reflectance mode in synchrotron reflection FTIR microspectroscopic work. More detailed methodology on slide preparation has been reported by Yu et al. (2003c, 2004a,b,c,d).

Synchrotron light source

The experiments were performed at the National Synchrotron Light Source in Brookhaven National Laboratory (NSLS-BNL; US Department of Energy, New York, NY, USA) and the Synchrotron Radiation Center (SRC; University of Wisconsin, Madison, WI, USA).

Synchrotron radiation-based Fourier transform infrared microspectroscopy

The spectroscopic images were recorded using a Nicolet Magna 860 FTIR (Thermo Nicolet Instruments, Madison, WI, USA) equipped with a Continuum IR microscope (Spectra Tech, Oak Ridge, TN, USA), mapping stage controller, 32 × objective and a mercury cadmium telluride (MCT-A) detector. The bench was configured with a
Use of synchrotron FTIR microscopy

Collimated synchrotron light beamline (U2B and U10B at the NSLS-BNL; port 031 at the SRC) which served as an external input to the Nicolet Magna 860. The modulated light was passed through the IR microscope to perform transmission or reflection microscopy. The spectra were collected in the mid-IR range of 4000–800 per cm at a resolution of 4 per cm with sixty-four scans co-added and an aperture setting of about 10 μm × 10 μm. The reasons for the chosen aperture size of 10 μm × 10 μm were: (1) the size was within cellular dimensions; (2) the 10 μm × 10 μm aperture size was the best for getting a good signal:noise ratio spectrum mapping of feed tissues. To minimise IR absorption by CO₂ and water vapour in the ambient air, the optics were purged using dry N₂. A background spectroscopic image file was collected from an area free of sample. The mapping steps were equal to aperture size. Stage control, data collection and processing were performed using OMNIC 6.0 (Thermo-Nicolet, Madison, WI, USA). Scanned visible images were obtained using a charge-coupled-device camera linked to the IR images (Yu et al. 2003c, 2004a,b,c).

In the microspectroscopic area mapping, the spatial information was obtained by translating the tissue along the x and y axes, and positioning different parts of the designated tissues in the synchrotron IR beam of the microspectroscope. The motorised computer-control stage was programmed to trace the designated areas in the tissues. After measuring the spectra of all parts of interest, the spectral information was related to the visible images. As a result, spectral datasets were formed with the xy surface, corresponding to the scanned area of the sample, and z direction, which contained the spectral information. Functional group images (such as amide I, aromatic compound, carbonyl ester) were generated by plotting the intensity of the synchrotron IR absorption bands as a function of xy position (Budevska, 2002). Different coverage of the sample with measurements could also be achieved by varying the step size and the dimensions of the image mask (aperture size).

Data analysis and chemical imaging

The spectral data of the feed tissue were collected, corrected with the background spectrum, displayed in the absorbance mode and analysed using OMNIC software 6.0 (Thermo-Nicolet, Madison, WI, USA). An automatic baseline correction was applied to generate the final spectra. The synchrotron FTIR absorbance was expressed as log (1/R) (The unit of the spectrum collected in the reflectance mode needed to be converted to absorbance units by log (1/R)). The data can be displayed either as a series of spectroscopic images collected at individual wavelengths, or a collection of IR spectra obtained at each pixel position in the image. Functional group bands were according to reports (Jackson & Mantsch, 1996; Himmelsbach et al. 1998; Wetzel et al. 1998; Miller et al. 2000; Wetzel, 2001; Miller, 2002). Chemical imaging of functional groups was determined by OMNIC software 6.0 (Thermo-Nicolet, Madison, WI, USA) or by IR spectral imaging software, version. 1.1.01 (Cytospec Inc., Croton-on-Hudson, NY, USA). Peak ratio images were obtained by the height or area under one functional group band (such as amide I; 1650 per cm) divided by the height or area under another functional group band (such as starch; 1025 per cm) at each pixel (pixel size 10 × 10 μm), which represented the biological component ratio intensity and distribution in the feed tissue (for example, protein:total starch ratio image and hemicellulose:total CHO ratio image). The false colour maps were used, which were derived from the area under particular spectral features.

