
Label-free biochemical imaging of neural organoids via deep learning-enhanced Raman microspectroscopy

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Abstract

Three-dimensional organoids have emerged as powerful models for studying human development, disease and drug response *in vitro*. Yet, their analysis remains constrained by standard imaging and characterisation techniques, which are invasive, require exogenous labelling and offer limited multiplexing. Here, we present a non-invasive, label-free imaging platform that integrates Raman microspectroscopy with deep learning-based hyperspectral unmixing for unsupervised, spatially resolved biochemical analysis of neural organoids. Our approach enables high-resolution mapping of cellular and subcellular structures in both cryosectioned and intact organoids, achieving improved imaging accuracy and robustness compared to conventional methods for hyperspectral analysis. This work establishes a versatile framework for high-content, label-free (bio)chemical phenotyping with broad applications in organoid research and beyond.

1 Introduction

The ability to image and accurately characterise the architecture and molecular composition of cells and tissues has been central to advancing biological and medical sciences. To this end, a broad array of techniques – ranging from histological staining and immunofluorescence to omics-based profiling – has been developed, each offering distinct insights across spatial and molecular scales^{1–6}. Nonetheless, a fundamental challenge persists: most available methods are inherently invasive and/or require molecular labelling.

These constraints become particularly limiting as scientific research shifts toward complex, three-dimensional (3D) tissue models, such as organoids. Over the past decade, neural organoids derived from pluripotent stem cells have emerged as powerful biological models of the developing human brain, recapitulating key features of early developmental dynamics and organisation^{7–10}. Compared

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to two-dimensional (2D) neural cell culture models, neural organoids exhibit enhanced complexity and three-dimensional structures, providing a more advanced representation of tissue architecture^{7,10}, cellular diversity^{11,12} and functional network formation¹³. Consequently, neural organoids have spurred new opportunities for studying human brain development, disease and drug response *in vitro*^{14–17}. Yet, despite growing interest, organoid research remains constrained by the limitations of conventional methods for tissue imaging and characterisation. There is a pressing need for techniques that can interrogate spatial organisation and dynamics in intact organoid systems under physiologically relevant settings, without prior knowledge of specific molecular targets or the introduction of exogenous labels.

Currently, immunofluorescence staining combined with fluorescence microscopy is widely used as a gold standard, reference technique for visualising the morphological organisation of 2D and 3D cellular models, owing to its high specificity and spatial resolution^{18,19}. In this approach, fluorophore-conjugated antibodies are used to selectively bind to and visualise specific target molecules²⁰. Despite its strengths, however, this technique has several inherent limitations. First, while protein labelling is generally well-established, detecting other biomolecular classes, such as carbohydrates, lipids and various small molecules (e.g. drugs and metabolites), often requires more specialised probes or labelling strategies, or is not possible altogether. Additionally, immunofluorescence staining involves multiple washing and permeabilisation steps, which can disrupt cellular architecture and compromise membrane integrity, potentially leading to artefactual results²¹. These steps may also displace small molecules such as drugs, metabolites and other transient compounds, obscuring their true spatial localisation *in situ* within cells and tissues. While other labelling strategies, such as fluorescent protein tagging, genetic modifications and dye-based staining, may help avoid these washing steps, they remain label-based (i.e. targets must be known and available), and issues with labelling non-proteins persist. Furthermore, regardless of the labelling strategy, the intrinsic spectral overlap of fluorescent markers constrains multiplexing, typically to four targets in standard fluorescence microscopy²², or seven in more advanced techniques^{23,24}. Moreover, genetic modifications used in live-cell imaging to express fluorescent protein reporters can be costly, time-consuming, and may induce poorly understood side effects, such as protein aggregation or disruption of ion transport dynamics. Enabling non-invasive, label-free analysis would thus be essential to expand the potential of neural organoids in neurodevelopmental research, disease modelling, drug screening and beyond.

Raman spectroscopy (RS), a technique within vibrational spectroscopy, offers a promising alternative for biochemical analysis. Unlike fluorescence-based imaging, RS provides molecular contrast through the analysis of the intrinsic inelastic light scattering of molecules, thereby enabling non-invasive, label-free chemical characterisation with minimal sample preparation^{25–27}. Consequently, RS has become a ubiquitous analytical technique across the life and physical sciences^{28–38}, with increasing use in analysing biological specimens such as cells and tissues^{39–44}. In the context of neurodevelopment, RS has been applied to profile neural cells at various stages of differentiation. Yet, most studies to date have focused on 2D cultures of individual cell types with RS measurements acquired at a single-cell level – e.g. distinguishing neural stem cells from neurons by identifying distinct cell-specific spectral signatures^{45–49}. The application of RS to complex three-dimensional cultured tissues containing multiple cell types, such as neural organoids, remains challenging due to their high biochemical heterogeneity, which hinders signal unmixing and interpretation. Recently, Bruno et al. explored the use of supervised machine learning to classify RS spectra from cortical organoids at four predefined maturation time points (weeks 6, 12, 16 and 20)⁵⁰. However, their approach focused on single-point prediction and did not explore the three-dimensional architecture, cellular interactions and functional complexity intrinsic to neurodevelopment. So far, the use of RS for spatially resolved analysis of neural organoid structure and maturation remains unexplored.

Our lab recently reported a technique based on RS for 3D biochemical imaging and drug distribution analysis in liver organoids⁵¹. The proposed methodology relied on the acquisition of ‘tissue phantoms’ – i.e. calibrated reference spectra from biomolecules of interest, which were used to guide the estimation of chemical concentration maps across the scanned area using ordinary least squares with non-negativity constraint⁵². This enabled the detection and visualisation of key biomolecules and drugs *in situ* in whole liver organoids. Nonetheless, the approach has two limitations. First, it assumes that target biomolecules are known, thereby restricting potential applications and undermining the label-free capabilities of RS. Second, the performance and robustness of ordinary least squares for chemical estimation can be compromised by complex mixtures, experimental noise and artefacts, and variations introduced during data calibration and preprocessing, leading to inaccurate qualitative and

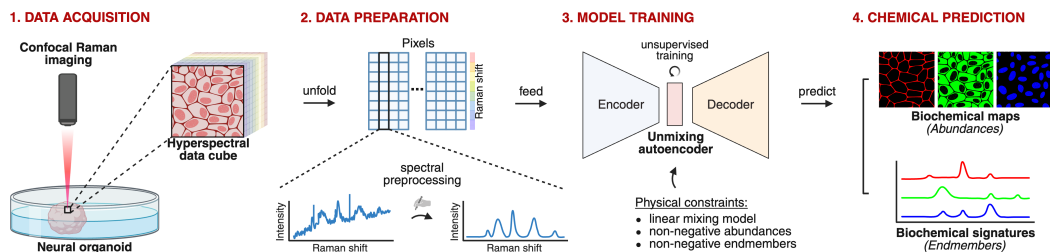


Figure 1: Schematic diagram of our organoid imaging pipeline combining Raman microspectroscopy and deep learning-enhanced hyperspectral analysis.

quantitative analysis. To address these limitations, we recently introduced a deep learning framework for RS chemometrics, using hyperspectral unmixing autoencoder neural networks, which enabled fully unsupervised, label-free (bio)chemical analysis with enhanced accuracy compared to established chemometric techniques⁵³.

