Peptidergic dense-core vesicles are involved in packaging and releasing neuropeptides and peptide hormones—critical processes underlying brain, endocrine and exocrine function. Yet, the heterogeneity within these organelles, even for morphologically defined vesicle types, is not well characterized because of their small volumes. We present image-guided, high-throughput mass spectrometry-based protocols to chemically profile large populations of both dense-core vesicles and lucent vesicles for their lipid and peptide contents, allowing observation of the chemical heterogeneity within and between these two vesicle populations. The proteolytic processing products of four prohormones are observed within the dense-core vesicles, and the mass spectral features corresponding to the specific peptide products suggest three distinct dense-core vesicle populations. Notable differences in the lipid mass range are observed between the dense-core and lucent vesicles. These single-organelle mass spectrometry approaches are adaptable to characterize a range of subcellular structures.

Organelles are one of the smallest structural units that influence the functional, morphological and biochemical characteristics of different cell types. Chemical analysis of individual organelles is challenging due to their attoliter volumes, the wide dynamic range of analyte concentrations and the need for sophisticated isolation procedures, thereby limiting our understanding of their chemical heterogeneity. Mass spectrometry (MS) imaging has begun being used for both cellular and some subcellular analyses in discovery-based studies but is limited in throughput and spatial resolution for organelle measurements. Alternatively, both cells and organelles of interest can be isolated and placed on a glass slide for subsequent MS measurement. Though we used this approach to assay the peptides within individual organelles, it was limited in throughput and difficult to automate due to the manual positioning of each individual organelle before measurement.

A recent enhancement to single-cell measurement involves scattering the cells of interest onto a microscope slide, determining their locations via fluorescence microscopy and then targeting selected locations with matrix-assisted laser desorption/ionization (MALDI) MS, allowing tens of thousands of cells to be assayed. Here we adapted the single-cell approach to single organelles, which required three enhancements: (1) improved object targeting approaches, (2) optimized analyte detection using high-resolution MS and (3) unsupervised data analysis workflows to characterize organelle heterogeneity. These advances allowed us to use MALDI Fourier-transform ion cyclotron resonance MS for the high-throughput simultaneous detection of both peptide and lipid species in 0.5- to 2-μm-diameter dense-core vesicles (DCVs) and electron lucent vesicles (LVs) isolated from the exocrine atrial gland (AG) (Fig. 1a–f) and red hemiduct of Aplysia californica, respectively. By developing single-organelle sampling techniques that can be conducted in a high-throughput manner, we observed subtypes of DCVs defined by their overlapping but distinct peptide content. We also identified a peptide prohormone not previously known to localize within DCVs, as well as large differences in the contents between the DCVs and LVs. Our method revealed post-translational proteolytic processing of AGPB1 (XP_012945142.1), AGPA1 (XP_012945143.1) and AGPA2 (XP_012945134.1) prohormones, with peptide structures validated by liquid chromatography–tandem mass spectrometry (LC–MS/MS). Additionally, a novel prohormone, AG Peptide D, was characterized (Supplementary Figs. 1 and 2 and Supplementary Table 1). Importantly, our workflow is extendable to multiple imaging modalities such as scanning electron microscopy (Supplementary Fig. 3) and instrumentation that uses a piezo linear stage and camera, providing an avenue for the analysis of targets that are smaller than the wavelength of light.

Results

Image-guided MALDI MS approach. To adapt single-cell approaches to objects that are tenfold smaller in diameter than cells measured previously, we created a method to enhance identification of micrometer-sized object locations while improving analyte detectability by minimizing the number of interfering chemicals that could reduce the ionization efficiency of our analytes of interest. Traditionally, single-cell targeting requires chemical labeling to mark objects of interest, but introduction of exogenous chemicals during single-organelle sample preparation is problematic due to the 1,000-fold less material present. Paraformaldehyde fixation is a hallmark step in most staining protocols, but paraformaldehyde fixation crosslinks DNA and peptides/proteins, rendering them poorly ionizable and undetectable by MS. By measuring the pixel properties produced by DCVs under brightfield conditions, we compiled sequences of image algorithms (Supplementary Fig. 4) that selectively identify the pixels produced by DCVs versus the background (Fig. 1b–d), allowing identification of the spatial locations of individual DCVs without the need for chemical labeling. The axial resolution of light microscopy limits this approach to the analysis of vesicles that are approximately 500 nm in diameter or...
larger. However, nonoptical approaches such as scanning electron microscopy can be applied, providing a resolution down to 10 nm, allowing targeting of nanometer-sized objects (Supplementary Fig. 3). A three-step approach was developed for isolation of a representative number of vesicles without loss of chemical detail, which is critical for the high-throughput preparation of organelle...
**Fig. 2** | Chemical heterogeneity of DCV populations: unsupervised data analysis, simultaneous acquisition of analyte classes and detection of novel Peptide D.  