Univariate and multivariate statistical approaches for chemical mapping

Statistical approaches to analyse spectral data collected with SR-FTIR microspectroscopy usually include uni- and multivariate statistical methods. The univariate methods of analysis consist of various mapping displays of spectral data. Usually researchers may select band intensities, integrated intensities, band frequencies, band intensity ratios etc., to construct false colour maps of the spectral data (Miller, 2002). The multivariate methods of data analysis create spectral corrections and maps by including not just one intensity or frequency point of a spectrum, but by utilising the entire spectral information. The methods include cluster analysis (Cytospec, 2004), which is a technique that clusters IR spectra in a map based on similarity with other spectra in the same map (Miller, 2002).

Hierarchical cluster analysis

An agglomerative hierarchical cluster analysis was used to perform a cluster analysis of IR spectra datasets and displayed the results of cluster analysis as images and as dendrograms (Cytospec, 2004) when a biological component band has more or less interference with other bands. For cluster analysis, Ward’s algorithm was used for clustering. Spectral distances were computed as D-values. In a dendrogram, spectral distances were given on the y axis. Horizontal lines illustrated the merging process, for example the combination of two clusters to a new cluster of spectra.

Statistical analysis

The statistical analysis was performed using SAS procedures (SAS Institute, Inc., 1998).

Applications of synchrotron radiation-based infrared microspectroscopy to feed science and animal nutrition

Application 1: chemical imaging of microstructures of feed tissues within cellular dimensions using synchrotron radiation-based transmission Fourier transform infrared microspectroscopy

The objectives of these studies (Yu et al. 2003c, 2004b) were to use this non-invasive bioanalytical technique (SR-FTIR) to chemically image the intrinsic structures of feed tissues at an ultraspacial resolution to reveal the plant-feed structural–chemical information. The experiments were performed at the beamline U10B of the NSLS-BNL (US Department of Energy, New York).

The results showed that with the extremely bright synchrotron light, the spectra with a high signal:noise ratio
were obtained from areas as small as 10 μm × 10 μm in the plant-feed tissue, which allowed us to ‘see’ feed tissue in a chemical sense on a cellular level (Figs. 2, 3 and 4). The ultraspatially resolved imaging of feed tissues by stepping in pixel-sized increments was obtained. Chemical intensities and distributions of biological components such as lignin, cellulose, protein, lipid and total CHO and their ratios could be mapped (Figs. 2, 3 and 4) (Yu et al. 2003c, 2004c). These images revealed the chemical information of feed intrinsic structure.

When a unique band had interference with other bands, cluster analysis gave satisfactory results and was conclusive in showing that it could discriminate and classify functional group differences existing in different structure regions of feed. For example, in maize tissue, spectral data (Fig. 5) show that the band at about 1515 cm⁻¹ has interference with part of the amide II band. A cluster image (Fig. 6) proved that the clusters in the pericarp region and endosperm region are completely different.

In conclusion from the studies (Yu et al. 2003c, 2004b), the SR-FTIR technique can provide chemical and functional characteristics of feed tissue at high ultraspatial resolutions. The SR-FTIR microspectroscopic images can generate spatially localised chemical information within cellular dimensions. This work clearly shows that imaging chemical and functional group distribution across feed tissue can be accomplished using FTIR microspectroscopy powered by synchrotron bright light. Such information could be analysed for clues of the plant intrinsic structures in relation to biodegradation characteristics in animals.

