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Rapid techniques for virus identification are more relevant today than ever. The conventional virus detection and identification strategies generally rest upon various microbiological methods and genomic approaches, which are not suited for the analysis of single virus particles. In contrast, the highly sensitive spectroscopic technique tip-enhanced Raman spectroscopy (TERS) allows a characterisation of biological nano-structures like virions on a single-particle level. In this study, the feasibility of TERS in combination with chemometrics to discriminate two pathogenic viruses, Varicella-zoster virus (VZV) and Porcine teschovirus (PTV), was investigated. In a first step, chemometric methods transformed the spectral data in such a way that a rapid visual discrimination of the two examined viruses was enabled. In a further step, these methods were utilised to perform an automatic quality rating of the measured spectra. Spectra that passed this test were eventually used to calculate a classification model, through which also a successful discrimination of the two viral species based on TERS spectra of single virus particles was realised with a classification accuracy of 91%.

1 Introduction

A quick virus identification, surveillance and diagnostics plays an increasingly important role in clinical medicine and public health, considering the growing globalization of travel and trade, bushmeat trafficking, climate change, as well as other factors that enhance the emergence and re-emergence of pathogenic viruses. The annually recurring worldwide flu seasons caused by the influenza viruses or the geographically limited, nevertheless devastating outbreak of the Ebola virus disease in West Africa in 2014 clearly exemplify the importance of a global-scale virus diagnosis and surveillance. In order to optimally counteract viral infections by means of fast clinical actions in infection control, prophylaxis and tailored antiviral treatments, the diagnosis of viruses should be performed as fast as possible.

Time-consuming and labour-intensive culture-based methods for virus isolation once were the mainstream in clinical virology, but are nowadays complemented by serological and genomic approaches like oligonucleotide microarrays, multiplex PCR and sequencing. Especially the advent of real-time monitoring of amplified PCR products can deliver rapid results, even within one hour. Additionally, virus isolation is still required to adapt primers and probes on a regular basis to highly variable viruses, e.g., in case of influenza surveillance. Reagents may not be available for all viruses; when they grow poorly or not at all in vitro systems, the generation of sufficient material to produce commercial test kits is complicated. Also, molecular or antigen-based testing would probably miss one of the agents in cases of dual infection.

Conversely, organism-specific reagents for the recognition of the pathogenic agents are not needed for direct methods like electron microscopy (EM). However, EM is often not able to identify a virus beyond the family level and requires lengthy and complex procedures for fixation and staining of the samples. Often, a complete dehydration is necessary. Another direct microscopy method applied for the analysis of viruses is atomic force microscopy (AFM). The advantages of AFM compared to EM are the simple sample preparation and the direct access to three-dimensional information without approximations via reconstruction. When combined with enhanced Raman spectroscopy, i.e., tip-enhanced Raman spectroscopy (TERS), not only topographic but also spectroscopic information of the virus at nanometre resolution can be obtained. Contrary to super-resolution microscopic methods like stimulated emission depletion (STED) microscopy or photoacti-
viation localisation microscopy (PALM). TERS requires no
target-specific fluorophores. A further advantage of TERS
(especially in view of samples with low viral particle concentra-
tion) is the intrinsically high lateral resolution of the AFM,
which allows a single-virus particle investigation.

In this work, the possibility for a fast virus detection and
identification of single virus particles by means of TERS is
presented. The Varicella-zoster virus (VZV), causing vario-
cella and shingles in humans, and the Porcine teschovirus
(PTV) were selected as model viruses. The discrimination of
the virus on the basis of TERS spectra takes place by means
of chemometric methods. In the first part, this study illus-
trates the possibility to visually discriminate the two exam-
ined species based on the spectral signal content. Therefore,
chemometric methods were used to emphasize the significant
differences for a fast visual decision. In the second part, the
developed method for extracting the crucial signal content is
adjusted and used for an automatic quality rating of the spec-
tra. Furthermore, the positively rated spectra were used to de-
velop a classification model using multivariate data analysis.
Both strategies allow a fast and successful species discrimina-
tion. To our knowledge, this is the first study coping with the
discrimination of virus species that is solely based on TERS
spectra combined with multivariate data analysis.

