Molecular Changes of Preclinical Scrapie Can Be Detected by Infrared Spectroscopy

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Infrared (IR) microspectroscopy was used to detect disease-associated molecular changes spatially resolved in cryosections of scrapie-infected tissue of the CNS. The results show that IR spectra can be used for the discrimination between normal and 263K scrapie-infected hamster nervous tissue not only in the terminal stage of the disease but also in early clinical and even in the preclinical stage at 90 d after oral infection. The nuclei of the cranial nerves located in the medulla oblongata were especially well suited for an early detection of the diseased state by IR microspectroscopy. The most prominent molecular changes indicated by the IR spectra were located between 1300 and 1000 cm⁻¹, a region that contains contributions primarily from carbohydrates and the phosphate backbones of nucleic acids but also from membrane constituents.

Key words: Fourier-transform infrared microspectroscopy; scrapie strain 263K; transmissible spongiform encephalopathy; spectral mapping; Syrian hamster; scrapie pathogenesis; medulla oblongata; cerebellum

Transmissible spongiform encephalopathies (TSE), such as scrapie in sheep and goats, bovine spongiform encephalopathy in cattle, and Creutzfeldt–Jakob disease in humans, are a family of fatal neurodegenerative disorders (Prusiner et al., 1998). Many aspects concerning this group of diseases are not yet understood and have been a matter of intense research. As was shown in a number of studies, histological and molecular differences in TSE-affected nervous tissue are manifold, reaching from deposition of pathological prion protein (PrPSc) (Hedge et al., 1999) as the hallmark of the disease to changes in protein expression, changes in the composition of membrane systems (Choi et al., 1998), alterations in gene regulation (Riemer et al., 2000), and changes in processes such as apoptosis (Fairbairn et al., 1994) and loss of cell populations. The interplay of many different biochemical changes accounts for the TSE-specific pathology. Only a few methods can detect changes in many different biomolecules at the same time during one measurement, and even fewer can accomplish this in situ. Among those are the vibrational spectroscopic methods, such as infrared (IR) and Raman spectroscopy. The nature of an IR spectrum of tissue is that of a fingerprint or pattern, revealing specific biochemical information contained in all IR-active molecules. Spatially resolved IR spectroscopy [using microscopes coupled to Fourier-transform IR (FTIR) spectrometers] can be used to produce IR spectral maps that match with histological maps, because each tissue structure possesses a distinct biochemical composition (Lewis et al., 1996; Kidder et al., 1997; Wetzel and LeVine, 1999). Methods of spectral classification can be used to differentiate between various tissue types, using the multidimensional structural information based on the sum of molecules at a specific location in the tissue (Lasch and Naumann, 1998), thereby adding biochemical information to known histological parameters. Here, we report on the use of FTIR microspectroscopy to study molecular alterations associated with TSE infection in sections of hamster brain from three anatomic regions: the dorsal motor nucleus of the vagus nerve (DMNV) with parts of the solitary tract nucleus (SolN), the nucleus of the hypoglossal nerve (HyN), and the interposed cerebellar nucleus (IntN). The DMNV, followed by the SolN, was recently identified to be the first region showing deposition of pathological PrPSc in hamsters orally challenged with scrapie (Beekes et al., 1998) and also in natural scrapie of sheep (van Keulen et al., 2000). To find out at which stage in the disease process spectral changes could be observed, we analyzed IR spectra at 90 d postinfection (d.p.i.), at 120 d.p.i., and in the terminal stage of orally transmitted scrapie in hamsters.