Application 2: ultrastructural–chemical makeup of yellow- and brown-seeded Brassica rape, explored with synchrotron reflection Fourier transform infrared microspectroscopy

*Brassica* rape including the yellow-seeded *Brassica rapa* and the dark-brown-seeded *B. napus* varieties are commonly grown in Western Canada. Yellow-seeded rape

![Fig. 2. Functional-group images of Harrington barley tissue within cellular dimensions from the pericarp (A), seed coat (B), aleurone layer (C) and endosperm (D). (1), Visible image; (2), chemical image; (3), chemical intensity ruler; (4), three-dimensional image; (5), spectra corresponding to the pixel at the cross-hair in the visible image. (a) Area under peaks centred at 1510 cm⁻¹ showing lignin distribution and intensity; (b) area under 1650 cm⁻¹ (amide I) showing protein distribution and intensity; (c) area under the peaks between 1180 and 1000 cm⁻¹ (carbohydrates); (d) area under peaks at 1246 cm⁻¹ indicating cellulosic material distribution and intensity; (e) area under peaks at 1738 cm⁻¹ (C=O ester linkage) showing lipid distribution and intensity. (From Yu et al. 2003c.)
has been reported to be lower in fibre than those of brown-seeded types (Stringam et al. 1974; Slominski et al. 1994; Simbaya et al. 1995). Rapeseed selection programmes have been directed toward increased yellow seed content to decrease fibre content and thereby increase nutritive value (Slominski et al. 1994; Simbaya et al. 1995). Several studies have focused on the total chemical composition of rapeseeds including total protein, CHO and dietary fibre components using traditional ‘wet’ chemical analysis (Slominski et al. 1994; Simbaya et al. 1995). No research has been carried out to study the ultrastructural chemical makeup features (chemical bonding and functional groups) of rapeseed tissue within cellular dimensions. These features are associated with rapeseed quality and nutritive value in man and animals.

The objective of this study (Yu et al. 2004a) was to use synchrotron reflection FTIR microspectroscopy as a non-destructive technique to explore the ultrastructural–chemical makeup (functional groups and bonding characteristics) of yellow- (*B. rapa* cv. Klondike) and brown-seeded (*B. napus* cv. Bounty) rape at an ultraspatial resolution. The investigated chemical bonding and functional groups included the NH$_{\text{+}}$OH group with a broad band at about 3298 per cm, CH groups (CH$_3$ asymmetric stretch at

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**Fig. 3.** Spectra of pericarp (A) and aleurone layer (C) of Harrington barley tissue selected from the corresponding area from the visible images, showing that similar morphological parts exhibit similar spectral characteristics and chemical composition. (B) Seed Coat, (D) endosperm. (From Yu et al. 2003c.)
Fig. 4. (a) Peak area ratio, area under the 1650 per cm band divided by the area under the peaks between 1180 and 1000 per cm at each pixel, representing protein:total carbohydrate ratio in Harrington barley seed. (b) Peak area ratio, area under the 1246 per cm band divided by the area under the peaks between 1180 and 1000 per cm at each pixel, representing hemicellulose:total carbohydrate ratio in Harrington barley seed. (c) Peak height ratio, representing protein:total starch ratio in Valier barley tissue, obtained by the height under the 1650 per cm band divided by the height under the peaks 1025 per cm at each pixel. (1), Chemical image; (2), chemical intensity ruler; (3), three-dimensional image; (4), visible image; (C) (4) shows the visible bright field image of the mapped area outlined by the lined box, from pericarp (A), seed coat (B), aleurone layer (C) and endosperm area (D); (5), spectra corresponding to the pixel at the cross-hair in the visible image. (From Yu et al. 2004c.)
Fig. 5. Spectra of pericarp and endosperm of maize tissue selected from pixels from the visible images, showing that the area under the 1515 per cm peak (aromatic compound) in the pericarp from lignin imaging (see left spectrum) covering part of the amide II band in the endosperm (right spectrum), indicating lignin imaging under the 1515 per cm peak having interference with part of the amide II band.