**a**, Deterministic CX was used to select 200 mass spectral features to improve data interpretation. CX decomposition was applied for feature selection where the best low-rank approximation was provided through singular value decomposition with a rank parameter $k = 150$, which was determined by the reconstruction errors with different rank parameters. Rank $k = 150$ has a reconstruction error less than 25% of the original dataset.  

**b**, ‘Statistical leverage scores’ for all spectral features were computed and plotted against the $m/z$ axis. Multiple detected AG peptides can be seen with high leverage scores and are annotated with blue triangles. The ‘Statistical leverage score’ for each feature indicates its influence on the best low-rank fit of the data matrix. Selecting the top 200 features ensured that CX decomposition closely approximated the best low-rank fit of the original data matrix while removing uninformative information.  

**c**, Results of $k$-means clustering of the dataset containing 200 selected features with the highest statistical leverage score.  

**d**, Violin plots visualizing a subset of the selected features and their relative distribution in each cluster, with known AG peptides marked.  

**e**, Simultaneous detection of PC(18:1/16:0), PC(18:1e/16:0) and PC(18:1e/18:1) with their corresponding sodiated adducts annotated.  

**f**, The novel prohormone, Peptide D, was identified by MALDI MS mass-match assignment of Peptide D [63–75] and Peptide D [132–162]. SVD, singular value decomposition.
samples. In this approach, a volatile, isosmotic-ammonium acetate buffer was deposited onto an indium tin oxide (ITO)-coated glass slide, into which a small aliquot of artificial sea water (ASW) solution containing the vesicles was deposited. The vesicles were allowed to sediment and adhere to the glass surface before the solution was aspirated, leaving a large number of visually intact vesicles distributed across the slide. After identifying the vesicles on the slide using the described pipeline, an image mask marks the vesicles distributed across the slide. After identifying the vesicles marked by clusters of maximal pixel intensity values, zero-pixel intensity, creating a binary image output (Fig. 1d). With the vesicles marked by clusters of maximal pixel intensity values, the binary image created from the masked image output can be used for identification of the spatial locations of vesicles on the slide for subsequent analysis via MALDI MS. A total of 123 LVs were measured from the same three biological replicates used in the DCV isolation. A total of 598 DCVs were analyzed and, after cross-refencing the single-DCV spectra with the LC–MS/MS data, over 50 mature full-length peptides were assigned by peptide mass fingerprinting to eight known prohormones and one novel prohormone (Supplementary Table 2). As a result, the peptides AGPA2 [36–69], AGPB1 [85–118], AGPA1 [117–152], AGPA1 [156–173] (or identical peptide AGPB1 [205–222]), N-terminal peptide AGPA1 [22–34] and the truncated C-terminal peptide AGPB1 [71–81] lacking the C-terminal Thr-Asn⁺ were detected, resulting in 39% and 28% coverage of AGPA1 and AGPB1 prohormones (Fig. 1g,h), respectively. Lastly, by using an m/z range of 150–4,500, we were able to perform simultaneous detection of both peptide and lipid species in individual DCVs. Three phosphatidylcholine (PC) lipid species detected by mass-match assignment include: PC(18:1/16:0), PC(18:1e/16:0) and PC(18:1e/18:1), with respective assignment mass errors of ±0.21 ppm and ±0.21 ppm (Fig. 2e). The three detected lipids were previously identified in A. californica neurons using MALDI MS.}