MATERIALS AND METHODS

Sample preparation and histology. All animal experiments were performed in accordance with European and German legal and ethical regulations. Twelve outbred Syrian hamsters (females) were challenged orally with 1–3 × 10⁷ 50% intracerebral lethal doses (LD₅₀,i.c.), corresponding to 1–3 × 10⁵ 50% oral lethal doses (LD₅₀,p.o.) of scrapie strain 263K as described previously (McBride et al., 2001). Twelve mock-infected hamsters of the same age were similarly fed normal brain homogenate and served as controls. At three time points, 90 d.p.i., 120 d.p.i., and in the terminal stage of the disease (150–160 d.p.i., varying between individuals), four infected and four control animals were killed with CO₂. Brains were frozen and stored at ~70°C. Cryosections were cut coronally, starting from the medulla oblongata. Two series, each containing five adjacent sections, were taken (Fig. 1). The plane of the first sequence (referred to as plane 1 in the following text) contained the HyN, the DMNV, and parts of the SolN, whereas a second sequence (plane 2) was cut through the cerebellar nuclei (Fig. 1). In each of the planes, the first and third sections (10 µm in thickness) were stained with 0.1% cresyl fast violet and 0.2% methylene blue, respectively. All second sections (10 µm) were thaw-mounted on 1 mm BaF₂ windows for FTIR microspectroscopy. For confirmation of scrapie pathology, all fourth and fifth sections (5 µm each) were stained for the prion protein with monoclonal antibody (mAb) 3F4 (Kausch et al., 1987) from cell culture (1.4 µg/ml; 1:100) (a kind gift from Dr. Hans Huser, Robert Koch-Institut, Berlin, Germany) and normal mouse serum (Dako Diagnostika, Hamburg, Germany) as controls, respectively, using a procedure adapted from Taraboulos et al. (1992). Biotinylated goat anti-mouse antibody
Figure 1. Coronal planes (schematic) investigated in this study. Plane I contained the hypoglossal nucleus, the DMNV, and parts of the SolN; plane II contained the cerebellar nuclei. The letters indicate the usage of the sections: a, Cresyl fast violet staining; b, FTIR microspectroscopy; c, methylene blue staining; d, mAb 3F4 immunostaining; and e, normal mouse serum immunostaining (control).

(Dako Diagnostika), peroxidase-conjugated Vectastain avidin–biotin complex kit (Vector Laboratories, Burlingame, CA), and 3,3'-diaminobenzidine as a substrate were used for detection of bound mAb 3F4. The immunostained sections were counterstained with hematoxylin.

Data acquisition and processing. An IFS28/B FTIR spectrometer (Bruker, Ettlingen, Germany) coupled to an IR microscope A 590 (Bruker) equipped with a mercury–cadmium–telluride detector, circular apertures, a 15× C assegrain objective, and a motorized stage (permitting the collection of spectra from a predefined grid of spots in a sample) and the Software OPUS 3.01 (Bruker) were used to take spectra in transmission mode from the tissue sections mounted to BaF2. Absorbance spectra were acquired in the spectral range of 4000–700 cm⁻¹ at 6 cm⁻¹ spectral resolution applying Happ–Genzel apodization and a zero filling factor of 4, yielding approximately one data point per wavenumber. The number of coadded interferograms per spectrum was set to 64 for “overview mapping” of larger tissue areas (4–6 mm²) using 100 μm aperture diameter and to 512 scans for detailed mapping measurements using an aperture diameter of 50 μm.

Spectra of poor quality showing too high an absorption of water vapor or peak intensities that were too low or too high were excluded from the data sets as described previously (Kneipp et al., 2000). Chemical maps were reconstructed using an OPUS 3.01 macro to calculate a so-called “protein/lipid ratio” (integrated intensity at 1700–1480 cm⁻¹) and the absorption bands of CH₂ and CH₃ groups (integrated intensity at 2800–2838 cm⁻¹). Both regions were used to calculate the protein/lipid ratio used for IR imaging (Fig. 3).

RESULTS
IR spectral characteristics of brain tissue
Figure 2 displays a typical IR spectrum of the DMNV region of a hamster brain section. Major absorption bands are indicated (see also Table 1). The shaded regions are the amide region (integrated intensity at 1700–1480 cm⁻¹) and the absorption bands of CH₂ and CH₃ groups (integrated intensity at 2800–2838 cm⁻¹). Both regions were used to calculate the protein/lipid ratio used for IR imaging (Fig. 3). For more specific band assignments, see Table 1.
as an example. They were reassembled using the ratio of the integral intensity of the amide bands and of the CH-stretching region of lipids (both indicated in Fig. 2). The nuclei are relatively rich in protein content and therefore can be easily distinguished from the surrounding lipid-rich white matter in the overview images. After confirmation of the position of the nuclei in the IR maps by comparison with the adjacent cresyl fast violet-stained sections and with a brain atlas (Knigge and Joseph, 1968; Frank-