Fig. 6. Cluster image of lignin clearly showing that the clusters in the pericarp region and endosperm region are different. Cluster analysis is as follows: (1) select spectral analysis region, 1538-94 to 1490-73 per cm; (2) distance method: D-values; (3) cluster method: Ward’s algorithm; (4) number of clusters (three in cluster image).
about 2961 per cm; CH$_3$ asymmetric stretch at about 2925 per cm; CH$_3$ symmetric stretch at about 2873 per cm; CH$_2$ symmetric stretch at about 2851 per cm), amide I at about 1650 per cm, carbonyl C=O ester stretch at about 1740 per cm, and hemicellulose at about 1240 per cm and total CHO at about 1180–900 per cm as well as their ratios (Table 1; Wetzel & LeVine, 1999; Jackson & Mantsch, 1996). The experiment was performed at the U10B station of the NSLS-BNL (US Department of Energy, New York).

The results showed two IR absorbed intensities of CH$_3$ asymmetric stretching of 0·061 and 0.056, CH$_2$ asymmetric stretching of 0·090 and 0·087, CH$_3$ symmetric stretching of 0·052 and 0·048, and CH$_2$ symmetric stretching of 0·045 and 0·043 for the yellow- and brown-seeded rape, respectively (Table 1). The ratios of total CH$_2$:CH$_3$, CH$_3$ asymmetric:CH$_3$ symmetric, CH$_2$ asymmetric:CH$_2$ symmetric and total CH asymmetric:CH symmetric were 1·06 and 1·13, 1·28 and 1·26, 2·90 and 3·08, 1·82 and 1·78, for the yellow- and brown-seeded rape, respectively. There were no differences between the two rapeseed types in the content and ratios of CH groups (CH$_2$ and CH$_3$) of the scanned areas, indicating that lipid chain length and branching is similar between the two seed types. The absorbed intensities of peak area of NH$^+$OH, total CH, carbonyl C=O, amide I, hemicellulose, and total CHO were 101·623 and 87·123, 8·522 and 7·406, 0·386 and 0·414, 39·200 and 31·045, 1·445 and 1·329, 36·450 and 24·199 ($P < 0·01$), for the yellow- and brown-seeded rape, respectively. There were significant differences in amide I and total CHO, indicating different microstructural protein (peptide C=O bonding) and CHO makeup between the two rapeseed types (Tables 2 and 3).

The results also show differences in the ratios of total NH$^+$OH:CHO (2·85 v. 3·84; $P < 0·01$), CH:CHO (0·23 v. 0·32; $P < 0·01$), amide I:NH$^+$OH (0·48 v. 0·37; $P = 0·07$), amide I:hemicellulose ($P = 0·09$), hemicellulose:CHO (0·039 v. 0·059; $P < 0·001$), and CHO:amide I (1·11 v. 0·84; $P = 0·051$) between the yellow- and brown-seeded rape.

| Table 1. Synchrotron-based Fourier transform infrared absorbed peak height of CH functional groups: CH$_3$ asymmetric stretching, CH$_2$ asymmetric stretching, CH$_3$ symmetric stretching, and CH$_2$ symmetric stretching vibrations in the microstructure (10 $\times$ 10 $\mu$m) of yellow- (Brassica rapa) and brown-seeded (B. napus) rape (Least squares means, standard errors and ranges) |
|-----------------|------------------|------------------|------------------|------------------|
| CH functional group (IR absorption units) | CH$_3$ asymmetric stretching | CH$_2$ asymmetric stretching | CH$_3$ symmetric stretching | CH$_2$ symmetric stretching |
| Wave number (per cm) | Mean | SE | Mean | SE | Mean | SE | Mean | SE |
| Yellow-seed rape (B. rapa) | 2958 | 50 | 2928 | 50 | 2873 | 50 | 2851 | 50 |
| $n$ | Mean | 0·061 | 0·004 | 0·090 | 0·015 | 0·052 | 0·004 | 0·045 | 0·012 |
| Range | 0·009 | 0·130 | 0·010 | 0·790 | 0·013 | 0·145 | 0·004 | 0·641 |
| Brown-seed rape (B. napus) | 36 | Mean | 0·056 | 0·005 | 0·087 | 0·018 | 0·048 | 0·005 | 0·043 | 0·014 |
| $n$ | Range | 0·021 | 0·106 | 0·018 | 0·309 | 0·013 | 0·100 | 0·004 | 0·227 |
| Pooled SEM | 0·0046 | 0·0162 | 0·0043 | 0·0132 |
| $P$ value | 0·46 | 0·91 | 0·52 | 0·92 |