2 Experimental

2.1 Sample preparation

Varicella-zoster virus (VZV) strain vOka (family Herpesvi-
dae) was amplified in human embryonic lung fibroblasts
(HEL). HELFs were cultured in Eagle minimal essential
medium (EMEM) complemented with 25 mM HEPES, 1 %
non-essential amino acids, 2 mM L-glutamin, 1 mg/100 ml
Ciprofloxacin and 10 % fetal bovine serum at 37 °C and 1 %
CO₂. Supernatants of infected cells showing complete lysis
were centrifuged for 15 min at 4000 rpm (3774 × g; Variofuge
3.0R) to remove cell detritus and were inactivated by UV-
irradiation (20 minutes). The inactivated viruses were diluted
serially in twofold steps with phosphate-buffered saline and
stored at -70 °C.

The Porcine teschovirus (PTV) type 1 strain Talfan (family
Picornaviridae) was amplified in porcine kidney cells PK-15
cultured in EMEM. For PTV preparation, PK-15 monolayers
in T75 Roux flasks were infected and incubated for approxi-
ately 48 hrs until complete lysis. Then, supernatant was cen-
trifuged at 4000 rpm (3774 × g; Variofuge 3.0R) to remove de-
tritus and further ultracentrifuged for 90 minutes at 100000 × g
(30000 rpm, rotor SW60T) to pellet PTV. The virus pellet was
resuspended in 100 μl PBS and inactivated by UV-irradiation
(20 minutes). Two-fold serial dilutions with PBS were stored
at -70 °C.

The resulting virus solutions of VZV and PTV were diluted
with sterile filtered, deionized water in a ratio of 1:10 and
1:100, respectively. A 18 × 18 mm glass cover slip, which
was cleaned with a mixture of concentrated nitric acid and
hydrogen peroxide (both from Roth) and stored under argon
atmosphere served as substrate. On these glass slides, 10 μl
of virus solution were dropped and dried under ambient condi-
tions. To remove residues of the companion matrix, the sam-
ple was washed with 100 μl deionized, sterile filtered water.
After a further drying step, the virus particles adsorbed on the
glass substrates and were ready for TERS measurement.

2.2 TERS setup

All TERS measurements were carried out with a transmis-
sion setup using a 180° back-scattering geometry. The TERS
setup consisted of a Raman spectroscopy (HR LabRam, Jobin
Yvon Horiba) coupled to an AFM (Nano Wizard, JPK In-
strument AG). The glass slide with the adsorbed virions
was fixed in the sample holder and illuminated from be-
low. An oil immersion objective with a magnification of 60 ×
(PlanApo, Olympus, NA 1.45) was used to focus the exci-
tation laser on the sample and the AFM tip. As TERS tips,
conventional AFM probes (NT-MDT, type: NSG10, average
ωResonance: 270 kHz, rTip: 6-10 nm), suitable for non-contact
and intermittent mode, were coated with 25 nm silver and
stored under argon atmosphere until usage. A krypton ion
laser (Innova 300c, U.S.A.) with a wavelength of 530.9 nm
and a power of 1 mW on the sample was used as excitation
and coupled to the inverse microscope (IX70, Olympus).

Sample scanning was done by a closed-loop XY piezo scan-
ning stage (P-734, Physik Instrumente, Germany). To com-
pensate for the z-tip movements the microscope objective was
synchronized via a z-pezo stage (PIFOC, Physik Instrumente,
Germany) with the tip vertical movement. The setup allows
to position the AFM tip with an accuracy of 0.5 nm. After
the scanning of the virion’s surface, distinct points in form of a
grid with a pre-defined dimension were selected. Then, tip
and sample were irradiated by the laser and from each point of
the grid, a TERS spectrum was recorded with an acquisition
time of 10 s.

2.3 Chemometric analysis

All data used in this study underwent an adequate prepro-
cessing that was implemented using the free software environ-
ment R. First, the measured spectra were interpolated and at
the same time truncated to a wavenumber region from 2000 to
400 cm⁻¹. After that, the data was normalised using the Sili-
con band at 520 cm⁻¹. This signal originates from the used tip
and is treated as internal standard. Therefore, it allows scaling
of the recorded TERS spectra. In a third step, the background
for each spectrum was estimated and then removed applying the sensitive non-linear iterative peak-clipping (SNIP) algorithm\textsuperscript{18}. All spectra were exposed to the same treatment.