**Fig. 3 | Comparison between two distinct vesicle types, LVs and DCVs.** a. Top, digitally inverted brightfield image showing LVs distributed across the glass slide. a Bottom, modification of the image-processing pipeline allows multiple morphologically distinct objects to be identified and added or removed for subsequent analysis. Green areas represent objects accepted for analysis. Magenta outline represents an object (here a spermatozoon) removed from analysis. A small number of spermatozoa (annotated) typically collected during red hemiduct LV sample preparation were used to demonstrate the effectiveness of our object filtering approach. A total of 123 LVs were measured from the same three biological replicates used in the DCV isolation. b. t-SNE was performed on the initial dataset for visualization of LVs and DCVs using all mass spectral features. c. Confusion matrix of the prediction on the test data using a threefold validation. d. Representative LV mass spectrum (red) overlaid on a representative DCV mass spectrum (gray). Representative spectra were not preprocessed. A subset of mass spectral features determined important via SHAP are annotated in the LV mass spectrum. e. Mass spectral feature contribution plot showing the importance of features to the vesicle classification task across m/z range 500–1,100 on the x axis. The y axis shows the mean absolute SHAP value for the corresponding m/z feature (normalized between 0 and 1). Gray dots represent features with elevated mean signal intensities in DCVs and pink dots represent features with elevated mean signal intensities in LVs. The different sized dots represent the respective P values (two-sided Wilcoxon rank-sum test) for each corresponding feature.
Differential packaging of peptides from the same prohormone into different vesicles was shown in the bag cell neurons of *A. californica* using immunogold staining approaches, although this has not been demonstrated in other *A. californica* cell types. Here, we considered whether the different peptide complements may be from differential proteolytic processing of AGPB1, AGPA1 and AGPA2 prohormones. Therefore, CX decomposition, an unsupervised statistical approach, was applied to select statistically important features while removing redundant and uninformative ones for improved downstream data interpretation (Supplementary Figs. 6 and 7). CX decomposition based on statistical leverage scores was used to select the top 200 features (Fig. 2a,b and Supplementary Fig. 8). Next, using the selected features, the DCVs were partitioned into three groups using the k-means clustering algorithm, for which the number of groups was determined by calculating the Within Cluster Sum of Squares (Fig. 2c). Significantly different features were obtained by performing a Wilcoxon rank-sum test for each individual cluster against the others (Supplementary Fig. 9). The violin plots (Fig. 2d) highlight selected mass spectral features and their distribution in the different DCV cluster types where known AG peptides are labeled. The N-terminal peptides, AGPB1 [71–81] and AGPA1 [22–34], and the novel Peptide D [63–75] (Fig. 2f), are shown to be defining features in Cluster 1 when compared with Cluster 2, whereas Cluster 3 demonstrates localization of multiple expected AG peptides from the prohormone C termini, including AGPA1 [156–173], Peptide D [63–75], AGPB1 [85–118] and a peptide cluster belonging to Peptide D [132–162] (Supplementary Fig. 10). The data indicate that the N- and C-terminal peptides from these two prohormones are differentially packaged into a subset of DCVs, whereas the bioactive peptides that initiate the egg-laying process, AGPA2 [36–69] and AGPB1 [85–118], are predominantly packaged into a separate subset of vesicles. An early pioneering study that quantified the amounts of AGPA1 [22–34], AGPA2 [36–69] and AGPB1 [85–118] in isolated intact granules from AG homogenates also suggested the possibility of packaging of AGPA1 [22–34] into a distinct class of secretory vesicle. Perhaps this differential packaging of N-terminal products in the AG would be akin to what is also observed in the bag cell neurons where N-terminal products are differentially packaged into a subset of vesicles. An early study showed that the AG contains a double membrane structure, with the inner membrane forming cristae-like structures that are analogous to those found within the mitochondrion. In contrast, DCVs from the AG contain a single membrane structure, with the inner membrane forming cristae-like structures that are analogous to those found within the mitochondrion. Due to the nonpeptidergic content of the LVs and the clear differences in ultrastructure between the LVs and DCVs, a machine learning model was trained for the differentiation between LVs and DCVs, focusing on the m/z range of 500–1,100. t-Distributed stochastic neighborhood embedding (t-SNE) was initially applied using the mass spectral features to visualize the data in a low-dimensional space (Fig. 3b). Next, gradient boosting trees were trained with a threefold validation, obtaining an accuracy of 98.6 ± 0.78% for the classification between LVs and DCVs (Fig. 3c). The most important features for the classification between LVs and DCVs (Fig. 3d) were selected via Shapley additive explanations (SHAP) through a previously described method, where a total of 97 features with nonzero mean SHAP values from the model output were selected, with 36 features putatively identified by mass-match assignment (Fig. 3e). The model provided an improved classification performance through retraining on the selected features (Supplementary Fig. 11), displaying the differential lipid profiles between the two vesicle classes (Supplementary Figs. 12, 13 and 14). The data show the strong presence of multiple sterol lipid species to be of importance to their classification. The absence of sterol lipid species correlates with uniformly curved membranes. Whereas when present in sufficient concentration, sterol lipid species will distribute asymmetrically in the lipid bilayer causing large, flat membrane regions that are separated by sharp curves in the membrane, creating negative membrane curvature. Interestingly, negative membrane curvature would promote the formation of the cristae-like invaginations that can be observed in the LVs.