| Table 2. Synchrotron-based Fourier transform infrared (FTIR) absorbed peak area and peak height (IR absorption units) of chemical functional groups in the microstructure (10 $\times$ 10 $\mu$m) of yellow- (Brassica rapa) and brown-seeded (B. napus) rape (from Yu et al. 2004a) (Least squares means, standard errors and ranges) |
|----------------|------------------|------------------|------------------|------------------|
| NH$^+$OH | CH | Carbonyl C=O ester |
| FTIR absorbed intensity | Peak area | Peak height | Peak area | Peak height | Peak area | Peak height |
| Approximate wave number (per cm) | Mean | SE | Mean | SE | Mean | SE | Mean | SE | Mean | SE |
| Yellow-seed rape (B. rapa) | 3298 | 6·228 | 0·393 | 0·025 | 8·522 | 0·883 | 0·088 | 0·014 | 1·187 | 44·089 | 0·011 | 0·790 | 0·024 | 0·862 | 0·002 | 0·035 |
| $n$ | Mean | 101·623 | 6·228 | 0·393 | 0·025 | 8·522 | 0·883 | 0·088 | 0·014 | 1·187 | 44·089 | 0·011 | 0·790 | 0·024 | 0·862 | 0·002 | 0·035 |
| Range | 0·021 | 221·703 | 0·025 | 0·890 | 0·025 | 0·890 |
| Brown-seed rape (B. napus) | 36 | Mean | 87·123 | 7·587 | 0·339 | 0·030 | 7·406 | 1·076 | 0·086 | 0·017 | 2·131 | 19·347 | 0·023 | 0·293 | 0·003 | 3·491 | 0·001 | 0·062 |
| $n$ | Range | 29·761 | 167·124 | 0·150 | 0·659 | 2·131 | 19·347 | 0·023 | 0·293 | 0·003 | 3·491 | 0·001 | 0·062 |
| Pooled SEM | 6·9412 | 0·0275 | 0·9842 | 0·0180 | 0·0792 | 0·0019 |
| $P$ value | 0·14 | 0·17 | 0·43 | 0·96 | 0·81 | 0·95 |
Harrington is a malting-type barley variety. As with most characteristics, tissue between Harrington (malting-type) and Valier differences in the ultrastructural matrix of endosperm tissue could be used for the prediction of rapeseed quality and nutritive value for man and animals and for rapeseed breeding programmes for selecting superior varieties of rapeseed for special purposes.

More detailed study is required to define the extent of the differences that exist between the yellow- (B. rapa) and the dark-brown- (B. napus) seeded rape. Such information could be used for the prediction of rapeseed quality and nutritive value for man and animals and for rapeseed breeding programmes for selecting superior varieties of rapeseed for special purposes.

Application 3: use of synchrotron radiation-based Fourier transform infrared microspectroscopy to identify chemical differences in the ultrastructural matrix of endosperm tissue between Harrington (malting-type) and Valier (feed-type) barley in relation to rumen degradation characteristics