Building on this work, we present here a pipeline for non-invasive, label-free biochemical imaging of neural organoids based on spontaneous Raman microspectroscopy and unsupervised autoencoder-based hyperspectral unmixing analysis. We show that our approach achieves improved organoid imaging compared to standard methods for RS analysis, as validated against fluorescence microscopy, and enables detailed morphochemical imaging and spatially resolved analysis in both cryosectioned and intact neural organoids with subcellular resolution.

2 Deep learning-enhanced Raman microspectroscopy

We begin by discussing the technical methodology behind our organoid imaging pipeline based on Raman microspectroscopy combined with autoencoder-based hyperspectral unmixing. Figure 1 provides an overview of our approach.

Raman measurements were collected from fixed neural organoid samples by performing point-wise imaging scans across predefined regions of interest. Volumetric scans were generated by acquiring sequential scans along the axis perpendicular to the imaging plane (e.g. sampling at different depths along the z -axis). Acquired data were preprocessed to remove non-specific signal contributions and spectral artefacts, such as cosmic spikes, autofluorescence baselines and dark noise (see Appendix 6.1). This was done using RamanSPy – an open-source Python toolbox for RS data analysis we recently developed⁵⁴.

After preprocessing, hyperspectral unmixing was performed to derive constituent spectral components (endmembers) and quantify their relative contributions (abundances) in each measurement^{55,56}. This was achieved using a custom autoencoder neural network model designed for hyperspectral unmixing. The model builds upon our previously reported approach⁵³, with substantial extensions and optimisations across the entire pipeline, spanning preprocessing, network architecture, training strategy and endmember estimation (see Appendix 6.1 for more details). Briefly, the autoencoder architecture included: (i) an encoder responsible for abundance estimation, which comprised a multi-branch spectral convolutional block followed by a five-layer fully connected network, and (ii) a one-layer decoder responsible for endmember identification. Our model incorporated relevant physical constraints in the representation learning process, namely, constraining the learnt endmembers and abundances to be non-negative through the use of appropriate activation functions and weight clipping, and constraining the mixing process to linear mixing through the use of a single-layer linear decoder. Models were trained on individual scans or datasets in an unsupervised manner by minimising a reconstruction loss based on the weighted sum of the mean squared error and spectral angle divergence⁵⁷ between input spectra and output reconstructions. The number of endmembers to extract for each scan/dataset was determined using principal component analysis (PCA)^{58,59}.

After model training, learnt endmember signatures and their corresponding abundances were derived, generating multi-channel abundance images/volumes with channels rendering chemical maps of the distribution of learnt endmembers. The obtained endmembers and abundances were examined to filter out components linked to non-tissue signals, such as background and noise. The remaining

tissue-specific endmembers were characterised via peak assignment, whereby prominent peaks were linked to chemical bonds to identify essential biomolecular species, such as nucleic acids, proteins, lipids and carbohydrates. Peak identification was guided by established reference materials⁴⁰. The corresponding abundance maps were visualised as morphochemical images showing the spatial distribution of the obtained chemical components to study organoids’ morphology and biochemical composition.

Further technical details regarding data acquisition, preprocessing and unmixing analysis are presented in Appendix 6.1.

3 Experimental validation against fluorescence microscopy

To validate that our method produces accurate morphochemical organoid reconstructions, we evaluated its performance against fluorescence microscopy and standard methods for hyperspectral RS analysis. To this end, defined tissue regions within two day-30 neural organoid cryosections were imaged using confocal Raman microspectroscopy, followed by immunostaining and fluorescence microscopy (Figure 2A). Raman imaging was performed first to avoid potential signal interference from staining molecules. To ensure morphological diversity, we selected areas containing both distinct neural rosette structures and surrounding neural tissue for downstream imaging (Figure 2B). Raman imaging was performed over areas of $600\text{ }\mu\text{m} \times 600\text{ }\mu\text{m}$ and $400\text{ }\mu\text{m} \times 400\text{ }\mu\text{m}$ from the two respective sections, with a pixel size of $2\text{ }\mu\text{m}$ (Figures 4C and 5C in Appendix 6.2). Subsequently, the organoid sections were stained for nuclei (DAPI), cell membranes (DiO), neural progenitor cells (SOX2) and neurons (TUI1), and fluorescence microscopy images were taken to validate and correlate with organoid morphology predicted by RS (Figures 4B and 5B in Appendix 6.2).

Using our autoencoder-based imaging pipeline, we analysed the acquired RS data from the first organoid section to infer its biochemical composition. Our method extracted four tissue-specific endmembers, which we associated with contributions from lipids, nucleic acids and proteins (Figure 4D–E in Appendix 6.2). Figure 2C shows a side-by-side comparison between the predicted abundance maps of the two endmembers related to nucleic acid-rich and lipid-rich species and the corresponding fluorescence microscopy images of nuclei (DAPI) and cell membranes (DiO), respectively. We observed that our RS imaging pipeline accurately reconstructed individual nuclei, aligning with the DAPI staining across the imaged tissue (Figure 2C (right), and Figure 4F in Appendix 6.2). It also effectively captured important membrane features, including both fine cellular membranes and prominent structures such as the apical domain of the visible neural rosette. Qualitatively, we observed strong correspondence between the lipid-rich component obtained with RS and the DiO signal around the apical domain of the rosette and with the TUI1 signal outside the rosette (Figure 4B in Appendix 6.2).

Having shown that our method accurately reconstructs key cellular components in cryosectioned neural organoid samples, we used the data collected from the second organoid section to compare the unmixing performance of our developed approach against a range of standard chemometric methods for multivariate analysis. This included methods for dimensionality reduction (principal component analysis (PCA)^{58,59}, non-negative matrix factorisation (NMF)⁶⁰), clustering (k-means clustering^{61,62}), and hyperspectral unmixing (N-FINDR⁶³ and vertex component analysis (VCA)⁶⁴ for endmember identification, applied together with non-negative least squares (NNLS)⁵² for abundance estimation). Our autoencoder-based pipeline derived six tissue-specific endmembers, which we characterised as lipid-, nucleic acid- and protein-rich species (Figure 5D in Appendix 6.2). Visual examination of the estimated abundance maps revealed major cellular and subcellular features, including membranes, nuclei, cytoplasmic regions and apical domains of neural rosettes (Figure 5E in Appendix 6.2). Compared to standard methods, our approach offered improved robustness to artefacts and noise and produced more discriminative and interpretable abundance maps, enabling better-delineated imaging reconstruction of organoid morphology (Figure 6 in Appendix 6.2).