**Discussion**

As the spatial resolution of MS advances from the single-cell level to include the single-organelle domain, new analytical approaches are being developed to unravel the complex microenvironments housed within single cells. Capillary micro-sampling techniques coupled to electrospray ionization (ESI)–MS have been developed for the identification of peptides in individual cells and in some subcellular compartments of cells. Though capillary micro-sampling ESI–MS techniques can identify peptides in subcellular compartments, they require fine-tuning of the capillary position and controlling the volume extracted. These requirements are critical to single-organelle extraction and pose challenges to repeated manual and throughput due to potential contamination from the cellular membrane, cytoplasm and other subcellular organelles. Alternatively, our image-guided MALDI MS approach provides direct single-organelle targeting using point-based image registration.

In conventional MS imaging, as spatial resolution increases, so does acquisition time. Therefore, to maintain the throughput required for characterizing large populations of individual organelles, we are not imaging slides containing the vesicles. Instead, the organelles themselves are isolated and selectively targeted using the MALDI laser. Our high-throughput, label-free, single-organelle approach enables simultaneous characterization of both the peptide and lipid contents of hundreds of intact 0.5–2-µm-diameter DCVs and LVs. Because our approach uses a combination of free ware and instrumentation that is commonly available to researchers, it is adaptable to many experimental designs, including other organelle types such as microvesicles, which can range from 100 nm up to 1 µm in diameter.

**Online content**

Any methods, additional references, Nature Research reporting summaries, source data, extended data, supplementary information, acknowledgements, peer review information; details of author contributions and competing interests; and statements of data and code availability are available at https://doi.org/10.1038/s41592-021-01277-2.

Received: 1 April 2021; Accepted: 18 August 2021; Published online: 30 September 2021

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**Methods**

General. All chemicals were obtained from Sigma-Aldrich unless specified otherwise.

**Vesicle isolation.** *A. californica* (150–200 g) were obtained from the National Resource for Aplysia and kept in a 14°C aquarium filled with Instant Ocean (Aquarium Systems). Animal euthanasia was performed in accordance with the AVMA Guidelines for the Euthanasia of Animals: 2020 Edition (Section S6.3.1.1 Noninhaled agents for immersion). Animals were anesthetized using a 333 mM MgCl₂ solution injected into the body cavity (50% volume/body).

For DCV isolation, the AG was isolated by manual dissection, and placed into a microcentrifuge tube containing 1 ml of ASW due to the holocrine release with a polypropylene Pasteur pipette, releasing intact DCVs into the ASW of 150 µl at temperature ranging from 100–150 °C for 8 min. Matrix deposition was followed by recrystallization using 5% particle size, Thermo Fisher Scientific) using solvent A for off-line desalting. Next, 30% tile overlap. The resulting tiles were stitched before exporting to TIFF-file format (ZEN 2.1 software). Scanning electron microscopy images were acquired using an FEI Quanta FEG 450 environmental scanning electron microscope (FEI). Images were acquired using an accelerating voltage of 10 kV, dwell time of 10 µs and working distance of 6.6 mm.

**Image-processing for vesicle recognition in microMS.** We utilized biological image analysis software, including ImageJ⁶ and CellProfiler⁷, for vesicle pixel recognition on the brightfield microscopy image. Importantly, other image-processing software can be used for this process as well. CellProfiler pipeline modules were created to selectively mask the DCVs and LVs on the brightfield microscopy image to remove the background and retain just information on the vesicles of interest as the foreground. Masking the identified vesicles creates a binary image marking the pixel locations of vesicles on the glass slide. The microMS software was used to translate the pixel locations on microscopy images to physical coordinates of the mass spectrometer’s stage. A 200-µm-distance filter was then applied in microMS (removing vesicles located closer than 200 µm to each other from the target list). Alternatively, ImageJ was used as a simple thresholding strategy by measuring the difference between the vesicle (foreground) and background pixel intensity, which allows a threshold to be set, leaving only objects of interest for recognition in microMS.