<table>
<thead>
<tr>
<th>Spectral region/frequency</th>
<th>Assignment</th>
</tr>
</thead>
<tbody>
<tr>
<td>3050-2800 cm⁻¹</td>
<td>Dominated by C—H stretching vibrations of fatty acids (mostly of membrane lipids)</td>
</tr>
<tr>
<td>3012 cm⁻¹</td>
<td>==C—H stretching of alkenes</td>
</tr>
<tr>
<td>2959 cm⁻¹</td>
<td>Antisymmetric stretching of &gt;CH₃ groups</td>
</tr>
<tr>
<td>2872 cm⁻¹</td>
<td>Symmetric stretching of &gt;CH₃ groups</td>
</tr>
<tr>
<td>2851 cm⁻¹</td>
<td>Symmetric stretching of &gt;CH₂ groups</td>
</tr>
<tr>
<td>1800-1500 cm⁻¹</td>
<td>Dominated by absorption of proteins</td>
</tr>
<tr>
<td>1738 cm⁻¹</td>
<td>&gt;C==O stretching vibration of ester carbonyl of phospholipids</td>
</tr>
<tr>
<td>1544 cm⁻¹</td>
<td>Amide I band of proteins</td>
</tr>
<tr>
<td>1466 cm⁻¹</td>
<td>C—H bending vibration of &gt;CH₂ groups of lipids, proteins, and nucleic acids</td>
</tr>
<tr>
<td>1390 cm⁻¹</td>
<td>&gt;C==O stretching vibration of COO⁻ groups</td>
</tr>
<tr>
<td>1300-1000 cm⁻¹</td>
<td>“Fingerprint region,” originating from characteristic C—O—P, C—O—C, and C—C vibrations of DNA/RNA, carbohydrates, lipids, and proteins and P==O stretching vibrations of nucleic acids and phospholipids</td>
</tr>
<tr>
<td>1240 cm⁻¹</td>
<td>Antisymmetric P==O stretching vibration</td>
</tr>
<tr>
<td>1080 cm⁻¹</td>
<td>Symmetric P==O stretching vibration</td>
</tr>
</tbody>
</table>

**Figure 3.** IR overview maps based on the protein/lipid ratio in the IR spectra (right) and cresyl fast violet stains of the adjacent tissue sections (left; scale bars, 500 μm) for orientation. Data obtained from one individual are shown as an example. A, Plane 1, containing the nuclei of the solitary tract, the DMNV, and the nucleus of the HypN. B, Plane 2, comprising the cerebellar nuclei and all cerebellar layers. The nuclei in the medulla oblongata (A) and the cerebellum (B) can be distinguished from the surrounding white matter by their high protein content. The insets in the photomicrographs (left) show the tissue area investigated by IR mapping. The grids in the IR maps indicate areas of detailed measurements that were performed for additional investigations.
lin and Paxinos, 1997), measurements at 50 μm spatial resolution were performed for specific nuclei in both coronal planes (see areas indicated in Fig. 3). In the images of plane 1, the HypN could be separated easily from the nuclei located more dorsally, such as the DMNV and SolN. The protein/lipid ratios of the DMNV and parts of the SolN (central and/or intermediate part) were very similar. Multivariate cluster analyses were performed with all spectra from each detailed measurement in planes 1 and 2. New, cluster analysis-based maps were then reconstructed as shown in Figure 4. A comparison with the topology in the
protein/lipid maps (Fig. 3A, grid) revealed that cluster analysis of all spectra obtained from plane 1 for each hamster had produced two major classes with the spectra of the HypN and DMNV/SolN, respectively (Fig. 4A). Furthermore, each cluster contained two spectral subclasses (1, 2 and 3, 4) as displayed in the dendrogram of Figure 4A. Generally, the subclasses could not be assigned to particular histological structures, with the exception of two spectra in class 3 of Figure 4A, which were identified as stemming from the central canal. Hierarchical clustering of the spectra from the detailed measurements in plane 2 (Fig. 4B) showed separation of cerebellar white matter (cluster 1) from those of the adjacent IntN spectra (cluster 2) and also from spectra of other gray matter structures (cluster 3). Spectra from the latter belonged to the cerebellar cortex (stratum granulosum) and to medullar nuclei, such as the vestibular nucleus (Fig. 4B, cluster 3). The results of cluster analyses performed in the same way with the 24 different brain samples were very similar to the examples shown in Figure 4.