Harrington is a malting-type barley variety. As with most barley varieties, its extent and rate of biodegradation in the rumen are high (Yu et al. 2003d). However, if ruminants consume rapidly degradable barley, the incidence of digestive disorders such as bloat and acidosis can increase. This reduces animal performance and can lead to death in acute cases (Nocek & Tamminga, 1991; Givens et al. 1993). Harrington barley has been recently developed as a specialised feed barley for ruminants (Gibson et al. 1994; Bowman et al. 2001). Its extent and rate of rumen degradation are low (Yu et al. 2003d), which should be beneficial to animal performance and health when animals are fed this variety (Yu et al. 2003d). Using two dairy models (Yu et al. 2003a), the predicted nutrient supply to dairy cows between Valier and Harrington barley also differed (Yu et al. 2003d). The hypothesis for this study was that the different degradation characteristics between the two barley varieties may be partly due to differences in the starch–protein matrix of the endosperm tissue. As we know, traditional ‘wet’ chemical analysis cannot detect such biological differences because the inherent seed structures that may inhibit digestion are destroyed during processing for analysis (Yu et al. 2003d). Chemical analysis also showed that the total chemical composition was similar between the two barleys.

The objective of this study was to use the SR-FTIR technique to identify differences in the structural makeup of Harrington and Valier barley endosperm tissues, so as to obtain a better understanding of the intrinsic structure of barley endosperm tissue in relation to rumen degradation characteristics. The experiment was performed at beamline port 031 in the SRC (University of Wisconsin, Madison). Photomicrographs of cross-sections of the endosperm tissue of Harrington and Valier barley are presented in Fig. 7. Harrington has larger starch granules that are less densely packed in the protein matrix, relative to Valier. As with traditional analytical methods, it is difficult to pinpoint exact differences. No specific chemical information on barley micro-endosperm tissue is obtained through such images. The results showed that Harrington had a wider range of starch:protein IR absorbance intensity ratio (1-406 to 10-119 v. 1-419 to 4-274), suggesting that it is more heterogeneous than Valier in endosperm chemical makeup. Valier had a lower starch:protein IR absorbance intensity ratio than Harrington (P<0-05; 4-121 (SE 0-576) v. 2-776 (SE 0-270); Fig. 8), which implies that the starch granules in Valier are more closely associated with the protein matrix. This closer association may prevent the starch granules from being rapidly degraded in the rumen.

The lower starch:protein ratio in the micro-endosperm tissue of Valier barley may provide a clue to its reduced effective degradability and the somewhat slower rate of rumen nutrient degradation observed in the in situ study (Yu et al. 2003d). McAllister et al. (1993), for example, suggested that the protein matrix surrounding starch granules is a major factor responsible for differences in the rate of ruminal starch digestion of cereal grains. Efficient starch digestion by ruminal micro-organisms requires an array of fibrolytic, proteolytic and amylolytic enzymes. The extent to which protein and structural CHO shield underlying starch granules from enzymic attack differs among cereal grains (Wang & McAllister, 2000). The greater association of the protein matrix with the

### Table 3. Synchrotron-based Fourier transform infrared (FTIR) absorbed peak area and peak height of chemical functional groups in the microstructure (10 × 10 μm) of yellow- (Brassica rapa) and brown-seeded (B. napus) rape (IR absorption units)

<table>
<thead>
<tr>
<th>FTIR absorbed intensity</th>
<th>Amide I</th>
<th>Hemicellulose</th>
<th>Carbohydrate</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Peak area</td>
<td>Peak height</td>
<td>Peak area</td>
</tr>
<tr>
<td>Approximate wave number (per cm)</td>
<td>Mean</td>
<td>SE</td>
<td>Mean</td>
</tr>
<tr>
<td>Yellow-seed rape (B. rapa)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>n</td>
<td>50</td>
<td>50</td>
<td>50</td>
</tr>
<tr>
<td>Mean</td>
<td>39.200</td>
<td>2.126</td>
<td>0.532</td>
</tr>
<tr>
<td>Range</td>
<td>4.14</td>
<td>66.933</td>
<td>0.053</td>
</tr>
<tr>
<td>Brown-seed rape (B. napus)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>n</td>
<td>36</td>
<td>36</td>
<td>36</td>
</tr>
<tr>
<td>Mean</td>
<td>31.045</td>
<td>2.591</td>
<td>0.440</td>
</tr>
<tr>
<td>Range</td>
<td>8.799</td>
<td>54.584</td>
<td>0.120</td>
</tr>
<tr>
<td>Pooled SEM</td>
<td>2.3699</td>
<td>0.0353</td>
<td>0.1266</td>
</tr>
<tr>
<td>P value</td>
<td>0.02</td>
<td>0.07</td>
<td>0.52</td>
</tr>
</tbody>
</table>
starch granules in the endosperm tissue of Valier barley may limit the access of ruminal microorganisms to the starch granules and thus reduce the rate and extent of degradation relative to that of Harrington barley. This study (Yu et al. 2004b) indicated that Harrington and Valier barley differ in their chemical-structural endosperm matrix makeup. The lower starch:protein SR-FTIR absorbance intensity ratio for Valier indicated a tighter association between starch and protein in the endosperm. This may partly explain why Valier exhibits a lower extent and rate of rumen degradation than Harrington barley.