**Matrix application.** Matrix deposition for MALDI MS analysis was performed using a glass sublimation apparatus (Wilmad-LabGlass) filled with 500 mM ammonium acetate solution was prespotted onto a glass substrate. Surface Resistivity 70–100Ω (Data Technologies). Then, 50 µl of vesicle-containing ASW solution was pipetted into the 100 µl of ammonium acetate solution on the ITO-glass slide for a total volume of 150 µl. The resulting vesicle solution was then rinsed with 500 µl ammonium acetate with simultaneous aspiration of the solution, leaving DCVs seeded across the slide. All steps were visually monitored using an inverted microscope. Each slide held DCVs from three animals (biological replicates) and a total of three slides (technical replicates) were prepared. For LV isolation, the procedure used for the DCVs was repeated using red hemiduct tissue, with the other steps being the same as for the DCV isolation; 123 LVs were analyzed from the same three biological replicates used for the DCV isolation.

**Imaging.** Brightfield images were acquired on an Axio Imager M2 (Zeiss) equipped with an AxioCam ICc 5, a ×0.63 camera adapter and a transmitted light visible light–light emitting diode lamp. Images were acquired in mosaic mode using a ×10 objective with 30% tile overlap. The resulting tiles were stitched before exporting to TIFF-file format (ZEN 2.1 software).

Scanning electron microscopy images were acquired using an FEI Quanta FEG 450 environmental scanning electron microscope (FEI). Images were acquired using an accelerating voltage of 10 kV, dwell time of 10 µs and working distance of 6.6 mm.

**Data preprocessing.** Data preprocessing was performed using Compass Data Analysis 4.4.2 (Bruker) and MALDI 2018b (MathWorks). Internal calibration using the exact mass of AG peptides was performed using AGFBI [71–81] (m/z 1.221 6878), AGPA1 [22–34] (m/z 1.396 7225), AGPA1 [156–173] (m/z 1.908 8761), AGPA2 [33–69] (m/z 3.922 9478) and AGPA1 [85–118] (m/z 4.031 0053). Peak picking and peak export for statistical analysis in MALAT 2018b were set to a signal-to-noise ratio of 5 with a relative intensity threshold of 0.01%. A nonuniform bin width was used for mass spectral alignment. For DCV data analysis, mass features were truncated at m/z 1,100 for downstream data analysis of only AG peptides. For LV and DCV vesicle classification, mass features in m/z 500–1,100 were selected for downstream data analysis. Internal calibration using AG peptides was not performed for the LV and DCV classification tasks. Using the target list provided by microMS, the pixel coordinates were used to find and crop the locations of individual vesicles across the original microscopy image. The corresponding mass spectra were matched with the appropriate vesicle for visual evaluation of corresponding single vesicles.

**CX decomposition and statistical analysis.** Unsupervised approaches such as principal component analysis (PCA) and, more generally, such as feature extraction or factorization, are valuable data analysis tools to enable field-specific interpretation of high-dimensional datasets. However, the interpretation is limited due to the complicated eigenspace obtained from PCA, detracting our further understanding of the feature space of the data matrix. CX matrix decomposition is designed to obtain low-rank decomposition of the data in terms of actual chemical compounds. Given an m × n data matrix A, the algorithm decomposes it into an m × c matrix X and a c × n matrix W, where c is expressed by c number of column vectors of the original data. The statistical leverage scores are used to rank and select the columns of C from A, which can be obtained by

$$ l_i = \sum_{v=1}^{k} v_i^2 $$

where $l_i$ is the leverage score for the jth column/feature, $v_i$ is the right singular vectors obtained by the singular value decomposition and $k$ is the rank to be selected. $X$ is then determined by minimizing the error:

$$ \min ||A - CX|| $$

The reconstruction error evaluation and the rank $k$ selection are provided as Supplementary Fig. 8. Based on the evaluation, the top 200 features are selected to form the columns of the matrix C. k-means clustering, and the Wilcoxon rank-sum test, were performed using the Python-based open source package SCANPY⁸. The stacked violin plots, shown in Fig. 2d, of the normalized signal intensities represent the identified peak features across the three clusters, with the vesicle type (or cluster) assignments obtained by k-means clustering. The y axis is the root-mean-squared-normalized peak intensity and each violin contains the distribution of the normalized intensities of the corresponding features in the x axis.