To quantitatively assess the performance of our autoencoder-based pipeline, we evaluated the image similarity between Raman-based reconstructions and fluorescence microscopy images in an overlapping organoid region. To this end, we spatially aligned both modalities and computed the Pearson’s correlation coefficient (Pearson) and structural similarity index measure (SSIM)⁶⁵ between RS-derived abundance maps of nuclei and membranes and fluorescence images of DAPI and DiO, respectively. We benchmarked the performance of our method against two conventional chemometric

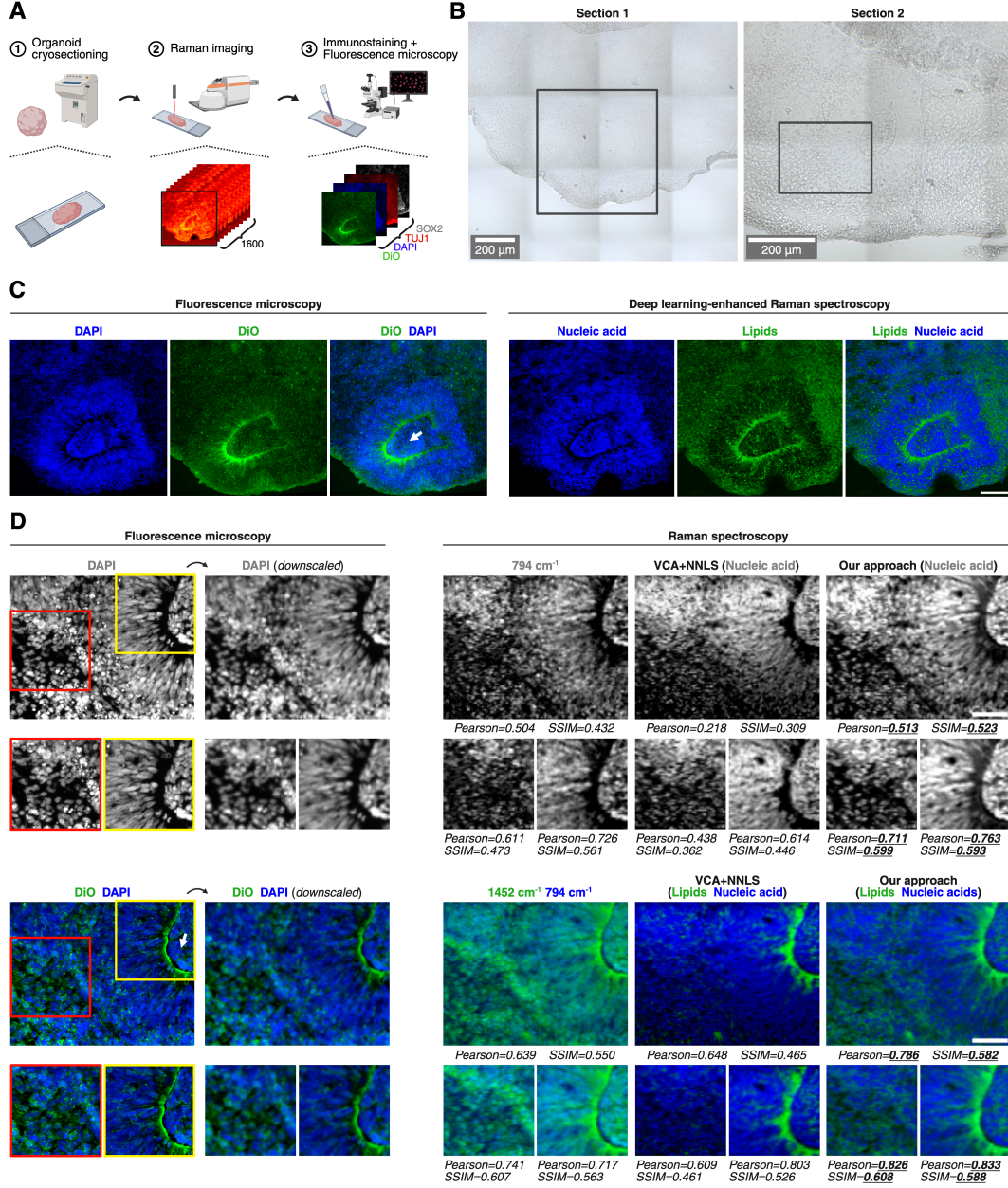


Figure 2: Deep learning-enhanced Raman microspectroscopy enables accurate morphochemical imaging of neural organoids as validated against fluorescence microscopy. **A**, Schematic diagram of our validation strategy, which involved the collection of paired Raman imaging and fluorescence microscopy data from neural organoid cryosections. **B**, Brightfield microscopy images of the two day-30 neural organoid cryosections imaged for validation. Marked areas show the regions used for validation in **C** and **D**, respectively. Scale bars: 200 μm . **C**, Imaging results on first organoid section. Comparison of fluorescence microscopy images of DAPI (nuclei) and DiO (membranes) (*left*), and RS-based abundance reconstructions of nucleic acid and lipids derived with our autoencoder pipeline (*right*). Arrow indicates the lumen of the present neural rosette. Scale bar: 100 μm . **D**, Imaging results on second organoid section. Comparison of fluorescence microscopy images of DAPI (nuclei) and DiO (membranes) (*left*), and RS-based abundance reconstructions of nucleic acid and lipids derived with three methods for Raman imaging, including univariate analysis, unmixing analysis with VCA and NNLS, and our autoencoder pipeline. Similarity metrics, namely Pearson's correlation coefficient (Pearson) and structural similarity index measure (SSIM), were performed with respect to the respective downscaled fluorescence images. The highest achieved similarity scores are indicated by underlined bold text. Arrow indicates the lumen of the present neural rosette. Scale bar: 50 μm .

approaches: (i) univariate analysis, and (ii) multivariate hyperspectral unmixing using VCA and NNLS. Univariate analysis was performed using the intensity profiles at 794 cm^{-1} and 1454 cm^{-1} , which we found to be indicative of nuclear and lipid-rich membrane regions, respectively (Figure 5D in Appendix 6.2).

As shown in Figure 2D, our autoencoder-based pipeline achieved the highest reconstruction accuracy across all evaluations – both quantitatively, with superior Pearson and SSIM scores, and qualitatively, with more interpretable and biologically relevant output. Nuclear regions were recovered with higher accuracy, with visually better-defined nuclei and lower non-specific signal. Membrane-associated structures were also more clearly delineated, enabling the visualisation of both large-scale features such as apical domains and individual cellular membranes.

Furthermore, our approach demonstrated improved robustness to experimental noise, maintaining consistent reconstruction quality under reduced denoising conditions. In comparison, the performance of conventional methods deteriorated markedly, yielding noisier, less interpretable spectral components and abundance profiles, or failing to converge to a solution during unmixing altogether (Figure 7 in Appendix 6.2).

4 Biochemical imaging of intact neural organoids at subcellular resolution

After demonstrating that our method achieves accurate subcellular imaging of 2D cryosectioned neural organoid tissue, we performed higher-resolution Raman imaging of a tissue region within an intact 3D neural organoid. The ability to resolve and visualise subcellular structures in intact organoid tissues in their native environment is critical for interrogating neurodevelopmental processes, such as nuclear positioning, polarity establishment and lumen formation, and studying their effect on brain tissue organisation, architecture and function.