After the preprocessing, the data is further evaluated in two different ways: For the visual virus discrimination, the data set was reduced by the low quality spectra based on a manual rating. For the automated discrimination based on multivariate data analysis, all spectra underwent an automatic quality rating explained below. After the rating, the automatically recognised low quality spectra were discarded and removed from the data set.

2.3.1 Manual quality rating The manually quality rated and reduced data set was then used to create the so called component maps for the visual discrimination of the examined virus particles. Details about the structure of these maps are given in the next section. In the maps, the presence or absence of characteristic signals (given in Tab. 1) is represented. To determine whether a certain spectrum showed a given signal, a peak was fitted around the expected wavenumber position in a spectrum. A criterion, the peak decision value \( x \), was set to decide, whether a peak counted as peak, or not.

\[
x(a, b) = m(a, b) + 1.5 \quad \text{if } I > x(a, b) \Rightarrow \text{Peak} \\
\text{if } I \leq x(a, b) \Rightarrow \text{no Peak}
\]

Equation 1 shows, how the decision value \( x \) for a subsection of a spectrum from wavenumber \( a \) to wavenumber \( b \) was determined. Thereby, \( m(a, b) \) denotes the arithmetic mean of the intensity values of a subsection of a spectrum from \( a \) to \( b \) inverse centimetres. A peak was recognized as valid signal if its intensity \( I \) exceeded \( x \). Fig. 1 illustrates this decision process. For every signal given in Tab. 1, an attempt was made to fit a peak in the range \([a, b]\) around the expected wavenumber position. The peak was not only fitted at its expected position but also in an individual allowed range to compensate for possible wavenumber shifts. In Tab. 1, the first column gives the expected wavenumber position for a peak as well as the allowed deviation. The range \([a, b]\) is derived from this deviation, i.e.,

\[
a = \text{expected wavenumber position} + \text{allowed deviation} \\
b = \text{expected wavenumber position} - \text{allowed deviation}
\]

If a peak was successfully fitted, its intensity value had to be higher than the decision value \( x \) for the considered range \([a, b]\). If the intensity value was higher than 1.5 times the mean intensity value over the examined wavenumber range, i.e., the decision value \( x \), the peak was considered as present. For each spectrum that is part of the analysis, the number of present signals was counted. Thereby, protein and lipid signals were counted separately to create a lipid and a protein map for each measured virus particle. This set of component maps was then used for a fast visual discrimination of the virus particles.

2.3.2 Automatic quality rating The above presented method determining the presence or absence of characteristic signals in a spectrum was adapted for an automatic quality rating of all measured spectra. For each spectrum, the number of present signals was determined without a distinction between lipid and protein signals. If a spectrum shows at least one of the expected protein and lipid signals, it was considered for further analysis. The method allows an objective and fast quality rating for the measured data.

2.3.3 Classification The automatically rated and sorted data set was used to train a classification model that is based on a Linear Discriminant Analysis (LDA). For this task, the R\textsuperscript{17} package “MASS”\textsuperscript{19} was used. The LDA is the method of choice for this study as it is fast and its power and suitability was proven in other experiments\textsuperscript{20,21}. The different sizes of the two virus classes was dealt with accordingly during the training process. Furtheron, a Leave-one-out cross-validation (LOOCV) was conducted to determine the quality of the trained model. The same analysis was performed for the same data undergoing a manual quality rating to compare both approaches and evaluate the advantages of an automatic quality rating.
Table 1 Band assignment of chosen signals