To directly compare spectral features in scrapie-affected brains with those of the controls, all single spectra of the DMNV/SolN and of the HypN from the plane 1 data sets and of IntN from the plane 2 data sets were extracted for all 24 individuals. Extraction was performed on the basis of the classification results shown in Figure 4. In this way, spectra from identical structures in healthy and diseased brains could be collated. The numbers of spectra per brain structure and infection stage are listed in Table 2.

**Comparison of average spectra**

In Figure 5A, normalized first derivatives of the average spectra of the DMNV/SolN, HypN, and IntN from terminally diseased and control hamsters are shown in the spectral region between 1300 and 1000 cm\(^{-1}\). First derivatives were calculated to enhance resolution of the spectral bands and to minimize slight baseline variations in the spectra. All peak maxima of the original absorption spectra appear as zero crossings in the first derivatives. It should be noted that the SDs in the groups of control animals were smaller than in those of the infected animals. The spectral region shown in Figure 5A contains a number of distinct spectral variations of the infected tissue. Differences in the spectra can be found at characteristic positions between 1300 and 1000 cm\(^{-1}\), where various types of molecules, such as carbohydrates, lipids, and nucleic acids, contribute to the spectral profile (Fig. 5A). The spectral alterations are caused by changes in the absorption bands from a number of different functional groups present in these molecules. Between 1200 and 1000 cm\(^{-1}\), spectral characteristics are dominated by \(\text{C}---\text{O}---\text{P}\) and \(\text{C}---\text{O}---\text{C}\) stretching vibrations of carbohydrates and lipids. For example, the differences at 1170 cm\(^{-1}\) (antisymmetric \(\text{C}---\text{O}---\text{C}\) stretching vibration) hint at compositional and structural changes of molecules containing \(\text{C}---\text{O}---\text{C}\) functional groups. Changes of the absorbance band resulting from the symmetric \(\text{P}---\text{O}\) stretching vibration of \(\text{PO}_3^-\) groups of nucleic acids and phospholipids located at \(\sim1080\) cm\(^{-1}\) (Liquier and Taillandier, 1996) were also observed. At \(\sim1240\) cm\(^{-1}\), the width of the peak of the antisymmetric \(\text{P}---\text{O}\) stretching vibration is slightly diminished in the DMNV/SolN and HypN of the infected samples. The prominent changes in band shape at \(\sim1060\) and 1040 cm\(^{-1}\) are probably caused by alterations in carbohydrates, because absorption bands at these frequencies can be assigned to complex sugar ring vibrations of these molecules (Parker, 1983). In the average spectra of the IntN, the spectral differences between infected and control hamsters are almost confined to this band (Fig. 5A).

Figure 5B displays spectral differences between infected and control brains for average spectra of the DMNV/SolN at the

**Table 2. Numbers of spectra per brain structure for different stages of infection as obtained from the data sets by cluster analysis from normal and infected animals**

<table>
<thead>
<tr>
<th>Brain structure (nucleus)</th>
<th>DMNV/SolN N</th>
<th>DMNV/SolN S</th>
<th>HypN N</th>
<th>HypN S</th>
<th>IntN N</th>
<th>IntN S</th>
</tr>
</thead>
<tbody>
<tr>
<td>Terminal stage</td>
<td>178</td>
<td>229</td>
<td>223</td>
<td>210</td>
<td>364</td>
<td>362</td>
</tr>
<tr>
<td>120 d.p.i.</td>
<td>107</td>
<td>119</td>
<td>122</td>
<td>184</td>
<td>335</td>
<td>319</td>
</tr>
<tr>
<td>90 d.p.i.</td>
<td>94</td>
<td>122</td>
<td>233</td>
<td>228</td>
<td>397</td>
<td>403</td>
</tr>
</tbody>
</table>

N, Normal; S, infected.
three different disease stages. In the control groups, the spectra of the DMNV/SolN are almost identical at all three stages (Fig. 5B, solid lines with light SD). In contrast, the average spectra of the diseased samples (Fig. 5B, dotted lines with dark SD) differ systematically between the investigated time points after infection. Variations in the spectra obtained from scrapie-infected DMNV/SolN are most prominent in the terminal stage. In the region between 1060 and 1040 cm⁻¹, the differences between healthy and infected sample spectra can be traced back even to 90 d.p.i. (Fig. 5B). They are small at 90 d.p.i. but clearly progress toward the terminal stage until a pronounced shoulder appears at ~1050 cm⁻¹.