To this end, we acquired a larger 400×346 Raman imaging scan taken with $1\text{ }\mu\text{m}$ pixel size from an intact day-10 neural organoid sample (Figure 3A). Using our Raman imaging pipeline, we analysed the data and identified six tissue-specific components. The spectral endmembers and corresponding abundance maps are presented in Figure 3B and Figure 3C, respectively. Endmember 1 (red) exhibited a lipid-rich profile with prominent peaks at 1077 cm^{-1} (C–C and C–O stretching, phospholipids), 1305 cm^{-1} , 1448 cm^{-1} and 1657 cm^{-1} . Its abundance map outlined membrane-like structures, along with more confined bright spots resembling lipid droplets. Endmember 2 (grey) was characterised by vibrations associated with both protein and lipid species, with peaks at 1000 cm^{-1} , 1124 cm^{-1} (C–C and C–N stretching), 1249 cm^{-1} , 1311 cm^{-1} , 1454 cm^{-1} and 1654 cm^{-1} . Its abundance was observed in regions partially overlapping with membranes, as well as in isolated domains, possibly indicating cytoplasmic or extracellular matrix compartments. Endmember 3 (blue) was marked by strong nucleic acid bands at 790 cm^{-1} and 1095 cm^{-1} , alongside protein-associated bands (1005 cm^{-1} , 1249 cm^{-1} , 1328 cm^{-1} , 1455 cm^{-1} and 1659 cm^{-1}). As in previous analyses, the abundance map revealed well-defined spherical compartments consistent with individual cell nuclei. Endmember 4 (green) presented a distinct protein-rich signature with peaks at 1005 cm^{-1} , 1251 cm^{-1} , 1454 cm^{-1} and 1662 cm^{-1} , along with vibrations associated with ring breathing modes in nucleic acid bases at 1573 cm^{-1} . The abundance map highlighted cellular-level compartments resembling cytoplasmic regions, as well as small punctate structures within nuclei with high signal intensity, potentially corresponding to biomolecular condensates. Endmember 5 (yellow) displayed a mixed profile comprising bands linked to proteins (1259 cm^{-1} , 1326 cm^{-1} , 1453 cm^{-1} and 1662 cm^{-1}) and nucleic acid (1095 cm^{-1}). This component appeared more diffusely distributed, with substantial spatial overlap with endmember 3. Endmember 6 (purple) presented a noisier signature, with carbohydrate-associated bands at 493 cm^{-1} (glycogen), 850 cm^{-1} (C–O–C skeletal mode, glycogen) and 938 cm^{-1} (C–C stretching), along with peaks linked to proteins and nucleic acids at 754 cm^{-1} and 1103 cm^{-1} . The abundance overlapped with endmember 4, particularly around nuclear condensates, potentially reflecting glycoproteins within cell nuclei.

To further inspect the spatial reconstructions produced by our imaging pipeline, we examined magnified fields from selected regions within the imaged organoid tissue (Figures 3D and 3E). The resulting reconstructions revealed consistent, well-defined cellular features, which delineated individual cells and subcellular structures resembling nuclei, cytoplasmic regions, cellular membranes and nuclear condensates.

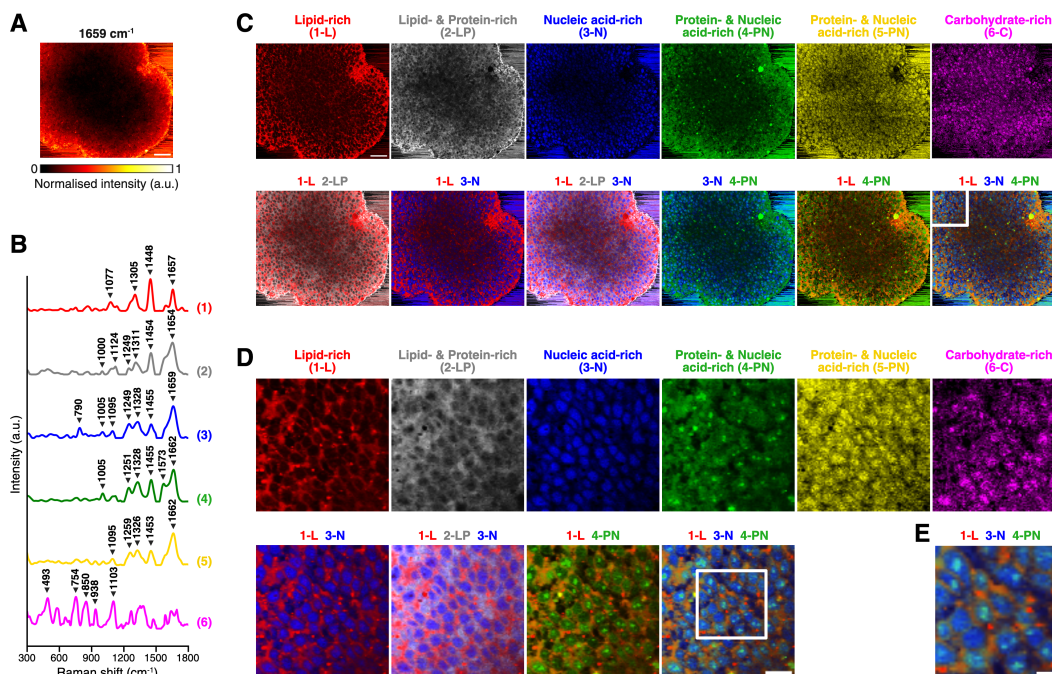


Figure 3: Biochemical imaging of intact neural organoids at subcellular resolution. **A**, Snapshot of collected Raman imaging data from an intact day-10 neural organoid sample, showing the intensity profile at the 1659 cm^{-1} band (proteins and lipids). Scale bar: $50 \mu\text{m}$. **B–E**, Results of hyperspectral unmixing analysis. **B**, Derived tissue-specific endmember signatures. Endmembers scaled for visualisation purposes (maximum intensity set to 1). **C**, Corresponding abundance maps. Scale bar: $50 \mu\text{m}$. **D**, Zoomed-in abundance images of a selected area (area marked in **C**, bottom right panel). Scale bar: $20 \mu\text{m}$. **E**, Further zoomed-in abundance image (area marked in **D**). Scale bar: $10 \mu\text{m}$.

These results demonstrate the effectiveness of our imaging platform in intact organoid tissues, allowing us to resolve complex biochemical heterogeneity and interrogate key morphological features at cellular and subcellular scales in an unsupervised, label-free manner.

5 Discussion

We have developed here a method for non-invasive, label-free organoid imaging based on Raman microspectroscopy combined with unsupervised autoencoder-based hyperspectral analysis. We showed that our approach is effective at discerning key biochemical components in unlabelled Raman spectroscopy measurements, enabling the visualisation of structural and morphological features in both cryosectioned and intact neural organoids and at different scales and resolutions.

By eliminating the need for destructive preparation steps and external labels, our workflow offers simpler sample preparation that does not compromise organoid integrity, with potential for live-organoid analysis. This opens new avenues for research into early neurodevelopment, but also in developmental diseases and drug discovery, where our approach could allow studying disease aetiology in neural organoid disease models or visualising drug distribution *in situ*. We note that our pipeline can also be applied to other types of samples, including other types of organoids and tissues. Beyond imaging, it can also be used as a feature extraction technique in RS for downstream applications – e.g. organoid phenotyping, disease classification, drug response modelling.