**Peptide sequencing by LC–MS/MS.** Peptide extracts (n = 3) were obtained by manually grinding entire AG tissue in 500 µl of acidified methanol followed by evaporation and reconstitution of each extract in 0.1% formic acid. A nanoElute (Bruker) ultra-high-pressure nano-flow chromatography system was coupled to a trapped ion mobility–quadrupole time-of-flight mass spectrometer (timsTOF Pro, Bruker) with a CaptiveSpray nano-electrospray ion source (Bruker) equipped with an external column oven. Mobile phases A and B were water with 0.1% formic acid (v/v) and acetonitrile with 0.1% formic acid (v/v), respectively. Samples were loaded onto a precolumn peptide trap (Acclaim PepMap 100 C18, 1 × 5 mm, 5-µm particle size, Thermo Fisher Scientific) using solvent A for off-line desalting. Next, the trap was placed in-line with the analytical column and peptide separation was performed at 40 °C with a uniform flow of 300 nl/min⁻¹ on a C18 ReproSil AQ column (Bruker FIITEN, P/N no 1842621: 150 mm x 75 mm, 1.9-µm particle size, pore size 120 Å) equilibrated at 2% B. A linear gradient of solvent B was applied as follows: 2–10% within 5 min, 10–50% in the next 115 min, followed by a washing step at 95% B and re-equilibration, during which data collection was not performed. The mass spectrometer was operated in parallel accumulation–serial fragmentation (PASEF) mode for peptide sequencing. The mass range for the precursor ion was set to m/z 100–1,700, ion mobility 1/Kₐ range 0.6–1.6 V cm⁻² s⁻¹. Fragmentation was performed with 10 PASEF scans, cycle time of 1.1 s, during which collision energy varied linearly between 20 and 59 eV depending on precursor 1/Kₐ value within the set range. Active dynamic exclusion of precursor ions was set to 0.4 min.

**Peptide identification by bioinformatics.** MS raw files were processed with PEAKS Online⁹ (Bioinformatics Solution) using the DeNovo database (DB) and post-translational modification (PTM) protocols, sequentially. Peptide sequence tags obtained by the DeNovo process (80% average local confidence score cut-off) were searched against the Aplysia ReSeq database available from the NCBI (GCF_000002075.1). The protein database was filtered to include proteins up to
1,000 amino acids long. Search parameters for DB included: parent mass error tolerance 20.0 ppm, fragment mass error tolerance 0.03 Da, no enzyme, digest mode unspecific. Next, PTMs were identified by searching the data for amidation, acetylation (K) and acetylation N terminus, pyro-glutamylation from (Q) and (E), phosphorylation (STY) and half-disulfide bridge. The false discovery rate was determined by decoy fusion method and the threshold set to 1% for peptides. Proteins with $-\log_{10} P > 20$ and at least one unique peptide are reported.

**Vesicle classification through machine learning.** We adapted a previously described machine learning strategy6 for vesicle type classification to differentiate between DCVs and LVs based on their lipid contents. Features in the m/z range of 500–1,100 were used for the classification task. Gradient boosting trees were trained with a threefold validation; in each fold of the model performance, the metrics were computed to obtain classification accuracies, confusion matrices and receiver operating characteristic curves (Supplementary Fig. 11). The most contributing features to the classification task were selected via SHAP, a game theory approach for model explanations, and were obtained through a Python implementation of SHAP. A total of 97 features with nonzero mean SHAP values from the output of trained models were selected and 36 were putatively annotated by searching against the LIPID MAPS30 database with a 7-ppm tolerance. Models were then retrained with the SHAP-selected features as well as the annotated features to verify the discriminative ability of those features. t-SNE using the cosine distance was used to visualize the lipid differences between DCVs and LVs in a low-dimensional space.

**Reporting Summary.** Further information on research design is available in the Nature Research Reporting Summary linked to this article.