Hierarchical clustering of spectra from diseased and control tissue

For the objective comparison of structure-specific spectra from each investigated individual, a multivariate method was used. Average spectra for each nucleus and each individual in the terminal stage of scrapie and the controls were subjected to cluster analysis, using the information contained in the frequency range 1480–950 cm⁻¹. Figure 6A shows the result of this cluster analysis as a dendrogram. A clear separation of the different nuclei (DMNV/SolN, HypN, and IntN) can be observed (Fig. 6A). Within these structure-specific classes, spectra from infected and control tissue are separated. One HypN spectrum from the infected individuals was misclassified and grouped together with the HypN spectra from the controls. The disease-induced spectral differences in the DMNV/SolN at the earlier disease stages (120 and 90 d.p.i.) were also investigated on the level of structure-specific average spectra of the individuals. Figure 6B shows the result of a cluster analysis of DMNV/SolN spectra of the infected animals at 90 and 120 d.p.i. and the corresponding controls. The spectra were normalized over the range 1300–1000 cm⁻¹, and the region 1050–1025 cm⁻¹ was used as input for cluster analysis. All spectra from the mock-infected controls of both stages appear as one group in the dendrogram, whereas another group is formed by the spectra from the infected animals. The spectra of 90 d.p.i. appear as one subgroup that also contains the spectrum of one of the 120 d.p.i. hamsters.

ANN classification of single spectra from diseased and control tissue

As a classification approach for single-point spectra, ANN analysis was applied to all mapping data sets. A feature extraction method based on univariate F values was used to identify spectral features that differed significantly between spectra of diseased and control DMNV/SolN, HypN, and IntN at each of the three infection stages. On the basis of these features, networks were trained and used for identification of independent test spectra as described in Materials and Methods. The classification result for each spectrum was compared with the scrapie/control status of the corresponding tissue sample. The numbers of correctly classified test spectra were used to determine identification accuracy (Table 3). The numbers of selected data points for network training differed for the structures and infection stages and are also given in Table 3. Table 3 shows an increase of the identification accuracy of the networks with incubation time. At 90 d.p.i., the majority of single spectra from all investigated structures were still classified correctly by the ANN analyses.

PrPSc immunocytochemistry

Cryosections adjacent to those investigated by FTIR microspectroscopy were stained with mAb 3F4, which binds to the prion protein. As was shown previously, the pathological isoform PrPSc can be distinguished from the cellular isoform PrPC by its morphological appearance (Beeke et al., 1998; McBride et al., 2001). The amount of PrPSc detected at each time point varied between individuals. At the terminal stage, massive deposits of PrPSc were found for the DMNV/SolN (Fig. 7A), HypN, and IntN. No PrPSc was detected in specimens from normal control animals (Fig. 7B). At 120 d.p.i., the accumulations were moderate in all structures and could be distinguished as single granules in the DMNV/SolN, IntN (Fig. 7C), and HypN (Fig. 7D). Deposition was more pronounced in the DMNV/SolN than in the HypN and IntN. At 90 d.p.i., PrPSc was undetected in the IntN and HypN. Some single neurons in the DMNV/SolN displayed small granular accumulations of the protein along their surface at this stage (Fig. 7E).
Table 3. Identification accuracy (in %) of ANN for spectra from three brain structures at three time points after inoculation

<table>
<thead>
<tr>
<th>Brain structure (nucleus)</th>
<th>DMNV/SoIN</th>
<th>HypN</th>
<th>IntN</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>N</td>
<td>S</td>
<td>N</td>
</tr>
<tr>
<td>Terminal stage</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>100%</td>
<td>100%</td>
<td></td>
<td>100%</td>
</tr>
<tr>
<td>120 d.p.i.</td>
<td>93.5%</td>
<td>95.8%</td>
<td>81.1%</td>
</tr>
<tr>
<td>90 d.p.i.</td>
<td>70.2%</td>
<td>77.0%</td>
<td>80.3%</td>
</tr>
</tbody>
</table>

The results are shown separately for spectra of control (N) and infected (S) animals. The numbers in brackets indicate the numbers of input data points used by the corresponding ANN.