<table>
<thead>
<tr>
<th>Wavenumber [cm(^{-1})]</th>
<th>assigned to</th>
<th>ref.</th>
</tr>
</thead>
<tbody>
<tr>
<td>720 ± 5</td>
<td>Phosphatidyl-choline(^a)</td>
<td>22-24</td>
</tr>
<tr>
<td>870 ± 5</td>
<td>Phosphatidyl-choline(^a)</td>
<td>23.25</td>
</tr>
<tr>
<td>950 ± 5</td>
<td>Phosphatidyl-choline(^a)</td>
<td>23.26</td>
</tr>
<tr>
<td>1735 ± 15</td>
<td>Lipid ester</td>
<td>25.27,28</td>
</tr>
<tr>
<td>830 ± 10</td>
<td>Phospholipid</td>
<td>22.25,27</td>
</tr>
<tr>
<td>1085 ± 10</td>
<td>Phospholipid</td>
<td>22.25,27</td>
</tr>
<tr>
<td>1085 ± 10</td>
<td>Phospholipid</td>
<td>22.25,27</td>
</tr>
<tr>
<td>1656 ± 24</td>
<td>Amide I</td>
<td>29</td>
</tr>
<tr>
<td>1315 ± 15</td>
<td>Amide III</td>
<td>29</td>
</tr>
<tr>
<td>655 ± 5</td>
<td>Cysteine</td>
<td>30–33</td>
</tr>
<tr>
<td>675 ± 5</td>
<td>Cysteine</td>
<td>30–33</td>
</tr>
<tr>
<td>690 ± 5</td>
<td>Cysteine/ Methionine</td>
<td>32,34</td>
</tr>
<tr>
<td>1180 ± 5</td>
<td>Histidine(^b)</td>
<td>31</td>
</tr>
<tr>
<td>1330 ± 5</td>
<td>Histidine(^b)</td>
<td>31</td>
</tr>
<tr>
<td>1500 ± 5</td>
<td>Histidine(^b)</td>
<td>31</td>
</tr>
<tr>
<td>1004 ± 5</td>
<td>Phenylalanine</td>
<td>31,33–36</td>
</tr>
<tr>
<td>1033 ± 5</td>
<td>Phenylalanine</td>
<td>31,33–36</td>
</tr>
<tr>
<td>1210 ± 5</td>
<td>Phenylalanine</td>
<td>31,33–36</td>
</tr>
<tr>
<td>902 ± 5</td>
<td>Proline</td>
<td>32</td>
</tr>
<tr>
<td>1015 ± 5</td>
<td>Tryptophan</td>
<td>31,33,36</td>
</tr>
<tr>
<td>1043 ± 5</td>
<td>Tryptophan</td>
<td>31,33,36</td>
</tr>
<tr>
<td>1340 ± 5</td>
<td>Tryptophan</td>
<td>31,33,36</td>
</tr>
<tr>
<td>1355 ± 5</td>
<td>Tryptophan</td>
<td>31,33,36</td>
</tr>
<tr>
<td>824 ± 5</td>
<td>Tyrosine</td>
<td>31,33,36</td>
</tr>
<tr>
<td>854 ± 5</td>
<td>Tyrosine</td>
<td>31,33,36</td>
</tr>
</tbody>
</table>

\(^a\) A combination of at least two of these three bands has to be present
\(^b\) All three bands have to be present in the spectrum