Application 4: using synchrotron radiation-based Fourier transform infrared microspectroscopy to reveal chemical features of feed protein secondary structures and their relation to protein values

Protein secondary structures include α-helix and β-sheet (Fig. 9; Carey, 1996; Mantsch & Chapman, 1996; Martin, 2002; Dumas, 2003). The relative percentage of the two may influence protein value. A high percentage of β-sheet may reduce the access of gastrointestinal digestive enzymes to the protein. Reduced accessibility results in poor digestibility and, as a result, low protein value. Feather is widely available as a potential protein supplement in the animal-feed industry. It is high in protein (84 %), but the digestibility is very low (5 %) (Fig. 10). Because of feather’s high protein content, it has been of interest in nutritional studies. Other commonly used protein feeds such as barley, oats and wheat have high protein value in terms of digestibility.

The objective of this study (Yu et al. 2004e) was to reveal the chemical features of feather protein secondary structures within amide I among different feed-protein sources. The results show that the secondary structure of feather protein differed from other feed-protein sources in terms of the line-shape and position of amide I (Fig. 11). The feather protein amide I band showed a peak at about 1630 per cm, which is consistent with the absorption peak of β-sheet protein amide I. However, barley, wheat and oats contain a lower percentage of β-sheet but a higher percentage of α-helix. With multi-component modelling or a fitting program with Lorentz function, the results showed that the secondary structure of feather protein contains a higher percentage of β-sheet. However, barley, wheat and oats contain a lower percentage of β-sheet but a higher percentage of α-helix. With multi-component modelling or a fitting program with Lorentz function, the results showed that the secondary structure of feather protein contains a higher percentage of β-sheet. However, barley, wheat and oats contain a lower percentage of β-sheet but a higher percentage of α-helix. With multi-component modelling or a fitting program with Lorentz function, the results showed that the secondary structure of feather protein contains a higher percentage of β-sheet. However, barley, wheat and oats contain a lower percentage of β-sheet but a higher percentage of α-helix.

The results indicated that ultraspatially resolved imaging of the feed-protein β-sheet:α-helix ratio was possible. Using the SR-FTIR technique, one could distinguish between secondary structures of protein amide I among the different feed-protein sources. The results show that the secondary structure of feather protein differed from other feed-protein sources in terms of the line-shape and position of amide I (Fig. 11). The feather protein amide I band showed a peak at about 1630 per cm which is consistent with the absorption peak of β-sheet protein amide I. However, the barley, oat and wheat protein sources showed a peak at about 1650 per cm, which is consistent with the absorption peak of α-helix protein amide I. These results indicate that the secondary structure of feather protein contains a higher percentage of β-sheet. However, barley, wheat and oats contain a lower percentage of β-sheet but a higher percentage of α-helix. With multi-component modelling or a fitting program with Lorentz function, the results showed that the secondary structure of feather protein contains a higher percentage of β-sheet. However, barley, wheat and oats contain a lower percentage of β-sheet but a higher percentage of α-helix.
Advantages and disadvantage of synchrotron radiation-based infrared microspectroscopy

The big advantage of using a synchrotron beam with FTIR spectroscopy to map the tissues is that diffraction limits and spatial resolution are achievable. SR-FTIR microspectroscopic imaging can generate spatially localised chemical information without the need to homogenise the sample as done in traditional analytical chemistry. It combines chemical and spatial information. The chemical information (qualitative and quantitative analytical results) can be linked to structural information.