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Competing Interests

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6 Supplementary Material

6.1 Extended methods

6.1.1 Raman microspectroscopy

Raman spectroscopy measurements were performed using a confocal Raman microscope (alpha300 R, WITec GmbH) equipped with a 532 nm solid-state laser (WITec GmbH). Laser power at the sample plane was set to about 48 mW for the two organoid cryosections or 32 mW for the intact sample. Backscattered light was collected through a water immersion objective – 20× (Zeiss W N-Achroplan, N.A.= 0.5) for organoid cryosections or 63× (Zeiss W Plan-Apochromat, N.A.= 1) for intact sample, and delivered via a 10 µm single-mode silica fiber to an imaging spectrograph (Newton, Andor Technology Ltd) with a 600 groove/mm grating and a thermoelectrically cooled CCD detector (−60°C). The system provided a spectral resolution of $\sim 2\text{--}3\text{ cm}^{-1}$ over the wavenumber range 0–3670 cm^{-1} . Spectra were acquired using an integration time of 0.5 seconds.

6.1.2 Spectral preprocessing of Raman imaging data

Acquired Raman spectroscopy measurements were exported as MATLAB files using the WITec Project FIVE Suite software and loaded in Python using RamanSPy⁵⁴. Following this, each scan was individually preprocessed using RamanSPy as follows: (1) spectral cropping whereby wavenumbers less than 300 cm^{-1} were removed; (2) cosmic rays removal, using the algorithm reported by Whitaker and Hayes⁶⁶, with a kernel size $m = 3$ and z -value threshold $\tau = 6$; (3) denoising with a third-order Whittaker smoother⁶⁷ with $\lambda = 10^3$; (4) baseline correction with asymmetric least squares⁶⁸ with $\lambda = 10^5$; and (5) global vector normalisation, where intensity values were divided by the ℓ^2 -norm of the spectrum with the highest norm in the given scan. Additionally, wavenumber calibration was performed on our cryosection scans by calculating the median position of the laser peak (Rayleigh scattering) and setting it to zero.

6.1.3 Hyperspectral unmixing analysis

Chemometric analysis was performed using a custom autoencoder neural network model designed for blind hyperspectral unmixing. In this context, Raman measurements $\mathbf{x}_i \in \mathbb{R}^b$ are treated as mixtures

$\mathbf{x}_i = \mathcal{F}(M, \alpha_i)$ of a set of n unknown endmember components $M \in \mathbb{R}^{n \times b}$ based on their relative abundances $\alpha_i \in \mathbb{R}^n$ in a given measurement \mathbf{x}_i .

To decompose a set of Raman measurements $\mathcal{X} \in \mathbb{R}^{m \times b}$ (i.e. a given scan) into endmember components M and their relative abundances $A \in \mathbb{R}^{m \times n}$, we train an autoencoder model \mathcal{A} consisting of an encoder module \mathcal{E} and a decoder module \mathcal{D} . The encoder was responsible for mapping input spectra \mathbf{x} into latent space representations $\mathbf{z} = \mathcal{E}(\mathbf{x})$, and the decoder was responsible for mapping these latent representations into reconstructions of the original input $\hat{\mathbf{x}} = \mathcal{D}(\mathbf{z}) = \mathcal{D}(\mathcal{E}(\mathbf{x})) = \mathcal{A}(\mathbf{x})$. The model was trained in an unsupervised manner by minimising the reconstruction error between the input \mathbf{x} and the output $\hat{\mathbf{x}}$. During this process, the model was guided to learn the endmember components M and their relative abundances A through the introduction of related physical constraints. Below, we provide additional details about the developed architecture and training procedure. For more information about hyperspectral unmixing, the reader is pointed to previous works by Keshava and Mustard⁵⁵ and Li et al.⁵⁶.

The encoder \mathcal{E} comprised two separate blocks applied sequentially. The first part was a multi-branch convolutional block comprising four parallel convolutional layers with kernel sizes of 5, 10, 15 and 20, designed to capture patterns at multiple spectral scales. Each convolutional layer contained 32 filters with ReLU activation, He initialisation⁶⁹ and ‘same’ padding. Batch normalisation⁷⁰ and dropout with a rate of 0.2⁷¹ were applied to each convolutional layer to improve training stability and generalisation. The outputs of the four convolutional layers were merged channel-wise through a fully connected layer to yield an output of dimension matching that of the input spectrum. The rationale behind this was to transform intensity values into representations that capture local spectral features (e.g. peak shape, width, local neighbourhood) and thus promote better generalisability. The second part of the encoder was a fully connected dimensionality reduction block, applied to learn patterns between the learnt spectral features. This block comprised a series of fully connected layers of sizes 256, 128, 64 and 32 with He initialisation and ReLU activation. Batch normalisation and dropout (rate of 0.5) were also applied at each fully connected layer. The block was followed by a final fully connected layer (Xavier uniform initialisation⁷²) that reduced the final 32 features to a latent space of size n . The number n was treated as a hyperparameter that encodes the number of endmembers to extract, with latent representations treated as abundance fractions. To improve interpretation, non-negativity was enforced in the latent space using a ‘softly-rectified’ hyperbolic tangent function $f(x) = \frac{1}{\gamma} \log(1 + e^{\gamma \tanh(x)})$, with $\gamma = 10$, as we previously reported⁵³. This ensured that latent values were constrained to the range $(0, 1)$ in accordance with the abundance non-negativity constraint, but does not enforce abundances to sum to one (abundance sum-to-one constraint).

The decoder \mathcal{D} consisted of a single fully connected layer (Xavier uniform initialisation) mapping the n -dimensional latent space representations back to the original spectral dimension b . In this layer, we used a linear activation and no bias term. Under this setup, the decoder mimics linear unmixing, where endmember signatures are encoded in the learnt weight matrix of the decoder layer⁵³. A non-negative kernel constraint was used to enforce the non-negativity of endmember signatures using weight clipping during training. To accelerate model training, the weight matrix in the linear decoder layer was initialised before training using endmember signatures derived with vertex component analysis⁶⁴.

Autoencoder models were trained for 5 epochs using the Adam optimiser⁷³ with a batch size of 32. The learning rate was set to 0.001. An exponential learning rate decay was used to ensure stable convergence, applying a factor of 0.9 to the learning rate after each epoch. The training loss \mathcal{L} was defined as a weighted sum of the mean squared error (MSE) and spectral angle divergence (SAD) between input spectra \mathbf{x} and output reconstructions $\hat{\mathbf{x}}$:

$$\mathcal{L}(\mathbf{x}, \hat{\mathbf{x}}) = \text{SAD}(\mathbf{x}, \hat{\mathbf{x}}) + \lambda \text{MSE}(\mathbf{x}, \hat{\mathbf{x}}), \quad (1)$$

where

$$\text{MSE}(\mathbf{x}, \hat{\mathbf{x}}) = \frac{1}{b} \|\mathbf{x} - \hat{\mathbf{x}}\|_2, \quad (2)$$

with b denoting the dimension of \mathbf{x} (i.e. the number of spectral bands), and

$$\text{SAD}(\mathbf{x}, \hat{\mathbf{x}}) = \arccos\left(\frac{\mathbf{x} \cdot \hat{\mathbf{x}}}{\|\mathbf{x}\|_2 \|\hat{\mathbf{x}}\|_2}\right). \quad (3)$$

The weighting factor λ was set to 1000.

The number of endmembers to derive was estimated individually for each scan of interest using principal component analysis. Specifically, we computed the cumulative explained variance as a function of the number of principal components and manually selected the number of components at the point where the increase in explained variance visually plateaued.