3 Results and discussion

For the investigation of virions based on tip-enhanced Raman spectroscopy (TERS), two virus species were chosen: the Porcine teschovirus (PTV) and the Varicella-zoster virus (VZV). Initially, AFM topography images of both virus samples was measured to find viral particles. In Fig. 2, example AFM topographic images of VZV (A) and PTV (B) are shown. After topographic imaging of the virions, the subsequent spectroscopic investigations were carried out. The combination of AFM with enhanced Raman spectroscopy, allowed to record Raman spectra of the surface of virions. After imaging the topography of the virus surface, distinct points in form of a grid were selected. The dimension of three grids per virus species can be found in the supporting information (SI-Tab. S1). At each grid point of a TERS spectrum was recorded. In Fig. 2, two spectra of each virus species are shown as examples. The band at 520 cm\(^{-1}\) is assigned to silicon\(^{37}\) from the TERS tip. The two spectra in the lower panel were recorded by spectroscopically mapping the PTV virions. To protect their genome, the approximately 30-nm PTV virions have an icosahedral capsid, which consists of protein capsomeres. The TERS spectra of PTVs thus exhibit protein-specific signals. As an example, the amide I and III bands (protein markers) are highlighted in green in the spectra. In the spectra of the 200-nm-sized VZV virion, protein-specific signals are visible as well. VZV virions also possess a capsid to protect the genome, which is further surrounded by a proteinaceous tegument and a lipid layer. Thus, VZV is an enveloped virus. In the lipid layer, proteins are tightly embedded, so that both categories of biomolecules can be detected in the TERS spectra. Some bands in the TERS spectra of VZV virions are therefore characteristic for lipids. In Fig. 2, these bands are marked in bluish colours. As can be clearly seen, the lipid bands are recognisable only in the TERS spectra of VZV virions, not in the PTV spectra. In addition to the labelled bands in Fig. 2, more characteristic TERS signals were used for the data evaluation and are listed in Tab. 1. For the evaluation of the TERS spectra, the signals were divided into two categories, protein and lipid. Tab. 1 does not present a complete assignment of lipids as well as proteins and TERS spectra contain many more signals. For example, bands of CH\(_2\) modes are present in both compounds and thus, are not specific to one of the two categories. In order to ensure an unambiguous assignment very specific marker bands of lipid and protein/amino acids were used for the data analysis. For instance, phenylalanine has an individual peak at 1004 ± 5 cm\(^{-1}\) (assigned to the ring breathing vibration\(^{31,33–36}\), which is considered as protein signal. In addition to individual peaks, band combinations were used to attain an unambiguous assignment. An example for a band combination are the two peaks at 1033 ± 5 cm\(^{-1}\)\(^{33–35}\) and 1210 ± 5 cm\(^{-1}\)\(^{131,33–36}\). These two peaks are also regarded as protein signal for phenylalanine, whereas both peaks have to be present in the TERS spectra in order to count as a valid signal. The simultaneous detection of certain bands was in some cases a premise to unambiguously categorize a spectrum. In particular this was valid for marker bands with close band positions like for instance phenylalanine and tryptophan. Here, the presence of a band at 1033 ± 5 cm\(^{-1}\) was not sufficient to identify phenylalanine but required a concurrent band...
Fig. 2 Left: AFM topography images (height profile) of a A: Varicella-zoster virus and of a B: Porcine teschovirus are visualised. The illustrated height profile was levelled by subtraction of the mean level and the scan lines were corrected by matching the height medians using the Gwyddion software (version 2.36). The AFM images were measured in intermittent mode and in air by using a silicon nitride AFM tip. C: Two exemplary untreated TERS spectra of each virus: Varicella-zoster virus (VZV) and Porcine teschovirus (PTV) are shown. The amide bands (protein markers) are marked in green colours. The lipid bands that only occur in the TERS spectra of VZV are labelled in blue colours.
at 1210 ± 5 cm⁻¹. A further example for band combinations to this molecule group have to be present in a TERS spectrum to be interpreted as lipid signal. The signals in unseparated rows in Tab. 1 must be simultaneously present in order to be recognised as signal.

3.1 Data evaluation with manual quality rating

Based on the assignment of specific Raman bands to the two categories, namely protein and lipid, component maps can be created. A grid-based TERS measurement of the virus surface was performed so that at each point of the grid a spectrum was recorded. A component map represents such a grid. Thereby, each square in the map represents the according measuring point on the grid and therefore a spectrum recorded at this location. Hence, in the maps, a value for both categories can be specified for each point. The grid dimensions are given in SI-Tab. S1. The resulting component maps are illustrated in Fig. 3. Thus, each point corresponds to a TERS spectrum and the colour reflects the number of bands and band combinations found for the respective category. For a better comparison of all measured TERS spectra, an intensity normalisation was carried out using the silicon peak. Before the bands of interest were fitted and accepted as peak using Equation 1, all spectra of the grid underwent a manual quality rating, i.e., spectra with insufficient quality (too low signal-to-noise ratio caused by, for example, poor feedback of the tip during the measurement) were sorted out. The corresponding pixels are illustrated in grey in the component map. If a point was coloured black, then a satisfying TERS spectrum was recorded at this position. However, the present bands couldn’t be unambiguously assigned to one of the two categories. For example, the vibration assigned to the C-H molecule group can be found in lipid as well as protein spectra. Thereby, this signal does not provide additional information to successfully discriminate the virus species as presented in this study.