**Data availability**
The data that support the findings of this study are publicly available via the Illinois Data Bank (https://doi.org/10.13012/B2IDB-5949772_V1).

**Code availability**
The code used in this study is available on GitHub (https://github.com/richardxie1119/DCV_DA).

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**Acknowledgements**
This work was supported by funding from the National Institute on Drug Abuse under award No. P30DA018310 (to J.V.S.) and the National Human Genome Research Institute under award No. R11HG010023 (to J.V.S.). The content is solely the responsibility of the authors and does not necessarily represent the official views of the awarding agencies.

**Author contributions**
J.V.S. conceived the project idea. S.S.R. and D.C.C. designed and performed the sample preparation approach. D.C.C. acquired microscopy images, developed the image-processing approaches and acquired the high-throughput MALDI MS data. Y.R.X. performed data analysis and contributed to the figures. E.V.R. performed LC–MS/MS peptide sequencing and bioinformatics. All authors wrote and approved the final manuscript. J.V.S. acquired funding and provided direction throughout the project.

**Competing interests**
The authors declare no competing interests.

**Additional information**
**Supplementary information** The online version contains supplementary material available at https://doi.org/10.1038/s41592-021-01277-2.

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**Peer review information** *Nature Methods* thanks Zhibo Yang and the other, anonymous, reviewer(s) for their contribution to the peer review of this work. Arunima Singh was the primary editor on this article and managed its editorial process and peer review in collaboration with the rest of the editorial team.

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Reporting Summary

Nature Research wishes to improve the reproducibility of the work that we publish. This form provides structure for consistency and transparency in reporting. For further information on Nature Research policies, see our Editorial Policies and the Editorial Policy Checklist.

Statistics

For all statistical analyses, confirm that the following items are present in the figure legend, table legend, main text, or Methods section.

- The exact sample size (n) for each experimental group/condition, given as a discrete number and unit of measurement
- A statement on whether measurements were taken from distinct samples or whether the same sample was measured repeatedly
- The statistical test(s) used AND whether they are one- or two-sided
  
  Only common tests should be described solely by name; describe more complex techniques in the Methods section.
- A description of all covariates tested
- A description of any assumptions or corrections, such as tests of normality and adjustments for multiple comparisons
- A full description of the statistical parameters including central tendency (e.g. means) or other basic estimates (e.g. regression coefficient) AND variation (e.g. standard deviation) or associated estimates of uncertainty (e.g. confidence intervals)
- For null hypothesis testing, the test statistic (e.g. F, t, r) with confidence intervals, effect sizes, degrees of freedom and P value noted
  
  Give P values as exact values whenever suitable.
- For Bayesian analysis, information on the choice of priors and Markov chain Monte Carlo settings
- For hierarchical and complex designs, identification of the appropriate level for tests and full reporting of outcomes
- Estimates of effect sizes (e.g. Cohen’s d, Pearson’s r), indicating how they were calculated

Our web collection on statistics for biologists contains articles on many of the points above.

Software and code

Policy information about availability of computer code

Data collection  Optical microscopy images were acquired using ZEN 2.0 Pro edition (Zeiss, Jena, Germany) software. Scanning electron microscopy images were acquired using FEI Quanta 400 (FEI Company, Hillsboro, Oregon). The microscopy images were processed using CellProfiler 3.1.8 (Broad Institute, Cambridge, Massachusetts), ImageJ 1.53c [NIH & Laboratory for Optical and Computational Instrumentation, University of Wisconsin] and MicroCNS (University of Illinois at Urbana-Champaign, Urbana, Illinois). The MALDI mass spectrometry data was acquired using GeMS version 2.7.2 (Bruker Corporation, Billerica, Massachusetts). The LC-MS/MS data was acquired using oTOFControl 6.2 (Bruker Corporation, Billerica, Massachusetts).