Image similarity analysis between RS abundance images and fluorescence microscopy images was performed using Pearson’s Correlation Coefficient (PCC) and Structural Similarity Index Measure (SSIM). Fluorescence images were first aligned to the corresponding RS images and then downsampled to match the RS image dimensions prior to analysis.

PCC was calculated as:

$$\text{PCC} = \frac{\sum_i (x_i - \bar{x})(y_i - \bar{y})}{\sqrt{\sum_i (x_i - \bar{x})^2} \sqrt{\sum_i (y_i - \bar{y})^2}}, \quad (4)$$

where x_i and y_i represent pixel intensities of the RS and fluorescence microscopy images, respectively, and \bar{x} , \bar{y} are their mean intensities.

The Structural Similarity Index Measure (SSIM) was computed as:

$$\text{SSIM}(x, y) = \frac{(2\mu_x\mu_y + C_1)(2\sigma_{xy} + C_2)}{(\mu_x^2 + \mu_y^2 + C_1)(\sigma_x^2 + \sigma_y^2 + C_2)}, \quad (5)$$

where μ_x , μ_y denote means; σ_x^2 , σ_y^2 variances; σ_{xy} covariance; and C_1 , C_2 constants.

Autoencoder model implementation, training, and evaluation steps were carried out using TensorFlow⁷⁴. Analysis with baseline chemometric methods was performed using RamanSPy⁵⁴.

6.2 Extended results

(see the following four pages)

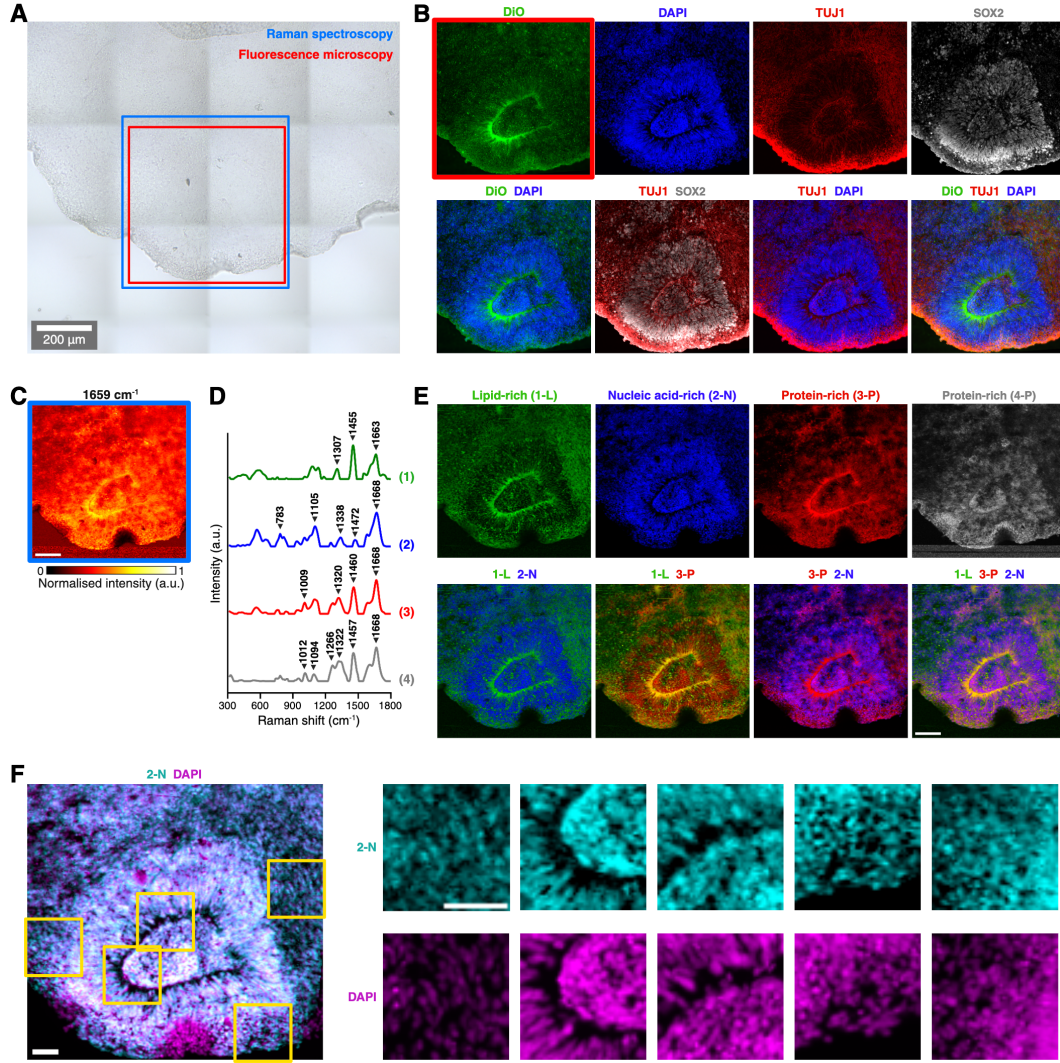


Figure 4: Extended results for Raman imaging scan of a first day-30 neural organoid cryosection. **A**, Brightfield image of a sectioned neural organoid tissue region imaged using Raman spectroscopy (blue) and fluorescence microscopy (red). Scale bar: 200 μm . **B**, Fluorescence microscopy images with markers for nuclei (DAPI), membranes (DiO), neural progenitors (SOX2) and neurons (TUJ1). **C**, Raman intensity profile at the 1659 cm^{-1} band (linked to proteins and lipids). Image size: 300 \times 300 pixels. Step size: 2 μm . Scale bar: 100 μm . **D–E**, Result of hyperspectral unmixing analysis. **D**, Derived tissue-specific endmember signatures. Endmembers scaled for visualisation purposes (maximum intensity set to 1). **E**, Corresponding abundance maps. Scale bar: 100 μm . Endmember 1 (green) presented a lipid-rich spectral signature with distinct peaks at 1307 cm^{-1} , 1455 cm^{-1} , and 1663 cm^{-1} . Its abundance was consistent with membrane structures, including cellular membranes and the apical domain of the visible neural rosette. The signal of this endmember showed a high degree of visual overlap with DiO and TOJ1. Endmember 2 (blue) displayed a nucleic acid-rich profile with a prominent band at 783 cm^{-1} , alongside vibrations linked to proteins (1338 cm^{-1} , 1472 cm^{-1} and 1668 cm^{-1}). Strong glass signal was also observed around 570 cm^{-1} and 1100 cm^{-1} . The abundance map revealed well-defined, spherical compartments, closely resembling cell nuclei, with increased density within the present neural rosette structure. Endmembers 3 (red) and 4 (grey) shared similar spectral signatures, dominated by protein-associated peaks around $\sim 1010 \text{ cm}^{-1}$, $\sim 1100 \text{ cm}^{-1}$, $\sim 1265 \text{ cm}^{-1}$, $\sim 1320 \text{ cm}^{-1}$, $\sim 1460 \text{ cm}^{-1}$, and $\sim 1670 \text{ cm}^{-1}$. Both endmembers were generally broadly distributed across the sample. Endmembers 3 showed increased signal at the apical domain of the present rosette, potentially suggesting structural proteins. The abundance of endmember 4 was increased within the rosette structure. **F**, Overlaid images of DAPI and endmember associated with nucleic acids (2-N). Scale bars: 25 μm .