On the left in Fig. 3, the component maps based on the protein signals for VZV (above) and PTV (below) are presented. In comparison, on the right, the component maps based on the lipid signals are visualised. It is obvious that a high protein content is present on the surface of both viruses derived from the detected protein signals in a large number of TERS spectra. This is consistent with the morphology of the PTV, whose outermost layer is constituted by the capsid’s proteins. In the case of the VZV, also protein signals can be expected, as proteins are embedded in a lipid envelope. Upon consideration of the lipid component maps of VZV, lipid-specific signals were found only in a few TERS spectra. This may be due to the hardly accessible lipid layer of VZV. The proteins incorporated there are packed very densely and thereby impede the measurements of the lipid signals. The TERS measurements were performed in the intermittent mode to reduce the tip-induced deformations of the viral particle, so that the TERS tip did not always reach the lipid layer of the VZV in between the protein spikes. In comparison, in the lipid component map of PTV virion no coloured points and therefore no lipid indicators are present. In agreement with the lack of lipid components in the PTV particle, no lipid-specific signal in the TERS spectra of the according PTV virion was identified. Therefore, a discrimination of the two viruses is possible based on the observation of protein and lipid contents.

The process described above is based on the manual decision regarding the quality of the spectra. Such a decision differs from person to person and is highly subjective. In addition, an automation of the quality rating process for a higher amount of data, as it is necessary for model building algorithms, is more than useful. A manual quality rating for more than 200 spectra is very time consuming and unsatisfying. The method for creating the protein and lipid maps was adopted and adjusted accordingly to allow its application for an automatic quality rating for all measured spectra. In the adjusted method, the distinction in protein and lipid signals was discarded and all found signals were counted for each spectrum. If a spectrum showed at least one of the signals from Tab. 1, it was regarded as high quality spectrum and considered for the automatic classification; otherwise, it was discarded. Consequently, it was possible to obtain an objective and fast quality rating for TERS spectra.

3.2 Classification

For the creation of the manual maps, two grids with each 144 single raw spectra were recorded. After the manual quality rating, 78 spectra for VZV and 128 spectra for PTV were analysed for the expected protein and lipid signals given in Tab. 1. To increase the robustness and the general character of the classification model obtained through a LDA, the data set was extended. For the analysis, six virus grids including 1507 raw spectra were measured (913 spectra of the VZV and 594 spectra of the PTV). For the sake of completeness, the protein and lipid maps for the additional grids are given in SI-Fig. S1.

After a successful automatic quality rating for the six grids, the cleared data set (then consisting of 1322 single spectra (802 for VZV, 520 for PTV)) was used to train a classification model based on a Linear Discriminant Analysis (LDA). Thereby, each class included spectra measured on three different particles of the according species. Furtheron, a leave-one-out cross-validation (LOOCV) was conducted to evaluate the classification model. The result of the LOOCV is presented in Tab. 2. The numbers on the diagonal of the table show the number of correctly assigned spectra to the according class. A classification accuracy can be determined by dividing the sum of the correctly assigned spectra by the number of spe-
Fig. 3 Component maps for the protein (left) and lipid (right) category of VZV grid: A (above) and PTV grid: a (below). The number of bands and band combinations found in the TERS spectra that are typical either for protein or lipids are visualised by the false colour scale. Each quadratic point of the component maps represents a TERS spectrum.
Table 2 Result of the LOOCV for the classification model based on LDA (A) with automatically rated spectra and an accuracy of approximately 91% and (B) with manually rated data and an approximate accuracy of 86%.

<table>
<thead>
<tr>
<th>A</th>
<th>reference</th>
<th>prediction</th>
<th>PTV</th>
<th>VZV</th>
</tr>
</thead>
<tbody>
<tr>
<td>PTV</td>
<td>512</td>
<td>8</td>
<td></td>
<td></td>
</tr>
<tr>
<td>VZV</td>
<td>108</td>
<td>694</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>B</th>
<th>reference</th>
<th>prediction</th>
<th>PTV</th>
<th>VZV</th>
</tr>
</thead>
<tbody>
<tr>
<td>PTV</td>
<td>346</td>
<td>16</td>
<td></td>
<td></td>
</tr>
<tr>
<td>VZV</td>
<td>109</td>
<td>423</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Further, it was shown that the automatic quality rating was used in the analysis (sum over the whole table). The analysis showed a classification accuracy of approximately 91% for the two examined species based on the automatically rated data set. For comparison, the same analysis was carried out for the same data with a manual quality rating. In this part of the analysis, 894 single spectra were taken into account. Thereby, 532 spectra represented the class of the Varicella-zoster virus and 362 spectra the class of the Porcine teschovirus, respectively. The accuracy for this analysis decreased to 86%.