The main disadvantage is the requirement for a bright synchrotron beam and special instrumentation and the difficulty in sample preparation for some feed materials.

Conclusions and implications

In conclusion, synchrotron-based IR microspectroscopy can be used as a new tool to reveal ultrastructural–chemical makeup and features within the feed tissue within cellular dimensions, in terms of spectroscopic characteristics, bonding and functional-group characteristics, and biological components ratio characteristics. With the SR-FTIR technique, chemical functional groups can be mapped to indicate chemical composition distribution and intensity.

Fig. 9. (a) Schematics of protein secondary structure (including α-helix and β-sheet) (from Dumas, 2003). (b) Protein structure determination with synchrotron radiation-based Fourier transform infrared technique. Peak positions of amide I are sensitive to the protein secondary structure (α-helix, β-sheet). For α-helix, the peak is at about 1648–1658 cm⁻¹. For β-sheet, the peak is at about 1620–1640 cm⁻¹ (from Mantsch & Chapman, 1996; Martin, 2002).

Fig. 10. Protein contents (%, III) and ruminant total digestibility (%, III) of selected feed-protein sources.
of feed tissues such as lignin, cellulose, protein, lipid and total CHO and their ratios. These images generate spatially localised chemical information and reveal the chemical information of feed intrinsic structure. The main disadvantage is the requirement for a bright synchrotron light source and special instrumentation and the difficulty in the preparation of some feed samples.

The implications from these studies are that, by using the SR-FTIR technique, we can chemically define the intrinsic structure and analyse individual parts with intact feed samples.

The line-shape and position of protein amide I from feather protein (a and b); from Harrington barley protein (c and d); from Superb wheat protein (e); from Derby oat protein (f), indicating different protein secondary structures between the two protein sources (α-helix at 1650 per cm and β-sheet at 1630 per cm).

Fig. 11. The line-shape and position of protein amide I from feather protein (a and b); from Harrington barley protein (c and d); from Superb wheat protein (e); from Derby oat protein (f), indicating different protein secondary structures between the two protein sources (α-helix at 1650 per cm and β-sheet at 1630 per cm).

Fig. 12. Multi-component amide I peaks modelling showing feather (a) containing about 88% β-sheet and 4% α-helix; barley (b) containing about 17% β-sheet and 71% α-helix; oats (c) containing about 2% β-sheet and 92% α-helix; wheat (d) containing about 42% β-sheet and 50% α-helix. The method of multi-component modelling is performed as follows: step 1, using secondary derivative ‘OMNIC’; identify amide I component peak frequencies; step 2, using multi-peak fitting program with Lorentz function in ‘Origin’, quantify the multi-component peaks area in protein amide I region. Lorentz multi-peaks fitting χ² values for feather, barley, oats and wheat were 0-00018 (R² 0-98666), 0-00013 (R² 0-99755), 0-00079 (R² 0-99581), and 0-00002 (R² 0-99744) respectively. (From Yu et al. 2004 e.)
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tissues, and we can compare the tissue (species or varieties) according to spectroscopic characteristics, functional groups, spatial distribution or chemical intensity within cellular dimensions. Also, we can relate feed-tissue intrinsic–chemical structures to feed quality, nutrient utilisation and digestive behaviours in animals. More research is needed on the ultrastructural–chemical makeup of feeds for a better understanding of feed digestive behaviour and nutrient utilisation in animals. The SR-FTIR technique will provide a new opportunity for feed and nutrition research.

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