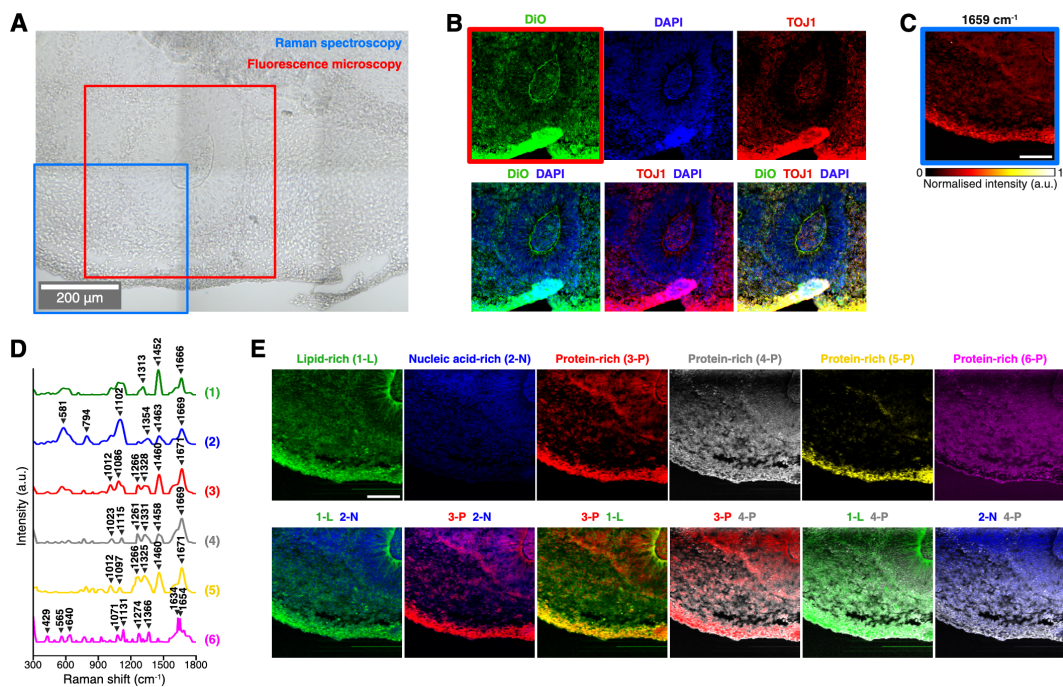


Figure 5: Extended results for Raman imaging scan of a second day-30 neural organoid cryosection. **A**, Brightfield image of a sectioned neural organoid tissue imaged using Raman spectroscopy (blue) and fluorescence microscopy (red). Scale bar: 200 μm . **B**, Fluorescence microscopy images with markers for nuclei (DAPI), membranes (DiO) and neurons (TUJ1). Due to partial tissue detachment during the staining process, a localised increase in fluorescence intensity was observed at the tissue edge. This peripheral region was identified as an artefact and was therefore excluded from subsequent analyses. **C**, Raman intensity profile at the 1659 cm^{-1} band (linked to proteins and lipids). Image size: 200 \times 200 pixels. Step size: 2 μm . Scale bar: 100 μm . **D–E**, Result of hyperspectral unmixing analysis. **D**, Derived tissue-specific endmember signatures. Endmembers scaled for visualisation purposes (maximum intensity set to 1). **E**, Corresponding abundance maps. Scale bar: 100 μm . Endmember 1 (green) exhibited a lipid-rich profile with prominent peaks at 1313 cm^{-1} , 1452 cm^{-1} , and 1666 cm^{-1} . The corresponding abundance map presented morphological architecture consistent with membrane-like structures, including individual cellular membranes and the apical domain of the present neural rosette. Endmember 2 (blue) displayed a strong nucleic acid band at 794 cm^{-1} , alongside several vibrations linked to protein species (1354 cm^{-1} , 1463 cm^{-1} and 1669 cm^{-1}), as well as bands attributed to the glass slide at (581 cm^{-1} and 1102 cm^{-1}). The abundance map revealed well-defined, spherical compartments, closely resembling cell nuclei. Endmembers 3 (red), 4 (grey) and 5 (yellow) exhibited similar spectral signatures, dominated by protein-associated peaks around $\sim 1015 \text{ cm}^{-1}$, $\sim 1100 \text{ cm}^{-1}$, $\sim 1265 \text{ cm}^{-1}$, $\sim 1330 \text{ cm}^{-1}$, $\sim 1460 \text{ cm}^{-1}$, and $\sim 1670 \text{ cm}^{-1}$. Endmember 4 showed a broad spatial distribution, likely corresponding to cytoplasmic or general background tissue signal. Endmembers 3 and 5, which showed a more pronounced 1460 cm^{-1} band, and an increased signal near the edge of the tissue section, likely due to tissue curling, as well as the basal domains of the present neural rosettes. Endmember 6 (magenta) presented a noisier signature, with prominent protein-associated peaks at 1131 cm^{-1} (C–N stretching), 1274 cm^{-1} , and 1366 cm^{-1} (tryptophan). Its spatial distribution showed substantial overlap with that of Endmember 4.