**DISCUSSION**

The results of this work demonstrate that changes in the IR spectra of the DMNV/SoIN, HypN, and IntN can be used to distinguish scrapie-infected from uninfected tissue. The procedure applied initially included identification of structure-specific spectra from various nuclei in the brain by cluster analysis and IR imaging, followed by extraction of these spectra for comparative purposes. The necessity of collating identical brain structures when comparing the spectral characteristics of normal tissue with those of disease-affected tissue can be illustrated by the comparison of average spectra from infected and control animals (Fig. 5). As proven by cluster analysis, the disease-related spectral variations were indeed smaller than the spectral differences between distinct histological structures (Fig. 6A).

Because one or more absorption bands correspond to a functional group characteristic for a class of molecules rather than for one specific compound, and because spectral variations occur in a number of different absorption bands, it can be concluded that superposition of multimolecular information is the basis of the disease-specific spectral changes. On the molecular level, the observed spectral differences are consistent with phenotypic features found, for example, in ultrastructural studies that occur in early stages of the disease (Liberksi et al., 1989; Jeffrey et al., 1995). These are, for example, microvacuolation, membrane proliferation, and structural and functional damage of mitochondria (Choi et al., 1998), in which changes in the content of phospholipids take place, as was suggested in this study by variations of the antisymmetric and symmetric P=O stretching vibrations of the polar head groups of the lipid and altered absorption of C—O—C stretching vibrations. Changes in absorption bands containing contributions from nucleic acids (ribose skeletal C—O—C and C—C stretching vibrations and P=O stretching vibrations) may also indicate changes in the DNA/RNA ratio caused, for example, by DNA decomposition during apoptosis (Fairbairn et al., 1994; Lucassen et al., 1995) and/or a changed RNA content as the result of upregulation or downregulation of genes (Diedrich et al., 1993; Riemer et al., 2000).

The first change of spectral features that could be observed for diseased DMNV/SoIN at 90 d.p.i. and that remained prominent until the terminal stage was a change in band shape between 1060 and 1040 cm⁻¹ (Fig. 5B). This indicates an altered absorption of ring vibrations of carbohydrates that could be assigned to the sugar moieties of nucleic acids, to changed content of metabolic sugar molecules in the cells, such as glucose, or to other events that have not yet been described. Glucose metabolism is known to change in intracerebrally infected hamsters (Gregoire et al., 1983) and in fatal familial insomnia, a human TSE (Cortelli et al., 1997). Little is known about the extent of these effects and which of them prevail. The above-mentioned spectral change at ~1050 cm⁻¹ is one of the features that was common to all investigated brain structures. However, this change was not equally expressed in all nuclei (Fig. 5A). These qualitative differences of spectral changes occurring in all structures, as well as qualitative absorbance differences (pointing to different molecular events taking place), were observed between the different nuclei. These findings are also reflected by the different heterogeneities between infected and control tissue spectra within each tissue structure in the dendrogram of Figure 6A.

The findings that spectral differences are clearly progressing (i.e., becoming more and more prominent) during the course of the disease in the average spectra (Fig. 5B) and that identification accuracy of ANN analyses increases with incubation time (Table 3) suggest that in the data sets obtained from the infected animals in earlier stages, a number of misclassified single spectra did not exhibit spectral features sufficient to make them distinguishable from those of control animals. This is similar to the well known spread of the histopathological features in scrapie, such as PrPSc deposition. Early in pathogenesis, this pathological form of the prion protein in the DMNV is confined to individual neurons (van Keulen et al., 2000), whereas in the terminal stage, it can be detected in and around almost every neuron of a nucleus. Disease-specific features in the IR spectra were very pronounced in the terminal stage, so that almost every point spectrum obtained from a nucleus during a mapping measurement (i.e., every spot in the mapped area) was classified correctly as “diseased” by the method applied in this study (Table 3). As could be shown by ANN analysis, even at 90 d.p.i., most of the spectra obtained from the DMNV/SoIN, HypN, and IntN still exhibited disease-specific IR spectral features.