Data analysis  Raw MALDI MS data were preprocessed using DataAnalysis 4.4.2 (Bruker Corporation, Billerica, Massachusetts) and in-house developed MATLAB scripts r2018b (MathWorks, Inc., Natick, Massachusetts). The data analysis pipeline was implemented in Python 3.7.1 (Python Software Foundation, Beaverton, Oregon) with open-source packages NumPy 1.15.2 (NumFOCUS, Inc., Austin, Texas), SciPy 1.5.3 (SciPy developers), scikit-learn 0.23.1 (scikit-learn developers), and single cell analysis package Scanpy 1.5.1. Plotting packages include matplotlib 3.3.2 (NumFOCUS, Inc., Austin, Texas) and seaborn 0.11.0. LC-MS/MS raw files were preprocessed using DataAnalysis 5.3 (Bruker Corporation, Billerica, Massachusetts) and PEAKS Online v1.3 (Bioinformatics Solutions Inc., Waterloo, Ontario). SignalP 5.0 (Bioinformatic Unit, Technical University of Denmark, Denmark) was used to predict sequence cleavages. Clustal Omega 1.2.4 (University College Dublin, Dublin, Ireland) was used for sequence alignment.

For manuscripts utilizing custom algorithms or software that are central to the research but not yet described in published literature, software must be made available to editors and reviewers. We strongly encourage code deposition in a community repository (e.g. GitHub). See the Nature Research guidelines for submitting code & software for further information.
Data

Policy information about availability of data
All manuscripts must include a data availability statement. This statement should provide the following information, where applicable:
- Accession codes, unique identifiers, or web links for publicly available datasets
- A list of figures that have associated raw data
- A description of any restrictions on data availability

The data that supports the findings of this study are publicly available via the Illinois Data Bank: https://doi.org/10.13012/B2IDB-5949772_V1

Field-specific reporting

Please select the one below that is the best fit for your research. If you are not sure, read the appropriate sections before making your selection.

☒ Life sciences
☐ Behavioural & social sciences
☐ Ecological, evolutionary & environmental sciences

For a reference copy of the document with all sections, see nature.com/documents/mr-reporting-summary-list.pdf

Life sciences study design

All studies must disclose on these points even when the disclosure is negative.

Sample size

To account for biological variability between animals we chose to use three biological replicates in our study. To minimize MALDI MS shot-to-shot variability and changes in instrumental conditions between analysis days, we placed dense-core vesicles from each of the three biological replicates on the same microscopy slide. A total of three microscopy slides were prepared using the three biological replicates and analyzed on separate days. By measuring 598 dense-core vesicles across the three microscopy slides, we believe we have mitigated any unwanted variation that may be attributed to biological variability between animals or variation attributed to changes in MALDI MS instrumentation conditions.

Data exclusions

No data was excluded from the study.

Replication

We performed the dense-core vesicle measurements using two types of mass analyzers (Fourier-transform ion cyclotron resonance mass spectrometer and time-of-flight mass spectrometer) where expected atrial gland peptides were detected using both types of mass analyzers. Every detected peptide by MALDI MS was structurally characterized using LC-MS/MS. Each experimental setup contained at least three biological replicates.

Randomization

Animals/Dense-core vesicles were not subjected to different treatments or interventions that would create the possibility of selection bias. Therefore, we believe randomization does not apply to this study.

Blinding

Group allocation due to different treatments or interventions did not occur during this study, which is why we believe blinding is not applicable to this study.

Reporting for specific materials, systems and methods

We require information from authors about some types of materials, experimental systems and methods used in many studies. Here, indicate whether each material, system or method listed is relevant to your study. If you are not sure if a list item applies to your research, read the appropriate section before selecting a response.

Materials & experimental systems

| n/a | Involved in the study |
|-----|-----------------------|
| ☒   | Antibodies           |
| ☒   | Eukaryotic cell lines |
| ☒   | Palaeontology and archaeology |
| ☒   | Animals and other organisms |
| ☒   | Human research participants |
| ☒   | Clinical data |
| ☒   | Dual use research of concern |

Methods

| n/a | Involved in the study |
|-----|-----------------------|
| ☒   | ChiP-seq |
|     | Flow cytometry |
|     | MRI-based neuroimaging |

Animals and other organisms

Policy information about studies involving animals. ARRIVE guidelines recommended for reporting animal research

Laboratory animals

Aplysia californica
| **Wild animals** | This study did not involve wild animals. |
|------------------|----------------------------------------|
| **Field-collected samples** | This study did not involve field-collected samples (the animals were from the Aplysia Research Facility, University of Miami). |
| **Ethics oversight** | Exempt from oversight because the animals used are mollusks / invertebrates. However, we followed published recommendations for the humane treatment of gastropod mollusks in research, including use of anesthesia. |

Note that full information on the approval of the study protocol must also be provided in the manuscript.