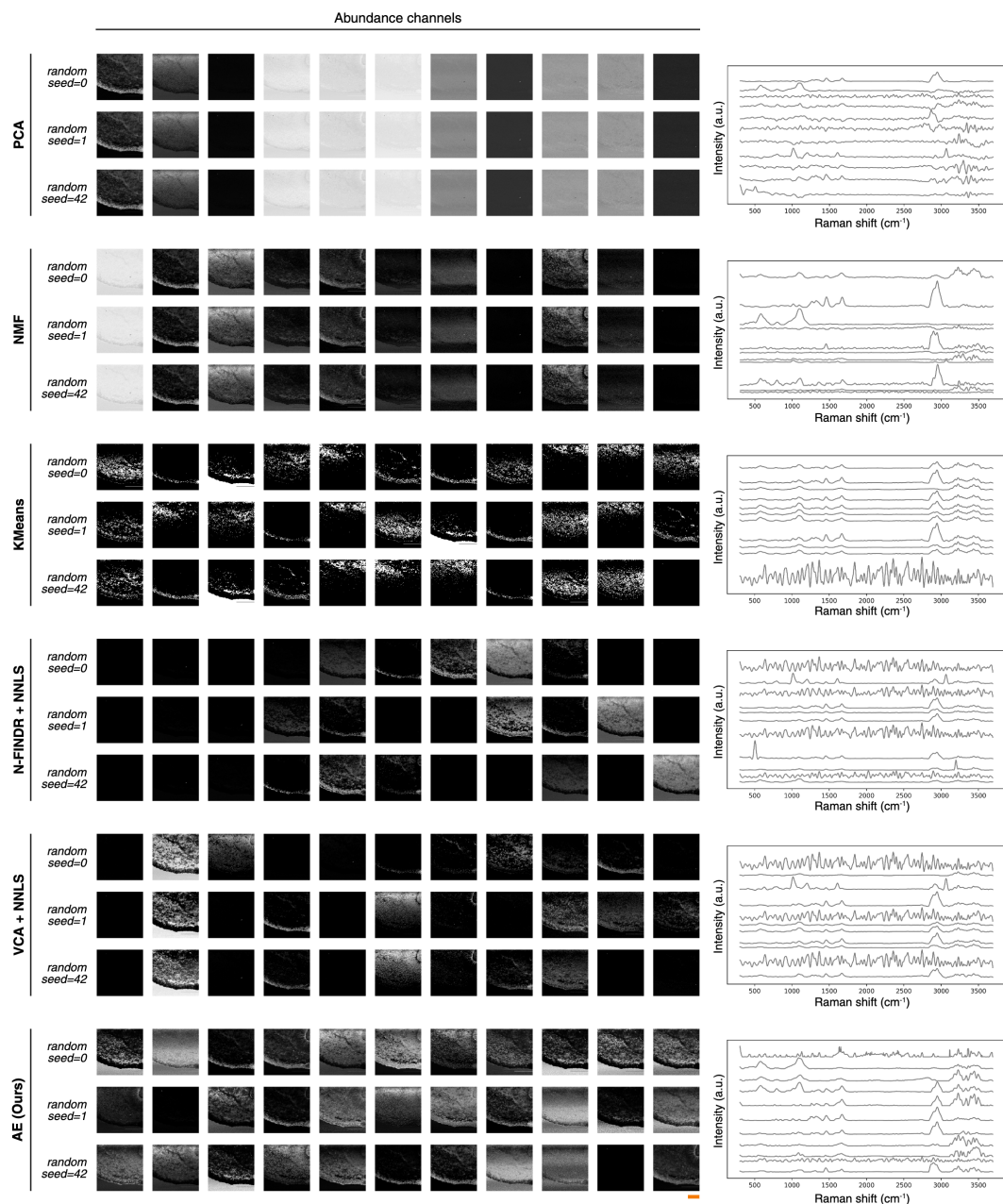


Figure 6: Comparative analysis of different chemometric methods on Raman imaging scan of a day-30 neural organoid cryosection. Methods include: principal component analysis (PCA), non-negative matrix factorisation (NMF), k-means clustering, N-FINDR + non-negative least squares (NNLS), vertex component analysis (VCA) + NNLS, our autoencoder (AE) pipeline. Abundance images were normalised for visualisation purposes. Endmember signatures displayed on the right correspond to the analyses performed with a random seed set to 42. Scale bar: 100 μm (bottom right, marked in orange).

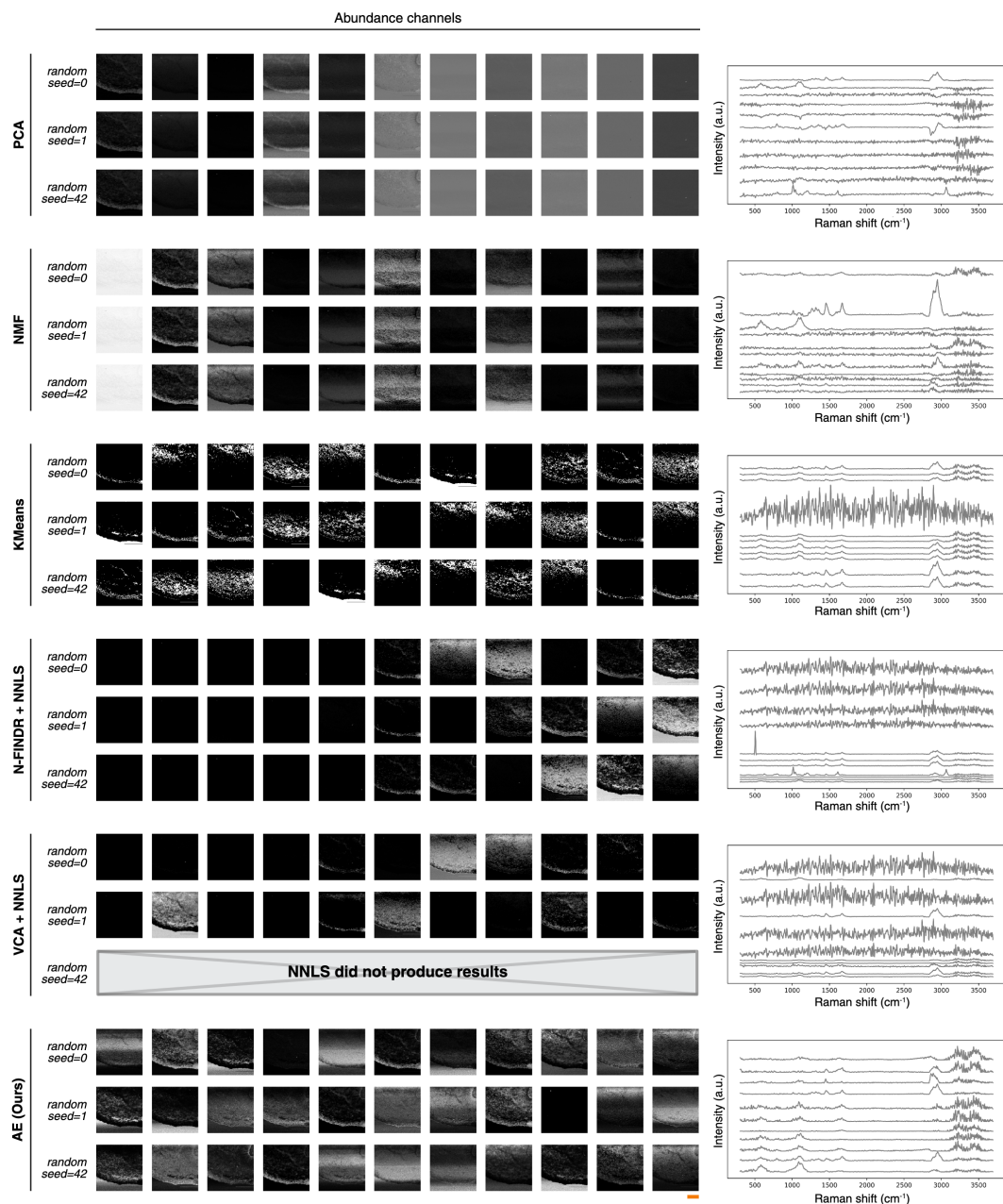


Figure 7: Comparative analysis of different chemometric methods on Raman imaging scan of a day-30 neural organoid cryosection in the presence of higher degrees of experimental noise. Higher noise levels were simulated by applying a more conservative denoising procedure with a second-order Savitzky-Golay filter with a window size of 7⁷⁵. Methods include: principal component analysis (PCA), non-negative matrix factorisation (NMF), k-means clustering, N-FINDR + non-negative least squares (NNLS), vertex component analysis (VCA) + NNLS, our autoencoder (AE) pipeline. One of the analyses with VCA+NNLS failed to converge, with the NNLS implementation terminating after exceeding the maximum number of iterations. Abundance images were normalised for visualisation purposes. Endmember signatures displayed on the right correspond to the analyses performed with a random seed set to 42. Scale bar: 100 μm (bottom right, marked in orange).

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