The manual quality rating discarded approx. 41% of the spectra whereas the automatic quality rating determined an unsatisfying quality for approx. 12% of the spectra. Despite the higher number of chosen spectra, the last mentioned method led to a decreased in-group variance for the considered spectra. Thus, the automatic quality rating, although being less strict than a manual quality rating, delivers a data set free of unusable spectra that behaves more stable in a classification task and leads thereby to improved general results.

In both cases, misclassifications were mainly due to the high variance in the spectra, this holds especially for the Varicella-zoster virus. This group showed a high standard deviation over all group spectra due to the highly variable morphology, the rather amorphous structure of the VZV virions and the fact that lipid and protein signals can occur in the spectra. In contrast, the PTV spectra showed a significantly lower in-group variance because of their simple structure. The different magnitudes of variance are reflected in both of the LOOCV results: The number of misclassified spectra of the PTV is a magnitude lower than the number of misclassified VZV spectra indicating a higher in-group variance. Such high variance for certain species was also observed by another working group and is thereby not an artefact of the presented measurements.

Furthermore, the variation of the TERS intensities can be explained by a different orientation of the molecules to the TERS tip as well as by the field gradient Raman (FGR) effect. The high lateral resolution of TERS results is much less averaging, consequently, the spectral variations are also related to locally specific interactions on the nanometre scale. Furthermore, at the apex of the tip (in the optimal case, there is a single nanoparticle), a field gradient can be induced in TERS measurement, so that a FGR effect may occur. Both effects result in intensity variations and significantly influence the spectroscopic analysis of complex structures. As previously mentioned, the VZV shows a high variability in its structure and is composed of lipids as well as proteins in comparison to the PTV. Thus, the interaction of the more complex virus structure with the mentioned effects can explain the higher variance in the TERS spectra of VZV.

Nevertheless, a discrimination of the two examined viruses based on a LDA is possible. In addition, the results indicate an improvement using an automatic quality rating for the spectral data and thereby a successful application of the introduced method.

4 Conclusion

Two different virus species were examined using tip-enhanced Raman spectroscopy and spectroscopic information was obtained from the virus surface. Due to the application of the TERS technique, the viruses were analysed on a single-particle level. Therefore, the possibility exists to refrain from pre-cultivation of viruses.

Through the assignment of characteristic lipid and protein bands, component maps of both categories were created. In the TERS spectra of the Varicella-zoster virus protein and lipid signals were identified. In comparison, for the Porcine teschovirus only protein signals were detected due to compositional differences. This is reflected in the component maps of both viruses that allowed a visual discrimination based on the spectroscopic data of the two examined species. Thus, the manual component maps represent a possibility to distinguish viruses for the untrained eye through detachment of the decision from spectroscopic expert knowledge.

Furthermore, the method used to create the above mentioned maps was adopted to perform an automatic quality rating of the TERS spectra. This step allows an objective view of the data as well as the inclusion of a high number of spectra and a reduction of the analysis time avoiding a manual quality rating by a spectroscopist. This is not only an achievement towards the automation but also to an user-independent data evaluation. The spectra which have at least one specific signal were positively rated, used to train a classification model based on a LDA, and evaluated by a leave-one-out cross-validation. For the two investigated virus species a classification accuracy of approximately 91% was achieved. Further, it was shown, that the automatic quality rating was
successfully applied to TERS spectra and thus increased the classification power of the trained model.

Thereby, the presented study illustrates two paths for successful discrimination of single viral particles based on their spectroscopic information. To our knowledge, this is the first time that a discrimination of viruses is achieved solely based on TERS spectra in combination with chemometric methods.

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