When average spectra were calculated for each group of infected and control animals, the SDs of the spectra were smaller for the control animals than for the scrapie-infected animals. This finding indicates that individual parameters seem to determine how rapidly and to what extent molecular changes have developed at a certain stage of the disease. Assuming that biochemical changes begin to be visible in the IR spectra of a specific cell type, it would be interesting to identify these histological substructures or cell types. Systematic comparisons of the spectra on the level of cell populations within the nuclei could then facilitate the detection of scrapie-specific features (such as events confined to neurons or proliferation processes in astrocytes) (Eklund et al., 1963) early in pathogenesis in situ without using immunocytochemistry but by comparing spectra of the identical cell populations of the DMNV/SoIN in infected and control brains.
reconstruction based on the results of cluster analyses revealed that the two subclusters within the two major spectral classes, DMNV/SolN and HypN (Fig. 4A), correspond to relatively small spots (comprising only a few spectra) that are equally distributed in the two nuclei. What is the histological basis of these subclusters? This question can only be addressed by acquiring spectra from single cells in scrapie-affected tissue with high spatial resolution by applying an IR synchrotron source (Jamin et al., 1998).

From histological studies of the 263K scrapie model in hamsters, it is known that the DMNV and the SolN are the first regions in the brain in which an orally induced scrapie infection can be detected by immunostaining of PrPSc deposits in paraffin sections. Much later in the disease process, other nuclei, among them HypN and IntN, are affected (Beekes et al., 1998). The results of our study match these observations: The comparison of average spectra from the DMNV/SolN in scrapie-infected and control animals revealed differences already at 90 d.p.i., whereas changes in the HypN and IntN could not be detected earlier than 120 d.p.i. (data not shown). Furthermore, differences between average spectra in the terminal stage were more pronounced in the DMNV/SolN than in the HypN and IntN (Fig. 5A). These findings are also in good accordance with results from immunostaining of the adjacent cryosections in the terminal stage and at 120 d.p.i. and with studies at early disease stages (McBride et al., 2001), when PrPSc deposition was always heavier in the DMNV and SolN than in the HypN. The extent and local detectability of the IR spectral variations (Fig. 5) obviously concur with the known sequence of PrPSc deposition. Thus, IR spectral changes provide a new biophysical parameter based on molecular markers that indicate the spread of scrapie pathology in the brain starting in the DMNV and SolN.

It is important to note that in contrast to immunocytochemistry, early detection of TSE by IR spectroscopy is not based on the spectral features of PrPSc but rather on changes in a number of different molecules. The spectral data indicate that a disease-specific change of carbohydrates, nucleic acids, and membrane constituents exists early in pathogenesis. The fingerprint-like nature of the scrapie-induced spectral variations greatly diminishes the probability of an identical multivariate pattern being observed in a different disease, such as pseudorabies, herpes simplex type 1, or reovirus serotype 3 (isolate T3C9) infections, which were shown to affect the DMNV in specific animal models (Card et al., 1990; Krinke and Dietrich, 1990; Morrison et al., 1991). However, to further assess the specificity of the IR method, other infections of the CNS of nonexperimental donors will have to be investigated in future studies. The potential sensitivity of the IR method compared with that of very sensitive PrPSc staining methods, especially with that of the paraffin-embedded tissue blot (Schultz-Schaeffer et al., 2000), can be discussed only when FTIR spectroscopic analysis on the single-cell level (as was discussed above) is established. The advantage of the IR spectroscopic method is that only frozen sections without any fixation or staining are required. Fast computerized methods can be used for identification of the diseased tissue, providing a method for the investigation of TSE pathogenesis that can also be developed into a rapid postmortem diagnostic screening method. Because TSE-related cell or tissue damage is most prominent in the brain and spinal cord, we focused on the IR investigation of samples from the CNS. Future IR experiments will also address peripheral organs involved in TSE pathogenesis, such as the peripheral nervous system and the lymphoreticular

Figure 7. PrPSc immunolabeling in sections adjacent to those investigated by FTIR microspectroscopy. PrPSc characteristically presents as granular accumulations of varying sizes. A. Massive PrPSc deposition in the DMNV/SolN at the terminal stage. B. The DMNV/SolN of a mock-infected control. C. An IntN at 120 d.p.i. D. A HypN at 120 d.p.i. E. A DMNV/SolN at 90 d.p.i. Arrowheads point to islets of granular accumulations. Scale bars: A–C, 40 μm; D, E, 10 μm.
system, along with samples suited for in vivo testing, such as blood or CSF.

REFERENCES


